BOWEL PREPARATION FOR COLONOSCOPY: COMPARISON OF MANNITOL AND SODIUM PHOSPHATE. RESULTS OF A PROSPECTIVE RANDOMIZED STUDY

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SUMMARY: Method: Eighty patients were prospectively randomized for precolonoscopic cleansing either with 750 ml of 10% mannitol (Group M) or 180 ml of a sodium phosphate preparation (Group NaP). Laboratory examinations before and after preparation on all patients included hemoglobin, hematocrit, sodium, potassium, phosphorous, calcium and serum osmolarity. A questionnaire was used to assess undesirable side effects and patient tolerance to the solution. The quality of preparation was assessed by the endoscopist who was unaware of the solution employed.

Results: Statistically significant changes were verified in serum sodium, phosphorous, potassium and calcium between the two groups, but no clinical symptoms were observed. There were no significant differences in the frequency of side effects studied. Six of the eight patients in Group NaP who had taken mannitol for a previous colonoscopy claimed better acceptance of the sodium phosphate solution. The endoscopic-blinded trial reported excellent or good bowel preparation in 85% prepared with sodium phosphate versus 82.5% for mannitol (p=0.37).

Conclusions: Quality of preparation and frequency of side effects was similar in the two solutions. The smaller volume of sodium phosphate necessary for preparation seems to be related to its favorable acceptance. Nevertheless, the retention of sodium and phosphate ions contraindicates the use of sodium phosphate in patients with renal failure, cirrhosis, ascites, and heart failure.


PATIENTS AND METHOD

The study was prospective and included 80 patients undergoing elective colonoscopy in the Colonoscopy Unit of the Coloproctology Division of Hospital das Clinicas, University of São Paulo during the period from April to July 1998. Only patients with a diagnosis of renal failure, liver disease with ascites, and severe congestive heart failure, and those undergoing emergency colonoscopies were excluded from this investigation. After an explanation about the study, all patients signed a consent agreement.

The patients were randomized by simple heads-or-tails allotment and
were given 10% mannitol (Group M) or sodium phosphate (Group NaP) for bowel preparation. Both solutions are currently produced at our pharmacy. Patients were instructed to follow a liquid diet without residues, starting after lunch on the day prior to the examination. Subjects in Group M were also instructed to take four tablets of bisacodyl (Dulcolax®, Boehringer De Angeli, São Paulo, SP) on the night prior to the exam. They were admitted on the morning of the colonoscopy and received two ml of metoclopramide intramuscularly prior to drinking 750 ml of 10% mannitol21. The sodium phosphate solution (monobasic sodium phosphate: 0.24 g/ml and dibasic sodium phosphate: 0.09 g/ml) was administered in two equal doses of 90 ml. The first dose on the afternoon prior to the examination (16:00h) and the second on the morning of the colonoscopy (07:00h), for a total of 180 ml. The colonoscopies were performed in the late morning.

The following data were recorded: sex, age, and variation of body weight (patients were weighed immediately before bowel prep and right after examination). Laboratory tests (hemoglobin level and hematocrit, serum Na, serum K, serum P, serum Ca, and serum osmolarity) were performed in all, immediately before and after bowel cleansing. Upon completion of bowel preparation and before colonoscopy, patients filled out a questionnaire with the objective of identifying side effects (nausea, vomiting, abdominal pain, abdominal distension, dizziness, perianal irritation) and degree of acceptance of the solution given.

Bowel preparation was evaluated by the endoscopist in a blinded-trial and graded as:
• **Excellent** – presence of clear fluid, without any fecal material
• **Good** – presence of cloudy fluid, but without residues.
• **Poor** – presence of residues.

The endoscopic diagnosis and complications were also recorded. Statistical analysis was based on Fisher and chi-square tests, with values of $p < 0.05$ considered statistically significant.

Forty of the 80 patients in the study received 10% mannitol for bowel preparation and the other 40 received sodium phosphate. Table 1 displays the data of the two groups. No statistically significant difference was noted regarding gender or age between the groups.

**RESULTS**

The biochemical analysis demonstrated a significant rise in serum sodium and phosphorous and a marked drop in potassium and calcium in
Group NaP compared to Group M (p = 0.0001). Nevertheless, there were no clinical symptoms. A slight increase occurred in hemoglobin (p=0.16), hematocrit (p=0.36), and serum osmolarity (p=0.74) in both groups, but these were without statistical significance. Weight loss (kg) in both groups was similar, 0.80 ± 1.4 for the Group NaP and 1.1 ± 1.2 for the Group M, without a statistically significant difference between them (p=0.42). Figure 1 illustrates weight variations and laboratory parameters between the two groups.

Bowel preparation was judged excellent or good in 85% of cases prepared with sodium phosphate and in 82.5% prepared with mannitol (p=0.37). A poor preparation was obtained in 15% and 17.5% in Group NaP and Group M respectively. In three patients who had been prepared with mannitol, it was impossible to complete the colonoscopy because of poor bowel cleansing. Table 2 displays patient distribution based on quality of bowel preparation and solution utilized.

Nausea and vomiting were the most frequent undesirable effects reported by patients of both groups. They occurred in 50% and 18% of those taking sodium phosphate and in 30% and 18% in those taking mannitol, respectively. Eight patients receiving sodium phosphate had been prepared for a previous colonoscopy with mannitol, and six of them claimed better acceptance of sodium phosphate. The Fisher test did not demonstrate a significant difference between the two groups for these variables (nausea, p = 0.11; vomiting, p = 1.00).

Distribution of endoscopic findings according to the type of the solution is displayed in Table 3. There was no statistically significant difference between the two groups (p > 0.01).

### DISCUSSION

The success of colonoscopy is directly related to the degree of bowel cleansing since it is based on perfect visualization of the colorectal mucosa. Thus, with improper cleansing, small lesions can be obscured by fecal residues in the lumen, impairing detection and treatment of various colonic diseases.

The ideal method for precolonscopy preparation should be effective, safe, easily administered, and well tolerated by the patient. Anterograde methods fulfill these requirements and have gradually replaced the conventional method, based on diet, laxatives, and enemas. Mannitol and polyethylene glycol (PEG) are currently the most commonly utilized oral solutions, each with inherent advantages and disadvantages1-6. Both solutions accomplish adequate colonic cleansing when properly administered.

The monosaccharide mannitol solution is almost non-absorbable in concentrations ranging from 5–20% and volumes of 500 to 750 ml. Its effect is due to the promotion of osmotic diarrhea with practically no absorption of water. Various studies have confirmed the effectiveness and safety of this solution for colonoscopic preparation and have achieved excellent or good results in over 90% of cases1,2. Because of the risk of explosion during electrocauterization resulting from fermentation of mannitol by hydrogen- and methane-producing intestinal bacteria, its use has been criticized and restricted in various centers in the United States and Europe3-5.

PEG preparations eliminate risk of explosion and are osmotically neutral, thus reducing the possibility of promot-
ing significant changes in fluids or electrolytes. For a proper cleansing, of a greater volume of PEG than mannitol must be ingested (about four liters). Drinking large volumes of liquids results in unacceptable bowel preparation in about 5–15% of patients, especially those with pharyngeal reflex problems, such as the elderly or those who have had strokes or neurological disturbances and children.

The present study was motivated by the good results obtained with the sodium phosphate solution for mechanical bowel preparation for elective colorectal surgery presented in the literature and by our own experience confirming better acceptance of sodium phosphate, especially because of the small volume utilized for colonoscopic bowel preparation. Some researchers have also demonstrated the advantages of sodium phosphate solution, especially comparing its use with the PEG solution.

In the present study, we compared the use of sodium phosphate and 10% mannitol, since the latter has been our method of choice for bowel preparation for colonoscopy for many years. We were not able to find a report of a similar investigation comparing sodium phosphate and mannitol, probably because of the preference for PEG in the majority of American and European medical centers.

The number of side effects was similar in the two groups, especially regarding nausea and vomiting. Despite more frequent bouts of nausea (50% vs. 30%) in the sodium phosphate group, the incidence of vomiting was identical in both groups (18%). Probably, the routine use of metoclopramide in patients receiving mannitol contributed to their lower incidence of nausea. In the literature, the incidence of nausea and vomiting resulting from sodium phosphate ranges from 11.2 to 44.3% and 5.4 to 8.6%, respectively. Frommer compared sodium phosphate and PEG in 486 patients submitted to colonoscopy and found a greater incidence of minor side effects, such as, nausea and vomiting in patients prepared with sodium phosphate. However, the majority of articles comparing the two solutions (PEG x NaP) mention a lower incidence of side effects with sodium phosphate.

The acceptance of ingesting one solution versus the other is difficult to compare, since acceptance is a subjective variable, perhaps reliably reported only by patients who have taken both solutions. In the present study, eight patients in the sodium phosphate group had been prepared with mannitol for previous examination, and six of them indicated better acceptance of sodium phosphate. The other two did not perceive a difference. Vanner et al. deemed sodium phosphate more easily acceptable than PEG in a prospective study including 102 patients, and 37 of them who had been previously prepared with PEG stated their preference for sodium phosphate. Other reports have demonstrated that there is no significant clinical difference in the incidence and intensity of side effects, but that swallowing a smaller volume of sodium phosphate solution compared to PEG is more easily accepted.

Hyperphosphatemia and hypocalcemia were the most common laboratory changes found in the sodium phosphate group, but there were no detectable clinical symptoms. Various articles have mentioned temporary hyperphosphatemia after the use of sodium phosphate. Although there are no reports of clinical problems due to increased serum levels of phosphate in normal adults, sodium phosphate should be avoided in patients with renal diseases. There was also greater retention of sodium ions, and greater loss of potassium in patients taking sodium phosphate, but without associated clinical symptoms in any of the cases. Nevertheless, this data obliges us to issue a warning restricting the use of sodium phosphate in patients with congestive heart failure or cirrhosis with ascites.

Concerning the quality of bowel preparation, both solutions obtained similar results, with excellent or good in 85% and 82.5% respectively for sodium phosphate and mannitol. Our data is similar to those obtained in other investigations studying the effectiveness of sodium phosphate for colonoscopy. Some researchers consider sodium phosphate more effective than PEG for bowel cleansing, justifying these results on the basis of the greater acceptance of sodium phosphate, which requires a markedly smaller volume than PEG. The examination was discontinued in three patients taking mannitol due to inadequate cleansing, because of vomiting in two of them.

The use of sodium phosphate has been associated with coarse macroscopic changes in the bowel mucosa. These changes range from friability and hyperemia to aphthoid injuries. However, we did not note these changes in any patient in the present study.

Based on the results of this series, we conclude that bowel preparation for elective colonoscopy may be equally effective and safe with either mannitol 10% or sodium phosphate solutions. However, due to the smaller volume necessary for adequate bowel cleansing, sodium phosphate seems to be better tolerated than mannitol. This is especially important in patients with difficulty in swallowing large volumes. Perhaps routine administration of an oral or parenteral antiemetic can further improve acceptance of sodium phosphate. However, retention of phosphate and sodium ions in patients with renal problems, liver diseases with ascites, and those with congestive heart failure makes the use of sodium phosphate solution inadvisable in patients with these complications.

Métodos: Oitenta pacientes foram prospectivamente randomizados para receber 750 ml de manitol a 10% (M) ou 180 ml de solução à base de fosfato de sódio (FS), como preparo intestinal para colonoscopia eletiva. Todos os pacientes foram submetidos a avaliação laboratorial (hemoglobina, hematócrito, sódio, potássio, fósforo, cálcio e osmolaridade sérica) antes e depois do preparo. Completado o preparo intestinal, antes da realização do exame, os pacientes foram avaliados por questãoário com a finalidade de identificar efeitos indesejáveis e tolerabilidade inerentes à solução empregada. A qualidade do preparo foi avaliada pelo colonoscopista, que desconhecia o tipo de solução empregada.

Resultados: A análise bioquímica demonstrou elevação significativa dos níveis séricos de sódio e fósforo no grupo do FS, bem como uma queda mais acentuada do pótassio e cálcio séricos neste grupo, mas nenhuma das alterações foi clinicamente sintomática. Não houve diferença significante na incidência de seis efeitos colaterais pesquisados. Seis de oito pacientes do grupo FS que em exame colonoscópico anterior haviam recebido manitol, manifestaram melhor tolerabilidade com a solução de FS. A qualidade do preparo foi considerada excelente ou boa em 85% dos casos preparados com FS e em 82,5% do grupo M (p=0,37).

Conclusão: As duas soluções foram similares quanto à qualidade do preparo e incidência de efeitos colaterais. O menor volume necessário para o preparo com FS parece estar relacionado com uma melhor tolerabilidade desta solução. No entanto, a retenção dos íons sódio e fosfato com o uso da solução de FS torna desaconselhável seu emprego em pacientes com insuficiência renal, cirrose e insuficiência cardíaca.


REFERENCES


Received for publication on the 26/04/99