Influence of Starters on Chemical, Biochemical, and Sensory Changes in Turkish White-Brined Cheese During Ripening

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ABSTRACT

Turkish White-brined cheese was manufactured using Lactococcus strains (Lactococcus lactis ssp. lactis NCDO763 plus L. lactis ssp. cremoris SK11 and L. lactis ssp. lactis UC317 plus L. lactis ssp. cremoris HP) or without a starter culture, and ripened for 90 d. It was found that the use of starters significantly influenced the physical, chemical, biochemical, and sensory properties of the cheeses. Chemical composition, pH, and sensory properties of cheeses made with starter were not affected by the different starter bacteria. The levels of soluble nitrogen fractions and urea-PAGE of the pH 4.6-insoluble fractions were found to be significantly different at various stages of ripening. Urea-PAGE patterns of the pH 4.6-insoluble fractions of the cheeses showed that considerable degradation of αs1-casein occurred and that β-casein was more resistant to hydrolysis. The use of a starter culture significantly influenced the levels of 12% trichloroacetic acid-soluble nitrogen, 5% phosphotungstic acid-soluble nitrogen, free amino acids, total free fatty acids, and the peptide profiles (reverse phase-HPLC) of 70% (vol/vol) ethanol-soluble and insoluble fractions of the pH 4.6-soluble fraction of the cheeses. The levels of peptides in the cheeses increased during the ripening period. Principal component and hierarchical cluster analyses of electrophoretic and chromatographic results indicated that the cheeses were significantly different in terms of their peptide profiles and they were grouped based on the use and type of starter and stage of ripening. Levels of free amino acid in the cheeses differed; Leu, Glu, Phe, Lys, and Val were the most abundant amino acids. Nitrogen fractions, total free amino acids, total free fatty acids, and the levels of peptides resolved by reverse phase-HPLC increased during ripening. No significant differences were found between the sensory properties of cheeses made using a starter, but the cheese made without starter received lower scores than the cheeses made using a starter. It was found that the cheese made with strains NCDO763 plus SK11 had the best quality during ripening. It was concluded that the use of different starter bacteria caused significant differences in the quality of the cheese, and that each starter culture contributed to proteolysis to a different degree.

(Key words: White cheese, starter, Lactococcus, ripening)

Abbreviation key: EtOH-i = ethanol-insoluble, EtOH-s = ethanol-soluble, FAA = free amino acids, HCA = hierarchical cluster analysis, PCA = principal component analysis, PTA-SN = 5% phosphotungstic acid-soluble nitrogen, RP-HPLC = reverse phase-HPLC, TCA-SN = 12% TCA-soluble nitrogen, WSN = water-soluble nitrogen.

INTRODUCTION

Cheese ripening is a complex and dynamic biochemical process that includes protein breakdown, fat hydrolysis, and lactose metabolism (El Soda et al., 1995; McSweeney and Sousa, 2000). Proteolysis is catalyzed by proteolytic enzymes from the coagulant, milk, and bacteria (starter, nonstarter, or secondary starter; Fox, 1989). The coagulant is mainly responsible for hydrolyzing the caseins to large and intermediate-size peptides (Law et al., 1992; Lane and Fox, 1997). The enzymes originating from starter (i.e., proteinases, peptidases) play a major role in formation of small peptides and the amino acids, which serve as precursors of flavor compounds in cheese (Urbach, 1997; Broome and Lim sowtin, 1998).

The primary role of starter bacteria is to produce lactic acid at a controlled rate; in addition, the bacteria affect the nonstarter microflora. Starter cultures are not used for many cheeses made in Turkey including Turkish White-brined cheese. In artisanal cheese production, the cheese is made without the deliberate addition of a starter culture; the indigenous flora of the milk contribute to ripening. Recently, the use of mixed-strain, mesophilic starter cultures, containing un-
known genera of lactic acid bacteria commenced in the manufacture of Turkish White-brined cheese. The quality of cheese manufactured with mixed starter cultures varies widely in terms of texture and flavor.

Cheeses made using a starter containing one or more defined Lactococcus strains usually have a uniform texture due to the known biochemical activities of their microflora during cheese manufacture and ripening (Powell et al., 2002). The organisms in mixed-strain starters used in the manufacture of the cheeses belong mainly to the genera Lactococcus or Lactobacillus (Hayaloglu et al., 2002). Lactococcus species, of which L. lactis ssp. lactis or L. lactis ssp. cremoris are the best known, are used in cheese manufacture – these species were also found to be the predominant flora in Turkish White-brined cheese by Karakus et al. (1992).

The objective of this study was to evaluate the influence of defined starter cultures on the gross composition, proteolytic profile, and other ripening characteristics of Turkish White-brined cheese during ripening. In addition, we attempted to standardize the manufacture and quality of the cheese using a defined-strain starter.

**MATERIALS AND METHODS**

**Lactococcal Strains**

*Lactococcus lactis* ssp. lactis UC317, *L. lactis* ssp. lactis NCDO763, *L. lactis* ssp. cremoris HP, and *L. lactis* ssp. cremoris SK11 were obtained from the culture collections of University College, Cork, Ireland; Fonterra Research Centre, Palmerston North, New Zealand; and Groningen Biomolecular Science and Biotechnology Institute, Department of Genetics, University of Groningen, Haren, The Netherlands, respectively. Before use, each strain was cultivated in M17 broth (Merck, Darmstadt, Germany) at 30°C for 24 h with 2 consecutive transfers (1%, vol/vol, inoculum). For starter propagation, the cultures were grown in reconstituted skim milk (10%, wt/vol, heated at 90°C for 30 min) as described by Fenelon et al. (2000).

**Cheese Making**

Turkish White-brined cheese was made in triplicate. In each trial, pasteurized (68°C for 10 min) milk was cooled to 32°C, divided into 3 equal parts and inoculated with a lactococcal culture at a level of 1% (wt/vol) as follows: 1) *L. lactis* ssp. lactis UC317 plus *L. lactis* ssp. cremoris HP (317HP); 2) *L. lactis* ssp. lactis NCDO763 plus *L. lactis* ssp. cremoris SK11 (763SK11); and 3) Starter-free (SF) cheese. Calcium chloride was added to the milk at a level of 0.2 g/L. During all stages of cheese making, precautions were taken to avoid cross-contamination. The inoculated milk (32°C) was held for about 30 min (until pH 6.30), and liquid calf rennet (Chr. Hansen, Copenhagen, Denmark) was added at a level of 1 g per 10 L of cheese milk (sufficient to coagulate the milk in 90 min). Following coagulation, the coagulum was cut into cubes (2 to 3 cm sides) and allowed to rest for 10 min. The curds were carefully transferred from the cheese vat into the molds. After 1 h of draining (without pressing), pressure was applied at room temperature (21°C) for 3 h or until whey drainage had stopped. Then, the weights were removed and the block of cheese cut into cubes of about 7 × 7 × 7 cm with a knife; the pieces, weighing 350 to 400 g each, were placed in brine (14% NaCl) for about 12 h at 21°C. After salting, the cheese blocks were packed in cans (16 × 8 × 8.6 cm) and covered with 14% NaCl brine. The cans, which contained about 1 kg of cheese, were closed hermetically and the cheese samples ripened at 6 to 8°C for 90 d.

**Analysis of Cheese Samples**

**Gross Composition.** Cheeses were analyzed at 1 d for moisture by the oven drying method at 102°C (IDF, 1982), salt by titration with AgNO₃ (Bradley et al., 1993), fat by the Van Gulik method (Ardo and Polychroniadou, 1999), and total protein by the Kjeldahl method (IDF, 1993). The pH of cheeses was measured in a slurry prepared by macerating 10 g of grated cheese in 10 mL of deionized water. Titratable acidity was determined as grams per 100 grams of lactic acid using the method described in AOAC (1995).

**Nitrogen Fractions.** Water-soluble nitrogen (WSN), 12% TCA-soluble nitrogen (TCA-SN), and 5% phosphotungstic acid-soluble nitrogen (PTA-SN) fractions were prepared by the methods of Kuchroo and Fox (1982), Polychroniadou et al. (1999), and Jarrett et al. (1982), respectively. The nitrogen content of the fractions was determined by the Kjeldahl method (IDF, 1993) and expressed as a percentage of total cheese nitrogen. All determinations were made in duplicate.

**Total Free Amino Acid Content.** The total concentration of free amino acids (FAA) in the cheeses was determined in triplicate by the method of Folkertsma and Fox (1992). Results were expressed as milligrams of Leu/gram of cheese. The standard Leu solutions were analyzed in triplicate and a standard curve was prepared.

**Total FFA.** The total concentrations of FFA in the cheeses was determined by titrating the acidity in the cheese fat with 0.05 N ethanolic KOH using the method of described by Nunez et al. (1986); results were expressed as the percentage of oleic acid in cheese fat (IDF, 1989).
Sample preparation for urea-PAGE and reverse phase-HPLC. After ripening for 1, 15, 30, 60, or 90 d, one cheese can from each vat was randomly selected and opened. The cheese block was divided into 4 equal parts with a knife, and 2 diagonal parts, about 200 g each, were grated to provide a representative sample. The pH 4.6-soluble and insoluble fractions were prepared using a slight modification of the method of Kuchroo and Fox (1982). A grated sample (20 g) was homogenized with 40 mL of distilled water for 5 min using a Colworth Stomacher 400 (Seward Laboratory, London, UK). The homogenate was adjusted to pH 4.6 using 1.0 M HCl, left to stand at room temperature for 30 min and the pH readjusted to pH 4.6, if necessary. The resultant homogenate was held at 40°C for 1 h, and then the pH 4.6-insoluble material was separated by centrifugation at 3000 × g for 30 min in a refrigerated (4°C) centrifuge (model RC5C, Sorvall, Wilmington, DE). The supernatant (pH 4.6-soluble fraction) was filtered through glass wool and Whatman No. 113 filter paper and the ethanol precipitated through Whatman No. 1 filter paper and frozen pending peptide analysis by reverse phase-HPLC (RP-HPLC) and further fractionation. The pellet (pH 4.6-insoluble fraction) was freeze-dried for analysis by urea-PAGE.

Absolute ethanol (46.67 mL) was added to 20-mL aliquots of the pH 4.6-soluble fraction to a final ethanol concentration of 70% (vol/vol). The resultant suspension was held for 30 min at room temperature (21°C) and then centrifuged at 3000 × g for 30 min at 20°C. The supernatant (containing the smaller, hydrophilic peptides that were soluble in ethanol) was filtered through Whatman No. 1 filter paper and the ethanol removed using a rotary evaporator (Bibby Sterilin Ltd., Stone, UK) at 30°C under vacuum. The pellet (containing the larger, more hydrophobic peptides that were insoluble in ethanol) was dispersed in distilled water and freeze-dried before analysis by RP-HPLC and urea-PAGE.

Urea-PAGE and densitometry. Urea-PAGE [4% C (cross-linking agent as percentage of total monomer), 12.5% T (total monomer), pH 8.9] of the pH 4.6-insoluble fraction of the cheeses was performed using a Protean II XI vertical slab-gel unit (Bio-Rad Laboratories Ltd., Watford, UK) according to the method of Andrews (1983), as modified by Shalabi and Fox (1987). Freeze-dried samples (10 mg) of the pH 4.6-insoluble fraction from each sample were dissolved in 1 mL of electrophoresis sample buffer and vortexed (and/or sonicated, if necessary) for 1 min and stored at −20°C until analyzed. Before electrophoresis, frozen samples were heated at 50°C for 5 min and then cooled to room temperature. A 7-μL aliquot of each sample was loaded onto the gel. Electrophoresis was performed through the stacking and separating gels at 280 and 300 V, respectively. The gels were stained directly by the method of Blakesley and Boezi (1977) with Coomassie Brilliant Blue G-250 and destained using distilled water. After destaining, gel slabs were digitized using a scanner (HP ScanJet software, ScanJet 6300C, Hewlett Packard, Palo Alto, CA). Scans of the electrophoretograms were used to quantify bands using densitometric software (Image Master TotalLab 1D Gel analysis v1.11 software, Amersham Pharmacia Biotech, Uppsala, Sweden). The casings and peptides were determined quantitatively by integration of peak volumes using the densitometer.

RP-HPLC. Peptide profiles of 70% ethanol-soluble and insoluble fractions of the pH 4.6-insoluble fraction of samples of Turkish White-brined cheese were determined by RP-HPLC using a Varian HPLC system (Varian Associates Inc., Walnut Creek, CA). The system comprised an autosampler (model 410), a ProStar solvent delivery system with 3 pumps (model 230), a ProStar programmable multiwavelength spectrophotometer (model 310) interfaced with a PC, onto which a Varian Star Workstation 5 software package was installed for system control and data acquisition. Nucleosil RP-8 (250 × 4 mm, 5 μm particle size, 300 Å pore size) analytical and guard columns (4.6 × 10 mm) (Catalal HPLC Ltd., Broxburn, UK) were used. The solvents were: (A) 0.1% (vol/vol) trifluoroacetic acid (sequencing grade; Sigma, St Louis, MO) in HPLC-grade deionized water (Milli-Q system, Waters Corp., Molsheim, France), and (B) 0.1% (vol/vol) trifluoroacetic acid in acetonitrile (HPLC grade, Lab-Scan Ltd., Dublin, Ireland) at a flow rate of 0.75 mL/min. Samples of freeze-dried 70% ethanol-soluble and insoluble subfractions of the pH 4.6-soluble fractions were dissolved in solvent A (10 mg/mL), filtered through a 0.45-μm cellulose acetate filter (Sartorius GmbH, Gottingen, Germany); a 40-μL aliquot of filtrate was injected onto the column. The samples were eluted initially with 100% solvent A for 5 min, then with a gradient from 0 to 50% solvent B (vol/vol) over 55 min, maintained at 50% solvent B (vol/vol) for 6 min, followed by a linear gradient from 50 to 60% solvent B (vol/vol) over 4 min, and finally with 60% solvent B (vol/vol) for 3 min. The column was washed with 95% solvent B (vol/vol) for 5 min, followed by equilibration with 100% solvent A for 5 min before the next injection. Elute was monitored at 214 nm.

Determination of individual FAA. The pH 4.6-soluble fractions were deproteinized by mixing with an equal volume of 24% (wt/vol) TCA; the mixture was allowed to stand 10 min and then centrifuged at 14,000 × g (Microcentaur, MSE, Fisher Scientific, Loughborough, UK) for 10 min. The supernatant was removed and diluted with 1.2 M sodium citrate buffer (pH 2.2) to give approximately 250 nmol of each amino acid. The samples were diluted 1:2 with norleucine, the internal

standard, to give approximately 125 nmol of each amino acid residue/mL of injection solution. Samples were then analyzed on a Beckman 6300 High Performance Amino Acid Analyser (Beckman Instruments Ltd., High Wycombe, UK) fitted with a 120 × 4 mm cation-exchange column (Na+ form). The individual amino acids were separated by ion-exchange chromatography with postcolumn derivatization with ninhydrin and visible colorimetric detection at 570 nm with the exception of proline, which was detected at 400 nm. Results were recorded using a Minichrom data handling system (VG Data Systems, Altrincham, UK) and expressed in milligrams/100 grams of cheese.

**Sensory evaluation.** The cheeses were graded after 1, 15, 30, 60, and 90 d of ripening by 5 graders (from the permanent staff of the Department of Food Engineering, Cukurova University, Turkey) who were experienced cheese tasters and familiar with Turkish White-brined cheese. They graded the cheeses for appearance (scale 0 to 20), body and texture (scale 0 to 35), odor (scale 0 to 10), and flavor (scale 0 to 35), according to the Turkish Standards for Turkish White-brined cheese, No 591 (TSE, 1995). Coded cheese samples were removed from the refrigerator about 1 h before evaluation and kept at room temperature. Approximately 400 g of cheese was presented to each panel member. Water and bread were also provided to the panelists to rinse their mouths between samples. Graders were requested to rank the samples in order according to their overall quality or general acceptability.

**Statistical Analyses**

A randomized complete block design which incorporated 3 treatments (cheeses A, B, or C), 5 ripening periods (1, 15, 30, 60, or 90 d), and 3 blocks (trials) was used to analyze the response variables relating to cheese composition, proteolysis, lipolysis, and sensory data. Analysis of variance was performed using the GLM procedure of SAS (SAS Institute, 1995) where the effect of treatment and replicates were estimated for response variables. Duncan’s multiple-comparison test was used as a guide for pair comparisons of treatment means. The level of significance of differences between treatments was determined at \( P < 0.05 \).

Data from urea-PAGE gels of the pH 4.6-insoluble fractions were analyzed using multivariate statistical techniques. Similar bands were recognized visually and matched as described by McSweeney et al. (1994) in the urea-PAGE gels. Volumes of corresponding bands were measured densitometrically and used as data for statistical analysis using principal components analysis (PCA). Reverse phase-HPLC chromatograms of the 70% ethanol-soluble or insoluble fractions were analyzed by multivariate statistical analyses. Data for multivariate statistical analysis of the RP-HPLC chromatograms were obtained by visually recognizing similar peaks in the chromatograms and using the peak heights as variables in the analysis. The peak heights were obtained by converting the corresponding chromatogram to an ASCII file. Principal components analysis was performed using the covariance matrix and varimax rotation, and hierarchical cluster analysis (HCA) was performed using Euclidean distance and average linkage without standardizing the variables (Pripp et al., 1999). Statistical analysis was performed using SPSS version 11.0 (SPSS Inc., Chicago, IL).

**RESULTS AND DISCUSSION**

**pH Changes During Cheese Manufacture**

Acid development is a very important criterion in the manufacture of cheeses ripened under brine due to the inhibitory effect of lactic acid on undesirable microorganisms, and curd stability during brining (Abd El-Salam et al., 1993; Bintis and Papademas, 2002). Changes in the pH of the cheese curd during cheese making are shown in Figure 1. The cheeses made from milk inoculated with a starter exhibited similar pH profiles during cheese manufacture. In SF cheeses, the pH of the curds after coagulation were the same as in the other cheeses, but after 60 min, there was no further acid development in SF cheeses until the end of cheese making. After 210 min, the pH of the 317HP and 763SK11 cheeses decreased sharply to 5.21 and 5.65, respectively. The pH of the cheeses made with a starter...
culture continued to decrease during preripening and reached pH 5.0 by the next morning. However, the pH of SF cheeses remained >5.0.

Gross Composition and pH

The chemical composition of Turkish White-brined cheeses during ripening is shown in Table 1. There were no significant differences in gross composition between the 2 cheeses made using different starter strains. However, SF cheeses were significantly different from 317HP or 763SK11 cheeses in terms of gross composition and curd acidity. Acidification with a starter culture influenced curd acidity, which caused differences in the syneresis of the curd during cheese making. The initial pH of the SF cheeses was 6.04, whereas that in the 763SK11 and 317HP cheeses was 4.97 and 4.84, respectively. The pH of the SF cheeses decreased during ripening (from 6.04 on d 1 to 5.19 on d 90), whereas the pH difference of cheese made using a starter culture was not significant (P > 0.05) during ripening. Because of the higher initial pH of the SF cheese compared with that of starter-added cheeses and the action of native microflora or nonstarter lactic acid bacteria on lactose in the SF cheese, a rapid decrease was seen in the SF cheese during ripening. No major differences were noted in terms of fat-in-DM content, with the level slightly lower in the SF cheeses than in other cheeses due to its higher moisture content. The mean moisture content of the SF cheese samples was significantly higher (P < 0.05) than that of the 763SK11 or 317HP cheese samples. The total protein content of the SF cheeses was significantly lower than that of the other cheeses during the ripening period. The difference can be attributed to the concentration effect on compositional parameters; the level of moisture was high in the SF cheese (Table 1). Differences in pH between the cheeses made with starter cultures and SF cheeses resulted in significant differences in the levels of cheese moisture or salt-in-moisture as well as lactic acid content. The salt-in-moisture content of all cheeses was higher than would be expected for Turkish White-brined cheese, d SF 763SK11 317HP Age of cheese, d SF 763SK11 317HP

Table 1. Chemical composition and pH values of Turkish White-brined cheeses at 1, 15, 30, 60, and 90 d.

<table>
<thead>
<tr>
<th>Age of cheese, d</th>
<th>pH</th>
<th>Titratable acidity</th>
<th>Moisture, %</th>
<th>Fat in DM, %</th>
<th>Salt-in-moisture, %</th>
<th>Protein, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>SF</td>
<td>763SK11</td>
<td>317HP</td>
<td>SF</td>
<td>763SK11</td>
<td>317HP</td>
<td>SF</td>
</tr>
<tr>
<td>1</td>
<td>6.04 ± 0.05a</td>
<td>4.97 ± 0.13b</td>
<td>57.39 ± 1.02a</td>
<td>50.81 ± 1.30b</td>
<td>50.18 ± 1.02a</td>
<td>12.78 ± 0.02a</td>
</tr>
<tr>
<td>15</td>
<td>5.59 ± 0.05a</td>
<td>4.94 ± 0.05b</td>
<td>57.56 ± 0.74a</td>
<td>50.87 ± 0.75b</td>
<td>50.11 ± 0.53a</td>
<td>12.84 ± 0.02a</td>
</tr>
<tr>
<td>30</td>
<td>5.48 ± 0.06a</td>
<td>4.81 ± 0.06b</td>
<td>58.63 ± 0.47a</td>
<td>51.74 ± 1.45b</td>
<td>51.16 ± 0.13b</td>
<td>12.91 ± 0.22a</td>
</tr>
<tr>
<td>60</td>
<td>5.33 ± 0.15a</td>
<td>4.80 ± 0.06b</td>
<td>49.04 ± 0.53a</td>
<td>49.51 ± 1.57b</td>
<td>49.53 ± 0.38b</td>
<td>12.95 ± 0.46a</td>
</tr>
<tr>
<td>90</td>
<td>5.19 ± 0.18a</td>
<td>4.78 ± 0.06b</td>
<td>51.76 ± 0.85a</td>
<td>52.89 ± 0.90b</td>
<td>51.11 ± 0.49b</td>
<td>12.95 ± 0.85b</td>
</tr>
</tbody>
</table>

Means in the same row having different letters are significantly different (P < 0.05).

Cheeses: SF = starter free; 763SK11 = L. lactis ssp. lactis NCDO763 plus L. lactis ssp. cremoris SK11; 317HP = L. lactis ssp. lactis UC317 plus L. lactis ssp. cremoris HP.

Titratable acidity expressed as percentage of lactic acid.
Figure 2. Formation of total free fatty acids during ripening of Turkish white-brined cheeses made with starter culture systems 317HP (■), 763SK11 (□), and SF (○). Cheeses: SF = starter free; 763SK11 = *L. lactis* ssp. *lactis* NCD0763 plus *L. lactis* ssp. *cremoris* SK11; 317HP = *L. lactis* ssp. *lactis* UC317 plus *L. lactis* ssp. *cremoris* HP.

brined cheese; however, protein and fat-in-DM content of the cheeses were in the normal ranges. No major differences were observed between the 763SK11 and 317HP cheeses in terms of pH or gross composition. The results confirmed the findings of Litopoulou-Tzanetaki et al. (1993), Karakus and Alperden (1995), Pappas et al. (1996), and Pappa and Anifantakis (2001).

**Total FFA**

The extent of lipolysis in the cheeses during ripening is shown in Figure 2. The total concentration of FFA in all cheeses increased during ripening, except at 15 d for cheeses made with a starter culture. At 1 and 15 d, the SF cheeses contained significantly lower levels of FFA than the other cheeses, whereas the level of FFA was higher in the 763SK11 cheeses than in the SF or 317HP cheeses after 60 d of ripening. The SF cheeses had the lowest level of lipolysis during aging. The results indicate that each starter culture caused a different level of lipolysis in cheese. This observation is in agreement with the results of Awad et al. (1999), who found that starter-free cheeses had the lowest FFA index, and starter strains contributed to lipolysis to different extents. Kiliç et al. (1998) reported that Tulum cheese made with a starter had a higher level of FFA than starter-free Tulum cheeses during ripening, probably due to lipase activity of starter bacteria.

**WSN.** The concentration of WSN as a percentage of total N in all cheeses increased significantly (*P* < 0.01) during ripening (Table 2). The level of WSN was significantly lower in SF cheeses at d 1 or 15 than in the 763SK11 and 317HP cheeses; however, WSN increased sharply at d 90 in SF cheeses, reaching a value of 23.68%, whereas this fraction in the 763SK11 and 317HP cheeses was 20.85 and 19.69%, respectively. At the end of ripening, the level of WSN was highest in SF cheeses, which is attributed to the proteolytic activity of nonstarter lactic acid bacteria that dominate toward the end of ripening (Crow et al., 1993; Moatsou et al., 1999), and to plasmin activity due to the high pH of cheese (Rank et al., 1985; Lynch et al., 1997; Michaelidou et al., 1998; Fox et al., 2000). The different starter cultures had no significant effect on the level of WSN in the cheeses at any stages of ripening, except at d 1 and 30. This indicates that the starter organisms do not make a direct contribution to the WSN content in cheese (Fox et al., 1993; Madkor et al., 2000).

**TCA-SN.** The content of TCA-SN in all cheeses increased linearly at a significant level (*P* < 0.01) during ripening (Table 2). The 317HP cheeses had the highest level of TCA-SN until 30 d of ripening; thereafter, this fraction was found at the highest level in 763SK11 cheeses. Differences in the level of TCA-SN between the cheeses were significant (*P* < 0.05) and as ripening advanced, these differences became greater. The observed differences in the level of TCA-SN between the cheeses were significant (*P* < 0.05) and as ripening advanced, these differences became greater. The observed differences in the level of TCA-SN in the 763SK11 and 317HP cheeses may be attributed to different proteolytic systems of the starters used (Law et al., 1992). A limited production of TCA-SN in the SF cheeses compared with the starter-ripened cheeses was evident during ripening, demonstrating the importance of starter lactococci in the formation of low molecular weight peptides and amino acids in cheese (O’Keeffe et al., 1976; Visser, 1977b; Rank et al., 1985).

**PTA-SN.** The formation of PTA-SN in the cheeses showed a similar trend to TCA-SN (Table 2). The level of amino acids in cheeses during ripening confirmed the results obtained from PTA-SN fraction as reported by Kawabata et al. (1997). Significant differences were found between the cheeses; that is, the formation of PTA-SN was lowest in SF cheese, and was highest in the 763SK11 cheeses during ripening, except at d 15 (Table 2). Visser (1977b) and Carmona et al. (1999) obtained similar results for starter-free cheeses; they noted that PTA-SN was formed at a very low level in these cheeses during ripening. The PTA-SN in the 763SK11 cheeses increased steadily during ripening and reached 3.2%. Use of starter bacteria in cheese manufacture resulted in the accumulation of free amino
Table 2. Soluble nitrogen fractions in Turkish White-brined cheeses.

<table>
<thead>
<tr>
<th>Age of cheese, d</th>
<th>SF</th>
<th>763SK11</th>
<th>317HP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total N, %</td>
<td>1</td>
<td>2.00 ± 0.00&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.71 ± 0.13&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>2.01 ± 0.10&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.71 ± 0.05&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>2.03 ± 0.07&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.70 ± 0.09&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>2.02 ± 0.05&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.52 ± 0.06&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>90</td>
<td>2.03 ± 0.13&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.37 ± 0.03&lt;sup&gt;ab&lt;/sup&gt;</td>
</tr>
<tr>
<td>Water-soluble N, % of total N</td>
<td>1</td>
<td>6.31 ± 0.43&lt;sup&gt;a&lt;/sup&gt;</td>
<td>7.88 ± 0.33&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>15</td>
<td>8.31 ± 0.11&lt;sup&gt;a&lt;/sup&gt;</td>
<td>10.46 ± 0.52&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>13.15 ± 0.74&lt;sup&gt;a&lt;/sup&gt;</td>
<td>10.56 ± 1.13&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td></td>
<td>60</td>
<td>14.61 ± 0.66&lt;sup&gt;a&lt;/sup&gt;</td>
<td>13.80 ± 0.82&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
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<td></td>
<td>90</td>
<td>23.68 ± 0.74&lt;sup&gt;a&lt;/sup&gt;</td>
<td>20.85 ± 0.72&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>12% TCA-soluble N, % of total N</td>
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<td>1.46 ± 0.16&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.97 ± 0.45&lt;sup&gt;b&lt;/sup&gt;</td>
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<td></td>
<td>15</td>
<td>2.58 ± 0.30&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.12 ± 0.25&lt;sup&gt;ab&lt;/sup&gt;</td>
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<td>30</td>
<td>4.58 ± 0.42&lt;sup&gt;a&lt;/sup&gt;</td>
<td>5.90 ± 0.44&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>5.83 ± 0.82&lt;sup&gt;a&lt;/sup&gt;</td>
<td>8.99 ± 1.09&lt;sup&gt;ab&lt;/sup&gt;</td>
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<td>7.56 ± 0.51&lt;sup&gt;a&lt;/sup&gt;</td>
<td>13.46 ± 0.80&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
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<td>5% phosphotungstic acid-soluble N, % of total N</td>
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<td>0.51 ± 0.05&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.84 ± 0.07&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>15</td>
<td>0.57 ± 0.11&lt;sup&gt;a&lt;/sup&gt;</td>
<td>0.75 ± 0.03&lt;sup&gt;ab&lt;/sup&gt;</td>
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<td>30</td>
<td>0.80 ± 0.12&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.37 ± 0.18&lt;sup&gt;b&lt;/sup&gt;</td>
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<td>60</td>
<td>0.98 ± 0.20&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.01 ± 0.40&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
<td></td>
<td>90</td>
<td>1.59 ± 0.27&lt;sup&gt;a&lt;/sup&gt;</td>
<td>3.22 ± 0.48&lt;sup&gt;b&lt;/sup&gt;</td>
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</table>

Means in the same row having different letters are significantly different (<i>P</i> < 0.05).

Cheeses: SF = starter free; 763SK11 = <i>L. lactis</i> ssp. <i;lactis</i> NCDO763 plus <i>L. lactis</i> ssp. <i>cremoris</i> SK11; 317HP = <i>L. lactis</i> ssp. <i;lactis</i> UC317 plus <i>L. lactis</i> ssp. <i>cremoris</i> HP.

acids, and as ripening proceeded, this difference became more evident, in agreement with the reports of several researchers, including Visser (1977b), Crow et al. (1993), Lane and Fox (1997), and Carmona et al. (1999).

**Total FAA**

The concentrations of total FAA in cheeses, as measured by the Cd-ninhydrin method, increased (Figure 3) gradually until 60 d, and then sharply at 90 d of ripening. The 763SK11 and 317HP cheeses contained significantly higher concentrations of FAA than did the SF cheeses during ripening. As expected based on the results for PTA-SN, a greater increase in the level of FAA was observed in 763SK11 cheese than in 317HP cheese during ripening, except at d 1. There was a 2-fold difference in the level of FAA between cheeses made with starter (763SK11 or 317HP) and the SF cheeses; however, at 90 d, the level of FAA in SF cheeses increased sharply and was close to that of the 317HP cheese. Increased levels of FAA in the SF cheeses may be attributed to the contribution of nonstarter lactic acid bacteria to the accumulation of amino acids toward the end of ripening (Lane and Fox, 1996).

**Individual FAA**

Free AA are released by the action of lactococcal peptidases on small- and medium-sized peptides and contribute directly to cheeses flavor or act as precursors for flavor compounds (McSweeney and Sousa, 2000). The extent of proteolysis in the cheeses (expressed as mg/100 g) was also monitored by determining the levels of individual FAA at d 60. As shown in Figure 4, amino acid profiles of the cheeses reflected both the use of a
Figure 4. Individual free amino acid concentrations at 60 d in Turkish white-brined cheeses made with starter culture systems 317HP (□), 763SK11 (■), and SF (▲). Cheeses: SF = starter free; 763SK11 = L. lactis ssp. lactis NCDO763 plus L. lactis ssp. cremoris SK11; 317HP = L. lactis ssp. lactis UC317 plus L. lactis ssp. cremoris HP.

Two bands with faster electrophoretic mobility than $\alpha_{\text{s1}}$-casein (f24–199) appeared in the electrophoretograms of the SF cheeses after 30 d of ripening (Figure 5). The region of $\alpha_{\text{s1}}$-casein degradation products on the gel was different in 763SK11 and 317HP cheeses after 60 d of ripening, perhaps due to differences in the specificity of cell envelope-associated proteinase. For example, L. lactis ssp. lactis NCDO763 and cremoris SK11 have a PIII-type enzyme (Visser, 1993), whereas L. lactis ssp. lactis UC317 and cremoris HP have PIII- and PI-type enzymes, respectively (Law et al., 1992; Tan et al., 1993).

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shortest ripening times (1- and 15-d-old cheeses) were located separately, and other groups for 30, 60, or 90 d of ripening for each cheeses separated as marked on Figure 6.

**RP-HPLC Peptide Profiles**

70% (vol/vol) ethanol-soluble fractions. Figure 7 shows the RP-HPLC peptide profiles of the ethanol-soluble (EtOH-s) fractions of the pH 4.6-soluble fractions of the 60- and 90-d-old cheeses. Chromatograms of the 70% EtOH-s fractions were similar to those of pH 4.6-soluble fractions of the cheeses. The 763SK11 cheeses showed higher concentrations of peptides eluting with retention times of 9 and 13 min than the other cheeses. These peaks were probably due to amino acids or low-molecular weight peptides produced by the action of starter strains (NCDO763 or SK11 strains). In our previous study (Hayaloglu et al., 2004), we observed that the use of NCDO763 or SK11 in Turkish White-brined cheese gave peaks with corresponding retention times. The peaks eluting with retention times of 26 or 33 min were common in the cheeses; however, their concentrations were different. Higher concentrations of peptides with retention times of 40 to 60 min were present in the EtOH-s fractions of SF cheeses than the

<table>
<thead>
<tr>
<th>Age of cheese, d</th>
<th>Residual Β-casein, %</th>
<th>Residual αs1-casein, %</th>
</tr>
</thead>
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<tr>
<td></td>
<td>SF 763SK11 317HP SF 763SK11 317HP</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>100 100 100 100</td>
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<td>60</td>
<td>89.6a 88.0b 83.5b</td>
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</tr>
<tr>
<td>90</td>
<td>84.3a 84.9b 80.1b</td>
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</table>

\(^{a,b,c}\)Means in the same row having different letters are significantly different (\(P < 0.05\)).  
\(^{1}\)Cheeses: SF = starter free; 763SK11 = *L. lactis* ssp. lactis NCDO763 plus *L. lactis* ssp. cremoris SK11; 317HP = *L. lactis* ssp. lactis UC317 plus *L. lactis* ssp. cremoris HP.

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**Table 3.** Mean values of residual Β-casein and αs1-casein in Turkish White-brined cheeses.  
Foote 1.  
cheeses made with a starter culture. These differences were confirmed by the use of multivariate statistical analyses; PCA or HCA on chromatographic data distributed the cheeses into 3 groups as shown in Figure 8. The 763SK11 cheeses, located on the positive side of PC1, had higher concentrations for peaks eluting with retention times of 9, 13, and 27 min (they could be considered hydrophilic peptides) than the other cheeses. Similar results were reported by Lane and Fox (1997), Shakeel-Ur-Rehman et al. (1999), and Poveda et al. (2003), who reported that the starter enzymes had a significant effect on the formation/breakdown of peptides soluble in 70% ethanol.

70% (vol/vol) ethanol-insoluble fractions. The RP-HPLC peptide profiles of the EtOH-i fractions of the 60- or 90-d-old cheeses are shown in Figure 9. Qualitative and quantitative differences were observed in the chromatograms of the cheeses, particularly in the hydrophobic region (from 45 to 60 min). No major differences were observed between the peptide profiles of EtOH-i fractions from 763SK11 or 317HP cheeses at
early retention times (5 to 20 min); it can be concluded that hydrophilic peptides were soluble in 70% ethanol, whereas the hydrophobic peptides were mainly in the EtOH-i fraction. Chromatograms of the EtOH-i fractions showed that some of the peptides present at higher concentrations were eluted in the hydrophobic region (40 to 60 min) in SF cheeses. Urea-PAGE electrophoretograms of the EtOH-i fractions of the cheeses showed that the 763SK11 or 317HP cheeses exhibited similar patterns on the gels, but SF cheeses showed clear differences from the others as shown in Figure 10. Comparing the urea-PAGE patterns of the EtOH-i fractions of the cheeses, we suggest that the starter enzymes played a significant role in the breakdown of these fractions during ripening (Visser, 1977a; Lau et al., 1991; Tan et al., 1993; Lee et al., 1996; Gomez et al., 1997; Rampilli et al., 1997). Principal component analysis and HCA obtained from the RP-HPLC of the EtOH-i fractions grouped the cheeses. Cheeses were divided into 2 groups; one including the 763SK11 and 317HP cheeses at 60 and 90 d of ripening period, whereas the other included SF cheeses (Figure 11). However, differences were observed between the SF cheeses at 60 and 90 d of ripening period; these were located on the opposite side of PC2.

Sensory Evaluation

The mean sensory scores of the experimental cheeses are shown in Table 4. A salty taste was usually pronounced for all samples during maturation, probably
due to the salty and acidic nature of this type of cheese. No significant differences were noted between the 763SK11 and 317HP cheeses during maturation in terms of all sensory scores. However, the graders reported that 763SK11 cheeses had more flavor or flavor intensity after 15 d of ripening and that their superiority increased as the cheese aged. Although _L. lactis_ ssp. _cremoris_ HP, which was present in the 317HP cheese, produces bitter peptides (Law et al., 1992), no bitter flavor was detected in the 317HP cheeses during aging. The SF cheeses received considerably lower values for all sensory scores than the other cheeses at all ripening periods. Firstly, their appearance and texture were not typical for white-brined cheeses; the SF cheeses had a few holes inside the molds due to presence of gas-producing bacteria (due to lack of a dominant starter). In addition, the SF cheeses received the lowest scores for flavor and odor \((P < 0.01)\), indicating that the use of a starter culture in cheese manufacture resulted in some improvements in flavor and aroma development. The lower level of flavor intensity in the SF cheeses may be correlated with lower levels of N fractions and FAA of the cheese than in the other cheeses as reported by several researchers (Urbach 1993; Tzanetakis et al., 1995; Urbach, 1997; Broome and Limsowtin, 1998; Madkor et al., 2000). A soft texture was noted in all cheeses toward the end of ripening; this may be linked to the degradation of \(\alpha_{s1}\)-casein (Lawrence et al., 1987; Fenelon et al., 1999). Although the 763SK11 and 317HP cheeses received similar sensory scores, the graders differentiated the 2 cheeses and awarded the 763SK11 cheese the best final score for both flavor and odor. The graders’ preference for the cheeses was in the order 763SK11 > 317HP >> SF.

**CONCLUSIONS**

The results clearly show that the use of a starter culture in the manufacture of Turkish White-brined cheese influenced the chemistry, biochemistry, and sensory characteristics of the cheeses during ripening period. However, the different starters used did not significantly influence the gross composition or the sensory attributes. Cheeses made with NCDO763 and SK11 in combination exhibited a significantly higher extent of
proteolysis (i.e., soluble N fractions, total and individual FAA contents, peptide profiles) and higher flavor intensity than the cheeses made with UC317 and HP in combination due to the different proteolytic enzyme systems of the strains used. In contrast, cheeses manufactured without a starter had lower concentrations of soluble N, total and individual FAA, and sensory scores. Considerable differences were found between the SF cheese and the starter cheeses (763SK11 or 317HP) during cheese making and aging. Results obtained demonstrated that the use of a starter is essential for cheese making.

In general, the urea-PAGE patterns and the RP-HPLC peptide profiles of the cheeses showed slight differences between the samples in the early ripening times (until 15 d); but the differences increased at the advanced stages of ripening. The highest concentrations of total or individual FAA were found in the 763SK11 cheeses; Leu, Glu, Phe, Lys, and Val were the principal FAA in all cheeses. Principal component analysis or HCA of the data obtained from the urea-PAGE patterns and from the RP-HPLC peptide profiles of the cheeses classified the samples according to ripening time.

**ACKNOWLEDGMENT**

This research was funded in part by Scientific Research Project Units of Cukurova University (Adana, Turkey).

**REFERENCES**


