Melt and rheological properties of Mozzarella cheese as affected by starter culture and coagulating enzymes

Rajiv Dave, Pragati Sharma, Donald Mcmahon

To cite this version:
Rajiv Dave, Pragati Sharma, Donald Mcmahon. Melt and rheological properties of Mozzarella cheese as affected by starter culture and coagulating enzymes. Le Lait, INRA Editions, 2003, 83 (1), pp.61-77. 10.1051/lait:2002050 . hal-00895360

HAL Id: hal-00895360
https://hal.archives-ouvertes.fr/hal-00895360
Submitted on 1 Jan 2003

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Melt and rheological properties of Mozzarella cheese as affected by starter culture and coagulating enzymes

Rajiv I. DAVEa*, Pragati SHARMAa, Donald J. McMAHONb

a Dairy Science Department, Minnesota-South Dakota Dairy Foods Research Center, South Dakota State University, Box 2104, Brookings, SD 57007, USA
b Department of Nutrition and Food Sciences, Western Center for Dairy Protein Research and Technology, Utah State University, Logan UT 84322, USA

(Received 6 February 2002; accepted 5 June 2002)

Abstract – Low moisture part-skim (LMPs) Mozzarella cheeses were made with single culture (SC) of Streptococcus thermophilus or mixed culture (MC) of Streptococcus thermophilus and Lactobacillus helveticus using 1× or 6× of cheese coagulants chymosin or Cryphonectria parasitica (CP). Cheeses were analyzed for total solids, fat, protein, ash, salt, and calcium on day 1. Changes in melt characteristics and proteolysis during storage (4 °C) were monitored on 1, 7, 15, and 30 days (d). After 30 d storage, melt area as measured by modified Schreiber test increased only by approximately 2 times in cheeses made with SC as against 3–4 times in the cheeses made with MC. On d 30, creep test showed that the fall in height decreased by approximately 4–5 times in the SC cheeses but approximately 7–9 times in the cheeses made with MC when compared with the values obtained on d 1 for corresponding cheeses. Melt characteristics on d 7 for cheeses prepared with MC almost corresponded to that of d 30 cheeses made with SC, suggesting faster ripening and increase in melt of cheeses made with MC. Soluble nitrogen was also higher in MC cheeses as compared to those made using SC only. The degradation of total αs-casein was higher in the chymosin cheeses and that of β-casein in the CP cheeses. The degradation of casein fraction f_{24-199} was higher in MC cheeses as compared to those prepared with SC only. After 30 d storage, the highest percentage breakdown of αs-casein was approximately 75% in cheeses made with MC using 6× chymosin and that of β-casein was approximately 50% in cheese samples prepared with MC using 6× coagulant from Cryphonectria parasitica. Meltability of Mozzarella cheese was better correlated to hydrolysis of β-casein and was comparable to soluble nitrogen but least to αs-casein.

Mozzarella cheese / melt / proteolysis / starter culture / coagulant concentration

Résumé – Fonte et propriétés rhéologiques de fromage Mozzarella en fonction du levain et des enzymes coagulantes utilisés. Des fromages Mozzarella obtenus à partir de lait partiellement écrémé et à faible teneur en humidité ont été fabriqués avec une culture pure (SC) de Streptococcus thermophilus ou avec une culture mixte (MC) de Streptococcus thermophilus et de Lactobacillus helveticus using 1× or 6× of cheese coagulants chymosin or Cryphonectria parasitica (CP). Cheeses were analyzed for total solids, fat, protein, ash, salt, and calcium on day 1. Changes in melt characteristics and proteolysis during storage (4 °C) were monitored on 1, 7, 15, and 30 days (d). After 30 d storage, melt area as measured by modified Schreiber test increased only by approximately 2 times in cheeses made with SC as against 3–4 times in the cheeses made with MC. On d 30, creep test showed that the fall in height decreased by approximately 4–5 times in the SC cheeses but approximately 7–9 times in the cheeses made with MC when compared with the values obtained on d 1 for corresponding cheeses. Melt characteristics on d 7 for cheeses prepared with MC almost corresponded to that of d 30 cheeses made with SC, suggesting faster ripening and increase in melt of cheeses made with MC. Soluble nitrogen was also higher in MC cheeses as compared to those made using SC only. The degradation of total αs-casein was higher in the chymosin cheeses and that of β-casein in the CP cheeses. The degradation of casein fraction f_{24-199} was higher in MC cheeses as compared to those prepared with SC only. After 30 d storage, the highest percentage breakdown of αs-casein was approximately 75% in cheeses made with MC using 6× chymosin and that of β-casein was approximately 50% in cheese samples prepared with MC using 6× coagulant from Cryphonectria parasitica. Meltability of Mozzarella cheese was better correlated to hydrolysis of β-casein and was comparable to soluble nitrogen but least to αs-casein.

Mozzarella cheese / melt / proteolysis / starter culture / coagulant concentration

1Published with the approval of the director of the South Dakota Agricultural Experiment Station as Publication Number 3290 of the Journal Series.
*Correspondence and reprints
Tel.: (605) 688-6648; fax: (605) 688-6276; e-mail: Rajiv_Dave@sdstate.edu
**Fromage Mozzarella / fonte / protéolyse / levain / concentration en enzyme coagulante**

1. INTRODUCTION

Mozzarella cheese has become one of the most popular cheese varieties in the USA. It is estimated that approximately 70% of Mozzarella cheese is used as an ingredient on pizza. Its usage is expected to grow as global interest and demand for pizza and other foods that use Mozzarella cheese increase [1, 21]. Since most of the Mozzarella cheese is used for pizza, its functionality is of utmost importance. The factors that will affect these characteristics will in turn affect the acceptability of Mozzarella cheese. In a survey, over 50% of pizza restaurants reported occasional to frequent problems in quality of which 67% related to melting of Mozzarella cheese [35].

The functional properties of Mozzarella cheese develop in two distinct, but interdependent phases. The first phase occurs during manufacture, when the basic curd structure is established and the second occurs during storage, when functionality and curd structure alter [38]. The functional properties attributed to melted Mozzarella cheeses are wide and varied, which are largely responsible for consumer perception [38]. Defects associated with Mozzarella cheese include a rubbery, tough texture, lack of flavor, paleness or green tint, inability to melt, and poor stretchability [15, 28, 41]. These functional properties of Mozzarella cheese are influenced by a multitude of factors that include cheese composition, especially the moisture and fat contents, pH, coagulating enzymes, starter culture, homogenization, cooking and stretching, salt content, and the changes occurring during aging and storage [11, 22, 26, 38]. Moisture in cheese acts as a plasticizer and also plays an important role in cheese functionality. The protein matrix/structure also largely affects cheese functionality. As the cheeses mature, casein undergoes both hydration and hydrolysis. This results in the weakening of the structure and an increase in melt of cheese [16]. Factors that affect the hydrolysis of different casein fractions may in turn affect the functional characteristics of Mozzarella cheese. Previous studies have shown that proteolysis in Mozzarella cheese is contributed both by
the starter culture [31, 32] and coagulating enzymes [9]. Different starter cultures have different proteolytic activity that can influence the functionality of Mozzarella cheese [18, 20, 31, 32]. *Lactobacillus helveticus* has been reported to give better functional properties, i.e. increase in melt, when used as starter culture with *Streptococcus thermophilus* as compared to the more traditional *Lb. delbrueckii* ssp. *bulgaricus* [31, 32]. Different coagulating enzymes are also known to have different specificity towards casein [13, 34, 48]. Protease from *Cryphonectria parasitica* is more proteolytic and specific towards β-casein compared to chymosin, which is less proteolytic and more specific towards αs-casein [48]. Bogenrief and Olson [4] and Olson [34] reported that meltability in Cheddar cheese was more closely related to β-casein hydrolysis compared to αs-casein hydrolysis. However, information on the relationship between meltability and type of proteolysis is scant for Mozzarella cheese, which is usually not stored for longer periods such as is Cheddar cheese. This study was therefore designed to gain an insight on the effects of starter culture and coagulating enzymes on the melting and rheological properties of Mozzarella cheese. The specific objectives were to study the effects of the type of proteolysis occurring in Mozzarella cheese as a function of starter culture and type and levels of coagulating enzyme and its correlation with rheological and the melting characteristics of Mozzarella cheese.

2. MATERIALS AND METHODS

2.1. Selection of starter culture

One of the objectives of this study was to investigate the contribution of the starter culture toward proteolysis occurring in the cheese and its effects on the melting and rheological characteristics. *S. thermophilus* is less proteolytic than lactobacilli and works symbiotically with the latter for faster acid production in yogurt [30, 37]. Therefore, a single culture (SC) of direct vat set (DVS) *S. thermophilus* (DS134) or a mixed culture (MC) of proteolytic strain of *Lb. helveticus* (R13V, Utah State University, Logan, UT, USA) as bulk culture and DVS *S. thermophilus* (DS134) were selected. For cheese making, *Lb. helveticus* was initially propagated in sterilized skim milk of 110 g·kg⁻¹ solids and with yeast extract, peptone, and glucose added to activate the culture. The final propagation to obtain bulk culture was in sterilized (121 °C for 15 min) skim milk. Prior to addition to the cheese milk, *S. thermophilus* (200 mg·L⁻¹ of cheese milk, as recommended by the supplier) was mixed with *Lb. helveticus* (10 g·L⁻¹). For the manufacture of SC cheeses the rate of starter addition was 800 mg·L⁻¹, because from preliminary trials using 200, 400, 600, 800, and 1000 mg·L⁻¹ of SC, it was determined that 800 mg·L⁻¹ SC was required in order to have the same rate of acid production in milk as that observed with MC used in this study.

2.2. Selection of appropriate enzyme level

Chymosin (Chymax® Extra, Chr. Hansen, Milwaukee, WI, USA) and *Cryphonectria parasitica* protease (Sure curd® 600, DSM Food specialties, USA) were selected as the two coagulating enzymes as they are known to have different specificity towards the casein fractions.

2.3. Cheese-making

This study had eight different treatments. Two different cultures, each with two types of coagulating enzymes and two different levels, were used as described earlier. Four replications were carried out on four separate occasions and the treatments were randomly assigned to cheese vats.
Raw milk procured from the Dairy Training and Research Unit (South Dakota State University, Brookings, SD, USA) was warmed to 40 °C and centrifugally separated (Model 100AE, De Laval Separator Co., New York, NY, USA). The separated milk was standardized to 1.6% fat, pasteurized (62.8 °C for 30 min), and stored at 4 °C until used. The standardized milk was warmed in a double-jacketed vat to 30 °C and 15 kg were transferred to cheese vats that were partly immersed in water in a bigger vat in order to have identical temperature treatments during cheese making. Milk in each of the vats was then warmed to 32 °C and the starter culture (MC or SC) was added at the rates described in Section 2.1. After a ripening time of 30 min, single strength (100 µL·kg⁻¹ cheese milk) chymosin or Cryphonectria parasitica protease was added to the cheese vats and the pH of milk at the time of renneting was between 6.36 to 6.45. After another 10 min, 6× (600 µL·kg⁻¹) chymosin or Cryphonectria parasitica protease was added. While adding the coagulating enzymes, the enzyme solution was diluted 1:40 times in case of single strength and 1:6 times in case of 6× enzymes (dilution factor for enzyme was different to keep the dilution effect of milk the same due to addition of enzyme-water solution in all the cheese vats). Curd was cut using 1.1 cm wire knives followed by heating for 10 min, and then cooking to a final temperature of 40 °C in 45 min. Approximately 1/3rd of the whey was drained and curds were gently stirred until pH of the whey was around 6.0. At this point, all of the whey was drained. During cheddaring, curds were turned every 20 min until pH was 5.1 to 5.3. The time to reach this pH varied in different vats according to the starter culture (single or mixed) used for preparing the cheese. Curd was then milled into cubes of 2 cm with a knife and salt added to the curd at the rate of 20 g·kg⁻¹ and left for 15 min with intermittent mixing. Salted curds were taken in a strainer for stretching and molding, and immersed in hot water (77 °C) containing 0.5% salt. The heat treatment was kept well controlled and identical in cheeses while stretching to obtain a similar rate of destruction of residual coagulant in all replicates. Curds were kneaded together, stretched, and re-immersed in hot water (77 °C) for a further 120 s. Cheese was stretched uni-directionally, folded, and stretched. It was immersed again into hot water (77 °C) for 30 s, and blocks were made that were immersed in chilled water. Cheese blocks were taken out after 30 min, cut into 4 parts, and vacuum packaged in Cryovac bags (Cryovac Division, W.R. Grace & Co., Duncan, SC, USA) using a Spiromac vacuum packaging machine (Sogevac®, Bourg-les-Valence, France) and stored at 4 °C. One of the blocks from each group of cheeses was randomly sampled for analysis on day 1, 8, 15, and 30.

2.4. Biochemical analysis

2.4.1. Cheese composition

Total solids and fat in cheese were determined using the Mojonnier method [2, 3], total protein using the macro-Kjeldahl method [2] and ash content by heating samples in the Muffle furnace at 535 °C [2]. Salt in cheese was determined using a sodium electrode attached to an ion analyzer (Model 350, Corning Medical, Medfield, MA, USA) [23]. Calcium was determined using inductively coupled plasma atomic emission spectroscopy [42]. pH in cheese was measured using a Ross® (Model 8163, Orion Research Inc., Beverly, MA, USA) combination spear tip pH electrode connected to a Corning pH meter (Model 320). Acid soluble protein in cheese was determined by the macro-Kjeldahl method after extracting it with Sharpe’s extract [24]. All proteins were calculated by multiplying the nitrogen content by a factor of 6.38.
Melt and rheology of Mozzarella cheese

2.4.2. Capillary elecrophoresis

The method described by Dave et al. [9] was used to prepare the acid-urea extract of cheese samples, and collected filtrates were stored (–20 °C) until applied to the capillary electrophoresis (CE) column. The “control” for identification of peaks was a milk sample. Capillary electrophoresis was performed on a PACE 2100 system (Beckman Instruments, Inc., Fullerton, CA, USA). The method used by Broadbent et al. [5] for Cheddar cheese was modified for use in Mozzarella cheese as described by Dave et al. [9]. The urea concentration (4 mol·L−1) and the polymeric additive hydroxy-propyl-methyl cellulose (HPMC, 1 g·L−1) were the same, but citric acid was used for sample digestion instead of phosphoric acid. For analysis, a 75 µm × 57 cm hydrophilic coated P1 capillary (Supelco, Bellefonte, PA, USA) was used with a field strength of 316 V·cm−1 (18 kV, 23 µA at 38°C). Run buffer and other operational conditions were identical, as reported by Dave et al. [9].

2.5. Meltability and rheological analysis

2.5.1. Modified Schreiber test

Melting of the cheeses was determined using a modified Schreiber test [29]. Preliminary trials indicated that single culture cheeses did not melt at 90 °C. Therefore the temperature of melting was increased to 125 °C in our study to melt experimental “control” cheeses and to have better comparison between the treatments. Area (mm2) of the melted cheese was measured using image-processing software (HL Image++98, Western Vision Software, Salt Lake City, UT, USA).

2.5.2. Creep test

Meltability of cheese was also measured by Creep test using a UW-melt-meter [45]. A sample of cheese 7 ± 0.1 mm thick parallel to the fiber axis and approximately 28.5 mm diameter was heated to 60 °C, and fall in height at 10, 20, and 30 s was measured under a constant force of 0.36 N. Meltability was expressed as percent change in the height of cheese [45].

2.6. Statistical analysis

The experiment was a factorial randomized complete block design with culture, enzyme, and the level of enzyme as the main effects, each having two different levels. Results of the compositional analysis were analyzed using Proc GLM. Data for capillary gel electrophoresis and acid soluble protein were also analyzed using Proc GLM, because the variance was not homogeneous if the data were analyzed using Proc mixed [39].

3. RESULTS AND DISCUSSION

3.1. Composition of cheese

The average moisture and protein content of cheeses made from SC were lower than in cheese made from MC (P ≤ 0.05), but the fat, ash, and salt in moisture content were higher (P ≤ 0.05) in SC cheeses (Tab. I). There were no differences (P > 0.05) in the salt and calcium contents of SC and MC cheeses. There were no differences (P > 0.05) in composition of cheeses made with different types and levels of coagulating enzymes. The differences in the moisture content of SC and MC cheeses could be due to the confounding effects of pH, i.e. the differences in the rate of acid production by these two starter cultures. It was observed that the pH drop below 5.5 was very slow in SC cheeses. This could be due to the acid injury to St. thermophilus restricting further acid development in SC cheeses. The total time (to drop a whey pH of 5.1 to 5.2) in SC cheeses was
approximately 2–3 h longer than MC cheeses. This was not observed in MC cheeses due to faster acid production by *Lb. helveticus* and its continued symbiosis with *St. thermophilus*. Differences in proteolysis and therefore melt and rheology of cheese can be anticipated due to the differences in moisture in non-fat substances of cheeses. Similar moisture in non-fat substances in cheese could have been obtained by keeping the same cheddaring time in SC and MC cheeses. SC cheeses took a longer time for acid production compared to MC cheeses, though we increased the rate of addition of starter in SC cheeses. It has been reported that calcium content and pH of cheese influence functionality of the cheeses [27]. Hence, to keep similar pH and calcium content in these cheeses, a longer cheddaring time was employed for SC cheeses and this might have led to lower moisture content in these cheeses. Thus, it was not possible to control all the parameters (pH, moisture, and calcium content) in a complex cheese system.

Further, the differences in composition, though small, were statistically significant (*P* ≤ 0.05) because of consistency among replicates. In previous reports [36, 47], cheeses with such differences in moisture (approximately 2% range) and in other compositional parameters [15] had similar meltability. Therefore, it is reasonable to believe that the differences in melt and rheology of SC or MC cheeses are not solely because of compositional differences.

The average pH of the experimental cheeses was 5.27 for MC and 5.45 for SC cheeses on d 1 (data not shown). These differences could be due to faster and continued growth of starter culture in MC cheeses during cheddaring and milling compared to SC cheeses, where a pH drop (after a pH of 5.5) was extremely slow (data not shown). This may have an effect on the amount of residual coagulating enzymes in the cheese, which is dependent on the pH of the cheese [8, 19]. pH of cheese influences proteolytic changes, which are considered to be the most important biochemical event during the ripening of many cheese varieties. Yun et al. [46, 50], however, did not observe any differences in proteolysis and meltability of Mozzarella cheeses milled at pH ranging from 5.40–5.10. According to these authors, differences in the milling pH alone will not result in variations in the functional characteristics or in proteolysis during storage.

### Table I. Composition\(^1\) (%) of single culture (SC) and mixed culture (MC) cheeses\(^2\).

<table>
<thead>
<tr>
<th>Cheese</th>
<th>Fat</th>
<th>Protein(^3)</th>
<th>Ash</th>
<th>Calcium</th>
<th>Salt</th>
<th>SM(^4)</th>
<th>Moisture</th>
<th>MNFS(^5)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SC</td>
<td>18.79(^a)</td>
<td>32.53(^a)</td>
<td>3.53(^a)</td>
<td>1.04</td>
<td>1.05</td>
<td>2.44(^a)</td>
<td>43.23(^a)</td>
<td>53.21(^a)</td>
</tr>
<tr>
<td>MC</td>
<td>18.37(^b)</td>
<td>32.94(^b)</td>
<td>3.27(^b)</td>
<td>1.01</td>
<td>0.97</td>
<td>2.12(^b)</td>
<td>45.64(^b)</td>
<td>55.91(^b)</td>
</tr>
<tr>
<td>LSD(^6)</td>
<td>0.218</td>
<td>0.276</td>
<td>0.087</td>
<td>0.092</td>
<td>0.096</td>
<td>0.226</td>
<td>0.393</td>
<td>0.448</td>
</tr>
</tbody>
</table>

\(^a, b\) Means in columns with no common superscripts differ \((P < 0.05)\).

\(^1\) Averages were not significant for the cheeses made with different types and levels of coagulating enzymes.

\(^2\) Mean averaged by culture (2 enzymes × 2 levels × 4 replicates).

\(^3\) Total protein.

\(^4\) Salt in moisture.

\(^5\) Moisture in non-fat substance.

\(^6\) Least significant difference.
3.2. Proteolysis in cheese

The capillary electrophoretograms of the mixed culture cheeses made with chymosin or protease from _Cryphonectria parasitica_ are illustrated in Figure 1. The percent corrected peak areas were used for calculating the hydrolysis of αs-casein and β-casein. The figure clearly showed that as the age of the cheese increased there was a corresponding decrease in the peak area for αs-casein and β-casein in the experimental cheeses. On d 1, the electrophoretogram of the cheeses was similar to that of milk, but as the cheeses aged there was a decrease in the peak area of αs-casein in case of chymosin cheeses and both caseins (β-casein and αs-casein) in case of cheeses made with CP protease. For chymosin cheeses, the disappearance of the αs-casein peak appeared to be well correlated with the appearance of the fraction 24–199 of αs1-casein (Figs. 1, 2 and 4). Also, it is evident from Figure 1 that the pattern of casein breakdown (as observed by several different peaks on electrophoretogram on d 30) was different in cheeses made with SC or MC using chymosin or CP protease.

### 3.2.1. Breakdown of casein fractions

Hydrolysis of αs-casein was significantly higher ($P \leq 0.05$) in cheeses made with chymosin compared to those made with CP protease (Tab. II and Fig. 2). Similarly, 6× cheeses also showed significantly higher ($P \leq 0.05$) hydrolysis of αs-casein compared to 1× cheeses (Tab. II; Fig. 2). Culture also had a significant effect ($P \leq 0.05$) on the degradation of αs-casein as did the interactions between culture×enzyme, culture×level, enzyme×level, culture×time, enzyme×time, level×time, and culture×enzyme×time. Effect of other higher order interactions was, however, not significant ($P > 0.05$) (Tab. II).
Among the various treatments, maximum hydrolysis of $\alpha_s$-casein was observed in MC cheeses made with $6 \times$ chymosin (approximately 70%), followed by SC cheeses made with $6 \times$ chymosin (approximately 40%) on d 30 (Fig. 2). In case of 1× cheeses, the degradation was lower than in 6× cheeses. Higher hydrolysis of $\alpha_s$-casein for 6× cheeses could be due to the higher concentration of the residual coagulating enzyme in these cheeses. Chymosin caused more degradation of $\alpha_s$-casein as compared to *Cryphonectria parasitica* protease. Also, the degradation of $\alpha_s$-casein was higher in MC cheeses compared to SC cheeses (Fig. 2). Results of our study are similar to other studies, where chymosin has been shown to hydrolyze $\alpha_s$-casein but was found to have almost no effect on $\beta$-casein [8, 10, 48, 49]. Chymosin predominantly cleaves leu-X and phe-X bonds but degrades $\alpha_s$-casein in cheese much more extensively than $\beta$-casein [44]. De Jong [10] reported negligible hydrolysis of $\alpha_s$-casein in chymosin-free cheeses. Faster proteolysis (Figs. 2 to 5) in MC cheeses could be attributed to their lower pH compared to SC cheeses because of greater residual chymosin activity at lower pH [8, 19, 25]. It may also be due to the activity of the starter culture, which is known to be proteolytically active during the initial stages of cheese ripening [12]. In addition, *Lb. helveticus* is more proteolytic than *St. thermophilus* [17, 18, 20, 31, 32]. Often, most of the proteolysis contributed by the starter culture occurs after the residual coagulant retained in the cheese first hydrolyzes casein [12, 19, 33].

Breakdown of $\alpha_{s1}$-casein results in the formation of two fractions viz. $f_{(1-23)}$ and $f_{(24-199)}$. Appearance of the fraction $f_{(24-199)}$ of $\alpha_{s1}$-casein was therefore monitored to countercheck if the hydrolysis of $\alpha_s$-casein was correlated with the formation and accumulation of the fraction $\alpha_{s1}$-casein $f_{(24-199)}$. Similar to the hydrolysis of $\alpha_s$-casein, significant differences ($P \leq 0.05$) were observed in the appearance of $\alpha_{s1}$-casein $f_{(24-199)}$ in cheeses made with different types of starter culture and enzymes at different levels (Figs. 2 and 4). Differences were also significant ($P \leq 0.05$) as the cheeses aged. All of the two-way interactions, namely culture×enzyme, culture×level, enzyme×level, culture×time, enzyme×time, and level×time, and the interactions between culture×enzyme×time were significant ($P \leq 0.05$) (Tab. II). Appearance of this fraction was the highest (Fig. 4) in MC cheese with 6× chymosin (average peak area of approximately 17) followed by that in SC cheeses with

**Figure 2.** Effect of starter culture and type and level of coagulating enzyme on $\alpha_s$-casein hydrolysis during storage (4 °C) of Mozzarella cheese. a = cheeses made with single culture and b = cheeses made with mixed culture. R1× (●) = cheeses made with single strength chymosin; R6× (▲) = cheeses made with six times concentration of chymosin; CP1× (▲) = cheeses made with single strength *Cryphonectria parasitica*; CP6× (●) = cheeses made with six times *Cryphonectria parasitica*. 
 Results also indicated that in the case of chymosin cheeses at 6× level, the appearance of the fraction was faster and the effect was again found to be higher in MC cheeses as compared to SC cheeses throughout the storage period. As the concentration of αs-casein decreased in the various cheeses, initially there was a corresponding increase in the appearance of αs1-casein f (24–199) (Figs. 2 and 4). However, due to the further hydrolysis of f (24–199) by residual chymosin or starter culture enzymes in some cheeses, the increase in the peak area was not as steep as it was initially, and in some cases even a decrease in the peak area was observed after 15 d storage period (Fig. 4b).

Hydrolysis of β-casein showed a similar pattern as that observed for αs-casein. There was a significant effect (P ≤ 0.05) of culture, enzyme, level, and age (Tab. II). In general, the hydrolysis of β-casein was higher in cheeses made with MC (approximately 23%), CP (approximately 24%), and 6× (approximately 21%) compared to those made with SC (approximately 11%), chymosin (approximately 9%), and 1× (approximately 12%), respectively (data not shown). Age of the cheese also had a significant effect on the hydrolysis of β-casein and the trend was

Table II. Analysis of variance for αs-casein, β-casein, soluble protein, meltability, and creep test.

<table>
<thead>
<tr>
<th></th>
<th>% αs-casein</th>
<th>% β-casein</th>
<th>% Soluble protein</th>
<th>Melt area (mm²)</th>
<th>Creep (30 s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Culture¹</td>
<td>72.45***</td>
<td>42.83***</td>
<td>220.98***</td>
<td>428.41***</td>
<td>468.33***</td>
</tr>
<tr>
<td>Enzyme²</td>
<td>227.45***</td>
<td>75.48***</td>
<td>5.38*</td>
<td>17.48***</td>
<td>0.24 NS</td>
</tr>
<tr>
<td>Culture×Enzyme</td>
<td>27.63***</td>
<td>59.15***</td>
<td>6.59*</td>
<td>19.02***</td>
<td>14.76***</td>
</tr>
<tr>
<td>Level ³</td>
<td>299.96***</td>
<td>24.54***</td>
<td>122.07***</td>
<td>10.54***</td>
<td>8.77***</td>
</tr>
<tr>
<td>Level×Culture</td>
<td>50.84***</td>
<td>14.48***</td>
<td>60.98***</td>
<td>4.40*</td>
<td>1.58 NS</td>
</tr>
<tr>
<td>Level×Enzyme</td>
<td>97.93***</td>
<td>27.74***</td>
<td>6.96***</td>
<td>0.35 NS</td>
<td>0.51 NS</td>
</tr>
<tr>
<td>Culture×Enzyme×Level</td>
<td>5.12*</td>
<td>14.07***</td>
<td>6.13**</td>
<td>2.14 NS</td>
<td>0.06 NS</td>
</tr>
<tr>
<td>Age⁴</td>
<td>30.55***</td>
<td>36.89***</td>
<td>73.57***</td>
<td>116.63***</td>
<td>376.39***</td>
</tr>
<tr>
<td>Age×Culture</td>
<td>5.44***</td>
<td>9.46***</td>
<td>30.65***</td>
<td>30.31***</td>
<td>10.49***</td>
</tr>
<tr>
<td>Age×Enzyme</td>
<td>2.09 NS</td>
<td>10.78***</td>
<td>2.61 NS</td>
<td>1.64 NS</td>
<td>0.23 NS</td>
</tr>
<tr>
<td>Age×Culture×Enzyme</td>
<td>0.82 NS</td>
<td>5.82***</td>
<td>3.22*</td>
<td>1.68 NS</td>
<td>1.10 NS</td>
</tr>
<tr>
<td>Age×Level</td>
<td>7.28***</td>
<td>2.79*</td>
<td>17.10***</td>
<td>1.90 NS</td>
<td>5.09***</td>
</tr>
<tr>
<td>Age×Level×Culture</td>
<td>1.49 NS</td>
<td>3.05*</td>
<td>8.70***</td>
<td>0.78 NS</td>
<td>0.49 NS</td>
</tr>
<tr>
<td>Age×Level×Enzyme</td>
<td>1.73 NS</td>
<td>1.57 NS</td>
<td>1.86 NS</td>
<td>0.39 NS</td>
<td>0.76 NS</td>
</tr>
<tr>
<td>Age×Culture×Enzyme×Level</td>
<td>0.79 NS</td>
<td>1.07 NS</td>
<td>2.26 NS</td>
<td>0.20 NS</td>
<td>0.39 NS</td>
</tr>
</tbody>
</table>

*Significant at P < 0.05; **significant at P < 0.01; ***significant at P < 0.001; NS not significant at P > 0.05.

¹ There were two types of culture (single or mixed culture).
² There were two types of enzymes (chymosin or sure curd).
³ There were two levels of additions (1× or 6×).
⁴ Cheeses were analyzed on d 1, 7, 15, and 30.
similar throughout storage as observed for α_5-casein. Also, the interactions between culture×enzyme, culture×level, enzyme×level, culture×time, enzyme×time, level×time, and culture×enzyme×time were significant (Tab. II). Among the treatments, maximum hydrolysis of β-casein was observed in MC cheeses made with 6× CP (approximately 51%) followed by that in MC cheeses made with 1× CP (approximately 22%) (Fig. 3b).

In contrast to α_5-casein, β-casein was hydrolyzed to a greater extent when CP protease was used as the coagulating enzyme, supporting previous observations of Yun et al. [48] and Bogenrief and Olson [4], who reported that coagulating enzyme from CP was more specific to β-casein hydrolysis in Mozzarella and Cheddar cheeses. Protease from CP is also heat sensitive (more than chymosin) but if it survives the heat treatment, it is highly
proteolytic [43]. Though this enzyme degraded both $\alpha_s$-casein and $\beta$-casein in pure solutions, it more specifically hydrolyzed $\beta$-casein in Gouda [43], Cheddar [4, 34] and Mozzarella [13, 48] cheeses.

Hydrolysis of $\beta$-casein in cheeses made with different types of culture followed the trends similar to those of $\alpha_s$-casein, though the extent of hydrolysis was lower. Cheeses made with MC showed much higher proteolysis compared to those made with SC cheeses, signifying the role of proteolytic enzymes of starter bacteria. It has been also reported that the retention of chymosin depends on the draining and milling pH, but retention of microbial coagulants was not influenced by the acidification rate during cheese making [8, 19]. Thus, the differences in the hydrolysis may not be pH-dependent because unlike chymosin, the retention of microbial coagulants is not pH-dependent [8, 19].

### 3.2.2. Soluble protein

Soluble protein increased in all the samples during storage (Fig. 5) and the differences were significant ($P \leq 0.05$) for culture, enzyme, level, and time. Interactions between culture×enzyme, culture×level, enzyme×level, culture×time, enzyme×time, level×time, and culture×enzyme×time were also significant (Tab. II). Soluble protein was significantly higher in MC cheeses compared to SC cheeses, and the differences were higher for the level of enzyme compared to the type of enzyme. Highest soluble nitrogen was found in MC cheeses made with 6× chymosin followed by MC cheeses made with 6× level of CP protease (Fig. 5b). Higher soluble nitrogen in MC cheeses can also be attributed to the complementary proteolytic effects of the starter culture, mainly $Lb. helveticus$ and the residual coagulating enzymes [18, 32].

The coagulating enzyme from chymosin and $Cryphoneuctria parasitica$ showed similar levels of soluble nitrogen initially; but towards the end of storage, higher soluble nitrogen was observed for cheeses made with chymosin. This could have happened because, though chymosin acts preferentially on $\alpha_s$-casein and CP protease on $\beta$-casein, they both hydrolyze total casein to a similar extent. Higher values of soluble nitrogen in the chymosin cheeses towards the end of storage may be attributed to the fact that chymosin and lactobacilli act synergistically and produce more free amino acid [18]. On the
contrary, Vanderpoorten and Weckx [43] observed that enzyme from *Cryphonectria parasitica* leads to higher NPN compared to chymosin. These differences in observations may be because the combined effect of the starter culture and the coagulating enzyme may not have been considered or because some of the enzyme may have been inactivated during cooking and stretching in our study. As described earlier, CP protease is more heat sensitive than chymosin [40]. Various researchers have reported that the coagulant is responsible for the formation of large peptides, while smaller peptides and free amino acids are produced by starter bacteria, possibly from peptides initially produced by the coagulant [33]. Further, *Lb. helveticus* is reported to have higher aminopeptidase, proline-aminopeptidase, and aminopeptidase-P activity [31, 32] and chymosin seems to complement this activity [17, 18]. These reasons may have also been responsible for the higher soluble nitrogen in the chymosin cheeses as compared to the CP cheeses in our study. Higher soluble nitrogen in 6× cheeses may be assumed to be a direct effect of the higher concentration of the coagulating enzyme on proteolysis.

3.3. Meltability of cheese

3.3.1. Schreiber test

Meltability of all the cheeses increased during storage (Fig. 6) and it was significantly (*P* ≤ 0.05) higher for MC cheeses than SC cheeses (Tab. II). Average increase in area for MC cheeses in 30 d was approximately 230% compared to approximately 80% for SC cheeses when they were compared with the average area of the SC cheeses on d 1 (Fig. 6). Effects of the type of enzyme and the level of addition were statistically significant (*P* ≤ 0.05), but the differences were not as great for SC or MC cheeses (Tab. II and Fig. 6). The average increase in area of chymosin and CP cheeses was approximately 135 and 170%, respectively (data not shown). Interactions between culture×enzyme, culture×level, enzyme×level, culture×time, enzyme×time, level×time and culture×enzyme×time were also significant (Tab. II).

Meltability also increased with an increase in the age of the cheese (*P* ≤ 0.05). Increase in meltability of cheese has been attributed to proteolysis occurring in cheese during storage. During storage, the protein matrix absorbs moisture that was
Melt and rheology of Mozzarella cheese

held within the fat serum channels. This allows the re-arrangement of proteins, which increases hydration in the surrounding matrix [6]. Then, as the proteins become more hydrated, they flow more easily when heated and thus increase the meltability of cheeses [26].

Overall, meltability of our cheeses was better correlated to the hydrolysis of $\beta$-casein ($r^2 = 0.72$) or soluble nitrogen ($r^2 = 0.76$) than to the hydrolysis of $\alpha_s$-casein ($r^2 = 0.36$) (data not shown), which is in agreement with Cheddar cheese studies of Bogenrief and Olson [4], and Creamer and Olson [7]. In our study, we observed that on d 30, the maximum hydrolysis of $\alpha_s$-casein was approximately 50% for SC cheeses and approximately 70% for MC cheeses made with 6× chymosin, but there were larger differences in meltability of these cheeses (Figs. 3 and 6). Additionally, cheeses made with CP protease mainly hydrolyzed $\beta$-casein and had an average increase in melt area of approximately 170% compared to 135% (data not shown) for the cheeses made with chymosin (which primarily hydrolyzed $\alpha_s$-casein). This implies that not just the hydrolysis of caseins is important, but the type of hydrolysis is also important. $\alpha_s$-Casein is the primary structural protein in cheese, however, breakdown of the same appears to be more related to the softness of the cheese than the flowing of cheese. Contrarily, breakdown of $\beta$-casein appears to improve the flow of cheese and therefore cheeses with greater $\beta$-casein breakdown showed increased melt in our study.

Higher level of coagulating enzymes resulted in increased melt of the cheeses (Fig. 6). This is due to the direct effect of higher proteolysis caused by the higher concentration and activity of coagulating enzymes retained in these cheeses.

Higher meltability of MC cheeses compared to SC cheeses may also be because of higher proteolysis caused by Lb. helveticus in the mixed culture [14]. Use of such proteolytic cultures has also been shown to increase the meltability of Mozzarella cheese compared to those prepared with less proteolytic starter cultures [31, 32].

Further, bacterial channels and their arrangement in a cheese protein matrix, along with the hydration of caseins, may have contributed to higher melt in the MC cheeses as previously explained by McMahon and Oberg [27] in describing the role of fat-serum channels. After stretching, the live starter bacterial population was approximately 2 logs higher in MC cheeses than in SC cheeses (data not shown). Thus, the bacterial cells may be acting as fillers in MC cheeses, and providing more proteolytic enzymes and resulting in an increase in melt.

### 3.3.2 Creep test

Creep test measures the deformation of cheese under constant force for a period of time. This test is a measure of the softness of cheeses unlike the Schreiber test, which measures the flowability of cheeses. Average values of decrease in height of the samples (after 30 s) are shown in Figures 7a and 7b.

Culture and type and level of enzymes had a significant effect ($P \leq 0.05$) on the softness of cheeses. MC cheeses showed a greater fall in height compared to cheeses made with SC, and the trend was the same throughout storage. On d 30 of storage, fall in height of MC and SC cheeses was approximately 73 and 43%, respectively, considering the original height of the samples as 100. MC and SC cheeses made with chymosin or CP protease exhibited a similar fall in height, whereas cheese with the two levels of enzymes had comparatively greater differences. Decrease in height observed with chymosin and CP cheeses was approximately 60%, but for 1× and 6× cheeses, it was approximately 54 and 62%, respectively (data not shown). Age of the cheese also had a significant effect on the softness of all cheeses. This may be attributed to the proteolysis occurring during
storage. Interactions between culture×enzyme, culture×level, enzyme×level, culture×time, enzyme×time, level×time, and culture×enzyme×time were also significant (Tab. II).

These observations highlighted a very important aspect of this study: softness and flowability of cheese are two different aspects. The type of coagulant (CP protease vs. chymosin) has comparatively less influence on the softness of cheese (Fig. 7), but it had greater influence on the meltability of the cheese (Fig. 6). Thus, cheeses made with chymosin were equally soft but were less flowable compared to the CP cheeses. The softness and flow are two different phenomena. We observed that when the intact caseins (either \(\alpha_s\)- or \(\beta\)-caseins) are hydrolyzed, it first affects the cheese protein structure and makes it softer, but the melting is not directly related to this cheese softness alone. Melting is largely governed by the flow of cheese, which appears to be better in cheeses made with coagulating enzyme that either speeds up proteolysis of \(\beta\)-caseins and/or generates smaller peptides from the first breakdown products of \(\alpha_s\)- or \(\beta\)-caseins, mainly by the starter culture proteinases and peptidases that make softer cheese to flow. De Jong [10] reported that the softening of cheeses was more often correlated to the amount of the intact \(\alpha_s\)-casein. This could possibly be because the breakdown products of caseins are largely water-soluble and do not contribute to the protein matrix.

Further, with an increased concentration of coagulating enzyme, there will be faster degradation of the already hydrolyzed larger protein fraction \(f_{24-199}\). This may have made cheese more flowable. The differences in MC and SC cheeses could be due to more activity of residual coagulating enzyme (due to confounding effects of pH in these cheeses, as explained earlier, or due to the combined action of starter culture and the coagulating enzymes) rendering faster proteolysis in MC cheeses (Fig. 5).

### 4. CONCLUSIONS

Cheeses made with MC had higher moisture content than those made with SC. This could be due to the faster rate of acid production in the former cheeses. There were, however, no significant differences
in moisture content of cheeses made with a particular type or level of coagulating enzyme. Chymosin cheeses showed greater hydrolysis of $\alpha_s$-casein, whereas *Cryphonectria parasitica* cheeses showed a greater hydrolysis of $\beta$-casein. Hydrolysis of both the caseins ($\alpha_s$ or $\beta$) was higher in MC cheeses and at 6× level of enzyme addition. Soluble nitrogen was also higher in MC cheeses and the differences were higher in 6× cheese compared to SC cheeses with 1× level of enzyme addition, respectively. Differences were, however, not very high for the type of enzyme.

Increase in the melt area as measured by the modified Schreiber test was again higher for MC cheeses and the 6× level of enzyme addition. CP cheeses showed a highest increase in melt area as against chymosin cheeses. Meltability as measured by Creep test indicated that MC cheeses softened to a greater extent compared to SC cheeses and that 6× cheeses softened better compared to 1× cheeses. Differences between chymosin and CP protease were, however, not significant. This is an important observation because it indicates that the type of proteolysis is important in determining the melt characteristics, as cheeses with similar softening may flow differently. Meltability of Mozzarella cheese was better correlated to hydrolysis of $\beta$-casein and was comparable to soluble nitrogen but least to hydrolysis of $\alpha_s$-casein.

ACKNOWLEDGEMENTS

The authors thank Dr. M. Strickland, Research Associate of Utah State University for her help with capillary electrophoresis. The authors are also indebted to Dr. K. Muthukumarappan of the Agricultural and Bio-Systems Engineering Department of South Dakota State University for use of the melt test facilities and for providing some insight at the time of concluding our results.

REFERENCES


[42] US Environmental Protection Agency, Method 6010a (Revision 1): Inductively
Melt and rheology of Mozzarella cheese


To access this journal online:
www.edpsciences.org