Effects of soaking and grinding conditions on anti-nutrient and nutrient contents of soy milk

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Abstract

Although soy milk is a very good source of nutrient with high biological value, the presence of anti-nutritional factors affects its nutrition quality and limits bioavailability of the nutrients. The effects of soaking duration and combination of soaking and grinding (hot or cold) on phytate, lipoxgenase, urease, trypsin inhibitor activity, protein solubility and other nutrient contents were investigated. Soaking alone at 55 and 60°C for different durations was found effective for the reduction of lipoxgenase activity. Combination of soaking, blanching (80°C for 10 min) and hot grinding (100°C) significantly (P<0.05) reduced urease activity, more than 80% phytate activity and deactivated trypsin inhibitor, but did not affect protein solubility. Meanwhile, protein solubility (10–15%) was increased due to hot grinding. Soy milk extracted from soaking at 55 and 60°C for 2, 4 and 6 h with hot grinding provided higher protein content compared to cold grinding. Increase in soaking temperature from 55 to 60°C increased the extracted solid content having a potential fraction of lipid. Increasing soaking time from 4 to 6 h did not show any significant difference in terms of phytate inhibition, urease activity reduction, trypsin inhibition and protein solubility except lipoxgenase activity. The results suggested that soaking of soybean at 60°C for 6 h and hot grinding (100°C) with blanching at 80°C for 10 min is the best for reducing anti-nutrient and retaining nutrient activity for soy milk and other soy-based products.

Introduction

Soybean foods have become increasingly popular since the US Food and Drug Administration (FDA) approved a health claim for the cholesterol-lowering effects of soy protein. Their claimed largely based on a meta-analysis of 38 clinical trials that reported significant decreased in total and low-density lipoprotein (LDL) cholesterol and triglycerides with soy protein intake (25 g/day) compared with animal protein consumption (Adlercreutz et al., 1995). Researches revealed that soy products could prevent heart disease, obesity, blood cholesterol, cancer, diabetes, kidney disease, osteoporosis and blood pressure (Garcia et al., 1997; Hassler, 1998; Liu, 1997; Riaz, 1999). Soy milk is a water extract of soybean, a grain legume and one of the oldest known food sources of the world of human beings. It is typically produced by grinding soaked soybeans with water. It is an inexpensive and convenient source of high quality protein. Soy milk is one of the most important traditional beverages that are consumed widely in Asian countries, including China, Japan, Korea, Singapore and Thailand. In recent decades, extensive evidence has indicated the strong relationship between soy food consumption and health-promoting effects. Soy milk provides a balanced nutrient combination, which is similar to cow’s milk, but free of cholesterol, gluten and lactose plus favorable phytochemical compounds linked to health. Among vegetarians, milk allergy patients or people with lactose intolerance, soy milk could be used as a daily alternative. As soy milk contained high amounts of protein, poly-unsaturated fatty acids, vitamins, minerals and phytochemicals, it could be easily used as a good source of nutrition food for malnourished people, especially in developing countries (Mazumder and Begum, 2016). In response to a gradual increase in sales and consumption, various new products have been introduced into the soy market. Some basic changes are made to the flavor and soybean source. However, the most recent innovations are focused on producing “functional soy milk”. Functional soy milk can be considered as soy milk that contains extra bioactive components and may help to enhance health or lower risk of diseases. Soybean is a good source of phenolic compounds with antioxidant properties and has an extraordinarily high amount of isoflavones, a group of phytosterogens that have been reported to possibly lower the risk of hormonal and age-related diseases. Among isoflavones, genistein being a powerful inhibitor of tyrosine carries additional small molecular modifiers, such as kinase activity in vitro (Akiyama et al., 1987). More importantly, genistein could act as an anti-oxidant and anti-browning agents in in vivo and in vitro (Mazumder and Hongsprabhas, 2016a). However, the presence of natural anti-nutrients, such as trypsin inhibitors (TI), lectins, phytic acids, and indigestible oligosaccharides, has limited the consumption of soybean and its products.

The presence of protease inhibitors, Kunitztrupsin inhibitor (KTI) and Browman-Birk inhibitor of

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chymotrypsin (BBI) results in the reduction of protein digestibility and thus limits the availability of essential amino acids such as trypsin (Dia et al., 2012; Monteiro et al., 2004). Phytic acid might reduce the bioavailability of proteins by binding to peptides and amino acids and thereby inhibits proteolytic enzyme action (Deak and Johnson, 2007). Moreover, lipoxygenases lead to the so-called beany flavor by catalyzing lipid hydroperoxidation and thus limit the consumption of soy products (Esteves et al., 2010; Felix et al., 2011). So, modifying the processing methods could be an effective way to improve the health-promoting bioactive components and/or reduce the undesired compounds originally present in soybeans in order to support functional soy milk product development. The soaking and heating process during soy milk extraction considerably destroys most of the anti-nutritional factors present in soybean and improves the digestibility of soy protein (Mazumder, 2016). This research aimed to evaluate the effects of soaking time-temperature and hot grinding on the anti-nutritional factors such as phytate, lipoxygenase, urease activity, protease and trypsin inhibitor, protein solubility and other nutrient content of soy milk.

Materials and Methods

Soy milk extraction

Soy milk was extracted by the modified method described by Mazumder (2016). Prior to grinding, the soybeans were soaked in 0.5% sodium bicarbonate solution at 55 and 60°C for 2, 4 and 6 h in a water bath (JSSB-30T, Korea). The ratio of soybean and water for soaking was 1:2. The beans were drained well after wards. After discarding water, the soaked soybean was dehulled manually with hands to remove unwanted substances before grinding. Hydrated beans were blanched in 0.5% sodium bicarbonate solution at 80°C for 10 min. The solution was then drained well and washed with potable water for three times. The blanched soybean was ground with addition of hot water (100°C) or cold water by using super mass colloidier and a basket centrifuge. The ratio of soybean to water was 1:4. Soy milk was obtained after filtering through double layers of cheese cloths. The soy milk was stored at -18°C for further studies. Soy milk prepared by different methods was analyzed for total solid, protein, fat, protein solubility, phytate, lipoxygenase, urease activity and trypsin inhibitor. Moisture content for total solid, protein and fat content was determined following AOAC (1999).

Determination of lipoxygenase (LOX) assays of soy milk

One hundred milliliter (100 ml) of soy milk was centrifuged at 25,000 x g for 20 min at room temperature. The supernatant was used for the crude enzyme extract. LOX activity was measured as the change in absorbance at 234 nm with a linolenic acid substrate solution according to Gokmen et al. (2002). A double beam spectrophotometer (Photolab 7600UV-VIS) and 1 cm path-length cuvette were used for enzyme measurement.

Determination of phytate activity of soy milk

Extraction of phytate was done for 1 h in a 20 ml vial with 0.5 N HCl in a ratio of 1:20 (w/v) while stirring. 0.5 ml of soy milk and 10 ml of 0.5 N HCl was used throughout. Approximately 2 ml of crude extract of each sample was centrifuged at 18,000 x g for 10 min in a micro-centrifuge. An aliquot of 1 ml of supernatant containing phytate was then filtered with a 1 ml tuberculin syringe and a 13-mm/0.45-µm syringe filter. Filtered samples could be stored at 4°C for several days prior to HPLC analysis.

Phytate was determined by the modified method as described by Rounds and Nielsen (1993). Elution of phytate for HPLC analysis was achieved by using a 30-min linear gradient of 0.01 M 1-methylpiperezaine at pH 4.0 and 0.5 M NaNO₃ in 0.01 M 1-methylpiperezaine at pH 4.0 with a flow rate of 1 ml/min. Wade’s color reagent consisting of 0.015% (w/v) FeCl₃ and 0.15% (w/v) 5-sulfosalicylic acid was prepared based on Wade and Morgan (1955) having a flow rate of 1 ml/min. Phytate eluted from the column were mixed with Wade’s reagents in a mixing tee with inline check valves for both eluents installed prior to the mixing tee to prevent backflow. The post column reaction was allowed to take place in 0.05 × 210 cm poly ether ketone tubing at a combined flow rate of 2 ml/min. The absorbance was monitored at 500 nm and the detector signals and/or phytate peaks were processed and integrated by chromatographic data acquisition system.

Determination of urease activity of soy milk

The urease activity was determined by an assay method described by Croston et al. (1955). The pH of the urease-urea reaction should maintain 6.8 to prevent a decrease in urease activity with increasing alkalinity. All reactions were carried out in a constant temperature bath at 40°C. The reagents used were 0.1 N HCl, 0.1 N NaOH, phosphate buffer of pH 6.8 made up of 0.025 mole of K₂HPO₄, 0.025 mole KH₂PO₄ and 0.8 g of glutathione per liter and the buffered urea solution was prepared daily by dissolving 6 g urea in 100 ml of buffer solution without the glutathione. Twenty milliliter (20 ml) of soy milk was added to 5 ml of the buffer solution containing glutathione, and was allowed to stand for 30 min at 40°C. Five milliliters (5 ml) of the buffered urea solution was then added to initiate the reaction. The pH of the reaction was maintained at approximately 6.8 by slowly adding 0.1 N HCl and using bromthymol blue in its green color range to indicate the desired end-point. At the end of 30 min, the reaction was terminated by rapidly adding additional 0.1 N HCl to a total of 10 ml or more. The system was then titrated with 0.1 N NaOH to pH 4.7. A control was
run parallel with each sample. For the control, the urease was inactivated by adding HCl to the sample before adding phosphate buffer and buffered urea solution. The difference between the control and sample in ml of 0.1 N HCl or its ammonia equivalent was taken as the urease activity of the meal.

**Determination of trypsin inhibitor in soy milk**

Trypsin inhibitor activity was determined according to the modified method described by Erlanger et al. (1961). Fifty milligram (50 ml) of soy milk and 5 ml of 0.1 M Tris-HCl buffer were mixed together to adjust pH 8.2. The solution was homogenized in an Erlenmeyer containing 20 mM CaCl₂ and agitated for 3 h followed by centrifugation for 3600 x g for 20 min. One hundred milliliter (100 ml) of soy milk was added with 450 µl of buffer and 50 µl of trypsin solution in a test tube and homogenized followed by keeping the suspension at room temperature for 10 minutes. 500 µl of the homogenate was transferred to a new test tube containing 500 µl of buffer and 500 µl of D, L-BAPNA solution. The solution was agitated for few minutes and left at room temperature for 10 minutes; the reaction was stopped by adding 300 µl of 60% acetic acid. Absorbance of the solution was determined at 410 nm using a spectrophotometer (Photolab 7600UV-VIS) and the results were converted in milligram of inhibited trypsin per gram of total protein in the sample.

**Determination of protein solubility of soy milk**

Protein solubility of soy milk was determined based on nitrogen (N) solubility (Mazumder, 2016). Fifty milliliter (50 ml) soy milk was centrifuged at 10,000 x g for 10 min at room temperature to separate solid and liquid. The suspension was pipetted and N solubility was determined by using Lowry method (Lowry et al., 1952). Protein solution (0.3 ml) was added in test tubes with 0.3 ml 2 M NaOH. The solution was heated at 100°C for 10 min and cooled at room temperature (25°C). Three milliliter (3 ml) of complex forming agent (2% sodium carbonate, 1% copper sulphate and 2% sodium potassium tartarate) was added to the solution. The solution was then mixed well. This solution was incubated at room temperature for 10 min. Then 0.3 ml of Folin-Ciocalteau solution (reagent solution) was added to each tube and incubated for 30 min (not more than 60 min). Absorbance of the solution was determined at 550 nm using a spectrophotometer (Photolab 7600UV-VIS). The absorbance was plotted against protein concentration to get a standard calibration curve and calculate percent nitrogen solubility.

**Results and Discussion**

**Effect of combination of soaking, blanching and hot grinding on lipoxygenase (LOX) content of soy milk**

Figure 1 illustrates the residual enzyme activities during soaking of soybean at different soaking conditions. Soaking at 55°C for 6 h showed around 85% reduction of LOX activities. Rapid inactivation was observed in soybean LOX at 60°C for different duration. Soaking at 60°C for 6 h was found sufficient to inactivate 100% of LOX's activity. As 100% LOX activity reduction was achieved by hot soaking, the effects of cold or hot grinding studies did not continue for LOX activity. The result suggested that soaking at 60°C for 6 h was sufficient to reduce 100% LOX activity. Bahceci et al. (2005) found that 60°C could reduce 90% LOX activity within the first 10 min, and the residual activity remained relatively stable during further 20 min.

Reduction of >90% LOX activity was recommended for optimum quality of vegetable during frozen storage. However, the extraction of soy milk requires 100% inhibition of LOX activity since soybean contained lot of anti-nutritional factors such as saponins, phospholipids, protease inhibitors, phytates and trypsin inhibitors (Giri and Mangaraj, 2012). Hence, blanching is an obvious pre-treatment during extraction of soy milk.

**Effect of combination of soaking, blanching and hot grinding on phytate content of soy milk**

Figure 2 illustrates the phytate activities during soaking of soybean at different soaking conditions and at combination of soaking, blanching and hot grinding. Only soaking at 55°C for 6 h showed around 21% reduction of phytase activity though it is not significantly (p>0.5) different with that at 55°C for 4 h. Although soaking at 60°C for 6 h showed 27% reduction of phytase activity, it is statistically similar with that at 60°C for 4 h. The result suggested for further heat treatment of soybean before extraction of soy milk. Blanching temperature was set at 80°C for 10 min based on Mazumder and Hongsprabhas (2016b) and Mazumder and Begum (2016). Combination of soaking, blanching (80°C for 10 min) and cold or hot grinding (100°C) showed rapid inactivation of soybean phytate. Soaking at 55 and 60°C for different time periods showed significant difference in phytate reduction, both in cold and hot grinding. Similarly, cold grinding and hot grinding also showed significant difference in phytate reduction. Maximum amount of phytate was reduced with a combined treatment of soaking (60°C for 6 h), blanching (80°C for 10 min) and hot grinding (100°C).
Most of the anti-nutrients in the beans and legumes are found in the skin, and many of them are water-soluble; they simply dissolve when soaked in water (Fernandes et al., 2010). In legumes and beans, soaking has been found to reduce phytate, protease inhibitors, lectins, tannins and calcium oxalate. Twelve-hour (12 h) soaking reduced the phytate content of peas by up to 9% (Bishnoi et al., 1994). However, the reduction of anti-nutrients might depend on the type of beans. In kidney beans, soybeans and faba beans, soaking reduces protease inhibitors only very slightly (Dhurandhar and Chang, 1990; Liu and Markakis, 1987; Sharma, 1992). Similarly, soaking is useful for leafy vegetables to reduce some of their calcium oxalate (Savage and Dubois, 2006).

It is very essential to reduce phytate in soy milk or soy food because phytate inhibits calcium absorption. It reduces iron, zinc, magnesium and calcium absorption, and resulted in poor iron absorption in soy foods (Mazumder and Hongsprabhas, 2016b; Schlemmer et al., 2009). But, vitamin C could increase the amount of iron absorbed from soy foods, although absorption rates are still low. Iron may be better absorbed from fermented soy foods like tempeh and miso than soy milk (Mazumder and Hongsprabhas, 2016b). Reduction of around 90% phytate activity will help to absorb calcium content of soy milk. To increase the calcium absorption, calcium-fortified soy milk should be supplied in the markets.

**Effect of combination of soaking, blanching and hot grinding on urease activity of soy milk**

The urease activity of the raw soybean was significantly higher (p<0.05) than that of the soybean soaked at 55 and 60°C for different time periods. Soaking at 55°C for 6 h showed maximum 45% reduction of urease activity (not significantly different with 55°C for 4 h, 44% reduction), while 60% reduction was observed for soaking at 60°C for the same time period (not significantly different with 60°C for 4 h, 58.5% reduction, Figure 3). This result suggests that soaking temperature and time were not sufficient to destroy 100% urease activity; further heat treatment was necessary to inactivate urease activity. Combination of blanching (80°C for 10 min) and soaking at 55 and 60°C for 4 and 6 h did not show any significant (p>0.05) different in urease activity. Cold grinding in combination with soaking and blanching did not reduce 100% urease activity. However, Combination of hot grinding (100°C), blanching (80°C for 10 min) and soaking at different time and temperature could significantly reduce 100% urease activity. Heat treatment was adequate to reduce urease activity that might be a good indicator of trypsin inhibition.

**Effect of combination of soaking, blanching and hot grinding on trypsin inhibitor content of soy milk**

Extraction under combination of soaking, blanching and hot grinding (100°C) significantly reduced more trypsin inhibitor in soy milk than extraction under combination of soaking, blanching and cold grinding (Figure 4). The heat treatment (100°C) used for hot grinding extraction was not sufficient for complete destruction or inactivation of trypsin inhibitor. The result suggests that high heat treatment or steam grinding rather than hot water grinding are more effective for soy milk extraction. Heat-treatment in an oven at 150°C for 30 min for raw soybean before soy milk extraction or UHT for extracted soy milk will completely destroy trypsin inhibitor. Similarly, inactivation of inhibitors to desirable levels could be achieved after autoclaving soybeans at 120°C for 18 minutes and 121°C for 10 minutes, respectively (Machado et al., 2008; Mendes et al., 2007). However, wet heat treatment at high temperature could cause protein denaturation with concomitant loss of its functionality and generates additional volatile organic compounds that are responsible for cooked or toasted off-flavors (Ha et al., 1992). It is also known that the denaturation of protein by heating increases hydrophobicity (Shung-Tang et al., 1997). However, steam injected hot grinding could be useful for extraction of soy milk that would reduce nutrient loss of soy milk as well as inhibit the maximum amount of trypsin inhibitor.
Soaking of soybeans in chemical solutions had a significant effect on the chemical composition and reduction in beany flavor in soy products (Khaleq et al., 1970). Soy milk prepared from soybeans soaked in sodium-bi-carbonate contained higher protein than soy milk prepared from beans soaked in water or sodium hydroxide (data not shown). Soybeans soaked in carbonate were easier to process than soaked in the other two solutions. However, sodium carbonate and sodium hydroxide had a significant effect on the reduction in beany flavor in soy milk (Giri and Mangaraj, 2012).

Soaking temperature and time with cold or hot grinding significantly affect the nutrient content of soy milk. Combination of soaking, blanching and hot water grinding significantly reduced carbohydrate content and increased the lipid with increasing extraction temperature from 55 to 60°C (Table 1). Maximum solids were extracted at 55 to 60°C. Protein recovery started to decrease slightly at more than 70°C. With higher extraction temperatures above 70°C, there was an indication that the insoluble carbohydrate fraction, with hydroscopic and swelling properties, might significantly contribute to inhibition of filtration and cause losses in the solid yield of soy milk. However, the effect of short-duration soaking was predominantly leaching of water-soluble carbohydrate accounted for as much as 60% of the solids contained in the soaked water. About 6% of the soaked water solids was lipid and the remainder was crude protein. About 50% of soaked water crude protein was non-protein nitrogen. Increasing the soaking temperature from 55 to 60°C significantly increases the extracted solid content. Increasing soaking time from 2 to 4 h, total solid content (protein and lipid) significantly increased; however, 4 to 6 h soaking time did not show any significant difference in terms of solid content.

### Table 1. Refractive index, total solids, protein, fat and yield of soy milk per 250 g soybean under combination of soaking (4 h), blanching (80°C for 10 min) and grinding conditions

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Hot water grinding (100°C)</th>
<th>Cold water grinding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Refractive index (%)</td>
<td>1.345±0.005</td>
<td>1.333±0.004</td>
</tr>
<tr>
<td>Total solids (%)</td>
<td>7.65±0.03</td>
<td>6.37±0.05</td>
</tr>
<tr>
<td>Protein (%)</td>
<td>4.46±0.07</td>
<td>4.02±0.05</td>
</tr>
<tr>
<td>Fat (%)</td>
<td>2.15±0.06</td>
<td>1.80±0.00</td>
</tr>
<tr>
<td>Yield (ml) per 250 g soybean</td>
<td>850±5.0</td>
<td>950±4.0</td>
</tr>
</tbody>
</table>

Mean ± standard deviation values in the same column with different superscripts are significantly different (p > 0.05).
Conclusions
Soy milk is a good source of protein, fat, and mineral contents. Soaking time and temperature along with hot or cold grinding reduced the anti-nutrient factors in soy milk. Soaking conditions did not considerably affect nutrient contents of soy milk; however, soaking temperature affected the nutrient content of soy milk. Hot grinding (100°C) was found sufficient to reduce 100% of urease activity, more than 85% phytate and trypsin inhibitor, and increase protein solubility. It is concluded that soaking at 60°C for 6 h in combination with hot grinding (100°C) with blanching at 80°C for 10 min is the best practice for producing high quality soy milk and other soy products.

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References


