Colonic preparation before colonoscopy in constipated and non-constipated patients: A randomized study

Lisandro Pereyra, Daniel Cimmino, Carlos González Malla, Mariano Laporte, Nicolás Rotholtz, Carlos Peczan, Sandra Lencinas, Silvia Pedreira, Hugo Catalano, Luis Boerr

AIM: To compare the efficacy of different doses of sodium phosphate (NaP) and polyethyenglicol (PEG) alone or with bisacodyl for colonic cleansing in constipated and non-constipated patients.

METHODS: Three hundred and forty-nine patients, older than 18 years old, with low risk for renal damage and who were scheduled for outpatient colonoscopy were randomized to receive one of the following preparations (prep): 90 mL of NaP (prep 1); 45 mL of NaP + 20 mg of bisacodyl (prep 2); 4 L of PEG (prep 3) or 2 L of PEG + 20 mg of bisacodyl (prep 4). Randomization was stratified by constipation. Patients, endoscopists, endoscopists’ assistants and data analysts were blinded. A blinding challenge was performed to endoscopist in order to reassure blinding. The primary outcome was the efficacy of colonic cleansing using a previous reported scale. Secondary outcomes were tolerability, compliance, side effects, endoscopist perception about the necessity to repeat the study due to an inadequate colonic preparation and patient overall perceptions.

RESULTS: Information about the primary outcome was obtained from 324 patients (93%). There were no significant differences regarding the preparation quality among different groups in the overall analysis. Compliance was higher in the NaP preparations being even higher in half-dose with bisacodyl: 94% (prep 1), 100% (prep 2), 81% (prep 3) and 87% (prep 4) (2 vs 1, 3 and 4, P < 0.01; 1 vs 3, 4, P < 0.05). The combination of bisacodyl with NaP was associated with insomnia (P = 0.04). In non-constipated patients the preparation quality was also similar between different groups, but endoscopist appraisal about the need to repeat the study was more frequent in the half-dose PEG plus bisacodyl than in whole dose NaP preparation: 11% (prep 4) vs 2% (prep 1) (P < 0.05). Compliance in this group was also higher with the NaP preparations: 95% (prep 1), 100% (prep 2) vs 80% (prep 3) (P < 0.05). Bisacodyl was associated with abdominal pain: 13% (prep 1), 31% (prep 2), 21% (prep 3) and 29% (prep 4), (2, 4 vs 1, 2, P < 0.05). In constipated patients the combination of NaP plus bisacodyl presented higher rates of satisfactory colonic cleansing than whole those PEG: 95% (prep 2) vs 66% (prep 3) (P = 0.03). Preparations containing bisacodyl were not associated with adverse effects in constipated patients.

CONCLUSION: In non-constipated patients, compliance is higher with NaP preparations, and bisacodyl is related to adverse effects. In constipated patients NaP plus bisacodyl is the most effective preparation.
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Key words: Colonic cleansing; Sodium phosphate; Poly- 
ethylenglicol; Bisacodyl constipation; Colonoscopy

Core tip: Colonoscopy has become the standard pro-
dure for the diagnosis and treatment of colon diseases. Ade-
quately bowel cleansing is essential for a high-quality 
effective and safe colonoscopy. In non-constipated 
patients, compliance is higher with sodium phosphate 
(NaP) preparations, and bisacodyl is related to adverse 
effects. In constipated patients NaP plus bisacodyl is 
the most effective preparation.

Pereyra L, Cimmino D, González Malla C, Laporte M, Rotholtz 
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INTRODUCTION

Colonoscopy has become the standard procedure for the 
diagnosis and treatment of colon diseases[10]. Adequate 
colonic cleansing is necessary for a proper evaluation of 
the entire colonic mucosa and therefore for achieving a 
high quality colonoscopy[19]. Sodium phosphate (NaP) is a 
small volume hyperosmotic solution that provides effec-
tive colonic cleansing in preparation for colonoscopy. In 
the past years the popularity of orally administered NaP 
has increased because of its superior tolerance by pa-
tients compared with large-volume cleansing agents such 
as polyethylene glycol electrolyte solutions[3-5]. Although it 
presents a safety profile similar to other colonic cleansing 
agents, serious adverse events have been reported when 
administered in high volume or in patients with contrain-
dications to NaP[9]. Polyethyleneglicol (PEG) solutions are 
the most commonly used laxatives for colonic cleansing 
because of their safety profile and lack of contraindica-
tion. However, unpleasant taste and large volume of 
P EG lead to poor compliance and result in patient dis-
atisfaction. The two aforementioned agents are the most 
frequently used for colonic cleansing in many counties 
and despite the significant heterogeneity between differ-
ent studies comparing them for colonic preparation, a 
systematic review showed similar adequate preparation 
ratios, 75% for NaP and 71% for PEG[5,8]. Numerous 
clinical trials have also assessed prokinetic (metoclo-
primide, cisapride and tegazerod)[10-13] and laxative agents 
(magnesium citrate and bisacodyl)[14-16] associated with 
standard or lower volumes of this colonic cleansing agents. 
Sharma et al[10] found that pretreatment with magnesium 
citrate or bisacodyl in addition to half-dose of PEG was 
associated with better preparation quality and patient sat-
satisfaction than full-dose of PEG. To our best knowledge, 
there is no study directly comparing whole and half-dose 
of PEG and NaP alone or in combination with bisacodyl 
in constipated and non-constipated patients. The aim of 
this study was to compare the efficacy and tolerability of 
whole doses of NaP and PEG and half-doses of those 
agents in combination with bisacodyl for colonic cleansing 
in constipated and non-constipated patients.

MATERIALS AND METHODS

This was a randomized, double-blind, four-arm study stratified by constipation. The study was carried out in 
accordance with the declaration of Helsinki. All patients 
included in the study signed an informed consent form. The human ethics committee from our institution ap-
proved the protocol.

Study population

All patients older than 18 years old who were scheduled 
for an elective outpatient’s colonoscopy were eligible for 
participating in the study and were randomized in a 6-mo 
period (June-December 2011). As safety issues about 
NaP solutions have emerged, we only included healthy 
patients following the United States Food and Drug 
Administration recommendation to avoid renal damage. 
Patients were excluded if they presented one or more of 
the following characteristics: age younger than 18 years 
old, were hospitalized for any reason, hypersensibility 
to any of the components of PEG, NaP or bisacodyl, 
were under more than one antihypertensive medication, 
presented history of diarrhea (more than 3 bowel move-
ments a day), acute or chronic renal failure, cardiovascular 
disease (history of myocardial infarction, congestive heart 
failure, unstable angina pectoris, unstable hypertension 
and/or cardiac arrhythmia), ascites, electrolyte imbalance 
(hyponatremia, hipokalemia, hipocalcemia, hipomagnes-
emia or hyperphosphatemia), inflammatory bowel disease, partial or subtotal colectomy, ileus or suspected intestinal 
obstruction and pregnancy or breastfeeding, childbearing 
potential without contraception.

Study design

Patients who met all the inclusion criteria and no exclu-
sion criteria were randomly assigned to receive one of 
the four colonic preparations according to a computer-
generated randomization list. Randomization was stratifi-
cation by constipation in order to make a subgroup analy-
sis of constipated and non-constipated patients at the 
end of the study. Constipation was defined according to 
Thompson et al[17] criteria. Allocation was concealed using 
same color, size and weight closed boxes. The nurses 
that provided the patients with colonic preparation, the 
endoscopy assistant that evaluated the preparation compli-
cance, tolerance and adverse reactions, the data analysts; 
and the endoscopists who evaluated bowel cleansing 
quality were blinded. If the patients had doubts about 
the preparation they could make a telephone call to a 
physician that was not blinded, was not present during
Preparation of colonoscopy involved bowel cleansing with different colonic cleansing quality. Patients were asked not to reveal their assigned preparation to the endoscopists. A kappa coefficient of agreement was used for this purposes. A kappa under 0.3 and a non-significant P value was considered as an adequate blinding. 

Results of colonoscopy with different colonic cleansing quality. Patients were asked not to reveal their assigned preparation to the endoscopists. A kappa coefficient of agreement was used for this purposes. A kappa under 0.3 and a non-significant P value was considered as an adequate blinding.

### Table 1 Bowel preparation quality grading score used by the endoscopists

| Grade | Description | Score |
|-------|-------------|-------|
| Excellent | No fecal matter or nearly none in the colon, small-to-moderate amounts of clear liquids | 57 ± 11.1 |
| Good | Small amounts of thin liquid fecal matter seen and easily suctioned, mainly distal to splenic flexure, small lesions may be missed, > 90% mucosa seen | 41 ± 49 |
| Fair | Moderate amounts of thick liquid to semisolid fecal matter seen and suctioned, included proximal to splenic flexure, small lesions may be missed, 90% mucosa seen | 50 ± 49 |
| Poor | Large amounts of solid fecal matter found, precluding a satisfactory study, unacceptable preparation; < 90% mucosa seen | NS |

### Table 2 Characteristics of the included patients (n = 346)

| Characteristics | Prep 1 | Prep 2 | Prep 3 | Prep 4 | P value |
|-----------------|-------|-------|-------|-------|---------|
| Patients        | 78    | 78    | 84    | 84    | NS      |
| Age (yr), mean ± SD | 59 ± 13.2 | 57 ± 11.1 | 60 ± 13.8 | 59 ± 10.9 | NS      |
| Sex             |       |       |       |       | NS      |
| Male            | 37 (47) | 40 (51) | 41 (49) | 45 (53) | NS      |
| Female          | 41 (53) | 38 (49) | 43 (51) | 39 (47) | NS      |
| Constipation    | 21 (27) | 16 (21) | 15 (12) | 24 (29) | NS      |
| Successful cecal intubation | 78 (100) | 78 (100) | 84 (100) | 84 (100) | NS      |

NS: Not significant.

### Quality of colonic cleansing

We obtained information about this outcome for 93% of patients. Knowing that 70% of colonic cleansings are excellent or very good (Table 1), a sample size of 88 patients in each group was calculated to detect a 20% difference in primary outcome with 80% of power at a standard level of significance $\alpha = 0.05$. Categorical variables were compared using the Fisher exact test or $\chi^2$ test. A P value of less than 0.05 was considered significant. Results were analyzed according to the intention-to-treat principle. Handling of loss to follow-up: We evaluated different assumptions about the incidence of events among participants lost to follow-up and the impact of those assumptions on the estimate of effect for the primary outcome. For this purpose, we used the RI_LTFU/FU as proposed by Akl et al.[18]. The RI_LTFU/FU is defined as the event incidence among those lost to follow-up relative to the event incidence among those followed up. The assumptions we evaluated by combining a range of RI_LTFU/FU values (1, 1.5, 2, 3.5 and 5) in the intervention group and control group.

### RESULTS

A total of 349 patients scheduled for outpatient colonoscopy participated in the study and were randomized to receive one of the four colonic cleansing preparations. Three patients were excluded post-randomization because they met one or more exclusion criteria, 15 patients failed to present to the procedure and 7 presented incomplete colonoscopy because of fixed angulations (4 patients) or colonic neoplasia (3 patients). Finally, of the 346 randomized patients, information about the primary outcome was obtained from 324 patients (93%) (Figure 1). There were no significant differences among the four preparation groups with respect to: age, sex, cecal intubation, and constipation (Table 2).

### Blinding challenge

There was no significant concordance between the endoscopists presumption and the colonic preparation group that the patients had been assigned to ($P = 0.56$, $\kappa = 0.019$). This observation reassures that the endoscopists were unaware of the assigned groups (blinding).

### Statistical analysis

Statistical analysis were performed using statistical software SPSS for windows 10.0. Knowing that 70% of colonic cleansings are excellent or very good (Table 1), a sample size of 88 patients in each group was calculated to detect a 20% difference in primary outcome with 80% of power at a standard level of significance $\alpha = 0.05$. Categorical variables were compared using the Fisher exact test or $\chi^2$ test. A P value of less than 0.05 was considered significant. Results were analyzed according to the intention-to-treat principle. Handling of loss to follow-up: We evaluated different assumptions about the incidence of events among participants lost to follow-up and the impact of those assumptions on the estimate of effect for the primary outcome. For this purpose, we used the RI_LTFU/FU as proposed by Akl et al.[18]. The RI_LTFU/FU is defined as the event incidence among those lost to follow-up relative to the event incidence among those followed up. The assumptions we evaluated by combining a range of RI_LTFU/FU values (1, 1.5, 2, 3.5 and 5) in the intervention group and control group.

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patients. The quality of colonoscopic visualization was similar in the four different groups (Figure 2A).

Results were dichotomized into satisfactory colonic cleansing (excellent and good) and unsatisfactory (fair and poor). Satisfactory preparations were achieved in similar proportion in the different groups: prep 1, 82%, prep 2, 80%, prep 3, 79% and prep 4, 78% (P > 0.05) (Figure 2B). Endoscopists thought that only 6% of all the patients in this study needed to repeat the study because of inadequate colonic preparation. This was also similar between different preparations: prep 1, 3.4%, prep 2, 4.7%, prep 3, 6.8% and prep 4, 6.8% (P > 0.05) (Figure 2C).

We conducted a separate analysis of constipated and non-constipated patients. In the non-constipated patients, we didn’t find differences in the quality of colonic cleansing (Figure 2B) but the necessity to repeat colonoscopy was more frequent in prep 4 compared to prep 1 (11% vs 2%, P < 0.05) (Figure 2C). In constipated patients, NaP plus bisacodyl preparation (prep 2) achieved higher rate of satisfactory colonic cleansing than those receiving whole dose of PEG (prep 3): 95% vs 66% (P = 0.03) (Figure 2B).

Compliance
Both preparations containing NaP, presented better compliance than those containing PEG. Preparation was completed by 94% of the patients in prep 1, 100% of patients in prep 2, 81% of patients in prep 3 and 87% of the patients in prep 4. Therefore, half-dose of NaP plus bisacodyl achieved the highest compliance (prep 2 vs 1, 3 and 4, P < 0.01) followed by full-dose of NaP (prep 1 vs 3 and 4, P < 0.05) (Figure 2D). In non-constipated patients, compliance was also higher in those preparations containing NaP compared to full-dose PEG: 95% (prep 1), 100% (prep 2) vs 80% (prep 3) (P < 0.05) (Figure 2D). In constipated patients compliance was similar between different preparations.

Tolerability
The preparation was reported as tolerable in 77% of the patients in prep 1, 81% in prep 2, 82% in prep 3 and in 84% in the prep 4, there was no significant difference between the different preparations (P > 0.05). There was also no significant difference in tolerability between preparations in constipated and non-constipated patients (Table 3).

Symptoms profile
The most frequent adverse effects reported were: nausea (33%), bloating (30%) and abdominal pain (23%). There were no significant differences among different groups with respect to: nausea, vomiting, chest pain, bloating and dizziness (Table 3). Abdominal pain was more frequent in patients that received both preparations containing bisacodyl, prep 1, 16%, prep 2, 27%, prep 3, 19%, prep 4, 28%, but this difference didn’t reach statistical significance in the overall analysis (P = 0.2) (Table 3). The patients receiving NaP and bisacodyl preparations (prep 2) presented more frequently poor sleep than the other groups (P < 0.05) (Table 3). In non-constipated patients, abdominal pain was more frequent in those preparations containing bisacodyl: prep 2 (31%) and prep 4 (29%); compared to those without it: prep 1 (14%) and prep 3 (20%) (P < 0.05) (Table 3). The symptoms profile was similar between different preparations in constipated patients.

Patient preferences
Only 21% of all the patients would refuse to take the same colonic preparation in the future and almost 37% would like to try a different preparation. This finding was similar in the different groups. There was also no significant differences in patients perception in different groups...
in constipated and non-constipated patients.

**Loss to follow up**

None of the different assumptions of incidence of events in loss to follow up patients changed significantly the estimate of the effect in the different outcomes.

**DISCUSSION**

There is a growing acceptance of colorectal cancer screening with colonoscopy. Its goal is to identify and remove neoplastic polyps; therefore a high-quality preparation that lends to a clear visualization is crucial. Inadequate colonic cleansing could lead to a diminished adenoma detection rate[19-21]. This has been recently shown to be the strongest predictor of interval colorectal cancer[22,23]. However none of the different preparation agents are ideal for colonic cleansing. They present historic rates for adequate cleansing that ranges from 70% to 82%[24-26]. Tolerability and side effects are probably the main issues and represent some of the most important reasons for patient’s refusal to the study[25]. In an attempt

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**Table 3 Symptoms profile of different preparations a (%)**

| Adverse effects | Overall | Non-constipated (n = 59) | Constipated (n = 59) | Overall | Non-constipated (n = 58) | Constipated (n = 20) | Overall | Non-constipated (n = 69) | Constipated (n = 15) | Overall | Non-constipated (n = 65) | Constipated (n = 20) |
|-----------------|---------|-------------------------|---------------------|---------|-------------------------|---------------------|---------|-------------------------|---------------------|---------|-------------------------|---------------------|
| Tolerability    | 62 (77) | 47 (60)                 | 15 (68)             | 63 (81) | 47 (81)                 | 16 (86)             | 69 (82) | 57 (83)                 | 12 (80)             | 71 (84) | 53 (82)                 | 18 (80)             |
| Nausea          | 27 (33) | 17 (29)                 | 10 (13)             | 30 (58) | 24 (41)                 | 6 (30)              | 26 (31) | 2 (32)                  | 4 (26)              | 25 (29) | 18 (28)                 | 7 (35)              |
| Vomiting        | 6 (7)   | 2 (3)                   | 4 (50)              | 3 (4)   | 6 (10)                  | 3 (15)             | 9 (11)  | 2 (3)                   | 1 (7)               | 6 (7)   | 5 (8)                   | 3 (15)              |
| Abdominal pain  | 13 (16) | 8 (14)                  | 5 (25)              | 21 (27) | 18 (33)                 | 3 (15)             | 16 (19) | 14 (20)                 | 2 (3)               | 13 (24) | 19 (29)                 | 5 (30)              |
| Bleeding        | 25 (31) | 17 (29)                 | 8 (36)              | 21 (27) | 15 (26)                 | 6 (30)             | 27 (32) | 22 (32)                 | 5 (33)             | 24 (28) | 16 (25)                 | 8 (40)              |
| Insomnia        | 10 (12) | 9 (15)                  | 1 (5)               | 17 (21) | 14 (24)                 | 5 (15)             | 5 (6)   | 4 (6)                   | 1 (7)               | 11 (13) | 10 (15)                 | 1 (5)               |
| Dizziness       | 12 (15) | 10 (17)                 | 2 (9)               | 7 (9)   | 6 (10)                  | 1 (5)              | 7 (8.5) | 5 (7)                   | 2 (13)             | 8 (9)   | 5 (8)                   | 3 (15)              |
| Chest pain      | 1 (1)   | 1 (2)                   | 0 (0)               | 2 (3)   | 1 (2)                   | 1 (5)              | 1 (1)   | 1 (1)                   | 0 (0)               | 1 (1)   | 0 (0)                   | 1 (5)               |

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A: Preparation quality score obtained with different preparations (no statistical difference between groups). Values are expressed as the percentage of patients. Prep 1, 90 mL of sodium phosphate (NaP); Prep 2, 45 mL of NaP followed by 20 mg of bisacodyl; Prep 3, 4 L of polyethylenglicol (PEG); Prep 4, 2 L of PEG followed by 20 mg of bisacodyl. B: Percentage of patients who had satisfactory and unsatisfactory colonic cleansing in the overall analysis and in the subgroup of constipated and non-constipated patients. Constipated patients obtained a higher rate of satisfactory colonic cleansing with prep 2 (45 mL of NaP followed by 20 mg of bisacodyl) when compared to preparation 3 (4 L of PEG) (prep 2 vs 3, P = 0.03); C: Endoscopist appraisal on the necessity to repeat colonoscopy due to inadequate preparation in the overall analysis and in the subgroup of constipated and non-constipated patients. Non-constipated patients assigned to prep 2 (45 mL of NaP followed by 20 mg of bisacodyl) needed to repeat colonoscopy due to inadequate preparation more often when compared to patients assigned to prep 1 (90 mL NaP) (prep 4 vs 1, P = 0.05); D: Compliance to different preparations in the overall analysis and in the subgroup of constipated and non-constipated patients. Prep 2 (45 mL of NaP followed by 20 mg of bisacodyl) vs 1 (90 mL NaP), 3 (4 L of PEG) and 4 (2 L PEG followed by 20 mg of bisacodyl), P < 0.05; prep 1 (90 mL NaP) vs 3 (4 L of PEG) and 4 (2 L PEG followed by 20 mg of bisacodyl), P < 0.05; prep 1 (90 mL NaP) and 2 (45 mL of NaP followed by 20 mg of bisacodyl) vs prep 3 (4 L of PEG), P < 0.05.

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[^1]: Prep 2 and 4 vs prep 1 and 3; ^2: prep 2 vs prep 1, 3 and 4. NS: Not significant.
to decrease these side effects, many studies have evaluated different doses of conventional preparation agents and pretreatment with prokinetics or laxative agents, but there is little information about the effect of these preparations in subgroups of constipated and non-constipated patients\[6\]. In this study we compared two of the most used colonic cleansing agents, PEG and NaP. As in past years, there has been a strong tendency to prepare patients with half doses of this previously mentioned agents associated with bisacodyl because of commercially available preparation kits. We decided to carry out a direct comparison between whole dose of PEG and NaP alone and half doses of these two agents associated with bisacodyl in constipated and non-constipated patients. Our studies main limitations include, single centre study and the use of non-validated scale for the evaluation of primary outcome (quality of colonic preparation) and patient related outcomes (tolerability, adverse events, preferences). Nevertheless, the randomized, double-blind, four-arm study design and the constipated and non-constipated subgroup analysis could provide useful information on how to manage patients that might undergo colonoscopy. Similar to the results reported by previous studies, almost 80% of patients presented to colonoscopy with satisfactory colonic cleansing (excellent or very good). We did not find any difference with respect to quality of colonic cleansing in the different groups, even in those with half doses of NaP and PEG. Preparation quality was also similar in different groups in non-constipated patients, but endoscopists thought that there was a greater necessity to repeat the study due to an inadequate colonic cleansing in prep 4 (half dose of PEG plus bisacodyl) compared to prep 1 (whole dose of NaP) (11% vs 2%, \(P < 0.05\)). Although this is a non-validated and subjective outcome; we think it’s interesting to know endoscopist perception, because it represents what they really do in the daily practice and is a patient important outcome. In constipated patients, preparations containing bisacodyl presented higher rates of satisfactory colonic cleansing: 95% (prep 2) and 85% (prep 4) vs 67% (prep 3) and 77% (prep 1). Only NaP plus bisacodyl reached a statistically significant difference compared to whole dose of PEG (95% vs 66%, \(P = 0.03\)). The prokinetic effect of the bisacodyl may explain the high rates of satisfactory colonic preparations. Even though a statistical significant difference was only obtained with NaP plus bisacodyl and not with PEG plus bisacodyl, we think that this may be related to the small sample size of the constipated patients subgroup. In the overall analysis, compliance was higher in groups with preparations containing NaP, reaching 100% in the half dose NaP plus bisacodyl group and 94% in the half dose of NaP in the overall analysis. Compliance was higher in groups with preparations containing NaP, reaching 100% in the half dose NaP plus bisacodyl group and 94% in the whole dose of NaP. In non-constipated patients, compliance with NaP preparations was higher than whole doses PEG preparation. We were not able to demonstrate higher compliance rates with NaP preparations in constipated patients. However, the observed tendency to higher compliance in these groups along with evidence of previous studies lead us to believe that we were unable to find statistically significant difference due to the small sample size. Tolerability (taste, nausea, etc) was similar in the different groups. Consequently, we believe that the differences in compliances were related to the volume of the preparations and probably not to tolerability. The most frequent adverse effect was nausea followed by bloating and abdominal pain. None of the different preparations were associated with an antimetic medication, so we do not know if nausea and probably tolerance could be optimized with this association. Bisacodyl has been previously associated with abdominal cramping. In this study both groups with preparations containing bisacodyl presented higher incidence of abdominal pain: prep 1, 16%, prep 2, 27%, prep 3, 19%, prep 4, 28%, but the difference was not statistically significant. The difference was statistically significant when we analyzed the subgroup of non-constipated patients: prep 1, 14%, prep 2, 31%, prep 3, 21%, prep 4, 29% \(P < 0.05\). Curiously, constipated patients that received preparation with bisacodyl did not have higher incidence of abdominal pain. We think that constipated patients can present a motility dysfunction that could be optimized with the administration of the bisacodyl and that could explain the difference perception of abdominal pain in constipated and non-constipated patients. In the overall analysis, the combination of NaP with bisacodyl was also associated with higher rates of poor sleep than other preparations. We did not find any previous reports of this association and we do not have a specific explanation for this finding. However, it seems that the bisacodyl adverse effects profile is different in constipated and non-constipated patients, suggesting that constipated patients are less affected by these effects. Although the evaluated preparations presented a high rate of satisfactory colonic cleansing, compliance and a low profile of side effects, almost 37% of all the patients when asked, would prefer to try a different preparation in next colonoscopy. This study shows that none of the preparations agents is ideal, and highlights the need to improve bowel cleansing methods not only to get high quality colonic cleansing, but also to achieve a higher adherence to colonoscopy screening and surveillance programs. In summary, the quality of colonic cleansing and side effects profile of evaluated preparations are different in constipated and none-constipated. In non-constipated patients, preparation quality is similar with whole or half doses of NaP or PEG, alone or in combination with bisacodyl and compliance is higher with NaP preparations. Bisacodyl addition is associated with a higher incidence of adverse events. In constipated patients, the combination of NaP with bisacodyl is the most effective preparation. In this subgroup of patients, bisacodyl addition is not associated with higher incidence of adverse effects as noticed in non-constipated patients.

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COMMENTS

Background
Colonoscopy has become the standard procedure for the diagnosis and treatment of colon diseases. Adequate bowel cleansing is essential for a high-quality effective and safe colonoscopy.

Research frontiers
Numerous clinical trials have assessed the efficacy of whole or low doses of sodium phosphate (NaP) and polyethylene glycol (PEG) alone or with bisacodyl. There is no information about which is the most suitable preparation regimen for constipated and noneconstipated patients.

Innovations and breakthroughs
Their randomized clinical trial compared the efficacy and tolerability of whole and half doses of NaP and PEG alone or associated with bisacodyl preparations and explored the different effect on constipated and nonconstipated patients.

Applications
Compliance was higher with NaP preparations in nonconstipated patients and the addition of bisacodyl was associated with higher incidence of adverse effects. Half-dose of NaP plus bisacodyl was the most effective preparation in constipated patients. Bisacodyl was associated with adverse effects in constipated patients as noticed in nonconstipated patients.

Peer review
This is a good study in which authors compare the efficacy of different doses of NaP and PEG alone or with bisacodyl for colonoscopy in constipated and nonconstipated patients. The results are interesting and suggest that in nonconstipated patients, compliance is higher with NaP preparations, and bisacodyl is related to adverse effects. In constipated patients NaP plus bisacodyl is the most effective preparation.

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