Bioavailability of isoflavone phytoestrogens in postmenopausal women consuming soya milk fermented with probiotic bifidobacteria

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We investigated the effects of consuming an isoflavone aglycone-enriched soya milk containing viable bifidobacteria on urinary isoflavone excretion and percentage recovery. Sixteen postmenopausal women were randomly divided into two groups to consume either fermented or non-fermented soya milk. Each group participated in a double-blind, crossover study with three 14 d supplementation periods, separated by a 14 d washout. Subjects ingested three daily dosages of isoflavone via the soya milk and collected four 24 h pooled urine specimens per supplementation period. Soya milks were prepared with soya protein isolate and soya germ, followed by fermentation with bifidobacteria. Isoflavone levels were quantified using HPLC. Non-fermented soya milks at 20, 40 and 80 mg isoflavone/200 ml contained 10%, 9% and 7% aglycone, respectively, with their fermented counterparts containing 69%, 57% and 36% aglycone (P<0.001). A trend to a greater percentage urinary recovery of daidzein and genistein was observed among women consuming fermented soya milk at a dosage of 40 mg isoflavone (P=0.13). A distinct linear dose response for the fermented soya milk group (R²=0.9993) compared with the non-fermented group (R²=0.8865) suggested less interindividual variation in isoflavone absorption. However, total urinary isoflavone excretion was similar for both groups (P>0.05), with urinary isoflavone recovery at approximately 31%. Increasing the isoflavone dosage correlated positively with its urinary excretion, but urinary percentage recovery of isoflavone was inversely related to dosage level. Hence, a modest dosage ranging from 20 to 30 mg/d may provide the most bioavailable source of isoflavone, regardless of whether it is via an aglycone-rich fermented soya milk or a glucoside-rich soya milk.

Isoflavone: Bifidobacteria: Soya milk: Bioavailability: Postmenopausal women

Diminishing levels of oestrogen caused by ovarian failure, inadequate to maintain oestrogen-dependent tissues, are associated with higher rates of chronic disease in postmenopausal women. Phytoestrogens, a group of non-steroidal plant-derived compounds, are structurally similar to oestrogen and thus able to exert weak oestrogenic effects (Mayr et al. 1995; Cassidy, 1996) and small-scale human clinical studies (Anderson et al. 1995; Kurzer, 2000). Soyabeans contain three types of isoflavone, each type being present in four chemical forms. Aglycone structures of daidzein, genistein and glycitein are those forms with an oestrogen-like configuration and functionality but are found in soya bean foods predominantly as malonyl-, acetyl- and β-glucoside conjugates (>90% of the total concentration; Murphy et al. 1999). Izumi et al. (2000) found that isoflavone aglycones were absorbed more quickly and in greater amounts than their glucosides in human subjects. Furthermore, Setchell et al. (2002) reported that isoflavone glucosides were not absorbed intact across the enterocytes of adults, their bioavailability requiring hydrolysis of the sugar moiety by intestinal β-glucosidases. This suggests that consuming isoflavone aglycone-rich soya foods may be more effective than consuming glucoside-rich products in preventing chronic disease. However, Richelle et al. (2002) discovered that the enzymic hydrolysis of isoflavone glucosides into aglycones in a soya drink (made from soya germ (SG)) before consumption did not enhance the absorption of isoflavones in postmenopausal women after a single dose; plasma and urine isoflavone pharmokinetics were similar for both aglycone- and glucoside-rich soya drinks. In contrast, Hutchins et al. (1995) and Slavin et al. (1998) reported that the fermentation of cooked soyabeans by Rhizopus oligosporus (tempeh) enhanced the bioavailability of daidzein and genistein in men and women over a 9 d feeding period (112 g tempeh/d) compared with the ingestion of non-fermented cooked soyabean pieces under identical conditions.

Abbreviations: AUC, area under the curve; CFU, colony-forming units; FS, fermented soya milk; ISTD, internal standards; NFS, non-fermented soya milk; SG, soya germ; SPI, soya protein isolate; UV/VIS, UV/visible.

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Intestinal bacteria play an important role in the ultimate bioactivities of isoflavones (Hendrich, 2002; Turner et al. 2003). Isoflavones are excreted in urine and found predominantly as glucuronide and sulphate conjugates. Owing to their phenolic nature, isoflavones undergo biotransformation at hydroxyl groups by mammalian UDP-glucuronosyltransferases and sulphotransferases in the intestinal mucosa and liver (Hendrich, 2002). To date, studies on the pharmacokinetics of isoflavones in human subjects have not investigated the effects of fermenting soya foods with bifidobacteria. Bifidobacteria constitute a major part of the normal microflora of the human intestine, their highest populations being found in the ileum and colon (Orrhage & Nord, 2000). We discovered that β-glucosidase-producing *Bifidobacterium animalis* Bb-12 hydrolysed isoflavone glucosides into bioactive and bioavailable aglycones when grown in soya milk (Tsangalis et al. 2002, 2003, 2004). In clinical studies, *B. animalis* Bb-12 has also been shown to effectively modulate intestinal microbial balance (Payle, 2002). Hence, the enrichment of isoflavone aglycone in soya milk by fermentation prior to consumption and the modulation of intestinal microflora via the ingestion of viable bifidobacteria may enhance the bioavailability of isoflavones consumed on a daily basis.

The objective of this study was to investigate the effects of the daily consumption of an isoflavone aglycone-enriched fermented soya milk containing viable populations of *B. animalis* Bb-12 on urinary isoflavone excretion and percentage recovery in postmenopausal women.

**Subjects and methods**

**Subjects**

Sixteen healthy postmenopausal women were recruited from the Melbourne metropolitan area from fifty-two interested volunteers. They were screened using a health-information and food-frequency questionnaire designed to exclude those who: had gastrointestinal disorders; had food allergies; had an alcohol intake greater than two standard drinks per day; regularly used prescription or non-prescription medication; had been on oestrogen-replacement therapy or antibiotics in the past 3 months; or had dietary patterns that were not representative of the general population (e.g. strict vegetarian, consumption of more than two servings of soya food per week). Participants were randomly allocated to one of two groups (eight subjects per group), to consume either fermented or non-fermented soya milk during the supplementation periods. The mean age of women in the fermented soya milk (FS) and non-fermented soya milk (NFS) group was 52·5 (SD 3·1) and 55·6 (SD 5·1) years, respectively. Five women in the FS group and six women in the NFS group had not had any menstrual bleeding for at least 12 months; the other women recruited had not menstruated for at least 6 months.

**Study design**

The study protocol was approved by the Human Research Ethics Committee of Victoria University (Melbourne, Victoria, Australia) and consisted of a randomised, double-blind, crossover design involving three 14 d soya milk supplementation periods separated by two 14 d washout periods. Subjects consumed either fermented or non-fermented soya milk containing three different concentrations of total isoflavone during each respective 14 d supplementation period. All batches of fermented soya milk contained a consistent population of *B. animalis* Bb-12, at 10⁸ viable cells/ml. A volume of 200 ml chilled (4°C) fermented soya milk was required per day, 100 ml before breakfast and before dinner. Each participant consumed a self-selected diet for the entire study and provided a 14 d weighed food and beverage record during each supplementation period. Subjects were advised not to alter their diet for the duration of the study and were asked to exclude foods containing soy as an ingredient, chickpeas, lentils, beans (all types), alfalfa, mung bean sprouts, fermented dairy products, probiotic supplements and alcoholic beverages. Daily energy, macronutrient and dietary fibre intake were quantified for each subject using Food Works version 3.01 nutrition software utilising a database of Australian foods (Xyris Software Pty Ltd, Highgate Hill, Queensland, Australia). Anthropometric measurements of height (m) and body weight (kg) were recorded and the BMI of each subject calculated (kg/m²) before and after each supplementation period.

**Collection and handling of urine specimens**

Each participant collected four 24 h pooled urine specimens per soya milk supplementation period: a day before beginning soya milk supplementation (baseline) and on days 4, 13 and 14. Twenty-four hour pooled urine specimens consisted of the second urination on the allocated day and those following, and included the first urination on the next day, and were passed into 3·2 litre sterile plastic bottles (Biolcorp Pty Ltd, Huntingdale, Victoria, Australia) containing 2 g ascorbic acid (Sigma Chemical Co., Castle Hill, New South Wales, Australia) as preservative against chemical degradation. Specimens were stored at 4°C or on ice until processed. The total volume of each 24 h pooled urine specimen was measured and a 100 ml aliquot taken for urinary isoflavone analyses. Prior to storage at −20°C, 1 ml 10 % (w/v) sodium azide (Labchem Pty Ltd, Auburn, Australia) was added to the aliquot as a preservative against microbial spoilage.

**Soya milk ingredients**

*Bifidobacterium animalis* Bb-12 was obtained from Chr Hansen Pty Ltd (Bayswater, Australia) and stored at −80°C in 12 % (w/v) sterile (121°C for 15 min) reconstituted skim milk supplemented with d-glucose (1 % w/v), yeast extract (0·5 % w/v) and glycerol (40 % v/v). Supro 159 soya protein isolate (SPI) was obtained from Protein Technologies International Australia (Kensington, Australia); this, according to manufacturer specifications, contained a minimum of 90 g protein, 4 g fat and 4 g or less soluble carbohydrates per 100 g. IsoLife SG was obtained from Soyl Health Pty Ltd (Sydney, Australia) and comprised a macronutrient composition of 40 g protein, 16 g fat and 25 g carbohydrate per 100 g. The total concentration of isoflavone in this brand of SPI and SG was 1·4 and 19·4 mg/g, respectively (Tsangalis et al. 2004). Vanilla flavour (in propylene glycol) was purchased from Essential Flavours and Ingredients (Rowville, Australia).

** Manufacture of soya milk**

Three soya milk formulations were manufactured, each containing a different ratio of SPI to SG (total of 4 % w/v soya ingredients), the calculation being based upon the concentration of isoflavone in each soya ingredient. SPI and SG at ratios of 9:4:9:6,
80:20 and 52:48 were used to manufacture soya milks containing approximately 20 (SPI/SG20), 40 (SPI/SG40) and 80 (SPI/SG80) mg isoflavone per 200 ml, respectively. For SPI/SG20, 748 g SPI and 52 g SG were reconstituted in 20 litres filtered water at a temperature of 40 °C, followed by heating with stirring at 70 °C for 30 min during which 80 ml vanilla flavour was added. The soya milk was then dispensed into four bottles in 5 litre quantities and autoclaved at 121 °C for 15 min. Two bottles of sterile soya milk (10 litres) were cooled to approximately 75 ± 5 °C, and 200 ml non-fermented product was dispensed into forty-five sterile, 200 ml high-density polyethylene bottles (Cospak Pty Ltd, Braeside, Australia) under aseptic conditions using a 100 ml Trubor bottle top dispenser (U-Lab Pty Ltd, Eltham, Australia). Aliquots of 100 ml from three randomly selected bottles of non-fermented soya milk were obtained under aseptic conditions and freeze-dried (50 ml aliquot) using a Dynavac FD300 freeze-drier (Rowville, Australia) for extraction of isoflavone and analysis using HPLC. Fourteen bottles of non-fermented soya milk were packaged and stored at 4 °C prior to distribution. The remaining 10 litres of sterile soya milk were cooled to 40 °C prior to inoculation with an active culture of B. animalis Bb-12. SPI/SG40 and SPI/SG80 were also manufactured in the same manner using SPI–SG 638 g:162 g and SPI–SG 416 g:384 g, respectively, per 20 litres of filtered water.

Bacterial growth media
Rehydrated MRS agar (De Mann et al. 1960) containing additional 1 % (w/v) d-glucose was prepared as per manufacturer instructions (Oxoid Ltd, West Heidelberg, Victoria, Australia) and autoclaved at 121 °C for 15 min. Filter-sterilised L-cysteine-HCl solution (5 % w/v) was added to the agar (0.05 % w/v final concentration) just prior to inoculation with bacteria to lower the oxidation-reduction potential of the media and enhance the growth of anaerobic bifidobacteria. d-Glucose and L-cysteine-HCl were purchased from Sigma.

Reconstituted SPI at a ratio of 24 g/600 ml filtered water supplemented with 1 % (w/v) food-grade glucose powder (Prahan Health Foods, Prahran, Australia) was prepared for the activation of B. animalis Bb-12, as described earlier (Tsangalis et al. 2004). A 600 ml batch was manufactured and dispensed into glass bottles in 20, 50 and 500 ml quantities and sterilised by autoclaving at 121 °C for 15 min.

Fermentation of soya milk with bifidobacteria
Bifidobacterium animalis Bb-12 was activated by three successive transfers. An inoculum level of 5 % (v/v) was used for strain activation, and incubation was at 37 °C for 20 h. Sterile soya milk (SPI/SG20, SPI/SG40 and SPI/SG80 10 litres) was inoculated with an active culture of B. animalis Bb-12 (5 % v/v). A volume of 200 ml inoculated soya milk was dispensed into forty-eight sterile 200 ml high-density polyethylene bottles under aseptic conditions using a 100 ml Trubor bottle top dispenser and incubated at 37 °C for 24 h. Three bottles were randomly taken from the batch at 0 and 24 h of incubation, and a 100 ml sample was withdrawn aseptically from each bottle for the enumeration of viable B. animalis Bb-12 populations (as described by Tsangalis et al. 2004), the remaining being stored at −20 °C. Frozen samples were freeze-dried (50 ml aliquot) using a Dynavac FD300 freeze-drier for the extraction of isoflavone and analysis using HPLC. The remaining forty-two bottles of fermented soya milk were equally packaged into three cartons and stored at 4 °C prior to distribution.

Isoflavone standards
Aglycone standards of genistein, daidzein and glycitein (synthetic) used for both soya milk and urinary isoflavone analyses were purchased from Sigma. β-Glucoside isomers of genistin, daidzin and glycitin were purchased from Indofine Chemical Co (Somerville, USA). Internal standards (ISTD) of flavone and benzophenone, used for the quantification of isoflavones in soya milk and urine, respectively, were obtained from Sigma. Mixed and single isoflavone standards used in the analysis of isoflavones in soya milk were prepared as described earlier (Tsangalis et al. 2002).

Reversed-phase HPLC apparatus and reagents
Chromatographic analyses were carried out on a Varian 9000 series high-performance liquid chromatograph (Varian Pty Ltd, Mulgrave, Australia) with auto-sampler (9100), solvent delivery system (9010), polychromatophotodiode array UV/visible (UV/VIS) detector (9065) and thermostatically controlled column compartment (Alltech Associates Pty Ltd, Box Hill, Australia). An Exsil (SGE International Pty Ltd, Ringwood, Australia) C18-ODS (250 × 4.6 mm internal diameter 5 μm) reversed-phase column was used to separate the isoflavone isomers. HPLC-grade methanol and acetonic acid were purchased from Merck Pty Ltd (Kilsyth, Australia) and trifluoro-acetic acid, glacial acetic acid, sodium acetate and ammonium acetate from Sigma. All reagents used for soya milk and urinary isoflavone extraction and HPLC analyses were filtered through a 0.5 μm membrane (Millipore, Bedford, USA), and mobile phases were degassed using nitrogen.

Extraction and HPLC analysis of isoflavones in soya milk
A sample of SPI powder (2 g), 1.5 g freeze-dried SPI/SG20, 1.0 g freeze-dried SPI/SG40 or 0.5 g either freeze-dried SPI/SG80 or SG powder was used in the extraction of isoflavone isomers (malonyl, acetyl, β-glucoside and aglycone forms), as described earlier (Tsangalis et al. 2002). HPLC gradient elution was composed of acetonitrile (solvent A) and 10 mM-ammonium acetate buffer containing 0.1 % trifluoro-acetic acid (solvent B), set at a flow rate of 1 ml/min (Setchell et al. 2001). After a 20 μL injection of sample or isoflavone standard on to the column (25 °C), solvent B was set at 100 % for 2 min, reduced to 40 % over 22 min followed by 40 % for 5 min, increased to 100 % over 6 min and finally increased to 100 % for 5 min prior to the next injection. The diode array UV/VIS detector was set at 259 nm, and the retention times (min) of isoflavone isomers were as follows: malonyldaidzin, 13.7; malonylglycitin, 14.0; malonylgenistin, 14.3; daidzin, 14.6; glycitin, 14.8; genistin, 16.2; acetylaidzin, 16.9; acetylglycitin, 17.1; acetylgenistin, 18.7; daidzin, 19.2; glycitin, 19.5; genistein, 21.7; and flavone (ISTD), 28.2. Retention times for aglycone and β-glucoside isoflavone isomers were determined using single standards, whereas those of malonyl- and acetyl-glucoside isomers were based on the retention times reported by Setchell et al. (2001) under similar HPLC conditions. Quantification of isoflavone isomers in soya
milk using multi-level calibration and an ISTD was as described by Tsangalis et al. (2002). Isoflavone concentrations are expressed as μmol/200 ml soya milk (calculated back to wet basis).

**Analysis of isoflavones in urine specimens**

Aliquots from 24 h pooled urine specimens were analysed for their daidzein, genistein and glycitein concentration. Urine samples were extracted for isoflavone and analysed in batches over consecutive days with all the specimens from one subject run in the same batch. Extraction of urinary isoflavones was performed described by Franke & Custer (1994), with modifications as follows. Frozen urine aliquots were equilibrated to room temperature, vortex-mixed and centrifuged (Sorvall RT7 refrigerated centrifuge; Newtown, USA) at 2700 g for 10 min. A volume of 20.2 ml clear supernatant was mixed with 5.0 ml 0.2 M acetic acid buffer (pH 4) and 120 μl benzophenone (10 mg/50 ml methanol) and filtered through a C18 solid-phase extraction column (Alltech) preconditioned with 5.0 ml methanol and 5.0 ml 0.2 M-acetate buffer (pH 4). After passing the entire urine sample through the column, the column was washed with 2.0 ml 0.2 M-acetate buffer (pH 4), and the isoflavones were eluted with 100 % methanol to give exactly 2.0 ml. A volume of 100 μl of this eluate was used for the analysis of unconjugated forms of daidzein, genistein and glycitein. The residual 1.9 ml of eluate was dried under a stream of nitrogen using a Pierce nine-needle evaporating unit (Pierce Biotechnology Inc., Rockford, USA) and then resuspended in 900 μl freshly prepared mixture comprising 10 ml 0.2 M-acetate buffer (pH 4), 150 mg ascorbic acid and 500 μl glucuronidase/sulphatase (sterile filtered crude solution isolated from Helix pomatia type HP-2S; Sigma). The hydrolysed sample (total of unconjugated and conjugated isoflavones) was then mixed with 1.0 ml 100 % methanol, centrifuged at 4000 g for 5 min, and 50 μl were injected on to the column (25°C). HPLC conditions were as described by Xu et al. (1994), with the diode array UV/VIS detector set at 259 nm to detect daidzein, genistein and glycitein. Six mixed standards containing daidzein, genistein and glycitein. The residual 1.9 ml of this eluate was used for the analysis of unconjugated forms of daidzein, genistein and glycitein. The residual 1.9 ml of eluate was dried under a stream of nitrogen using a Pierce nine-needle evaporating unit (Pierce Biotechnology Inc., Rockford, USA) and then resuspended in 900 μl freshly prepared mixture comprising 10 ml 0.2 M-acetate buffer (pH 4), 150 mg ascorbic acid and 500 μl glucuronidase/sulphatase (sterile filtered crude solution isolated from Helix pomatia type HP-2S; Sigma). The hydrolysed sample (total of unconjugated and conjugated isoflavones) was then mixed with 1.0 ml 100 % methanol, centrifuged at 4000 g for 5 min, and 50 μl were injected on to the column (25°C). HPLC conditions were as described by Xu et al. (1994), with the diode array UV/VIS detector set at 259 nm to detect daidzein, genistein and glycitein. Six mixed standards containing daidzein, genistein and glycitein (50, 100, 150, 250, 350 and 450 ng/50 μl) were used for the quantification of isoflavones. Benzophenone (ISTD) was added to each isoflavone standard at a concentration of 600 ng/50 μl. Single standards were also prepared for peak identification, and the retention times (min) of isoflavone isomers were as follows: daidzein, 15.5; glycitein, 16.8; genistein, 19.8; and benzophenone (ISTD), 28.8. Isoflavone concentrations were calculated with respect to the ISTD and multiplied to represent the entire pooled urine specimen.

**Statistical analysis**

Non-fermented and fermented SPI/SG20, SPI/SG40 and SPI/SG80 were manufactured on three occasions, with three bottles of soya milk randomly selected from each batch and their isoflavone content and viable B. animalis Bb-12 populations analysed in duplicate. Hence, populations of B. animalis Bb-12 in fermented soya milks consumed during each supplementation period are reported as a mean with standard deviation of six replicates, and the concentrations of each isoflavone isomer in non-fermented and fermented soya milks consumed during the entire study are presented as a mean with standard deviation of eighteen replicates. For the FS and NFS groups, concentrations of daidzein, genistein and glycitein in 24 h pooled urine specimens were analysed in duplicate and are presented as a mean with standard deviation of sixteen replicates. To find significant differences in body weight, BMI, nutrient intake, viability of B. animalis Bb-12 and isoflavone levels in soya milk, and urinary isoflavone excretion, means were analysed using ANOVA and 95 % CI with Microsoft Excel Stat Pro, as described by Albright et al. (1999). ANOVA data with a P < 0.05 were classified as statistically significant (two-sided test).

**Results and discussion**

**BMI and body weight**

The mean with standard deviation BMI [body weight] of the NFS and FS groups (n = 8) at baseline (i.e. a day before the first supplementation period) were 29.3 (SD 6.2) kg/m² [71.3 (SD 13.4) kg] and 24.6 (SD 5.4) kg/m² [61.9 (SD 11.3) kg], respectively, with no significant difference in either BMI (P = 0.13) or body weight (P = 0.15) between the two groups. No significant fluctuations in body weight were observed during the entire span of supplementation and washout periods for the NFS and FS groups (P > 0.05). Hence, the mean BMI of the NFS and FS groups after the final supplementation period were 28.9 (SD 5.6) and 24.3 (SD 5.3) kg/m², respectively, similar to their respective baseline values (P > 0.05).

**Intake of dietary nutrients, viable bifidobacteria and isoflavones**

The mean daily energy intake for women in the FS and NFS groups ranged between 5698.8 (SD 1478.9) and 6445.9 (SD 1568.6) kJ/d, with no significant differences in daily energy intake between the two groups during the supplementation periods (P = 0.9). The mean daily intake of macronutrients for the FS and NFS groups during supplementation are shown in Fig. 1. Both groups of women consumed an average of 20 % energy in the form of protein, 44 % energy as carbohydrates and 36 % as fat (total intake of polysaturated, monounsaturated and saturated fats) per day during supplementation. There were no significant differences in protein (P = 0.8), carbohydrate (P = 0.7), fat (P = 0.9) and dietary fibre (P = 0.9) intake (g/d) between the FS and NFS groups during supplementation periods.

According to the protein content of SPI and SG, non-fermented SPI/SG20, SPI/SG40 and SPI/SG80 comprised 6.96, 6.40 and 5.28 g protein per 200 ml serving, respectively. The concentration of carbohydrate (and fibre) in non-fermented SPI/SG20, SPI/SG40 and SPI/SG80 was 0.42 g (0.01 g), 0.66 g (0.05 g) and 1.13 g (0.12 g) per 200 ml serving, respectively. The total concentration of polysaturated, monounsaturated and saturated fats in non-fermented SPI/SG20, SPI/SG40 and SPI/SG80 was 0.38, 0.51 and 0.78 g per 200 ml serving, respectively. Each of the soya milks contributed relatively little to the mean intake of macronutrients ingested per day by women in the FS and NFS groups, being approximately 8.5 %, 0.4 %, 1.0 % and 0.3 % of the daily intake of protein, carbohydrate, fat and fibre, respectively. Fermented SPI/SG20, SPI/SG40 and SPI/SG80 may have contained a lower level of protein, carbohydrate, fat and fibre than their non-fermented counterparts, as each of these nutrients could have been metabolised as a growth substrate during the fermentation of bifidobacteria (Kamaly, 1997; Hou et al. 2000).

Viable populations of B. animalis Bb-12 in fermented SPI/ SG20, SPI/SG40 and SPI/SG80 (200 ml serving) consumed by...
women in the FS group during supplementation ranged from 7.60 (SD 0.06) to 8.87 (SD 0.08) \log_{10} \text{CFU/ml} per ml. This level of viable bifidobacteria exceeded the minimum level considered to be of therapeutic dosage, at 6 \log_{10} \text{CFU/ml} (Gomes & Malcata, 1999). Such high numbers of bifidobacteria compensate for the possible reduction in number of viable cells during passage through the stomach and intestine. There were no significant differences between \textit{B. animalis} Bb-12 populations found in fermented SPI/SG20, SPI/SG40 and SPI/SG80 \((P=0.07)\). Furthermore, similar populations of \textit{B. animalis} Bb-12 were ingested during each supplementation period \((P=0.08)\).

Mean concentrations of isoflavone isomers in non-fermented and fermented SPI/SG20, SPI/SG40 and SPI/SG80 ingested per day (via a 200 ml serving) during soya milk supplementation are shown in Table 1; each value is expressed as \mu\text{mol} so that it can be compared with the levels of isoflavone (\mu\text{mol/d}) excreted in urine. Non-fermented SPI/SG20, SPI/SG40 and SPI/SG80, which were formulated to contain 20, 40 and 80 mg isoflavone/200 ml, were equivalent to 64-11, 101-80 and 172-28 \mu\text{mol} aglycone constituents per 200 ml, respectively. The highest daily dose of isoflavone via SPI/SG80 was about 1-7 and 2-6 times the isoflavone concentration of SPI/SG40 and SPI/SG20, respectively \((P<0.001)\). Non-fermented SPI/SG20, SPI/SG40 and SPI/SG80 comprised 10 \%, 9 \% and 7 \% of its isoflavone level as unconjugated aglycone, respectively. Fermentation of SPI/SG20, SPI/SG40 and SPI/SG80 with \textit{B. animalis} Bb-12 significantly increased the proportion of aglycone structures to 69 \%, 57 \% and 36 \% of total isoflavone, respectively \((P<0.001)\). Fermented SPI/SG40 and SPI/SG80 contained the highest concentration of aglycone structures at 57-74 and 61-67 \mu\text{mol}/200 ml, significantly higher than the other soya milks \((P<0.001)\). However, changes in isoflavone isomer composition were less apparent in SPI/SG80 after fermentation. In an earlier study (Tsangalis \textit{et al.} 2004), we reported that the \beta-glycosidase levels produced by \textit{B. animalis} Bb-12 during fermentation were insufficient to hydrolyse high levels of isoflavone glucoside in the range found in SPI/SG80. Furthermore, SPI/SG80 comprised the greatest concentration of glycitein, malonylglycitin and acetylglycitin (approximately 35 \% of total isoflavone), which we found to be poorly hydrolysed by \beta-glycosidase during soya milk fermentation with \textit{B. animalis} Bb-12 (Tsangalis \textit{et al.} 2004). The high concentration of glycitein and its glucosides in SPI/SG80 was a result of the greater proportion of SG powder in the soya milk formulation. SG is a concentrated source of isoflavone naturally found in the soyabean’s hypocotyl, the only part of the bean that contains glycitein and its glucosides at a concentration higher than both daidzein and genistein and their glucosides (Kudou \textit{et al.} 1991).

**Urinary isoflavone excretion after consumption of fermented and non-fermented soya milk**

Interindividual variation in intestinal microflora has been shown to affect the bioavailability of isoflavones (Hendrich, 2002). The influence of diet on the bioavailability of isoflavones has also been observed (Setchell \textit{et al.} 1984; Kelly \textit{et al.} 1993; Lampe \textit{et al.} 1998). The ingestion of viable bifidobacteria to modulate intestinal microflora and its possible effect on the bioavailability of isoflavones has not been studied. Changes in intestinal microflora occur from birth to old age; with young adult women having a different intestinal microbial ecology from those women in their postmenopausal years (Ballongue, 1993). Mitsuoka (1984) reported that the number of \textit{Bifidobacterium} in adult stools, particularly those of the elderly, was significantly lower than the populations of \textit{Bifidobacterium} found in the stool samples of children. Therefore, owing to the relevance of isoflavone-rich soyabean foods to postmenopausal women and considering the variation in intestinal microflora after different stages of life, we limited the inclusion of subjects to women between the ages of 45 and 65 years. Furthermore, we segregated those women consuming fermented soya milk from those consuming the non-fermented product, as we expected changes in intestinal microbial composition in the group of women consuming bifidobacteria.

Mean concentrations of isoflavone excreted in urine by the FS and NFS groups at 4, 13 and 14 days of soya milk supplementation are shown in Table 2, with the concentrations representing the sum of unconjugated forms of daidzein, genistein and glycitein and their deconjugated glucuronide and sulphate isomers. The proportion of unconjugated isoflavone metabolites excreted in urine by the FS and NFS groups ranged between 0.6 \% and 3.0 \% of total urinary isoflavone. Similarly, Zhang \textit{et al.} (2003) found that the percentage of aglycone forms excreted in urine by young adult women after the ingestion of non-fermented soya milk was 4–5 \% of total isomers, with daidzein and genistein glucuronides representing 73 \% and 71 \% of total daidzein and genistein excreted in urine, respectively. As shown in Table 2, women from both the FS and NFS groups excreted the greatest level of isoflavone (38–53 \mu\text{mol/d}) during supplementation with SPI/SG80, which contained the highest dose of isoflavone (Table 1). The level of isoflavone excreted in urine during supplementation with SPI/SG80 was significantly greater than the levels excreted during supplementation with SPI/SG20 and SPI/SG40 for both the FS and NFS groups on day 4 \((P<0.001)\), 13 \((P<0.001)\) and 14 \((P<0.001)\) (Table 2). Hence, a higher ingested dose of isoflavone via SPI/SG80 appeared to enhance the amount of isoflavone absorbed and subsequently excreted during soya milk supplementation. Likewise, Xu \textit{et al.} (1994) reported that total urinary isoflavones increased significantly with increasing dose \((P<0.05)\).

The ingestion of fermented soya milk did not appear to enhance the urinary excretion of isoflavone at any of the isoflavone
Table 1. Concentration of isoflavone isomers (μmol/200 ml serving) in non-fermented and fermented soya milks SPI/SG20, SPI/SG40 and SPI/SG80
(Mean values with their standard deviations; n 18)

<table>
<thead>
<tr>
<th>Isoflavone isomer</th>
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<td>99·85c</td>
<td>8·27</td>
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</table>

SPI/SG20, soya milk containing 20 mg isoflavone/200 ml; SPI/SG40, soya milk containing 40 mg isoflavone/200 ml; SPI/SG80, soya milk containing 80 mg isoflavone/200 ml.

Table 2. Urinary excretion of total isoflavone (μmol/d) by two groups of eight women during supplementation of fermented and non-fermented soya milks SPI/SG20, SPI/SG40 and SPI/SG80
(Mean values with their standard deviations; n 169)

<table>
<thead>
<tr>
<th>24 h urine collection</th>
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<tr>
<td>Baseline</td>
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<tr>
<td>Day 14</td>
<td>19·91± 9·76</td>
<td>22·75± 13·19</td>
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</table>

SPI/SG20, soya milk containing 20 mg isoflavone/200 ml; SPI/SG40, soya milk containing 40 mg isoflavone/200 ml; SPI/SG80, soya milk containing 80 mg isoflavone/200 ml.

Day 14 includes two segregated groups consuming either the fermented or non-fermented soya milk.

† Concentration of isoflavone in 24 h pooled urine specimen.
‡ Mean total of malonyl-, acetyl- and glucoside isomers.
§ Mean total of three respective isomers.
* Inoculated with active culture of Bifidobacterium animalis BB-12 and incubated for 24 h at 37°C.
** Mean total of malonyl (B), acetyl (G) and glucoside (S) isomers.
according to the AUC of the appearance and disappearance concentrations in the plasma, which appeared to be due to the hydrolytic actions of intestinal β-glucosidases. To date, there is no evidence to indicate that ingesting isoflavones in an aglycone-rich configuration enhances their bioavailability after either a single dose or consistent daily consumption.

Following the collection of a urine specimen at day 0, we allowed 3 d of soya milk intake for women to reach a steady state of urinary isoflavone excretion and compared this with 24 h pooled urine specimen collections at the completion of supplementation. There were no significant differences between the levels of total urinary isoflavone and the individual isomers (daidzein, genistein and glycitein) excreted on days 4, 13 and 14 of supplementation with fermented and non-fermented SPI/SQG20, SPI/SQG40 and SPI/SQG80 (P > 0.05). Hence, a steady level of isoflavone absorption and urinary excretion appeared to have been achieved at each isoflavone dose for both fermented and non-fermented soya milk. Setchell et al. (2003) stated that optimum steady-state serum isoflavone concentrations would be expected from modest intakes of soya foods consumed regularly throughout the day rather than from a single, highly enriched product. Each subject in our study was required to consume the daily 200 ml serving of soya milk as two portions, before breakfast and before dinner.

The positive correlation between the mean urinary excretion of total isoflavone per day versus the mean amount of total isoflavone ingested per day during supplementation with non-fermented and fermented soya milk is shown in Fig. 2(a) and 2(b), respectively. According to the computed lines of regression, there was a clearer linear relationship between isoflavone excretion and dose among fermented soya milks (R² = 0.9993) compared with the linearity observed for non-fermented soya milks containing equivalent dosages of isoflavone (R² = 0.8865). Even though the consumption of fermented soya milk did not enhance urinary isoflavone excretion (P > 0.05) (Table 2), it possibly reduced interindividual variability in isoflavone absorption and urinary excretion, indicated by the high linearity in dose response (Fig. 2(b)). This is reflected in Table 2, with values of standard deviation for urinary isoflavone excretion generally higher among the NFS group at each isoflavone dose.

The distinct linear dose response for the FS group (Fig. 2(b)) may have been due to the higher proportion of aglycone structures ingested via the fermented soya milks or the consumption of viable B. animalis Bb-12. Turner et al. (2003) stated that the composition of intestinal microflora appears to play a key role in the intestinal biotransformation and absorption of isoflavones and may cause significant interindividual variability. Those women consuming the fermented soya milk may have established a more consistent intestinal microflora during supplementation by increasing the populations of β-glucosidase-producing bifidobacteria and adjusting intestinal microbial balance in favour of other saccharolytic enzyme-producing lactic acid bacteria. Hence, a greater variation in gut microflora between women consuming the non-fermented soya milks may have caused the poorer linearity in dose response (Fig. 2(a)). Furthermore, non-fermented soya milks probably contained a greater concentration of oligosaccharide compared with fermented soya milk; oligosaccharides are metabolised by B. animalis Bb-12 during fermentation (Tsangalis & Shah, 2004). Soyabean oligosaccharides are classified as prebiotics, which stimulate the growth of bifidobacteria and other lactic acid-producing bacteria in the intestinal tract (Masai et al., 1987). This may have enhanced intestinal microbial metabolic activity in some subjects in the NFS group, potentially influencing the extent of isoflavone biotransformation and absorption. Not to be discounted are the deglycosylation of isoflavones by mammalian intestinal glucosidases (Day et al. 1998; Setchell et al. 2001) and the extent of their effect on interindividual variation in isoflavone absorption within the FS and NFS groups. Furthermore, Rowland et al. (2000) reported that the dietary intake of carbohydrate and fat also influenced the intestinal biotransformation of isoflavones. However, since both the FS and NFS groups had a similar daily intake of carbohydrate (P = 0.7) and fat (P = 0.9), we believe that dietary intake had a lesser influence on interindividual variation in urinary isoflavone excretion within each group and the difference in linearity in isoflavone dose response shown in Fig. 2.

Bioavailability of daidzein, genistein and glycitein from fermented and non-fermented soya milk

The bioavailability of daidzein, genistein and glycitein appears to be dependent on the extent of their biotransformation in the intestinal tract. The deglycosylation of isoflavones by mammalian and microbial glucosidases results in the release of the aglycone form (Heinonen et al. 2002), which can undergo further structural changes. Ioannou et al. (1995) reported that the reductive metabolism of daidzein results in the formation of equol or 0-desmethylangolensin. Reductive metabolism transforms genistein...
into dihydrogenistein and 6'-OH-O-desmethylangolensin (Joannou et al. 1995). Recently, Heinonen et al. (2002) reported that glycitein also underwent reductive metabolism to form structures dihydroglycitein and the equol analogue of glycitein (6-OH-eq). Consequently, pharmacokinetic studies have shown that there are differences in the extent of absorption and urinary excretion of daidzein, genistein and glycitein in human subjects, with large interindividual variability (Xu et al. 1994; King & Bursill, 1998; Lu & Anderson, 1998; Richelle et al. 2002).

The mean concentrations of daidzein, genistein and glycitein (total of conjugated and unconjugated forms) excreted in urine by the FS and NFS groups at 4, 13 and 14d of soya milk supplementation are shown in Table 3, and the percentage of ingested dose of daidzein, genistein and glycitein recovered in urine is shown in Fig. 3. On days 4, 13 and 14, the urinary excretion of daidzein was significantly greater than that of glycitein and genistein during supplementation with both fermented and non-fermented SPI/SG20, SPI/SG40 and SPI/SG80 (P < 0.001; Table 3). Similarly, Richelle et al. (2002) reported that after the ingestion of an SG beverage, postmenopausal women excreted greater concentrations of daidzein in urine, followed by glycitein and genistein, with a urinary recovery of daidzein of approximately 53%, compared with 34% and 19% for glycitein and genistein, respectively. Furthermore, Zhang et al. (2001) fed men and women (between the ages of 19 and 35 years) SG and found that the urinary recovery of daidzein was higher than that of glycitein (P > 0.05), although the recovery of daidzein and glycitein was significantly greater than that of genistein (P < 0.05). Fig. 3 shows that the percentage recovery of daidzein in urine was significantly greater than that of both glycitein and genistein during supplementation with both fermented and non-fermented soya milk at all three isoflavone dosages (P < 0.001). The percentage recovery of glycitein in urine was also greater than that of genistein during supplementation with fermented and non-fermented SPI/SG20, SPI/SG40 and SPI/SG80 (P > 0.05; Fig. 3). Considering the profound differences in gut microflora between subjects from different countries and cultural backgrounds, as well as from different age groups (Mitsuoka, 1984), there appeared to be a consistent pattern in the urinary excretion and percentage recovery of isoflavone isomers found in our study and in the studies by Zhang et al. (2001) and Richelle et al. (2002). That is, the greatest proportion of isoflavone in urine was found as daidzein, the smallest proportion as genistein, and the proportion as glycitein found somewhere in between.

According to the urinary isoflavone profiles shown in Table 3 and Fig. 3, the urinary recovery of genistein appears to be less than that of daidzein and glycitein. This is consistent with previous studies by Xu et al. (1994, 1995) and Lu et al. (1996), substantially reporting a higher urinary recovery of daidzein compared with genistein. Pharmacokinetic studies by King & Bursill (1998) also found the urinary recovery of daidzein (62%) to be significantly greater than that of genistein (22%; P < 0.001). However, King & Bursill (1998) concluded that the bioavailabilities of daidzein and genistein were similar considering that the ratio of the plasma AUC for genistein and daidzein was equal to the ratio of the concentration of the respective isoflavones in the soya meal ingested. Setchell et al. (2001) administered a 50 mg dose of daidzein, daidzin, genistein or genistin to four groups of premenopausal women and found that the systemic bioavailability of genistein (mean AUC of 454 μg/ml per h) was much greater than that of daidzein (mean AUC of 294 μg/ml per h). Nevertheless, the results from previous urinary and plasma pharmacokinetic studies only involved the administration of a single dose. Over a period of daily soya isoflavone ingestion, the bioavailability of daidzein, genistein and glycitein may begin to differ depending on the response of intestinal microflora to the presence of these compounds. The lower concentrations of genistein in urine may be the result of its greater susceptibility to microbial breakdown in the intestinal tract. Xu et al. (1995) reported that genistein was more susceptible than daidzein to bio-transformation by intestinal bacteria during anaerobic incubations of human faecal samples. However, as we did not measure concentrations of these isoflavones in timed plasma samples, we cannot assume a reduced bioavailability of genistein based on percentage urinary recovery alone. Genistein may have been absorbed more extensively than suggested by its urinary recovery but may have undergone greater metabolism and/or excretion by non-renal mechanisms.

Women consuming both the fermented and non-fermented SPI/SG80 excreted significantly greater levels of daidzein (ranging from 28 to 37 μmol/d) on days 4, 13 and 14 of supplementation, compared with the consumption of SPI/SG20 and SPI/SG40 (Table 3). The greater level of daidzein and its glucosidic conjugates in SPI/SG80 (Table 1) appeared to account for the increased absolute absorption and urinary excretion of this isomer in the FS and NFS groups. However, the percentage recovery of daidzein in urine tended to be greater for women consuming fermented and non-fermented SPI/SG20 (57% and 63% of dose, respectively; P > 0.05) (Fig. 3). Concentrations of daidzein and its glucosides at approximately 22.6 μmol (5.7 mg) aglycone constituents per 200 ml, found in SPI/SG20, appear to be less susceptible to microbial breakdown (reductive metabolism) in the gastrointestinal tract and more bioavailable according to percentage recovery of dose. Lower doses of daidzein may be absorbed more effectively in the proximal gastrointestinal tract, reflecting more complete absorption. With higher doses of daidzein, the fraction not absorbed proximally may be absorbed distally in the gastrointestinal tract, where there are greater populations of intestinal bacteria possibly biotransforming daidzein into equol, dihydrodaidzein and other metabolites prior to absorption, accounting for the lower percentage recovery of dose for SPI/SG40 and SPI/SG80 (Fig. 3). There was a trend towards an increased excretion and urinary recovery of equal during supplementation with SPI/SG80 (P > 0.05), especially in the FS group (results not shown). Limited or saturable absorption of daidzein in the proximal gastrointestinal tract may be the reason why higher ingested doses of daidzein pass into the ileum and colon. This may be influenced by the efficacy of mammalian glucosidases present in the proximal intestinal tract to hydrolyse glucosidic isomers of daidzein into a bioavailable aglycone configuration, as there are considerably lower populations of bacteria in this region compared with the ileum and colon (Turner et al. 2003).

In contrast to the urinary excretion of daidzein, Table 3 shows that there were no significant differences in the urinary excretion of genistein during supplementation with fermented and non-fermented SPI/SG20, SPI/SG40 and SPI/SG80 at day 4, 13 and 14 (P > 0.05). In this case, increasing the dose of genistein was not associated with increased urinary excretion of this isomer, due possibly to its greater susceptibility to intestinal microbial breakdown in comparison to daidzein (Xu et al. 1995); alternatively, this could reflect a limited or saturable absorptive capacity in.
Caused the similar percentage recovery of genistein in urine between fermented and non-fermented SPI/S20, SPI/S40 and SPI/S80, in the range of 14–17 % of ingested dose (P>0.05; Fig. 3). In comparison, Hutchins et al. (1995) reported that the mean urinary recovery of genistein among a group of men consuming fermented soyabeans (tempeh) was only 1.9 % at a dosage of 44.5 μmol genistein/d, equivalent to the dosage of fermented SPI/S80 (44.4 μmol genistein/d). The mean urinary recovery of daidzein from the same group was 9.7 % at a dosage of 21.3 μmol daidzein/d (Hutchins et al. 1995), considerably lower than the mean urinary recovery of daidzein among the FS group (57 % of ingested dose) ingesting an equivalent dosage of daidzein via SPI/S20 (23.3 μmol daidzein/d). The greater percentage recovery of daidzein and genistein in our study may have been due to the greater proportion of unconjugated daidzein in fermented SPI/S20 (73 % of dose) and unconjugated genistein in fermented SPI/S80 (53 % of dose). According to Hutchins et al. (1995), tempeh comprised 22 % and 10 % of its total daidzein and genistein concentration, respectively in an unconjugated form. Alternatively, soya milk has a greater proportion of simple isoflavone glucosides (β-glucoside isomers) than do less processed soya foods, which are possibly easier to hydrolyse into bioavailable aglycones via mammalian and microbial glucosidases (King & Bursill, 1998).

Even though the urinary excretion of glycitein resembled a linear response to dose in the FS group, there were no significant differences at day 4, 13 or 14 to suggest that increasing the dosage level of glycitein significantly enhanced the absorption and urinary excretion of this isomer (Table 3). As with the urinary recovery of daidzein, a significantly greater percentage recovery (in the range of 30–35 %) was observed when ingesting the lowest level of glycitein via fermented and non-fermented SPI/S20.
non-fermented SPI/SG40 (urinary recovery of 41%; SG40 (48% of dose) compared with women consuming fermented SPI/SG40 tended to be greater among women consuming non-fermented SPI/SG40 (urinary recovery of 41%; $P=0.13$). Similarly, the percentage recovery of daidzein in urine also tended to be greater among women consuming fermented SPI/SG40 (48% of dose) compared with women consuming non-fermented SPI/SG40 (urinary recovery of 41%; $P=0.14$).

In an earlier crossover study, Hutchins et al. (1995) assigned an adequate 12 d washout period between each of their 9 d feeding periods. Since our study involved daily soya milk ingestion for up to 14 d, we decided to implement a similar time period to our washout phase as used by Hutchins et al. (1995). Previous pharmacokinetic studies involving a single serving of soya food generally used shorter washout periods of 5–6 d (King & Bursill, 1998; Richelle et al. 2002), which we believed may not have been adequate considering the longer exposure to isoflavones and the potential for enterohepatic recirculation. Nevertheless, urinary isoflavones were detected on day 0 (baseline) of each of the isoflavone dosages ingested (Table 2), suggesting that the 14 d washout period implemented in the crossover design was not long enough to fully excrete circulating isoflavone metabolites in each of the women. Alternatively, isoflavones detected in baseline urine specimens may have been the result of women unknowingly ingesting foods or ingredients containing isoflavones, which were supposed to be excluded during the supplementation and washout periods. This might have been clarified had the subjects recorded their dietary intake during the washout periods in order to assess whether the presence of isoflavones at baseline was the result of physiological or dietary reasons. Of the individual isolomers, low concentrations of both daidzein and genistein were detected at baseline for both the FS and NFS groups (Table 3). In contrast, glycitein was not detected in baseline urine specimens (Table 3), 14 d appearing to be an adequate period to wash out glycitein from the circulatory system. Then again, if dietary intake during the washout phases was the reason for the presence of isoflavones in the baseline urine specimens, glycitein may not have been detected because it is found at considerably lower levels than daidzein and genistein in conventional soya foods and ingredients found in Australia (King & Bignell, 2000).

From a health perspective, isoflavones may be of greater benefit to those women who are able to retain biologically active isoflavone metabolites in their circulatory system for longer periods of time after discontinuing the intake of soya foods. In the NFS group, the woman who excreted the greatest level of isoflavone after two 14 d washout periods also had the highest BMI, at approximately 39 kg/m². Isoflavones in an aglycone form are lipid soluble; hence, women with a higher percentage of body fat may have a greater tendency to retain isoflavones.

Conclusions

The fermentation of each soya milk with B. animalis Bb-12 enriched the level of bioavailable isoflavone aglycone. Additionally, each soya milk was able to support the viability of B. animalis Bb-12 during refrigerated storage at levels able effectively to modulate intestinal microbial balance. There was, however, no strong evidence to suggest that fermenting soya milk with bifidobacteria improved the bioavailability of isoflavone in postmenopausal women over 14 d of daily soya milk ingestion. Levels of total isoflavone (unconjugated and conjugated forms) excreted in urine and urinary isoflavone recovery was similar for both groups of women consuming either fermented or non-fermented soya milks. Nevertheless, the ingestion of fermented soya milk appeared to reduce the interindividual variation in isoflavone absorption. Of the individual isolomers, the percentage recovery of daidzein and genistein in urine tended to be greater among women consuming fermented soya milk at a daily dosage of 40 mg isoflavone. Increasing the dosage of isoflavone correlated positively with the urinary excretion of isoflavone, but the urinary recovery of isoflavone was inversely related to dosage level. Hence, a modest isoflavone dosage ranging from 20 to 30 mg aglycone constituents per day may provide the most bioavailable source of isoflavone. Finally, according to the isoflavone profiles excreted in urine, there is no preliminary evidence to indicate that ingesting soya milk fermented with bifidobacteria will enhance the potential health benefit of isoflavone compared with an equivalent non-fermented soya milk. Studies investigating the physiological effects of the concomitant ingestion of isoflavone and probiotic bifidobacteria on biomarkers of disease risk in postmenopausal women are currently underway.

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References

Gomes AMP & Malcata FX (1999) Bifidobacterium spp. and...


