Full Length Research Paper

Quantification of lactic acid bacteria and bifidobacteria in goat milk based yoghurts with added water-soluble soy extract

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This work was carried out with the objective of evaluating the microbiological aspects, during 29 days of storage, through the quantification of viable cells and probiotic bacteria enumeration, of strawberry flavored yoghurts produced with goat milk with water-soluble soy extract and Bifidobacterium animalis ssp. lactis probiotic culture during processing. Determination of the viable cell count during yoghurt storage showed that the number of lactic bacteria found was below the expected only in the treatment in which there was water-soluble soy extract addition, not presenting microbiological viability. The other yoghurt treatments were viable during storage. The enumeration of Bifidobacterium animalis ssp. lactis showed that the yoghurt treatments presented microbiological viability during storage. We concluded that the addition of water-soluble soy extract interfered negatively with the production of goat milk based yoghurts.

Key words: Yoghurt, water-soluble soy extract, lactic acid bacteria, Bifidobacterium.

INTRODUCTION

Yoghurt is a very popular fermented milk product produced by lactic acid fermentation of milk by addition of a starter culture containing Streptococcus thermophilus and Lactobacillus delbrueckii ssp. bulgaricus. It is a very versatile product that suits all palates and meal occasions (Isleten and Karagul-Yuceer, 2006). Besides the traditional culture, strains of other probiotic organisms, such as lactobacilli and bifidobacteria, have been used in fermented products as potential health promoters. The supplementation of fermented products with probiotic bacteria becomes beneficial by providing better use of the lactose, anticarcinogenic activity and intestinal infection control. Probiotics are referred to as "live microorganisms, which when administered in adequate amounts confer a health benefit on the host" (FAO/WHO, 2001; Allgeyer et al., 2010).

Strains of L. acidophilus and of Bifidobacterium lactis predominate in commercial probiotic products (Tabasco et al., 2007). The presence of multiple species in these products makes the differential enumeration between probiotic bacteria and starter cultures necessary. Numerous means have been proposed for selectivity and differential enumeration of lactobacillus and bifidobacteria in bacterial population mixtures.

In relation to other types of milk, goat milk presents advantages such as smaller size fat globules, low allergenic properties (Martín-Diana et al., 2003), a balance of essential amino acids, high levels of calcium, selenium, phosphate and rich in vitamins A and B. However, goat milk is deficient in folic acid and vitamin D. Furthermore, a certain therapeutic value in human nutrition has been attributed to goat milk (Alférez et al.,
The use of goat milk as an excellent food source is undeniable. It has beneficial effects on health maintenance, physiological functions, in the nutrition of children and elderly people, and according to some authors, can be consumed without negative effects of suffering from cow milk allergy. This highlights the market potential of goat milk (Ribeiro and Ribeiro, 2010).

The elaboration of the yoghurts based on goat milk, can present significant alterations in its rheological properties, such as low consistency and a tendency towards whey separation (syneresis). Therefore, to obtain satisfactory results with fermented goat milk products, the addition of stabilizers is recommended. The enrichment of the dry matter content and/or the protein content are standard measures used to avoid syneresis and to improve the texture of the yoghurt. As such, the soy proteins, specifically, the water-soluble soy extract (WSSE), deserves prominence to improve the nutritional value of the product and to affect the gel structure formation of the yoghurt (Silva et al., 2012).

The nutritional deficiency of goat milk can be improved by the lactic fermentation process. According to Hugenholtz (2008), many lactic bacteria seem to produce some vitamins, where the fermented product is enriched as a result of the bacterial production. The fermented milk products are reported as containing high amounts of folates, as a result of the additional folate production via bacteria. Therefore, from the choice of viable starter cultures, the natural fortification of a milk product can be undertaken. Silanikove et al. (2010) reported that the quality and safety of goat milk are optimized to ensure consumer confidence owing to the growing interest in existing and new goat dairy products worldwide.

Based on the above, this research had the objective of evaluating the growth of total lactic acid bacteria and B. animalis ssp. lactis in goat milk based yoghurts with added water-soluble soy extract and probiotic culture during 29 days of refrigerated storage.

**MATERIALS AND METHODS**

**Characterization of the raw material**

The milk used in the experiment was through the milking of Saanen breed females. After milking, the milk was cooled immediately to 5°C, transferred to previously sanitized polypropylene milk cans and transported to the dairy plant where the processes were conducted.

The physiochemical analyses were carried out with milk samples in triplicate, to proof its quality. It consisted of the determination of the pH by direct potentiometry in a digital pH meter (Instituto Adolfo Lutz, 1985), titratable acidity, density, fat and total soluble solids percentage (Brazil, 2006).

**Preparation of the yoghurt**

The methodology used in this study for development of the yoghurt was based on that described by Tamime and Robinson (1991) as shown in Figure 1. The yoghurts were prepared and identified with letters according to their processing specifics (addition of WSSE adjusted to the protein level of the milk at a concentration of 20%, resulting in the supplementation of 14.8 g/L of water-soluble soy extract and B. animalis ssp. lactis probiotic culture (2%), as expressed below:

1. Yoghurt A: without the WSSE addition, without addition of probiotic culture.
2. Yoghurt B: addition of WSSE (20%), without the probiotic addition.
3. Yoghurt C: without addition of WSSE with added probiotic culture.
4. Yoghurt D: with WSSE (20%) addition with added probiotic culture.

After analysis of the milk quality, WSSE was added in B and D treatments. Later, all the treatments were pasteurized at 80°C for 30 min and then cooled to 43°C. The different combinations of bacterial cultures: S. thermophilus, L. delbrueckii ssp. bulgaricus and B. animalis ssp. lactis, were inoculated. The final fermentation point was determined when the yoghurts reached pH 4.6 and were then removed from the oven and cooled to 15°C for the addition of the strawberry pulp and stored at 4°C. The microbiological analyses of the products were conducted on the 1st, 8th, 15th, 22nd and 29th day, post-production.

**Viable lactic bacteria count during storage at 4°C**

The lactic bacteria count was conducted by the pour plate method, with an overlay, adding 1 mL of diluted inoculum and adding a small amount of MRS agar in the Petri dishes. After solidification of the medium, an overlay was added, seeking the creation of a 15% CO2 atmosphere, followed by incubation at 30°C for 5 days. After the required incubation time, the count was conducted in Petri dishes that presented between 25 and 250 colonies (Tebaldi et al., 2007).

**Probiotic count during storage at 4°C**

For the enumeration of B. animalis ssp. lactis the MRS medium was used supplemented with 1% raffinose, 0.05% lithium chloride and 0.05% cytoene, sterilized by filtration in a 0.45 mm membrane. There was no use of antibiotics and the inoculation technique used was the pour plate, with overlay. After the inoculation, the Petri dishes were incubated inverted in jars containing anaerobiose generator (Anaerobac) at 45°C for 72 h (Tabasco et al., 2007).

**Statistical analysis**

The experiment was conducted in a completely randomized design (CRD), in a 2 x 2 x 5 factorial (2 soy extract concentrations, 2 treatments with probiotic culture addition at 5 storage times). Three repetitions were carried out for each treatment. The effects of the different treatments were appraised by variance analysis (ANOVA), followed by the Scott-Knott Test, to 5% of significance, to express the differences in significant cases. The evaluation of the storage time of the yoghurts was analyzed through linear regression, after transformation of the data into logarithms (log). The variance analyses, test of averages and linear regression were carried out in the R software (R Development Core Team, 2009).

**RESULTS AND DISCUSSION**

**Characterization of raw material**

The results of the physiochemical analysis of the goat
milk used in the yoghurt treatments are 6.68 from pH, 0.17 g lactic acid /100 mL of product from titratable acidity, 1.02 g L⁻¹ from density, 3.1% from fat and 11.5% from total soluble solids. These results were similar to those found by Lora et al. (2006). The average value found for the goat milk acidity was 0.17 g lactic acid/100 mL of product. Such an average is within the standards required by the legislation (Brazil, 2000a).

The average fat value obtained in the raw milk by the Gerber method was above the minimum (3%) required by the legislation for whole milk, density of the goat milk was within the norms of the current legislation (Brazil, 2000a).

**Viable lactic acid bacteria count**

Table 1 presents the average values (CFU/mL) of the traditional lactic bacteria (viable cells of *S. thermophilus* and *L. delbrueckii* ssp. *bulgaricus*) counts of the strawberry flavor goat milk based yoghurts with added WSSE and of the probiotic culture during the storage period of 29 days at 4°C.

The viable lactic cell count remained 2.6 x 10⁷ to 1.8 x 10⁷ for the control yoghurt. The maintenance of the number of viable cells during the 29 day storage period under cooling at 4°C met the values established by the
Table 1. Average count of the number of viable lactic acid bacteria from goat milk yoghurt samples with added WSSE and probiotics during storage time (CFU/ml).

<table>
<thead>
<tr>
<th>Yoghurt</th>
<th>Day 1</th>
<th>Day 8</th>
<th>Day 15</th>
<th>Day 22</th>
<th>Day 29</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>2.6 x 10^6 a</td>
<td>2.3 x 10^6 a</td>
<td>1.8 x 10^6 a</td>
<td>2.5 x 10^6 a</td>
<td>1.8 x 10^6 a</td>
</tr>
<tr>
<td>B</td>
<td>3.2 x 10^6 b</td>
<td>2.2 x 10^6 b</td>
<td>5.0 x 10^6 b</td>
<td>4.1 x 10^6 b</td>
<td>3.0 x 10^6 b</td>
</tr>
<tr>
<td>C</td>
<td>9.0 x 10^6 c</td>
<td>4.8 x 10^6 c</td>
<td>1.15 x 10^7 a</td>
<td>3.3 x 10^6 a</td>
<td>3.1 x 10^7 a</td>
</tr>
<tr>
<td>D</td>
<td>7.3 x 10^6 c</td>
<td>5.0 x 10^6 c</td>
<td>4.1 x 10^6 b</td>
<td>8.3 x 10^6 c</td>
<td>2.5 x 10^6 b</td>
</tr>
</tbody>
</table>

Means followed by the same letter do not differ statistically among themselves, by Scott-Knott Test Average, at 5% probability. Treatment: A (without addition of WSSE and without addition of probiotic culture), B (with addition of WSSE and without addition of probiotic culture), C (without addition of WSSE and with addition of probiotic culture), D (with addition of WSSE and with added probiotic culture).
Figure 2. Regression model for the viable cell count (log CFU / mL) versus storage time at 4°C, in yoghurts. Treatment: A (without addition of WSSE and without addition of probiotic culture), B (with addition of WSSE and without addition of probiotic culture), C (without addition of WSSE and with addition of probiotic culture), D (with addition of WSSE and with added probiotic culture).

Table 2. Average viable cell count of *B. animalis* ssp. *lactis* in goat milk yoghurt samples with added WSSE and probiotic culture during the storage time (CFU/ml).

<table>
<thead>
<tr>
<th>Yoghurt</th>
<th>CFU (ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Day 1</td>
</tr>
<tr>
<td>C</td>
<td>1.6 x 10^8</td>
</tr>
<tr>
<td>D</td>
<td>1.1 x 10^8</td>
</tr>
</tbody>
</table>

Means followed by same letter do not differ statistically among themselves, by Scott-Knott Test Average, to 5% probability. Treatment: C (without addition of WSSE and with addition of probiotic culture), D (with addition of WSSE and with added probiotic culture).

centration (osmotic pressure), dissolved oxygen (especially for the *Bifidobacterium* ssp.), amount inoculated, incubation temperature and storage temperature duration (Lourens-Hattingh and Viljoen, 2001; Silva, 2007).

The average enumeration of viable *B. animalis* ssp. *lactis* cell colonies in the yoghurt treatments can be seen in Table 2. The average viable cell count of the *B. lactis* probiotic microorganism remained between 1.6 x 10^5 and 3.1 x 10^7 CFU/mL for the yoghurt C, without WSSE addition, and the average count was 1.1 x 10^8 to 2.7 x 10^7 for the yoghurt D, with addition of water-soluble soy extract. From the ANOVA results, it can be observed that the average bifidobacteria count was not significantly influenced by the addition of soy extract. It is worth noting that the average viable cell count of the *B. animalis* ssp. *lactis* probiotic microorganism in the treatment without soy extract addition was slightly higher, however not significant, during the storage time of the yoghurts in comparison with the treatment with addition of this product. These results were contrary to the expected, because, according to Tamine et al. (1995), the soy contains prebiotics, such as rafinose and stachyose that are natural bifidobacteria growth promoters. It was
observed that the time influenced the bifidobacteria count in a significant way, according to the graph in Figure 3. The quality of a probiotic product is usually determined by the level, viability and amount of the probiotic cells in the food. This has been proposed as guarantee of the beneficial effects to human health (Schessler, 2009). Silva (2007) verified viability when evaluating species of *Bifidobacterium* and *L. acidophilus* probiotic bacteria in yoghurt development with the probiotic inulin.

According to Technical Regulation of Isolated Bioactive and Probiotic Substances with Allegation of Functional and/or Health Properties, Resolution RDC No.2, January 2002, probiotic is understood as live microorganisms capable of improving the intestinal microbial balance producing beneficial effects to the health of the individual (Brazil, 2002). Some benefits related to probiotic use are: stabilization of the intestinal function, reduction of infection by *Helicobacter pylori*, that is associated to gastritis and peptic ulcers and lactose intolerance symptoms relief. Other advantages in the use of these microorganisms are: the conservation of the milk through the production of lactic acid and possibly antimicrobial compounds; the production of flavor and aroma compounds and other metabolites that will provide a product with sensorial properties desirable to the consumer; the improvement of the nutritional value of the food, for instance, through the liberation of free amino acids or vitamin synthesis; the provision of therapeutic properties and special prophylaxis for cancer prevention and control of serum cholesterol levels. The potential benefits are resultant from the bacterial development and action during the fermented food production (Parvez et al., 2006).

**Conclusions**

This study showed that the viable cell count in the yoghurts was satisfactory without addition of water-soluble soy extract and also for those with added *B. animalis* ssp. *lactis* probiotic culture during the period appraised, therefore, the yoghurts were considered in accordance with the current law. However, the treatment with only added water-soluble soy extract was not shown viable according to the legislation; therefore it cannot be considered a yoghurt in that respect. The addition of the *B. animalis* ssp. *lactis* probiotic culture in the yoghurts with goat milk was shown viable during the storage time.

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