Fructo-oligosaccharides and iron bioavailability in anaemic rats: the effects on iron species distribution, ferroportin-1 expression, crypt bifurcation and crypt cell proliferation in the caecum

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Abstract

The present study investigated the effects of fructo-oligosaccharides (FOS) on the bioavailability of Fe from ferric pyrophosphate (FP), a water-insoluble compound, in Fe-deficient anaemic rats that were subjected to a Hb repletion assay. Male Wistar rats (n = 64) were fed adequate or low (8 mg/kg) Fe diets for 15 d followed by 1 or 2 weeks of Fe repletion with diets providing 35 mg Fe/kg as ferrous sulphate (FS), FP or FP that was mixed with 7.5% FOS in the form of yacon flour or Raftilose P95 (RAF), a purified source of FOS. The effects of FOS were observed within the 1st week of the repletion period. Fe bioavailability was improved by FOS supplementation, as measured by Hb regeneration efficiency and hepatic Fe stores, which were more pronounced in the RAF group. Moreover, RAF supplementation resulted in a higher biological value relative to that of the FP group. FOS supplementation resulted in caecal enlargement, in addition to acidification and Fe species redistribution in the caecal contents relative to the control rats. These effects occurred concomitantly with decreased ferroportin (FPN)-1 expression in the caecal mucosa, which was similar in magnitude to that observed in the FS group. Caecum mucosal morphometry was influenced by FOS supplementation, whereas crypt fission and cell proliferation were highest in the caecum of the RAF group. These results reinforce the effects of FOS as Fe bioavailability enhancers in anaemic rats that are sustained by early changes in their caecal environment (decreased mucosal FPN-1 expression and increased Fe absorbability, crypt fission and cellularity).

Key words: Fructo-oligosaccharides: Iron: Bioavailability: Caecum: Rats

The type and amount of dietary fibre that reaches the large intestine and the extent to which it is fermented by intestinal bacteria (i.e. its fermentability) are important factors for understanding its overall effects. In general, the bacterial fermentation of non-digestible carbohydrates affects the following: (1) the luminal environment, by influencing the composition and/or activity of the gut microbiota and the physico-chemical properties of the intestinal content(1,2), and (2) the intestinal epithelium, by regulating the gut hormone response and/or altering the intestinal barrier function, including the stimulation of mucus production and the regulation of cell proliferation, migration and survival, cellular permeability, and the mucosal immune system(3–6). However, other factors could also modulate the fermentative process including the digestibility of food matrix components (such as resistant proteins, lipids and some micronutrients), intestinal motility and inter-individual differences in gut microbiota composition(5,7–9).

In recent years, several studies have also emphasised the effects of inulin-type fructans (ITF), such as fructo-oligosaccharides (FOS) and inulin, on the intestinal absorption of minerals. The acidification of intestinal contents from SCFA production by bacterial ITF fermentation increases mineral solubility and absorbability(10,11). Furthermore, dietary ITF seem to affect mineral absorption in the large intestine by altering the permeability of tight junctions and the gene expression of mineral transporters such as calbindin-D 9k (for Ca), divalent metal transporter (DMT-1) (for Fe) and zinc transporter 1 (ZnT1) (for Zn), supporting an effect on both paracellular and transcellular absorption(12–14). In addition, changes in the mucosal architecture of the intestine as a result of increases in both the cellularity and number of

Abbreviations: CT, control; FOS, fructo-oligosaccharides; FP, ferric pyrophosphate; FPN, ferroportin; FS, ferrous sulphate; HRE, Hb regeneration efficiency; ID, Fe deficient; ITF, inulin-type fructans; RAF, Raftilose P95; RBV, relative biological value; YF, yacon flour.

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crypts are factors that could contribute to an increase in the absorptive surface for mineral uptake(3,4,7,15). These effects are reflected in the improved bioavailability of minerals, as demonstrated by the increased retention in their target tissues(7,12–18).

We have previously demonstrated the positive effects of short-chain ITF (FOS), a rapidly fermentable fibre, on the bioavailability of Ca(7,15–17), Mg(7,15,16) and Fe(17,18) in rats fed diets containing purified and non-purified ITF sources. In the present study, we used a Hb depletion–repletion method resembling the one proposed by the AOAC(19) to investigate the effects of FOS on Fe bioavailability in anaemic rats. The effects of FOS as Fe bioavailability enhancers were confirmed and such effects were sustained by early changes in the caecal environment (decreased mucosal ferroportin (FPN)-1 expression and increased Fe absorbability and crypt bifurcation and cellularity).

**Experimental methods**

The experimental protocol was approved by the Commission on Ethics in Animal Experiments of the Faculty of Pharmaceutical Sciences at the University of São Paulo (FCF/USP) (CEEA 213/2009 FCF-USP) according to the guidelines of the Brazilian College on Animal Experimentation.

**Iron deficiency protocol**

Female Wistar rats (n 12) were obtained from the colonies for Animal Experimentation at FCF/USP, and each of them was breast-feeding six to eight male pups. These rats were housed in plastic cages with ripcurl and given an Fe-deficient powder diet(19) (8 mg Fe/kg; n 10 female rats) or an AIN-93M diet(20) (2 female rats) for 21 d.

On weaning day, a total of eighty-eight male rats initially weighing 54–58 g were transferred to individual stainless-steel wire-mesh metabolism cages (to limit coprophagy) weighing 54–58 g were transferred to individual stainless-steel wire-mesh metabolism cages (to limit coprophagy) and an Fe-deficient powder diet (ID group; n 80) or an AIN-93G diet(20) (control (CT) group; n 8) for 14 d (depletion period). During this period, ten ID rats were selected to determine their body weight and Hb concentration (g/l). Rats were fed modified AIN-93G diet during the repletion period, and at the end of this period (week 2), the mean Hb concentration of this group was 140 (sd 22) g/l.

Rats were anaesthetised through the intraperitoneal route with a 1:2 (v/v) mixture of ketamine (10 mg/kg body weight; Vetaset; Fort Dodge Animal Health) and xylazine (25 mg/kg body weight; Virbaxil 2%, Virbac). After anaesthesia, blood was collected from each rat and the liver was perfused through the subhepatic vein with a NaCl solution (9 g/l) to drain blood out of the organ. The liver was then removed, rinsed with saline, weighed and stored at −20°C until analysis. The caecum was cut adjacent to the ileocaecal valve and the caecocolonic junction, weighed (whole caecum) and placed in a Petri dish with ice(22). The pH of caecal content was measured in situ by inserting an electrode (UP-25; Denver Instrument, Denver) through the ileocaecal junction. The caecum was then cut open along the small curvature and aliquots of the caecal contents were collected, frozen in liquid N2 and adequately stored at −20°C for Fe fractionation analysis. The caecal wall was flushed clean in ice-cold saline, blotted dry and weighed (to determine the caecal wall weight). The weight of the caecal contents was calculated by taking the difference between the weights of the whole caecum and those of the caecal wall. The caecal mucosa was scraped off with a glass slide before being snap-frozen in liquid N2 and stored at −80°C until analysis. In addition, tissue samples from the caecal wall were fixed in formaldehyde solution and stored in 70% ethanol.

**Table 1. Formulation of the experimental diets**

<table>
<thead>
<tr>
<th>Ingredients (%)</th>
<th>FS</th>
<th>FP</th>
<th>RAF</th>
<th>YF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Casein*</td>
<td>20.9</td>
<td>20.9</td>
<td>20.9</td>
<td>20.9</td>
</tr>
<tr>
<td>Fibre</td>
<td>5.0</td>
<td>5.0</td>
<td>5.0</td>
<td>5.0</td>
</tr>
<tr>
<td>Soyabean oil†</td>
<td>7.0</td>
<td>7.0</td>
<td>7.0</td>
<td>7.0</td>
</tr>
<tr>
<td>Cystine</td>
<td>0.3</td>
<td>0.3</td>
<td>0.3</td>
<td>0.3</td>
</tr>
<tr>
<td>Choline bitartrate (41:1 % choline)</td>
<td>0.25</td>
<td>0.25</td>
<td>0.25</td>
<td>0.25</td>
</tr>
<tr>
<td>Vitamin mixture‡</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
<td>1.0</td>
</tr>
<tr>
<td>Mineral mixture§</td>
<td>3.5</td>
<td>3.5</td>
<td>3.5</td>
<td>3.5</td>
</tr>
<tr>
<td>Sucrose</td>
<td>10.0</td>
<td>10.0</td>
<td>2.5</td>
<td>2.5</td>
</tr>
<tr>
<td>Maize starch</td>
<td>52.05</td>
<td>52.05</td>
<td>51.45</td>
<td>46.35</td>
</tr>
<tr>
<td>Yacon flour¶</td>
<td>–</td>
<td>–</td>
<td>8.1</td>
<td>–</td>
</tr>
<tr>
<td>Yacon flour†</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>15.7</td>
</tr>
</tbody>
</table>

FS, ferrous sulphate; FP, ferric pyrophosphate; RAF, Raftilose P95; YF, yacon flour.

* Casein = 85 % protein (N x 6.25).
† Cargill Agrícola S/A.
‡ AIN-93-XX vitamin mixture(20).
§ Modified from a mineral mixture (AIN-93G-MX(20)); Fe₃(PO₄)₂ at 4.12 g Fe/kg or FeSO₄·H₂O at 2.71 g Fe/kg mix was utilised in the mineral mix of the experimental diets.
¶ Origin Activa Food International (Clariant), 93 % inulin-type fructans (ITF).
† São Sebastião Farm (São Paulo), 48 % ITF, 49 % fructose, 11 % total dietary fibre and 4 μg Fe/g.
Chemical composition of yacon flour and experimental diets

The diet fed to the CT group was formulated according to the AIN-93G diet(20). In the RAF and YF diets (Table 1), maize starch, sucrose and dietary fibre were quantitatively substituted, taking into consideration the carbohydrate content of the FOS sources. The Raftilose P95 used in the present study was donated by Clariant S/A. According to the analysis certificate, this product contains approximately 93% ITF-containing molecules (GFn) with an average polymerisation degree of 4 (FOS). The FOS in RAF were obtained by the partial hydrolysis of inulin, which was extracted from chicory roots (Cichorium intybus). Yacon tuberous roots were harvested 8 months after cultivation, and they were obtained from the Company of General Warehouses of São Paulo and processed properly as described in previous studies(15,18). The roots were autoclaved (121°C, 20min), freeze-dried (Liotechc Inústria e Comércio Ltda) and ground to obtain flour.

The total fructans and fructose in YF were analysed according to the method of Steegmans et al.(121) using an enzymatic, spectrophotometric kit (R-Biopharm AG) for glucose and fructose determination with hexokinase, glucose-6-phosphate dehydrogenase and phosphoglucose isomerase. The total dietary fibre was determined by enzymatic–gravimetric method(24). Fe concentrations in the diets and in the YF were determined by atomic absorption spectrophotometry (AAnalyst 100; Perkin Elmer) using a hollow cathode lamp at 283·4 nm and a 0·2 nm slit after wet digestion (HNO3–H2O2, 5:1; v/v). The working standard solution was prepared with FeCl3 (Tritisol; Merck). Yacon flour analysis identified 48% total fructans, 49% fructose, 11% total dietary fibre and 4 μg Fe/g. A certified reference material (lyophilised bovine liver, SRM1577c, National Institute of Standards and Technology) was used to check Fe recovery (Fe value = mean 188 (sd 14) μg/g, n 6) (certified value, 197 (sd 6) μg/g).

Iron bioavailability, hepatic iron concentration and iron fractionation analyses in the caecal contents

To determine Hb concentration during the depletion period, blood was obtained by tail puncture. At the time of killing during the repletion period, blood was obtained from the abdominal aorta. Blood Hb concentration was determined with a commercial kit (reference no. 43, Labtest Diagnostica) using the cyanide Hb method(25) and a commercially available control material (Labtest Diagnostica). The Hb Fe pool was calculated according to the following equation, assuming that 67% of the body weight is blood and that Hb contains 0·335% Fe(26):

Hb Fe pool (mg) = \frac{\text{body weight (g) × Hb (g/l) × 6·7 × 0·335}}{10000}

Values obtained in the Hb Fe pool and Fe intake calculations were used to estimate the Hb regeneration efficiency (HRE), according to the following equation(26):

HRE (%) = \frac{(Hb \text{ Fe pool (final)} - Hb \text{ Fe pool (initial)}) \times 100}{\text{Fe intake (mg)}}

The bioavailability of Fe from FS (FS group) was considered as the reference and was therefore assigned a value of 100%(18,27). Thus, the relative Fe bioavailability, or biological value (RBV) of each FP group (FP, RAF and YF groups), was determined at weeks 1 and 2 of the repletion period as follows:

RBV (%) = 100 \times \frac{\text{HRE test group}}{\text{HRE FS group}}

The total Fe in the left lateral lobe of the liver and the caecal contents was determined as described previously for dietary Fe analysis. Caecal contents were subjected to a sequential solubilisation to analyse the intestinal Fe species distribution, as described by Simpson et al.(28), which was modified from Tessier’s method(29). In brief, approximately 1 g of the sample was subjected to extraction under stirring followed by centrifugation (10 000 g for 30 min). The supernatant was collected and combined with the washed pellet that was obtained by mixing the demineralised water after sample re-centrifugation. Each fraction of the extraction process corresponded to the supernatant added to its wash. The resulting pellet was subjected to the next extraction step. Fe concentrations were determined in the following five fractions: exchangeable (soluble in 1M-magnesium chloride; for 1 h at room temperature); carbonate-bound (acid soluble; 1M-sodium acetate, pH 5·0-adjusted with acetic acid, for 4 h at room temperature); oxide-bound (soluble in 0·04M-hydroxy-lamine in acetic acid; for 16 h at 95°C); organic matter (soluble after treating with 0·02M-nitric acid and H2O2 (pH 4·8 adjusted with nitric acid); for a total of 5 h at 85°C); residual (the remaining pellet)(28,29). Fe concentrations were obtained directly from calibration curves that were prepared from the solution components used in different extraction steps. The analytical precision for each extraction step was evaluated by subjecting ten subsamples of a caecal content pool to the sequential procedure described above. The pool was obtained by homogenising the caecal contents of male Wistar rats (n 10; body weight approximately 200 g) fed commercial chow (664 mg Fe/kg). CV of 10·5% (for the organic matter fraction) or lower (6·8, 9·4, 6·3 and 9·7% for exchangeable, carbonate-bound, oxide-bound and residual fractions, respectively) were obtained. A comparison of the sum of Fe concentrations in each fraction was consistent with the total Fe concentrations (r2 0·98; Fe value = mean 13·76 (sd 1·15) vs. 13·95 (sd 1·14) μmol/g, respectively).

Immunoblotting

N-frozen samples of caecal mucosa were homogenised (T10 basic Ultra Turrax; IKA Works, Inc.) in T-PER Tissue Protein Extraction Reagent (Pierce Technology) containing
protease and phosphatase inhibitors (Pierce Technology). The homogenates were centrifuged at 10 000g for 5 min, and total protein concentrations in the supernatants were determined using the bicinchoninic acid assay (Pierce Technology). Equal amounts of total protein (50 μg) were denatured by boiling for 5 min in 4× SDS Laemmli buffer (200 mm-Tris–HCl pH 6·8, 8% (v/v) SDS, 40% (v/v) glycerol, 20% (v/v) β-mercaptoethanol and 0·4% (v/v) bromophenol blue), separated by 12·5% SDS–PAGE (Fisher Scientific) and then transferred onto nitrocellulose membranes (Whatman). The blots were pre-incubated in Blocker Casein Buffer (Pierce Technology) for 2 h at room temperature and then incubated with 1:500 (v/v) diluted rabbit polyclonal anti-FPN1 (Abcam Anti-logy) for 2 h at room temperature and then incubated with 1:5000 (v/v) diluted peroxidase-labelled anti-rabbit secondary antibodies (Sigma Chemical Company), and the immunoreactive band signals were visualised with an enhanced chemiluminescence kit (GE Healthcare). ImageQuant 400 equipment (GE Healthcare) and Quantity One Basic software (BioRad Laboratories) were used for scanning the photo-images and for densitometry analysis, respectively. Each membrane was re-probed with 1:50 000 (v/v) diluted peroxidase-labelled anti-β-actin monoclonal antibody (Sigma Chemical Company). FPN protein expression data were normalised to corresponding β-actin values and expressed as arbitrary units.

Caecal histology

To perform a histological examination, the fixed tissue fragments were embedded in paraffin and approximately 5 μm-thick sections were obtained and stained with haematoxylin and eosin. In each crypt, the cells in the left-hand column were counted (cells/hemicrypt) from the bottom to the top of the caecal crypt under an optical microscope. At least thirty hemicyclets per rat were assessed, and only crypts that were cut lengthwise were considered. To determine the percentage of bifurcating crypts, the crypts with an indentation at the base or those presenting longitudinal fission (one crypt mouth and two bases) were considered. The calculation was performed by determining the percentage of bifurcating crypts (only crypts cut lengthwise) per microscopic field. To estimate the total number of crypts per microscopic field, the obliquely sectioned crypts were also considered. At least twenty microscopic fields per rat were analysed. To evaluate the metaphase index, each rat was administered a single intraperitoneal injection of vincristine sulphate (Tecnocris, Zodiac Produtos Farmaceuticos S/A; 1 mg/kg body weight) 2 h before being killed. The metaphase index was determined by counting the number of cells arrested at metaphase under a light microscope (Nikon). Approximately 2000 cells per rat were counted in longitudinally sectioned crypts using an ocular grid (Zeiss Integration Eyepiece 1 Kpl 8; Carl Zeiss) at 1000X magnification. The metaphase index was determined as the ratio of the number of arrested metaphases: the total number of cells.

Statistical analysis

Statistical analysis was performed using Prism 5 for Windows (version 5.00, 2007; Graphpad Software, Inc.). All tests were performed by assuming the bilateral hypotheses and a 5% significance level. Descriptive statistics were initially used to evaluate the means and standard deviations of the studied variable. During the depletion period, the mean values of the CT and ID groups were compared using an unpaired t test. During the repletion period, the variable means of the groups were compared using an ANOVA. Tukey’s post hoc test was used to identify where significant differences occurred. A non-parametric Kolmogorov–Smirnov test was used to verify the normality of the observations, and when the normality hypothesis was rejected, ANOVA was substituted with non-parametric Kruskal–Wallis and Dunn’s post hoc tests. The observed power was 85–95% for most tests.

Results

Total food and iron intake, body weight gain, iron bioavailability parameters and hepatic iron concentrations

At the end of the depletion period, the blood Hb concentrations and body weights of the ID group were significantly reduced in comparison with those of the CT group (59 (SD 6) vs. 122 (SD 19) g/L, P < 0·0001, and 82 (SD 18) vs. 104 (SD 5) g, P=0·05, respectively). During the repletion period, no significant differences were observed in the total food intake among the groups (112 and 260 g at weeks 1 and 2, respectively, in the FS group). Moreover, there were no differences in total Fe intake (5·5 and 13 mg at weeks 1 and 2, respectively, in the FS group), considering that the different diets had similar Fe concentrations, as measured by atomic absorption spectrophotometry (49·4, 44·9, 45·3 and 47·0 mg Fe/kg diet in the FS, FP, RAF and YF diets, respectively). There were no differences in body weights among the groups at the end of week 1, but at week 2, YF supplementation resulted in a higher weight gain in comparison with that in the other groups (P<0·05).

Blood Hb concentrations, Hb Fe pool, HRE values, RBV and hepatic Fe concentrations during the repletion period are shown in Fig. 1. In general, the effects of FOS in anaemic rats were observed within the 1st week of the repletion period. Hence, considering that blood Hb concentrations were similar among the groups at the start of this period (P=0·78), changes in Hb concentrations in FOS-fed rats at week 1 were greater than those in the FP group (P<0·01) and similar to those in the FS group. At week 2, there were no differences in blood Hb concentrations among the groups (Fig. 1(A)). Moreover, no changes were observed in the Hb Fe pool among the groups throughout the repletion period (Fig. 1(B)).

The efficiency of Hb recovery reflects the ratio of dietary Fe conversion into Hb: the amount of ingested Fe over the course of the repletion period. In the present study, only the RAF group exhibited higher HRE values than the FP group and had HRE values similar to those of the FS group at week 1 of the repletion period (P < 0·05; Fig. 1(C)). The Hb
concentration observed at the end of each week during the repletion period (weeks 1 and 2) in the FS group was considered as the reference. Hence, the FS group was used as a reference to express the bioavailability of Fe from FP (FS is assigned a 100% value) (RBV). The RBV of the FP in the RAF group were higher than those of the FP in the FS group (111% after the 1st week and 109% after the 2nd week). These values were significantly higher than those in the FP group at week 1 ($P < 0.05$; Fig. 1(D)). Moreover, there was no significant difference in the RBV between rats provided with either dietary FOS source at week 1.

As expected, hepatic Fe concentrations were lower in the FP group than in the FS group at week 1 of the repletion period, whereas FOS supplementation led to the recovery of hepatic Fe stores to levels comparable to those observed in the FS group ($P < 0.01$; Fig. 1(E)). In addition, there were no differences in hepatic Fe concentrations between the two FOS groups.

**Effects of fructo-oligosaccharides on caecal parameters during recovery from anaemia: early changes in iron species distribution, mucosal ferroportin-1 expression, morphometry and crypt cell proliferation**

Because the effects of FOS on Fe bioavailability had already been observed at week 1 of the repletion period, the intestinal changes in response to the supplementation of these fermentable fibres were evaluated. The effects of FOS supplementation on caecal parameters (caecum weight, caecal content pH values and Fe species distribution in the caecal contents) are summarised in Table 2. FOS supplementation resulted in heavier caeca (total, wall and contents), irrespective of the dietary FOS source supplemented, when compared with those of the FP group ($P < 0.05$). Moreover, the caecal content pH was decreased by FOS supplementation, but only the values of the RAF group were significantly different from those of the CT rats ($P < 0.05$). Accordingly, although the
total Fe concentrations in the caecal contents were not significantly different among the groups, RAF supplementation redistributed Fe from the carbonate-bound fraction (which is sensitive to pH changes) to the exchangeable (more bioavailable) and oxide-bound (susceptible to reduction) fractions (Fig. 2). Thus, the concentrations identified for exchangeable Fe were higher in rats fed the RAF diet ($P=0.01$) than in those fed the YF and CT diets (an increase from 2 to 5 % relative to the total Fe concentrations in the CT and RAF rats, respectively (Fig. 2)). Similarly, the levels of Fe bound to oxides were significantly increased by RAF supplementation ($P<0.05$). Fe concentrations in the oxide-bound fraction represented 6–7 % of those in the CT rats and increased to 17 % in RAF rats (Fig. 2). The levels of Fe bound to organic matter were lower than their detection limits, representing a negligible fraction of the total amount of Fe in the caecal contents (which are therefore not shown in Fig. 2). Fe concentrations in the residual fraction were higher than those observed in any of the preceding extractions (Fig. 2) and were not different among the groups (Table 2).

The expression of FPN-1, the transporter responsible for cellular Fe efflux, was increased in the caecal mucosa of the FP group ($P=0.04$; Fig. 3), which accounted for the highest expected absorption efficiency in rats from the FP group.

### Table 2. Caecal parameters (organ weight, pH values, total iron concentrations and iron species distribution in the caecal contents) of iron-deficient rats fed diets containing ferrous sulphate (FS) or ferric pyrophosphate (FP) supplemented with 7.5 % inulin-type fructans in the form of Raftilose P95 (RAF) or yacon flour (YF) during the 1st week of recovery of Hb

(Mean values with their standard deviations, $n=8$)

<table>
<thead>
<tr>
<th>Analyses</th>
<th>FS</th>
<th>FP</th>
<th>RAF</th>
<th>YF</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dietary treatments</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td><strong>Caecum wet weight (g)</strong></td>
<td>2.5$^{a,b}$</td>
<td>1.9$^a$</td>
<td>6.3$^b$</td>
<td>4.5$^{ab}$</td>
<td>0.02</td>
</tr>
<tr>
<td><strong>Caecum wet weight (g/100 g)</strong></td>
<td>2.0$^a$</td>
<td>1.2$^a$</td>
<td>3.8$^a$</td>
<td>2.6$^a$</td>
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<tr>
<td><strong>Caecal wall weight (g)</strong></td>
<td>0.39$^{a,b}$</td>
<td>0.37$^a$</td>
<td>0.84$^b$</td>
<td>0.62$^{ab}$</td>
<td>0.01</td>
</tr>
<tr>
<td><strong>Caecal wall weight (g/100 g)</strong></td>
<td>0.32$^{a,b}$</td>
<td>0.23$^a$</td>
<td>0.52$^b$</td>
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<td></td>
</tr>
<tr>
<td>Weight (g)</td>
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<td>5.4$^b$</td>
<td>3.0$^{ab}$</td>
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<tr>
<td>Weight (g/100 g)</td>
<td>1.7$^a$</td>
<td>0.9$^a$</td>
<td>3.3$^b$</td>
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<td>7.1$^a$</td>
<td>6.9$^b$</td>
<td>7.0$^{ab}$</td>
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<td>Total Fe ($\mu$mol/g)</td>
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<td>2.7$^a$</td>
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<td>2.6$^a$</td>
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<tr>
<td><strong>Fe species distribution ($\mu$mol/g)</strong></td>
<td></td>
<td></td>
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<tr>
<td>Exchangeable</td>
<td>0.08$^a$</td>
<td>0.07$^a$</td>
<td>0.15$^b$</td>
<td>0.09$^a$</td>
<td>0.0002</td>
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<td>Carbonate-bound</td>
<td>0.58$^a$</td>
<td>0.60$^a$</td>
<td>0.12$^b$</td>
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<td>&lt;0.0001</td>
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<tr>
<td>Oxide-bound</td>
<td>0.20$^a$</td>
<td>0.20$^a$</td>
<td>0.50$^b$</td>
<td>0.29$^a$</td>
<td>0.0003</td>
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<td>ND</td>
<td>ND</td>
<td>ND</td>
<td></td>
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<tr>
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<td>1.86</td>
<td>2.07</td>
<td>2.11</td>
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</table>

ND, not detected.

$^{a,b}$ Mean values within a row with unlike superscript letters were significantly different ($P<0.05$).

$^*$ Data were analysed using ANOVA and Tukey’s post hoc tests.

$^†$ Data were analysed using non-parametric Kruskal–Wallis and Dunn’s post hoc tests.

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**Fig. 2.** Effects of fructo-oligosaccharides on the distribution of iron species in caecal contents during the 1st week of recovery of Hb in anaemic rats. Iron species distribution in the caecal contents of iron-deficient rats fed diets containing ferrous sulphate (FS), ferric pyrophosphate (FP) or FP supplemented with 7.5 % inulin-type fructans in the form of Raftilose P95 (RAF) or yacon flour (YF). The fractions are as follows: exchangeable ( ); carbonate-bound ( ); oxide-bound ( ); residual ( ). Values are means ($n=8$), with standard deviations represented by vertical bars.
that still remained anaemic at week 1 of the repletion period. Moreover, there were no differences in FPN-1 expression between FOS-fed rats and FS rats (Fig. 3).

Caecal enlargement was in agreement with the histological changes observed in FOS-fed rats; there was a noticeable increase ($P<0.05$) in the number of cells per crypt (Fig. 4(A)), the number of crypts per field (Fig. 4(B)) and the percentage of bifurcating crypts per field (Fig. 4(C)) when compared with those in the CT rats. Consistent with these findings, cell proliferation as assessed by the $2h$ accumulation of vincristine-arrested metaphases per crypt was increased by FOS supplementation ($P<0.05$), and this effect was more pronounced in the RAF group (Fig. 4(D)).

**Discussion**

The primary strategies for correcting Fe deficiency in populations are dietary modification or diversification to improve Fe intake and bioavailability; Fe supplementation and Fe fortification of foods; and biofortification by plant breeding (31). Although the efficiency of Fe absorption increases as Fe stores become depleted, the amount absorbed from foods, especially when diets are low in meat, fish, fruit and vegetables, is not enough to prevent Fe deficiency in many women and children, especially in developing countries (31–32). For instance, the primary cause of increasing Fe deficiency in Brazil is that the consumption of food items considered to be Fe sources has continually decreased (33). Fe deficiency in Brazil is that the consumption of food items containing ferrous sulphate (FS, $\alpha$), ferric pyrophosphate (FP, $\beta$) or FP supplemented with 7.5% inulin-type fructans in the form of Raftilose P95 (RAF, $\gamma$) or yacon flour (YF, $\delta$). Values are means ($n$ 4), with standard deviations represented by vertical bars. Mean value was significantly different from that of the YF group ($P<0.05$; non-parametric Kruskal–Wallis and Dunn’s post hoc tests). a, $\beta$-Actin, immunoblotting.

Fig. 3. Effects of fructo-oligosaccharides on ferroportin-1 (FPN-1) expression in the caecal mucosa of anaemic rats during the 1st week of recovery of Hb. FPN-1 expression in the caecal mucosa of iron-deficient rats fed diets containing ferrous sulphate (FS, $\alpha$), ferric pyrophosphate (FP, $\beta$) or FP supplemented with 7.5% inulin-type fructans in the form of Raftilose P95 (RAF, $\gamma$) or yacon flour (YF, $\delta$). Values are means ($n$ 4), with standard deviations represented by vertical bars. *Mean value was significantly different from that of the FS group ($P<0.05$; non-parametric Kruskal–Wallis and Dunn’s post hoc tests).<ref>

Fig. 4. Changes in the caecal mucosa architecture of iron-deficient rats fed fructo-oligosaccharides during the 1st week of recovery of Hb. The number of cells (A), number of crypts (B), percentage of bifurcating crypts (C) and metaphase index in the crypts (D) in the caecum of iron-deficient rats fed diets containing ferrous sulphate (FS, $\alpha$), ferric pyrophosphate (FP, $\beta$) or FP supplemented with 7.5% inulin-type fructans in the form of Raftilose P95 (RAF, $\gamma$) or yacon flour (YF, $\delta$). Values are means ($n$ 5), with standard deviations represented by vertical bars. Mean values with unlike letters were significantly different ($P<0.05$; non-parametric Kruskal–Wallis and Dunn’s post hoc tests).

Discussion

The primary strategies for correcting Fe deficiency in populations are dietary modification or diversification to improve Fe intake and bioavailability; Fe supplementation and Fe fortification of foods; and biofortification by plant breeding (31). Although the efficiency of Fe absorption increases as Fe stores become depleted, the amount absorbed from foods, especially when diets are low in meat, fish, fruit and vegetables, is not enough to prevent Fe deficiency in many women and children, especially in developing countries (31–32). For instance, the primary cause of increasing Fe deficiency in Brazil is that the consumption of food items considered to be Fe sources has continually decreased (33). Indeed, the search for new food standards, proposals for food distribution and knowledge about the dietary composition must be the researcher’s target.

The effects of dietary fibre or fibre-rich diets on mineral absorption have been discussed for many years and appear to fundamentally depend on whether they are observed in vitro or in vivo studies (34,35). On the one hand, some fibres (e.g. cellulose, hemicelluloses and pectin) or fibre-associated substances (e.g. phytates and phenolics) negatively affect the absorption of minerals by binding to or entrapping them in the lumen of the small intestine (35). On the other hand, these negative effects seem to be reduced if the fibres are fermented in the large intestine. In this context, the positive effects of a group of fermentable fibres (namely ITF (FOS and inulin)) on the intestinal absorption and bioavailability of some minerals have been repeatedly demonstrated in animal (7,12–18) and human (36,37) studies. Most of these studies focused on the bioavailability of Ca and Mg. Nevertheless, the number of studies concerning the bioavailability of trace elements (Fe, Cu and Zn) is proportionally lower, and these studies have thus far yielded contradictory results. In particular, there is some evidence in rats (18,38,39) and pigs (15,14,40,41) that Fe bioavailability is positively affected. However, these effects have not been consistently demonstrated in human studies (42). To our knowledge, there are no studies that have compared the effects of purified and non-purified sources of ITF on Fe bioavailability and on histological changes in the large intestine of rats subjected to an experimental model of Fe-deficiency anaemia. We had previously demonstrated the improved bioavailability of Fe from FP (as measured by Hb repletion efficiency) by supplementing diets with 7.5% ITF from either RAF (a purified source)
or yacon tuberous roots (a non-purified source) in anaemic rats (18). In the present study, we confirmed these findings and demonstrated that early changes in the caecal environment (luminal Fe species redistribution, decreased mucosal FPN-1 expression, and morphological alterations) could be related to Fe bioavailability improvement by FOS supplementation.

Fe in foods is released into the lumen as haem or as non-haem ferric complexes (43,44). However, non-haem Fe is most affected by the food matrix composition and the physicochemical conditions in the intestinal lumen (44). Among the various oxidation states of Fe, divalent Fe (Fe2+) and trivalent Fe (Fe3+) are more stable in aqueous media, and they occur naturally in foods (44). The two forms are soluble at the pH of the stomach; nonetheless, Fe3+ precipitates more rapidly than Fe2+ because of the alkalinisation of the alimentary bolus in the duodenum during the digestive process. However, Fe precipitation can be inhibited by complexing agents in foods, and these agents include amino acids, carboxylic acids, carbohydrates and phosphates (43,44). Thus, factors that affect the luminal pH or chelating properties of certain dietary components can also affect their redox potential and interfere with their solubility (43,44).

In this context, physico-chemical changes in the caecal environment resulting from the bacterial fermentation of non-digestible carbohydrates could contribute to alterations in mineral speciation and thus mineral absorbability (10,11,18,45). In the present study, the lower luminal pH observed in the caecum of FOS-fed rats could have contributed to Fe mobilisation from the carbonate-bound to the exchangeable and oxide-bound fractions. Only 10% of dietary Fe is absorbed in the small intestine (primarily in the duodenum), which indicates that significant amounts of Fe are recovered in the luminal content of the large intestine (46). Thus, our findings indicate that FOS fermentation could have increased the fraction of Fe available for uptake in the caecum by approximately 15%.

Complementarily, mucosal FPN-1 expression in the caecum was decreased in FOS-fed rats at a magnitude similar to that observed in the FS group. Some studies have also demonstrated that, under specific circumstances, the proximal colon may significantly contribute to Fe absorption (13,40,47,48). In fact, feeding rats with an Fe-deficient diet induces an increase in the gene activity of the microbiota (1,8,50–52). Butyrate is recognised as the primary source of energy for colonocytes, playing an important role in the stimulus of cell division in the intestinal mucosa (51,52). We had previously demonstrated increased caecal SCFA production by FOS supplementation and found YF to be more butyrogenic than RAF (18). However, the yacon roots used in that study had lower fructan contents than those used in the present study (18% (18) vs. 48%), most probably because of the increased fructan depolymerisation catalysed by fructan 1-exohydrolase (53), which led us to use a larger amount of YF in the formulation of experimental diets (41.7% (18) vs. 15.7%). Thus, although the fructan content was the same (7.5%) in the different diets (RAF and YF), the dietary fibre (soluble and insoluble) concentrations (as measured by the enzymatic−gravimetric method (24)) in the additional YF used for diet formulation may have contributed to the higher fermentability observed in the YF group in that study (18). In fact, in the present study, besides exhibiting a lower caecal content pH, RAF rats also had heavier caecae compared with YF rats. Factors other than dietary fibre fermentability may also directly influence the mucosal growth of the caecum, such as increased intestinal content viscosity (43,44).

In addition, caecum weight can be influenced by stool water content because FOs remain in solution in the chyme and contribute to increased osmotic pressure, resulting in increased water flow to the intestinal lumen (16,17,54).

Bacterial fermentation has also been implicated in morphological changes in the intestinal epithelium of FOS-fed animals. In this context, germ-free rats were found to exhibit an increase in intestinal weights, but exhibited no proliferative effects in the intestinal mucosa in response to fibre consumption, confirming that the fermentation products (SCFA) are responsible for trophic effects on the intestinal mucosa (5,55). Thus, that FOS-containing diets indirectly affect mucosal morphology (with increases in the number and depth of crypts and crypt bifurcation) in the rat large intestine has previously been demonstrated by us (7,45) and other authors (5,55). Moreover, the increased number of caecal crypts was supported by the present data on crypt cell proliferation and bifurcation, which are complementary mechanisms to increase tissue mass and were stimulated upon FOS supplementation. The production of new crypts by fission is a process that is thought to be related to the expansion of the crypt stem cell population and is increased in postnatal development, during intestinal recovery from injury and in crypts isolated from adenomas and hyperplastic polyps (50,56,57). Thus, FOS fermentability must be addressed with some caution, because the observed effects may, on the one hand, be interesting for promoting mineral absorption and bioavailability by increasing the absorptive surface and contributing to the improvement of mineral solubility in the caecum. On the other hand, the intestinal changes caused by this type of carbohydrate (rapidly fermentable) have been linked to the promotion of intestinal tumour development (57,58). Furthermore, some studies in rats have demonstrated detrimental effects on the intestinal barrier function after FOS supplementation in Ca-restricted diets (5,59). Thus, from a nutritionist’s point of view, a dietary prescription for fermentable fibres should consider not only the type and amount of fibre but also whether the consumed fibre is given in a purified or non-purified form. In addition, the composition of the food...
matrix, in which the fibre is present, is another important factor to be taken into consideration.

In conclusion, dietary FOS supplementation improves Fe bioavailability in anaemic rats by changing luminal Fe solubility, most probably as a result of bacterial fermentation in the cecum. These effects were more pronounced on using a purified source (RAF) than on using a non-purified source (YF) of fructans. These effects, if confirmed in humans, might contribute to the formulation of specific diets for individuals with Fe deficiency. The changes observed in the caecal tissue of FOS-fed animals must be evaluated with caution because increased cell proliferation and crypt fission have been implicated in the carcinogenic process within the large intestine.

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The authors’ contributions are as follows: A. R. L. was responsible for the study concept and design and for the acquisition, analysis and interpretation of the data; E. H. S. G. and E. D. C. were responsible for the acquisition and analysis of the data; E. P. A. was responsible for the acquisition, analysis and interpretation of the data; C. C. was responsible for the study concept and design, acquired, analysed and interpreted the data, and obtained funding.

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