Evaluation of the Effect of Starter Culture on the Quality of White Soft Cheese (Gibna Beyda)

Article · October 2016

3 authors, including:

Hassan Ali Mudawi
University of Khartoum

36 PUBLICATIONS 10 CITATIONS

Some of the authors of this publication are also working on these related projects:

- Design modification of wheat roller mill to grind sorghum View project
- The effect of malting conditions and mashing methods on the production of non-alcohol sorghum bee View project

All content following this page was uploaded by Hassan Ali Mudawi on 18 April 2018.

The user has requested enhancement of the downloaded file.
Evaluation of the Effect of Starter Culture on the Quality of White Soft Cheese (Gibna Beyda)

Mudawi, H.A.a, Khairalla, L.M.b and El Tinay, A.H.c

a. University of Khartoum, Dep. of Food Science & Technology.
b. Food Processing Research Centre, Khartoum-North.
c. University of Khartoum, Dep. of Food Science & Technology.

Abstract
This investigation was carried out to study the effect of starter culture on the quality of white soft cheese (Gibna Beyda), yield and coagulation time. White soft cheese was made from fresh cows milk (heated at 65°C for 30 minutes, then cooled to 35°C), then starter culture was added at different concentrations (1.5, 2.0, 2.5%) in addition to CaCl₂ (0.02%), NaCl (2%) and rennet. Chemical analysis, microbial analysis and sensory evaluation were carried out. The coagulation time for cheese made without addition of starter culture (control) was 189 min. and the yield was 9.61%. The obtained results indicated that the use of starter culture at concentration of 2.5% showed reduction in coagulation time to 64 min. and increase in yield to 13.51%. The effect of different concentrations of starter culture on the chemical composition of white soft cheese revealed that the moisture content showed a significant increase from 56.15% to 69.17%, acidity increased from 0.22% to 0.32%, fat content decreased from 27.60% to 21.00%, protein content decreased from 15.00% to 9.40%, total solids decreased from 43.77% to 31.80%. The total bacterial count increased from 2.3 × 10⁴ to 4.1 × 10⁶, the yeast count increased from 8.5 × 10³ to 1.7 × 10⁵. The flavor and overall acceptability were affected by the addition of starter culture. The best flavor and best overall acceptability scores were obtained in cheese made with 1.5% starter culture, while the least score to flavor was obtained in cheese made with 2.5% starter culture.
Introduction

Milk is a complex biological fluid, secreted by the mammary glands in lactating mammals. The major constituents of milk include water, lactose, fats, protein, minerals, vitamins and enzymes. The unique composition and properties of milk make it an ideal food for the mammalian neonate (Tannahil, 2008). When the supply of fresh milk is in surplus, the producers try to make the best use of it to avoid falling down of milk prices. So milk can be separated into cream or processed into cheese, butter or yoghurt. Approximately one third of the world milk production is used in cheese manufacture (Tannahil, 2008). The manufacture of cheese is a form of milk preservation as milk is highly perishable (El Owni and Hamid, 2007). Cheese is highly nutritious food, with many diverse flavours and textures (Winstein, 2017 and Suliman et al., 2013).

Cheese can be used as a snack or as a part of dish or prepackaged convenience food, supplies abundant quantities of proteins, fat and calcium, which are essential for good health and growth (Quinee, 2004). The individual characteristics of each type of cheese are due to the type of milk, the microbial starter culture and the manufacturing procedure used (El Khider et al, 2012).

Traditionally, the white cheese is processed in the Sudan by natural fermentation induced by microorganisms either present in the raw milk or from the surrounding environments. Starter cultures are not usually used in the processing of Sudanese white cheese. (Ahmed, 1997). Various starter culture types are used in the dairy industry for cheese making. The lactic acid bacteria are used to induce lactic fermentation which is very essential in the manufacture of cheese and fermented milk products (Kats and Pollan, 2012). The use of starter culture in the production of cheeses encourages whey separation, inhibits the growth of pathogenic bacteria, generates some aroma compounds and increases the degree of ripening, though the use of starter culture is not common (Nour El Daim and El Zubeir, 2010 and El Zubeir et al, 2014).

The objectives of the study are to study the effect of different concentrations of starter culture on the chemical composition and microbial quality of white cheese. It is also meant to determine the optimum starter concentration that produces the best quality cheese.

Raw materials

The raw materials used in this study include: Fresh cow milk brought from University farm, thermophilic lactic starter culture Streptococcus thermophiles brought from biotechnology laboratory, Faculty of Agricultures, media, namely potato dextrose agar (PDA) and plate count agar, brought from Food Research Centre, rennet, commercial salt (NaCl) and calcium Chloride, bought from Samabel Chemical Company.

Cheese manufacture

Twenty liters of milk were heated to 65°C for 30 minutes, cooled to 35°C and then salted by two types of salts (2% w/w sodium chloride and 0.02% w/w calcium chloride). The milk was then divided into four batches of 5 liter each. The first batch was kept without starter culture (as control), the other three batches were treated with starter culture at the concentration of 1.5%, 2%, 2.5%, respectively. Rennet tablets were dissolved in 20-ml distilled water and the formed solution was divided into four batches of 5 ml. each. The mixture was hand stirred for five minutes using a spoon. The batches were then incubated at 40°C and left to develop a curd. After coagulation the curd was cut with an ordinary stainless steel knife to allow for whey separation. The curd was poured into small wooden molds lined with cheese cloth, pressed and left over night.

In the following day, brine solution was prepared by adding salt (6% w/w) to distilled water, heated to 65°C for 30 min and then cooled to 35°C. The fresh curd was immersed into brine solution for 48 hours. Cheese was then transferred to plastic containers and stored at room temperature, for chemical and
microbial analysis, as also previously done by El Owni and Hamid (2007).

**Chemical analysis of milk and cheese**

Chemical analyses were carried out for raw milk and the cheese. Titratable acidity was determined according to AOAC (1990). Fat content was determined by Gerber Method according to AOAC (1990). Total solids content was determined according to the modified method of AOAC (1990). The protein content was determined by Kjeldahl method as described by AOAC (1990). The moisture content of cheese was determined by the method described by Nour El Diam and El Zubeir (2010).

Before carrying out the microbial examination the glassware were sterilized in a hot dry oven at 160°C for 3 hours. Culture media were prepared: plate count agar to determine the total bacterial count and potato dextrose agar (PDA) for the enumeration of yeasts and molds. Then the serial dilution was prepared. The total viable count was carried out using the pour plate method as described by Kats and Pollan (2012). The mold and yeast enumeration was carried out according to Kats and Pollan (2012).

**Sensory evaluation**

Ten untrained panelists have tested the produced cheese to evaluate taste, color, flavor, body and texture, and overall acceptability.

**Statistical analysis**

The data were analyzed using (ANOVA) in completely randomized design. Mean separation was carried out using Duncan multiple range test according to statistical analysis system suggested by Bylund, (2005).

**Results and discussion**

**Coagulation time**

As shown in Table 1 the coagulation time for the applied procedure varied between 64 and 75 minutes with an average of 70 minutes, while coagulation time in traditional method varied between 120 and 180 min. depending on the quantity of added salt prior to renneting (Robinson, 2009).

The addition of starter culture in milk in this study and reduction of the quantity of added salt to 2% (w/w) has resulted in shortening of the coagulation time as compared to the traditional procedure, Asher (2015) obtained similar result.

**Yield**

Table 1 presents the yield of white soft cheese made using different concentrations of starter culture. The highest yield (13.5%) was obtained in cheese made from the milk treated with 2.5% starter culture while the lowest yield (9.61%) was obtained from the cheese made without starter culture addition. The effect of addition of starter culture in increasing the yield is clear in these results and the results of other authors (Nour El Diam and El Zubeir, 2010; El Owni and Hamid 2007 and Abdalla et al., 2012).

**The effect of starter culture on chemical composition of cheese**

**Moisture content**

The data in Table 2 presents the effect of starter culture (0.0, 1.5, 2.0 and 2.5%) on the moisture content of white soft cheese. The effect of addition of starter culture was found to be highly significant (P<0.01). The highest moisture content (69.17%) was obtained in cheese treated with 2.5% (w/w) starter culture and the lowest moisture content (56.15%) was obtained from cheese made without addition of starter culture. The moisture content of the cheese was relatively similar to the values 61.2, 54.2 and 61.6% which were obtained by Warsama et al. (2006).

**Titratable acidity**

The data in Table 2 presents the effect of the different rates of starter culture (0.0, 1.5, 2.0 and 2.5%) on the acidity of white soft cheese. The acidity of cheese was highly significantly (P<0.01) affected by the addition of starter cultures. The highest acidity (0.32%) was
obtained in cheese made with addition of 2.5% starter culture, while the lowest acidity (0.22%) was obtained in cheese made without starter culture (zero starter culture) Bylund (2005) and Asher (2015) and Abdalla et al. (2013) obtained similar results. Increasing the rate of starter culture increased the acidity. Fox (2003) reported that the titratable acidity was in the range of 0.162 to 1.895% when they manufactured Domiati cheese using calf rennet. These values were relatively similar to that obtained in this study.

**Fat content**

The data in Table 2 shows the effect of the different rates of starter culture (0.0, 1.5, 2.0 and 2.5%) on the fat content of white soft cheese. The fat content was highly significantly (P<0.01) affected by the addition of starter culture. The highest fat content (27.6%) was obtained in the cheese made without adding starter culture (zero starter culture) while the lowest fat content (21.0%) was found in cheese made with addition of 2.5% starter culture. The addition of starter culture decreased the fat content. Asher (2015) found that the average fat content of commercial Gibna-Beyda was 14%, while Idris and Alhassan (2010) reported a fat content of 12.65%. These values were lower than those obtained in this study. Obviously the fat content of the cheese is related to the fat content of the original milk as well as to the moisture content of the cheese. The variation in the fat content in this study was due to the addition of starter culture, which increased the moisture content, thus decreased the total solids and fat content, Nour El Diam and El Zubeir (2010) obtained a similar result.

**Total solids content**

Table 2 showed that the total solids content was highly significantly (P<0.01) affected by the addition of starter culture. The highest total solids content (43.77%) was found in cheese made without addition of starter culture and the lowest (31.8%) was found in cheese treated with 2.5% starter culture. The average of the total solids content was 36.75%. These values were significantly different from those obtained by Ibrahim (1999) who reported a range of 39.4 – 45.5%, Bylund who reported a range of 40.3 – 46.7% and Asher who reported a range of 38.8 – 44.9%. These values are higher compared to that obtained in this study, and the difference was claimed to be due to the increase of moisture content which affect the curd concentration. Winstein (2017) obtained values similar to that obtained in this study.

**Total protein content**

The data in Table 2 presents the effect of starter culture (0.0, 1.5, 2.0 and 2.5%) on the total protein content of white soft cheese. The total protein content of cheese was highly significantly (P<0.01) affected by the addition of starter culture. The highest total protein content (15.0%) was obtained in the cheese made without addition of start culture, while the lowest protein content (9.41%) was obtained in the cheese made with 2.5% starter culture, average value was 12.36%. Ador kour (1992) found that the average total protein of fresh white soft cheese was 13.77%, Fox (2003), Tannahil (2008) and Winstein (2017) reported total protein content in the range of 10.4 – 16.6%, these values were relatively similar to those obtained in this study. Abdel Razig (1996) reported that the total protein of fresh white cheese was 17.79%, and Bylund (2005) reported a value of 16.9% for total protein of fresh white cheese, these values are higher than the value obtained in this study.

**Microbial quality**

**Total bacterial count**

The results in Table 3 present the effect of different rates of starter culture (0.0, 1.5, 2.0 and 2.5%) on the total bacterial count of white soft cheese. The total bacterial count was found to be highly significantly (P<0.01) affected by the addition of starter culture. The highest bacterial count (4.1 106) was obtained in cheese used 2.5% starter culture and the lowest bacterial count was obtained in cheese made without adding starter culture. The results obtained in this study disagreed with that of El Owni and Hamid (2009) who found that the viable bacterial count of the cheese made in the laboratory (3.5×108) was higher than the viable bacterial count of the fresh market cheese (9.5×105). The obtained results agreed with that of Doyle (2007) who used
Yeast and mould count

The results in Table 3 shows the total yeast and mould count. The highest yeast count was obtained in the cheese made with addition of 2% starter culture and the lowest was found in cheese made without adding starter culture. The results in Table 3 revealed the absence of mould (no mould growth). The results obtained in this study were relatively similar to those obtained by El Owni and Hamid (2009) who found that the average yeast count in the fresh cheese varied between 3 104 /g and 8.9 106/g and the mould count varied between 1 102/g and 3 102/g.

Organoleptic properties of white soft cheese

The results in Table 4 present the organoleptic properties of white soft cheese made by addition of different rates of starter culture (0.0, 1.5, 2.0 and 2.5%). Generally all cheese samples to which starter culture was added at any rate scored well. Cheese samples made with addition of 1.5% starter culture had the best score, followed by the cheese made using 2% starter culture, compared to control cheese and cheese made by 2.5% starter culture. According to the panelists who judged the quality of cheese made in this study, the cheese is of high quality, it has good color and smooth but firm body and texture with better consistency and good flavor. However, the obtained results disagreed with those obtained by El Zubeir et al. (2014) who found that the cheese developed acceptable flavor and acid taste after two weeks of storage, that was attributed to low bacterial count of heat treated milk which resulted in slower acidity and flavor development. The results in this study agreed with the results obtained by Alcamo (2003) and Doyle (2007). Fresh cheese from heated milk was superior to that made from raw milk with regard to body and texture, while during ripening raw milk cheese was superior to that made from heated milk. Asher (2015) reached the same.

The results obtained in this study are in agreement with those obtained by Fox (2003) who reported that good quality fresh cheese may be obtained from pasteurized whole milk, 1% Streptococcus lactis and 0.75% Enterococcus as starter culture.

Table 1: The coagulation time and yield of cheese samples made by different concentrations of starter cultures

<table>
<thead>
<tr>
<th>Sample</th>
<th>Coagulation time (min)</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A - Control</td>
<td>189</td>
<td>9.61</td>
</tr>
<tr>
<td>A1</td>
<td>76</td>
<td>11.82</td>
</tr>
<tr>
<td>A2</td>
<td>70</td>
<td>12.33</td>
</tr>
<tr>
<td>A3</td>
<td>64</td>
<td>13.51</td>
</tr>
</tbody>
</table>

A: Cheese made without addition of starter culture (control).
A1: Cheese made with addition of 1.5% starter culture.
A2: Cheese made with addition of 2.0% starter culture.
A3: Cheese made with addition of 2.5% starter culture.

Table 2: The effect of different concentrations of starter culture on the chemical composition of white soft cheese

<table>
<thead>
<tr>
<th>Component (%)</th>
<th>0 %</th>
<th>1.5 %</th>
<th>2.0 %</th>
<th>2.5 %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture</td>
<td>56.15d</td>
<td>60.11c</td>
<td>61.11b</td>
<td>69.17a</td>
</tr>
<tr>
<td>Fat</td>
<td>27.60a</td>
<td>26.30b</td>
<td>25.00c</td>
<td>21.00d</td>
</tr>
<tr>
<td>Acidity</td>
<td>0.22d</td>
<td>0.25c</td>
<td>0.29b</td>
<td>0.32a</td>
</tr>
<tr>
<td>Protein</td>
<td>15.00a</td>
<td>14.00ab</td>
<td>13.70b</td>
<td>8.40d</td>
</tr>
<tr>
<td>Total solids</td>
<td>43.77a</td>
<td>40.11b</td>
<td>39.17c</td>
<td>31.80d</td>
</tr>
</tbody>
</table>

Means within the raw bearing the same superscript letters are not significantly different (P>0.05).
Table 3: The effect of different concentrations of starter culture on the total microbial count of white soft cheese

<table>
<thead>
<tr>
<th>The concentration of starter culture</th>
<th>Microbial count</th>
<th>0.0 %</th>
<th>1.5 %</th>
<th>2.0 %</th>
<th>2.5 %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteria</td>
<td>2.3*10⁴b</td>
<td>4.1*10⁵b</td>
<td>4.4*10⁵b</td>
<td>4.1*10⁶a</td>
<td></td>
</tr>
<tr>
<td>Yeast</td>
<td>8.5*10⁴b</td>
<td>6.0*10⁵a</td>
<td>1.7*10⁵b</td>
<td>1.2*10⁵b</td>
<td></td>
</tr>
<tr>
<td>Mould</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

Means within the same raw bearing the same superscript letters are not significantly different (P≤0.05).

Table 4: The effect of different concentrations of starter culture on organoleptic quality of white soft cheese

<table>
<thead>
<tr>
<th>The concentration of starter culture</th>
<th>Parameters</th>
<th>0.0 %</th>
<th>1.5 %</th>
<th>2.0 %</th>
<th>2.5 %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color</td>
<td>8.37a</td>
<td>8.70a</td>
<td>8.07a</td>
<td>8.05a</td>
<td></td>
</tr>
<tr>
<td>Flavor</td>
<td>5.87b</td>
<td>8.07a</td>
<td>7.57a</td>
<td>6.20b</td>
<td></td>
</tr>
<tr>
<td>Texture</td>
<td>6.57c</td>
<td>7.97a</td>
<td>7.30a</td>
<td>5.87c</td>
<td></td>
</tr>
<tr>
<td>Acceptability</td>
<td>6.84c</td>
<td>8.18a</td>
<td>7.71b</td>
<td>6.74c</td>
<td></td>
</tr>
</tbody>
</table>

Means within the same raw bearing the same superscript letters are not significantly different (P>0.05).

Conclusion

It was found that the use of starter culture in the processing of white soft cheese caused reduction in coagulation time and increased the yield with increasing rate of starter culture. The chemical composition of white soft cheese (fat, protein, total solids) increased with decreasing the rate of starter culture except for the acidity and the moisture content. Also, it was found that the Sudanese cheese showed the highest quality and the least microbial count at the lower rate of starter culture addition. Hence

It is recommended to use a low concentration of starter culture to produce white soft cheese of good constitution and high acceptability. Furthermore, future studies on the subject area should study the effect of different types of starter cultures on chemical composition and quality of white soft cheese.

References


Kats, Sandor Ellix; Pollan, Michael (2012). The Art of Fermentation an In-depth Exploration of Essential Concepts and Processes from around the World. Vermont: Chelsea Green Publishing


