The protective effect of supplemental calcium on colonic permeability depends on a calcium phosphate-induced increase in luminal buffering capacity

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Abstract
An increased intestinal permeability is associated with several diseases. Previously, we have shown that dietary Ca decreases colonic permeability in rats. This might be explained by a calcium-phosphate-induced increase in luminal buffering capacity, which protects against an acidic pH due to microbial fermentation. Therefore, we investigated whether dietary phosphate is a co-player in the effect of Ca on permeability. Rats were fed a humanised low-Ca diet, or a similar diet supplemented with Ca and containing either high, medium or low phosphate concentrations. Chromium-EDTA was added as an inert dietary intestinal permeability marker. After dietary adaptation, short-chain fructo-oligosaccharides (scFOS) were added to all diets to stimulate fermentation, acidify the colonic contents and induce an increase in permeability. Dietary Ca prevented the scFOS-induced increase in intestinal permeability in rats fed medium- and high-phosphate diets but not in those fed the low-phosphate diet. This was associated with higher faecal water cytotoxicity and higher caecal lactate levels in the latter group. Moreover, food intake and body weight during scFOS supplementation were adversely affected by the low-phosphate diet. Importantly, luminal buffering capacity was higher in rats fed the medium- and high-phosphate diets compared with those fed the low-phosphate diet. The protective effect of dietary Ca on intestinal permeability is impaired if dietary phosphate is low. This is associated with a calcium phosphate-induced increase in luminal buffering capacity. Dragging phosphate into the colon and thereby increasing the colonic phosphate concentration is at least part of the mechanism behind the protective effect of Ca on intestinal permeability.

Key words: Calcium: Phosphate: Short-chain fructo-oligosaccharides: Intestinal permeability

The intestinal tract is lined by a monolayer of epithelial cells, which is the largest and most important barrier between the body’s internal milieu and the hostile external environment. Tight contact between the epithelial cells prevents access of luminal digestive enzymes, toxins, antigens and enteric microbiota to underlying tissue compartments(1–3). Intestinal barrier dysfunction is suggested to be associated with the pathogenesis of a variety of intestinal diseases, including inflammatory bowel disease, coeliac disease, post-infectious irritable bowel syndrome and food allergy(1–5).

Interestingly, dietary components can influence the epithelial barrier, in particular intestinal permeability, and hence possibly modulate disease development. We are interested in the effect of dietary Ca, since it has been shown in several controlled studies that Ca is important for protection against intestinal infections with food-borne bacterial pathogens, both in rats and in human subjects(6–9). In addition, Ca displayed cytoprotective effects in several studies in the field of colon carcinogenesis by precipitating cytotoxic surfactants, such as secondary bile acids(10,11). Furthermore, supplemental Ca attenuated the development of colitis in HLA-B27 transgenic rats, which was associated with the prevention of a colitis-related increase in intestinal permeability due to Ca(12). In a previous study(13), we identified that the effect of dietary Ca on intestinal permeability is located in the colon. At present, however, it is still not clear how Ca exerts its effect on intestinal permeability, since it has been shown in several controlled studies that Ca is important for protection against intestinal infections with food-borne bacterial pathogens, both in rats and in human subjects(6–9). In addition, Ca displayed cytoprotective effects in several studies in the field of colon carcinogenesis by precipitating cytotoxic surfactants, such as secondary bile acids(10,11). Furthermore, supplemental Ca attenuated the development of colitis in HLA-B27 transgenic rats, which was associated with the prevention of a colitis-related increase in intestinal permeability due to Ca(12). In a previous study(13), we identified that the effect of dietary Ca on intestinal permeability is located in the colon. At present, however, it is still not clear how Ca exerts its effect.
protective effect on colonic permeability. Upon dietary intake, an insoluble calcium phosphate complex is formed in the small intestine, both in rats\(^{(14)}\) and in humans\(^{(15)}\). In this way, Ca prevents phosphate from being absorbed in the small intestine, and drags phosphate into the colon. In the colon, solubilisation of this complex increases the buffering capacity of the luminal contents, which protects the intestinal mucosa from being injured by an acidic pH due to microbial fermentation\(^{(16)}\). This mechanism might play a role in the effect of Ca on intestinal permeability, and it implies that dietary phosphate intake might be of importance for the effect of Ca on permeability.

In the present study, we investigated the potential effect of dietary phosphate on Ca-induced lowering of intestinal permeability, by applying high, medium or low phosphate concentrations in a high-Ca diet. Inert chromium-EDTA (CrEDTA) was added to the diets as an established marker for intestinal permeability\(^{(17,18)}\). After dietary adaptation for 10 d, short-chain fructo-oligosaccharides (scFOS) were introduced in the diets. The intestinal fermentation of scFOS leads to a luminal organic acid load, which challenges the buffering capacity. We hypothesise that intestinal permeability will increase on a high-Ca diet with concomitant low dietary phosphate levels because of a decreased buffering capacity.

Materials and methods

Experimental design: animals and diets

The experimental protocol was approved by the animal welfare committee of Wageningen University and Research Centre (Wageningen, The Netherlands). Specific pathogen-free outbred male Wistar rats (WU; Harlan, Horst, The Netherlands), 8 weeks old and with a mean body weight of 288 g at the start of the experiment, were housed individually in metabolic cages. Animals were kept in a temperature- and humidity-controlled environment in a 12 h light–12 h dark cycle. Rats (ten animals per dietary group) were fed a purified ‘humanised’ Western diet in restricted quantities (16 g/d). Restricted feeding was necessary to prevent scFOS-induced differences in food consumption as observed earlier\(^{(19)}\) and hence differences in vitamin, mineral and CrEDTA intake. Demineralised drinking-water was supplied ad libitum. The reference diet (low-Ca, medium-phosphate; LCaMP), which has been used as a reference diet in our previous experiments\(^{(12,13)}\), contained (per kg): 200 g acid casein, 326 g maize starch, 172 g glucose, 160 g palm oil, 40 g maize oil, 50 g cellulose, 2 g CrEDTA (see below) and 5·16 g CaHPO\(_4\),2H\(_2\)O (corresponding to 30 mmol Ca/kg diet; Sigma-Aldrich, St Louis, MO, USA). Vitamins and minerals (other than Ca) were added to the diets according to AIN-93\(^{(20)}\). The concentration of vitamins and minerals was increased by 20% to ensure adequate intake during restricted feeding. The diets had a high fat content to mimic the composition of a Western human diet. Dietary phosphate mainly originates from calcium phosphate and the protein source of the diet (casein: about 40 mmol phosphate/kg diet). The experimental diets were supplemented with 90 mmol CaHPO\(_4\),2H\(_2\)O/kg diet (high-Ca, high-phosphate; HCaHP), or with 90 mmol CaCl\(_2\),2H\(_2\)O/kg diet (high-Ca, medium-phosphate; HCaMP), at the expense of glucose. The high-Ca, low-phosphate diet (HCaLP) was supplemented with 90 mmol CaCl\(_2\),2H\(_2\)O/kg diet, and the casein of this diet was replaced by whey protein isolate as a low-phosphate protein source (BiPRO, about 5 mmol phosphate/kg diet; Davisco Foods International, Inc., Eden Prairie, MN, USA). Thus, the experiment consisted of four different diets: LCaMP diet (reference diet; 30 mmol Ca and 70 mmol phosphate/kg diet) and HCaLP diet (positive control since this diet has been used in our previous experiments, showing the effects of Ca on intestinal permeability\(^{(12,13)}\); 120 mmol Ca and 160 mmol phosphate/kg diet), HCaMP diet (120 mmol Ca and 70 mmol phosphate/kg diet) and HCaLP diet (120 mmol Ca and 35 mmol phosphate/kg diet). Inert CrEDTA was added to all diets to quantify intestinal permeability\(^{(18)}\). CrEDTA solution was prepared as described elsewhere and subsequently freeze-dried\(^{(21)}\). To check the complete formation and stability of the CrEDTA complex, the prepared CrEDTA solution was passed through a cation-exchange resin column (Chelex 100 Resin; Bio-Rad, Hercules, CA, USA). No uncomplexed Cr\(^{3+}\) ions were present. Rats were fed the experimental diets for 10 d, after which all diets were supplemented with 60 g scFOS/kg diet (6% w/w; Raffinose\(^{®}\) P95; Orafti, Tienen, Belgium) at the expense of glucose. Food intake was recorded daily and animal weight twice every week. At experimental day 20, rats were anaesthetised with isoflurane and killed. The caecum was excised and caecal contents were collected.

Measurement of intestinal permeability

Total 24 h urine samples were collected at experimental days 10 and 20. For the CrEDTA measurement, urine was acidified with 50 g TCA/l, centrifuged for 2 min at 14 000 g and the supernatant was subsequently diluted with 0·5 g CsCl/l. Then, chromium was analysed by inductively coupled plasma-atomic emission spectrophotometry.

Analyses of faeces and caecal contents

All faeces were collected during the last 3 d of experimental feeding without scFOS (pre-scFOS; experimental days 8, 9 and 10), and during the last 3 d of the scFOS supplementation period (experimental days 18, 19 and 20). Faeces and caecal contents were freeze-dried and subsequently ground to obtain homogeneous powdered samples. Ca and P were quantified in the faeces of the pre-scFOS period. To this end, faeces were treated with 50 g TCA/l, centrifuged for 2 min at 14 000 g, diluted with 0·5 g CsCl/l, and analysed by inductively coupled plasma-atomic emission spectrophotometry. Buffering capacity was also determined in the faeces of the pre-scFOS period. To this end, pools of the freeze-dried faeces (2 g) were reconstituted with double-distilled water to 15% dry weight. The quantities of hydrochloric acid required to decrease the pH to 5 in these samples were measured, as described earlier\(^{(16)}\). For the cytotoxicity assay, faecal water was prepared by reconstituting freeze-dried
facess with double-distilled water to 25 % dry weight as described previously[22]. The cytotoxicity of the faecal water of the scFOS supplementation period was determined by potassium release of a human erythrocyte suspension after incubation with faecal water, as described earlier[23], and validated earlier with intestinal epithelial cells[24]. Cytotoxicity was calculated and is expressed as a percentage of maximal lysis. Total lactic acid was determined in caecal contents using a colorimetric enzymatic kit (Enzyplus; BioControl Systems, Inc., Bellevue, WA, USA), as described earlier[25].

Statistical analysis

All results are expressed as mean values with their standard errors. The predefined comparisons of interest were the LCaMP diet v. the HCaHP diet to study the effect of Ca (these two diets have been compared in our previous studies, showing the effects of Ca on intestinal permeability[12,13]), and the HCaHP diet v. the HCaMP diet and the HCaMP diet v. the HCaLP diet to study the effects of phosphate. Statistics were done by using one-way ANOVA or Kruskal–Wallis test, depending on the normality of the data. If differences were significant, this was followed by Student’s t test (for normally distributed data) or Mann–Whitney U test (for non-normally distributed data) to identify the significant dietary effects. Differences were considered statistically significant when P<0·05 (all two-sided). Statistical analyses were conducted with GraphPad Prism version 5.01 (GraphPad Software, Inc., La Jolla, CA, USA).

Results

Animal growth and food intake

No diet-induced differences in animal growth and food intake were observed before supplementation with scFOS: all rats consumed the provided 16 g/d as intended. However, animal growth was affected by the different diets during scFOS supplementation. Rats from the HCaLP group gained less weight during supplementation with scFOS than those from the HCaMP group (Fig. 1(a), P=0·006). The other dietary groups did not differ with respect to growth. The results for food intake were similar: HCaLP-fed rats had a more decreased food consumption than those fed the HCaMP diet during scFOS supplementation (Fig. 1(b), P=0·02).

Faecal baseline characteristics before short-chain fructo-oligosaccharide supplementation

Daily output of faeces, based on dry weight, was increased due to Ca supplementation (Table 1; P<0·001), in accordance with previous work[7,14]. To check whether the dietary interventions indeed affected the baseline Ca and phosphate levels in the colonic lumen before scFOS supplementation, we measured total Ca and P in the faeces (Table 1). Indeed, Ca concentration was clearly higher in the Ca-supplemented group (HCAHP; P<0·001) compared with levels in the LCaMP group. P levels in the faeces also corresponded well with the dietary intervention. Faecal P was increased in rats

Table 1. Effect of diet on faecal baseline characteristics before short-chain fructo-oligosaccharide supplementation

<table>
<thead>
<tr>
<th></th>
<th>LCaMP</th>
<th>HCaHP</th>
<th>HCaMP</th>
<th>HCaLP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Faecal dry weight (g/d)</td>
<td>1·02±0·02</td>
<td>1·31±0·02</td>
<td>1·37±0·03</td>
<td>1·38±0·04</td>
</tr>
<tr>
<td>Ca excretion (μmol/d)</td>
<td>78±10</td>
<td>1051±34</td>
<td>1098±19</td>
<td>887±19</td>
</tr>
<tr>
<td>P excretion (μmol/d)</td>
<td>65±6</td>
<td>733±21</td>
<td>350±17</td>
<td>116±8</td>
</tr>
</tbody>
</table>

* Mean values were significantly different from those of rats on the LCaMP diet (P<0·0001).
† Mean values were significantly different from those of rats on the HCaMP diet (P<0·0001).
‡ Mean values were significantly different from those of rats on the HCaHP diet (P<0·0001).

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Dietary effects on caecal lactate levels

To investigate whether fermentation of scFOS is disturbed due to acid accumulation, lactate was quantified in caecal contents. Ca supplementation in a high-phosphate background (HCaHP) resulted in significantly lower lactate levels in caecal contents compared with rats fed the LCaMP diet (P=0.006; Fig. 4(a)). Lactate levels were similar in the HCaMP group compared with rats fed the HCaHP diet, while these levels were higher in rats on the HCaLP diet compared with the HCaMP group (P=0.01; Fig. 4(a)). Interestingly, lactate levels are correlated with the colonic permeability results (Spearman’s r 0.45, P=0.006). The dietary effects on caecal pH show comparable results, since the pH in the caecum of the HCaHP group was significantly higher than the pH of the caecal contents of LCaMP-fed rats (6.9 (SEM 0.2) v. 5.8 (SEM 0.1), respectively, P=0.0008). No differences in caecal pH were observed in the other dietary groups (HCaMP, 7.2 (SEM 0.2); HCaLP, 6.8 (SEM 0.3)).

Dietary calcium and phosphate influence luminal cytotoxicity

To investigate whether faecal water cytotoxicity, reflecting colonic mucosal exposure to luminal irritants, plays a role in the dietary effects on colonic permeability, the cytotoxicity assay was performed with faecal water from the period after scFOS supplementation. Cytotoxic activity of faecal water was lower in rats fed the HCaHP diet (P=0.006; Fig. 4(b)) compared with those fed on the LCaMP diet. In HCaLP-fed rats, luminal cytotoxicity was increased compared with rats fed the HCaMP diet (P=0.01; Fig. 4(b)).

The effect of dietary calcium and phosphate on intestinal permeability

The main outcome of the present study is the dietary effect on intestinal permeability, measured by urinary CrEDTA excretion. Dietary Ca already decreased intestinal permeability before the colon was challenged with scFOS; however, this was not the case in rats fed the HCaLP diet. Urinary CrEDTA expressed as a percentage of dietary intake was 5.9 (SEM 0.5) in the LCaMP group, 4.2 (SEM 0.3) in the HCaHP group (P=0.0007, compared with LCaMP), 4.1 (SEM 0.2) in the HCaMP group and 6.5 (SEM 0.5) in the HCaLP group (P=0.0002, compared with HCaMP). Importantly, after scFOS supplementation, dietary Ca only prevented the fermentation-induced increase in intestinal permeability when rats were fed a diet containing high or medium levels of phosphate (HCaHP, P=0.049 compared with LCaMP; Fig. 3), but not when rats were fed the low-phosphate diet (P=0.04 compared with HCaMP; Fig. 3).

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associated with a decrease in luminal buffering capacity. If dietary phosphate levels were low, and this effect was observed with luminal acidification through scFOS in the diet. Indeed, the protective effect of Ca, in a situation of enhanced buffering capacity affects intestinal permeability, and is part of the protective effect of a high-Ca diet on intestinal permeability. It becomes even more important. Interestingly, scFOS induce an increase in colonic permeability, low phosphate intake also adversely affected food intake and animal weight after scFOS supplementation.

The protective effect of dietary Ca on intestinal permeability has been shown consistently in several studies using different animal models. It is of relevance that Ca can improve intestinal permeability both in a healthy situation and during intestinal infection and colitis development. In the latter two studies, the prevention of an increase in intestinal permeability by Ca was associated with an enhanced resistance to the development of intestinal disease. The present study emphasises the potential of Ca to decrease intestinal permeability in a healthy condition; however, it also shows that dietary phosphate intake has to be taken into account.

Since it is still unknown how Ca exerts its beneficial effect on gut permeability, we aimed to investigate whether a calcium-phosphate-induced increase in luminal buffering capacity is involved. By acidifying the colon with scFOS in the present study, the capacity to buffer the intestinal contents becomes even more important. Interestingly, scFOS induce an increase in intestinal permeability, but it is not proven that this is associated with luminal acidification caused by scFOS, although this seems likely. There are indications from in vitro studies that the extracellular pH is important for paracellular permeability. We aimed to show experimental support for the hypothesised effect that the luminal buffering capacity affects intestinal permeability, and is part of the protective effect of Ca, in a situation of enhanced luminal acidification through scFOS in the diet. Indeed, the protective effect of Ca on colonic permeability was impaired if dietary phosphate levels were low, and this effect was associated with a decrease in luminal buffering capacity.

Fermentation already starts in the caecum of rats; however, as has been described earlier, it clearly continues in the rat colon, showing the relevance of faecal sampling. Therefore, the similarity of the effects of scFOS in both the caecum and colon of rats and human colonic fermentation supports the use of the rat as an appropriate model to study possible subsequent permeability alterations. Moreover, intestinal pH values and faecal phosphate levels are similar in rats and humans, indicating that diet-modulated buffering capacity in the colon is comparable between these species. These results show that the protective effect of dietary Ca is at least partly due to its role as a carrier of phosphate into the colon. Subsequently, phosphate is responsible for the increase in colonic buffering capacity.

Ca supplementation in combination with high or medium dietary phosphate prevented a fructo-oligosaccharide-induced accumulation of lactate in the caecal contents, which was associated with the decrease in intestinal permeability. During rapid fermentation of easily fermentable carbohydrates, lactate accumulation can occur when micro-organisms that utilise lactate are inhibited. These micro-organisms might be inhibited by an acidic pH due to a compromised buffering capacity. It is therefore likely that alterations in microbiota composition or activity are responsible for the changes in caecal lactate levels. Interestingly, intestinal micro-organisms, for example Escherichia coli, are able to induce an increase in intestinal permeability. Therefore, buffering capacity might also indirectly influence intestinal permeability by modulating the gut microbiota.

An alternative explanation for the dietary effects on intestinal permeability, besides modulation of luminal buffering capacity, might be the influence of supplemental Ca on luminal cytotoxicity. The present study demonstrates that a lack of phosphate counteracts the inhibiting effect of Ca on cytotoxicity. Clearly, both Ca and phosphate are needed to precipitate cytotoxic components, which can irritate the intestinal epithelium and subsequently modify permeability. Modulation of luminal cytotoxicity might also change the microbiota, which can thereby exert an influence on epithelial integrity. We have shown earlier that Ca affects the gut microbiota. The results of the present study suggest that phosphate is necessary for the effect of Ca, but not...
determinative, i.e. as long as phosphate intake is sufficient (≥ 70 mmol/kg diet), the effect of Ca is not dependent on phosphate.

Impairment of the gut mucosal barrier by scFOS has been shown earlier in association with a decreased resistance to intestinal infection (39,28,38). These harmful effects were decreased when Ca was supplemented to the diet (31). The present study emphasises again that it is important to take care of a sufficient Ca intake when non-digestible carbohydrates are consumed. Furthermore, the present results show that dietary phosphate levels should also be taken into consideration. The adverse effects of low phosphate intake on food consumption and animal weight, which emerged during scFOS supplementation, support this observation.

The Ca and phosphate content of the rat diets in the present study is nutritionally relevant for the human diet. In general, human dietary Ca intake in the Western world ranges from 600 to 1100 mg daily (39). The Ca concentration of the low-Ca diet corresponds to a daily Ca intake of 600 mg in humans, while the Ca-supplemented diets provided more than the general habitual dietary Ca intake (comparable to 2-4 g daily), which is not unrealistic when taking Ca supplements (32). In addition, the animal diets contained phosphate levels of approximately 160 mmol/kg diet (high-phosphate), 70 mmol/kg diet (medium-phosphate) and 35 mmol/kg diet (low-phosphate), which are in the range of human intake of about 40 mmol phosphate daily (80 mmol/kg diet), assuming that humans have a daily dry food intake of about 500 g (39).

In conclusion, the present study shows that the protective effect of dietary Ca on intestinal permeability is impaired if phosphate levels are low, and this is associated with the effects on luminal buffering capacity. This is particularly important when consuming a diet with non-digestible and rapidly fermentable carbohydrates such as scFOS. So, phosphate levels should be sufficient in the diet to accomplish the protective effect of Ca on intestinal permeability. It cannot be excluded that changes in luminal buffering capacity also indirectly modulate intestinal permeability, for example, by the effects on the intestinal microbiota. The present study encourages follow-up studies, particularly in human subjects, to further explore and apply the potential to modulate gut barrier integrity by relatively simple dietary interventions.

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