Soluble maize fibre affects short-term calcium absorption in adolescent boys and girls: a randomised controlled trial using dual stable isotopic tracers

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Abstract

Soluble maize fibre (SCF) has been found to significantly improve bone mineral density and strength in growing rats compared with several other novel prebiotic fibres. The objective of the present study was to investigate the effect of SCF on Ca absorption and retention in pubertal children by studying the potential absorption mechanisms of the intestinal microbiota. A total of twenty-four adolescent boys and girls (12–15 years) participated in two 3-week metabolic balance studies testing 0 g/d SCF (control (CON) treatment) and 12 g/d SCF (SCF treatment) in a random order by inclusion in a low-Ca diet (600 mg/d). Fractional Ca absorption was measured at the end of the two intervention periods using a dual-stable isotope method. Diet composites and faecal and urine samples were collected daily and analysed for Ca content. Ca retention was calculated as dietary Ca intake minus Ca excretion in faeces and urine over the last 2 weeks. Microbial community composition in the faecal samples collected at the beginning and end of each session was determined by 454 pyrosequencing of the PCR-amplified 16S ribosomal RNA gene. Fractional Ca absorption was 12% higher (41 mg/d) after the SCF treatment compared with that after the CON treatment (0·664 (SD 0·129) and 0·595 (SD 0·142), respectively; P=0·02), but Ca retention was unaffected. The average proportion of bacteria of the phylum Bacteroidetes was significantly greater in the participants after the SCF treatment than after the CON treatment. These results suggest that moderate daily intake of SCF, a well-tolerated prebiotic fibre, increases short-term Ca absorption in adolescents consuming less than the recommended amounts of Ca.

Key words: Prebiotics: Calcium absorption: Microbiota: Soluble maize fibre

Adolescence is an important stage of life for bone health that provides a unique opportunity to maximise mineral retention and prevent the risk of osteoporosis-related fractures later in life. Because Ca is becoming increasingly deficient in the diet due to a decrease in milk consumption1, there is strong interest in functional food ingredients for enhancing Ca utilisation. Non-digestible oligosaccharides, including fructo-oligosaccharides3,4 and galacto-oligosaccharides5, have been reported to be associated with increased Ca absorption. In animal studies, these carbohydrate compounds have been found to be associated with increases in bone mineral density (BMD)6–8 as well as improved bone geometry and strength9,10. The increase in Ca absorption and subsequent improvements in bone health are possibly mediated by microbial fermentation. Bongers & Van den Heuvel11 suggested that SCFA produced by microbial fermentation can produce an acidic intestinal environment ideal for increasing the solubility and transcellular absorption of Ca.

We had compared eight novel fibres for their effects on Ca utilisation and bone properties in a growing rat model12. Soluble maize fibre (SCF) was found to have effects similar to those of soluble fibre dextrin and confer greater bone health benefits when compared with resistant starch (RS), pullulan, polydextrose, inulin and Synergy1®; ORAFTI. SCF increased bone mineral content by 7.6%, BMD by 2.9%, femoral cortical area by 19.6%, cortical thickness by 22.4% and peak breaking force by 88.8%. Compared with controls, SCF resulted in significant increases in faecal content weight and total SCFA, acetate and propionate production, which, in part, indicate the involvement of intestinal microbes.

Factors that enhance bone density and bone mineral content have the potential to contribute to a skeleton that

Abbreviations: BMD, bone mineral density; CON, control; IGFBP3, insulin-like growth factor-binding protein 3; NTX, cross-linked N-telopeptides of type I collagen; OTU, operational taxonomic units; rRNA, ribosomal RNA; RS, resistant starch; SCF, soluble maize fibre.

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will resist fractures later in life. Prebiotics may provide a novel means of influencing mineral bioavailability to improve peak bone mass. Therefore, the objective of the present study was to better understand the role of SCF in humans by examining its effects on Ca absorption and retention as well as changes in gut microbiota composition in adolescent boys and girls consuming habitually low daily Ca amounts.

Subjects and methods

Subjects

Healthy adolescent girls and boys were recruited from local schools, community centres and Indiana extension offices and by sending direct mails to the surrounding areas. A total of fifteen boys, aged 13–15 years, and nine girls, aged 12–14 years, participated in the metabolic studies. Screening questionnaires were used to determine eligibility based on a brief medical history, maturational age, physical activity and habitual dietary intake assessed with a 6 d diet record. Adolescent girls and boys who were generally healthy and were willing to consume the foods provided as part of the controlled diets were eligible to participate in the study. Exclusion criteria included abnormal liver or kidney function, malabsorptive disorders, anaemia, smoking, history of using medications influencing Ca metabolism (steroids and thiazide diuretics), body weight outside the 5th–95th BMI percentile for age, regular consumption of illegal drugs, non-prescription drugs, or any kind of contraceptives, and pregnancy. The subjects were not permitted to take nutritional supplements while participating in these studies and were asked to discontinue use before coming to the camp. The present study was conducted according to the guidelines laid down in the Declaration of Helsinki, and all procedures involving human subjects were approved by the Institutional Review Board of Purdue University. Written informed consent was obtained from all subjects. The present study was registered at ClinicalTrials.gov ID: NCT01571440.

Study design

The present study, designed to provide a summer camp environment, consisted of two 3-week balance studies separated by a 7 d washout period. A double-blind, cross-over design was used in which the participants received two treatments in a randomised order: 12 g SCF (SCF treatment) or 0 g SCF (control (CON) treatment). The study staff used a simple technique for randomisation such that the treatment orders were drawn, for each participant, from a hat containing equal numbers of the two possible randomisation schemes. Studies were structured as summer camps where adolescent boys and girls went on field trips, participated in recreational activities and performed a variety of educational activities. The participants were housed in University Residence Halls at Purdue University during the camp duration.

Diet

Controlled diets were provided throughout both camp sessions and consisted of foods typically eaten by adolescent children such as spaghetti, hamburgers, sandwiches and potato chips. The participants were assigned to one of five energy levels (7322, 8786, 10 042, 11 297 and 12 552 kJ) based on the estimated energy requirements calculated using the Harris–Benedict equations. The diets were designed to maintain body weight and to contain constant amounts of key nutrients. The controlled diets were provided as a 4 d cycle menu with three meals and two snacks daily. On average, the diets contained 14% protein, 33% fat, 53% carbohydrate, 5 μg vitamin D, 1100 mg P, 2300 mg Na and 600 mg Ca. The basal diet contained 15 g of fibre and the intervention product contributed an additional 0 or 12 g SCF. This yielded a total dietary fibre content of 15 and 27 g for the CON and SCF treatments, respectively. SCF was present in Welch’s® fruit snacks (WELCH Foods, Inc.) and divided into two 0 or 6 g fibre doses provided at lunch and dinner. The SCF ingredient (PROMITOR® Soluble Corn Fiber) provided by Tate & Lyle is a fermentable, non-digestible carbohydrate and contains 70% (minimum) soluble dietary fibre.

Anthropometric parameter and bone measurements

Anthropometric parameters including weight, sitting height, bitrochanteric width, waist circumference and hip circumference were measured during the first session of the camp. Standing height was measured at the beginning of the first session using a wall-mounted stadiometer, and weight was monitored in the morning on each day with an electronic digital scale to ensure that it remained stable throughout the sessions. Bone mineral content and BMD were measured by dual-energy X-ray absorptiometry (GE Lunar) during one balance period to determine the bone status of the participants. Bone measurements of the whole body, spine, forearm and both hips were taken.

Measurement of hormones and biochemical markers of bone metabolism

A fasting, baseline blood sample was collected on the 1st day of the camp for the determination of general blood chemistry to verify the clinical profile and health of the participants. A second fasting sample was collected at the end of the camp to measure the biochemical markers of bone dynamics and hormones related to Ca and vitamin D metabolism. The concentrations of serum total Ca, phosphate, total alkaline phosphatase and creatinine and urinary creatinine and phosphate were measured by routine chemistry. The concentration of serum parathyroid hormone 1–84 was determined using a two-site immunoassay (CV ± 7.1%) (Nichols Institute Diagnostics). The concentrations of insulin-like growth factor 1 and insulin-like growth factor-binding protein 3 (IGFBP3) were measured by ELISA (Diagnostics Systems Laboratories, Inc.). The concentration of leptin was determined by RIA (EMD Millipore) and that of sclerostin by ELISA (TECO-medical AG). The concentrations of urinary cross-linked N-telopeptides of type I collagen (NTX) were measured by ELISA (Ostex International). The concentration of serum
intravenous dose.

Sample collection and analysis

All urine and faecal samples were collected from day 1 to day 21 of each balance period and pooled as 24 h collections. Ca content in the diets and faecal and urine samples was measured using inductively coupled plasma optical emission spectrometry (Optima 4300 DV, Perkin Elmer Instrument) as described previously. All faecal samples were frozen and processed later for the determination of Ca content. Urine samples were refrigerated and also analysed later for the determination of total Ca content. The procedure followed for faecal sampling for the microbiological analysis is described below.

Evaluation of compliance

The participants were supervised during activity, meal and sample collection periods by trained counsellors for 24 h each day. Unconsumed food from meals was collected and its amount recorded. Urine sample collection compliance was evaluated by measuring creatinine excreted in the urine by an enzymatic colorimetric assay (COBAS Integra, Roche Diagnostics). Faecal sample collection compliance was assessed by polyethylene glycol recovery in the faeces. Each participant was given 3 g of polyethylene glycol (E3350; Dow Chemical Company) divided into 1 g doses at breakfast, lunch and dinner. The percentage of polyethylene glycol recovery was measured in 24 h faecal collections by a turbidimetric assay and was used as a basis to exclude participant data in case of poor compliance.

Evaluation of gastrointestinal symptoms

The presence of stomach noises, flatulence, bloating and abdominal pain among the participants was evaluated daily using a short questionnaire. The severity of gastrointestinal symptoms was assessed daily by self-reports using a scale of 1–10 (0 = none and 10 = very severe). The questionnaire was created during the second 3-week treatment, which did not allow for validation before use and resulted in a shorter assessment period (18 consecutive days).

Fractional calcium absorption test

During the last week of each session, following an overnight fast, the participants underwent a Ca absorption test carried out using methods described previously. On the morning of the test, phlebotomists inserted catheters and drew 10 ml baseline venous samples. Immediately after the blood draw, the participants consumed a breakfast consisting of an English muffin, scrambled eggs, butter and jam. The meal contained 150 mg Ca from 2 % milk plus 15 mg 44Ca (96·5 % enrichment), a stable non-radioactive isotope. The oral isotope was administered as a liquid calcium chloride (44CaCl2) that was added to the milk and allowed to equilibrate overnight.

Following breakfast, the participants were not allowed to consume any food, but were allowed to drink ad libitum deionised water. A second stable isotope, 43Ca (5 mg, 60·2 % enrichment) also given as calcium chloride, was administered intravenously 1 h after the consumption of breakfast and the oral isotope. A final blood draw was performed 3 h after the administration of the intravenous dose, after which the catheters were removed and the participants were served lunch.

Absorption and retention calculations

Changes in fractional Ca absorption over 48 h following Ca isotope administration were measured using two 24 h urine pools with a modified version of the published methodologies. Urine samples collected in 24 h pools for 2 d (0–24 h and 24–48 h) after the absorption test were analysed for 44Ca and 43Ca enrichment by high-resolution inductively coupled plasma MS (Finnegan Element 2, Thermo Scientific). To more specifically measure increases in absorption due to bacterial fermentation in the colon, two separate 24 h pooled analyses were used instead of using the more typical 48 h cumulative urine analysis. This method is based on previous results indicating that a urine collection period of at least 36 h is required to detect an effect of prebiotics. Previously, our laboratory group had found that separating urine samples into 12 h pools was helpful in identifying the 24–48 h time frame as being important for measuring lower gut absorption. Fractional Ca absorption (equation 1) was calculated as an enrichment ratio of the administered isotopes (43Ca and 44Ca) to another naturally occurring isotope (42Ca) in the 0–24 and 24–48 h urine collections. The Δ excess value was calculated as the presence of 44Ca and 43Ca relative to 42Ca in the urine at the end of each 24 h period minus the baseline abundance ratios of 44Ca and 43Ca relative to 42Ca. These values were corrected for the natural abundance of each isotope (44Ca = 0·02083, 43Ca = 0·00135) and doses (mg) given:

\[
\text{Fractional Ca absorption} = \left( \frac{\text{44Ca } \Delta \text{excess}}{\text{43Ca } \Delta \text{excess}} \times \frac{0.02083}{0.00135} \right) \times \left( \frac{\text{intravenous dose}}{\text{oral dose}} \right).
\]

Balance data were used to calculate Ca retention (equation 2) by subtracting 24 h Ca excretion in urine and faeces from 24 h dietary Ca intake. The first week of each 3-week study was considered to be the equilibration period for the participants to get acclimated to Ca intake and fibre treatment, while the remaining 2 weeks were considered to be the experimental period. Balance was calculated based on as many of the 14-d present in the experimental period as possible, so long as it allowed calculations to begin and end on days the faecal samples were collected. The total Ca balance over this period was divided by the number of consecutive days between the faecal sample collection periods to obtain the average daily Ca balance. Daily urinary Ca excretion values used in the balance calculations were corrected for variation.
in the timing of collections and incomplete collections by adjusting 24 h urinary Ca values for daily creatinine excretion (equation 3). Urinary excretion of creatinine remains relatively constant over 24 h, making this metabolite a standard marker of urine sample collection compliance. Ca retention was calculated using both uncorrected and corrected values for urinary Ca excretion. Apparent Ca absorption (equation 4) was determined as the difference between Ca intake and faecal Ca excretion, while net Ca absorption efficiency (equation 5) was calculated as Ca intake minus Ca faecal excretion, divided by Ca intake:

\[
\text{Ca retention} = \text{dietary Ca intake} - \text{urinary Ca excretion},
\]

\[
\text{Corrected 24 h urinary Ca excretion} = \frac{(24 \text{ h urinary Ca (mg))} / (24 \text{ h creatinine (mg))}}{\text{average 24 h creatinine excretion for the balance period (mg))},}
\]

\[
\text{Apparent Ca absorption} = \frac{\text{dietary Ca intake} - \text{faecal Ca excretion}}{\text{dietary Ca intake}},
\]

\[
\text{Net Ca absorption efficiency} = \frac{\text{dietary Ca intake} - \text{faecal Ca (mg)} \times 100}{\text{dietary Ca intake}}.
\]

Faecal sample processing and DNA extraction

Faecal microbial community composition and structure were determined in samples collected at the beginning and end of each session for each participant. After frozen faecal samples were weighed and completely thawed at 4°C, sterilised double-distilled water (twice the weight of the faecal samples) was added to the samples and the samples were homogenised in a stomacher. Faecal slurries were stored at −20°C until DNA extraction. DNA was extracted from 50–100 mg of faecal material using the FastDNA SPIN Kit for Soil (MP Biochemicals) as described previously. DNA quality was checked using a 0.7% agarose gel and a NanoDrop 1000 spectrophotometer (Thermo Scientific), and then DNA was quantified using a NanoDrop 3500 fluorospectrometer (Thermo Scientific).

Determination of microbial community composition using pyrosequencing

The phylogenetic diversity of bacterial communities was determined using 16S ribosomal RNA (rRNA) gene sequences obtained using 454 FLX titanium chemistry and Roche Genome Sequencer (454 Life Sciences-Roche) and primers that amplify the V3–V5 region of the 16S rRNA gene. Multiple samples were run and differentiated using 10bp-tagged forward primers. The initial PCR of faecal sample extracts was carried out using high-fidelity Phusion DNA Polymerase (NEB) and amplicons were gel-purified (QIAEX II Gel Extraction Kit, Qiagen). At the Purdue Genomics facilities, purified amplicons were quantified by fluorometry after staining using the PicoGreen DNA Assay Kit (Invitrogen) and by quantitative PCR, and equimolar amounts were used for 454 FLX titanium chemistry sequencing.

The 16S rRNA gene sequence dataset was analysed using the Quantitative Insights Into Microbial Ecology (QIIME) pipeline (http://qiime.org) that utilises a multi-software approach to perform quality filtering, operational taxonomic unit (OTU) picking, taxonomic assignment, alpha diversity (bacterial diversity within each sample) and beta diversity (bacterial similarities and differences among the samples) measures. With the QIIME pipeline, sequences were split into sample groups using identification tags on the primers and low-quality sequences were removed from the dataset. The quality of the sequences was considered low if the Q score was <25 and length was <400 or >1000bp or ambiguous sequences or mismatches were found in the primer sequence. OTU were picked using the uclust method, and representative OTU sequences were aligned using PyNast (http://biocore.github.io/pynast/) within the QIIME pipeline. ChimeraSlayer, a component of the QIIME pipeline, was used to identify and remove chimeric sequences from the OTU table. Final taxonomic assignments for representative OTU were made using the Greengenes dataset version 4feb2011 (http://greengenes.lbl.gov) (97% sequence similarity) and Ribosomal Database Project (RDP) (http://rdp.cme.msu.edu) classifier at 80% confidence.

Statistical analyses

Statistical analyses were carried out using SAS (version 9.2; SAS Institute, Inc.). The baseline characteristics of the female and male participants were compared using t tests. The Wilcoxon rank-sum test was used to assess differences in non-parametric gastrointestinal symptoms. Pearson’s correlations were used to examine potential associations between the change in fractional Ca absorption in the 24–48 h urine pools (absorption with the SCF treatment minus absorption with the control treatment) and the differences in Ca balance and vitamin D status, baseline anthropometrics, and measures of bone density and strength. A general linear model was used to assess the effect of SCF on fractional Ca absorption. The model controlled for the phase (first v. second 3-week camp session) and order of treatments and accounted for the cross-over design by nesting participant identifier (id) within the order of treatments (random-effect variable). The data were analysed separately for each time period (0–24 and 24–48 h). Similar analyses were carried out for Ca balance. Using published means and standard deviations for fractional Ca absorption reported in adolescent children, we determined that a sample size of twenty-four children would provide sufficient power (80%) to observe a 5-9% difference in fractional Ca absorption assuming an α-error of 0.05 and a standard deviation of 9.6%. P values <0.05 were considered statistically significant for all statistical tests.

Alpha diversity rarefaction analysis was carried out to estimate total sequence coverage of the community. OTU tables were used to perform alpha biodiversity estimations (e.g. Shannon and Chao1 indices), and within-subject diversity was assessed by pairwise comparisons of the CON
v. SCF samples. Beta diversity analysis of the OTU and phylogenetic datasets by ‘Fast UniFrac’ (phylogenetic distance measure) allowed for the comparison of bacterial community composition between the samples(25). Wilcoxon rank-sum tests were used for pairwise comparisons of the proportions of members of bacterial taxa (all taxonomic levels tested) at the end of the CON v. SCF treatments. A P value < 0.05 was used as the cut-off for statistical significance in bacterial genera analyses. Pearson's correlations and Spearman's rank correlations were used to detect associations between the difference in fractional Ca absorption between the treatments (24–48 h) and the difference in the proportions of members of bacterial genera after each treatment. Bacteria in these correlations were those with a genera average value > 0.001 (= 0.1 %), which included the following bacterial taxa: Bifidobacterium; other Coriobacteriaceae; Bacteroides; Barnesiella; Butyrivimonas; Parabacteroides; Prevotella; Alitipes; other Rikenellaceae; Enterococcus; Lactobacillus; other Lactobacillaceae; Streptococcus; Clostridium; Eubacterium; Mogibacterium; Blautia; Anaerostipes; Coprococcus; Dorea; other Lachnospiraceae; Roseburia; other Clostridiales; other Peptostreptococcaceae; Sporacetigenitum; Acetibacter; Butyriviccuc; Faecalibacterium; Oscillibacter; other Ruminococcaceae; Ruminococcus; Subdoligranulum; Dialister; other Clostridia; Catenibacterium; Coprococcus; other Erysipelotrichaceae; Turicibacter; other Firmicutes; other bacteria; Escherichia/ Shigella; Pseudomonas; Actinomyces; other Streptococcaceae; Anaerofisit; Anaerococcus.

Results

A total of twenty-four adolescents (nine girls and fifteen boys) participated in the present study during the summer of 2010. Among these, three participants did not undergo the fractional Ca absorption test in both sessions and one participant attended only one of the camp sessions. Therefore, twenty-one and twenty-three participants were included in the fractional Ca absorption and Ca balance analyses, respectively. Analyses of bone biomarkers and bacterial community proportions included twenty-three participants. The participants evaluated in the present study were ethnically diverse with a distribution of eleven Asian, six Hispanic, one Black and six multi-racial (other) teenagers. The characteristics of the participants including age, anthropometrics, physical parameters and bone measures are reported as means and standard deviations for girls and boys separately in Table 1. Girls and boys had similar physical characteristics; statistically significant differences were observed only in the percentage of lean mass (lower in girls than in boys, P= 0.009) and the percentage of fat mass (higher in girls than in boys, P=0.01) in the study cohort. The mean habitual intakes of Ca and total fibre and soluble fibre were 768 (SD 403) mg/d and 12 (SD 4) g/d and 3-4 (SD 1-1) g/d, respectively.

Gastrointestinal symptoms

No significant differences were observed in gastrointestinal symptom severity between participants on the SCF treatment and those on the CON treatment over the 18d observation period. Self-reported mean daily gastrointestinal symptom scores during the SCF and CON treatments were 0.1 (SD 0.3) v. 0.3 (SD 0.8) for bloating, 0.6 (SD 0.8) v. 1.1 (SD 1.7) for flatulence, 0.3 (SD 0.5) v. 0.4 (SD 0.7) for abdominal cramping, and 0.2 (SD 0.7) v. 0.3 (SD 0.9) for stomach noises.

Fractional calcium absorption, calcium balance and bone biomarkers

Fractional Ca absorption was based on the analysis of isotopes excreted in the urine, which was collected between 0 and 24 and between 24 and 48 h after oral and intravenous isotope administration (Fig. 1). Compared with that observed after the CON treatment, mean fractional Ca absorption did not differ during the first 24 h, but was significantly higher after the SCF treatment (0.595 (SD 0.142) v. 0.664 (SD 0.129), respectively, P=0.02) at 24–48 h. The difference in mean fractional Ca absorption of 0.069 occurring 24–48 h after the treatment represented an 11.6 % increase in absorption with the SCF treatment. A general linear model identified significant effects of treatment on fractional Ca absorption at 24–48 h (P=0.02), but not at 0–24 h (P=0.09). There was no significant effect of treatment order on fractional Ca absorption. Although carry-over effects may have resulted from the 1-week washout period being too short for gut microbiota proportions to return to baseline levels, this finding suggests that the occurrence of carry-over effects was unlikely. There were no significant effects of treatment on net Ca absorption, net Ca absorption efficiency, faecal Ca excretion or Ca retention (Table 2). Values obtained for faecal sample collection compliance, as measured by polyethylene glycol recovery, were 76.0 (SD 12.9) and 78.4 (SD 15.8) % for the SCF and CON treatments, respectively.

The concentrations of bone turnover markers (Table 3) were measured to explore possible intervention effects on bone turnover. When compared as changes from the baseline, no differences in the concentrations of serum alkaline phosphatase, P, Ca, parathyroid hormone 1–84, leptin,

| Table 1. Baseline characteristics of the participants (Mean values and standard deviations) |
|---------------------------------|------|------|
| Females (n 9) | Males (n 15) |
| Age (years) | 13.3 (SD 1.0) | 13.5 (SD 0.9) |
| Weight (kg) | 59.9 (SD 13.2) | 61.1 (SD 11.8) |
| Height (cm) | 157.3 (SD 4.9) | 164.9 (SD 8.2) |
| BMI (kg/m²) | 24.1 (SD 4.0) | 22.4 (SD 3.1) |
| BMI percentile (%) | 80.0 (SD 16.4) | 74.7 (SD 19.2) |
| Tanner score average | 3.8 (SD 0.7) | 2.4** (SD 0.9) |
| Lean mass (%) | 61.2 (SD 5.2) | 71.1** (SD 9.5) |
| Fat mass (%) | 35.1 (SD 5.6) | 25.1* (SD 10.0) |
| Total body BMD (g/cm²) | 1.07 (SD 0.11) | 1.04 (SD 0.11) |
| Total body BMC (g) | 2115 (SD 329) | 2316 (SD 424) |
| Total spine BMD (g/cm²) | 1.09 (SD 0.13) | 1.04 (SD 0.14) |
| Femoral neck BMD (g/cm²) | 1.03 (SD 0.18) | 1.05 (SD 0.15) |

BMD, bone mineral density; BMC, bone mineral content. Mean value was significantly different from that of the females: *P< 0.05; **P< 0.01 (t test).
insulin-like growth factor, IGFBP3 and sclerostin and urinary NTX, Ca and P were observed after the SCF treatment.

The mean 25-hydroxyvitamin D status values after the SCF and CON treatments were 65·2 (SD 18·8) and 59·1 (SD 15·9) nM, respectively, which were not significantly different. No significant associations were observed between differences in vitamin D status and those in fractional Ca absorption (between 24 and 48 h) or net Ca absorption efficiency.

The anthropometric parameters and dietary intake variables of the participants that might influence the effect of SCF on lower gut Ca absorption were evaluated. The difference in fractional Ca absorption in the 24–48 h urine collections between participants on the SCF treatment and those on the CON treatment was not correlated with height ($r = 0·112$, $P = 0·63$), body surface area ($r = 0·012$, $P = 0·96$), weight ($r = 0·022$, $P = 0·92$), habitual dietary fibre ($r = 0·150$, $P = 0·54$) and Ca ($r = 0·012$, $P = 0·96$) intakes, Tanner stage ($r = 0·131$, $P = 0·57$) or BMI ($r = 0·074$, $P = 0·75$).

**Changes in bacterial community composition**

Roche 454-Titanium pyrosequencing of forty-six faecal samples yielded a total of 1 793 821 16S rRNA sequences with an average of 19 498 (SD 7126) (range: 8211–41 212) sequences per sample. The number of sequences generated for each participant did not differ significantly when compared by treatment ($P > 0·05$) or the time of faecal collection (baseline v. end samples) ($P > 0·05$). Firmicutes was the most dominant phylum with an average of 89·4% followed by the phyla Bacteroidetes (5·1%), Actinobacteria (4·9%) and Proteobacteria (0·5%) (Fig. 2). Detailed comparisons of microbial community composition in these adolescents have been reported elsewhere$^{26}$, and herein we only report differences in community composition after the SCF treatment.

At the phylum level, regardless of the inclusion of SCF in the clinical diet, the average relative proportion of members of the phylum Bacteroidetes significantly increased and that of members of the phylum Firmicutes decreased by the end of each camp session (Fig. 2). The community composition of participants on the SCF treatment was significantly different from that of participants on the CON treatment at the family level (Fig. 3). There were significantly higher proportions of Porphyromonadaceae ($P = 0·02$) and other Clostridiales ($P = 0·009$) and significantly lower proportions of Peptostreptococcaceae ($P = 0·04$) after the SCF treatment. At the end of each 3-week camp session, there were four genera and three ‘other’ groups that had average proportions that differed significantly ($P < 0·05$) between participants on the SCF treatment and those on the CON treatment (Table 4).

Table 2. Effect of soluble maize fibre (SCF) treatment on calcium absorption and retention in adolescent boys and girls*  
(Mean values and standard deviations)

<table>
<thead>
<tr>
<th></th>
<th>SCF</th>
<th>CON</th>
<th>P</th>
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</thead>
<tbody>
<tr>
<td>Stable isotope analysis†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fractional Ca absorption (0–24 h)</td>
<td>0·522 0·110</td>
<td>0·497 0·108</td>
<td>0·09</td>
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<tr>
<td>Fractional Ca absorption (24–48 h)</td>
<td>0·664 0·129</td>
<td>0·595 0·142</td>
<td>0·02</td>
</tr>
<tr>
<td>Balance analysis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca intake (mg/d)</td>
<td>606 29</td>
<td>604 25</td>
<td>0·69</td>
</tr>
<tr>
<td>Urinary Ca (mg/d)</td>
<td>77 56</td>
<td>65 36</td>
<td>0·11</td>
</tr>
<tr>
<td>Faecal Ca (mg/d)</td>
<td>318 108</td>
<td>312 106</td>
<td>0·77</td>
</tr>
<tr>
<td>Net Ca absorption efficiency (%)</td>
<td>47 18</td>
<td>48 17</td>
<td>0·75</td>
</tr>
<tr>
<td>Ca retention (mg/d)</td>
<td>212 117</td>
<td>227 101</td>
<td>0·42</td>
</tr>
</tbody>
</table>

CON, control.  
* General linear model that includes treatment, order and phase.  
† For fractional Ca absorption calculations, $n = 21$ (three subjects did not undergo both fractional Ca absorption tests and were, therefore, excluded from the analysis).  
‡ For balance calculations, $n = 23$ (one subject completed only one treatment phase).
Table 3. Serum and urinary bone turnover marker concentrations at baseline and after treatment with 0 and 12 g soluble maize fibre (SCF)*

(Mean values and standard deviations, n 23)

<table>
<thead>
<tr>
<th>Bone biomarkers</th>
<th>Baseline</th>
<th>SCF</th>
<th>CON</th>
</tr>
</thead>
<tbody>
<tr>
<td>Serum</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alkaline phosphatase (U/l)</td>
<td>237·29 125·80</td>
<td>235·57 135·45</td>
<td>235·00 132·5 0·50</td>
</tr>
<tr>
<td>Ca (ng/l)</td>
<td>105·1</td>
<td>4·1</td>
<td>102·3 4·0</td>
</tr>
<tr>
<td>Creatinine (ng/l)</td>
<td>8·5</td>
<td>1·1</td>
<td>8·5 1·2</td>
</tr>
<tr>
<td>P (ng/l)</td>
<td>48·6</td>
<td>5·4</td>
<td>47·6 5·9</td>
</tr>
<tr>
<td>PTH (pg/ml)</td>
<td>18·20</td>
<td>7·34</td>
<td>21·23 11·27</td>
</tr>
<tr>
<td>Leptin (ng/ml)</td>
<td>10·17</td>
<td>7·77</td>
<td>8·65 7·61</td>
</tr>
<tr>
<td>IGF-1 (ng/ml)</td>
<td>296·53</td>
<td>53·97</td>
<td>266·81 52·30</td>
</tr>
<tr>
<td>IGFBP3 (ng/ml)</td>
<td>3879·88 554·63</td>
<td>3648·99 572·71</td>
<td>3651·26 560·23 0·46</td>
</tr>
<tr>
<td>Sclerostin (ng/ml)</td>
<td>0·45</td>
<td>0·23</td>
<td>0·41 0·17</td>
</tr>
<tr>
<td>Urine</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NTX (mmol BCE/l)</td>
<td>2973·47 1788·00</td>
<td>5378·81 4632·47</td>
<td>5323·75 4538·67 0·76</td>
</tr>
<tr>
<td>Ca (mg/l)</td>
<td>39·1</td>
<td>27·9</td>
<td>34·3 25·1</td>
</tr>
<tr>
<td>P (mg/l)</td>
<td>522·7</td>
<td>205·1</td>
<td>778·5 399·7</td>
</tr>
<tr>
<td>Creatinine (mg/l)</td>
<td>586·9</td>
<td>218·5</td>
<td>903·0 457·2</td>
</tr>
</tbody>
</table>

CON, control; PTH, parathyroid hormone 1–84; IGF-1, insulin-like growth factor 1; IGFBP3, insulin-like growth factor-binding protein 3; NTX, cross-linked N-telopeptides of type I collagen; BCE, bone collagen equivalents.

* One participant did not complete both treatment phases.

† Significance is based on differences from the baseline (SCF minus baseline v. CON minus baseline).

Significant increases in proportions after the SCF treatment were observed for the genera Parabacteroides (P < 0·003), other Clostridiales (P = 0·04) and other Ruminococcaceae (P < 0·03), while significant decreases in proportions were observed for Enterococcus (P < 0·03), Anaerofastis (P < 0·05), Coprococcus (P < 0·03) and other Peptostreptococcaceae (P = 0·002). There was also an increase in the proportions of Bifidobacterium, Alistipes, Anaerococcus, Catenibacterium and other Clostridia, but the increase in the proportions was not significant. Similarly, decreases in the proportions of Robha, other Streptococcaceae, Clostridium, Sporacetigenium, Turicibacter and other TM7 genera Incertae sedis were observed after the SCF treatment, but the difference in community proportion after the CON treatment was not significant. Fast UniFrac with jackknife analysis revealed no differences in community structure due to the overall treatment, despite the data being processed using different classification criteria (e.g. OTU v. phylogeny) (data not shown).

Correlations between the proportions of members of bacterial genera and fractional calcium absorption

In an exploratory analysis, Pearson’s correlations and Spearman rank-sum analyses were used to identify bacteria having potential influence on the Ca absorption response. Changes (those with the SCF treatment minus those with the CON treatment) in fractional Ca absorption measured in the 24–48 h urine collections were negatively correlated (decreases in the proportions of members of bacterial genera as Ca absorption with the SCF treatment increased) with the proportions of Actinomyces, Pseudomonas of the phylum Actinobacteria and other Erysipelotrichaceae of the phylum Firmicutes. Conversely, changes in fractional Ca absorption were positively correlated (increases in the proportions of members of bacterial genera as Ca absorption with the SCF treatment increased) with the proportions of the Bacteroidetes member Bacteroides as well as Butyrivibrio, Oscillibacter and Dialister of the phylum Firmicutes (Table 5).

Fig. 2. Comparison of the average relative proportions of members of bacterial phyla in participants at the beginning (B) and end (E) of the clinical sessions where diets included soluble maize fibre (SCF) v. control (CON). Only phyla representing > 0·5 % of the total community during at least one treatment are depicted. Values are means, with their standard errors represented by vertical bars. * Mean value was significantly different from that of the B sample within each phylum (P < 0·05). Significant differences in bacterial phyla between participants on the SCF treatment and those on the CON treatment at E were not apparent.
Discussion

Significant increases in fractional Ca absorption were observed in adolescent boys and girls consuming SCF. Broadly, the consumption of SCF resulted in lower proportions of faecal bacteria of the phylum Firmicutes and greater proportions of bacteria of the phylum Bacteroidetes. These changes may influence bone health as increases in fractional Ca absorption were negatively correlated with the proportions of members of the genera *Actinomyces* and *Pseudomonas* and positively correlated with those of the genera *Bacteroides*, *Butyricicoccus*, *Oscillibacter* and *Dialister*. Despite the evidence for an acute increase in fractional Ca absorption via faecal microbiota, no effect of SCF on Ca balance or biochemical markers of bone turnover was observed.

In the present study, daily consumption of 12 g SCF for 21 d by adolescent girls and boys increased fractional Ca absorption by approximately 12%. This effect was similar to that observed in a previous study reporting a 10.8% difference in true Ca absorption in adolescent boys consuming 15 g/d inulin–oligofructose for 9 d compared with the control group17. Using dual-stable isotope methods, increases in fractional Ca absorption relative to those observed after control treatments were found to range from 3 to 18% in young girls and boys consuming 8 g/d inulin–oligofructose mix (Synergy 19). In the present study, habitual fibre intake, specifically of soluble fibre, was low. This may have influenced the response to SCF intervention observed in these teenagers, as positive effects of inulin-type fructans on Ca absorption were only observed when the habitual basal diet lacked fermentable fibres28.

The 12% increase in fractional Ca absorption observed in the present study occurred between 24 and 48 h as the effect was significant when measuring the second 24 h urine pool (24–48 h) after the administration of stable Ca isotopes. No significant difference was observed in isotope enrichment in the urine samples collected during the first 24 h. This finding is supported by the literature that suggests that microbial involvement and lower gut absorption are not realised until 24 h after isotope administration17. Other studies have not measured fractional Ca absorption at various time intervals, as has been done in the present study, to evaluate the effects of prebiotics on Ca absorption over time.

Urinary Ca excretion was non-significantly 18% higher after the SCF treatment compared with that after the CON treatment, reflective of a greater fractional Ca absorption efficiency. An increase in urinary Ca excretion could contribute to the lack of a beneficial effect of SCF on net Ca absorption and net Ca balance. However, because the increase in urinary Ca excretion was not significant, it is more likely that differences in methods explain the inconsistencies observed in the benefits conferred by SCF. Fractional Ca absorption reveals
small changes in isotopic tracer levels over a short period of time and Ca balance is a more crude measure of total Ca content over a longer period of time. We powered the study for fractional Ca absorption, a more sensitive measure than Ca retention, with respect to measurement variability compared with Ca retention and net Ca absorption. For example, faecal Ca excretion is highly variable, requiring a sample size of thirty-four subjects to detect a 61 mg difference in Ca retention with an α-error of 0.05, 80% power and a standard deviation of the difference of 122 mg/d between the participants. Despite the lack of power to detect an effect on Ca retention, it is possible that the SCF treatment exerted an effect on bone strength. Weaver et al.\(^\text{(12)}\) reported an increase in the peak breaking force of the femur, but no change in retention in rats consuming 4% SCF by weight. These two methods have a large difference in the percentage of standard deviation, i.e. 9.7% for breaking force and 41.3% for Ca retention. This change in bone strength suggests that the SCF intervention perturbed Ca balance in order to affect bone balance.

Consistent with a beneficial effect of fibre consumption on bone, in a long-term study carried out in adolescent boys and girls, consumption of 8 g/d of an inulin-type fructan mixture was found to result in a significantly increased BMD and a persistent effect on fractional Ca absorption from 8 weeks (8.5%) to 1 year (5.9%) of daily consumption with no deviation, i.e. 9.7% for breaking force and 41.3% for Ca retention. This change in bone strength suggests that the SCF intervention perturbed Ca balance in order to affect bone balance.

Table 5. Correlations* between calcium absorption and proportions of bacterial genera that may affect lower gut mechanisms (n 21)†

<table>
<thead>
<tr>
<th>Generation/Phylum</th>
<th>Correlation coefficient (P value)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Actinomyces</td>
<td>-0.483 (0.027)</td>
</tr>
<tr>
<td>Pseudomonas</td>
<td>-0.473 (0.003)</td>
</tr>
<tr>
<td>Butyricimonas</td>
<td>-0.464 (0.039)</td>
</tr>
<tr>
<td>Oscillibacter</td>
<td>-0.565 (0.008)</td>
</tr>
<tr>
<td>Diaster</td>
<td>0.619 (0.003)</td>
</tr>
<tr>
<td>Other Erysipelotrichaceae</td>
<td>-0.463 (0.034)</td>
</tr>
</tbody>
</table>

* Pearson's correlations and Spearman's rank correlations. † Data are shown for correlations that were significant (P < 0.05), while analysis included correlations for all genera making up >0.001 (=0.1%), which included the following bacterial taxa: Bifidobacterium; other Coriobacteriaceae; Bacteroides; Banesiella; Butyrinimonas; Parabacteroides; Prevotella; Alistipes; other Rikenellaceae; Enterococcus; Lactobacillus; other Lactobacillaceae; Streptococcus; Clostridium; Eubacterium; Megasphaera; Blautia; Anaerostipes; Capnocytophaga; Dorea; other Lachnospiraceae; Roseburia; other Clostridiales; other Peptostreptococcaceae; Sporoclostridium; Arthrobacter; Butyricoccus; Faecalibacterium; Oscillibacter; other Ruminococcaceae; Ruminococcus; Subdoligranum; Diaster; other Clostridiales; Catenibacterium; Coprobacillus; other Erysipelotrichaceae; Turibacter; other Firmicutes; other bacteria; Escherichia/Shigella; Pseudomonas; Actinomyces; other Streptococcaceae; Anaerofustus; Anaerococcus.
towards a greater increase in the proportion of *Bifidobacterium* suggest that the inclusion of SCF in the diet increased the proportion of beneficial bacteria capable of fermenting RS and producing SCFA. The interesting finding that the proportions of members of the phylum Bacteroidetes increased significantly on both the SCF and CON treatments was more likely due to a response to the control diet rather than to a carry-over effect of the SCF treatment. This is supported by the lack of a significant order effect on study outcomes and is further supported by the control diet providing more dietary fibre than the participants’ habitual diets. Significant increases in the proportions of members of the family Bacteroidaceae have been reported in adult men after the consumption of SCF as a part of their regular diet, but curiously in this particular study the proportion of *Bifidobacterium* significantly decreased compared with that in the control group. Conversely, healthy adults consuming four increasing doses of galacto-oligosaccharides (0–10 g) were found to exhibit dose-dependent increases in the proportion of bifidobacteria after 3 weeks of consumption. In addition to the previously mentioned genera, changes in the proportions of *Butyrivibrio, Oscillibacter, Dialister* and *Dialister* were also significantly correlated with increased Ca absorption in the present study. These genera have previously been identified for their production of butyrate, a short-chain fatty acid critical for maintaining gut health and immune function. However, SCF did not alter net Ca retention. Therefore, the present study cannot resolve whether the balance method is too insensitive to determine the benefits of improved Ca absorption or whether the benefits decline with chronic feeding. Further work is required to assess whether SCF influences bone mineralisation in the long term and how microbial interplay modulates the individual response to prebiotic treatments.

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C. M. Weaver serves on the Advisory Board of Pharmavite.

The authors’ contributions are as follows: C. M. Weaver, C. H. N. and B. R. M. designed the research; B. R. M., C. H. N. and C. M. Whisner conducted the research; C. M. Whisner, C. H. N., L. D. M. and G. P. M. analysed the data; C. M. Whisner, C. H. N. and C. M. Weaver wrote the paper; C. M. Weaver had primary responsibility for the final content. All authors read and approved the final manuscript.

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