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Protein profile of dairy products: Simultaneous quantification of twenty bovine milk proteins

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1	Protein profile of dairy products: Simultaneous quantification of twenty bovine milk
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### 24

### 25 ABSTRACT

26

While proteomic techniques allow the identification and relative quantification of thousands of 27 proteins in a single run, methods for absolute quantification remain laborious. In this study, a 28 newly developed multiple reaction monitoring (MRM) method using liquid chromatography 29 30 mass spectrometry (LC-MS) that enables the simultaneous quantification of twenty key milk proteins is presented. The selected proteins comprise all individual caseins, the major whey 31 proteins and most well-known milk fat globule membrane (MFGM) proteins. For validation, 32 the twenty milk proteins in raw milk, raw cream, raw milk Emmental cheese and whey, were 33 quantified as well as in eighteen commercial heat-treated dairy products. The method 34 presented is ideally suited for various applications, for example, the comparison of the 35 protein patterns in raw milk of cows at different stages of lactation or of different breeds. 36 37

#### 38 1. Introduction

39

Proteins form a major class of milk components, comprising over 400 different types 40 41 (Lu, 2013). They are present over a broad concentration range and can be grouped into three main classes (Casado, Affolter, & Kussmann, 2009): (i) the casein micelle proteins 42 (CasMPs, 80–85%), organised as supramolecular, dynamic structures called casein micelles, 43 that entrap colloidal calcium phosphate (McMahon & Oommen, 2013); (ii) the whey proteins 44 45 (WPs, 13–18%), dissolved in the water phase; and (iii) proteins which are associated within the milk fat globule membrane (MFGMPs, 1–2%), a phospholipid bilayer that embeds 46 47 proteins that protects the fat globules from coalescence and lipolysis (Bauman, Mather, Wall, & Lock, 2006; Dewettinck et al., 2008). 48

Milk proteins are of high value from a technological point of view as well as for their 49 beneficial physiological effects (Supplementary material, Table S1). Therefore, the analysis 50 of the different individual protein profiles in milk and dairy products is of high interest, not only 51 52 for the dairy industry but also for nutritional research as well as applied biotechnology. 53 Typical methods for the quantification of individual proteins involve ELISA or Western blot techniques. However, these methods require the availability of specific antibodies and a 54 significant amount of time and effort, as the possibility of multiplex assays is restricted. 55 Other techniques utilise high-performance liquid chromatography (HPLC; Schwendel et al., 56 57 2017) or two-dimensional gel electrophoresis with the subsequent application of 58 densitometry, dyes, fluorophores or radioactivity (Turner, MacDonald, Back, & Thomson, 2006) to attain the necessary sensitivity and resolution for protein quantification. However, 59 due to a lack of individual certified milk protein standards, these quantifications are rarely 60 61 absolute and proteins, which are insoluble or present in low concentrations are not detected. Recently, a few laboratories developed multiple reaction monitoring (MRM) methods 62 using liquid chromatography mass spectrometry (LC-MS) for the quantification of major milk 63 proteins such as  $\beta$ -lactoglobulin and caseins (Lutter, Parisod, & Weymuth, 2011) or certain 64 individual MFGMPs (Affolter, Grass, Vanrobaeys, Casado, & Kussmann, 2010; Fong & 65

Norris, 2009). By targeted fragmentation of sample derived peptides as well as selected signature peptides and subsequent monitoring of their specific ions simultaneously, the MRM procedure enables the quantification of sample peptides based on their corresponding signal intensities. Therefore, MRM methods are very sensitive and increase selectivity to a level required when complex mixtures such as food matrices are analysed (Lange, Picotti, Domon, & Aebersold, 2008).

72 Up to now, MRM methods were developed covering only a few major milk proteins 73 (Le, Deeth, & Larsen, 2017). To the best of our knowledge, no method is currently available for the absolute and simultaneous quantification of minor as well as the most abundant 74 bovine milk proteins. Therefore, we developed a MRM method for the individual 75 76 quantification of twenty key milk proteins at once by applying the absolute quantification (AQUA) strategy (Kirkpatrick, Gerber, & Gygi, 2005). This new method allows the 77 simultaneous quantification of the CasMPs ( $\alpha_{S1}$ -,  $\alpha_{S2}$ -,  $\beta$ - and  $\kappa$ -casein) and the casein-78 79 associated lipoprotein lipase, six key WPs ( $\beta$ -lactoglobulin,  $\alpha$ -lactalbumin, proteose peptone 3, serum albumin, lactoperoxidase and lactoferrin), and the most abundant MFGMPs 80 81 (butyrophilin, xanthine dehydrogenase/oxidase, adipophilin, lactadherin, platelet glycoprotein 82 4 (CD36), polymeric immunoglobulin receptor, fatty acid binding protein, fatty acid synthase and glycoprotein 2). Designations, abbreviations, and most known functions of those milk 83 proteins are listed in Supplementary material, Table S1. 84

85 The MRM method developed was validated by quantifying the twenty proteins in dairy 86 products produced from raw milk and in commercial heat-treated dairy products. The dairy products raw milk (RM), raw cream (RC), Swiss Emmental (EM)—a raw milk cheese, and 87 sweet whey (W) are particularly suitable for the evaluation of the method, since they 88 89 represent all three different milk fractions, thus having different protein profiles. The investigated commercial dairy products comprise pasteurised and ultra-pasteurised milk (MI), 90 ultra-high temperature (UHT) and ultra-pasteurised cream (CR), yoghurt (pasteurised, YOG), 91 92 buttermilk (pasteurised, BM), quark (pasteurised, Q) and cottage cheese (pasteurised,

93 COTC), each from three different manufacturers are listed, including the specifications of the
94 indicated preservation processes (Supplementary material, Table S4).
95
96 2. Materials and methods
97
98 2.1. Materials

99

100 RM was obtained from the cheese dairy Uettligen (Bern, Switzerland). The RC was separated by centrifugation from the RM at 10 °C and 2000  $\times$  g for 15 min. The W was 101 collected after the production of Tilsit cheese (from milk that was heated at 44.5 °C) at 102 Agroscope (Bern, Switzerland). The commercial dairy products were bought from the 103 supermarkets Migros and Coop (Switzerland, Supplementary material, Table S4). LC-MS 104 grade water was purchased from VWR International (Dietikon, Switzerland); LC-MS 105 hypergrade acetonitrile (ACN), formic acid (FA), ammonium bicarbonate (ABC) and the 106 107 reference proteins used for spiking [ $\alpha$ -casein ( $\alpha_{s1}$  +  $\alpha_{s2}$ -casein),  $\beta$ -lactoglobulin,  $\alpha$ -108 lactalbumin, serum albumin, lactoferrin] from Merck (Zug, Switzerland); and trypsin Gold (MS-Grade) from Promega (Dübendorf, Switzerland). 109

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111 2.2. Sample preparation for sodium dodecyl sulphate polyacrylamide gel electrophoresis
112 (SDS-PAGE)

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The total protein content of each dairy product was calculated from the total nitrogen (TN) content (raw milk dairy products) or from the TN minus the non-protein nitrogen (NPN) (heat-treated dairy products) determined by Kjeldahl according to the ISO/IDF standard method ISO 8968-3:2007/IDF 20-3:2007 (ISO, 2007) and multiplied by a conversion factor of 6.38 (Supplementary material, Table S4). A precise quantity of dairy product containing 2 mg of total proteins was placed in a 1.5 mL Eppendorf tube and precipitated with 1 mL of icecold acetone. The resulting pellets were dissolved in 200 µL Tris-HCI (100 mmol L<sup>-1</sup>, pH 7.5,

121	1% sodium dodecyl sulphate (SDS)); 100 $\mu L$ of the sample (clear phase) was mixed with 20
122	$\mu$ L of the sample buffer 6× (Tris-HCl 350 mmol L <sup>-1</sup> , pH 6.8, SDS 10%, glycerol 50%,
123	Dithiothreitol (DTT) 100 mmol $L^{-1}$ ) with bromophenol blue and heated at 95 °C for 5 min.
124	Then, 3.6 $\mu$ L (30 $\mu$ g of protein) of each sample and 5 $\mu$ L of the molecular weight marker
125	(Benchmark <sup>™</sup> Prestained Protein Ladder; Thermo Fisher Scientific, Reinach, Switzerland)
126	were separated by SDS-PAGE (15% polyacrylamide) and stained with colloidal Coomassie
127	Blue, as previously described (Egger et al., 2016).
128	
129	2.3. In-gel tryptic digestion
130	
131	Pieces of polyacrylamide gel containing protein bands of interest were excised from
132	gels (15% polyacrylamide), washed and digested with trypsin as described in Kopf-Bolanz et
133	al. (2012).
134	
135	2.4. Isotopically labelled peptides as internal standards
136	
137	The twenty isotopically labelled AQUA peptides were manufactured by Thermo Fishe
138	GmbH (Ulm, Germany) according to the provided sequences (Table 1). The last AA of the
139	tryptic peptides was labelled with <sup>13</sup> C and <sup>15</sup> N, thereby producing a mass shift between the
140	AQUA and the native peptides of +10 for arginine, +8 for lysine and +6 for valine. The
141	labelled AQUA peptides were produced as lyophilised trifluoroacetic salts, which were
142	dissolved in sample solution (5% ACN, 0.1% FA in water), thereby resulting in a
143	concentration of approximately 50 pmol $\mu L^{-1}$ . Following the accurate determination of the
144	soluble concentrations by AA analyses (phenylthiocarbamyl (PTC)-derivatisation, as
145	described by Kopf-Bolanz et al., 2012), a specific quantity of each AQUA peptide dilution was
146	mixed to produce a peptide-mix-solution containing the labelled peptides of CASA1, CASA2,
147	CASB, CASK, LACB, LALBA, FABP, PAS 6/7 and PIGR at a concentration of 0.2 pmol $\mu L^{\text{-1}}$
148	and the labelled peptides of LPL, BSA, LPO, LF, BTN, XDH, ADPH, CD36, PP3, FAS and

GP2 at a concentration of 0.02 pmol  $\mu$ L<sup>-1</sup>. These concentration ratios have proven to be favourable, since the signals of the native peptides in milk and the AQUA peptides differed by no more than a factor of 10, if 5  $\mu$ L of the AQUA peptide-mix-solution was injected simultaneously with each sample. A twenty-fold concentrated solution of the AQUA peptidemix was stored at –80 °C for up to a maximum of 8 weeks.

154

#### 155 2.5. Sample preparation for protein quantification

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The total protein content of each sample was determined as described above 157 (Supplementary material, Table S4). The EM was grated, and subsequently, precisely 2 mg 158 of total protein (RM, RC, W, EM, YOG, Q, and COTC), or 60 µL of product (MI, CR and BM) 159 respectively, were added into a 1.5 mL Eppendorf tube. For EM and RC, 50 µL of digestion 160 buffer (10% ACN, 25 mM ABC in micro filtered H<sub>2</sub>O) were added and placed in a sonication 161 bath for 30 min, helping the matrix to dissolve and liberate the proteins. The proteins were 162 163 precipitated by an addition of 1 mL of ice-cold acetone and the tube was let on ice for 1 h. The suspensions were centrifuged at 4 °C and 18,000  $\times$  g for 20 min, the acetone was 164 carefully discarded, and the resulting pellets were air-dried for at least 30 min. The pellets 165 166 were resolubilised in 1 mL of digestion buffer by vortex and sonication in a warm water bath (40 °C). Fifty  $\mu$ L of protein solution (2  $\mu$ g  $\mu$ L<sup>-1</sup>) were mixed with 40  $\mu$ L of digestion buffer 167 followed by addition of 10  $\mu$ L of trypsin solution (0.2  $\mu$ g  $\mu$ L<sup>-1</sup> in 0.1% Tris, pH 9, specific 168 activity >15,000 u mg<sup>-1</sup>). After overnight incubation at 37 °C, 10 µL of the digested protein 169 solution were diluted in 990 µL of sample solution (5% ACN, 0.1% FA in water). For the 170 quantification of the high concentrated proteins in milk (LACB, LALBA, PP3, CASA2, CASK, 171 172 CASA1, CASB), 10 µL of this (100× diluted) digest solution was injected into the LC-MS, corresponding to 0.1 µg of total proteins. For the guantification of the lower concentrated 173 proteins (FAS, TRFL, XDH, BTN, PIGR, LIPL, ADPH, FABP3, GP2, CD36, PAS6/7, BSA and 174 PERL), 10 µL of the undiluted digest solution was injected (corresponding to 10 µg of total 175 protein). With each sample-injection, 5 µL of the internal standard peptide-mix-solution (see 176

above) were simultaneously injected. To minimise the experimental error, exactly the same 177 sample was prepared and analysed multiple times (technical replicates) and each replicate 178 179 was measured three times.

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2.6. Analysis by liquid chromatography tandem mass spectrometry 181

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Peptides were separated on a Rheos 2200 HPLC (Flux Instruments, Reinach, 183 184 Switzerland) equipped with a XTerra MS C18 column (3.5 µm, 1.0 mm × 100 mm) and a guard column (XTerra MS C18 VanGuard Cartridge, 3.5 µm, 2.1 mm × 10 mm; both 185 columns: Waters, Baden-Dättwil, Switzerland) with a flow rate of 80 µL min<sup>-1</sup> for 30 min. A 186 gradient from 5%–60% solution B (ACN, 0.1% FA) in solution A (H<sub>2</sub>O, 0.1% FA) was applied 187 in the first 15 min, increased to 95% in the next 5 min, and returned to the initial conditions 188 within the 21<sup>st</sup> min for a 9-min re-equilibration. The column temperature was maintained at 25 189 °C. 190

191 The Rheos 2200 HPLC was coupled directly to a LTQ linear ion trap mass spectrometer (QQQ-MS, Thermo Scientific, Reinach, Switzerland) using an electron spray 192 ionisation (ESI) interface. The HPLC eluent of the first 3.5 min and the last 17 min were 193 194 diverted to waste. ESI conditions were as follows: source voltage 4000 V, capillary voltage 5 V, tube lens 150 V, capillary temperature 275 °C, sheath gas flow 20 arbitrary units and 195 auxiliary gas flow 10 arbitrary units. The ion trap mass spectrometer was operated in a 196 197 positive ion mode. The MRM included liquid chromatography-tandem mass spectrometry (LC-MS/MS) runs with 2-4 segments (1-8 min) and 2-6 scan events. The following MRM 198 conditions were included: full scan range 260–1500 m/z, isolation width 2 m/z, normalised 199 200 collision energies 35.0, collision gas helium, activation time 30 ms and activation Q 0.250. The retention time (r.t.) of the signature peptides, peptides m/z and transitions, as 201 shown in Table 1. The resulting peaks were integrated using the quantitative software 202 LCquan (version 2.8) from Thermo Scientific. The performance of the LC-MS/MS-system 203 was reviewed before and in between each batch of measurements by injection of a BSA

solution (10  $\mu$ L, final concentration 10 fmol  $\mu$ L<sup>-1</sup>, tryptic digested peptides of BSA), separated 205 with the identical gradient and flow rate on the same XTerra MS C18 column/guard column-206 207 system as peptide separation of the samples occurred. The ESI settings were the same as 208 for the peptides samples. The MS setup was a full-scan range from 300 to 1100 m/z in one segment with a start delay of 1.9 min. The MS settings included isolation width 1 m/z, 209 210 normalised collision energy 35.0, collision gas helium, activation Q 0.250 and activation time 30 ms. The measurements were conducted in the positive ion mode and the resulting 211 212 spectra were evaluated by an identification search with Mascot v 2.2.04 (Matrix Science Inc., Boston, MA) using the UniProt Database (search parameter settings: MS/MS ion search; 213 trypsinisation; variable modifications: deamination, pyroglutamic acid, oxidation; average 214 mass values; unrestricted protein mass; mass and fragment mass tolerance: ± 0.8 Da; 215 maximum missed cleavage: 1; instrument type: ESI-TRAP). The performance was assessed 216 by means of a BSA sequence-coverage of at least 15%. 217

218

#### 219 2.7. Recovery experiments

220

221 For recovery experiments, the protein content of six commercially available standard proteins (CASA1, CASA2, LACB, LALBA, BSA and LF) was determined by measuring TN 222 and NPN with Kjeldahl (ISO, 2007). A solution containing a determined amount of the 223 standards in digestion buffer was prepared. The concentration of each of the proteins was 224 quantified in the solution (Supplementary material, Table S3), as well as in 60 µL milk using 225 226 the developed MRM method. Increasing quantities of the prepared protein solution were added to the basis of 60 µL of milk (0-50 µL in 10 µL steps, corresponding to 0-5 aliguots in 227 Supplementary material, Fig. S3). The proteins of the gradually spiked samples were 228 229 quantified with the MRM method. For each of the six proteins, the recovery was calculated by 230 the ratio of the added amount of protein and the quantity determined by MRM in the spiked 231 sample after subtraction of the amount found in the milk.

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233

#### 3. **Results and discussion**

- Establishment of a multiple reaction monitoring (MRM) method for the simultaneous 3.1. 235 quantification of twenty bovine milk proteins 236
- 237

To define the quantifiable protein set, the proteins of the four raw milk dairy products 238 RM, RC, EM and W were separated by SDS-PAGE (Fig. 1b). Polyacrylamide gel pieces 239 were manually excised from the most intense bands and prepared for in-gel tryptic digestion. 240 241 The proteins were identified by mass spectrometry based on matching MS/MS spectra with in-silico generated spectra using the MASCOT database (Kopf-Bolanz et al., 2012). Thus, 242 MS data for the most abundant milk proteins were obtained, which included the r.t. of the 243 tryptic peptides, mass-to-charge ratios and their MS/MS features. The twenty milk proteins 244 245 were selected for quantification, due to their technological importance, biological function or nutritional value. The selected proteins include all CasMPs (CASA1, CASA2, CASB, CASK, 246 LPL), six major WPs (LACB, LALBA, PP3, BSA, LPO, LF) and a selection of MFGMPs (BTN, 247 XDH, ADPH, PAS 6/7, CD36, PIGR, FABP, FAS, GP2) (Supplementary material, Table S1). 248 The peptide data obtained by protein identification provided the necessary information 249 for selecting proteotypic signature peptides suitable for absolute quantification by MRM. The 250 selection of the signature peptides was based on the uniqueness of the tryptic peptide 251 sequence, the absence of reported post-translational modification sites (UniProt Database), 252 and their ionisation efficiency. Furthermore, the selection was restricted to tryptic peptides up 253 to triply charged ones with a mass-to-charge-ratio between 350 to 1000 m/z. In the first step, 254 255 for each protein, three highly detectable peptides that met these conditions were selected 256 and their suitability for quantification by MRM was tested. In a second step, out of the three 257 peptides, one signature peptide was selected for each protein on the basis of peak shape, r.t. and resolution. To increase the specificity for each signature peptide, the two most intense 258 transitions with the best resolution were chosen for monitoring and subsequent quantification 259 with the LCquan software (Thermo Scientific). The sequence of the selected signature 260

261 peptides as well as their corresponding r.t. and the specific m/z value for the peptides and 262 the fragments are listed in Table 1.

263

264 3.2. Method validation

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266 3.2.1. Specificity and selectivity

To assure specificity and selectivity of the developed MRM method, particular 267 attention was given to the selection of the proteotypic peptides and transitions to ensure that 268 269 there was no extended interaction with the matrix. The use of two transitions with at least ten data points per peak for the identification and quantification of each protein increased the 270 specificity of the method. Moreover, the labelled peptides were spiked as internal standards 271 into every single experiment just before co-injection to circumvent ion suppression arising 272 from the interaction with the sample matrix. Time segmentation was applied to optimize dwell 273 time and S/N ratio for predefined sets of transitions, thereby resulting in improved sensitivity 274 275 with minimum length of the MRM method. As the extracted ion chromatograms in 276 Supplementary material, Fig. S2 indicate all peaks were well separated and were therefore 277 easy to integrate. The displayed data was extracted directly from MRM experiments of commercially available buttermilk and cream samples. 278

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280 3.2.2. Precision

To review the precision of the quantification method, the coefficients of variation (CV) 281 were determined for each protein, measuring a RM sample in six biological replicates, each 282 283 injected three times (eighteen measurements). The CVs for CasMPs and WPs ranged between 5% and 15% (Supplementary material, Table S2), which are typical for such 284 methods (Brönstrup, 2004; Yang et al., 2007). Higher CVs were obtained for a few MFGMPs 285 (ADPH, PAS 6/7, CD36, PIGR, 7 to 33%). This is most likely due to the low concentration of 286 287 these proteins in milk. The quantification area of these minor proteins was close to the quantitation limit, thereby increasing the error. Due to the much higher concentrations of 288

CasMPs in milk, it was not possible to increase the injection volume to improve quantification
quality of those MFGMPs. As most MFGMPs are more concentrated in cream, the CV
declines for these minor proteins, getting into the typical range of CV around 15% (data not
shown).

293

294 3.2.3. Accuracy and recovery

Recovery experiments were conducted for six selected commercially available 295 296 proteins (CASA1, CASA2, LACB, LALBA, BSA and LF) through a gradual increase in concentration of the spiked protein in a milk sample (section 2.7). The determined recovery 297 values of the proteins in milk and the corresponding linear regression curves are plotted in 298 Supplementary material, Fig. S3. The recovery rate ranged from 91% to 105% 299 (Supplementary material, Table S3). The CVs of the recovery rates varied for the high 300 abundant CasMPs between 6.6% (CASA1) and 7.9% (CASA2), for the lower abundance 301 proteins between 9.8% and 26.7% (LF, BSA, LALBA and LACB). The higher variability could 302 303 be explained by difficulties to obtain a homogenous mixture of the spiked protein with the raw 304 milk sample.

305

#### 306 3.2.4. Range and linearity

The linearity of the labelled AQUA peptide quantification was determined over at least 307 four orders of magnitude  $(10^{-3}-10 \text{ pmol})$ , and the linear regression curves found had an R<sup>2</sup> of 308 at least 0.96 for each single peptide (Supplementary material, Fig. S4). The upper limit of 309 linearity was not determined due to the high cost and limited quantity of the labelled AQUA 310 peptides. The linearity in the response of the native peptides was determined on an RM 311 312 sample. Linearity is demonstrated for the most abundant milk proteins (all CasMPs, LACB, 313 PP3; Supplementary material, Fig. S1). For minor milk proteins, it was not possible to obtain sufficient data points, since the injection of higher amounts of protein led to a saturation of 314 the column with CasMPs. 315

316

317 3.2.5. Detection and quantitation limits

318	As the twenty proteins quantified in this method are present in milk in a broad
319	concentration range, the limit of detection and quantitation were different for each individual
320	protein. For the most abundant proteins, a detection and quantitation limit of 0.0001 $\mu g$ was
321	found, which corresponded to 0.001 $\mu$ g of total injected protein. For the low concentration
322	proteins, the detection limits ranged between 0.1 and 1 $\mu$ g of total injected protein and the
323	quantitation limits between 1 and 10 $\mu$ g of total injected proteins (Supplementary material,
324	Table S2). For the AQUA peptides, the limits of detection and quantitation were 0.0001 pmol
325	and 0.001 pmol, respectively. Hence, to achieve a precise quantification for each protein, the
326	same samples were injected twice with two different amounts of total proteins in the column
327	(0.01 ug for LACB, LALBA, PP3, CASA2, CASK, CASA1, CASB and 10 ug for FAS, TRFL,
328	XDH, BTN, LALBA, PIGR, LIPL, ADPH, FABP3, GP2, CD36, PAS6/7, BSA and PERL).
329	
330	3.3. Simultaneous quantification of twenty bovine milk proteins in raw milk and
331	commercial heat-treated dairy products
332	
333	3.3.1. Raw milk dairy products
334	To investigate the quantification method in practice, the amounts of the twenty
335	proteins were determined in the four raw milk dairy products already used for the method
336	development. RM, RC, EM and W represent the different fractions of milk and, thus, are rich

337 in milk proteins belonging to different classes. The results for the major CasMPs and WPs in

338 RM measured with the MRM method were compared with previously obtained results from

339 literature (Table 3), for which however the methods used were not always clearly described.

- 340 Moreover, the concentration of proteins in milk depends on many factors and differs
- according to the course of lactation, udder health, supply of energy and crude protein, feed,
- 342 season, environmental temperature and breed. Therefore, concentrations of the different milk
- proteins are often indicated as a range (Eigel et al., 1984; Swaisgood, 1993; Table 3).

The side-by-side comparison showed that the results for seven of the eight proteins 344 (CASA1, CASA2, CASB, CASK, LACB, LALBA, BSA and LF) were in the expected range 345 346 (Table 3). Only the amount of BSA detected in RM deviated from the published values. 347 However, this deviation was not observed in heat-treated samples (Table 4). The reasons for the deviation found in RM could lie in the nature of the protein, as BSA has several lipid-348 binding sites (Spector, John, & Fletcher, 1969) as well as 35 cysteines, of which only one 349 sulfhydryl group is free (Chevalier, Hirtz, Sommerer, & Kelly, 2009). It is therefore possible 350 351 that the tight folding of the native BSA, held together by the disulphide bridges, makes it more resistant to tryptic hydrolysis. Since there is also evidence that BSA in its native form is 352 involved in protein complexes in non-heat-treated skim milk (Chevalier et al., 2009), a lower 353 susceptibility of the protein to precipitation and tryptic digestion may also result from a 354 possible interaction of BSA with lipids or other native proteins present in the RM. 355

To the best of our knowledge, no literature data in RM are available for LPL, PP3 and 356 LPO as well as all investigated MFGMPs (BTN, XDH, ADPH, PAS 6/7, CD36, PIGR, FABP, 357 358 FAS, GP2). The milk proteins in the four products analysed, RM, RC, EM and W, were attributed to the three classes, CasMPs, WPs and MFGMPs and given in g 100 g<sup>-1</sup> of total 359 protein (Fig. 1a) and the individual concentrations are listed in Table 2. As displayed in Fig. 360 1, the protein profiles of the four products correspond well with the expected class of proteins 361 on the basis of their manufacturing technology. RM comprised 80.3% CasMPs, 13.5% WPs, 362 363 and 1.8% MFGMPs, which is in good accordance with the existing literature (Fox, 2011). RC 364 had a higher proportion of MFGMPs (4.8%), due to the high content of native fat globules and contained 64.7% CasMPs, 16.2% WPs. EM contained mainly CasMPs (100.3%) and 365 only minor amounts of the other classes of proteins, namely 0.9% WPs and 0.5% MFGMPs. 366 367 And as expected, whey contained mostly WPs (84%) and only 1.6% CasMPs and 1.5% MFGMPs (Fig. 1a). The protein distribution was qualitatively confirmed by SDS-PAGE (Fig. 368 1b). 369

370

#### 371 3.3.2. Commercial heat-treated dairy products

In addition to the four raw milk products, the protein profiles of eighteen commercially 372 available heat-treated dairy products with different fat contents were examined, including MI 373 374 3.5–3.9% fat, CR 35% fat, YOG 3.5–4% fat, BM 0.5% fat, Q 0.1% fat, and COTC 4–4.5% fat, 375 each product obtained from three different manufacturers. The average total amount of CasMPs, WPs and MFGMPs was calculated by the sum of the individual proteins belonging 376 377 to these protein classes and plotted for each type of product (Fig. 2a). The deviations and the 378 fact that total protein amounts that were below the expected 100% can be explained by the 379 different origin of the products, and are therefore no measure for the quality of the experiments. A representative example for the individual protein pattern of each dairy 380 product is shown in the SDS-PAGE in Fig. 2b. 381

382

383 3.3.3. Comparison of the protein concentrations found in raw and commercial heat-treated
384 dairy products

A one-to-one comparison between the determined protein concentrations in the raw 385 386 milk products and those found for the heat-treated dairy products might be tempting, but is 387 not fully appropriate, since the investigated products are purchased at different times from different suppliers (Supplementary material, Table S4). Hence, the milk used for these 388 products originates from different animals, possibly of different breeds, which were most 389 likely at different stages of the lactation cycle and receiving different feeds with varying 390 391 energy levels and crude protein content. Moreover, the milk was processed by different 392 manufacturers and has thus undergone a variety of different processing steps besides heattreatment. Nevertheless, taking these restrictions into account, it is interesting to consider a 393 brief and cautious comparison of the measured values. 394

Remarkably, in comparison with the RM and the raw milk dairy products, the content of the twenty proteins found in the heat-treated dairy products were lower, with some exceptions. As indicated before, the most obvious explanation are differences in the concentration of these proteins in the milk of origin and that processing and storage might affect protein conformation and stability and possibly lead to protein degradation (Deeth &

Lewis, 2017; García-Risco, Ramos, & López-Fandiño, 2002). Another explanation is the 400 potential emergence of glycated lysyl residues due to the presence of reducing 401 402 carbohydrates during heat-treatment and storage as a first step of the Maillard reaction 403 (Metha & Deeth, 2015). As a result, peptide bonds might be blocked for cleavage by trypsin, 404 therefore preventing a precise quantification by MRM, leading to a substantial decrease in concentration of individual proteins in dairy products that have undergone a more severe 405 thermal treatment. However, the determined quantities for the individual proteins in 406 407 pasteurised milk compared with high-pasteurised milk as well as in high-pasteurised cream, compared with UHT cream, do not support this hypothesis (Supplementary material, Fig. S5). 408 Furthermore, the observed reduction in quantity was not higher for the proteins when the 409 selected signature peptides ended with a c-terminal lysyl residue compared with the ones 410 with a c-terminal arginyl residue (Table 1). Nevertheless, the heat-treated dairy products of 411 one kind shared a similar protein pattern, while the concentration of specific proteins varied 412 in the products between the different manufacturers (Supplementary material, Fig. S5). Since 413 414 RM is used as a starting material in the manufacturing process for all of these dairy products, 415 the amount of individual proteins in the RM could be decisive for the quality of the product.

416

Casein micelle proteins. In the four raw milk products analysed, RM, RC, EM and W, 417 amounts of the individual CasMPs were in accordance with previous results (Tables 3 and 418 419 4). In RC, CasMPs had a lower concentration but the same ratio compared with RM. In EM 420 however, the ratio of CasMPs was different from RM. CASA1, CASB and CASK were higher than CASA2 and LPL. LPL, which is loosely attached to the casein micelles, is most probably 421 partially released into whey during cheese manufacturing (Table 2). In the case of CASA2, 422 423 possible hydrolysis of the indicator peptide by bacteria during cheese ripening might explain the finding. As expected, in W, only traces of CasMPs were found. Surprisingly, with the 424 exception of the COTC samples, the total amount of CasMPs in heat-treated dairy products 425 was lower than that in the RM, possibly due to a lower concentration of CasMPs in the milk 426 of origin and the degradation during heating, processing and storage (Meltretter, Schmidt, 427

Humeny, Becker, & Pischetsrieder, 2008). In fermented dairy products such as BM, YOG
and Q, a possible hydrolysis of the indicator peptides by the added starter cultures could also
contribute to the lower protein concentrations. The quantities of the individual CasMPs in the
heat-treated samples did not follow a clear pattern that could be explained by the
technological transformations and no clear tendency could be deduced.

433

Whey proteins. In line with expectations, the highest concentration of all serum 434 proteins was found in W, while most of them were absent in the EM (Table 2). PP3 were 435 found to be slightly more concentrated in RC than in RM and MI (Table 4), thereby 436 suggesting that it might be partially associated with the MFGM (Table 2), as previously 437 described by Dewettinck et al. (2008). This is also supported by the results for PP3 in CR 438 and BM and the observed low concentration in Q (Table 4). Remarkably, in the experiments 439 reported here, LF behaved neither as a real WP nor a real MFGMP, since it was found in 440 similar concentrations in all four milk fractions (RM, RC, EM and W) and was only slightly 441 442 increased in BM and YOG. In the literature, the localisation of LF is controversially 443 discussed: some publications associate LF with the MFGM and some with the milk serum fraction (Casado et al., 2009). LPO is present in all investigated products at similar levels 444 445 with a higher concentration in W. Its activity is used as an indicator for high temperature treatment, for example, for ultra-pasteurisation (Fox & Kelly, 2006). With the exception of W. 446 BSA levels detected in raw milk products were below the expected values reported from 447 448 literature (Table 3). In contrast to that, the values found in heat-treated samples (MI, BM, CR, YOG and Q) were higher and matched the expected values. As mentioned before, the 449 reason for this finding might lie in the tight folding of the protein or a possible interaction of 450 451 the protein with lipids or other proteins present in raw milk, thereby interfering with the precipitation and hydrolysis by trypsin. 452

453

454 *Milk fat globule membrane proteins.* During butter manufacture, an important part of 455 the MFGM is released into the serum phase. Therefore, it is not surprising that BM contains

the highest concentration of MFGMPs of all investigated dairy products (Table 4, Fig. 2). As
the higher fat content of cream suggests, MFGMPs were also found in high concentrations in
RC and CR (35%) (Fig. 1a, Table 2 and Fig. 2, Table 4, respectively). However, thus far, only
a few methods were available for the quantification of MFGMPs and thus the quantity of the
individual MFGMPs in different milk products is mostly unknown in literature.

In the products investigated here, the most prominent MFGMPs were BTN, XDH and 461 PAS 6/7. BTN is a protein involved in the formation and stabilisation of the MFGM (Robenek 462 463 et al., 2006) and had highest levels in RC and BM. However, XDH, a protein that amplifies the antibacterial effect of LPO, and PAS 6/7, which is known to have anti-infectious 464 properties (Fox & Kelly, 2006; Mather, 2000), were most prominent in BM and CR. Moreover, 465 BTN and XDH varied the most in the different CR samples. A possible reason for this 466 variance might be the fact that BTN and XDH form a high molecular weight aggregate 467 induced by heat-treatment (Ye, Singh, Taylor, & Anema, 2002). The content of PAS 6/7 468 differed the most between the BM samples of different manufacturers (Supplementary 469 470 material, Fig. S5). Since, during the process of homogenisation or butter making, the MFGMs 471 are destroyed and not entirely rearranged around the newly formed droplets, the dissolved proteins transferred to the serum are more vulnerable to degradation by enzymes, 472 473 microorganisms and heat. This indicates that the observed high variance of MFGMPs in 474 commercial heat-treated products might not only be attributed to different manufacturers and milk origin. Remarkably, the concentration of CD36, FABP, FAS and GP2 was comparatively 475 476 high in W (Table 2). As these are non-transmembranous MFGMPs, they might have been liberated in the W during the manufacture of cheese. 477

478

#### 479 **4.** Conclusions

480

In the last decade, more and more nutritional- and health-related aspects of milk
proteins have been discovered (Supplementary material, Table S1), making it more relevant
for research and for dairy manufacturers to define dairy products as detailed as possible to

satisfy the increasing awareness of the consumer for health and nutrition. While methods for
relative quantification of major milk proteins date back to 1944 (Warner, 1994), methods for
quantification of the minor milk proteins are rare.

487 The LC-MS method presented in this study provides a simultaneous absolute quantification of twenty bovine milk proteins without elaborate or time-consuming sample 488 preparations. The selected proteins belong to all three main milk protein classes, CasMPs, 489 WPs, and in particular MFGMPs, for which a quantification method was lacking so far. The 490 491 MRM results for RM are in good accordance with literature data. Unfortunately, the precise quantification of BSA was not yet satisfying in RM and needs to be further investigated, 492 possibly by adding a heat-treatment during sample preparation leading to protein unfolding 493 and reshuffling of the disulphide bridges, as the BSA concentrations in heat-treated milk 494 samples were within the expected range (Table 4). 495

The method achieves a good precision in RM for proteins present at higher 496 concentrations (CVs of 5–15%; Supplementary material, Table S2). For the minor MFGMPs 497 498 (ADPH, PAS 6/7, CD36, PIGR), the results obtained showed higher CVs (up to 33%) due to 499 the low individual protein concentration in milk, being close to the quantitation limit. The accuracy of the method was checked by recovery experiments and ranged between 91% 500 501 and 105%. In the future, the method can be applied for screening purposes assessing the 502 individual protein content of milk produced under different feeding regimes, at different 503 lactation stages or between different breeds. Moreover, the method is suited for the 504 quantification of specific milk proteins in the whole range of different dairy products and can as such be used for the evaluation of bioactive effects, since most known bioactive peptides 505 are embedded in the sequence of major milk proteins. Therefore, the presented 506 507 straightforward quantification method is a valuable and convenient tool for the simultaneous guantification of the minor and most abundant milk proteins, and will contribute to a better 508 understanding of the impact of protein composition on the technological and nutritional 509 quality of milk and dairy products. 510

511

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520	
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### **Figure legends**

**Fig. 1.** Panel a: quantity (g 100 g<sup>-1</sup> total protein) of casein micelle proteins (CasMPs), whey proteins (WPs) and milk fat globule membrane associated proteins (MFGMPs) in raw milk (RM), raw cream (RC), Swiss Emmental cheese (EM) and whey (W), determined by multiple reaction monitoring. Mean values and standard deviations were determined for twelve RM (in technical duplicates, each measured three times), for RC, EM and W, each sample was measured thrice in technical triplicates. Panel b: SDS-PAGE showing the protein pattern for each raw dairy product. For abbreviations for the proteins, see Table 1.

**Fig. 2.** Panel a: average amount (g 100 g<sup>-1</sup> total protein) of casein micelle proteins (CasMPs), whey proteins (WPs) and milk fat globule membrane associated proteins (MFGMPs) in heattreated milk (MI), cream (CR), yoghurt (YOG), buttermilk (BM), quark (Q) and cottage cheese (COTC) determined by multiple reaction monitoring. Mean values and standard deviations for every group of dairy products were determined for three products from different manufacturers, each in technical triplicates, measured thrice. Panel b: SDS-PAGE showing a representative protein pattern for each type of dairy product. For protein abbreviations, see Table 1. Data of the signature peptides for the quantification of twenty bovine milk proteins by multiple

### reaction monitoring (MRM).<sup>a</sup>

		Signature peptide sequence (Internal standard)	r. t. (min)	Pep. charge	Pep. <i>m/z</i>	Frag. <i>m/z</i>	Frag. <i>m/z</i>
Casein micelle	e proteins	(Internal standard)	()	onargo			11#2
CASA1	$\alpha_{s_1}$ -Casein	YLGYLEQLLR	10.8	2	634.6	992.1	771.9
0/10/11		YLGYLEQLLR*	10.0	-	639.6	1002.1	781.9
CASA2	α <sub>s2</sub> -Casein	FALPQYLK	9.3	2	490.7	648.8	761.9
		FALPQYLK*				656.8	769.9
CASB	β-Casein	GPFPIIV	11.2	1	742.5	625.8	441.6
		GPFPIIV*		748.5	625.8	447.6	
CASK	κ-Casein	YIPIQYVLSR	10.0	2	627.1	488.6	976.1
		YIPIQYVLSR*		632.1	493.6	986.1	
LPL	Lipoprotein lipase	EPDSNVIVVDWLSR	10.6	2	815.3	624.6	875.0
		EPDSNVIVVDWLSR*		820.3	624.6	885.0	
Whey proteins	3						
LACB	β-Lactoglobulin	ALPMHIR	6.7	2	419.6	327.4	653.8
		ALPMHIR*			424.6	332.4	663.8
LALBA	α-Lactalbumin	VGINYWLAHK	8.8	2	601.4	523.1	932.1
		VGINYWLAHK*			605.4	527.1	940.1
PP3	Lactophorin (proteose peptone 3)	LPLSILK	9.0	2	392.7	335.9	573.7
		LPLSILK*			396.7	339.9	581.7
BSA	Bovine serum albumin	LGEYGFQNALIVR	9.7	2	740.8	814.0	685.8
		LGEYGFQNALIVR*			745.8	824.0	695.8
LPO	Lactoperoxidase	ASEQILLATAHTLLLR	10.4	3	584.6	611.8	498.6
		ASEQILLATAHTLLLR*			587.9	616.8	503.6
LF	Lactoferrin	YLTTLK	6.8	2	370.1	462.6	277.3
		YLTTLK*			374.1	470.6	277.3
Milk fat globul	e membrane proteins						
BTN	Butyrophilin (Subfamily 1 Member A1)	EIPLSPMGEDSASGDIETLHSK	8.6	3	772.4	887.9	1036.6
		EIPLSPMGEDSASGDIETLHSK*			775.1	891.9	1040.6
XDH	Xanthine dehydrogenase/oxidase	TNLSSNTAFR	6.7	2	556.4	782.8	896.0
		TNLSSNTAFR*			561.4	792.8	906.0
ADPH	Adipophilin (adipophilin differentiation	VANLPLVSSTYDLVSSAYISRK	10.4	3	795.6	994.1	1107.3
	related protein ADRP)	VANLPLVSSTYDLVSSAYISRK*			798.3	998.1	1111.3
PAS 6/7	Lactadherin	NIFETPFQAR	9.5	2	611.9	996.1	719.8
		NIFETPFQAR*			616.9	1006.1	729.8
CD36	Platelet glycoprotein 4	VAIIDTYK	7.7	2	461.9	752.9	639.7
		VAIIDTYK*			465.9	760.9	647.7
PIGR	Polymeric immunoglobulin receptor	SPIFGPEEVTSVEGR	9.2	2	482.5	530.6	608.7
		SPIFGPEEVTSVEGR*			487.5	535.6	613.7
FABP	Fatty acid binding protein	SIVTLDGGK	7.2	2	445.4	689.8	590.6
		SIVTLDGGK*			449.4	697.8	598.6
FAS	Fatty acid synthase	IPALQDGR	6.4	2	435.5	378.9	659.7
		IPALQDGR*			440.5	383.9	669.7
GP2	Glycoprotein 2 (zymogen granule	DSTISVEENGVSAESR	7.2	2	840.8	549.6	1078.1
	membrane)	DSTISVEENGVSAESR*			845.8	559.6	1088.1

<sup>a</sup> Amino acids (AAs) are abbreviated according to the IUPAC-IUB Joint Commission on

Biochemical Nomenclature (JCBN) standard; an asterisk indicates isotopically labelled by <sup>13</sup>C

and <sup>15</sup>N contained in the last amino acid (AA) of each peptide. Abbreviations are: r.t.,

retention time; Pep., peptide; Frag., fragment.

### Table 2

Quantity of twenty individual milk proteins in raw milk, raw cream, Swiss Emmental cheese

Protein	Product	Product								
	Raw milk		Raw crea	Raw cream		al	Whey			
	Amount	σ	Amount	σ	Amount	σ	Amount	σ		
Casein mic	elle protein	S								
CASA1	31.1	1.8	23.3	1.1	36.4	3.1	1.00	0.04		
CASA2	11.1	1.2	9.5	0.4	11.6	0.9	0.12	0.01		
CASB	31.2	3.7	25.8	0.9	42.5	2.3	0.38	0.02		
CASK	6.9	1.0	6.0	0.2	9.6	0.7	0.07	0.004		
LPL	0.052	0.006	0.047	0.005	0.066	0.005	0.025	0.005		
Whey proteins										
LACB	10.2	1.3	10.6	1.1	0.39	0.04	58.9	1.6		
LALBA	2.0	0.3	3.3	0.6	0.16	0.05	18.7	1.0		
PP3	1.0	0.1	2.0	0.2	0.064	0.009	4.2	0.2		
BSA	0.02	0.01	0.016	0.005	0.003	0.001	1.9	0.5		
LPO	0.040	0.022	0.046	0.008	0.032	0.04	0.171	0.024		
LF	0.25	0.052	0.25	0.02	0.20	0.013	0.19	0.03		
Milk fat glo	bule memb	rane proteins								
BTN	1.15	0.19	3.35	1.69	0.21	0.11	0.44	0.22		
XDH	0.146	0.03	0.578	0.04	0.073	0.008	0.374	0.08		
ADPH	0.00025	0.0001	0.0069	0.003	0.0036	0.0001	0.00008	0.00005		
PAS 6/7	0.02	0.01	0.93	0.13	0.27	0.04	0.25	0.04		
CD36	0.022	0.006	0.177	0.08	0.023	0.008	0.055	0.05		
PIGR	0.31	0.12	0.36	0.17	0.05	0.06	0.02	0.01		
FABP	0.03	0.02	0.17	0.07	0.05	0.01	0.43	0.2		
FAS	0.027	0.007	0.083	0.005	0.023	0.004	0.051	0.006		
GP2	0.023	0.003	0.035	0.005	0.005	0.001	0.071	0.007		

and whey determined by multiple reaction monitoring (MRM). <sup>a</sup>

<sup>a</sup> Mean values (g 100 g<sup>-1</sup> total protein) and standard deviations were determined on twelve raw milk samples (3.5% total protein), each measured three times, one raw cream sample (2.2% total protein), one Swiss Emmental cheese sample (30.3% total protein), and one whey sample (0.5% total protein) in technical triplicates, measured three times. Protein abbreviations are listed in Table 1.

Comparison of the concentration of twenty milk proteins in raw milk determined by multiple

Protein Concentration in g 100 g<sup>-1</sup> protein Concentration in g L<sup>-1</sup> MRM Literature data MRM Literature data Walstra et al., 2006 Tremblay Swaisgood, Eigel et al., Kuczynska et al., 2003 1995 1984; et al., 2012 Swaisgood, 1993 Casein micelle proteins CASA1 32 11.36 10.0 11.9 12-15 31.1 CASA2 3-4 11.1 8.4 4.02 2.6 3.1 CASB 31.2 26 11.33 9.3 9.8 9-11 CASK 6.9 9.3 2.5 3.3 3.5 2-4 LPL 0.052 -0.019 -Whey proteins LACB 10.2 9.8 3.7 3.2 3.2 2-4 2.68-4.12 0.6-1.7 LALBA 2.0 3.7 0.74 1.2 1.73-2.06 1.2 PP3 1.0 0.37 0.3 -BSA 0.02 1.2 0.4 0.12-0.2 0.01 0.4 0.4 LPO 0.040 0.015 LF 0.25 -0.09 0.1 0.19-0.33 \_ \_ Milk fat globule membrane proteins BTN 1.15 0.42 XDH 0.146 0.054 ADPH 0.00025 0.0001 \_ -PAS 6/7 0.0078 0.02 -CD36 0.022 0.0079 PIGR 0.31 0.11 FABP 0.03 0.01 FAS 0.01 0.027 GP2 \_ 0.023 0.0086

reaction monitoring (MRM) with published literature data. <sup>a</sup>

<sup>a</sup> Means determined by multiple reaction monitoring (MRM) were on twelve raw milk samples, each measured three times. Data from Walstra, Wouters, & Geurts (2006) are approximate composition and those from Swaisgood (1995) are averaged values. Protein abbreviations are listed in Table 1.

Average amount of twenty individual milk proteins in groups of heat-treated dairy products

determined by multiple reaction monitoring (MRM). <sup>a</sup>

Protein	Product											
	MI		CR		YOG		BM		Q		COTC	
	Amoun	tσ	Amount	σ	Amount	σ	Amount	σ	Amount	σ	Amount	σ
Casein m	icelle prot	eins										
CASA1	20.90	2.54	13.98	0.28	19.14	1.79	20.58	1.65	23.76	2.89	24.61	1.5
CASA2	10.37	0.63	7.88	0.98	9.93	1.35	8.17	0.58	12.08	0.81	15.75	0.51
CASB	25.31	1.43	16.85	0.57	25.80	2.94	18.32	1.06	30.64	2.55	35.93	2.76
CASK	6.43	0.26	4.43	0.60	4.76	0.35	4.29	0.105	5.89	0.48	7.68	0.18
LPL	0.05	0.005	0.04	0.005	0.05	0.003	0.04	0.003	0.06	0.007	0.06	0.0004
Whey pro	teins											
LACB	9.23	1.06	7.74	0.97	9.38	1.44	10.96	0.53	12.69	1.70	2.03	0.23
LALBA	1.26	0.35	0.43	0.22	1.53	0.33	2.05	0.25	1.85	0.20	0.43	0.06
PP3	0.84	0.05	1.97	0.35	0.81	0.11	2.29	0.06	0.25	0.07	0.46	0.008
BSA	0.21	0.10	0.30	0.028	0.37	0.04	0.27	0.005	0.47	0.02	0.18	0.02
LPO	0.05	0.003	0.03	0.003	0.08	0.01	0.08	0.008	0.09	0.007	0.05	0.004
LF	0.33	0.03	0.26	0.07	0.38	0.080	0.41	0.008	0.30	0.037	0.32	0.012
-	obule mer	mbrane pro	oteins									
BTN	0.35	0.05	1.95	0.43	0.33	0.02	2.26	0.09	0.22	0.018	0.36	0.038
XDH	0.22	0.05	1.00	0.20	0.23	0.028	1.31	0.08	0.22	0.018	0.23	0.015
ADPH	0.0009	0.0001	0.0023	0.001	0.0006	0.0001	0.0014	0.0001	0.0007	0.0001	0.0003	0.00004
PAS 6/7	0.30	0.05	1.48	0.24	0.36	0.04	3.55	0.71	0.11	0.016	0.28	0.02
CD36	0.07	0.005	0.25	0.063	0.07	0.02	0.24	0.026	0.08	0.012	0.06	0.005
PIGR	0.22	0.024	0.34	0.10	0.59	0.05	0.35	0.101	0.13	0.025	0.41	0.23
FABP	0.09	0.005	0.17	0.04	0.09	0.02	0.33	0.05	0.25	0.034	0.12	0.036
FAS	0.02	0.003	0.12	0.04	0.01	0.003	0.13	0.005	0.02	0.005	0.02	0.006
GP2	0.02	0.01	0.06	0.02	0.03	0.001	0.06	0.005	0.04	0.002	0.02	0.003

<sup>a</sup> Abbreviations are: MI, milk (3.5–3.9% fat); CR, cream (35% fat); YOG, yoghurt (3.5–4% fat): BM, buttermilk (0.5% fat); Q, quark (0.1% fat); COTC, cottage cheese (4–4.5% fat). Values are in g 100 g<sup>-1</sup> total protein; mean values and standard deviations for every group of dairy products were determined for three products from different manufacturers, each in technical triplicates, measured thrice. Protein abbreviations are listed in Table 1.



(b)

(a)



(b)



Figure 2