Non-digestible oligosaccharides and calcium absorption in girls with adequate calcium intakes†

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Non-digestible oligosaccharides such as inulin and oligofructose have been shown to consistently increase calcium absorption in experimental animals, but data in humans are less clear-cut. The objective of this study was to assess the effect of 8 g/d of oligofructose or a mixture of inulin and oligofructose on calcium absorption in girls at or near menarche. A total of fifty-nine subjects were studied using a balanced, randomized, cross-over design. They received, in random order, 8 g/d placebo (sucrose), oligofructose or the mixture inulin+oligofructose for 3 weeks, separated by a 2-week washout period. Throughout the study, subjects consumed a total of approximately 1500 mg/d dietary calcium, by adding two glasses of calcium-fortified orange juice to their diet. Four grams of placebo, oligofructose or the mixture inulin+oligofructose was added to each glass of orange juice immediately before it was consumed. At the end of each 3-week adaptation period, calcium absorption was measured, using a dual stable isotope technique, from the cumulative fractional excretion of an oral and an intravenous tracer over 48 hours. Calcium absorption was significantly higher in the group receiving the inulin+oligofructose mixture than in the placebo group (38.2 ± 9.8% vs. 32.3 ± 9.8%; P < 0.01), but no significant difference was seen between the oligofructose group and the placebo group (31.8 ± 9.3% vs. 31.8 ± 10.0%; P = NS). We conclude that modest intakes of an inulin+oligofructose mixture increases calcium absorption in girls at or near menarche.

Calcium absorption: Non-digestible oligosaccharides: Oligofructose: Inulin: Prebiotics: Stable isotope

Introduction

Maintenance of an adequate calcium intake at or near puberty is essential for the development of optimal peak bone mass (Matkovic, 1992) An adequate calcium intake during this pivotal time period is vital for optimum bone mineral accretion (Chan, 1991; Nieves et al. 1995). Interventions aimed at preventing the morbidity and mortality associated with osteoporosis may, therefore, best be aimed at this vulnerable age group (Matkovic, 1992). Despite the importance of an adequate calcium intake during this period, self-selected diets of children during this stage typically provide insufficient calcium (Eck & Hackett-Renner, 1992), leading to inadequate calcium retention (Abrams & Stuff, 1994).

There has been increasing interest in the effect of prebiotic non-digestible oligosaccharides as modifiers of mineral absorption in animals (Brommage et al. 1993; Delzenne et al. 1995; Rémesy et al. 1993) and humans (Roberfroid, 1999; Van Loo et al. 1999). These compounds resist digestion by human alimentary enzymes, and undergo fermentation in the large intestine (Van Loo et al. 1999). It is speculated that the volatile short-chain fatty acids produced by this fermentation lower cecal pH, increase calcium concentration in the liquid phase of the cecal contents, and increase colonic absorption of calcium (Greger, 1999; Schulz et al. 1993). They may also have trophic effects in the gastrointestinal tract, which may increase calcium absorption either in the colon or throughout the entire gut (Chonan & Watanuki, 1995; Greger, 1999; Rémesy et al. 1993).

The most widely studied non-digestible oligosaccharides...
in humans are inulin and oligofructose, which are naturally occurring components of the Western diet (Van Loo et al. 1995; Moshfegh et al. 1999). Human studies examining their effect on calcium absorption have been contradictory. Two studies have shown a beneficial effect of 40 g/d inulin (Coudray et al. 1997) or 15 g/d oligofructose (van den Heuvel et al. 1999) on calcium absorption in young men and adolescent boys on modest calcium intakes. A third study, however, found no effect of 15 g/d of inulin, oligofructose or galacto-oligosaccharides on calcium absorption in a similar population (van den Heuvel et al. 1998). These discrepant findings may be due to methodological differences (Coudray & Fairweather-Tait, 1998) or due to a type II error, as all the studies were relatively small.

All three previous studies involved male subjects on moderate calcium intakes (approximately 800 mg/d), where calcium retention can be increased significantly by increasing calcium intake (Matkovic & Heaney, 1992). In children at or near puberty calcium retention appears to reach a maximum at an intake of about 1200 mg/d; calcium intakes above this threshold do not seem to lead to increased calcium retention (Matkovic & Heaney, 1992). It is not known whether non-digestible oligosaccharides will increase calcium absorption at higher levels of calcium intake. Further, because previous studies have examined the effect of relatively large doses of these ingredients on calcium absorption, it is unclear whether significant increases in calcium absorption can be achieved at lower intakes.

The object of this study was, therefore, to examine the effect of relatively modest intakes of oligofructose and a mixture of oligofructose and inulin, on calcium absorption in a large sample of girls at or near menarche, with calcium intakes approximating the currently recommended dietary intake.

Materials and methods

Sixty healthy girls, 11.0–13.9 years of age, were recruited from the greater Houston area by public advertisement. Subjects were interviewed by a pediatric dietician prior to enrolment, and calcium intake was estimated using a food frequency questionnaire. Only subjects with a habitual calcium intake between 500 and 1400 mg/d were considered eligible for the study. Subjects were excluded from the study if they had chronic gastrointestinal disease, renal failure, or disorders of calcium homeostasis; were taking prescription medications (including oral contraceptives); smoked; or weighed more than the ninetieth percentile for age.

Subjects were studied using a randomized, double-blind, cross-over design. Subjects were randomized to receive two 4-g servings of non-digestible oligosaccharides daily for 3 weeks and two 4-g servings of placebo daily for 3 weeks, separated by a 2-week washout period. The oligosaccharides and placebo were given in random order, and investigators were blinded to the treatment assignment. Two similar protocols were carried out simultaneously. In Protocol I, the non-digestible oligosaccharide used was chicory oligofructose (Rafitlose® P95, Orafti, Tienen, Belgium); Protocol II used an inulin+oligofructose mixture (Rafitlose® Synergy1, Orafti, Tienen, Belgium). In both protocols, the placebo (sucrose) was packaged and presented in a manner that was identical to that of the oligosaccharides.

Subjects were instructed to maintain a calcium intake aimed at providing a total of approximately 1200–1300 mg/d during the study period by consuming an 8-ounce glass of calcium-fortified orange juice with breakfast and with the evening meal, and an 8-ounce glass of milk, calcium-fortified orange juice or a serving of yogurt with the midday meal. The placebo (sucrose), oligofructose or the inulin+oligofructose mixture were provided as packets containing 4 g of carbohydrates. One packet was added to both the morning and evening glasses of orange juice and gently mixed until it dissolved completely. The same dietary regime was continued during the 2-week washout period, but without oligosaccharides or placebo added to the glasses of orange juice. During the third week of each study period, subjects kept a weighed dietary record involving two weekdays and one weekend day. Calcium intake was calculated from these records by a pediatric dietician using the Nutrition Data System for Research (version 4.02, University of Minnesota, Minneapolis, MN).

At the end of each 3-week study period, the subjects were admitted to the Metabolic Research Unit of the Children’s Nutrition Research Center. Calcium absorption was measured using a double stable isotope method. All used and unused packets of the placebo, oligofructose or the inulin+oligofructose mixture were returned, and compliance was measured by a packet count.

Measurement of calcium absorption

Calcium absorption was measured using a modification of a previously validated dual-isotope methodology (Yergey et al. 1994; Yergey et al. 1990). On the morning of the study, subjects were admitted to the Metabolic Research Unit of the Children’s Nutrition Research Center, and a baseline urine sample was collected. Subjects consumed a low-calcium breakfast and an 8-ounce glass of calcium-fortified orange juice to which was added one packet of placebo, oligofructose or the inulin+oligofructose mixture and 10 µg of 46Ca. The oligosaccharide was added immediately prior to consumption of the drink, and the 46Ca 18–24 h earlier. Immediately after breakfast, 1.5 mg of 42Ca was infused intravenously over 3–5 min. The midday meal contained approximately 400 mg calcium in the form of calcium-fortified orange juice, milk or yogurt. The evening meal contained another serving of calcium-fortified orange juice, 10 µg of 46Ca, and a packet of oligosaccharide or placebo. Immediately after the first dose of calcium isotopes was administered, a complete 48 h urine collection was begun. After the evening meal, subjects were discharged to their homes, where they completed the urine collection.

Stable isotopes were purchased from Oak Ridge National Laboratories (Oak Ridge, TN) as the carbonate salts. Aqueous solutions were prepared as previously described (Eastell et al. 1989) by the Pharmacy Department of Texas Children’s Hospital, and tested for sterility.
Statistical analysis

Based on previous studies, we had anticipated a mean calcium absorption of approximately 30%, with a standard deviation of 8%. We considered a 5% difference in calcium absorption to be the smallest clinically significant difference, so thirty subjects were required to have a 90% power (\(\beta = 0.10\)) to detect this difference at a statistical significance of \(P<0.05\) (\(\alpha = 0.05\)).

The effect of oligofructose or the inulin+oligofructose mixture on calcium absorption was assessed using paired \(t\)-tests, using StatView v4.5.1 for Macintosh (Abacus Concepts Inc., Berkeley, CA). Statistical significance was taken at \(P<0.05\). All data are presented as mean (standard deviation) unless otherwise stated.

Ethical considerations

The study received ethical approval from the Institutional Review Board of Baylor College of Medicine. Informed written consent was obtained from the subjects (where age-appropriate) and their parents.

| Table 1. Demographic data for the subjects in Protocol I (placebo v. oligofructose) and Protocol II (placebo v. Synergy1) |
|---|---|---|
| **Number** | Protocol I | Protocol II | \(P\)-value‡ |
| **Age (years)** | 30 | 29 | 0.12 |
| **Weight (kg)** | 12.1 (0.7) | 11.8 (0.8) | 0.14 |
| **Height (cm)** | 42 (9) | 46 (9) | 0.24 |
| **Ethnicity (%)†** | 152 (8) | 154 (6) | 0.53§ |
| **African-American** | 27% | 7% | 0.08§ |
| **Hispanic** | 13% | 28% | |
| **Caucasian** | 57% | 52% | |
| **Asian/others** | 3% | 14% | |
| **Pubertal status** | | | |
| **Premenarcheal (%)** | 67% | 59% | 0.53§ |

* Mean (SD).
† Totals may not equal 100, due to rounding.
‡ Unpaired \(t\)-test, unless otherwise stated.
§ Chi-squared test.

Results

Of the sixty subjects recruited, a total of fifty-nine completed the study. One subject in Protocol II defaulted from the second visit, and it was not possible to re-arrange her admission, so she was excluded from analysis. This was not related to intolerance of the study product. Demographics of the populations for the two protocols were similar (Table 1). The majority of subjects were Caucasian and premenarcheal.

Compliance with non-digestible oligosaccharide

Compliance was good, with a mean of 95.4% of the expected number of packets of the mixture inulin+oligofructose taken (SD 7.3%, range 71%–100%, median 100%), and 95.2% of the expected number of packets of oligofructose taken (SD 7.9%, range 68%–100%, median 100%). All but four subjects achieved a compliance rate of at least 80% of the expected number of packets of oligosaccharides (two in Protocol I, and two in Protocol II). There was no difference in compliance between oligofructose and of the inulin+oligofructose mixture (unpaired \(t\)-test, \(P=0.59\)).

Calcium intake

There were no significant differences in calcium intake between the placebo and the oligofructose or the inulin+oligofructose mixture phases of the study, either for Protocol I (oligofructose 1524 (265) mg/d vs. placebo 1611 (326) mg/d; \(P=0.34\)) or Protocol II (inulin+oligofructose mixture 1525 (282) mg/d vs. placebo 1495 (280) mg/d; \(P=0.50\)).

Calcium absorption

In Protocol I, there was no significant difference in fractional calcium absorption on placebo (31.8 (9.3%)) or on oligofructose (31.8 (10.0%); \(P=0.75\)). In Protocol II, calcium absorption on the inulin+oligofructose mixture was 38.2 (9.8%), significantly greater than on placebo (32.3 (9.8%), \(P=0.007\)). Calcium absorption during the placebo period did not differ significantly between Protocol I and Protocol II (\(P=0.97\)). The absolute calcium absorption (the product of fractional calcium absorption and dietary calcium intake) was significantly increased by the inulin+oligofructose mixture (575 (148) mg/d vs. 485 (154) mg/d, \(P=0.004\)), but not by oligofructose (489 (169) mg/d vs. 490 (153) mg/d; \(P=0.99\)), in comparison with placebo.

Urinary calcium excretion

Calcium excretion during the 48 h study period was variable, and did not differ significantly between groups. In Protocol I, calcium excretion was 71 (48) mg/d during the placebo period and 79 (50) mg/d during the oligofructose period (\(P=0.26\)). In Protocol II, the corresponding values were 65 (54) mg/d for placebo and 71 (50) mg/d for the inulin+oligofructose mixture (\(P=0.92\)).
Estimated calcium balance

Net calcium balance was estimated by subtracting measured urinary calcium excretion and an estimate of endogenous fecal excretion, 1.4 mg/kg (Abrams et al. 1991) from the product of the fractional calcium absorption and the dietary calcium intake. This was similar during the sucrose and oligofructose periods of Protocol I (430.7 (153.4) mg/d v. 438.5 (169.0) mg/d; \( P \)-value=0.87). In Protocol II, estimated calcium balance was significantly greater during the inulin+oligofructose mixture period than the sucrose period (511.4 (151.4) mg/d v. 421.1 (154.0) mg/d; \( P \)-value=0.004).

Discussion

In this study, we examined the effect of 21 days’ adaptation to modest amounts of two non-digestible oligosaccharides, oligofructose and an inulin+oligofructose mixture, on calcium absorption in girls at or near puberty who were consuming a diet containing the recommended dietary allowance of calcium. Calcium absorption, measured using a dual stable isotope method, was significantly higher when subjects consumed 8 g/d of the inulin+oligofructose mixture than whilst consuming placebo. No significant benefit was seen from 8 g/d of oligofructose.

The dual stable isotope method of evaluating calcium absorption has been discussed in detail elsewhere (Yergey et al. 1990; Yergey et al. 1994); and a 24 h urine collection is adequate to allow accurate measurement of calcium absorption (Yergey et al. 1994). Indeed, in some populations, as brief a collection as 12 h may be adequate (Hillman et al. 1988). In normal circumstances, very little calcium absorption occurs in the colon (Hillman et al. 1988); however, oligosaccharides may significantly increase colonic absorption of calcium (Van Loo et al. 1999). It has been argued that a longer urine collection is required to capture this late, colonic, phase of absorption (Barger-Lux et al. 1989; van den Heuvel et al. 1998). For this reason, we extended the urine collection in our study to 48 h after administration of the intravenous tracer, or at least 36 h after administration of the second dose of oral calcium tracer.

Three studies had evaluated the effects of different oligosaccharides on calcium absorption in humans, with conflicting results. Coudray et al. (1997) showed in a study of nine men that 40 g/d of inulin significantly increased calcium absorption, measured by a metabolic balance, from 21.3 % (SD 12.5 %) to 33.7 % (SD 12.1 %). Van den Heuvel et al. (1998), however, found no effect of 15 g/d inulin, oligofructose or galacto-oligosaccharides on calcium absorption in young men, using a dual-isotope tracer method. One criticism of this study was that urine was only collected for 24 h, potentially missing the late colonic phase of absorption (Courdary & Fairweather-Tait, 1998). Indeed, a subsequent study by the same group (van den Heuvel et al. 1999), using a 36 h urine collection, showed that 15 g/d oligofructose significantly increased calcium absorption from 47.8 % (SD 16-4) to 60.1 % (SD 17-2). Our data show a significant increase in calcium absorption in response to the consumption of 8 g/d of an inulin+oligofructose mixture, but no beneficial effect of 8 g/d of oligofructose. This suggests that this mixture may be a more potent promoter of calcium absorption than oligofructose.

The present study differs from previous studies in a number of important aspects. All previous studies have been very small, involving only nine to twelve male subjects, and used rather high intakes of oligosaccharides (15–40 g/d) (Coudray et al. 1997; van den Heuvel et al. 1998; van den Heuvel et al. 1999). The dose of oligosaccharides used in the current study was only 8 g/d. This compares to a typical dietary intake of 2.6 g/d of inulin and 2.5 g/d of oligofructose in the Western diet (Moshfegh et al. 1999), and well below the amount of oligosaccharides that may cause abdominal symptoms (Briet et al. 1995).

Previous studies have evaluated subjects with calcium intakes in the order of 800 mg/d, well below the RDA for this age group of 1300 mg/d (Institute of Medicine Food and Nutrition Board’s Standing Committee on the Scientific Evaluation of Dietary Intervals, 1997). Work by Matkovic & Heaney (1992) has suggested that during the stage of life approaching and at the achievement of puberty, net calcium balance increases with increasing calcium intake, to a maximum of about 1200 mg/d. Beyond this threshold, further increases in calcium intake do not improve calcium balance. Despite the fact that our subjects averaged total daily calcium intake that achieved and even surpassed this threshold, a mixture of inulin+oligofructose significantly increased their absorption of calcium. It is not clear from our data whether the additional amount of absorbed calcium was utilized for bone mineral production; however, we did not find an increase in urinary calcium excretion that would have negated the increase in calcium absorption. The absolute increase in calcium absorption due to consumption of a mixture of inulin+oligofructose was approximately 90 mg/d, which is, clinically, highly significant. If even part of this additional calcium was utilized for bone mineral production, it could lead to a significant increase in peak bone mineral density during this critical period. Preliminary data from ovariectomized rats show that not only did galacto-oligosaccharides increase calcium absorption, but the additional calcium absorption resulted in an increased bone mineral mass (Chonan et al. 1995). If this was the case in our study population, the results could have significant public health implications.

The effects of non-digestible oligosaccharides on more sophisticated measures of calcium metabolism and on bone mineral accretion rates will require further study, and were not the purpose of this study. Our findings, however, show that regular intake of modest amounts of a mixture of inulin+oligofructose significantly increases calcium absorption in girls at or near menarche, with adequate or high calcium intakes, without any compensatory increase in urinary calcium excretion.

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References


