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PROTEOLYSIS AS A FUNCTION OF DISTANCE FROM SURFACE TO CENTRE IN A SMEAR-RIPENED IRISH FARMHOUSE CHEESE

A.S. MANE and P.L.H. McSWEENEY*
School of Food and Nutritional Sciences, University College Cork, Cork, Ireland
*Corresponding author: p.mcsweeney@ucc.ie

ABSTRACT

This study focused on proteolysis in an Irish farmhouse smear-ripened cheese by serial slicing (0.41 mm/slice) the first 2 cm from surface towards the centre of the cheese. Urea-polyacrylamide gel electrophoretograms confirmed higher proteolysis in the outer layers than at the centre. Free amino acid (FAA) analysis confirmed decrease in proteolytic activity from surface to centre. Peptides produced at depths 0.41 mm and 20.5 mm were 720 and 427 from $\alpha_s$-casein; 691 and 337 from $\alpha_l$-casein; 807 and 453 from $\beta$-casein; 180 and 109 from $\kappa$-casein. The study confirms higher proteolytic activity at surface due to action of enzymes of the smear microbiota, than at the centre of cheese and identified the agents responsible for production of many peptides.

Keywords: Gubbeen, smear-ripened cheese, proteolysis, surface, mass spectrometry
1. INTRODUCTION

Smear-ripened, or bacterial surface ripened, cheeses are semi-soft varieties with an orange-red rind. As the name suggests, smear ripened cheeses (e.g., Gubbeen, Reblochon, Tilsit, brick and Taleggio) develop on their surfaces during ripening a viscous, microbial red-orange smear composed of bacteria and yeast (MOUNIER et al., 2005; GOBETTI et al., 1997). Towards the end of the ripening of the mature red-smear cheese surface, Gram-positive bacteria, yeasts (Debaryomyces hansenii) are predominant microorganisms present (BOCKELMANN et al., 2002; BRENAN et al., 2004; RITSCHARD et al., 2018). This microbial complexity is mainly responsible for the development of the characteristic flavours in the cheeses (CORSETTI, et al., 2001; VALDÈS-STAUER and SCHERER, 1997). During the first days of ripening of smear-ripened cheeses acid-tolerant yeasts grow and metabolize lactic acid produced by the lactic starter cultures to CO₂ and H₂O, thereby increasing the pH and producing growth factors, such as pantothenic acid, that encourages the growth of the smear bacteria (ELISKASES-LECHNER and GINZINGER 1995; PRILLINGER et al., 1999; VALDES-STAUER et al., 1997; WYDER and PUHAN, 1999). Debaryomyces hansenii is the dominant yeast found in Tilsit cheese (CHURCHILL et al., 2003; COGAN et al., 2014). The yeasts isolated from the surface of Limburger cheese grew between pH 3.3 and 8.5 (KELLY and MARQUARDT, 1939; REPS, 1993) and Mycoderma yeasts isolated from the surface of Brick cheeses grow from pH 3 to 8 (IYA and FRAZIER, 1949; REPS, 1993). Presence of smear organisms on the surface of smear-ripened cheeses is associated with development of its colour, flavour and characteristic aroma (ADES and CONE 1969; RATTRAY and FOX, 1999). Although Brevibacterium linens was considered in the past as the dominant bacterium on red-smear cheese, recent studies have shown a number of other species, particularly of genera Arthrobacter, Brevibacterium, Corynebacterium, Microbacterium and Rhodococcus (REPS, 1993; ELISKASES-LECHNER and GINZINGER 1995; SEILER, 1986; VALDES-STAUER et al., 1997) are more important at the surface. Several bacterial species more recently isolated during the ripening of the smear cheeses include Argococcus casei, Arthobacter arilaitensis, Arthobacter bergerei, Brachybacterium alimentarium, Brachybacterium tyroformentans, Corynebacterium casei, Microbacterium gubbeenese, Mycetocola reblochoni, Staphylococcus succinus subsp. casei and Staphylococcus equorum subsp. linens (BORA et al., 2007; BORA et al., 2008; BRENAN et al., 2001; IRLINGER et al., 2005; PLACE et al., 2003; SCHUBERT et al., 1996).

Species of bacteria occasionally isolated from smear cheese include Staphylococcus spp., Brevibacterium aurantiacum in very high numbers, and relatively high numbers of Psychrobacter spp. C. casei, C. variabile, and M. gubbeenese. The presence of S. saprophyticus in certain varieties (e.g., Gubbeen) distinguishes them from other varieties (Livarot, Reblochon, and Limburger) (COGAN et al., 2014; REA et al., 2007).

Proteolysis is the key biochemical process in cheese during ripening (FOX, 1993). The complex proteolytic breakdown from surface towards the centre of smear-ripened cheese is a result of the combined action of bacterial and yeast proteolytic enzymes. Proteolysis in this type of cheese depends directly on the extent of penetration of the smear microbiota in the cheese curd and their proteolytic activity and decreases inwards from the surface towards the centre (CHURCHILL et al., 2003). Considerable differences in degree of proteolysis are observed, depending on depth from surface to centre of the cheese (CHURCHILL et al., 2003). Smear cheese microbiota (bacteria, moulds and yeasts) and their enzyme systems plays a key role in proteolytic, lipolytic and free amino acid catabolic activities in the cheese during ripening (FOX, 1993). Yeasts present in the surface
smear, *D. hansenii* and *G. candidum*, exhibit a strong extracellular proteolytic and or peptidolytic activity (BAUR et al., 2015). An *in situ* study by BOUTROU et al. (2006) found *G. candidum* from the surface of the cheese participates in primary proteolysis along with chymosin at pH 5.5 to 6 (CHEN and LEDFORD, 1972; GUEUEN and LENOIR, 1976) during the proteolysis. Rapid hydrolysis of α-caseins at the surface, in presence of *G. candidum* and surface microbiota of the cheese was observed also by BOUTROU et al. (2006).

Proteolytic enzymes of one of these organisms, *B. linens*, were studied by RATTRAY and FOX (1998). The purified proteinase enzyme from *B. linens* was most active at pH 8.5 and at 50°C with a molecular mass 120 kDa and peptides produced by these proteinases from α-, and β-caseins were identified (RATTRAY and FOX, 1996, 1997, 1998).

However, there is relatively little literature describing in detail proteolysis and breakdown of caseins in smear-ripened cheese. The current study attempts to characterise proteolysis in a smear ripened variety of Irish farm-house cheese Gubbeen from the surface towards the centre. The study gives an insight to the detail of the proteolysis in smear cheese providing a deeper and more comprehensive insight into the metabolic activity of microbial ecosystems of these varieties.

2. MATERIALS AND METHODS

Commercially ripe samples (300 g) of Gubbeen, an Irish farmhouse smear cheese, product of Gubbeen Farmhouse Products Ltd., Gubbeen House, Schull, Co. Cork. Ireland), were obtained in triplicate from the local market (G1, G2 and G3), made generally as outlined by MOUNIER et al. (2005). These were cut in half along the diameter of the cheese wheel. One portion was crumbled and mixed thoroughly (with rind) for compositional analysis and remaining part was used for serial slicing and proteolytic analysis (Fig. 1 A, B, C and D).

2.1. Serial slicing of cheese and peptide extraction

The cheese. Four or five blocks from each cheese were sliced separately and slices taken at the same depth were later pooled. A tissue chopper (McIlwain tissue chopper; Model TC752, Campden Instruments Ltd., Loughborough, Leics, England) with a double end razor blade (Wilkinson Sword, Edgewell Personal Care, Shelton, CT, USA) was set to give a thickness of 410 mm (0.410 mm) per slice. The blade was cleaned with 70% ethanol before and after each sectioning. Cheese blocks were sliced frozen, at thickness of 0.41 mm/slice, to a depth of 2 cm (Fig 1 A, B, C and D). These slices were collected on aluminium foil sheet with a sterile scalpel and an artist’s brush. Slices from multiple blocks from a single cheese were pooled to get a sufficient quantity (10 mg) for analysis and stored frozen in a micro-centrifuge tube. The pooled slices were weighed out (10 mg), then dispersed in 10 mM citrate buffer (pH 5.5), teased using a needle and vortexed (RotaMixer, model # 8768; Hook and Tucker Instruments Ltd., Croydon, England) for 1 min (repeated if required) for uniform mixing. The micro-centrifuge tubes were held in a pre-heated water bath at 40°C for 15 min and centrifuged (Micro-Centaur Centrifuge, MB010.CX1.5, Sanyo, Loughborough, Leics, UK) at 9,600 g for 10 min, cooled for 30 min at 4°C to remove any fat from the supernatant. The supernatant was separated and sufficient 10 % trichloroacetic acid (TCA) solution was added to the supernatant to give final concentration of 2 % TCA in the extracts, which were centrifuged again at 6,700 g for 10
min and cooled at 4°C for 15 min. The clear supernatant was separated and stored frozen in aliquots at -20°C until use. The aliquots were thawed and filtered through 0.22 µm cellulose acetate filters (Sartorious GmbH, Gottingen, Germany) for peptide analysis by mass spectrometry (Q-ToF LCMS) as described below.

**2.2. Physico-chemical analysis**

Compositional analysis of the cheeses comprised of determination of moisture content using an oven-drying method (IDF, 1982). Protein and nitrogen content of the cheeses and pH 4.6-soluble extracts were determined by the macro-Kjeldahl method (N x 6.38; IDF, 1986). Percentage of fat was determined by the Gerber method (IIRS, 1955) and percentage of NaCl was measured by a titrimetric method using potentiometric end-point determination as described by FOX (1963). Each experimental analyses were run in triplicate for each cheese sample. Statistical analyses were performed using R® 16 (R version 3.4.0; the R Foundation for Statistical Computing, University of Auckland, Auckland, New Zealand). Differences in means in between the batches and/or depth were tested by analysis of variance (one way-ANOVA) at significance level, α, of 0.05 (P value ≤ 0.05), throughout the study.

**2.3. Proteolytic analysis**

The pH of a cheese slurry (made from 25 g cheese and 50 g of deionised water) was measured using a calibrated pH meter. The same slurry was used to prepare pH 4.6-soluble and -insoluble fractions of cheeses at all time points as described by KUCHROO and FOX (1982).

Proteolysis in the slices taken from surface to centre of smear-ripened cheeses was studied by urea-polyacrylamide electrophoresis (urea-PAGE) of freeze-dried samples (10 mg/mL) of cheese slices from surface to centre (ANDREWS, 1983; SHALABI and FOX 1987; O’MAHONY et al., 2005). The gels were stained with Coomassie Brilliant Blue G250 (BLAKESLEY and BOEZI, 1977) and de-stained by several distilled water washes.
Individual free amino acids (FAAs) content was determined according to FENELON et al. (2000). Frozen extracts of cheese slices taken at depths 0.41 mm, 1.64 mm and 20.5 mm were dispersed in 10 mM citrate buffer, pH 5.5, in 2% TCA, and were used for the analysis of all three batches.

Peptide analysis from cheese extracts (Section 2.1) were conducted at Conway Institute of Biomedical and Biomolecular Sciences, University College Dublin, Ireland, on a quadrupole Orbitrap (Q-Exactive, Thermo Scientific, Waltham, MA, USA) mass spectrometer equipped with a reversed-phase Nano LC UltiMate 3000 HPLC system (Dionex LC Packings, now Thermo Scientific Waltham, MA, USA). Peptide samples were loaded onto C18 reversed phase columns (10 cm length, 75 µm inner diameter) and eluted with a linear gradient from 2 to 97% acetonitrile containing 0.5% acetic acid in 60 min at a flow rate of 250 nL/min. The injection volume was 5 µl. The mass spectrometer was operated in data dependent mode, automatically switching between MS and MS2 acquisition. Survey full scan MS spectra (m/z 300-2000) were acquired in the Orbitrap. The twelve most intense ions were sequentially isolated and fragmented by higher-energy C-trap dissociation.

2.4. Peptide identification

Raw data from the mass spectroscopy (Orbitrap Q-Exactive) were processed using MaxQuant version 1.5.5.1 (COX and MANN, 2008; TYANOVA et al., 2016a), incorporating the Andromeda search engine (COX et al., 2011). MS/MS spectra were matched to a bovine proteins custom database of previously identified entries (1,059) in the milk proteome in order to identify peptides and proteins. All searches were performed with unspecific digest. The database searches were performed with no fixed modification but with acetylation (protein N terminus) and oxidation (M) as variable modifications. Mass spectra were searched using the default setting of MaxQuant, namely a false discovery rate of 1% at the peptide level. For the generation of ion intensities for peptide profiles, signals of corresponding peptides in different nano-HPLC MS/MS runs were matched by MaxQuant applying a mass accuracy of at least 20 ppm and a maximum time window of 1 min (COX et al., 2014). The Perseus statistical software (version 1.5.5.3) (TYANOVA et al., 2016) was used to analysis the peptide ion current intensities.

3. RESULTS

3.1. Physico-chemical composition

The physico-chemical composition of three batches (G1, G2 and G3) of commercial smear-ripened cheese was determined (Table 1). pH decreased from 6.3 to 5.7 from surface to centre of the cheese, respectively, in all the batches, as was also reported by MOUNIER et al. (2005). Levels of moisture, fat, salt (NaCl), fat in dry matter (FDM), moisture in non-fat solids (MNFS), protein and nitrogen content in cheese and pH 4.6-soluble nitrogen (SN) as a percentage of total nitrogen (TN), are reported in Table 1. Moisture levels were 44 to 46% in agreement with the literature (MOUNIER et al., 2005; REA et al., 2007; COGAN et al., 2014). Salt (% NaCl) was ~2.1%, with no significant variation between batches. The % fat values were similar in all the batches (47 %) as were the values of % FDM (84% to 86%), %MNFS (83% to 87%). The gross composition was observed to be typical of smear ripened
cheese (COGAN et al., 2014). Crude protein (N x 6.38) content of cheese showed a slight variation between the batches G1, G2 and G3 (Table 1).

Table 1. Physico-chemical compositional analysis of mature smear-ripened cheese after ripening.

<table>
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<th>% Moisture</th>
<th>%Fat</th>
<th>%Salt</th>
<th>%N</th>
<th>(%Prt)</th>
<th>%pH4.6SN/TN</th>
<th>%MNFS</th>
<th>%FDM</th>
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<td>G1</td>
<td>44.11±0.5a</td>
<td>47±0.01a</td>
<td>2.13±0.05b</td>
<td>2.93±0.05a</td>
<td>19.67±0.34a</td>
<td>9.73±0.11b</td>
<td>83.21±0.95a</td>
<td>84.09±0.76a</td>
</tr>
<tr>
<td>G2</td>
<td>44.31±1.4a</td>
<td>47±0.02a</td>
<td>2.03±0.05a</td>
<td>3.36±0.15b</td>
<td>21.41±1.00b</td>
<td>6.56±0.66a</td>
<td>83.61±2.81a</td>
<td>84.43±2.22a</td>
</tr>
<tr>
<td>G3</td>
<td>46.27±0.7a</td>
<td>47±0.01a</td>
<td>2.07±0.05a</td>
<td>3.59±0.07b</td>
<td>22.07±0.48b</td>
<td>6.41±0.17a</td>
<td>87.33±1.32a</td>
<td>86.85±1.15a</td>
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Samples G1, G2 and G3 are separate batches of smear-ripened cheese used for compositional analysis, where %N= Nitrogen%, (%Prt)= crude protein% in smear-ripened cheese, total SN= pH 4.6-soluble nitrogen (SN) as a percentage of total nitrogen, %FDM= fat in dry matter and % MNFS=moisture in non-fat solids, represented by mean ± standard deviation and different superscript letters represent significant difference (P<0.05).

3.2. Proteolysis

The ratio between nitrogen soluble at pH 4.60 and total nitrogen (pH 4.6 SN/TN) (%) is an index of proteolysis and, in smear-ripened cheese, is mainly produced by the action of chymosin, plasmin, lactocepins (LacCeps) and enzymes from the smear microbiota on the caseins (ARDO et al., 2017). Levels of pH 4.6 SN/TN were higher in batch G1 as compared to batches G2 and G3 (Table 1). Overall % pH 4.6 SN/TN indicated a normal proteolysis at the end of the ripening in the cheese and values from the current study were comparable to those reported by COGAN et al. (2014) a range of smear-ripened cheeses.

Proteolysis from surface to the centre of the cheese was also studied by urea-PAGE (Fig. 2). Samples for the gel were prepared from freeze dried slices (0.41 mm/ slice) of smear-ripened cheese from surface to centre (2 cm). Extensive breakdown of αs1- and β-casein was observed in topmost layers up to 1.23 mm, followed by normal patterns of proteolysis typical of internal bacterially ripened varieties, closer to the centre. Action of plasmin was prominent for the break-down of β-caseins into γ-caseins at depth less than 1 cm from the outer surface, as was also observed in other smear ripened cheese varieties like Tilsit at early stages (day 6) of ripening (BOUTROU et al., 1999, 2005, 2006; GUIZANI et al., 2002; CHURCHILL et al., 2003). Extensive degradation of αs1-casein at the surface (Fig. 2) was likely caused by the action of chymosin together with the enzymes from the smear microbiota. Levels of degradation decreased from surface to center of the cheese, due to reduced penetration of smear enzymes. The high extent of degradation of αs1-casein at the surface despite the higher pH (which would reduce chymosin action) suggests a major role of enzymes from smear organisms. The results are similar to those observed by CHURCHILL et al. (2003) in Tilsit cheese.
Profiles of free amino acids analysed from extracts from cheeses dispersed in 10 mM citrate buffer at pH 5.5 are as shown in Fig. 3. Total amino acid contents observed were 107.19 µg/mg at 0.41 mm, 33.59 µg/mg at 1.64 mm and 33.78 µg/mg at 20.5 mm depth of cheese. Levels of individual amino acids were also found to be higher at the surface than towards centre (Fig. 3). Values at 0.41 mm for glutamic acid were 21.60 µg/mg, glycine were 16.5 µg/mg, proline were 16.56 µg/mg and of alanine were 10.03 µg/mg (Fig. 2). The outer layers of the cheese showed high levels of glutamic acid, glycine, alanine, tyrosine, and proline only particular to the surface of the smear cheese. Higher concentrations of threonine, lysine, phenylalanine and valine were found towards the inner layers of smear-ripened cheese than at the surface. Values were similar also to amino acid profiles of mould ripened cheeses (ZARMPOUTIS et al., 1997).
Figure 3. Free amino acid analysis. Concentration (µg /mg) of free amino acids in 10 mM citrate buffer pH 5.5 dispersed extracts from 0.41 mm, 1.64 mm and 20.5 mm slices of smear-ripened cheese.

3.3. Peptide profiles at different depths of the cheese

Qualitative differences were found in peptides produced at all depths. Mass spectroscopic analysis of peptides in 10 mM citrate extracts at pH 5.5 showed a large difference between the numbers of peptides produced at the outer surface as compared to the centre (depth of ~2 cm) (Fig 4). In case of the individual caseins, the number of β-casein-derived peptides was highest at all the depths, followed by peptides derived from αs1-casein, αs2- and κ-casein.

Figure 4. Total caseins at depths in smear ripened cheese. Graph representing the number of casein derived-peptides as a function of distance from surface of cheese (-O-). Bars indicate number of peptides derived from αs1-, αs2-, β- and κ-casein.
Identification of peptides during ripening was a major objective of this study. So far, very little has been reported in the literature about identification of peptides from smear cheese. The first 150 peptides with highest values of relative intensity at each depth, obtained from LCMS data were considered and sorted into peptides from α-casein, αs-casein, β-casein and κ-casein at depths 0.41 mm, 9.02 mm and 20.05 mm and were plotted on the primary structure of the proteins (Fig. 5 A, B, C and D). Identified peptides were compared with known cleavage specificities of chymosin, plasmin and lactocepins (SINGH et al., 1994, 1995, 1997; BREEN et al., 1995; FERNANDEZ et al., 2005; UPADHYAY et al., 2006; ARDO et al., 2017). The peptides were also compared with the cleavage specificities of proteinases of B. linens on αs- and β-casein found by RATTRAY and FOX (1993). However, cleavage site matches were not found in first 150 peptides from the extracts of smear cheese.

A total of 5176 α-casein fragments were produced by hydrolysis during ripening. Peptides derived from α-casein only at 0.41 mm were observed in regions 121-157 and 165-198. Those derived only at 9.02 mm and 2.05 mm and were found at regions 79-117. The region hydrolysed common to all depths, (0.41 mm, 9.02 mm and 2.05 mm) was 1-40 (Fig. 5 A). All these regions were highly susceptible to enzymatic hydrolysis for action of chymosin and lactocepins (ARDO et al., 2017) and proteinases from surface smear microbiota. The common fragment, α-CN (f1-23) produced by the action of chymosin (FOX and McSWEENEY, 1996), was present at all depths. Other major chymosin-derived peptides, α-CN (14-23), α-CN (25-36/38), α-CN (f17-23/24), α-CN (f 24-34) and α-CN (f 25-34), were found at all depths. Peptides produced by action of lactocepins and/or chymosin at their N- or C- termini (ARDO et al., 2017) included αs-CN (f30-38), α-CN (f31-37/38), α-CN (f33-40), α-CN (f154-164), α-CN (f165-193), α-CN (f169-189/190/197/198), α-CN (f170-189/190/192/198) and α-CN (f180-198/199). Lactocepin-derived peptide fragments found at all the depths were α-CN (1-8/9/13/16/17), α-CN (f7-13), α-CN (f10-16), α-CN (f122-139,142/143/144) and α-CN (131-139/142/148). Most common peptides from α-casein present only at surface were α-CN (f122-139/142/143), α-CN (f 124-142/143/144), α-CN (f 125-144) and α-CN (f 150-155/156/157) and those present only at the center were α-CN (f191/193-199). Peptides from α-casein present at all the depths were α-CN (f1-23), α-CN (f14-23), α-CN (f 24-34) and α-CN (131-139/142/148).

A total of 4744 peptides were produced by hydrolysis of α-casein during ripening. Peptides derived from α-casein only at depth 0.410 mm were from region 68-98, those common to depths 0.410 mm and 9.02 mm were from the regions 98-114 and 175-207 and peptides common at depths 9.02 mm and 2.05 mm were found in regions 89-99, 102-115 and 181-207 (Fig. 5 B). Peptides produced at all the depths were mainly due to the action of plasmin, lactocepins (ARDO et al., 2017), chymosin (McSWEENEY et al., 1994) and mainly at the surface by proteinases from the smear microbiota. Fragments produced by the action of plasmin and lactocepins at their N-termini were α-CN (f115-125) and α-CN (f115-126). Action of plasmin and lactocepins (ARDO et al., 2017) on α-casein helped to produce fragments α-CN (f80-88), α-CN (f150-161/162), α-CN (f151-156/158/161/162/163) and α-CN (f196-208), which were also found in smear ripened cheeses at all depths. Most common peptides from α-casein present only at surface were α-CN (f68-74/88/94), α-CN (f71-94/98), α-CN (f73-86/88), α-CN (f75-88/94) and α-CN (f87-95) and those present only at the center were, α-CN (f176-181/182), α-CN (f186-197), α-CN (f188-199/202), α-CN (f191-197/199/203) and α-CN (f194-203). Peptides from α-casein present at all the depths were α-CN (f115-125), α-CN (f151-156/158/161/162/163) and α-CN (f196-208).
Figure 5 A. The primary structure of bovine αs1-casein, showing the peptides in 10 mM citrate buffer pH 5.5-dispersed extracts of Gubbeen cheese slices taken at depth 0.41 mm, 9.02 mm and 20.05 mm, together with known cleavage sites of by action of unknown agent, plasmin (Pla), chymosin (Chy), or Lactocepins (LacCeps).
0.41 mm \( \alpha_{s2} \)-CN Unknown agent; Plasmin; Chymosin; LacCeps.

9.02 mm \( \alpha_{s2} \)-CN Unknown agent; Plasmin; Chymosin; LacCeps.

20.05 mm \( \alpha_{s2} \)-CN Unknown agent; Plasmin; Chymosin; LacCeps.

Figure 5 B. The primary structure of bovine \( \alpha_{s2} \)-casein, showing the peptides in 10 mM citrate buffer pH 5.5-dispersed extracts of Gubbeen cheese slices taken at depth 0.41 mm, 9.02 mm and 20.05 mm, together with known cleavage sites of by action of unknown agent, plasmin (Pla), chymosin (Chy), or Lactocepins (LacCeps).
0.41 mm β-CN Unknown agent; Plasmin; Chymosin; LacCeps.

9.02 mm β-CN Unknown agent; Plasmin; Chymosin; LacCeps.

20.05 mm β-CN Cheese: Unknown agent; Plasmin; Chymosin; LacCeps.

Figure 5 C. The primary structure of bovine β-casein, showing the peptides in 10 mM citrate buffer pH 5.5-dispersed extracts of Gubbeen cheese slices taken at depth 0.41mm, 9.02 mm and 20.05 mm, together with known cleavage sites of by action of unknown agent, plasmin (Pla), chymosin (Chy), or Lactocepins (LacCeps).
Figure 5 D. The primary structure of bovine κ-casein, showing the peptides in 10 mM citrate buffer pH 5.5-dispersed extracts of Gubbeen cheese slices taken at depth 0.41 mm, 9.02 mm and 20.05 mm, together with known cleavage sites of by action of unknown agent, plasmin (Pla), chymosin (Chy), or Lactocepins (LacCeps).

A total of 5740 peptides were produced by hydrolysis of β-casein during ripening (Supplementary Data Table 4). Peptides derived from β-casein found only at depth 0.410 mm were produced from regions 130-161 and those produced only at depth 9.02 mm and 20.05 mm were from regions 39-52 and 69-119. Peptides commonly produced at all depths were found at region 1-6, 164-190 and 193-207 (Fig. 5 C). Peptide fragments derived from action of only plasmin on N- and C-termini were β-CN (f22-28), β-CN (f94/97-105), β-CN (f106-119) and β-CN (f129-137) with action of proteinases from the smear microbiota. Similarly, peptide fragments produced by the action of only lactocepins were β-CN (f1-
during the study at depths 0.41 mm, 9.02 mm and 20.05 mm.

- Peptides from β-casein present only at the surface were β-CN (f 1-14), β-CN (f 53/57-68), β-CN (f 94-124/128), β-CN (f 95-125/126/127/128/129), β-CN (f 129/130/132-137/139/142), β-CN (f 132/142-161) and β-CN (f 144-162/163/189) and those present only at the center were, β-CN (f 22/23-28), β-CN (f 25-33), β-CN (f 70-77), β-CN (f 74-86), β-CN (f 78/80/84-91), β-CN (f 94-102/105), β-CN (f 95/95-119) and β-CN (f 201-204). Peptides from α-casein present at all the depths were β-CN (f 22-28), β-CN (f 166/176-182), β-CN (f 166/168/192) and β-CN (f 166/191/192-209).

- Most common peptides from β-casein present only at surface were β-CN (f 1-14), β-CN (f 53/57-68), β-CN (f 94-124/128), β-CN (f 95-125/126/127/128/129), β-CN (f 129/130/132-137/139/142), β-CN (f 132/142-161) and β-CN (f 144-162/163/189) and those present only at the center were, β-CN (f 22/23-28), β-CN (f 25-33), β-CN (f 70-77), β-CN (f 74-86), β-CN (f 78/80/84-91), β-CN (f 94-102/105), β-CN (f 95/95-119) and β-CN (f 201-204). Peptides from α-casein present at all the depths were β-CN (f 22-28), β-CN (f 166/176-182), β-CN (f 166/168/192) and β-CN (f 166/191/192-209).

A small number of peptides from κ-casein was found at low concentrations during ripening (Fig. 5 D). Fragments produced from pH 4.6-soluble extract, by action of lactocepins at their N- or C- termini (ARDO et al., 2017) were κ-CN (f 33-41), κ-CN (f 78-85) and κ-CN (f 96-103/105). κ-CN (f 96-105) was produced by chymosin cleaving at its C-terminus (ARDO et al., 2017). More detailed study on enzyme specificities of the smear ripened cheese can be a future area of research.

4. CONCLUSION

The gross composition of smear ripened cheese was observed to be typical of the cheese variety. Overall % pH 4.6 SN/TN in all three batches G1, G2 and G3, indicated a normal proteolysis at the end of the ripening in the cheese. Extent of proteolysis was observed by urea-PAGE which showed extensive breakdown of α- and β-casein essentially from surface to 1.23 mm, followed by normal patterns of proteolysis typical of bacterially ripened cheeses, towards the centre. Free amino acid analysis of surface layers of the cheese showed high levels of glutamic acid, glycine, alanine, tyrosine, and proline, whereas higher concentrations of threonine, lysine, phenylalanine and valine were found towards the inner layers of smear-ripened cheese. A total of 5614 peptides were identified during the study at depths 0.41 mm, 9.02 mm and 20.05 mm.

Most common peptides from α-casein present only at surface were α-CN (f 122-139/142/143), α-CN (f 124-142/143/144), α-CN (f 125-144) and α-CN (f 150-155/156/157) and those present only at the center were α-CN (f 191/193-199). Most common peptides from α-casein present only at surface were α-CN (f 68-74/88/94), α-CN (f 71-94/98), α-CN (f 73-86/88), α-CN (f 75-88/94) and α-CN (f 87-95) and those present only at the center were, α-CN (f 176-181/182), α-CN (f 186-197), α-CN (f 188-199/202), α-CN (f 191-197/199/203) and α-CN (f 194-203). Most common peptides from β-casein present only at surface were β-CN (f 1-14), β-CN (f 53/57-68), β-CN (f 94-124/128), β-CN (f 95-125/126/127/128/129), β-CN (f 129/130/132-137/139/142), β-CN (f 132/142-161) and β-CN (f 144-162/163/189) and those present only at the center were, β-CN (f 22/23-28), β-CN (f 25-33), β-CN (f 70-77), β-CN (f 74-86), β-CN (f 78/80/84-91), β-CN (f 94-102/105), β-CN (f 95/95-119) and β-CN (f 201-204).

Common peptides from α-casein at all the depths were α-CN (f 11-23), α-CN (14-23), α-CN (f 24-34) and α-CN (131-139/142/148); from α-casein at all the depths were α-CN (f 115-125), α-CN (f 151-156/158/161/162/163) and α-CN (f 196-208); β-casein at all the depths were β-CN (f 22-28), β-CN (f 166/176-182), β-CN (f 166/168/192) and β-CN (f 166/191/192-209).
(f166/191/192-209) and those from κ-casein were κ-CN (f33-41), κ-CN (f78-85) and κ-CN (f96-103/105). κ-CN (f96-105).
Study of individual proteolytic enzymes and their effect on the caseins of smear ripened cheese and detailed study of microbiological aspects will be an interesting area for further research.

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REFERENCES


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