# Berberine for diarrhea in children and adults: a systematic review and meta-analysis

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## Abstract

**Background:** Diarrhea is a ubiquitous digestive system disease, leading to loss of fluid and electrolytes, and may be life-threatening, especially in children and adults who are immunosuppressed or malnourished. Berberine has a broad-spectrum antibiotic activity and is very widely used to treat diarrhea in China. No systematic review has been carried out to evaluate the evidence presented in clinical trials. The aim of this study was to assess the effectiveness and safety of berberine in diarrhea treatment among children and adults. **Methods:** Seven databases and two clinical trial registries were searched on 1 September 2019. Randomized controlled trials were included, where participants were diagnosed (first diagnosed) as having diarrhea according to clear diagnostic criteria. Berberine alone or in combination with Western medication as intervention were included. Subgroup analyses were conducted based on children or adults, acute or persistent diarrhea, infectious or noninfectious and treatment courses. Primary outcomes were clinical cure rate and duration of diarrhea. The GRADE tool was used to assess the guality of evidence.

**Results:** A total of 38 randomized controlled trials were included involving 3948 participants (including 27 trials on 2702 children) were included. Compared with antibiotics, berberine plus antibiotics showed better results in both adults and in children in general, especially when given for 7 days or 3 days in acute infectious diarrhea of children. Compared with the control groups, using berberine alone or in combination with montmorillonite, probiotics, and vitamin B increased the clinical cure rate of diarrhea. The use of berberine alone or berberine had significantly better laboratory indicators (isoenzyme, inflammatory factors, myocardial enzyme, and fecal trait) and fewer systemic symptoms than the no berberine groups. Overall, 22 of 27 trials on children used berberine as an enema. No deaths and serious adverse events were reported. The quality of evidence of included trials was moderate to low or very low. The impact of different dosages, frequencies and treatment durations on the outcomes was not evaluated due to insufficient number of trials.

**Conclusion:** This review demonstrated that berberine was generally effective in improving clinical cure rates and shortening the duration of diarrhea compared with control groups. No severe adverse event was reported. However, there is still a lack of high-quality evidence for evaluating the efficacy and safety of berberine.

**Trial registration:** PROSPERO CRD42020151001 (available from http://www.crd.york.ac.uk/ PROSPERO/).

Keywords: antibiotic, berberine, diarrhea, inflammatory factors, isoenzyme, myocardial enzyme

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#### Introduction

Diarrhea is defined as the passage of three or more loose or liquid stools per day (or more frequent passage than what is typical for the individual).<sup>1</sup> It is a ubiquitous digestive system disease. Globally, there are nearly 1.7 billion cases of childhood diarrhea disease every year.<sup>2</sup> Continued diarrhea can cause loss of fluid and electrolytes, and may even become life-threatening, especially in children and adults who are immunosuppressed or malnourished.3 Although not life-threatening for adults, diarrhea still has a high incidence.1 Diarrhea usually has intestinal tract noninfectious symptoms that could be caused by improper diet and malnutrition, and infectious symptoms that are usually caused by a variety of bacterial, viral, and parasitic organisms.<sup>4</sup> According to the World Gastroenterology Organisation Global guidelines,1 the etiological diagnosis of diarrhea usually is based on medical history, symptoms, routine testing, and especially stool testing. In many countries, the goal of diarrhea treatment is to relieve symptoms and avoid complications.5 Commonly used treatments include probiotics,6 zinc,7 lactose-free formula,8 antibiotics, and antidiarrheal agents (such as montmorillonite).9

ingredient The main of berberine is C20H19NO5, an isoquinoline alkaloid belonging to the structural class of proto-berberines extracted from traditional Chinese herbal medicines, mainly rhizoma coptidis and rhizoma phellodendri,<sup>10,11</sup> which have been used in China for more than 2000 years treating diarrhea.<sup>12</sup> Pharmacological studies using doses of berberine (50-200 mg/kg per time, one time a day)<sup>13</sup> substantially exceed those likely to occur in humans (0.1-0.3 g per time, three times a day) have demonstrated that berberine has a broad-spectrum antibiotic activity, including Vibrio cholera, Shigella and Pseudomonas14-16 and have approved that berberine has advantages in treating intestinal bacterial infections, including bacterial dysentery and viral infections.<sup>10,17-20</sup> Escherichia coli, Klebsiella pneumoniae and Proteus mirabilis can also be suppressed by berberine.<sup>21</sup> In addition, berberine has broad prospects in the treatment of secretory diarrhea, it can inhibit the hypersecretion of electrolytes caused by cholera toxin.<sup>13</sup> It has effects on reducing inflammation, inhibition of gastrointestinal motility, ameliorates impaired gastrointestinal function, and reducing intestinal secretion and exudation.<sup>21</sup> Berberine can relax intestinal smooth muscle and prolong the residence time of intestinal contents to fully digest and absorb the intestinal contents to treat functional diarrhea.<sup>22</sup> Therefore, berberine has a significant effect on infectious or noninfectious, secretory or exudative diarrhea.<sup>21</sup> In clinical practice, berberine is sometimes combined with Western medicine in the treatment of diarrhea, such as montmorillonite, antibiotics, probiotics, vitamin B or anisodamine. However, there are no systematic reviews that have assessed efficacy and safety.

This systematic review aims to evaluate the effectiveness and safety of berberine alone or in combination with Western medicine in diarrhea treatment, and to provide evidence for clinical practice in this specific area.

#### Methods

This review was registered in PROSPERO (CRD42020151001), and the protocol is available at http://www.crd. york.ac.uk/PROSPERO/.

#### Eligibility criteria

This review included randomized controlled trials (RCTs) enrolling participants needed to be diagnosed (first diagnosed) with diarrhea according to clear diagnostic criteria, regardless of sex, age, and type of diarrhea (infectious, noninfectious, acute, and chronic). Diarrhea caused by drugs (such as antibiotics) or diseases (such as cancer) were excluded. All randomized controlled studies, regardless of language or publication date or state, were screened. Acute diarrhea was defined as a duration of diarrhea less than 14 days, while diarrhea was considered as persistent if lasting longer than 14 days.<sup>23</sup> Usage of other Western medicine was permitted as long as they were antibiotics, fluid therapy (including oral rehydration salts, intravenous rehydration, electrolyte supplementation and correction of acid-base balance disorders), antipyretic, adjustment of diet/diet guidance/nutritional guidance, supplementation of trace elements and calories. Secondary diarrhea was excluded from this review. Based on realworld clinical practice, to include combinations from clinical practice to a large extent, we categorized interventions based on different combinations (berberine + montmorillonite; berberine + Bifidobacterium subtilis; berberine + montmorillonite + vitamin B; berberine + Bifidobacterium lacto*bacillus* triple viable + montmorillonite; berberine +

montmorillonite + anisodamine) in this systematic review.

#### Search strategy

Published studies were comprehensively searched in the following databases from their inception to April 2019: PubMed, the Cochrane Library, EMBASE, Chinese National Knowledge Infrastructure Database (CNKI), VIP, Wanfang, and SinoMed. We also searched for two clinical trial registration networks (ClinicalTrials.gov and Chinese Clinical Trial Registry). The detailed search strategy is provided in the Appendix.

## Study outcomes

The primary outcome measures were clinical cure rate (as defined in each original study) and duration of diarrhea (day). Secondary outcomes included stool frequency (number of depositions per day), stool output (g or ml/kg per day), fecal trait improvement, absenteeism, intensity of antibiotic use, stool routine examination, stool bacterial culture, duration of hospitalization, duration of other symptoms (such as vomiting and fever), quality of life, recurrent diarrhea, laboratory indicators, death (all-cause and diarrhea-related), adverse events, and adverse drug reactions.

## Study selection

The retrieved literature was imported into NoteExpress, and duplicate records were removed. Title/abstract screening and full-text screening were conducted by two reviewers (MK Yu and CH L) independently.

Any disagreement between the two reviewers was resolved through discussing with a third reviewer (or a senior author, YT Fei).

## Data extraction

Data extraction was conducted by two pairs of reviewers (YM and CL, DP and QH) using a standardized, pre-piloted data extraction form, including authors information, characteristic of participants, details of interventions and controls, outcomes, and information related to study design. Two authors in each group independently and duplicated extracted data from each trial, cross-checked the data. Discrepancies were solved by discussing within the pair of reviewers or arbitrated by the senior author (YF) if necessary.

## Quality assessment and publication bias

We assessed each included study's risk of bias based on a modification of the Cochrane Risk of Bias tool, which consists of the following aspects: random sequence generation (selection bias); allocation concealment (selection bias); blinding (performance bias and detection bias); incomplete outcome data (attrition bias); selective reporting of outcomes and other bias. This modified tool has four response options for each aforementioned aspect: 'low risk of bias,' 'probably low risk of bias,' 'probably high risk of bias,' and 'high risk of bias'.24 Publication bias was assessed using the funnel plot when there were more than 10 studies available for a certain outcome/comparison. In addition, we used the Grading of Recommendation, Assessment, Development, and Evaluation (GRADE) system to grade the quality of evidence for primary outcomes.<sup>25</sup>

## Data synthesis

Review Manager 5.3 was used. Continuous outcomes were analyzed by standardized mean difference (MD). We assessed the dichotomous variable by relative risk [(RR), Mantel–Haenszel RR]. The *p*-value and 95% confidence interval (CI) were obtained.

For those publications that failed to report any critical information, we tried to contact the corresponding author or first author to obtain the information *via* email or telephone calls.

# Subgroup analysis and investigation of heterogeneity

The heterogeneity was assessed according to  $I^2$  statistic. When heterogeneity was not significant  $(I^2 < 50\%)$ , we used a fixed-effect model to synthesize the data. When heterogeneity was significant  $(I^2 \ge 50\%)$ , the random-effects model was used.

The following potential sources of heterogeneity (ages; different drugs of berberine combined; a degree of diarrhea; route of administration; duration of treatment; type of primary disease; time of observation; drug dosage) were explored in subgroup analyses. We performed a sensitivity

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#### Figure 1. Study flow diagram.

CBM, SinoMed Database; CNKI, Chinese National Knowledge Infrastructure Database; VIP, VIP Chinese Science and Technique Journals Database; WanFang, Wanfang Database.

analysis to assess the robustness of the meta-analysis by excluding trials with poor methodological quality (those with insufficient randomization methods and trials with selective reporting bias).

#### Results

In total, we identified 10,010 studies through the searches, and a total of 38 studies were included (Figure 1).

## Study characteristics

Overall, 38 RCTs involving 3948 participants<sup>10,26-62</sup> were included in this review. A total of 11 studies<sup>10,28,32,33,37,39,40,42,44</sup> were conducted on adults (1246 of 3948, 31.56%), aged 16–76 years, with the remaining 27 studies<sup>26,27,29–31,34,35,37,38,41,43,45–62</sup> conducted on children (2702 of 3948, 68.4%)

aged from 2 months to 12 years. A total of 22 studies<sup>29,34,35,37,38,41,45-49,51-57,59-62</sup> involved taking a berberine enema (all in children), and the other studies<sup>10,26-28,30-33,36,39,40,42-44,50,58</sup> involved taking berberine orally. Sex was not reported in two studies.<sup>33,43</sup> In the remaining 36 studies,<sup>10,26–32,34–42,44–62</sup> the proportion of males to females was about 1:1.2. Overall, 31.6% studies were for infectious diarrhea, while 15.8% were mixed and 52.6% were not reported. A total of 19 studies were treating acute diarrhea, while 4 studies<sup>28,36,44,62</sup> were mixed acute and persistent, 632,33,37,39,48,54 exclusively with persistent diarrhea, and 13 studies<sup>25,27,29-31,34,35,38,45,49,52,57,61</sup> did not specify. The principle investigators of 37 studies came from a Western medicine hospital, and only 1 study came from a traditional Chinese medicine hospital. In the included studies that used antibiotics, 23 studies did not report the cause of diarrhea, 10

studies were due to virus of bacteria infection, 3 studies were due to bacteria infection, and 2 studies were virus infection.

The sample size ranged from 30 to 279 patients for each study, with a mean of 104 patients. Overall, 34 studies<sup>26–31,34–38,40–62</sup> had two arms, 3 studies had three arms,<sup>32,33,39</sup> and 1 trial had five arms.<sup>10</sup> There were 14 studies that mentioned the inclusion criteria,<sup>10,28,33,39,42–44,48,51–53,58–60</sup> and the remaining studies did not. The dosage and the frequency of oral berberine was 0.05–0.5 g and 2–3 times per day, while the dosage and the frequency of enema berberine was 10–20 mg and 1–2 times per day (Table 1).

## Risk of bias in included studies

All studies were considered high risk of bias, mainly due to unclear concealment (36 of 38, 94.7%), no blinding (37 of 38, 97.3%) and selective reporting of outcomes (11 of 38, 28.9%; Figure 2).

## **Effect estimates**

Data analysis was performed according to the comparison. We could not summarize data or perform meta-analyses due to inclusion of just one study.

## Primary outcomes

A total of 35 studies reported clinical cure rate, with a largely similar definition, that was the symptoms of diarrhea disappeared, and fecal trait and the stool frequency returned to normal within 48h or 72h or 100h (as defined in each original study) after treatment.

## Clinical cure rate

Comparison 1: berberine versus no berberine. Overall, a meta-analysis from 17 studies<sup>26-39,41-44</sup> showed that compared with no berberine groups, treatment with berberine resulted in a more significant number of patients being cured of diarrhea (RR 1.37, 95% CI 1.27–1.47,  $I^2=5\%$ , n=1684, 17 trials, moderate certainty evidence; Figure 3). A funnel plot for these 17 studies appeared to be symmetrical, indicating no severe publication bias (Figure 4). In the subcomparison, that is, berberine + antibiotics versus antibiotics, there is insufficient evidence to prove that berberine has an advantage in treating children with diarrhea on day 3.28,29 One study37 found that berberine does not have an advantage in treating children with persistent infectious diarrhea on day 3. Other studies have reported advantages of berberine in treating diarrhea.<sup>26,28,30,31,36,39,41</sup> In berberine versus antibiotics, a study<sup>43</sup> reported that berberine could cure more children with infectious diarrhea compared with antibiotics on day 3. But there was no statistically significant between the experiment group and control group for adults in other studies.42,44 In berberine + probiotics *versus* probiotics, three studies<sup>32,33,39</sup> showed that berberine has an advantage for adults with persistent diarrhea. In berberine + intestinal mucosa protection + antibiotics versus intestinal mucosa protection + antibiotics, the cure rate of berberine groups was significantly higher than the control group for children with diarrhea.34,38

A sensitivity analysis exploring the effect of randomization, allocation concealment (Appendix 1), and selective reporting bias (Appendix 2) did not change the result of the overall meta-analysis significantly, although it changed the results of some subgroup analyses.

Comparison 2: berberine + montmorillonite versus no berberine + montmorillonite. Overall, 13 studies<sup>45–57</sup> evaluated clinical cure rates. Treatment with berberine and montmorillonite resulted in a more significant number of children being cured of acute diarrhea (RR 1.70, 95% CI 1.43 to 2.02,  $I^2$ =55%; n=1146, 13 trials, verylow-certainty evidence, Appendix 3). On visual inspection, the funnel plot was asymmetric, with most studies centered together on the upper left (Appendix 4).

In the subcomparison (berberine + montmorillonite + antibiotics *versus* antibiotics), 12 studies reported that berberine combined with montmorillonite and antibiotics had an advantage for children with diarrhea than antibiotics alone.<sup>45–49,51,57</sup> In berberine + montmorillonite *versus* antibiotics, compared with the antibiotics group, the cure rate of berberine combined with montmorillonite and antibiotics was significantly higher for adults in acute diarrhea.<sup>50</sup>

Table 1. Ch	aracteristics	of the included (	38 studies.							
Study ID	Sample T: (F), C: (F)	Mean/range age (T/C)	Population	Acute/ persistent/ both/NR	Cause of diarrhea	Course of diarrhea (d)	Intervention	Route of administration	Control intervention	Outcome measures
Berberine v	ersus no berbe	rine								
Yang <sup>26</sup>	T: 38 (16) C: 38 (17)	T: 14.6 m C: 14.9 m	Children	Acute	Virus or bacterial infection	Ш	1. BE: 0.2g, 3times/d, for 7 d 2. AMC: 30 mg/kg, 2times/d, for 7 d*	ю. d	AMC: 30 mg/kg, 2 times/d, for 7 d*	10
Li <sup>27</sup>	T: 35 (17) C: 35 (16)	T: 2y C: 1y	Children	л Х	л Х	ж Z	1. BE: 0.2g, 3times/d, for 7 d 2. LOF: 200 ml, 1 times/d, for 7 d*	o. d	LOF: 200 ml, 1 times/d, for 7 d*	(1)
Lin <i>et al.</i> <sup>28</sup>	T: 49 (23) C: 49 (21)	T: 36.61y C: 37.03y	Adult	Both	Bacterial infection	T: $6.36 \pm 2.58d$ C: $6.44 \pm 2.63d$	1. BE: 0.3 g, 3 times/d 2. LOF: 500 ml 1 times/d*	p.o.	LOF: 500 ml 1 times/d*	19B
Wei and Li <sup>29</sup>	T: 15 C: 15	T: 3–13 y C: 3–13 y	Children	NR	NR	NR	<ol> <li>BE: 10 mg/kg, 1 times/d, for 3 d</li> <li>Antibiotic*</li> </ol>	p.r.	Antibiotic*	(]
Wang <sup>30</sup>	T: 45 (21) C: 45 (20)	T: 1.68 <i>y</i> C: 1.65 <i>y</i>	Children	х К	Virus or bacterial infection	ЖZ	1. BE: 0.2g, 3times/d, for 7 d 2. LOF: 200 ml, 1 times/d, for 7 d*	o. d	LOF: 200 ml, 1 times/d, for 7 d*	(1)
Cheng et al. <sup>31</sup>	T: 29 (13) C: 29 (14)	T: 1.60 y C: 1.64 y	Children	R	Virus or bacterial infection	ЖZ	1. BE: 0.2g, 3times/d, for 7 d 2. LOF: 200 ml, 1 times/d, for 7 d*	o. d	LOF: 200 ml, 1 times/d, for 7 d*	(1) $(1)$ $(1)$
Luo <sup>32</sup>	T1: 35 (16) T2: 31 (13) C: 33 (17)	T1: 37.5y T2: 32.1y C: 34.4y	Adult	Persistent	л Х	T1: 12.3 ± 1.1m T2: 10.5 ± 2.3m C: 11.9 ± 1.0m	1. BE: 0.2g, 3times/d, for 21d 2. BLT: 0.63g, 2times/d, for 21d	p.o.	BLT: 0.63 g, 2 times/d, for 21 d	126
Ye <sup>33</sup>	T1: 30 T2: 30 C: 30	T1: 19–67 <i>y</i> T2: 19–67 <i>y</i> C: 19–67 <i>y</i>	Adult	Persistent	х Х	ш Z	1. BE: 0.2g, 3times/d, for 14d 2. BLT: 0.63g, 2times/d, for 14d	o. d	BLT: 0.63 g, 2 times/d, for 14 d	(12)
Zhou <sup>34</sup>	T: 100 (48) C: 98 (53)	T: 1.4y C: 1.3y	Children	R	Virus or bacterial infection	Х Х	1. BE: <1y 0.05 g; >1 y 0.1g, 3 times/d 2. MN: <1y 1g, 3 times/d >1y 2g, 3 times/d 3. Antibiotic*	ч	1. MN: <1y 1g, 3times/d >1y 2g, 3times/d 2. Antibiotic*	<b>(</b> ]
Huang et al. <sup>35</sup>	T: 50 C: 50	T: 3-10y C: 3-10y	Children	Acute	NR	Х	1. BE: 0.1g, 3times/d, for 7 d 2. Amp:50 mg/kg, 2times/d, for 7 d*	p.r.	Amp:50 mg/kg, 2 times/d, for 7 d*	Ē
										(Continued)

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Study ID	Sample T: (F), C: (F)	Mean/range age (T/C)	Population	Acute/ persistent/ both/NR	Cause of diarrhea	Course of diarrhea (d)	Intervention	Route of administration	Control intervention	Outcome measures
Fu and Tan <sup>36</sup>	T: 53 C: 53	T: 18–60 y C: 18–60 y	Adult	Both	Virus or bacterial infection	≤21 d	1. BE: 0.3 g, 3 times/d, for 5 d 2. Antibiotic*	р.o.	Antibiotic*	(1)(E)
Huang and Zhang <sup>37</sup>	T: 40 (21) C: 40 (27)	T: 18.6 m C: 19.8 m	Children	Persistent	Viral infection	ЖZ	1. BE: 10 mg/kg, 2times/d, for 3d 2. BLT: <12 m 0.59-1 g, 3 times/d; 12-72 m 0.5g, 2 times/d, for 3 d*	р.г.	BLT: <12 m, 0.59-19, 3 times/d; 12-72 m0.59, 2 times/d, for 3 d*	(1)(3)(4)(0)
Li <sup>38</sup>	T: 60 C: 60	T: 4 m-2 y C: 4 m-2 y	Children	NR	NR	R	<ol> <li>BE: 20 mg/kg, 2times/d, for 3 d</li> <li>MN: NR, for 3d*</li> </ol>	Ľ. d	MN: NR, for 3d*	19
Hu <sup>39</sup>	T1: 20 T2: 20 C: 20	T1: 20-58y T2: 20-58y C: 20-58y	Adult	Persistent	NR	>2m	1. BE: 0.2g, 3 times/d, for 14d 2. BLT: 0.63g, 2 times/d, for 14d	р.о.	BLT: 0.63 g, 2 times/d, for 14 d	10
Zhao <sup>40</sup>	T: 146 C: 133	T: 14-63y C: 14-63y	Children	Acute	NR	NR	1. BE: 0.2–0.3 g, 3 times/d, for NR 2. LOF: 500ml, 1 times/d, for 3 d	р.о.	LOF: 500 ml 1 times/d, for 3 d	200
Xu <sup>41</sup>	T: 84 C: 84	T: 4 m-2 y C: 4 m-2 y	Children	Acute	Bacterial infection	≤14 d	<ol> <li>BE: 20 mg/kg, 2times/d, for NR</li> <li>Antibiotic*</li> </ol>	p.r.	Antibiotic*	(1)(2)
Zhang <sup>42</sup>	T: 30 (14) C: 30 (14)	T: 18y-65y C: 18y-65y	Adult	Acute	NR	≰2d	1. BE: 0.2g, 3 times/d, for 7 d	ю.	CPFX: 0.2g, 3times/d, for 7d.	(78) (5)
Dang <sup>43</sup>	T: 57 C: 45	T: 2y-12y C: 2y-12y	Children	Acute	NR	T: 48.4 ± 2.5h C: 47.9 ± 2.1h	BE: 2-3y 0.1g, 4-6y 0.1-0.157g, -9y 0.15g- 0.2g, 9-12y 0.2g-0.25g, 3 times/d, for 3d*.	Ъ.o.	CEC: 20 mg- 4o mg/kg, 3times/d, for 3d*	
Wu et al. <sup>44</sup>	T: 80 (35) C: 80 (30)	T: 44y C: 47y	Adult	Both	NR	Acute: <5 d Chronic: 2 m-1 y	BE: 0.5g, 2 times/d*	p.o.	Fur: 0.1g, 2 times/d*	8 (E)
Khin-Maung and Nyunt- Nyunt-Wai <sup>12</sup>	T: 45 (21) C: 45 (23)	T: 38.8 <i>y</i> C: 37.2 <i>y</i>	Adult	Acute	Bacterial infection	T: 10.6h C: 12.8h	BE: 0.1g, 4 times/d, for 3d*	р.о.	Tet: 500 mg, 4 times/d, for 3d*	2)(6)
Berberine +	montmorillor	nite <i>versus</i> no ber	-berine + mont	morillonite						
Liu <sup>45</sup>	T: 45 (22) C: 45 (21)	T: 1.5 y C: 1.6 y	Children	R	Virus or bacterial infection	ЖZ	<ol> <li>MN: NR, 1 times/d, for 3 d.</li> <li>BE: 10 mg/kg, 1 times/d, for 3 d</li> <li>Antibiotic*</li> </ol>	ь.г.	Antibiotic*	Ð
										(Continued)

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Table 1. (Co	ntinued)									
Study ID	Sample T: (F), C: (F)	Mean/range age (T/C)	Population	Acute/ persistent/ both/NR	Cause of diarrhea	Course of diarrhea (d)	Intervention	Route of administration	Control intervention	Outcome measures
Wang <sup>46</sup>	T: 40 (15) C: 30 (10)	T: 5 m–6 y C: 5 m–6 y	Children	Acute	ц К	2-7 d	1. MN: <1 1.5g, >1 y 3g, 2 times/d, for 3d. 2. BE: 10–20 mg/kg, 2 times/d, for 3d. 3. Antibiotic*	ц.	Antibiotic*	Ð
Jin <sup>47</sup>	T: 60 (18) C: 56 (20)	T: 2 m–3 y C: 2 m–3 y	Children	Acute	Viral infection	Х	1. MN: 3g, 2times/d, for 3d. 2. BE: 0.1g, 2times/d, for 3d 3. Antibiotic*	p.r.	Antibiotic*	(1)
Huang <sup>48</sup>	T: 32 (15) C: 24 (11)	Т: 17.68 ± 1.64 п С: 16.82 m	n Children	Persistent	ш Z	T: 3.54 ± 0.43 m C: 3.32 ± 0.62 m	1. MN: <1 31g, -3y 3-4.5g, 2 times/d, for 5d. 2. BE: <1 0.05-0.1g, 1-3y 0.1-0.2g, 2 times/d, for 5d. 3. Antibiotic*	Г.	Antibiotic*	Ð
Qiu <sup>49</sup>	T: 52 (22) C: 36 (16)	T: 6 m–3 y C: 6 m–3 y	Children	N Х	ш Z	R	1. MN: <1 1.5g, 1-3y 2g, >3y 3g, 1 times/d, for 3d. 2. BE: 10-15 mg/kg, 1 times/d, for 3d. 3. Antibiotic*	Ŀ.d	Antibiotic*	Ð
Qin <i>et al.</i> <sup>50</sup>	T: 60 C: 46	T: 18y C: 18y	Adult	Acute	Bacterial infection	К	1. MN: 3g, 3times/d, for 3d. 2. BE: 0.2–0.3g, 3times/d, for 3d.	b.o.	NOR: 0.3g, 3times/d, for 3d*	<b>1</b> (E)
Gan <sup>51</sup>	T: 51 (27) C: 51 (30)	T: 5–23 m C: 5–24 m	Children	Acute	к Z	T: 2–13d C: 2–14d	1. MN: <1 1.5g ≋1 y 3g, 2 times/d, for 5d. 2. BE: 10–15 mg/kg, 2 times/d, for 5d 3. Antibiotic*	Г d	Antibiotic*	Ð
Guo <sup>52</sup>	T: 30 (11) C: 28 (8)	T: 5 m–2.5 y C: 5 m–2.5 y	Children	R	N N	К	1. MN:1.5–3g, 1 times/d, for 5 d. 2. BE: 0.1–0.2g, 2 times/d, for 5 d.	p.r.	Antibiotic*	Θ
Wang <i>et al.</i> <sup>53</sup>	T: 48 [22] C: 42 [19]	T: 6–38 m C: 5–40 m	Children	Acute	Virus or bacterial infection	≤14 d	1. MN: <1 1.5g, 1-3y 2g, >3y, 3g, 2times/d, for 5d. 2. BE: 10-15 mg/kg, 2 times/d, for 5d. 3. Antibiotic*	Ŀ.d	Antibiotic*	Ð
Zhu <sup>54</sup>	T: 46 C: 56	T: 6 m – 30 m C: 6 m – 30 m	Children	Persistent	ж	T: 15.85 ± 2.43 d C: 16.30 ± 1.87 d	1. MN: 3g, 2times/d, for NR. 2. BE: 0.1g, 2times/d, for NR. 3. Antibiotic*	Ъ.г.	Antibiotic*	<b>(</b> ]

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(Continued)

Table 1. (C	ontinued									
Study ID	Sample T: (F), C: (F)	Mean/range age (T/C)	Population	Acute/ persistent/ both/NR	Cause of diarrhea	Course of diarrhea (d)	Intervention	Route of administration	Control intervention	Outcome measures
Shi <sup>55</sup>	T: 35 (19) C: 33 (20)	T: 6–38 m C: 5–40 m	Children	Acute	х Х	ж Z	1. MN: <11.5g, 1-3y 2g, >3y 3g/times, 1 times/d, for 3d. 2. BE: 10-15 mg/kg, 1 times/d, for 3d. 3. Antibiotic*	с. с	Antibiotic*	$\odot$
Wan and Qin <sup>56</sup>	T: 40 C: 40	T: 3 m-10 y C: 3 m-10 y	Children	Acute	Virus or bacterial infection	From hours to days	1. MN: 3m-1y 1.88 g1, -5y 1.88-3.76 g6, -10y 3.76-7.52 g, 2times/d, for 3 d. 2. BE: 10 ml-15 ml, 1-2 times/d, for 3 d. 3. Antibiotic*	с с	Antibiotic*	$\overline{\mathbf{O}}$
L:157	T: 63 C: 57	T: 12.11m C: 12.11m	Children	NN	Virus or bacterial infection	T: 4, 95 $\pm$ 1.68 d C: 6.27 $\pm$ 2.87 d	1. MN:0.3 g, 2 times/d, for 3 d. 2. BE: 0.1 g, 2 times/d, for 3 d. 3. Antibiotic*	ц. Ч	Antibiotic*	(1)
Berberine	+ Bifidobacteriu	m subtilis versus	no berberine +	- Bifidobacteriu	m subtilis					
Geng and Liu <sup>58</sup>	T: 49 C: 49	T: 16y-76y C: 16y-76y	Adult	Acute	N N	3-14 d	1. BS: 0.5g, 3 times/d, for 3 d 2. BE: 4 g+ 100 ml water, 3 times/d, for 3 d*	ю. Ф	MN: 3g, 3 times/d, for 3d*	<b>1</b> 3E
Berberine	+ montmorillon	ite + vitamin B ve	ersus no berbei	rine + montmo	rillonite + v	vitamin B				
Lu <sup>59</sup>	Т: 49 С: 43	T: 6 m-3 y C: 6 m-3 y	Children	Acute	N	и Z	<ol> <li>MN: 3g, 2times/d, for 3d.</li> <li>BE: 10 mg/kg, 2times/d, for 3d.</li> <li>VB: NR, 2times/d, for 3d.</li> <li>Antibiotic*</li> </ol>	Ľ.d	Antibiotic*	$\overline{\mathbf{O}}$
Yiéo	T: 81 C: 81	T: 0-7y C: 0-7y	Children	Acute	ж Х	≤14 d	1. MN: <11.5g, 1-3y 2g, >3y 3g, 2times/d, for 3d. 2. BE: 10 mg/kg, 2times/d, for 3d. 3. VB:NR, 2times/d, for 3d. 4. Antibiotic*	ц.	1. Antibiotic* 2. MN: <1 1.5g, 1-3y 2g, >3y 3g, 1 times/d, for 3 d.	080
										(Continued)

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Table 1. (C	ontinued)									
Study ID	Sample T: (F), C: (F)	Mean/range age (T/C)	Population	Acute/ persistent/ both/NR	Cause of diarrhea	Course of diarrhea (d)	Intervention	Route of administration	Control intervention	Outcome measures
Berberine -	+ montmorillor	ite + Bifidobacter	ium lactobacill	<i>us</i> triple viable	ersus no	berberine + mont	morillonite + <i>Bifidobacterium</i>	lactobacillus tripl	e viable	
Han <sup>61</sup>	T: 82 [36] C: 78 [33]	T:6m-2y C:6m-2y	Children	х Х	х х	ж Z	1. BLT: <1 y 0.21 g; 1-2 y 0.21-0.42 g, 2 times/d, for 3d. 2. BE: 10-15 mg/kg, 2 times/d, for 3 d. 3. MN: <1 y 1.5 g; 1-2 y 3 g, 2 times/d, for 3 d. 4. Antibiotic*	ц ц	Antibiotic*	Θ
Berberine -	+ montmorillor	nite + anisodamine	e <i>versus</i> no bei	rberine + mon	tmorillonite	e + anisodamine				
Yu <sup>62</sup>	T: 78 (35) C: 78 (32)	T: 3 m-20 m C: 2 m-18 m	Children	Both	ж Z	3 d–2 m	<ol> <li>MN:3g, 2times/d, for 3d.</li> <li>BE: 0.12g, times/d, for 3d.</li> <li>AD: 3-5 mg, 2times/d, for 3d.</li> </ol>	ц.	None	$(\overline{-})$
*liquid thera Outcome me absenteeism (12) serum in and diarrhea AD, anisodar ciprofloxacin tetracycline;	py, adjustment o asures: () clinic asures: () clinic () intensity of a flammatory facto flammatory facto flammatory facto flammatory facto hydrochloride; d VB, vitamin B; y,	t diet/diet guidance/r al cure rate (as ment antibiotic use; (7) sto rs: (3) serum myocar e vents; adverse dru e events; advulanat icillin and clavulanat , day; F, female; Fur, year.	nutritional guidar nutritionad by the stuc ol routine examir dial enzymes; ( greaction); ( <b>b</b> ; s furazolidone; LO furazolidone; LO	nce, supplementa dy author); (2) du ation; (8) stool t extorrent diarri tool output, mea tool output, mea t) evofloxacin; N )F, levofloxacin; N	tion of trace of diarration of diarration of diarration of diarration cultures acterial cultures repeated in g or to be sured in g or to be there in e. E berberine, E MN, montmores di NN, montm	elements and calorie rhea (day); ③ stool f irre; ④ duration of hu pisodes of diarrhea i pisodes of diarrhea d BLT, Bitidobacterium ( illonite; M, male; m,	s. requency, measured as the numbe spitalization: ((() duration of other adfined period (as mentioned b quality of tite. <i>actobacillus</i> triple viable; BS, <i>Bifidi</i> month; NR, not reported; NOR, no	er of depositions per - symptom such as vo y the study author]; - obacterium subtilis; C orfloxacin; p.o., orally	day;	urovement; (5) m isoenzymes; :ath: all-cause cefaclor; CPFX, :rine group: Tet,



**Figure 2.** Risk of bias graph: review of author's judgements about each risk of bias item presented as percentages across all included studies.

A sensitivity analysis exploring the effect of selective reporting bias [Appendix 3(a)] did not change the result of the overall meta-analysis significantly.

Comparison 3: berberine + Bifidobacterium subtilis versus *no berberine* + Bifidobacterium subtilis: berberine + montmorillonite + vitamin B versus no berberine + montmorillonite + vitamin B: berberine + Bifidobacterium lactobacillus triple via*ble*+*montmorillonite* versus *no berberine*+ Bifidobacterium lactobacillus triple viable + montmorillonite: berberine + montmorillonite + anisodamine versus no berberine + montmorillonite + anisodamine. Compared with the control groups, using berberine in combinations with probiotics,58 vitamin B59,60 and Bifidobacterium *lactobacillus* triple viable<sup>61</sup> increased the clinical cure rate of diarrhea (Appendices 5–7). There was no statistically significant benefit using berberine combined with montmorillonite and anisodamine<sup>62</sup> (Appendix 8).

# Duration of diarrhea

*Comparison 1: berberine* versus *no berberine; berberine* + *montmorillonite* versus *no berberine* + *montmorillonite.* Overall, five studies evaluated the duration of diarrhea. The use of berberine alone<sup>10,32,33</sup> or berberine combined with montmorillonite<sup>47,57</sup> has an advantage for diarrhea (low-certainty evidence) (Appendices 9 and 10). One

of the studies<sup>10</sup> measured the mean number of the duration of acute diarrhea in adults on day 3 (mean of 37.4h in the berberine group *versus* 44.9h in the control group).

# Secondary outcomes

Stool frequency, stool output and fecal trait. The use of berberine  $alone^{37}$  or berberine combined with montmorillonite<sup>58</sup> may improve the stool frequency, fecal trait, stool output (mean of 2.51 in the berberine group *versus* 5.01 in the control group)<sup>10</sup> (Appendices 11–13).

Stool routine examination, stool bacterial culture and duration of hospitalization. The use of berberine alone<sup>42,44</sup> (very-low-certainty evidence) or berberine combined with montmorillonite and vitamin  $B^{60}$  might have no effect on the improvement of the stool bacterial culture and the stool routine examination (Appendices 14–16).

There was evidence of benefit using berberine or berberine combined with montmorillonite<sup>28,38</sup> (very-low-certainty evidence), vitamin B (mean of 2.5 d in the berberine group *versus* 5.0 d in the control group)<sup>60</sup> (Appendices 17 and 18). There was significant heterogeneity ( $I^2 = 99\%$ ). Due to the insufficient studies, we did not perform a subgroup analysis based on age and the route of administration to explain the statistical differences between the trials.

Study or Subgroup	Experime Events	ntal <u>Total Ev</u>	Contro <u>/ents</u>	ol <u>Total</u>	Weight	Risk Ratio <u>M-H, Fixed, 95% CI</u>		Risk Ratio M-H, Fixed, 95% Cl
1.1.1 Berberine + anti	ibiotics vs	antibiotics	s - chil	dren d	iarrhea oi	n day 3		
Li W2017	17	35	16	35	3.5%	1.06 [0.65, 1.75]		
VVel H2016 Subtotal (95% CI)	9	15	8	15	1.8%	1.13 [0.60, 2.11]		
Total events	26	50	24	50	3.370	1.00 [0.7.5, 1.00]		
Heterogeneity: Chi <sup>2</sup> =	0.02. df = 1	(P = 0.89)	): I <sup>2</sup> = 0	1%				
Test for overall effect:	Z = 0.40 (P	= 0.69)						
1.1.2 Berberine + anti	ibiotics vs	antibiotics	s - chil	dren d	iarrhea oi	n day 7		
Cheng H2015	20	29	10	29	2.2%	2.00 [1.14, 3.49]		
Huang YC2012 Wong 7H2016	34	50	20	50	4.4%	1.70[1.15, 2.51]		
Yang C72017	23	38	16	38	3.5%	1.38 [0.87 2.18]		
Subtotal (95% CI)		162		162	14.5%	1.53 [1.23, 1.91]		•
Total events	101		66					
Heterogeneity: Chi <sup>2</sup> =	2.27, df = 3	(P = 0.52)	); I² = 0	1%				
Test for overall effect:	Z = 3.76 (P	= 0.0002)						
1 1 3 Berherine + anti	ihiotice ve	antibiotics	e . acu	to info	ctious dia	rrhea in children d	nn daw 3	
Xu YF2006	80	84	66 - 100 - 1	84	14.5%	1.21 [1.07. 1.37]	in day 5	
Subtotal (95% CI)		84		84	14.5%	1.21 [1.07, 1.37]		◆
Total events	80		66					
Heterogeneity: Not ap	plicable							
Test for overall effect:	Z = 3.10 (P	= 0.002)						
1.1.4 Berberine + anti	ibiotics VS	antibiotic	s - chi	ldren v	with persi	stent infectious dia	arrhea on dav 3	
Huang H2010	23	40	15	40	3.3%	1.53 [0.95, 2.48]		
Subtotal (95% CI)		40		40	3.3%	1.53 [0.95, 2.48]		
Total events	23		15					
Heterogeneity: Not ap	plicable							
Test for overall effect:	Z=1.74 (P	= 0.08)						
1.1.5 Berberine + anti	ibiotics VS	antibiotic	s - adu	itts wi	th acute a	nd persistent infe	ctious diarrhea on day 3	
Lin RX2016	40	49	20	49	4.4%	2.00 [1.39, 2.87]		
Subtotal (95% CI)		49		49	4.4%	2.00 [1.39, 2.87]		
Total events	40		20					
Heterogeneity: Not ap	plicable							
Test for overall effect:	Z = 3.75 (P	= 0.0002)	•					
1.1.6 Berberine + anti	ibiotics vs	antibiotics	s - adu	lts wit	h acute ai	nd persistent, infe	ctious and non-infectious diarrhea on day 5	
Fu JQ2011	36	53	25	53	5.5%	1.44 [1.03, 2.02]	······································	
Subtotal (95% CI)		53		53	5.5%	1.44 [1.03, 2.02]		
Total events	36		25					
Heterogeneity: Not ap	plicable	- 0.04						
restion overall ellect.	Z = 2.10 (F	- 0.04)						
1.1.7 Berberine VS ar	ntibiotics -	children v	vith inf	ectiou	s diarrhea	a on day 3		
Dang GL2011	35	57	18	45	4.4%	1.54 [1.02, 2.32]		
Subtotal (95% CI)		57		45	4.4%	1.54 [1.02, 2.32]		
Total events	35		18					
Test for overall effect:	plicable 7 = 2.03 (P	= 0.04)						
restion overall ellect.	2 - 2.00 (1	- 0.04)						
1.1.8 Berberine VS ar	ntibiotics -	adults wit	h infec	ctious	diarrhea d	on day 7		
Zhang HF2015	24	30	17	30	3.7%	1.41 [0.98, 2.02]		
Subtotal (95% CI)		30		30	3.7%	1.41 [0.98, 2.02]		
Total events	24 nliaohla		17					
Test for overall effect:	7 = 1.88 (P	= 0.06)						
		0.00)						
1.1.9 Berberine VS ar	ntibiotics-a	dults with	acute	e and p	ersistent,	infectious and no	n-infectious diarrhea on day 5	
Wu J2001	56	80	51	80	11.2%	1.10 [0.88, 1.37]		<b>—</b>
Subtotal (95% CI)	50	80	51	80	11.2%	1.10 [0.88, 1.37]		
Heterogeneity: Not an	oc plicable		51					
Test for overall effect:	Z = 0.84 (P	= 0.40)						
4.4.40 Post								
1.1.10 Berberine + pr	obiotics VS	s probiotio	cs-adu	its per	sistent di	arrhea		
HU YX2009	19	20	14	20	3.1%	1.36 [1.00, 1.84]		
Ye J2013	20	30	16	30	3.5%	1.38 [0.92, 2.22]		
Subtotal (95% CI)		85		83	10.2%	1.40 [1.13, 1.75]		
Total events	66		46					
Heterogeneity: Chi <sup>2</sup> =	0.11, df = 2	(P = 0.95	); I² = 0	1%				
Test for overall effect:	Z = 3.02 (P	= 0.003)						
1.1.11 Berberine + int	testinal mu	cosal pro	tection	n + ant	ibiotics V	S intestinal mucos	al protection + antibiotics-Children with diarrhea	
Li YL2010	53	60	37	60	8.1%	1.43 [1.15, 1.78]		_ <b></b>
Zhou Y2013	89	100	68	98	15.1%	1.28 [1.11, 1.49]		
Subtotal (95% CI)		160		158	23.2%	1.34 [1.18, 1.51]		◆
Total events	142		105					
Heterogeneity: Chi <sup>2</sup> =	0.67, df = 1	(P = 0.41)	);  ² = 0 1\	1%				
rest for overall effect:	∠= 4.58 (P	~ 0.00001	0					
Total (95% CI)		850		834	100.0%	1.37 [1.27, 1.47]		◆
Total events	629		453					
Heterogeneity: Chi <sup>2</sup> =	17.98, df =	17 (P = 0.	39); I² =	= 5%				0.5 0.7 1 1.5 2
Test for subgroup diffect:	∠ = 8.53 (P	≤ 0.000001 hi≇ = 1 2.04	1) 3 df = 1	10 /0 -	017\18-	79.4%	F	avours [control] Favours [experimental]
reactor adoutoup dille	crences. O		2. MI = 1	. o (r =	9.10.1°=	AV.4 /V		





Figure 4. Berberine versus no berberine clinical cure rate funnel plot.

Duration of other symptoms, isoenzyme, inflammatory factors and myocardial enzyme. Using berberine also significantly had better laboratory tests results [isoenzyme-CK<sup>30,31</sup> (low-certainty evidence), CK-MB<sup>27,30,31</sup> (very-low-certainty evifactors-INF- $\alpha^{26,27,30,31}$ dence), inflammatory (very-low-certainty evidence), IL-6<sup>26,27,30,31</sup> (vervlow-certainty evidence), IL-10<sup>26,30,31</sup> (very-lowcertainty evidence) and myocardial enzyme-ALT<sup>26,27,31</sup> (low-certainty evidence), AST<sup>27,30</sup> (low-certainty evidence), LDH30] and fewer systemic symptoms<sup>37</sup> (such as vomiting,<sup>40</sup> heating<sup>40</sup>) than the control groups (Appendices 19-30).

*Adverse events.* The most commonly reported adverse effect was vomiting and rash, but were not serious. No deaths were reported in any of the included studies.

# Discussion

# Summary of our results

We conducted extensive literature searches and identified 38 studies (3948 participants, 27 pediatric trials with 2702 children) for analysis. Compared with antibiotics, berberine plus antibiotics showed better results in both adults and in children in general, especially when given for 7 days or 3 days in acute infectious diarrhea of children. enema in 2306 (2306 of 2702, 85.3%) children. We examined the use of berberine alone or berberine combined with montmorillonite, probiotics, vitamin B or anisodamine for diarrhea. Overall, compared with the control groups, using berberine alone or in combinations with montmorillonite, probiotics, and vitamin B increased the clinical cure rate of diarrhea. The use of berberine alone or berberine combined with montmorillonite may reduce the duration of hospitalization, improved the stool frequency and fecal trait. There was no evidence for the improvement of the culture of the stool bacteria due to use of berberine or berberine combined with montmorillonite and vitamin B. Using berberine also significantly had better laboratory tests results (isoenzyme, inflammatory factors and myocardial enzyme) and fewer systemic symptoms than the control groups. There was no evidence that these interventions had caused death or severe side effects. No study reported industry conflict of interest. However, the quality of evidence of included trials was moderate to low or very low.

Berberine was used orally in all adults and as

# Strengths and limitations

This is the first systematic review to provide evidence for the efficacy and safety of using berberine alone or in combination with Western medicine in diarrhea treatment. Berberine is the major active component of rhizoma coptidis and rhizoma phellodendron, which has been the most well-acknowledged traditional Chinese herb for diarrhea for over 2000 years. We used a modification of the Cochrane Risk of Bias Tool, which uses judgements of 'probably high/low risk of bias' to reduce the occurrence of 'unclear' to assist transparent judgements of the readers.<sup>23</sup> We applied GRADE criteria to determine the certainty in the estimate of effect for our primary outcomes and plausible subgroups.<sup>25</sup> We also evaluated the robustness of the meta-analysis by sensitivity analysis.

There are some limitations to our review. Excessive heterogeneity came to our notice in some of the comparisons; however, we did not have sufficient data and information to identify the source of heterogeneity. We contacted the authors of those studies that were missing critical information but did not receive any useful information from them. Missing essential information such as the course of the disease, absenteeism, the intensity of antibiotic use, quality of life and recurrent diarrhea, may reduce the guiding significance of the clinical practice. Because there is no blind design, most studies have produced a particular bias that affects the quality of evidence. The insufficient number of included studies in some comparisons affected the reliability of the results. We planned subgroup analysis based on dosage and treatment duration, however, the number of trials in each stratum was so limited that subgroup analysis could not be conducted as planned. Thus, the impact of different dosages, frequencies and treatment durations on the outcomes was not evaluated.

# Implications for practice

The current data are promising but inconclusive. According to available evidence, berberine has a good effect on infectious, noninfectious, acute and persistent diarrhea. It can also be used to treat adults and children with diarrhea. There is also some evidence for the efficacy of berberine combined with montmorillonite in the treatment of diarrhea in children. The use of berberine and berberine in combination with montmorillonite appears to be an adjunctive therapy for diarrhea. Although some studies have demonstrated the efficacy of berberine and montmorillonite in combination with anisodamine, vitamin B or probiotics for diarrhea; however, due to the small number of studies and participants, we are not certain of this. Berberine enema treatments for childhood diarrhea were very outstanding. Montmorillonite slows down the peristaltic speed of the intestinal wall, improves the microcirculation of intestinal, and enhances the immunity of body.<sup>63</sup> Vitamin B promotes metabolism, repairs the gastrointestinal mucosa and improves the function of the digestive system. Probiotics adjust the balance of intestinal flora, inhibit the production of endotoxins by harmful bacteria and maintain the normal physiological function of human intestines.37 Berberine combined with montmorillonite and vitamin B or probiotics can improve dyspepsia, reduce vomiting and restore the normal physiological function of the intestinal mucosal barrier.64

Because of the low bioavailability of berberine,<sup>38</sup> it needs to be given three times a day for adults.<sup>65</sup> Although it adds extra burden to the patients, it only has a high drug concentration in the intestine, and its blood drug concentration is almost negligible.<sup>38</sup> Thus, berberine can reduce the risk of systemic side effects.

Acute diarrhea is often due to viruses or bacteria which are resistant to some antibiotics.<sup>66</sup> Overall, 15 studies (Table 1) included in our review were infectious diarrhea; however, no pathogen was reported. We are unable to tell whether there were misuse of antibiotics. We suggest, in clinical practice, antibiotics should be prescribed according to local regulations to avoid as much as possible misuse and overuse (see Table 2 for summary of primary outcomes of RCTs on berberine for diarrhea).

Taking account of the over 30 years' usage of berberine for diarrhea as an over-the-counter antimicrobial in a large Chinese population, the very low marketing price, the effectiveness we found, as well as the low incidence of adverse events, which was also addressed in a pharmacology review,<sup>10</sup> berberine should be an attractive intervention for diarrhea.

The dosage of oral berberine for adults or children based on literature is consistent with the user's manual, and for adults, is 0.1-0.3g per time, three times a day,<sup>67</sup> and for children, is based on the weight and age of the children (Table 3).<sup>67</sup> Although we found that berberine

Certaint	V assessm	ent					No. of partic	cipants	(studies)		Effect	Certaintv
No. of studies	Study design	Risk of bias	Incon- sistency	Indire- ctness	Impre- cision	Other conside- rations	Berberine	No berberine	Relative (95% CI)	Absolute (95% CI)		
berberir	ne <i>versus</i> n	io berberine	e: clinical cı	ure rate								
30	rand- omized trials	serious <sup>a</sup>	not seri- ous	not seri- ous	not serious	not sus- pected	629/850 [74.0%]	453/834 [54.3%]	<b>RR 1.37</b> [1.27– 1.47]	<b>201 more per</b> <b>1000</b> (from 147 more to 255 more)	⊕⊕⊕⊖ MODERATE	CRITICAL
Berberii	re + mont	tmorillonite	versus no t	berberine +	montmor	illonite: clinica	ıl cure rate					
13	rand- omized trials	serious <sup>a</sup>	serious <sup>b</sup>	not seri- ous	not serious	publication bias strongly suspected <sup>d</sup>	429/612 [70.1%]	214/534 (40.1%)	<b>RR 1.70</b> (1.43– 2.02)	<b>281 more per</b> <b>1000</b> (from 172 more to 409 more)	⊕○○○ VERY LOW	CRITICAL
Berberii	he + montr	morillonite	+ vitamin B	<i>versus</i> no b	erberine +	- montmorillon	ite + vitamin [	B: clinical cui	re rate			
2	rand- omized trials	serious <sup>a</sup>	not seri- ous	not seri- ous	seri- ous <sup>c</sup>	not sus- pected	109/130 (83.8%)	70/124 [56.5%]	<b>RR 1.49</b> [1.26- 1.77]	<b>277 more per</b> <b>1000</b> (from 147 more to 435 more)	⊖ ⊖⊕⊕ ⊕	CRITICAL
Berberii	ne <i>versus</i> r	no berberin	e: duration	of diarrhea								
5	rand- omized trials	serious <sup>a</sup>	not seri- ous	not seri- ous	seri- ous <sup>c</sup>	not sus- pected	65	63	I	MD <b>3.25</b> <b>lower</b> (3.64 lower to 2.85 lower]	⊕⊕_ Low	CRITICAL
Berberii	he + montr	morillonite	<i>versus</i> no b	erberine + r	nontmorill	lonite: duration	ו of diarrhea					
5	rand- omized trials	serious <sup>a</sup>	not seri- ous	not seri- ous	seri- ous <sup>c</sup>	not sus- pected	123	113	1	MD <b>1.63</b> <b>lower</b> [2.14 lower to 1.13 lower]	⊕⊕⊖ Low	CRITICAL
CI, confia <sup>a</sup> All the tr is difficul <sup>b</sup> There is <sup>c</sup> For diche	lence intervarials had a h t to blind pe significant s otomous out	al; MD, mean nigh risk of pe rrsonnel and f statistical heth tcomes, the th	difference (d rformance bi participants. erogeneity in otal number o	ays); RR, risk as not blindin dicating by a l of events is les	ratio. g the partici arge /² valu€ ss than 300;	ipants. Methodolc e. for continuous ou	ogical quality of utcomes, the tot	these trials wa tal population s	is graded as 'f size is less tha	nigh risk of bias' du un 400; or pooled re	ue to the design o	ıf comparison ə effects.

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Table 3.	The	instructions	of oral	berberine	for	children <sup>6</sup>	7
Table J.	IIIC	mon uctions	UI UI at	Der Der Infe	101	cinturen.	

Age	Weight	Dosage
1–3years	10–15 kg	0.05–0.1 g
4-6years	16–21 kg	0.1–0.15 g
7-9years	22–27 kg	0.15–0.2 g
10–12 years	28–32 kg	0.2–0.25 g

can be used by enema from the included literature, no instruction could be found in the user's manual. However no obvious adverse event was reported, either. Merits and risks of berberine enema need to be further explored.

#### Implications for research

A large sample and multicenter well-designed clinical studies on berberine in specific population (children/adults), disease condition (infectious/noninfectious; acute/chronic) and treatment protocol (alone or in combination, orally taken or enema) are needed since berberine has shown therapeutic evidence or potentials in all above circumstances with seemingly low safety risk, and preclinical research evidence support. Economic analysis should be provided to guide practices in different countries or regions.

#### Conclusion

In conclusion, the current data are promising but inconclusive. Berberine alone or combined with montmorillonite may have benefit for diarrhea. There are few studies evaluating necessary outcomes in berberine for diarrhea, especially for absenteeism, the intensity of antibiotic use, quality of life and recurrent diarrhea. Due to the risk of bias and concern of publication bias, the above evidence was moderate to low or very low. A large sample and multicenter well-designed clinical studies are needed.

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#### Author contributions

YT Fei and MK Yu conceived and designed the review. MK Yu and FL Bu drafted the protocol.

MK Yu and CH Liang were responsible for the searching, screening and selecting studies. CH Liang, Y Ming, D Pan and Q He participated in data extraction and assessed study quality. MK Yu and CZ Li made forms and pictures and performed the statistical analysis. MK Yu and XI Jin drafted the manuscript. XJ Jin revised the article in its entirety. JP Liu, HB Du, SB Liang, RX Hu, Paul Little and Yanhong Jessika Hu were all involved in critically revising the manuscript. XI Jin, Yanhong Jessika Hu and Paul Little were responsible for the refinement of the content and language of our article. HJ Cao re-checked and analyzed the article data. All authors have read and approved the final manuscript. All authors approved the final version of the article, including the authorship list.

## **Conflict of interest statement**

The authors declare that there is no conflict of interest.

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## **PRISMA 2009 checklist statement**

The authors have read the PRISMA 2009 Checklist, and the manuscript was prepared and revised according to the PRISMA 2009 Checklist.

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#### Supplemental material

Supplemental material for this article is available online.

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# Search strategy

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