1	Evaluation of production of Cheddar cheese from micellar
2	casein concentrate
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14 ABSTRACT

15 The objective of this study was to evaluate the production of Cheddar cheese using micellar casein concentrate (MCC), a novel milk ingredient powder with a high casein 16 17 content (~92%). Four types of Cheddar cheese were manufactured and ripened for 180 18 days from the following starting materials: standardized control milk (control), skim 19 milk with cream (SC), reconstituted MCC with cream (MC) and reconstituted low-heat 20 skim milk powder with cream (PC). Only minor differences were found in composition 21 between treatments, but MC cheese showed higher levels of proteolysis compared to 22 other treatments, linked to significantly higher plasmin and chymosin activities. No 23 differences were observed in hardness between treatments (60, 120 and 180 days), but 24 the springiness and cohesiveness of MC and PC cheeses were significantly higher than 25 that of the control and SC cheeses at 60, 120 and 180 days. In conclusion, the use of 26 casein-dominant dairy streams has the potential for production of Cheddar cheese with 27 tailored functionality.

28 **1. Introduction**

29 Global milk production was approximately 843 million tonnes in 2018 (FAO, 30 2019); among dairy products (i.e., cheese, casein and butter), cheese production used 31 the highest proportion of milk. Whey protein powder, production of which is 32 traditionally associated with the manufacture of cheese, represents 57% of the market 33 for global protein supplement for exercising and nutrition, while there is a huge 34 market for whey protein powders in infant formula. Cheese output is increasing at a 35 rate of 2% yearly while the demand for whey protein has been growing at 6-7% vearly (Hoogwegt Group, 2019). Nowadays, whey protein with high quality is 36 37 required as it can be used to manufacture a range of food ingredients or products with 38 nutrition and functional properties (Boland, 2011). Kelly (2019) reported that the 39 ultrafiltration (UF) properties of sweet and acid whey are influenced by their high or low pH, respectively. However, microfiltration (MF) permeate made directly from 40 41 skim milk is considered to be an ideal whey source for whey protein ingredients production. Therefore, recovering whey protein from milk rather than cheese whey 42 43 could improve the whey quality and is an option for whey protein ingredients 44 manufacture.

During membrane filtration of skim milk, native micellar casein is concentrated
in the retentate, which could be recovered and concentrated to produce micellar
casein concentrate (MCC). This is a novel dairy ingredient powder with a high casein
fraction of 85-95% of total protein which may be used in functional and nutritional

applications (Crowley et al., 2018). As the traditional way to manufacture Cheddar
cheese is followed by whey protein manufacture, the concept of recovering whey
protein before Cheddar cheese manufacture is of interest.

52 In terms of the protein ingredients being used for Cheddar cheese manufacture, 53 low heat skim milk powder (LHSMP) can be used to enhance cheese yield giving a 54 constant cheese production throughout the year (Freeman et al., 1970). LHSMP is 55 produced from skim milk using low temperatures during manufacture and is mostly 56 used for condensed milk, UHT-treated fluid milk and ice-cream (Augustin & 57 Margetts, 2003). Unlike reconstituted medium- and high-heat skim milk powder that have impaired rennet coagulation ability, the rennet coagulation properties of 58 59 LHSMP are good as the whey protein is not highly denatured (Ménard et al., 2005). With only physical separation processing, the micellar casein in MCC may have 60 61 better rennet coagulation properties compared to LHSMP. The rennet coagulation 62 properties and cheese-making potential of MCC may be better than those of LHSMP. 63 Two main factors that influence cheese yield are lactation and seasonality. The 64 gross composition of bovine milk varies with the stage of lactation. Kuchtík et al. 65 (2008) reported that protein and casein content increase and lactose content decreases through the lactation. After 200 days lactation, cows are in the late lactation stage, 66 which requires the udder tissue to repair and recover for next lactation. During late 67 68 lactation, milk yield decreases dramatically, which may influence cheese yield, unless milk is standardised. Also, at specific times of the year, the milk volume 69 70 decreases, and the composition and rennet coagulation properties of milk are

71	significantly influenced (Freeman et al., 1970; O'Brien et al., 1999). Both cheese
72	yield and manufacturing efficiency are influenced by seasonal variation of milk
73	protein and fat composition (Barbano & Sherbon, 1984).
74	To solve the problem of low cheese yield, one solution is adding LHSMP to
75	low-protein milk to increase the protein composition (Freeman et al., 1970). Another
76	possibility may be making Cheddar cheese with MCC that is manufactured as a co-
77	product of high-quality whey protein.
78	This study aimed to evaluate the production of Cheddar cheese from micellar
79	casein concentrate, with LHSMP used for comparison. The consequence of this
80	manufacture was evaluated concerning composition, proteolysis, texture and

81 functionality.

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83 **2** Materials and Methods

84 2.1 Preparation of cheese milk

Micellar casein concentrate (MCC) powder was obtained from Teagasc
(Moorepark, Fermoy Co. Cork, Ireland); whereby pasteurized bovine skim milk was
microfiltered at 50 °C using 0.14-µm TAMI Isoflux ceramic membranes (TAMI,
Lyons, France). MF was performed in a batch mode involving two diafiltration steps
to produce a final 3X MF retentate (liquid MCC) which was evaporated at 65 °C
using a pilot plant single-effect falling-film evaporator (Anhydro F1 Laboratory;

91 Copenhagen, Denmark). This was followed by spray-drying of the evaporated liquid 92 MCC in a single-stage spray dryer (Anhydro Laboratory Spray Dryer, SPX Flow 93 Technology, Denmark) equipped with nozzle atomisation at inlet and outlet 94 temperatures of 185 and 85°C, respectively. The protein content (91.6% casein) of 95 MCC was determined by the Kjeldahl method (IDF, 1986). MCC powder and low 96 heat skim milk powder (LHSMP; WPNI=6.0) (Uelzena, Uelzen, Germany) were 97 rehydrated using a Silverson mixer at 55 °C for 2 h.

Raw milk was obtained from a local farm in Cork, Ireland. Skim milk and cream 98 99 were separated from raw milk using a separator. Composition of milk samples was 100 measured by MilkoScanTM Mars (Foss, Hilleroed, Denmark) and casein content was 101 calculated by multiplying the casein percentage of protein of the milk sample (an estimated percentage of 78% for milk and LHSMP milk and 91.6% for MCC). 102 103 Control milk was prepared by combining raw milk and skim milk to a casein: fat 104 ratio of 0.7:1. Both reconstituted MCC and reconstituted LHSMP were made up to 105 the same casein level as skim milk. Reconstituted MCC and LHSMP and skim milk 106 were blended with cream and lactose according to the same standardization ratio 107 (casein: fat of 0.70:1.00) and lactose content (~5%) as control milk to obtain three more vats: skim milk with cream (SC), reconstituted MCC milk with cream (MC) 108 109 and reconstituted LHSMP milk with cream (PC). All milk samples were pasteurized at 72 °C for 15 s (Microthermics Inc., Raleigh, NC, USA) and stored at 4 °C until 110 111 analysis and cheese-making.

113	Dynamic oscillatory analysis (small amplitude oscillatory measurement) of
114	renneted control milk, SC milk, MC milk and PC milk was performed using a Peltier
115	concentric cylinder geometry, which comprised of an aluminium conical rotor [42.01
116	mm (h) by 28.02 mm (d)], on an AR-G2 controlled stress rheometer (Waters TA
117	Instruments, Leatherhead, Surrey, UK). Aliquots (25 mL) of each sample were pre-
118	warmed in a water bath at 32°C for 15 min, and 80 μL of a 1:10 (v/v) dilution of
119	Maxiren TM (Chr Hansen A/S, Hørsholm, Denmark) was added. The sample was
120	placed immediately in the preheated cup (32 $^{\circ}$ C), the frequency of oscillation was set
121	at 0.6283 rad s ⁻¹ , and the storage modulus, G' of the sample was recorded
122	continuously as a function of time at a low-amplitude shear strain (0.01) over 90 min.
123	Each sample was analysed in triplicate (Ibáñez et al., 2015).

124 2.3 Cheese Manufacture

125 Cheddar cheeses were made from 20 L control, SC, MC and PC milks which 126 were prepared, pasteurised and stored at 4 °C overnight before cheese-making. 127 Cheddar-type cheeses were manufactured according to a standard Cheddar cheese 128 manufacture protocol (Fox et al., 2000). An aliquot of 6 g (0.03% w/w) of Cheddar cheese starter culture (R604Y Chr. Hansen Ltd., Little Island, Co. Cork, Ireland) was 129 added into each sample and allowed ripen for 30 min, followed by 0.09% (v/w) of 1 130 mol L⁻¹ CaCl₂ and 60 IMCU L⁻¹ rennet being added to all samples. Whey was drained 131 when the pH dropped to 6.2 and curd was cheddared. When the pH decreased to 5.4, 132

133	the curd was milled into small pieces and salted at a level of 2.5% (w/w) NaCl. The
134	curds were then wrapped in cheesecloth and moved to 2-kg circular moulds, which
135	were pressed at a pressure of 2.5 kg cm ⁻² for 14 h. The cheeses were then removed,
136	vacuum-packed and ripened at 8 $^\circ C$ for 180 days.
137	2.4 Compositional analysis
138	Gross composition of cheeses was analysed at 14 days old. Moisture was
139	determined by oven-drying method (IDF 1982), protein (%N×6.38) by the Kjeldahl
140	method (IDF 1986) and salt was analysed by titration with AgNO ₃ (Fox, 1963). The
141	pH was measured at 14, 30, 60, 120 and 180 days of ripening by using a calibrated
142	pH probe placed in contact with dry grated cheeses. All results were determined in
143	triplicate.

144 2.5 Microbiological analysis

Enumeration of starter lactic acid bacteria (LAB) was performed using LM 17
agar plates (Terzaghi and Sandine 1975), incubated for 3 days at 30 °C. Enumeration
of non-starter lactic acid bacteria (NSLAB) was performed on Rogosa agar plates
(Rogosa and Mitchell 1951), incubated anaerobically for 5 days at 30 °C.
Enumeration of LAB and NSLAB were performed in duplicate after 60 and 90 days
of ripening.

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154	Urea-polyacrylamide gel electrophoresis (urea-PAGE) of cheese samples was
155	used to study the proteolysis of α_{S1} - and β -casein (CN) during ripening using the
156	method of Andrews (1983) with modifications of Shalabi and Fox (1987). The pH
157	4.6-soluble and insoluble fraction were prepared (Kuchroo & Fox, 1982), and peptide
158	profiles of pH 4.6-soluble fractions filtered through 0.22 -µm cellulose acetate filters
159	(Sartorious GmbH, Gottingen, Germany) were analysed by reverse-phase high-
160	performance liquid chromatography (HPLC) using an ultra-performance liquid
161	chromatography (UPLC) Waters Acquity UPLC H-Class Core System (Waters,
162	Milford, MA, USA), with a Waters Acquity UPLC TUV Detector (dual-wavelength;
163	Waters) operated by Empower 3 software (Waters Corp., Milford, MA, USA),
164	following the method of Mane et al. (2019).
165	For determination of free amino acids (FAA), frozen pH 4.6-soluble fractions
166	were de-proteinised by mixing equal volumes of 24% (w v ⁻¹) trichloroacetic acid

167 (TCA) and sample and following the method of Fenelon and Guinee (2000).

168 2.7 Plasmin activity

169 The plasmin activity of cheese samples at 180 days of ripening was measured 170 using the coumarin peptide method (Richardson & Pearce, 1981). A standard curve 171 of the emission intensity at 460 nm was constructed using 7-amido-4-methyl 172 coumarin (AMC), and results expressed in nmol AMC min⁻¹ mL⁻¹, which was 173 defined as one unit of plasmin activity.

175 Grated cheese samples (50 mg) were extracted by dissolving cheese in 1 mL of 0.1 M trisodium citrate, followed by 30 minutes incubation at 37 °C. Fat was 176 177 separated by centrifugation at 1000 g (Sigma 1-16K, Harz, Germany) for 1 min, and 178 the aqueous layer was used for analysis. An aliquot of 70 µL citrate dispersion of cheese was incubated with 30 μ L of 1 mg mL⁻¹ aqueous solution of a synthetic 179 180 heptapeptide substrate (Pro-Thr-Glu-Phe-[NO₂-Phe]-Arg-Leu) in 400 µL 0.1 M 181 sodium formate buffer, at 37 °C, pH 3.2, for 24 h. The mixture was heated at 70 °C 182 for 10 min to stop the reaction, followed by centrifugation at 16,000g for 10 min, and 183 the supernatant was used for UPLC analysis. Substrate and product levels were 184 determined using the reversed-phase HPLC system described above, following the 185 method of Hurley et al. (1999).

186 2.9 Texture profile analysis

187 Texture profile analysis was performed using a Texture Analyzer TA-XT2i 188 (Stable Micro Systems, Godalming, Surrey, UK) at 60, 120, and 180 days of ripening. 189 Cheese samples were cut into 20 mm height, 20 mm diameter cylinders and kept at 190 4 $^{\circ}$ C overnight. Cheese cylinders were compressed to 25% of the initial height in two 191 continuous compressions with a speed of 1 mm s⁻¹. Hardness, cohesiveness and 192 springiness were measured (Truong et al., 2002), and five cheese samples were 193 measured for each treatment.

194

Meltability was analyzed using the Schreiber meltability test as described by
Altan et al. (2005). Cheese samples were heated at 232 °C for 5 min, and meltability
was measured as the percentage increase in diameter of the original samples.
Analyses were performed in triplicate at 60, 120, 180 days of ripening.

200 2.11 Dynamic small amplitude oscillatory rheology

The rheological properties of meltability of cheese after 180 days ripening were analysed with a controlled stress AR-G2 rheometer (TA Instruments, Waters LLC, Leatherhead, Surrey, UK) according to the method of Ibáñez et al. (2015). Storage modulus (G'), loss modulus (G''), and loss tangent (LT) were measured during heating. The maximum LT (LT_{max}) and the temperature where LT=1, which are indicators of melting, were also recorded. Each sample was analysed in triplicate.

207 2.12 Colour measurements

Colour values were measured using a Konika-Minolta colourimeter CR400 (Konika-Minolta Optics Inc., Osaka, Japan) at 14, 30, 60, 120, 180 days of ripening. The measurement used the CIELAB system based on illuminant D65 and a visual angle of 2°. Five random readings were taken on fresh-cut cheese at 20°C. The Eluclidean distance between the colour of control cheese and that for other treatments was calculated by $\Delta E^*_{ab} = \sqrt{(\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2}$ where $\Delta L^* = L^*_{sample} - L^*_{control}, \Delta a^* = a^*_{sample} - a^*_{control}$ and $\Delta b^* = b^*_{sample} - b^*_{control}$ (Mahy et al., 1994).

216 Control, SC, MC and PC cheeses were made in three independent trials, 4×3 217 blocks. Statistical analysis was carried out using one-way analysis of variance 218 (ANOVA) with R project for Windows version i386 3.4.0 to establish significant 219 differences between samples. The probability level used for statistical significance 220 was P < 0.05.

221

222 **3** Results and Discussion

223 3.1 Rennet coagulation properties

224 The rennet coagulation properties of milk samples are shown in Table 1. During 225 the first 10 min, G' value of all the treatments was low and constant, after which the 226 G' value of all samples increased, with the MC and PC samples exhibiting a slower rate of increase compared to other samples (Fig. 1). The gel strength (G' value) at 90 227 min of MC and PC samples was significantly (P < 0.05) lower than that of other 228 229 samples. The gel strength of MC and PC kept increasing and did not reach a constant 230 level during the analysis, while control and SC reached stable values at 45 min. The 231 loss tangent at 90 min of MC sample was significantly (P < 0.05) higher than that of 232 the other treatments.

233 Martin et al. (2010) reported that, during the manufacture of milk protein 234 concentrate (MPC), casein micelles were not damaged; however, the minerals removed through diafiltration and decreased calcium ionic strength could depress rennet coagulation. The rennet coagulation ability of reconstituted MPC could be restored by adding extra calcium ions. The production of MCC involves two diafiltration steps, which may reduce soluble calcium levels; thus, 0.09% (v/w) of 1 mol L^{-1} CaCl₂ was added to all of the batches before cheese manufacture.

240 *3.2 Cheese composition and pH*

241 The composition of cheese at 14 days of ripening and pH of cheese samples at 242 14, 30, 60, 120 and 180 days of ripening are shown in Table 2; the results presented 243 are means of data from the three trials. Guinee et al. (2007) stated that different casein 244 to fat ratio can influence the protein and fat content of cheese. As the casein: fat ratio 245 of milk was adjusted to 0.7 for all samples, no statistically significant differences 246 (P>0.05) were found in cheese protein and fat contents between all treatments, and 247 the use of MCC or LHSMP for Cheddar cheese manufacture did not affect these 248 parameters.

The moisture content of PC cheese was significantly (P < 0.05) higher than that of SC and MC cheese; no significant differences (P > 0.05) were found for moisture content between control, SC and MC cheese. The salt content of MC cheese was significantly (P < 0.05) higher than that of control and SC cheese. The pH of MC cheese was significantly (P < 0.05) higher than that of other treatments, the reason for which is not clear. No significant differences (P > 0.05) were found for moisture in non-fat substances (MNFS) and fat in dry matter (FDM) for all treatments; MNFS is an important parameter for Cheddar cheese quality, so it is important that all batches of
cheese sample had similar levels of MNFS (Fox et al., 2000; Bogenrief & Olson,
1995).

259 3.3 Proteolysis

260 Table 2 shows the level of pH 4.6-SN/TN (%) of control, SC, MC and PC cheeses at 180 days of ripening; pH 4.6-SN/TN is an index of proteolysis (Sousa et al., 2001). 261 262 Significantly (P<0.05) higher pH 4.6-SN/TN levels were found in MC cheese than in 263 SC and PC cheese. In addition, the levels of pH 4.6-SN/TN were significantly (P<0.05) 264 lower in PC than control cheese. The value (23%) found at 180 days of ripening in this 265 study was comparable with previous studies on Cheddar cheese (Lucey et al., 2005; 266 O'Mahony et al., 2005). The main agent producing pH 4.6-soluble nitrogen is the 267 coagulant (O'Keeffe et al., 1978), while Farkye & Fox (1992) reported that plasmin, 268 the principal indigenous milk proteinase, is also important for primary proteolysis in 269 Cheddar cheese during ripening.

Urea-PAGE electrophoretograms of cheese samples during ripening are shown in Fig. 2. All cheese samples showed break-down of β- and α_{S1} -CN, but lower levels of intact β-and α_{S1} -CN were apparent during ripening in MC cheese. Due to the action of plasmin, β-CN is hydrolysed to β-CN (f29-209), β-CN (f106-209) and β-CN (f108-209) (Eigel et al., 1984). The levels of β-CN (f106-209) and β-CN (f108-209) of MC cheese were higher than other treatments during ripening, which suggests higher plasmin activity. Also, in Fig. 2, the MC cheese showed the lowest level of intact α_{S1} -CN level, followed by PC cheese, control and SC cheese from 60 to 180 days of ripening; at 180 days of ripening, no intact α_{S1} -CN was found in MC and PC samples. Residual chymosin in cheese hydrolyses α_{S1} -CN to α_{S1} -CN (f24-199) and α_{S1} -CN (f102-199) and, as the residual chymosin activity increases, the breakdown of α_{s1} -CN increases (Sheehan et al., 2008). The faster breakdown of α_{S1} -CN of MC and PC samples may indicate higher residual chymosin activity in these samples.

283 The peptide profiles of the pH 4.6-soluble extracts for control, SC, MC and PC cheeses at 180 days of ripening were generated by ultra-performance liquid 284 285 chromatography (Fig. 3). The highest peak areas were observed at the retention time of 1 to 5 min for all treatments. The peak area of peptides that were eluted later (28-45 286 287 min retention time) was higher in MC sample compared to the other treatments. Peptides in the pH 4.6-soluble extracts reflect the effect of proteinases and peptidases 288 289 of starter (Fox & McSweeney, 1997). The number of peptide peaks for all treatments 290 was similar, which suggests that the proteolysis for all treatments broadly followed 291 similar pathways, but differences in peak area indicate that there was more extensive 292 proteolysis in MC cheese compared to the other treatments.

The individual FAA levels of all treatments at 120 and 180 days of ripening are shown in Fig. 4 (A and B). The major FAA determined in cheese were glutamic acid, valine, leucine, phenylalanine, histidine and lysine; Bansal et al. (2009) also reported that glutamic acid, valine, leucine, phenylalanine histidine and lysine are the principal FAA in Cheddar cheese after 180 days of ripening. No significant differences (*P*>0.05) were found between treatments at 120 days of ripening. At 180 days of ripening,

299	significantly (P<0.05) higher concentrations of threonine, serine, glycine, alanine,
300	valine, methionine, isoleucine, phenylalanine, histidine and proline were found in
301	control and SC cheese than that of MC and PC cheese. No significant differences
302	(P>0.05) were found between treatments in the levels of aspartic acid, glutamic acid,
303	leucine, tyrosine and arginine. From 120 to 180 days of ripening, the concentrations of
304	glutamic acid, alanine, leucine, tyrosine and phenylalanine increased significantly
305	(P < 0.05) for all treatments, while only control and SC cheese had a significant increase
306	in the concentration of threonine, glycine, valine and isoleucine. The total FAA levels
307	of all treatments at 120 and 180 days of ripening are shown in Fig. 4 (C). No significant
308	differences (P >0.05) were found between all treatments at 120 days of ripening, but
309	significantly higher (P <0.05) levels of total FAA were found in control cheese than in
310	MC and PC cheeses at 180 days of ripening. No significant differences (P >0.05) were
311	found between control and SC cheese at 180 days of ripening.
312	Fox and McSweeney (1996) reported that peptidases of starter and non-starter lactic
313	acid bacteria are the principal agents releasing FAA in Cheddar cheese during ripening.
314	No significant (P >0.05) differences were found for the numbers of starter bacteria and
315	NSLAB between treatments in this study (result not shown). Therefore, the release of
316	the FAA by starter and NSLAB should not have been an influence in this regard, and
317	so the reasons for this difference is not clear.

321	The plasmin activity of MC cheese was significantly ($P < 0.05$) higher than that of the
322	other cheese batches (Table 3) which is consistent with the result of urea PAGE
323	electrophoresis. The plasmin activity of PC cheese was significantly ($P < 0.05$) lower
324	than that of the other cheese samples. As the pH increases, the hydrolysis of β -CN
325	increases since plasmin has an alkaline optimum pH (Watkinson et al., 2001).
326	Aaltonen and Ollikainen (2011) reported that in diafiltration, with the removal of
327	whey protein, the concentration of inhibitors of both plasmin and plasminogen
328	activators decreases, which promotes the conversion from plasminogen to plasmin and
329	thus increases plasmin activity. In this study, MCC was made using microfiltration,
330	which may have enhanced the plasmin activity of retentate by removing inhibitors in
331	the permeate, resulting in the higher plasmin activity and more β -CN breakdown.MC
332	cheese showed significantly (P<0.05) higher residual chymosin activity compared to
333	the other treatments (Table 3), which is consistent with the faster breakdown of α_{S1} -CN
334	in that cheese. No significant differences (P>0.05) were found between the residual
335	chymosin activities of control and SC cheese, so the recombination of fat and skim milk
336	did not affect the retention of coagulant during cheese manufacture. The residual
337	chymosin activity of PC cheese was significantly (P<0.05) lower than that of control
338	cheese. The MC cheese apparently retained more coagulant compared to control, SC
339	and PC cheeses; the percentage of retained chymosin in cheese curd depends on the pH

340 at the curd-cutting stage, pH at whey drainage, cooking temperature and method341 (Hurley et al., 1999).

342 3.5 Texture profile analysis

No significant changes (P>0.05) were found in the hardness of MC and PC 343 cheeses from 60 to 180 days of ripening (Fig. 5). The hardness of control cheese 344 significantly (P<0.05) decreased between 60 and 120 days of ripening, while that of 345 346 SC cheese significantly (P < 0.05) increased. At 60 days of ripening, the hardness of control cheese was significantly (P < 0.05) higher than that of SC and PC cheeses. No 347 348 significant differences (P>0.05) were observed in the hardness of the cheeses made 349 by different treatments at 120 and 180 days of ripening. Chevanan and 350 Muthukumarappan (2007) reported that the contents of calcium, phosphate, and 351 residual lactose affect the texture profile of Cheddar cheese. The cheese-milk of SC, 352 MC and PC samples was reconstituted according to the composition of control 353 cheese-milk, which would not influence lactose content. In this study, extra calcium 354 chloride was added to all treatments, but the addition of calcium chloride does not 355 cause significant changes to the texture of ripened Cheddar cheese (Soodam et al.,

356 2015).

From 60 days to 180 days of ripening, the springiness and cohesiveness of MC cheese decreased, and the springiness and cohesiveness of MC and PC cheeses were significantly (P<0.05) higher than that of control and SC cheese at 60 and 180 days of ripening. At 120 days of ripening, the springiness of MC cheese was significantly 361 (P<0.05) higher than that of the other treatments and the cohesiveness of MC and PC 362 cheeses was significantly (P<0.05) higher than that of control and SC cheese. Everard 363 et al. (2006) found that higher pH of Cheddar cheese was associated with increased 364 springiness and cohesiveness; the pH of MC cheese was significantly (P<0.05) higher 365 than that for the other treatments (Table 2), which may explain the increases in springiness and cohesiveness. O'Mahony et al. (2005) reported that, as levels of 366 367 secondary proteolysis increase, the charged groups released from peptides would 368 associate with free water, which may lead to increased cohesiveness and springiness 369 during ripening.

370 *3.6 Meltability*

371 Meltability (percentage increase in diameter) of Cheddar cheeses was 372 determined by the Schreiber method; results are shown in Fig. 6. The four types of 373 cheeses showed increases in meltability between 60 and 180 days. At 60 and 120 374 days of ripening, no significant differences (P>0.05) for meltability were found 375 between control and SC cheese. Significantly (P < 0.05) higher meltability was found 376 in control cheese compared to SC cheese. At 60, 120 and 180 days of ripening, the meltability of MC cheese was significantly higher (P < 0.05) than that of cheese from 377 378 the other treatments. At 120 and 180 days of ripening, the meltability of PC cheese 379 was significantly lower (P < 0.05) than that of cheese from the other treatments. 380 The lower meltability of PC cheese may be caused by the changes of casein in

381 protein during drying of milk powder. Moiseev et al. (2017) reported that Mozzarella

382	cheese made with reconstituted non-fat milk powder has lower meltability than the
383	control, due to the drying process of non-fat milk powder decreasing the stability and
384	dispersity of casein micelles and promoting demineralization of calcium salts. Dave
385	et al. (2003) found that the meltability of Mozzarella cheese is related to the
386	breakdown of (α_{S1} -CN and α_{S1} -CN (f24-199)) and breakdown of intact β -casein,
387	while Bogenrief and Olson (1995) reported that hydrolysis of β -casein increases the
388	meltability of Cheddar cheese. Hydrolysis of β -casein is primarily related to the
389	action of plasmin (Eigel et al., 1984) and, although both MC and PC cheeses were
390	made from powder, and showed a higher breakdown of α_{S1} -casein and β -casein, the
391	meltability of MC cheese was significantly higher than that of PC.

392 *3.7 Dynamic small amplitude oscillatory rheology*

393 The results of the rheology of meltability of all treatments at 180 days of ripening are shown in Table 4. Higher LT value indicates a higher extent of melting 394 (Lucey et al., 2003). MC cheese showed significantly higher (P<0.05) LT_{max} values 395 396 than PC cheese, followed by control and SC cheese. The temperature at LT_{max} of MC 397 and PC cheeses was significantly higher (P < 0.05) than that of control and SC cheese. 398 The LT_{max} is an indicator of cheese meltability (Mounsey & O'Riordan, 1999); the more thermal energy needed to melt cheese, the higher the LT_{max} temperature of 399 400 cheese will be.

401 In the Schreiber test for cheese meltability, MC cheese also exhibited the402 highest percentage increase in diameter (Fig. 4). There were differences between

403 Schreiber melting test and dynamic small amplitude rheology for control, SC and PC 404 cheese. Cooke et al. (2013) stated that differences between results generated by those 405 tests may be linked to the more complete fat melting during the higher temperature 406 of the Schreiber melting test. MC cheese exhibited the lowest temperature when 407 LT=1 compared to the rest of the treatments (P < 0.05). At the point of LT=1, cheese 408 is considered to start transforming from a solid to a viscous form (Gunasekaran & 409 Ak, 2002). MC cheese thus commenced melting at the lowest temperature and had 410 the highest meltability compared to the other treatments, which may be linked to the 411 more extensive proteolysis of casein.

412 *3.8 Cheese colour*

413 The colour values of samples at 180 days of ripening are shown in Table 5. 414 Whiteness (L* values) of control, SC cheese was significantly higher (P < 0.05) than 415 that of MC cheese at 180 days of ripening. The greenness of MC and PC cheeses was 416 significantly lower (higher a* values) (P < 0.05) than that of control or SC cheese at 417 180 days of ripening. The yellowness (b* values) of MC cheese was significantly 418 lower (P < 0.05) than that of SC cheese at 180 days of ripening. Ibáñez et al. (2015) 419 reported that the whiteness of cheese may be related to proteolysis; lower whiteness 420 of MC cheese may thus be attributed to higher proteolysis. The ΔE^*_{ab} values are also shown in Table 5. Sharma (2003) reported that ΔE_{ab}^* of above 2.3 leads to a just 421 422 noticeable difference (JND) and, on this basis, overall, no JND was found between

424 between the cheeses made from protein ingredient and control and SC cheese.	423	control cheese and the other treatments. Overall, no visible difference was found
	424	between the cheeses made from protein ingredient and control and SC cheese.

426 **Conclusions**

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427 Cheddar cheeses were made from standardised cheese milk, skim milk with cream, reconstituted MCC with cream and reconstituted LHSMP with cream according to the 428 429 same casein, fat and lactose content. Only minor differences in composition were found with cheese made from reconstituted MCC with added cream compared to control 430 431 cheese. The cheese manufactured with MCC had significantly higher (P < 0.05) levels of pH 4.6-SN/TN than cheese manufactured with skim milk or LHSMP with cream. 432 433 The level of intact β -and α_{S1} -CN of MC cheese was lower than rest treatments, 434 consistent with significantly (P < 0.05) higher plasmin and chymosin activity. No significant difference (P>0.05) were found in hardness between all treatments. 435 Significantly higher (P < 0.05) springiness and cohesiveness were found in cheese 436 437 manufactured from MCC and LHSMP powder, meltability and maximum loss tangent 438 in MC cheese were significantly (P < 0.05) higher than that of the other treatments. No 439 overall differences of colour were found between all treatments. Use of novel caseindominant dairy streams such as MCC has potential for production of Cheddar cheese 440 441 with tailored functionality. The results of this study suggest that using reconstituted 442 LHSMP with cream for the manufacture of Cheddar cheese may result in changes in

443	functionality, while using reconstituted MCC with cream for the manufacture of
444	Cheddar cheese may be more feasible.
445	
446	

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