BMJ Open

Different doses of prucalopride in treating chronic idiopathic constipation: a meta-analysis and Bayesian analysis

Tao Yang,1 Kaili Wang,2 Yibo Cao,3 Jianxia Wen,4 Shizhang Wei,5 Haotian Li,5 Xiangdong Yang,5 Tianbao Xiao7

ABSTRACT

Objective This study aims to explore the incremental benefit of different doses of prucalopride in treating chronic idiopathic constipation (CIC).

Methods PubMed, EMBASE, MEDLINE, Cochrane Library, Chinese Biomedical Database, China National Knowledge Infrastructure, VIP medicine information and Wanfang databases were comprehensively searched up to March 2020. Prospective trials with different doses of prucalopride versus placebo were selected. The frequency of spontaneous bowel movements (SBMs) per week and the treatment-emergent adverse events (TEAEs), such as headache, arrhythmia, diarrhea, dizziness, nausea and vomiting, were first synthesised in a meta-analysis. The probability of optimal dose of prucalopride was then ranked by random-effects within Bayesian analysis.

Results 14 high-quality randomised controlled trials with 4328 patients were ultimately included. SBMs per week increased significantly after using 1 mg (OR: 2.40, 95% CI 1.32 to 4.37), 2 mg (OR: 2.55, 95% CI 1.93 to 3.36) and 4 mg (OR: 2.51, 95% CI 1.92 to 3.28) prucalopride. Bayesian analysis demonstrated 1 mg dose obtained the maximum SBMs per week (OR: 3.31, 95% credible interval 1.72 to 6.16, probability rank=0.70) indirectly compared with 2 mg and 4 mg doses. TEAEs were higher significantly in 2 mg (risk ratio (RR): 1.20, 95% CI 1.09 to 1.33) and 4 mg (RR: 1.14, 95% CI 1.07 to 1.22) prucalopride. The 1 mg dose did not reach statistical significance (RR: 1.17, 95% CI 0.94 to 1.44).

Conclusions The study concludes that 1 mg dose at commencement could be safer in treating CIC and that 2 mg prucalopride could be more efficacious in terms of SBMs per week outcome receiving.

PROSPERO registration number CRD42019136679.

INTRODUCTION

One of the most complicated disorders of the digestive system is chronic idiopathic constipation (CIC). The global prevalence of CIC in adults was estimated at 14% (95% CI 12% to 17%).1 Age, particularly over 65 years, and female gender could be potential risk factors for increased prevalence.2 CIC has the ability to curtail quality of life and increase psychological distress significantly.3 Furthermore, these harmful effects are inclined to be more widespread in patients with inability to coordinate bowel movements than in patients with constipation with slow transition.4 A community survey about spending in CIC management demonstrates this annoying disease enormously increases social and economic burden.5 6 Management methods are also hard and arduous strikingly for both gastrointestinal doctors and surgeons worldwide. Although laxatives (both stimulant and osmotic) have been widely employed in clinical practice, evidence on long-term efficacy in clinical setting remains inadequate.7 Furthermore, the weakened gastrointestinal motility and impairment of the intestinal nervous system (INS) precipitate the occurrence of CIC evidently.8 Therefore plausible treatments for CIC are suggested to focus on the neurons in the INS and improve their physiological excitability.9 Although many novel pharmacological attempts have been made, such as cisapride, tegaserod and lubiprostone, these attempts failed to earn popularity among clinical physicians and surgeons and failed to be included as first-line treatments, possibly due to remarkable side effects, possibly due to remarkable side effects (eg, cardiovascular...
problems). Prucalopride was then investigated in clinical trials.

Prucalopride, a highly selective serotonin 5-hydroxytryptamine 4 (5-HT4) agonist, was introduced to the market in 2009. The pharmacological mechanism of prucalopride is to stimulate the receptors of 5-HT4, which are diffusely concentrated on the enteric neurons. The excitatory neurotransmitter acetylcholine is then released by the 5-HT4 agonists and promotes mucosal secretion. Prucalopride has been advocated by the US Food and Drug Administration and the European Union for symptomatic treatment of CIC, especially in patients who have failed to adequately achieve symptom relief following stimulant or osmotic laxatives. Previous randomised controlled trials (RCTs) have reported prucalopride doses of 0.5 mg, 1 mg, 2 mg and 4 mg compared with placebo accelerated colonic transit and spontaneous bowel movements (SBMs) per week prominently. However, there are rare guidelines or expert opinions to explore the incremental benefit of prucalopride. The administration of doses in clinical practice is empirical in most cases. It remains a conundrum whether dose-dependent relationship exists in using prucalopride.

This article attempts to define the clinical usefulness and safety of different doses of prucalopride in treating patients with CIC. We intend to provide the optimal dose of prucalopride for gastrointestinal physicians in clinical work and employed both pairwise meta-analysis and Bayesian analysis to address these problems as much as possible.

MATERIALS AND METHODS

Patient and public involvement

No patients were involved.

Protocol registration

We have registered this protocol in PROSPERO (International Prospective Register of Systematic Reviews) in May 2019 (https://www.crd.york.ac.uk/PROSPERO).

Eligibility criteria

The Cochrane Handbook for Systematic Reviews of Interventions and the Preferred Reporting Items for Systematic Reviews and Meta-Analyses statement were used in this study. Studies that meet the following inclusion criteria were included: (1) the design of the study was a prospectively randomised controlled trial; (2) CIC was histologically confirmed; and (3) at least one of the investigated outcomes were reported in the original research. Exclusion criteria were as follows: (1) irrelevant studies and duplicate literature; (2) unavailable data in the literature; and (3) letters, reviews, comments, case reports and meta-analyses.

Search methodology

PubMed, EMBASE, MEDLINE, Cochrane Library, Chinese Biomedical Database, China National Knowledge Infrastructure, VIP medicine information system and Wanfang electronic databases were comprehensively searched up to March 2020. The search terms were a combination of medical subject headings (MeSH) terms and the following free words: (constipation/chronic constipation/functional constipation) and (prucalopride/Resolor) and (random/randomized/RCTs/clinical trial). In addition, we manually searched for other potential and relevant references. There were no limitations in the language of all publications. Taking the PubMed database as an example, the search strategies are as follows:

- #1 constipation (MeSH).
- #2 constipation* af.
- #3 chronic constipation af.
- #4 functional constipation af.
- #5 #1 OR #2 OR #3 OR #4.
- #6 prucalopride (MeSH).
- #7 prucalopride af.
- #8 Resolor af.
- #9 #6 OR #7 OR #8.
- #10 #5 AND #9.
- #11 random* af.
- #12 #10 AND #11.

Study selection

Two investigators (TY and LKW) filtered the original studies independently. If the literature meets the eligibility criteria, the two investigators will further read the full text to screen the study. Any discrepancies were addressed by discussion or third party consensus.

Data extraction and analysis

All data were collected independently by two investigators (TY and BYC) from eligible RCTs using a standardised form. The following information was extracted: (1) study identification, including author name and publication year; (2) country where the study was conducted; (3) study subjects, number of participants and male to female ratio; (4) settings of trial arms and specific dose of prucalopride; (5) the primary outcome was SBMs per week, while the secondary outcome was treatment-emergent adverse events (TEAEs), defined as the total complications following prucalopride application, including headache, diarrhoea, dizziness, nausea, vomiting and so on, and were collectively reported in original trials and diagnosed using the same criteria. If insufficient details were reported, the authors were contacted for further information. Disagreements between two investigators were addressed by discussion and consensus.

Quality assessment

The Cochrane Collaboration’s tool for assessing risk of bias and the Jadad scoring system were employed for quality evaluation. Randomisation, double blindness and dropout/withdrawal management were the main components of the Jadad scale. Any disagreements during
assessment were resolved by iteration, discussion and consensus.

**Statistical analysis**

**Traditional pairwise meta-analysis with STATA**

All original data were performed using STATA V.13.0 software for Windows. Q test and F test were used to evaluate heterogeneity before pooled effects. The fixed-effect and random-effect models were based on the result of Q test and F test. A fixed-effect model was adopted if $I^2$ was $<50\%$ and $p>0.1$. Otherwise, a random-effect model was chosen. For the outcomes, the proportion of patients receiving SBMs per week was set as the dichotomous variable, and the pooled analysis of SBMs was expressed as unadjusted OR. The TEAE was also a dichotomous variable, and we intend to apply risk ratio (RR) for synthesised effect. The 95% CI was calculated simultaneously within pooled OR and RR. Possible sources of heterogeneity were explored by meta-regression performed via a fixed-effect model; the the restricted maximum likelihood (REML) estimation method proposed by Harbord et al.\(^4\) was applied in meta-regression. Sensitivity analysis was carried out by omitting each trial one by one. Possible publication bias was determined by Egger’s regression asymmetry test.\(^20\) Meanwhile, contour-enhanced funnel plot was applied to distinguish detailed reasons for publication bias.\(^21\)

**Optimal dose of prucalopride ranked by Bayesian analysis**

Bayesian framework provided a multiple-treatment meta-analysis model because it allows a feasible and practical estimation of direct and indirect comparisons based on probabilities, ranks or predictions.\(^22\)\(^23\) In the current study, different doses of prucalopride (1 mg, 2 mg and 4 mg) and placebo were set as individual treatment groups. A random-effects approach was applied within a Bayesian framework with the GeMTC package in R (V.3.6.0). The Markov chain Monte Carlo method was used for pooled estimates. To achieve posterior distribution, three independent Markov chains were run simultaneously. Running for 20000 adaptations and 100000 updates per chain, the updating frequency of chain per one update was set as 10. Running lengths were extended if the Markov chains had not converged.\(^24\) Probabilities of being superior were obtained from the posterior distribution. Ranks were obtained according to the estimated OR and RR. After posterior distributions were ensured, 95% credible interval (CrI) was derived from the 2.5th and 97.5th percentiles. The probability of the optimal dose of prucalopride was ranked using the surface under the cumulative ranking (SUCRA), which ranges between 0% and 100%.\(^22\) A parameter to rank different doses of prucalopride based on their probability of ranking first, second or third was used.

**RESULTS**

**Study selection outcome**

There were 532 relevant articles retrieved ultimately, among which 108 were duplicate articles. In total, 202 studies were excluded by title and abstract screening due to the studies being reviews, conference abstracts, animal experiments and case reports; this resulted in 162 articles. After examining the abstracts and full texts, 148 articles were further excluded. The meta-analysis finally included 14 studies\(^13\)\(^15\)\(^25\)\(^35\) that fulfilled the inclusion criteria (figure 1).

**Study characteristics**

In total, 4328 patients were included in the 14 studies,\(^13\)\(^15\)\(^25\)\(^35\) The mean sample size was 309 participants, ranging between 37\(^27\) and 713\(^15\) patients. Table 1 displays the main characteristics of the 14 studies. Overall, 14 studies were published between 2001 and 2015 years, and 12 studies\(^13\)\(^15\)\(^25\)\(^30\)\(^33\)\(^35\) were conducted in Western countries and 2 in Asian countries.\(^31\)\(^32\) With respect to the treatment protocols, seven trials\(^26\)\(^28\)\(^31\)\(^35\) had one-arm design, five trials\(^13\)\(^15\)\(^25\)\(^27\)\(^30\) had two-arm setting, and two trials\(^14\)\(^29\) had three-arm design. One trial\(^28\) used 0.5 mg prucalopride (PRU) versus placebo, 1 mg prucalopride was applied in 4 trials,\(^14\)\(^26\)\(^27\)\(^29\) 12 trials,\(^13\)\(^15\)\(^25\)\(^27\)\(^29\)\(^35\) employed 2 mg prucalopride versus placebo, and 6 trials\(^13\)\(^15\)\(^25\)\(^28\)\(^30\) made use of 4 mg prucalopride versus placebo. There were diverse durations, ranging from 1 to 24 weeks. To retrieve data and combine outcomes consistently, our study analysed SBMs per week and total TEAEs after using 1 mg, 2 mg and 4 mg doses of prucalopride.
### Table 1 Main information of the studies included in the meta-analysis

<table>
<thead>
<tr>
<th>Study identification</th>
<th>Publication year</th>
<th>Region</th>
<th>Total patients (n)</th>
<th>M/F (n)</th>
<th>Treatment protocols</th>
<th>Treatment duration (weeks)</th>
<th>Clinical endpoints</th>
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<tr>
<td>Bouras et al&lt;sup&gt;25&lt;/sup&gt;</td>
<td>2001</td>
<td>USA</td>
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<td>74</td>
<td>NR</td>
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<td>4</td>
<td>Whole gut transit time, HR-QOL, oro-facial transit time, rectal sensitivity and psychological state.</td>
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<td>The Netherlands</td>
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<td>35</td>
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<td>10</td>
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<tr>
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<td>USA</td>
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<td>12</td>
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<td>Europe (multicentre)</td>
<td>370</td>
<td>NR</td>
<td></td>
<td>2 mg vs placebo</td>
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</table>

F, female; HR-QOL, health-related quality of life; M, male; NR, not reported; SBMs/week, spontaneous bowel movements per week.

### Study quality assessment

Methodological quality assessment and outline of the 14 studies are presented in figure 2A,B. Randomised sequence generation and allocation concealment were identified adequately in all trials. These trials were placebo-controlled with either single or double blinding, except one study<sup>25</sup>.
Significant heterogeneity in 4 mg

Test results revealed there was no statistically significant heterogeneity in 1 mg (figure 4A), moderate heterogeneity in 2 mg ($I^2=60.5\%$ and $Q$ test $p=0.005$; figure 4B) and no significant heterogeneity in 4 mg ($I^2<50\%$ and $Q$ test $p>0.1$; figure 4C) prucalopride versus placebo. Thus, a fixed-effect model was applied for pooled analysis in 1 mg and 4 mg groups. A random-effect model was selected for 2 mg pooled analysis. In the pooled meta-analysis, the results showed that the incidence of TEAEs increased significantly after using 2 mg prucalopride ($Z=3.57$, $p=0.000$; RR: 1.20, 95% CI 1.09 to 1.33) and 4 mg prucalopride ($Z=3.82$, $p=0.000$; RR: 1.14, 95% CI 1.07 to 1.22). Meanwhile, 1 mg prucalopride did not reach statistical significance after pooled analysis ($Z=1.42$, $p=0.154$; RR: 1.17, 95% CI 0.94 to 1.44). Findings are shown in figure 4, along with RR.

Bayesian analysis for optimal dose of prucalopride
Bayesian methodology was used to determine the optimal dose of prucalopride in the treatment of CIC through multiple-treatment meta-analysis models. Different doses of prucalopride reported in order of SBMs per week and TEAEs ranked according to SUCRA are shown in figure 5. We reported the consistency model results of SBMs per week and TEAEs after confirming there was no statistically significant inconsistency and the three Markov chains converged (potential scale reduction factor=1). SBMs per week and TEAEs of 1 mg, 2 mg and 4 mg prucalopride reached statistical significance compared with placebo. After statistical comparison, we revealed that 1 mg prucalopride demonstrated the maximum frequency of SBMs per week (OR: 3.31, 95% CI 1.72 to 6.16, table 2; probability rank=0.70, figure 5A) indirectly compared with 2 mg and 4 mg prucalopride. In terms of TEAEs, 1 mg prucalopride also holds the highest incidence (RR: 1.72, 95% CI 1.02 to 2.92, table 2; probability rank=0.47, figure 5B) compared with 2 mg and 4 mg prucalopride. Meanwhile, 4 mg prucalopride has the second probability sequence (probability rank=0.33; figure 5B).

Sensitivity analysis
We carried out sensitivity analysis to verify the robustness of our results. The results are shown in figure 6. We performed sensitivity analysis via the leave-one-out procedure each time, and the outcomes of the sensitivity analysis showed that exclusion of any study did not significantly account for heterogeneity, demonstrating our results were robust to some extent.

Publication bias
Egger’s test and contour-enhanced funnel plot were employed to assess potential publication bias. First,
education and develop an effective pharmacotherapy in the management of CIC. The results of the pooled analysis are presented in figure 3, along with OR.

TEAEs following different doses of prucalopride versus placebo
Before pooled analysis, heterogeneity across studies was tested conventionally. The results of the heterogeneity test revealed there was no statistically significant heterogeneity in 1 mg (figure 4A), moderate heterogeneity in 2 mg ($I^2=60.5\%$ and $Q$ test $p=0.005$; figure 4B) and no significant heterogeneity in 4 mg ($I^2<50\%$ and $Q$ test $p>0.1$; figure 4C) prucalopride versus placebo. Thus, a fixed-effect model was applied for pooled analysis in 1 mg and 4 mg groups. A random-effect model was selected for 2 mg pooled analysis. In the pooled meta-analysis, the results showed that the incidence of TEAEs increased significantly after using 2 mg prucalopride ($Z=3.57$, $p=0.000$; RR: 1.20, 95% CI 1.09 to 1.33) and 4 mg prucalopride ($Z=3.82$, $p=0.000$; RR: 1.14, 95% CI 1.07 to 1.22). Meanwhile, 1 mg prucalopride did not reach statistical significance after pooled analysis ($Z=1.42$, $p=0.154$; RR: 1.17, 95% CI 0.94 to 1.44). Findings are shown in figure 4, along with RR.

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Publication bias
Egger’s test and contour-enhanced funnel plot were employed to assess potential publication bias. First,
Figure 3  Forest plot for SBMs per week following different doses of prucalopride. (A) 1 mg prucalopride vs placebo; (B) 2 mg prucalopride vs placebo; and (C) 4 mg prucalopride vs placebo. SBMs, spontaneous bowel movements.
Figure 4  Forest plot for TEAEs following different doses of prucalopride. (A) 1 mg prucalopride versus placebo; (B) 2 mg prucalopride versus placebo; and (C) 4 mg prucalopride versus placebo. RR, risk ratio; TEAEs, treatment-emergent adverse events.
Egger's test was applied to assess potential publication bias for pooled SBMs per week and TEAEs as the results are quantitative. The results of the Egger's test revealed there was no publication bias in both synthesised analyses (pSBMs per week = 0.929, pTEAEs = 0.483; figure 7A,B). Then, contour-enhanced funnel plot, which added conventional milestones in levels of statistical significance (p<0.01, p<0.05, p<0.1 or p>0.1) to funnel plots, was used to distinguish detailed reasons for publication bias. The results indicated many missing studies are of higher statistical significance (p<0.01; figure 7C,D), which suggested the origin of asymmetry may be more likely to be due to undetected factors rather than publication bias. Finally, we traced the original research again, speculating that studies with a small sample size and with ITT analysis missing in many studies may account for the undetected factors rather than publication bias. Finally, we traced the original research again, speculating that studies with a small sample size and with ITT analysis missing in many studies may account for the undetected factors rather than publication bias. Overall, the findings of the current study acknowledge that patients with CIC can be treated with prucalopride efficaciously, whether 1 mg, 2 mg or 4 mg doses are used. The certainty of conclusion from the current study is mainly reflected in the following two aspects. First, the proportion of SBMs per week was employed as the cardinal endpoint in the current study for pooled meta-analysis. According to the definition of SBMs per week based on the Rome criteria, SBMs per week can incorporate anorectal assessments of rectal evacuation and subjective symptoms completely. In the current study, SBMs per week increased significantly after using 1 mg (OR=2.40, 95% CI 1.32 to 4.37), 2 mg (OR=2.55, 95% CI 1.93 to 3.36) and 4 mg (OR=2.51, 95% CI 1.92 to 3.28) prucalopride. The results demonstrated that patients with CIC may benefit from prucalopride using different doses. Second, additional indicators of CIC, such as the Patient Assessment of Constipation Quality of Life and the Patient Assessment of Constipation Symptoms, also got ameliorated according to original researches.

| Table 2 Bayesian analysis of SBMs/week and TEAEs following different doses of prucalopride |
|-------------------------------------------------|---------------------------------|---------------------------------|---------------------------------|
| 1 mg OR (95% CI)                                  | RR (95% CrI)                    | RR (95% CrI)                    | RR (95% CrI)                    |
| 1.27 (0.63 to 2.48)                              | 1.72 (1.02 to 2.92)             | 1.61 (1.28 to 2.06)             | 1.67 (1.23 to 2.40)             |
| 2 mg OR (95% CI)                                  |                                 | 1.03 (0.57 to 1.80)             | 0.97 (0.67 to 1.33)             |
| 4 mg OR (95% CI)                                  |                                 |                                 | 1.07 (0.63 to 1.82)             |
| Placebo OR (95% CI)                               |                                 |                                 |                                 |

Significant results are in bold.

- Treatment; SBMs/week, OR (95% CrI); TEAEs, RR (95% CrI).
- 95% CrI, 95% credible interval; RR, risk ratio; SBMs/week, spontaneous bowel movements per week; TEAEs, treatment-emergent adverse events.
Third, a prospective, randomised, cross-over study with 2 mg prucalopride versus polyethylene glycol 3350 + electrolytes (PEG3350) also demonstrated prucalopride had obvious advantages over PEG3350 in promoting high-amplitude propagating contractions in patients with CIC. All supporting evidence mentioned above demonstrates prucalopride should be an effective pharmacotherapy in the management of CIC. We tried to provide the optimal dose of prucalopride in CIC management through Bayesian framework models. The results revealed that 1 mg, 2 mg and 4 mg prucalopride reached statistical significance compared with placebo, which is consistent with a previous pairwise meta-analysis. Moreover, we revealed 1 mg prucalopride obtained the maximum frequency of SBMs per week (OR: 3.31, 95% CrI 1.72 to 6.16, probability rank=0.70) indirectly compared with 2 mg and 4 mg doses. A previous study showed prucalopride applied at 4 mg dose in the experiment arm did not provide a dose-dependent benefit in SBMs per week over 2 mg dose (4 mg 28.4% vs 2 mg 30.9%). Another study also failed to achieve higher frequency of SBMs per week with a dose of 4 mg over the 2 mg dose (4 mg 24% vs 2 mg

Figure 6  Sensitivity analysis for (A) spontaneous bowel movements per week and (B) treatment-emergent adverse events.

Figure 7  Publication bias assessment. (A) Egger’s test for SBMs per week; (B) Egger’s test for TEAEs; (C) contour-enhanced funnel plot for SBMs per week; and (D) contour-enhanced funnel plot for TEAEs. SBMs, spontaneous bowel movements; TEAEs, treatment-emergent adverse events.
Considering all these, we arrived at the conclusion that there is no incremental benefit in prucalopride application and that 1 mg dose could be safer in treating CIC.

TEAEs related to prucalopride consumption were evaluated in this meta-analysis. When reported, headache, nausea, diarrhoea and abdominal pain are the most common TEAEs following prucalopride consumption. However, the severity of these TEAEs was slight or mild mainly during the first 24 hours of taking and lasts ephemerally. Our findings revealed the incidence of TEAEs increased significantly after using 2 mg dose (RR: 1.20, 95% CI 1.09 to 1.33) and 4 mg dose (RR: 1.14, 95% CI 1.07 to 1.22). Meanwhile, the 1 mg dose did not reach statistical significance after pooled analysis (RR: 1.17, 95% CI 0.94 to 1.44). Bayesian framework analysis showed the 1 mg dose holds the highest incidence (RR: 1.72, 95% CrI 1.02 to 2.92, probability rank=0.47) compared with the 2 mg and 4 mg doses indirectly. Meanwhile, the 4 mg dose has the second probability sequence (RR: 1.61, 95% CrI 1.28 to 2.06, probability rank=0.33) and the 2 mg dose ranked the lowest (RR: 1.67, 95% CrI 1.23 to 2.40, probability rank=0.2). Clearly, the probability rank of 1 mg, 2 mg and 4 mg doses is lower than 50%. TEAEs of 5-HT4 receptor agonists have received considerable attention, particularly cardiovascular adverse events. An earlier study suggested that no consistent or clinically relevant TEAE divergences were noted in ECG components versus baseline and placebo groups, even in 88% of volunteers with a history of cardiovascular diseases. A thorough QT study, based on the ICH E14 guidance, also demonstrated no evidence of QT prolongation or proarrhythmic potential for prucalopride application. This evidence suggests prucalopride has good tolerance in cardiovascular system and overall TEAEs.

Figure 8 Univariate meta-regression analysis: (A) spontaneous bowel movements per week of 2 mg prucalopride and (B) treatment-emergent adverse events of 2 mg prucalopride. RR, risk ratio.

Table 3 Meta-regression analysis of 2 mg SBMs per week (OR) and TEAEs (RR)

<table>
<thead>
<tr>
<th>Covariates</th>
<th>2 mg SBMs per week, OR (11 studies)</th>
<th>2 mg TEAEs, RR* (11 studies)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coefficient 95% CI† P value Tau2</td>
<td>Coefficient 95% CI† P value Tau2</td>
</tr>
<tr>
<td>Univariate analysis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethnicity in different regions</td>
<td>0.70 0.14 to 1.27 0.020 0.023 0.24</td>
<td>0.03 to 0.46 0.002 0.006</td>
</tr>
<tr>
<td>Total sample size</td>
<td>0.00 −0.001 to 0.004 0.332 0.095 0.08</td>
<td>−0.55 to 0.71 0.786 0.021</td>
</tr>
<tr>
<td>Courses of taking prucalopride</td>
<td>2.09 −1.01 to 5.20 0.144 0.025 0.14</td>
<td>−0.35 to 0.63 0.514 0.026</td>
</tr>
<tr>
<td>ITT analysis</td>
<td>0.10 −1.10 to 1.30 0.856 0.114 0.01</td>
<td>−0.37 to 0.38 0.953 0.024</td>
</tr>
<tr>
<td>Multivariate analysis</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethnicity in different regions</td>
<td>0.70 −0.03 to 1.45 0.058 NA 0.31</td>
<td>0.07 to 0.55 0.017 NA</td>
</tr>
<tr>
<td>Total sample size</td>
<td>0.00 −1.97 to 1.97 0.999 NA 0.37</td>
<td>−0.33 to 1.07 0.688 NA</td>
</tr>
<tr>
<td>Courses of taking prucalopride</td>
<td>−0.04 −0.34 to 0.25 0.723 NA 0.09</td>
<td>−0.03 to 0.22 0.108 NA</td>
</tr>
<tr>
<td>ITT analysis</td>
<td>0.05 −0.78 to 0.89 0.873 NA −0.04</td>
<td>−0.25 to 0.18 0.688 NA</td>
</tr>
<tr>
<td>Omnibus test for moderators</td>
<td>0.897 0.051 0.108 0.003</td>
<td></td>
</tr>
</tbody>
</table>

Significant results are in bold.
ITT, intention-to-treat; NA, not applicable; RR, risk ratio; SBMs, spontaneous bowel movements; TEAEs, treatment-emergent adverse events.
The current study exerts more attention to the incremental benefit of prucalopride. Previous work has provided critical findings that prucalopride is able to treat CIC effectively with mild, transitory and slight by-effects. However, the dose-related trends in efficacy assessment and TEAEs have not been explored. In this study, the results revealed that there is no incremental benefit in prucalopride application and 1 mg dose is suggested to be given at commencement. Furthermore, clinical evidence from China with 313 samples was included in the current study. The study of Zou et al could be an eventful supplement to current evidence and fill the gap in knowledge about use of prucalopride in China, even in the Asian region. Finally, the current study is the most extensive meta-analysis based on 14 well-designed RCTs with 4328 patients, providing more powerful evidence in favour of the integrated analysis.

Potential limitations of this integrated analysis should be considered. Fourteen included trials with variable duration (from 1 to 24 weeks) measured endpoints by different criteria, such as colonic transit time, rectal sensitivity, anorectal function and SBMs per week, increasing the risk of measurement bias. Meanwhile, undetected bias predicted by contour-enhanced funnel plot showed studies with a small sample size and with IIT analysis missing may account for potential bias. These factors may have a potential impact on our conclusions. Meta-regression by univariate and multivariate analyses found different patient ethnicities (Asian and Western) were a potential covariate causing moderate heterogeneity ($I^2=60.5\%$) and de-escalate the validity of the results of the current analysis.

All in all, this is the largest meta-analysis with 4328 patients from 14 high-quality RCTs and provides pivotal evidence that prucalopride given at 1 mg dose can manage CIC effectively. Furthermore, no incremental benefit was detected in prucalopride application. When accepting the conclusions of this study, the methodological limitations should be considered at the same time. It is widely recognised that CIC is a chronic disease of intestinal dysfunction and requires long-term medication, so more RCTs of larger scale and with longer-term efficacy assessments are needed to balance the risk–benefit profile of prucalopride in the treatment of CIC.

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**Acknowledgements** Our greatest acknowledgement goes to the authors who made detailed data available for this meta-analysis and all our colleagues in this study for their hard work.

**Contributors** TY, LKW and XJW performed the search and drafted the manuscript. TY and BYC performed the data extraction and analysed the data. TY, TX and DXY designed the study and amended the original draft.

**Funding** The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

**Competing interests** None declared.

**Patient and public involvement** Patients and/or the public were not involved in the design, conduct, or reporting, or dissemination plans of this research.

**Patient consent for publication** Not required.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Data availability statement** All data relevant to the study are included in the article or uploaded as supplementary information.

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