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American Gastroenterological Association-American College of Gastroenterology Clinical Practice Guideline: Pharmacological Management of Chronic Idiopathic Constipation

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INTRODUCTION: Chronic idiopathic constipation (CIC) is a common disorder associated with significant impairment in

quality of life. This clinical practice guideline, jointly developed by the American Gastroenterological Association and the American College of Gastroenterology, aims to inform clinicians and patients by providing evidence-based practice recommendations for the pharmacological treatment of CIC in adults.

METHODS: The American Gastroenterological Association and the American College of Gastroenterology formed a

multidisciplinary guideline panel that conducted systematic reviews of the following agents: fiber, osmotic laxatives (polyethylene glycol, magnesium oxide, lactulose), stimulant laxatives (bisacodyl, sodium picosulfate, senna), secretagogues (lubiprostone, linaclotide, plecanatide), and serotonin type 4 agonist (prucalopride). The panel prioritized clinical questions and outcomes and used the Grading of Recommendations Assessment, Development, and Evaluation framework to assess the certainty of evidence for each intervention. The Evidence to Decision framework was used to develop clinical recommendations based on the balance between

the desirable and undesirable effects, patient values, costs, and health equity considerations.

RESULTS: The panel agreed on 10 recommendations for the pharmacological management of CIC in adults. Based

on available evidence, the panel made strong recommendations for the use of polyethylene glycol, sodium picosulfate, linaclotide, plecanatide, and prucalopride for CIC in adults. Conditional

recommendations were made for the use of fiber, lactulose, senna, magnesium oxide, and lubiprostone.

DISCUSSION: This document provides a comprehensive outline of the various over-the-counter and prescription

pharmacological agents available for the treatment of CIC. The guidelines are meant to provide a framework for approaching the management of CIC; clinical providers should engage in shared decision making based on patient preferences as well as medication cost and availability. Limitations and gaps in the evidence are highlighted to help guide future research opportunities and enhance the care of patients with chronic

constipation.

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KEYWORDS: fiber; polyethylene glycol; magnesium oxide; lactulose; docusate; bisacodyl; senna; sodium picosulfate; lubiprostone; linaclotide; plecanatide; prucalopride

SUPPLEMENTARY MATERIAL accompanies this paper at http://links.lww.com/AJG/C894

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EXECUTIVE SUMMARY

Chronic idiopathic constipation (CIC) is a common clinical diagnosis that affects approximately 8%–12% of the US population (1). Nonpharmacological therapies often represent the initial steps in management and may include dietary recommendations (such as increased fluid intake and increased dietary fiber) and behavioral changes (such as exercise). Pharmacological treatment may include the use of over-the-counter (OTC) or prescription medications, such as polyethylene glycol (PEG), secretagogues, or prokinetic agents (2). This joint evidence-based guideline from the American Gastroenterological Association (AGA) and the American College of Gastroenterology (ACG) aims to provide recommendations for the pharmacological management of CIC in adults.

HOW TO READ THESE GUIDELINES

Table 1 provides an overview of each guideline recommendation along with the associated certainty of evidence and the strength of recommendation. Additional information about the background, methods, evidence reviews, and detailed justifications for each recommendation is provided after Table 1 for readers wishing to read the full guideline. Corresponding forest plots for each intervention and evidence profiles which provide a synthesis of the evidence as well as Evidence to Decision (EtD) framework tables that summarize the panel's detailed judgments supporting each recommendation are provided in the Supplementary Document (see Supplementary Digital Content 1, http://links.lww.com/ AJG/C894). Each recommendation is accompanied by clinical practice considerations (based on the collective experience of the panel members) that are meant to help guideline users implement the recommendations. The term "recommend" was used to indicate strong recommendations, and the term "suggest" was used to indicate conditional recommendations. The interpretation of certainty of evidence and implications of strong and conditional recommendations for healthcare providers, patients, and policymakers are presented in Tables 2 and 3, respectively. For all the recommendations, the alternative approach was management of CIC without the intervention.

INTRODUCTION

Description of the health problem

CIC is a common clinical diagnosis that affects approximately 8%–12% of the US population (1). CIC is a lower gastrointestinal (GI) tract disorder of gut-brain interaction and can be associated with symptoms such as infrequent and incomplete defecation in the absence of mucosal or structural abnormalities (2,3). The medical costs related to the management of constipation are estimated to be between approximately \$2,000 and \$7,500 US dollars per patient per year, and the effects on quality of life can be similar to those associated with conditions such as chronic obstructive pulmonary disease, diabetes, and depression (4,5). Nonpharmacological interventions often represent the initial step

in management and may include dietary (such as increased fluid intake and increased dietary fiber) and behavioral changes (such as exercise). Pharmacological treatment may include the use of PEG, secretagogues, and prokinetic agents (2,6,7). Overall, a significant proportion of patients with CIC are not satisfied with their treatment and may use multiple OTC medications, followed by prescription medications before they have improvement in their symptoms (8–10).

Objective of the review and guideline

The AGA and ACG jointly developed this systematic review and clinical guideline to provide evidence-based recommendations for the pharmacological management of CIC in adults.

Target audience

The target audience for these guidelines includes primary care, internal medicine, family medicine, and gastroenterology healthcare providers; patients; and policymakers. The recommendations in this document are not intended to be used as the standard of care. Instead, they can be used to guide the management of adult patients with CIC. Although no single recommendation can encompass every individual circumstance and context, it can be used to address the benefits and harms of treatments and support the processes of shared decision making so that patients are treated based on their values and preferences.

METHODS

Overview

This document represents the official recommendations of the AGA and ACG. These recommendations were developed using the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) approach.

Organization and panel composition

The guideline panel members from the AGA and ACG were selected based on their clinical and methodological expertise. Each member underwent a vetting process that required disclosing all conflicts of interest. The panel included 3 guideline committee members specializing in general gastroenterology, motility, and primary care. Panel members comprising the evidence review team (divided into 3 subcommittees) included gastroenterologists with expertise in CIC, 1 senior methodologist, and 3 junior methodologists. All included interventions were divided among the 3 subcommittees (see Supplementary Table 1, Supplementary Digital Content 1, http://links.lww.com/AJG/ C894). The senior methodologist supervised the evidence synthesis for all the interventions across the 3 subcommittees. Members of the guideline committee helped review all the synthesized evidence, contributed to discussion, and helped develop the clinical decision support tool. A librarian assisted with designing and executing the relevant literature searches. An executive committee of members of the AGA and ACG were

Table 1. Summary of recommendations and implementation considerations

Strength of recommendation	Certainty of evidence
Conditional	Low
Strong	Moderate
Conditional	Very low
	Conditional

Table 1. (co	ontinued)
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Table 1. (continued)		
Recommendations	Strength of recommendation	Certainty of evidence
Recommendation 4: In adults with CIC who fail or are intolerant to OTC therapies, the panel suggests the use of lactulose over management without lactulose Implementation considerations • Bloating and flatulence are dosedependent and common side effects, which may limit its use in clinical practice	Conditional	Very low
Stimulant laxatives		
Recommendation 5: In adults with CIC, the panel recommends the use of bisacodyl or sodium picosulphate short term or as rescue therapy over management without bisacodyl or sodium picosulphate Implementation considerations • Short-term use is defined as daily use for 4 wk or less. While long-term use is probably appropriate, data are needed to better understand tolerance and side effects • This is a good option for occasional use or rescue therapy in combination with other pharmacological agents for CIC • The most common side effects are abdominal pain, cramping and diarrhea. The panel suggests starting at a lower dose and increasing the dose as tolerated	Strong	Moderate
Recommendation 6: In adults with CIC, the panel suggests the use of senna over management without senna Implementation considerations • While the trials were conducted for 4 wk, longer term use is probably appropriate, but data are needed to better understand tolerance and side effects • The dose evaluated in trials is higher than commonly used doses in practice. The panel suggests starting at lower dose and increase if no response • Abdominal pain and cramping may occur with a higher dose of senna	Conditional	Low
0		

Secretagogues

Table 1. (continued)		
Recommendations	Strength of recommendation	Certainty o evidence
Recommendation 7: In adults with CIC who do not respond to OTC agents, the panel suggests the use of lubiprostone over management without lubiprostone <i>Implementation considerations</i> • Can be used as a replacement or as an adjunct to OTC agents • Duration of treatment in trials was 4 wk, but the drug label does not provide a limit • Nausea may occur; however, the risk of nausea is dose-dependent and seems to be lower when taken with food and water	Conditional	Low
Recommendation 8: In adults with CIC who do not respond to OTC agents, the panel recommends the use of linaclotide over management without linaclotide Implementation considerations • Can be used as a replacement or as an adjunct to OTC agents • Duration of treatment in trials was 12 wk but the drug label does not provide a limit • May be associated with side effects of diarrhea leading to discontinuation of treatment	Strong	Moderate
Recommendation 9: In adults with CIC who do not respond to OTC agents, the panel recommends the use of plecanatide over management without plecanatide Implementation considerations • Can be used as a replacement or as an adjunct to OTC agents • Duration of treatment in trials was 12 wk, but the drug label does not provide a limit • May be associated with side effects of diarrhea leading to discontinuation of treatment	Strong	Moderate
5-HT ₄ agonist		
Recommendation 10: In adults with CIC who do not respond to OTC agents, the panel recommends the use of prucalopride over management without prucalopride <i>Implementation considerations</i> • Duration of treatment in trials was 4–24 wk but the drug label does not provide a limit • Can be used as a replacement or as an adjunct to OTC agents • May be associated with side effects of	Strong	Moderate

The implementation considerations are based on the collective experience of the panel members, and evidence may not be available for each of the implementation considerations.

5-HT₄, serotonin type 4; CIC, chronic idiopathic constipation; OTC, over-the-counter; PEG, polyethylene glycol.

responsible for oversight of this collaborative guideline (S.S., W.D.C., L.C., A.L.).

Management of conflict of interest and guideline funding

Panel members disclosed all potential conflicts of interest. Conflicts were managed according to AGA and ACG policies, the National Academy of Medicine, and Guidelines International Network standards (11–13). Panel members determined to have a potential conflict of interest with a specific intervention or agent were allocated to a subcommittee that did not include the specific intervention(s). Development of this guideline was wholly funded by the AGA and ACG with no support from the industry.

Scope

The guideline panel and evidence review team formulated clinically relevant questions on the pharmacological therapies for CIC in adults. The last position paper by the AGA on CIC included guidance on clinical evaluation, diagnostics tests, and medical and surgical management (6). This document does not specifically address considerations related to special populations such as those with malignancy, pregnancy, or opioid-induced constipation.

Formulation of clinical questions and determining outcomes of interest

PICO format. For each guideline question, the evidence review team conducted a systematic review. The systematic review was based on specific Population, Intervention, Comparison, Outcome (PICO) questions developed by the guidelines committee and approved by the Boards of both organizations. A protocol guided the systematic review process and is registered at the international Prospective Register of Systematic Reviews website (CRD42021254673). In summary, we included individual randomized controlled trials (RCTs). We also considered the trials with multiple arms and included comparisons where the only difference between the 2 groups was the intervention of interest. If a study included multiple treatment arms but only 1 comparison group, we combined the treatment arms when appropriate so that the comparison group was not counted twice in the meta-analysis.

Population. The population of interest was adults (18 years or older) diagnosed with CIC. We excluded studies where individuals were diagnosed with other similar conditions such as opioid-induced constipation or constipation due to other medical conditions such as hypothyroidism and celiac disease. We also excluded studies in patients with irritable bowel syndrome with constipation (IBS-C) because the pharmacological management of IBS-C was covered in another recent AGA (14) and ACG guideline (15).

Intervention. We included the following interventions: fiber: psyllium, bran, methylcellulose, and inulin; osmotic or surfactant laxatives: PEG, magnesium oxide (MgO), lactulose, and docusate; stimulant laxatives: bisacodyl, senna, and sodium picosulfate; secretagogues: lubiprostone, linaclotide, and plecanatide; and serotonin type 4 (5-HT₄) agonist: prucalopride. We considered studies that included the above interventions, regardless of the dose or route of administration. We included studies in which the intervention duration was at least 4 weeks. We analyzed all the interventions separately.

Comparison. The comparison group included placebo, no intervention, or standard of care. We excluded studies that

headache, abdominal pain, nausea,

and diarrhea

Table 2. Interpretation of the certainty of effects using the GRADE framework

Certainty of evidence	Definition
High	We are very confident that the true effect lies close to that of the estimate of the effect.
Moderate	We are moderately confident that the true effect lies close to that of the estimate of the effect. There is a possibility that it is substantially different.
Low	Our confidence that the true effect lies close to that of the estimate of the effect is low. The true effect may be substantially different from the estimate of the effect.
Very low	Our confidence that the true effect lies close to that of the estimate of the effect is very low. The true effect is likely substantially different from the estimate of the effect.

GRADE, Grading of Recommendations Assessment, Development, and Evaluation.

compared different doses or frequencies of the same drug and did not include a comparison group that did not receive the drug. We also excluded studies that compared different pharmacological agents for CIC, and there was no placebo group.

Outcomes. We considered the following outcomes: complete spontaneous bowel moments (CSBMs) per week; spontaneous bowel movements (SBMs) per week; responder rate, defined as CSBM per week of equal or greater than 3 and increased by at least 1 from baseline; diarrhea (adverse event) leading to discontinuation of treatment; serious adverse events; global relief outcome; quality-of-life scores (using the Patient Assessment of Constipation-Quality of Life, or PAC-QOL); and stool form. We considered the outcomes of CSBMs per week, SBMs per week, and adverse events leading to discontinuation of medication as the critical outcomes.

Search strategy

A comprehensive search was conducted on the following data-bases: EMBASE, MEDLINE, Cochrane, Scopus, Web of Science, ClinicalTrials.gov, Centre for Reviews and Dissemination, and PubMed. We also reviewed the reference sections of available systematic reviews (7,8,16,17) and updated the searches if a recent systematic review was available. The literature was first searched on May 15, 2021, and the search was updated on November 5, 2022. The search terms used can be found in Appendix 1 (see Supplementary Digital Content 1, http://links.lww.com/AJG/C894).

Study selection, data collection, and analysis

Searches from all the databases were combined in bibliographic software (EndNote) (18), and duplicates were removed. Two reviewers screened the titles and conducted a full-text review of the eligible studies (using a reference software Covidence), and a consensus was reached on inclusion (19). Any conflicts were resolved with the help of a senior member of the team. Data were extracted from each study, including study characteristics, such as year of publication, study site, study population, dose and frequency of intervention, comparison group, outcomes and methods for risk-of-bias assessment. Meta-analysis was conducted when more than 1 study contributed data for the same intervention and outcome. We combined the dichotomous

outcomes to obtain a relative risk (RR) and 95% confidence interval (CI). We used the mean difference (MD) to pool the continuous outcomes (20). For the meta-analysis, we used the generic inverse variance method of weighting and applied the random-effects model. We assessed the statistical heterogeneity by using the I^2 index and χ^2 statistic. We used funnel plots to assess the small study effect and publication bias when at least 10 studies were available in a pooled analysis. We used RevMan (21) software for all the statistical analyses. We used the Cochrane Risk of Bias tool to assess the risk of bias in the included studies (22). This tool assesses the risk of bias in the following domains: sequence generation, allocation concealment, blinding, incomplete outcome data, selective reporting, and other biases (22).

Certainty of the evidence

We used the GRADE approach to assess the certainty of evidence for the effect of the intervention on each outcome using the software GradePro (23). The GRADE approach considers factors such as study design, risk of bias, inconsistency, indirectness, imprecision, and risk of publication bias to rate the certainty of evidence as high, moderate, low, or very low (24) (Table 2). The results of certainty assessment are reported in evidence profiles available in the Supplementary Document (see Supplementary Digital Content 1, http://links.lww.com/AJG/C894) for all the interventions included in this review.

Development of recommendations

The process of translation of evidence into guideline recommendations followed the GRADE EtD framework (25) and was achieved by discussion during virtual meetings of the guideline committee. The EtD framework considers the certainty of evidence, balance of benefits and harm, patient values and preferences, feasibility, acceptability, equity, and resource use (25). All 10 EtD tables are presented in the Supplementary Document (see Supplementary Digital Content 1, http://links.lww.com/AJG/C894). Consensus was reached for all the recommendations among the group. The interpretation of strength of recommendations is summarized in Table 3.

Document review

The guideline underwent expedited internal and external peer review. The guideline document was revised to address pertinent comments, but no changes were made to the direction or strength of recommendations.

Recommendations

The literature search yielded 993 titles, and a total of 726 titles and abstracts were screened after the duplicates were removed. Of 54 studies reviewed with full text, 28 studies were included in evidence synthesis and 14 studies were excluded (Figure 1). Supplementary Table 2 (see Supplementary Digital Content 1, http://links.lww.com/AJG/C894) gives the reason for exclusion of studies. The summary estimates for effect of a specified intervention on each of the prespecified outcome are included as forest plots and evidence profiles for each PICO question and are available in the Supplementary Document (see Supplementary Digital Content 1, http://links.lww.com/AJG/C894).

Table 1 provides a summary of the recommendations presented in this guideline. More detailed information regarding the medication indication, dosing, availability, and mechanism of action is summarized in Table 4.

Table 3. Interpretation of a strong and conditional recommendation

	Implications	Strong recommendation	Conditional recommendation
	For patients	Most individuals in this situation would want the recommended course of action and only a small proportion would not.	The majority of individuals in this situation would want the suggested course of action, but many would not.
	For clinicians	Most individuals should receive the intervention. Formal decision aids are not likely to be needed to help individuals make decisions consistent with their values and preferences.	Different choices will be appropriate for individual patients consistent with his or her values and preferences. Use shared decision making. Decision aids may be useful in helping patients make decisions consistent with their individual risks, values, and preferences.
	For policy makers	The recommendation can be adapted as policy or performance measure in most situations.	Policy making will require substantial debate and involvement of various stakeholders. Performance measures should assess whether decision making is appropriate.

FIBER

Recommendation 1: In adults with CIC, the panel suggests the use of fiber supplementation over management without fiber supplements (conditional recommendation, low certainty of evidence).

Implementation considerations

- Dietary assessment is important to determine total fiber intake from diet and supplements.
- Fiber supplements can be used as first-line therapy for CIC, particularly for individuals with low dietary fiber intake.
- Among the evaluated fiber supplements, only psyllium appears to be effective (with very limited and uncertain data on bran and inulin).
- Adequate hydration should be encouraged with the use of fiber.
- Flatulence is a commonly observed side effect with the use of fiber.

We evaluated evidence for fiber supplementation in the form of bran, inulin, psyllium, and methylcellulose. The overall certainty of evidence for the use of fiber in the management of CIC compared with management without fiber was low. No studies were found on the use of methylcellulose, but data on bran, inulin, and psyllium are outlined below.

Bran

Summary of evidence. One randomized study conducted in Italy with a cross-over design evaluated the efficacy and safety of bran in 29 patients with chronic nonorganic constipation (26). Participants received either a ground bran (6.6 g) or an identical-looking preparation of placebo 3 times a day for two 4-week periods (26).

Benefits and harms. Based on this 1 small study, bran may lead to an increase in SBMs per week; however, the CI was wide and included a possible null effect (MD 1.30, CI -0.98 to 3.58) (26). Only 1 adverse event was noted in the treatment group compared with no events in the placebo arm (RR 2.79, CI 0.12-62.48) (26). There were no data on CSBMs per week, responder rates, diarrhea, global relief, quality of life, or stool form.

Certainty in evidence of effects. We are very uncertain about the effects of bran. The overall certainty of evidence for bran is very low because of concerns about the adequacy of randomization and allocation concealment as well as very serious imprecision (26).

Inulin

Summary of evidence. Two studies assessed the effect of inulin for the treatment of CIC (27,28). The first study was a randomized placebo-controlled study conducted in Brazil involving 60 female participants aged 18–65 years with at least 3 months of constipation and <3 bowel movements per day (27). Twenty-eight patients were in the fiber group and 32 in the placebo group (27). Participants were given 4 days to adapt to the mixture of inulin and partially hydrolyzed guar gum or placebo before beginning a 3-week treatment with either 15g/d of inulin or 15 g/d of maltodextrin (placebo) that was divided into 3 sachets of 5 g each (27). The second study was conducted in Belgium on participants aged 50–70 years who were randomized to 7.5 g of inulin sachet and placebo for 28 days (28). The inulin and placebo groups included 25 participants each. The duration of follow-up was 28 days.

Benefits and harms. All included studies did not contribute data for our outcomes of interest. Based on one small study, treatment with inulin had little to no effect on SBMs per week (MD -0.75, CI -2.60 to 1.10) and responder rate, defined as >3 CSBMs per week (RR 1.21, CI 0.83-1.74) (27). Regarding side effects, no serious adverse events were reported, although a minor side effect of flatulence occurred more frequently in the inulin group, according to one study (28). There were no data on CSBMs per week, diarrhea, serious adverse effects, quality of life, or stool form.

Certainty in evidence of effects. We are very uncertain about the effects of inulin. The certainty of evidence for the effect of inulin on SBMs and the responder rate was very low and low, respectively, because of concerns about risk of bias and imprecision because of the small number of included participants in the study.

Psyllium

Summary of evidence. Three RCTs conducted between 1986 and 1995 studied psyllium vs placebo (29-31). Psyllium was evaluated in a small study from the United States that included 22 adult participants with CIC confirmed by prospectively administered stool diaries (29). Constipation was defined as <3 bowel movements per week. A 4-week baseline phase was followed by 8 weeks of double-blinded treatment, during which patients received either 5 g twice daily of psyllium or placebo, followed by a 4-week washout period (29). The second study included 201 participants with functional constipation between 18 and 70 years (30). Participants were randomly allocated to either Regulan (a refined hydrophilic mucilloid derived from psyllium seed husks) or matching placebo for 14 days (30). Participants received one sachet (containing 3.6 g of active ingredient) of either psyllium or placebo 3 times daily (30). A third study included 35 participants with constipation who were randomized to receive either

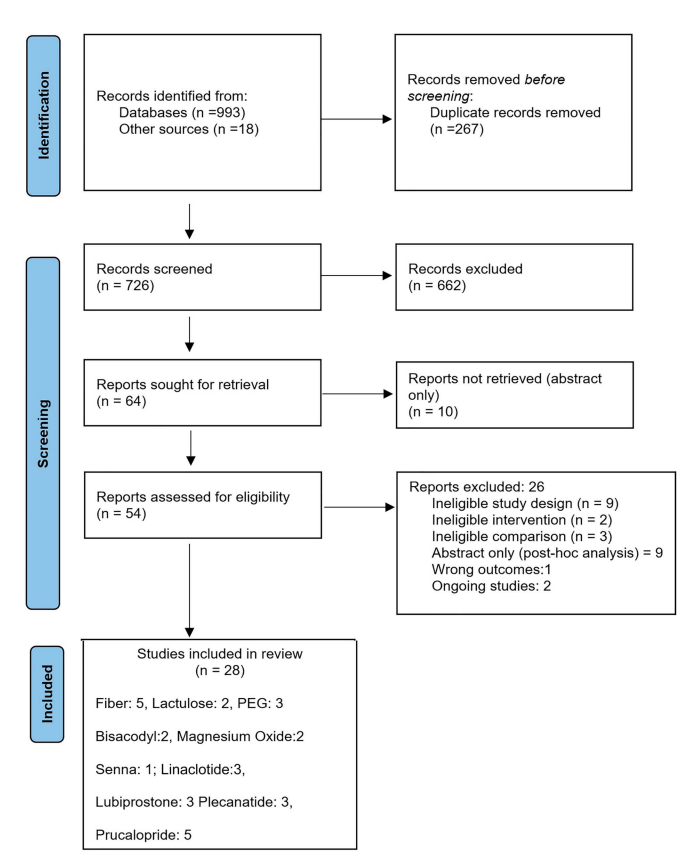


Figure 1. PRISMA flow diagram. Six studies on fiber supplements included studies on the use of insulin (1 study), bran (1 study), and psyllium (3 studies). One study addressed magnesium oxide and senna in the same trial, so the total number of included studies is 28 and not 29. PEG, polyethylene glycol.

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What medications can be used to treat chronic idiopathic constipation?	Mechanism of action	Recommended initial dose	Guidance for dose titration	Maximum dose	Estimated monthly cost, USD ^a	Additional comments
Fiber	Soluble fiber traps water in the intestine and softens stool Insoluble fiber increases stool bulk	The Academy of Nutrition and Dietetics ^a recommends 14 g/1,000 kcal intake per day Total daily fiber intake (dietary + supplement) 20–30 g/d	Per response to symptoms and side effects Common side effects include bloating and abdominal discomfort	Usually no benefit to increasing total fiber intake over 25–30 g	<\$50	Ensure adequate hydration as fiber intake increases No clear evidence that soluble or insoluble fiber is more effective Soluble fiber includes psyllium, inulin, oats, fruit, barley, and legumes Insoluble fiber includes wheat bran, methylcellulose, wheat, rye, and other grains
Polyethylene glycol	Osmotic laxative	17 g daily	Per symptom response and side effects Common side effects include bloating, abdominal discomfort, and cramping	No clear maximum dose	\$10-\$45	Response to PEG has been shown to be durable over 6 mo
Magnesium oxide	Osmotic laxative	400–500 mg daily	Per symptom and response and side effects	No clear maximum dose. Prior studies used 1,000–1,500 mg daily	<\$50	Use with caution in patients with renal insufficiency and in pregnancy
Lactulose	Osmotic laxative	15 g daily	Per symptom response and side effects Bloating and flatulence may be limiting if preexisting symptoms or at higher doses	No clear maximum dose. May cause hypernatremia and hypokalemia if patients experience significant diarrhea	<\$50	Only osmotic agent studied in pregnancy
Bisadocyl and picosulfate	Stimulant laxative	Bisacodyl 5 mg daily	Per symptom response and side effects Side effects limited by cramping and abdominal discomfort	10 mg orally daily	<\$50	Recommended for short-term use or rescue therapy Prolonged or excessive use can cause diarrhea and electrolyte imbalance Long-term safety and efficacy unknown
Senna	Stimulant laxative	8.6–17.2 mg daily	Per symptom response and side effects Side effects most commonly cramping and abdominal discomfort	No clear maximum dose. Often recommended maximum is 4 tablets twice per day	<\$50	Also present in many laxative teas, where dose may be difficult to calculate Long-term safety and efficacy unknown

Table 4. Overview of interventions for the pharmacological management of chronic idiopathic constipation

Table 4. (continued)

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What medications can be used to treat chronic idiopathic constipation?	Mechanism of action	Recommended initial dose	Guidance for dose titration	Maximum dose	Estimated monthly cost, USD ^a	Additional comments
Lubiprostone	Intestinal secretagogue acting on chloride channel type 2 in the gut that increases chloride secretion	24 μg BID	Per symptom response Diarrhea may occur in a subset of patients, leading to discontinuation	24 μg BID	\$374	May have benefit for abdominal pain. Also approved for the treatment of IBS-C at a dose of 8 µg BID
Linaclotide	Intestinal secretagogue acting on guanylate cyclase-C, which activates CFTR in the gut to increase chloride secretion	72–145 μg daily	Per symptom response Diarrhea may occur in a subset of patients, leading to discontinuation	290 μg daily	\$523	May have benefit for abdominal pain. Also approved for the treatment of IBS-C
Plecanatide	Intestinal secretagogue acting on guanylate cyclase-C, which activates CFTR in the gut to increase chloride secretion	3 mg daily	Per symptom response Diarrhea may occur in a subset of patients, leading to discontinuation	3 mg daily	\$526	Also approved for the treatment of IBS-C
Prucalopride	5-HT ₄ agonist	1–2 mg daily	Per symptom response Headaches and diarrhea may occur in a subset of patients, leading to discontinuation	2 mg daily	\$563	May have additional benefit for abdominal pain

⁵⁻HT₄, serotonin type 4; BID, twice a day; IBS-C, irritable bowel syndrome with constipation; PEG, polyethylene glycol; USD, US dollar.

^aThe given cost accommodates the extent of generic and prescription medications and may not be the exact cost. In addition, the given cost is not the cost-effectiveness of the medication, but a probable cost per month.

celandine-aloe vera-psyllium or placebo (31). Capsules of 500 mg contained the active ingredients celandine, aloe vera, and psyllium in the ratio of 6:3:1. The initial dose was 1 capsule per day, taken with water at bedtime, and increased to 3 capsules per day depending on the response (31).

Benefits and harms. Based on the meta-analysis of data from 3 studies (29–31), the use of psyllium may lead to an increase in SBMs per week (MD 2.32, CI 0.86–3.79). Combined data from 2 studies showed that the use of psyllium may increase global relief symptoms (RR 1.86, CI 1.49–2.30), but there was little to no difference in stool consistency (MD -1.08, CI -1.33 to 0.83) (30,31). In absolute terms, psyllium was associated with 391 per 1,000 more individuals with global relief (from 223 more to 591 more). One study (30) examined withdrawal from the study because of diarrhea but with only 3 events; the summary estimate was too imprecise to make any conclusive statement (RR 0.47, CI 0.04–5.06); and no serious adverse events were reported in either arm. There were no data on CSBMs per week, responder rate, diarrhea, quality of life, or stool form.

Certainty in evidence of effects. The certainty of evidence for SBM, serious adverse events, and global relief was low and for diarrhea, was very low. We rated down certainty of evidence because of high risk of bias (concerns about methods of randomization and allocation concealment in 2 studies and high attrition in another study), indirectness, and imprecision.

Discussion

Fiber can be divided into soluble and insoluble fiber. Wheat bran, an insoluble fiber, is produced when the hard outer fiber of the wheat kernel is removed during the refining process. Inulin is a naturally occurring polysaccharide present in many plants and most often extracted from chicory. Inulin is a fructan and is considered both a soluble fiber and a prebiotic, meaning that it can stimulate the growth or activity of intestinal bacteria (32) that are believed to promote good gut health. Psyllium is also considered to be a soluble fiber and may also have prebiotic potential (33). Fiber has been recognized as important for normal laxation, primarily because it increases stool weight, and this has the secondary effect of reducing transit time. Fiber increases stool weight by its presence but also by increasing the water held by the fiber, as well as increasing bacterial mass from fermentation. However, inulin does not increase stool weight to the extent that wheat bran and psyllium do, but does undergo extensive fermentation (32).

Fiber is often recommended as a therapy to supplement dietary intake of fiber. However, as the studies above indicate, there are different formulations and types of fibers that have been evaluated and the included studies did not quantify the intake of dietary fiber. At least 2 studies were conducted with fiber that contained other ingredients such as milk or aloe vera that may additionally influence laxation. All studies are 30-40 years old, and the number of participants in the studies has been small and most are primarily conducted in women. Most of the included studies on fiber supplementation did not report on relevant patient important outcomes. The best data exist for psyllium, but even those are of low quality. In addition, wheat bran can exist as a finely ground powder that can decrease stool water content and harden stool (34). The chief side effect of fiber supplementation seems to be flatulence. In individuals with mild-to-moderate symptoms of constipation, especially who consume diets deficient in fiber, a trial of fiber supplementation is warranted because it is low-risk, low-cost, and easily accessible. In general, chronically constipated patients and nonconstipated

persons drink similar amounts of fluid on a daily basis. However, when individuals are placed in quartiles based on daily fluid intake, those in the lowest quartile for fluid intake are more likely to be constipated. Thus, efforts to increase fluid intake should be focused on those with low levels of fluid intake (35). Standard doses of fiber supplements are typically taken with 8–10 ounces of fluid (36).

OSMOTIC LAXATIVES

Recommendation 2: In adults with CIC, the panel recommends the use of PEG compared with management without PEG (strong recommendation, moderate certainty of evidence).

Implementation considerations

- A trial of fiber supplement can be considered for mild constipation before PEG use or in combination with PEG.
- Response to PEG has been shown to be durable over 6 months.
- Side effects include abdominal distension, loose stool, flatulence, and nausea.

Polyethylene glycol

Summary of evidence. Three randomized, placebo-controlled trials studied the effect of PEG on constipation (37-39). Two of these studies were multicenter trials conducted in Italy and enrolled participants aged 18-70 years with chronic constipation defined in accordance with the Rome criteria, that is, less than 2 bowel movements per week for at least 12 months or the presence of 2 or more of the following symptoms: <3 bowel movements per week, straining at defecation, sense of incomplete evacuation, and hard stools on at least 25% of occasions (37,38). In both trials, the treatment consisted of 17.5 g of PEG with electrolytes as a granular preparation or placebo dissolved in 250 mL of water taken twice daily. One study included 55 participants, and the treatment period lasted 8 weeks (38). The second study (37) included 70 participants and had 2 consecutive periods in which all participants received the active treatment, PEG for 4 weeks, and then those who responded were randomized to receive either PEG or placebo for 20 weeks. The third study was conducted in the United States and included 304 participants with chronic constipation based on modified Rome criteria, where participants reported <3 bowel movements per week for at least 3 months and one or more of the following: straining, lumpy or hard stools, and sensation of incomplete evacuation in >25% of defecations (39). Participants were randomized in a 2:1 ratio to PEG 3350 (n = 204) at a dose of 17 g or placebo (n = 204) at a dose of 18 g or placebo (n = 204) at a dose of 18 g or placebo (n= 100) mixed in 8 ounces of liquid once daily for 6 months.

Benefits and harms. PEG likely results in an increase in CSBMs per week compared with placebo (MD 2.90, CI 2.12-3.68), based on one study (39), and SBMs per week (MD 2.30, CI 1.55-3.06), based on meta-analyzed data from 3 studies (37-39). Across 2 studies, a higher rate of individuals met the responder definition, one study defined responder as normalization of bowel moments (37,38) and other defined based on the US Food and Drug Administration (FDA) end points (40), compared with placebo (RR 3.13, CI 2.00–4.89); PEG was associated with 312 more per 1,000 (from 146 more to 569 more). Diarrhea was noted more commonly in the treatment arm (158 more per 1,000, from 6 fewer to 896 more). A higher proportion of participants had global relief of symptoms with PEG compared with placebo with 454 per 1,000 in the PEG group (from 159 more to 948 more). There were no data for the following outcomes: stool form and quality of life. Two studies examined serious adverse events, but the number of

events was very small, and no conclusive statements could be made about risk of serious adverse events with the use of PEG (RR 0.47, CI 0.16-1.33).

Certainty in evidence of effects. The certainty of evidence for CSBMs, SBMs, responder rate, and global relief was moderate (because of imprecision). The certainty of evidence for serious adverse events was low because the CI includes both low and high risk of serious adverse events. The overall certainty of evidence for PEG was moderate.

Discussion

PEG is a long-chain polymer of ethylene oxide, which acts as an osmotic laxative. PEG is approved at a dose of 17 g daily for the treatment of occasional constipation by the FDA in the United States and is widely available OTC. Two of the studies used PEG with electrolytes given twice daily (37,38) while the other larger study evaluated the efficacy of PEG 3350 without electrolytes administered once daily (39). The 2 studies of PEG 3350 electrolytes measured the frequency of SBMs per week, but not CSBMs per week (37,38), while the PEG 3350-only study measured CSBM and SBM frequency along with other outcomes (39). Despite the differences in the PEG preparations, doses, and treatment durations, the studies all demonstrated that PEG was associated with a greater efficacy in increasing CSBMs, SBMs, responder rate, improvements in stool form, straining, and global relief compared with placebo, but not abdominal pain. Although PEG is approved by the FDA for the treatment of occasional constipation and not CIC, it has been shown to be efficacious in individuals with CIC for up to 6 months (39). There are additional treatment trials comparing the efficacy of PEG with tegaserod, prucalopride, and lactulose, in which PEG demonstrated a similar or greater efficacy in individuals with CIC than these other medications (41,42), although these trials used different primary end points.

There were no differences in side effect profiles observed between PEG and placebo, although data are limited. Individuals treated with PEG may experience bloating, flatulence, and diarrhea. These effects are consistent with and expected from laxative therapy, and most of these events were mild or moderate. However, PEG is widely available without the need for a prescription and is relatively inexpensive. It is, therefore, reasonable to use PEG earlier in the algorithm for the management of CIC, either after a trial of fiber supplementation or in combination with fiber supplementation.

Recommendation 3: In adults with CIC, the panel suggests the use of MgO over management without MgO (conditional recommendation, very low certainty of evidence certainty).

Implementation considerations

- The trials were conducted for 4 weeks, although longer term use is probably appropriate.
- The panel suggests starting at a lower dose, which may be increased if necessary.
- Avoid use in patients with renal insufficiency due to risk of hypermagnesemia.

Magnesium oxide

Summary of evidence. Two randomized, placebo-controlled trials evaluated the use of MgO for the management of CIC

(43,44). Both trials were completed in Japan. The dose of MgO studied was 1.5 g/d for 4 weeks. The 2 trials randomized a total of 47 participants to MgO and 47 participants to the placebo arm, and 93% of the participants were females. At baseline, participants randomized to the placebo group had 4.6 SBMs per week. Those randomized to MgO had 4 SBMs per week at baseline.

Benefits and harms. Compared with placebo, treatment with MgO may increase the number of CSBMs per week (MD 4.29, 95% CI 2.93–5.65) and SBMs per week (MD 3.59, 95% CI 2.64–4.54). Participants treated with MgO achieved a higher treatment response compared with placebo (RR 3.93, 95% CI 2.04–7.56). In absolute terms, 499 more participants per 1,000 might respond to MgO (from 177 to 1,000 more). There was little to no difference in the degree of diarrhea leading to treatment dose change or discontinuation between the 2 study groups (RR 1.07, 95% CI 0.65–1.74). Participants treated with MgO may have better quality-of-life scores as measured by PAC-QOL (MD 16.23, 95% CI 11.44–21.01) and better stool consistency based on the Bristol Stool Form Scale (MD 1.89, 95% CI 1.44–2.33).

Certainty in evidence of effects. The certainty of evidence was very low for the outcomes of CSBMs per week and SBMs per week because of concerns related to inconsistency, indirectness, and imprecision. The certainty of evidence was moderate for the outcomes of responder rate and adverse events due to diarrhea because of imprecision and inconsistency, respectively. The outcomes of quality of life and stool form were rated as low certainty because of indirectness and imprecision. The overall certainty of evidence for MgO was very low.

Discussion

Magnesium is a naturally occurring element that plays an important role in a wide range of biological and biochemical processes (45). Within the lumen of the GI tract, nonabsorbed magnesium creates an osmotic gradient, which leads to net secretion of water and electrolytes, which can exert a beneficial effect on constipation-related symptoms. MgO dosing in the available RCTs was 1.5 g/d. Although not studied in RCTs, lower MgO doses of 500 mg/d to 1 g/d are often used in clinical practice. Only MgO has been evaluated in RCTs; the bioavailability and clinical efficacy of other formulations of magnesium (e.g., citrate, glycinate, lactate, malate, sulfate) for CIC are unknown. Data on adverse effects of MgO from the available trials are limited. The available data suggest no increased reports of diarrhea with MgO compared with placebo (43). Systemic regulation of magnesium levels is maintained by renal excretion (46). Therefore, hypermagnesemia is more likely to occur in individuals with significant renal impairment and magnesium supplements should be avoided in those with a creatinine clearance of <20 mg/dL (47).

The combination of efficacy, tolerability, availability of OTC, and low cost make MgO an attractive first-line option for individuals with CIC. Limitations to consider include the small number of clinical trials and included participants with CIC, all trials being conducted in Japan, formulations other than MgO not being evaluated, the dose of MgO used in trials being higher than

that typically used in clinical practice, and no long-term effectiveness or harms data being available.

Recommendation 4: In adults with CIC who fail or are intolerant to OTC therapies, the panel suggests the use of lactulose over management without lactulose (conditional recommendation, very low certainty of evidence).

Implementation considerations

 Bloating and flatulence are dose-dependent and common side effects, which may limit its use in clinical practice.

Lactulose

Summary of the evidence. Two RCTs studied the efficacy of lactulose syrup for the treatment of CIC in elderly participants (48,49). One multicenter study performed in the Netherlands included 103 participants who were regularly taking laxatives for the treatment of chronic constipation (49). The initial dose of 15 mL of either 50% lactulose syrup or 50% glucose syrup was administered daily for a total of 3 weeks. The daily dose was reduced by half after 3 consecutive days with defecation, but if no defecation occurred for more than 48 hours, the dose was doubled. If defecation occurred on 3 consecutive days with the doubled dose, the dose was reduced back to 15 mL (49). The second study, conducted in the United States, included 55 constipated participants (48). Participants received 30 mL daily of either 50% lactulose syrup or 50% glucose syrup, taken at bedtime for 12 weeks (48).

Benefits and harms. Based on one study, lactulose may have little to no effect on SBMs per week (MD 0.35, CI -0.91 to 1.61). A second study, however, showed that lactulose may be associated with a large increase in global relief (RR 2.42, CI 1.29–4.54, in absolute terms, 473 more per 1,000, CI 97 more to 1,000 more). Across the 2 studies, there was a higher rate of individuals taking lactulose who met the responder definition (defined as >1 SBM from baseline in 1 study and lack of need of other laxatives in other study) compared with placebo. Lactulose was associated with 267 more per 1,000 (from 108 more to 471 more) responders. The studies did not report on CSBMs per week, diarrhea, serious adverse events, quality of life, or stool form.

Certainty in evidence of effects. The certainty of evidence for SBMs was very low because of risk of bias (unclear methods of randomization and blinding) and imprecision. The certainty of evidence for global relief and responder rate was low because of concerns for risk of bias and imprecision. The overall certainty of evidence for lactulose was very low.

Discussion

Lactulose is β -galactosido-fructose, a synthetic disaccharide not digested in the small intestine that exerts an osmotic laxative effect in the colon to promote peristalsis. It is approved by the FDA in the United States for the treatment of constipation at a dose of 10–20 g (15–30 mL or 1–2 packets) daily and is widely available in other countries. The dose may be increased to 40 g (60 mL or 2–4 packets) daily if needed. There were significant limitations in the 2 RCTs of lactulose; both trials were conducted over 40 years ago, included relatively small numbers of elderly participants, and did not report the diagnostic criteria for constipation (48,49). In the US study, most participants were women living in a nursing home and medical facility and the mean age was in the mid-

80s (48). Bowel movement frequency and the severity of symptoms improved to a greater degree in the lactulose group compared with the glucose group. Interestingly, the most dramatic finding was the decrease in impactions and need for enemas in individuals receiving lactulose. The other study from the Netherlands did not report demographics of the patient population.

There were minimal data on adverse events from the 2 published studies (48,49); however, bloating and flatulence (which are dose-dependent) are considered very common side effects of lactulose in clinical practice, which limit its use. Some brands of lactulose may be expensive, although generic lactulose is generally low cost. Lactulose can be considered if symptoms of CIC have failed to improve with fiber and OTC laxatives, and individuals do not experience significant bloating or abdominal pain with lactulose use. The use of lactulose in mildly constipated, noninsulindependent patients with diabetes mellitus type 2 may not lead to increase in blood sugar levels (50).

STIMULANT LAXATIVES

Recommendation 5: In adults with CIC, the panel recommends the use of bisacodyl or sodium picosulfate (SPS) short term or as rescue therapy over management without bisacodyl or SPS (strong recommendation, moderate certainty of evidence).

Implementation considerations

- Short-term use is defined as daily use for 4 weeks or less. While long-term use is probably appropriate, data are needed to better understand tolerance and side effects.
- This is a good option for occasional use or rescue therapy in combination with other pharmacological agents for CIC.
- The most common side effects are abdominal pain, cramping and diarrhea. The panel suggests starting at a lower dose and increasing the dose as tolerated.

Summary of evidence. The panel considered studies that evaluated bisacodyl and SPS, which are mechanistically related, for the management of CIC. Of note, SPS tablets/drops are not available for use in the United States; however, they are approved for use in Europe. In the United States, SPS is available in combination with other laxatives and used for bowel preparation before colonoscopy. Given their common mechanism of action and limited number of trials on these drugs in treatment of CIC, the data from available trials were pooled in calculations of estimates of effect. A total of 2 studies were included. One of these studies was a multicenter randomized double-blinded placebo-controlled trial in the United Kingdom where recruited participants were assigned in a 2:1 ratio to bisacodyl (n = 247) vs placebo (n = 121) and treated for 4 weeks (51). The other study examined effects of SPS in a multicenter double-blinded placebo-controlled RCT conducted in Germany, and participants were randomized in a 2:1 ratio to SPS (n = 229) or placebo (n = 133) and treated for 4 weeks (52).

Benefits and harms. Based on meta-analyzed data from 2 studies, SPS likely leads to a large increase in CSBMs per week (MD 2.54, 95% CI 1.07–4.01) and SBMs per week (MD 4.04, 95% CI 2.37–5.71) and improved the consistency of stool on the Bristol Stool Form Scale (53) (MD 2.4 points higher, 95% CI

2.07–2.73) and PAC-QOL scores (MD 0.65, 95% CI 0.50–0.80) compared with placebo. Furthermore, the use of SPS leads to higher responder rates (RR 2.60, 95% CI 2.05–3.30) and an increased proportion of individuals with global relief (RR 1.75, 95% CI 1.48–2.07). In absolute terms, of 1,000 individuals treated with SPS, there would be 359 more responders (236 more to 516 more) and 357 more with global relief (228 more to 509 more). Use of SPS may increase the proportion of individuals who experience diarrhea compared with placebo (RR 8.76, 95% CI 4.99–15.39). One study reported serious adverse events but with only 3 events, results were very imprecise (RR 0.24, 95% CI 0.02–2.67). The rate of diarrhea leading to discontinuation of treatment was higher in the SPS group compared with placebo (RR 8.76, 95% CI 4.99–15.39).

Certainty in evidence of effects. The certainty of evidence was rated as moderate for the following outcomes: CSBM and SBM frequency, responder rate, global relief, and stool consistency (because of risk of bias). The certainty of evidence was very low for the outcomes of diarrhea and serious adverse events (because of risk of bias, indirectness, and imprecision) and low for quality of life (because of risk of bias and indirectness). The overall certainty of evidence for bisacodyl was moderate.

Discussion

Bisacodyl and SPS are converted in the gut into the same active metabolite, bis-(phydroxyphenyl)-pyridyl-2-methane (BHPM). Bisacodyl is converted into BHPM by small bowel and colonic mucosal deacetylase enzymes while SPS is converted into BHPM by colonic bacteria desulfate enzymes. BHPM acts directly on the colonic mucosa to stimulate colonic peristalsis and secretion. Similar to other stimulant laxatives (e.g., senna), use of antibiotics can potentially decrease the efficacy of SPS because they may affect colonic bacteria that produce the active metabolite of the drug (54).

Initial dosing in the available RCTs was 10 mg orally for bisacodyl and SPS, although dose reduction was permitted. At this dose, adverse effects were common (see below), and therefore in clinical practice, 5 mg orally is often used initially. Although not studied in RCTs, bisacodyl is also available as a rectal suppository (10 mg). The onset of action is typically 6–12 hours for the oral tablet while the suppository works within 30–60 minutes.

The most common adverse effects for bisacodyl and SPS were diarrhea and abdominal pain. For bisacodyl at the initial starting dose of 10 mg compared with placebo, diarrhea occurred in 53.4% vs 1.7%, respectively, while abdominal pain occurred in 24.7% vs 2.5%, respectively (51). Most adverse events occurred in the first week of treatment. For SPS at the initial starting dose of 10 mg compared with placebo, diarrhea was reported by 31.8% vs 4.5%, respectively, while for abdominal pain, it was reported by 5.6% vs 2.2%, respectively (52). Bisacodyl and SPS are contraindicated in individuals with ileus, intestinal obstruction, severe dehydration, or acute inflammatory conditions in the bowel.

Although effective, side effects are common, and the panel recommended the use of bisacodyl and SPS for a short term or rescue therapy. The long-term effectiveness of these agents has not been studied.

Recommendation 6: In adults with CIC, the panel suggests the use of senna over management without senna (conditional recommendation, low certainty of evidence).

Implementation considerations

- While the trials were conducted for 4 weeks, longer term use is probably appropriate, but data are needed to better understand tolerance and side effects.
- The dose evaluated in trials is higher than commonly used doses in practice. The panel suggests starting at a lower dose and increase if there is no response.
- Abdominal pain and cramping may occur with a higher dose of senna

Senna

Summary of evidence. One placebo-controlled RCT examined the safety and efficacy of senna in the management of CIC (44). Participants were randomly assigned to 1 g of senna (n = 30) or placebo (n = 30) and treated for 28 days.

Benefits and harms. Participants treated with senna may have higher CSBMs per week (MD 7.60, 95% CI 5.90–9.30) and SBMs per week (MD 7.6, 95% CI 6.42–8.78) compared with the placebo group. The response rate might be higher in the senna-treated group compared with placebo (RR 5.25, 95% CI 2.05–13.47), 567 more per 1,000 in the senna group (from 140 to 1,000 more). The quality-of-life scores may be higher in the senna group compared with placebo (MD 7.80, 95% CI 1.40–14.20). Participants taking senna might have higher rates of diarrhea, 175 more per 1,000 (from 100 fewer to 1,000 more). No participants in the senna and placebo arms experienced a severe treatment-related adverse event.

Certainty in evidence of effects. The certainty of evidence was low for the outcomes of CSBMs per week, SBMs per week, and quality of life as the panel rated down because of concerns for indirectness and imprecision. The certainty of evidence for responder rate was moderate because of imprecision. The overall certainty of evidence for senna was low.

Discussion

Senna is a natural derivative of the senna plant. Sennosides A and B are sequentially metabolized by the gut microbiota to the active metabolites, rheinanthrone and rhein, which stimulate the production of prostaglandin E2 and secretion of chloride ions leading to attendant changes in colonic peristalsis and luminal water content (55,56). Over 90% of sennosides and their metabolites are excreted in the feces (56). Dosing in the single RCT published to date was 1 g by mouth daily for 4 weeks, which is higher than that typically used in clinical practice. It is notable that while no details were provided, 83% of participants randomized to senna reduced their daily dose during the trial (44). Most commercially available senna products contain 8-9 mg per tablet. Rigorous dose ranging data with senna are currently not available. In the clinical trial by Moshita et al. (44), no participants experienced a severe treatment-related adverse event. However, as Moshita et al. (44) did not provide rates for the mild treatment-related adverse events, the fact that 83% of participants with CIC randomized to senna engaged in dose reduction raises concerns about potential adverse events such as abdominal pain, cramping, or diarrhea with the higher dose of senna. There are no long-term safety studies with senna in humans. Sennosides are not recommended in pregnant women because chemically similar substances have been found to exert weak genotoxic effects in animals, although the supporting evidence is controversial (57).

The combination of efficacy, impact on quality of life, availability OTC, and low cost makes senna an attractive first-line option for individuals with CIC. Limitations to consider include the following: only a single, small RCT from Japan supports its efficacy; the dose of senna used in the trial is higher than that typically used in clinical practice; there are no long-term effectiveness data; and very limited short-term and no long-term harms data are available.

SECRETAGOGUES

Recommendation 7: In adults with CIC who do not respond to OTC agents, the panel suggests the use of lubiprostone over management without lubiprostone (conditional recommendation, low certainty of evidence).

Implementation considerations

- Can be used as a replacement or as an adjunct to OTC agents.
- Duration of treatment in trials was 4 weeks, but the drug label does not provide a limit.
- Nausea may occur; however, the risk of nausea is dosedependent and seems to be lower when taken with food and water.

Lubiprostone

Summary of evidence. Three 4-week randomized double-blinded placebo-controlled trials evaluated the use of lubiprostone for the management of CIC (58-60). The studied dose of lubiprostone was 24 µg twice daily, and studies were conducted in the United States and Japan. Lubiprostone is a chloride channel activator, resulting in increased intestinal fluid and accelerated GI transit. **Benefits and harms.** The pooled data showed that lubiprostone resulted in an increased number of SBMs per week compared with placebo (MD 1.98, 95% CI 1.17-2.79). The data for the outcome of CSBMs per week were not available. Use of lubiprostone in adults with CIC may increase responder rates (RR 1.67, 95% CI 1.36–2.06), 226 more per 1,000 (from 122 to 358 more). Individuals may also be at increased risk of diarrhea leading to discontinuation of the treatment compared with placebo (RR 5.30, 95% CI 1.53–18.44), 28 more per 1,000 (from 4 more to 115 more). There was little to no difference in serious adverse events; however, the CI around the summary estimate was wide, and increased risk of serious adverse events could not be ruled out (RR 1.22, 95% CI 0.62-2.42). The data on quality of life from the available studies were not reported. Stool form, using a 0- to 4point scale (very loose to very hard, where a lower score is better), was evaluated in 2 studies and improved in the lubiprostone group (MD 1.09 lower, 95% CI 0.16-2.03 lower). Finally, the rate of global relief, evaluated in one study using a 0- to 4-point scale (not effective to very effective, where higher is better), was higher in the lubiprostone group (MD 0.75, 95% CI 0.42–1.08 higher). *Certainty in evidence of effects.* The certainty of evidence was moderate for the outcome of SBMs per week (because of imprecision) and low for the remainder of the outcomes (because of very serious imprecision). The overall certainty in evidence for lubiprostone was low.

Discussion

Lubiprostone, a bicyclic fatty acid derived from prostaglandin E1 that increases intestinal chloride secretion by activating type 2 chloride channels on epithelial cells, is approved by FDA for treating CIC at a dose of 24 µg 2 times daily. For IBS-C, the approved dose is 8 µg 2 times daily. Lubiprostone improved stool frequency and consistency as well as abdominal discomfort and bloating, which is a bothersome symptom in some individuals with CIC (60,61). Among individuals who respond, these effects generally manifest within 2 days. The efficacy in persons 65 years and older is comparable with the overall study population. Lubiprostone accelerates small intestinal and colonic transit in healthy people (62), should be taken with meals, and is contraindicated in individuals with known or suspected mechanical GI obstruction. Observed in 35% of individuals, nausea was the most common adverse event and typically mild or moderate, but led to discontinuation of therapy in only 5% of individuals (60). The risk of nausea is dose-dependent and seems to be lower when taken with food and water (63).

Systemic absorption of oral lubiprostone is negligible. Individuals with moderate or severe hepatic insufficiency should receive a lower dose (i.e., 8 µg twice daily).

Recommendation 8: In adults with CIC who do not respond to OTC agents, the panel recommends the use of linaclotide over management without linaclotide (strong recommendation, moderate certainty of evidence).

Implementation considerations

- Can be used as a replacement or as an adjunct to OTC agents
- Duration of treatment in trials was 12 weeks, but the drug label does not provide a limit.
- May be associated with side effects of diarrhea leading to discontinuation of treatment

Summary of evidence. Three 12-week randomized double-blinded placebo-controlled trials evaluated the use of linaclotide for the management of CIC (10,64). The studied doses of linaclotide were 145 and 290 μg daily, and all studies were conducted in the United States and Canada. The following dose has also been studied (72 μg) (65). Linaclotide is a guanylate cyclase-C agonist, which increases cyclic guanosine monophosphate concentrations resulting in luminal chloride and bicarbonate secretion, thereby increasing intestinal fluid and accelerating GI transit.

Benefits and harms. The use of linaclotide leads to increases in the number of CSBMs per week (MD 1.37, 95% CI 1.07–1.95) and SBMs per week (MD 1.97, 95% CI 1.59–2.36), improves stool consistency (MD 1.25, 95% CI 1.1–1.39 higher), and increases the rates of global relief (RR 1.96, 95% CI 1.63–2.35). The use of linaclotide might lead to a large increase in responder rates compared with placebo (RR 3.14, 95% CI 1.68–5.88), 119 more per 1,000 (from 38 to 271 more). However, participants treated with linaclotide might be 3 times more likely to have diarrhea leading to treatment discontinuation compared with placebo (RR 3.35, 2.09–5.36), 83 more per 1,000 (from 38 to 154 more). The use of linaclotide might improve the PAC-QOL scores compared with placebo (64); however, data could not be pooled.

Certainty in evidence of effects. We rated the certainty of evidence as high for outcomes of CSBMs per week, SBMs per week, stool form, and global relief and moderate for the responder outcome and diarrhea (rating down for imprecision). The outcome of serious adverse events was rated down to low because of very serious imprecision. The overall certainty of evidence for linaclotide was moderate.

Discussion

Linaclotide is a guanylate cyclase-C agonist FDA-approved for the treatment of CIC at a dose of 72 μg or 145 μg daily. The 290 μg daily dose is approved for IBS-C, recognizing that CIC and IBS-C overlap and are often indistinguishable in practice (66). Linaclotide is also approved in many other countries. Linaclotide has been demonstrated to improve abdominal symptoms of bloating, discomfort, and pain in IBS-C trials (67). Because of its effect on abdominal discomfort, pain, and bloating, it may be useful in individuals with these coexisting symptoms. Patients should be instructed to take linaclotide without food, at least 30 minutes before the first meal of the day. Linaclotide is contraindicated in individuals with known or suspected mechanical GI obstruction.

The use of linaclotide might be associated with diarrhea leading to discontinuation or dose reduction; however, this was not very common (in one study, 4.7% discontinued the medication because of diarrhea) (68,69). The most common reasons for discontinuation over the first year of treatment were loss of efficacy and insurance coverage barriers related to obtaining prescription refills and not discontinuations because of adverse events, in a retrospective analysis at a large health system (70). Descriptively, there were no clear differences in outcomes among individuals older than 65 years in clinical trials, although the sample size was too small to support formal analysis on differences in outcomes related to age.

Recommendation 9: In adults with CIC who do not respond to OTC agents, the panel recommends the use of plecanatide over management without plecanatide (strong recommendation, moderate certainty of evidence).

Implementation considerations

- Can be used as a replacement or as an adjunct to OTC agents
- Duration of treatment in trials was 12 weeks, but the drug label does not provide a limit.
- May be associated with side effects of diarrhea leading to discontinuation of treatment

Plecanatide

Summary of evidence. Three 12-week randomized double-blinded placebo-controlled trials evaluated the use of plecanatide for the management of CIC (71–73). The studied dose of plecanatide was 3 mg/6 mg daily, and all studies were conducted in the United States and Canada. Plecanatide is a guanylate cyclase-C agonist, which increases cyclic guanosine monophosphate concentrations resulting in luminal chloride and bicarbonate secretion, thereby increasing intestinal fluid and accelerating GI transit.

Benefits and harms. The pooled data showed that the use of plecanatide in adults with CIC leads to an increase in the number of CSBMs per week (MD 1.1, 95% CI 0.85–1.35) and SBMs per week (MD 1.66, 95% CI 1.37–1.94) and improves the quality-of-life scores. The intervention group had increased responder rates,

defined as ≥3 CSBMs per week and ≥1 CBSM over baseline for ≥9 of 12 weeks including ≥3 of the last 4 weeks, compared with placebo (RR 1.78, 95% CI 1.46–2.18), 88 more per 1,000 (from 52 to 134 more). Participants treated with plecanatide might have higher rates of reported diarrhea leading to treatment discontinuation (RR 5.39, 95% CI 2.40–12.11), 27 more per 1,000 (from 9 to 69 more). The use of plecanatide might improve stool consistency based on the Bristol Stool Form Scale compared with placebo (MD 0.83, 95% CI 0.6–1.05).

Certainty in evidence of effects. The certainty of evidence was high for outcomes of CSBM and SBM frequency and QOL and moderate for diarrhea, leading to treatment discontinuation, serious adverse events, and stool form. The panel rated down certainty of these outcomes because of imprecision. The overall certainty in evidence for plecanatide was moderate.

Discussion

Plecanatide is a pH-dependent guanylate cyclase-C agonist approved by the FDA for CIC at a dose of 3 mg daily taken with or without food. Plecanatide is also approved at the same dose for IBS-C. Plecanatide may have beneficial concurrent effects with relief in abdominal pain based on indirect evidence from IBS-C trials (74). Individuals using plecanatide might be at higher risk of diarrhea leading to discontinuation of medication; however, the absolute risk seems small (68). Descriptively, there were no clear differences in outcomes among individuals older than 65 years in clinical trials, although the sample size was too small to support formal analysis on differences in outcomes related to age.

5-HT₄ AGONIST

Recommendation 10: In adults with CIC who do not respond to OTC agents, the panel recommends the use of prucalopride over management without prucalopride (strong recommendation, moderate certainty of evidence).

Implementation considerations

- Duration of treatment in trials was 4–24 weeks, but the drug label does not provide a limit.
- Can be used as a replacement or as an adjunct to OTC agents
- May be associated with side effects of headache, abdominal pain, nausea, and diarrhea

Prucalopride

Summary of evidence. Five 12-week randomized double-blinded placebo-controlled trials evaluated the use of prucalopride (2 mg daily) for the management of CIC (75–79). The studies were conducted in the United States, Europe, and the Asia-Pacific region. The 4 mg dose has also been studied (75). Prucalopride is a selective, high-affinity 5-HT $_4$ agonist that promotes neurotransmission by the enteric neurons resulting in stimulation of the peristaltic reflex, intestinal secretions, and GI motility.

Benefits and harms. Compared with placebo, prucalopride resulted in an increased number of CSBMs per week (MD 0.96, 95% CI 0.64–1.29). SBMs per week was not studied in any of the included study. Responder rates, defined as ≥3 CSBMs per week, were higher in the prucalopride group (RR 2.37, 95% CI

1.97-2.85) with 165 more responders per 1,000 (range 117-222 more). An alternative responder end point, deemed alternative end point A, defined as ≥ 3 CSBM per week and ≥ 1 CBSM over baseline for \geq 75% of study weeks, was higher in the prucalopride group (RR 2.51, 95% CI 1.97-3.21) with 109 more responders per 1,000 (range 70-160 more). The rates of diarrhea leading to treatment discontinuation might be higher in the prucalopride group compared with placebo (RR 3.00, 95% CI 1.89-4.78). The occurrence of serious adverse events was low; however, the CI around the summary estimate was imprecise and included a possible increased risk. PAC-QOL, where lower scores are better, improved in 4 studies in the prucal opride group compared with placebo (MD 0.32 lower, 95% CI 0.41-0.23 lower). Definitions and scales used to assess stool form varied widely across studies and could not be pooled. Global relief was reported in 4 studies and defined as those who felt that treatment was extremely or quite a bit effective, and the responder rates were higher in the prucalopride group compared with placebo (RR 2.09, 95% CI 0.15-3.0).

Certainty in evidence of effects. We rated the certainty of evidence as high for outcomes of CSBM frequency, responder rate, and alternative end point A and moderate for diarrhea, leading to treatment discontinuation, serious adverse events, quality of life, and global relief (because of small event rates and wide CIs around the summary estimates). The overall certainty in evidence for prucalopride was moderate.

Discussion

For prucalopride, a selective agonist of serotonin 5-HT $_4$ receptors, the recommended dose is 2 mg once daily in adults and 1 mg daily in individuals with severe renal impairment (i.e., creatinine clearance <30 mL/min) (80). The efficacy in persons 65 years and older is comparable with the overall study population. Besides increasing bowel frequency, prucalopride also improved constipation symptoms, abdominal symptoms, quality of life, and satisfaction with treatment vs placebo assessed with the PAC instrument (81). Arguably, these effects are at least partly explained by the ability of prucalopride to induce and increase the amplitude of colonic high-amplitude propagated contractions (73,82). Such high-amplitude propagated contractions propagate colonic contents (83).

The most frequent, generally transient, side effects are headache, abdominal pain, nausea, and diarrhea (84). In most of the individuals who reported headache and diarrhea, this side effect occurred within the first week of treatment and typically resolved within a few days. Five percent of individuals discontinued prucalopride because of side effects. Cardiovascular adverse events were not more common than placebo. In a safety database of 4,476 subjects, 4 individuals attempted suicides and 2 completed suicides, both of whom had discontinued prucalopride more than 1 month before the event. The label cautions patients and clinicians to be alert to unusual changes in mood and behavior and suicidal ideation. It is, however, unclear what the mechanism of action is or whether there is a causal association between the use of prucalopride and risk of suicide (85). No drug-associated risks of miscarriage, major birth defects, or adverse maternal or fetal outcomes have been identified. Prucalopride is contraindicated in patients with intestinal perforation or obstruction, Crohn's disease, ulcerative colitis, and toxic megacolon/megarectum.

LIMITATIONS AND EVIDENCE GAP

An important limitation of this body of evidence was that clinical trials did not uniformly evaluate interventions for patient important outcomes on efficacy, adverse effects, and tolerability. Importantly, there was a paucity of data for the most commonly used treatments of CIC such as fiber, lactulose, senna, and docusate. There was also variability in the definition of inclusion criteria, efficacy, and tolerability outcomes, as well as variance in acceptable clinical trial length by regulators over time. Most of the included studies followed the patients for the short term, and the safety and tolerance of these medications in the long term is not well studied. Future research is needed to assess the long-term safety of these medications and to assess whether the patients develop tolerance to these medications over time. In our systematic review, we compared individual drugs against placebo arms and did not aim to inform the relative efficacy of pharmacological agents. Network meta-analysis is an appropriate statistical method to facilitate indirect comparison against a common comparator such as placebo or other active treatment (8). However, for the reasons stated, readers should be cautioned on the limitation of indirect comparisons to support substantive claims on superiority or inferiority to inform care or policy decisions.

This guideline is limited to covering pharmacological interventions for the treatment of CIC in otherwise healthy adults and does not apply to pediatric populations or to individuals who are pregnant or with opioid-induced constipation or malignancy. The evidence on the management of constipation during pregnancy has been reviewed in a recent publication that discusses the safety of almost all the pharmacological agents assessed in this guideline (86). This guideline does not review anorectal evacuation disorders that were evaluated in a recent ACG guideline (87) and an AGA review (80). We also did not assess the efficacy of dietary fiber including fruit-based laxatives in CIC, which was evaluated in a recent systematic review (8). Other interventions not included in this review include lifestyle modifications, such as increasing water and physical activity, and other pharmacological agents, such as elobixibat, mizagliflozin, naronapride, tegaserod, tenapanor, or velusetrag, or the efficacy of surgical interventions for the management of CIC. We did not assess the evidence on the use of probiotics for the treatment of CIC, but it has been synthesized elsewhere in a recent systematic review (88). Although we considered the cost of pharmacological agents evaluated in this guideline during the evidence to decision-making process, we did not perform formal cost-effectiveness analyses and refer the audience to recently published evidence that addresses this topic (89). There was no patient representative in the guideline development panel, which is a limitation for this study.

IMPLEMENTATION, COST, AND HEALTH EQUITY CONSIDERATIONS

This document provides a comprehensive outline of the various OTC and prescription pharmacological agents available for the treatment of CIC. The guidelines are meant to provide a template for approach to management and practitioners should engage in shared decision making based on the preference of patients and cost and availability of the medications. Although the recommendations in this guideline were based on available evidence, the implementation considerations included suggestions from the collective experience of the expert panel and may not be based on

evidence. Most of the medications assessed in this guideline document are readily available; however, some of them are still available only in brand name formulations because generic formulations do not exist. As a result, it is important to consider the out-of-pocket expenses for patients that may depend on prescription coverage with various insurance plans (89). Prior authorization might be required for some of the medications (90). The clinical decision support tool is available from the website of AGA.

PLANS FOR UPDATING

Considerable resources are expended for the development of guidelines, and keeping guidelines up to date is a challenging process. Future update of this guideline will depend on the availability of new evidence on the existing interventions and new intervention. We hope to incorporate the advances in the technological platforms and models of guideline development in the future updates without duplication or reproduction of the current guideline document.

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CONFLICTS OF INTEREST

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