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Fermentation of calcium-fortified soya milk does not appear to enhance acute calcium absorption in osteopenic post-menopausal women

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Abstract
Ageing women may choose to drink soya milk to reduce menopausal symptoms. As fermentation enriches soya milk with isoflavone aglycones, its beneficial qualities may improve. To reduce osteoporotic risk, however, soya milk must be Ca enriched, and it is not known how fermentation affects Ca bioavailability. A randomised crossover pilot study was undertaken to compare the Ca absorption of fortified soya milk with that of fermented and fortified soya milk in twelve Australian osteopenic post-menopausal women. The fortified soya milk was inoculated with Lactobacillus acidophilus American Type Culture Collection (ATCC) 4962 and fermented for 24 h at 37°C. Ca absorption from soya milk samples was measured using a single isotope radiocalcium method. Participants had a mean age of 54·8 (SD 12·3) years, with mean BMI of 26·5 (SD 5·5) kg/m2 and subnormal to normal serum 25-hydroxyvitamin D (mean 62·5 (SD 19·1) nmol/l). Participants consumed 185 kBq of 45Ca in 44 mg of Ca carrier. The mean fractional Ca absorption (α) from soya milk and fermented soya milk was 0·64 (SD 0·23) and 0·71 (SD 0·29), respectively, a difference not of statistical significance (P=0·122). Although fermentation of soya milk may provide other health benefits, fermentation had little effect on acute Ca absorption.

Key words: Calcium: Soya milk: Fermentation: Post-menopausal women

Soya milk, as a protein-rich drink containing isoflavones, is increasingly consumed in developed countries. Potential benefits include relief from hot flushes, improved lipid profiles, protection against oxidative damage to DNA and, in particular, maintenance of bone health1–3. Long-term consumption of isoflavones can have bone-sparing effects due to attenuation of bone loss3,4. Little is known on whether fermentation of soya milk will also affect Ca absorption from the small intestine.

Natural soya milk contains approximately 20 mg Ca/100 ml compared with cows’ milk which contains approximately 120 mg Ca/100 ml. Commercially available soya milk is now fortified to the same level as cows’ milk by adding fortificants such as calcium phosphate or carbonate. Not all Ca fortificants, however, are equivalent5. The bioavailability rather than the total content of Ca in soya milk is thus an important issue. Ca bioavailability is improved by the presence of high amounts of soluble Ca in food6,7 and by facilitating ionisation of Ca in the digestive system.

One way to potentially enhance the biological activity and nutritional value of soya milk is through fermentation with probiotics. The fermentation of soya milk in vitro with β-glucosidase-producing probiotic bacterial strains allows acetyl-glucoside and β-glucoside isoflavones to undergo enzymatic hydrolysis into biologically available aglycone structures and also increases Ca solubility8. Aglycones are absorbed faster and in greater amounts than their corresponding glucosides9,10.

In addition, probiotics are a living microbial food supplement which may have beneficial effects on symptoms of lactose intolerance, atopic disorders and coeliac disease, and they are useful in the treatment of diarrhoea, ulcerative colitis and irritable bowel syndrome11. Claims are also

Abbreviations: ATCC, American Type Culture Collection; CFSM, Ca-fortified soya milk.

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made for cholesterol-lowering effects, anti-carcinogenic actions and improved immune function\(^{12}\).

Our hypothesis is that fermented soya milk may have greater Ca bioavailability as measured by fractional Ca absorption than an otherwise equivalent non-fermented soya milk.

Post-menopausal women are at high risk of osteoporosis following increased bone loss. In the present study, we have investigated whether fermentation of fortified soya milk improves hourly fractional Ca absorption. A well-established crossover radiisotope method was used to compare Ca absorption from Ca-fortified soya milk v. fermented Ca-fortified soya milk in osteopenic but otherwise healthy Australian post-menopausal women.

**Experimental methods**

**Calcium fortification of soya milk**

The Ca-fortified soya milk (CFSM) used in the present study is widely sold throughout Australia (So Good; Sanitarium Health Foods, NSW, Australia). It is made from soya protein isolate (4\%) and has been fortified with a ‘proprietary’ phosphate of Ca to achieve Ca content similar to that of cows’ milk (120 mg/100 ml).

**Labelling of the fortificant with \(^{45}\)Ca after soya milk manufacture**

The fortificant present in the CFSM was labelled by adding 1 \(\mu\)g of high-specific activity \(^{45}\)CaCl\(_2\) to a 20 ml amount of soya milk, yielding a tracer concentration of approximately 185 kBq. Labelled CFSM was vortexed continuously for 1 min and then heat-treated (90°C for 30 min) before storing at 4°C for 24 h to allow for Ca exchange. The labelled CFSM was then either given after the 24 h exchange as the test drink or fermented before consumption with *Lactobacillus acidophilus* American Type Culture Collection (ATCC) 4962.

**Bacteria**

A pure culture of *L. acidophilus* ATCC 4962 was obtained from the Victoria University Culture Collection (Werribee, VIC, Australia). Purity was checked by Gram staining before storage at \(-80°C\) in 40\% glycerol.

**Fermentation of calcium-fortified soya milk**

The probiotic culture *L. acidophilus* ATCC 4962 was activated through three successive transfers in de Man Rogosa Sharpe broth\(^{13}\) at 37°C for 20 h using a 2\% inoculum. The labelled CFSM was aseptically inoculated with a 1\% (v/v) inoculum, incubated at 37°C for 24 h and stored for a maximum period of 48 h at 4°C before consumption.

**Human study protocol**

Twelve osteopenic but otherwise healthy post-menopausal women aged 50–68 years were recruited by advertisement and screened by telephone interview. Women were included if they were post-menopausal non-smokers diagnosed as osteopenic (i.e. with bone mineral density T-score \(-1\) to \(-2.5\) as measured by dual-energy X-ray absorptiometry); were otherwise healthy (no chronic disease by self-report; including gastrointestinal, kidney, liver, parathyroid or CVD); were not taking medications or antibiotics affecting Ca absorption; and had not taken hormone replacement therapy during the preceding 12 months. The participants were required to be lactose tolerant and not allergic to soya. Each completed an eating habit questionnaire (extracted from a FFQ; Australian Cancer Council, VIC, Australia) to assess dietary Ca intake. They were asked to avoid any Ca supplements for at least 4 weeks before and during the study. The present study was conducted according to the guidelines of the 1995 Declaration of Helsinki as revised in Edinburgh, 2000, and all procedures were approved by the Southern Health Human Research and Ethics Committee (project number 07013A). Written informed consent was obtained from all the participants.

**Test milk drinks for the acute pilot study**

Soya milk for the acute study had a tracer concentration of 185 kBq/dose. Each dose comprises 20 ml of the test drink containing a microgram amount of \(^{45}\)CaCl\(_2\) (Amersham Biosciences, Rydalmere, NSW, Australia) in a total 44 mg of Ca carrier (20 mg as \(^{45}\)Ca and 24 mg as \(^{40}\)Ca present in the CFSM). Immediately after ingestion of the test soya milk, 200 ml of distilled water was consumed.

**Study design**

The participants arrived at Monash Medical Centre after an overnight fast and were tested for Ca absorption on two separate occasions. Treatments were randomised and given in a crossover design study with a minimum washout between tests of 3 weeks. At each test, the subjects consumed either radiolabelled CFSM or radiolabelled and fermented CFSM. At each visit, bioelectrical impedance was determined (SFB7; Impedimed, Brisbane, QLD, Australia). A 10 ml venous blood sample was collected from an antecubital vein for a baseline sample and for measurement of serum 25-hydroxyvitamin D. Participants then consumed the 20 ml test drink immediately followed by the consumption of 200 ml distilled water. Blood samples were collected after 60 min as described by Nordin et al\(^{14}\). These were centrifuged at 3500 \(g\) for 10 min, and the activity of \(^{45}\)Ca in 1 ml plasma aliquots was measured using a liquid scintillation counter (Wallac 1410; Perkin Elmer Life Sciences, MA, USA). Fractional Ca (\(\alpha\)) was then calculated\(^{15}\).
Calculation and statistical analysis

The sample of twelve women recruited into this crossover design pilot study provides a probability of 90% that a treatment difference can be detected at the 5% level of significance (two sided), if the true difference between the treatments is 15%. This calculation is based on the assumption that the within-patient sd of the response variable is 10%. Data from the present study were compared by Student’s paired t test. SPSS for Windows (version 11.5; SPSS Australasia Limited, Melbourne, VIC, Australia) was used for statistical analyses. A P value of <0.05 was considered as significant.

Results

Comparison of calcium absorption from calcium-fortified soya milk v. fermented calcium-fortified soya milk

Women in the study had a mean age of 54.8 (SD 12.3) years and were overweight (mean BMI 26.5 (SD 5.5) kg/m²). Mean serum 25-hydroxyvitamin D was 62.6 (SD 19.1) nmol/l. Most women had normal levels, but 38.5% women had vitamin D insufficiency (serum 25-hydroxyvitamin D < 50 nmol/l).

The mean fractional Ca absorption (α) values for fermented CFSM compared to CFSM were 0.71 (SD 0.29) and 0.64 (SD 0.23), respectively. The mean fractional Ca absorption (α) of the fermented CFSM was approximately 10% higher compared with that of CFSM, a difference not of statistical significance (P = 0.122). The individual differences in fractional Ca absorption between fortified soya milk and fermented fortified soya milk in the participants are shown in Fig. 1.

Discussion

In vitro studies indicate that fermentation of soya milk with some probiotics may enhance Ca solubility and bioavailability\(^{18,16}\). To date, no other studies have examined the effects of fermenting CFSM on Ca absorption in human subjects. In the present study, the participants were not vegetarian, and most of them (77%) rarely consumed soya milk or soya products. Around one-third (38.5%) of them regularly had Ca and vitamin D supplements. All performed only low to moderate physical activity. Habitual intake of Ca (by self-report) was moderate, and Ca supplementation was avoided during the study. Ca intake by the participants is, thus, unlikely to have affected the present results; moreover, from the crossover design of the study, each participant acted as their own control.

The study was based on the single isotope radiocalcium absorption test, a robust, well-validated measure of Ca absorption\(^{14}\). This method would be applicable to other drinks fortified with Ca. The test can be completed over a short-time period, allowing absorption from a segment of small intestine to be followed via a sharp peak of radioactivity\(^{17}\). The rate of Ca absorption measured by this method correlates strongly with that measured in balance studies and correlates very highly with double isotope Ca absorption tests\(^{18}\). It is important, however, when employing this method, to use a small Ca load (e.g. 44 mg as here). Higher loads increase absorption time and will reduce test sensitivity. The larger the carrier dose, the more interference during the Ca absorption diffusion process and the less valid the single-isotope procedure\(^{14}\). The labelling of the fortificant with \(^{45}\)Ca after soya milk manufacture was shown to have a tracer distribution pattern very similar to that when the fortificant was labelled before the soya milk manufacture, provided a heat treatment was applied\(^{19}\). In the present study, the soya milk fortificant was also labelled before the fermentation. No studies have indicated negative effects of probiotics on the availability of the \(^{45}\)Ca radioisotope during Ca absorption, although a recent study found that Ca\(^{2+}\) plays a positive catalytic role for human gut colonic bacteria\(^ {20}\).

We have shown that fermenting CFSM with L. acidophilus ATCC 4962 did not improve fractional Ca absorption in twelve osteopenic post-menopausal women. The insignificant effect of fermentation on Ca bioavailability observed may in part reflect our choice of CFSM for fermentation. We have previously demonstrated that the fractional Ca absorption (α) from the CFSM used in the present study is comparable to that of cows’ milk\(^ {19}\). The fortificant present in this CFSM may already be in its most absorbable form so that fermentation in this case does not signifi cantly improve acute Ca absorption. Optimum Ca absorption (α) was observed 1 h after ingestion of the unfermented soya milk. It remains possible that fermentation will improve Ca absorption in other soya milk drinks where other methods of Ca fortification have been used. It would thus be valuable to repeat the present study with other types of commercially available fortified soya milk.
Even without change in Ca bioavailability, fermentation may have health benefits as it significantly increases aglycone content (increasing daidzein, glycitein and genistein)\(^{(23)}\). Fermentation also increases the solubility of Ca by decreasing soya milk pH. Moreover, the phytase enzyme produced by some probiotics will hydrolyse phytic acid and IP6-generating myo-inositol with reduced numbers of phosphate groups (IP3–IP5)\(^{(21,22)}\) causing a beneficial effect on Ca bioavailability. In the present study, the CFSM was made from soya protein isolate rather than from a whole soya bean, and even before fermentation, it had minimal phytic acid content.

Fig. 1 shows the individual fractional Ca absorption from CFSM to fermented CFSM. Four of the twelve postmenopausal women absorbed Ca from the fermented CFSM better than from the non-fermented CFSM (32, 28, 69 and 31%, respectively). These women may have come from the approximate one-third of the population who are ‘equol producers’, a mechanism known to facilitate Ca absorption\(^{(25)}\). In further studies, it would be advised to assess equol-producing status via 24 h urine excretion\(^{(25)}\). Although our acute study indicates that fermentation of CFSM has no effect on Ca bioavailability under conditions of acute absorption from the small intestine, it cannot rule out the possibility that fermented CFSM facilitates slower Ca absorption from the large intestine. Post hoc analysis suggests that for adequate statistical power, 174 subjects would be needed for the present study per treatment group. Our pilot study may thus have been underpowered to detect any small difference in bioavailability between the two test drinks. It does not exclude a greater difference with fermentation in different soya milk drinks fortified by other methods. Our findings do not therefore preclude possible benefits on long-term consumption of fermented CFSM on bone health and Ca balance.

In summary, during this acute pilot study, there was no significant improvement on fractional Ca absorption from the ingestion of CFSM fermented with \textit{L. acidophilus} ATCC 4962 in osteopenic post-menopausal women. Limitations of this randomised crossover pilot study include the sample size of twelve, the use of a test soya milk with relatively high Ca bioavailability in its non-fermented state and the comparison of small-intestinal Ca absorption in the acute setting only, therefore looking at neither differences in colonic Ca absorption nor longer-term Ca accretion with the two test soya milk drinks. To observe a significant improvement in fractional Ca absorption with this particular brand of soya milk, a much larger sample size study would be required. The effects of fermentation might be observed more readily with soya milk of different composition (e.g. whole bean rather than soya protein isolate based) and Ca fortification system. Fermentation of CFSM may also contribute to the potential cumulative long-term benefits on Ca bioavailability and bone health. A longer-term study, over at least 6–12 months, looking at markers of bone turnover and bone mineral densitometry, may be needed to test this hypothesis further.

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