

# Effect of mono or co-culture of EPS-producing Streptococcus thermophilus strains on the formation of acid milk gel and the appearance of texture defects

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1	Effect of mono or co-culture of EPS-producing Streptococcus thermophilus strains on
2	the formation of acid milk gel and the appearance of texture defects
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### 23 ABSTRACT

25	One acidifying (ST1) and two texturing strains (ST2 and ST3) of Streptococcus thermophilus
26	were used as pure or co-cultures to identify and understand their effects on the structuring of
27	acid milk gels and on the appearance of texture defects, i.e., syneresis and graininess.
28	Symbiosis between specific texturing and acidifying strains reduced acidification time and
29	increased exocellular polysaccharide (EPS) content. The texturing strain could
30	simultaneously produce low and high molar mass EPS and their distribution in mass and/or in
31	number were influenced by the proportion of acidifying to texturing strain used. The results
32	of this study suggest that the high molar mass EPS contributes to acid gel firmness, but less
33	so compared with the acidification rate. The ability of strain ST3 to prevent texture defects,
34	specifically graininess, did not depend on the acidification kinetics or final EPS content, but
35	rather on the structural properties of EPS and/or the bacterial chain morphology.
36	

- 38 1. Introduction
- 39

40 Fermented milk such as yoghurt is one of the most popular dairy products in many 41 countries. In the literature, set-type fermented milk is defined as a viscoelastic gel formed 42 from the aggregation of casein micelles due to milk acidification (Tamime, Robinson, & 43 Latrill, 2001). For manufacturers, a set-type acid milk gel must have a firm body, smooth 44 texture, thick mouthfeel and structural stability. The most common texture defects in milk 45 gels are spontaneous syneresis and graininess. Spontaneous syneresis, or wheying-off, 46 indicates the separation of serum on the yoghurt surface, and graininess usually refers to 47 lumpiness or granular texture due to the presence of large protein aggregates that range in 48 size from 1 to 5 mm (Folkenberg, Dejmek, Skriver, & Ipsen, 2006; Lucey & Singh, 1997). 49 The three main factors determining gel texture and the appearance of defects are the milk 50 composition, the manufacturing process and the starter cultures (Lucey & Singh, 1997; 51 Sodini, Remeuf, Haddad, & Corrieu, 2004). A positive correlation between graininess and 52 high content of denatured whey proteins was reported in the literature (Lucey, 2004; Nguyen 53 et al., 2018a; Remeuf, Mohammed, Sodini, & Tissier, 2003). However, the effect of the 54 starter culture on the graininess has not been clarified.

55 Starter cultures can influence milk gel texture by their acidification rate and/or by 56 their exocellular polysaccharide (EPS) production. EPS from "texturing" or "ropy" cultures 57 are well known to improve the gel viscosity and reduce syneresis (Folkenberg et al., 2006; 58 Marshall & Rawson, 1999; Mende, Mentner, Thomas, Rohm, & Jaros, 2012). Some authors 59 have also observed a reduction in graininess when using a ropy strain or a high-level EPS-60 producing culture, but no further details on the mechanism of this phenomenon have been 61 reported (Hassan, Ipsen, Janzen, & Qvist, 2003; Küçükçetin, Weidendorfer, & Hinrichs, 62 2009). In a previous study (Nguyen et al., 2018a), the effects of heat treatment, milk

63 composition and two EPS-producing starter cultures on the microstructure and the texture 64 defects of acid milk gel have suggested that the presence of denatured whey proteins is 65 critical for grain formation. Gel viscosity and texture defects appeared to be culture-66 dependent and were correlated with the amount, molar mass and intrinsic viscosity of the EPS produced. Furthermore, the gel microstructure obtained from the two cultures showed 67 68 major differences in the location of bacterial cells in the gels, suggesting that simultaneously 69 with the EPS effect, the length of bacterial cell chains could also interfere with the formation 70 of the protein network during acidification due to steric hindrance. However, the different 71 acidification kinetics of the two cultures used might influence gel structuring.

72 In industrial applications, commercial starter cultures for yoghurt fermentation usually 73 consist of several strains of Streptococcus thermophilus, which are chosen for their specific 74 properties such as fast initial acidification or texture promoting capacity. The interaction 75 between different S. thermophilus strains could affect the total EPS production, but different 76 trends were observed and no details on the acidification kinetics or EPS characterisation have 77 been reported (Folkenberg et al., 2006). In addition, the influence of co-cultures of S. 78 thermophilus strains on milk gelation has not yet been studied. The aim of this study was to 79 identify the effects of mono or co-culture of EPS-producing S. thermophilus on the formation 80 of acid milk gels and the appearance of texture defects. For this purpose, three S. 81 thermophilus strains were used alone or in combination: one for its fast acidification 82 characteristics and the other two for their texturing properties. Hence, seven starter cultures 83 were formed with different ratios between acidifying and texturing strains to study the 84 correlations between the gel texture characteristics (e.g., firmness, viscosity, syneresis, 85 graininess), the kinetics of acidification and gelation, but also the quantity and structural 86 features of EPS produced.

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88	2. Materials and methods
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90	2.1. Solvents and reagents
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92	Low-heat skim milk powder (Spray 0) was purchased from Ingredia (France). The
93	three S. thermophilus strains were provided by Chr. Hansen A/S, Arpajon (France). Lactose
94	monohydrate and trichloroacetic acid (TCA) were from GPR Rectapur, VWR chemicals
95	(Belgium). D(+)Glucose monohydrate and 1 N sodium hydroxide (NaOH) were from Merck
96	(Germany). Phenol, LiNO <sub>3</sub> and NaN <sub>3</sub> were from Sigma (USA). Ethanol (95%, $v/v$ ) was from
97	TechniSolv, VWR chemicals (France). Concentrated sulphuric acid (95%, w/w) was from
98	Normapur, VWR chemicals (France).
99	
100	2.2. Starter cultures
101	
102	Three different frozen lyophilised strains of S. thermophilus ST1, ST2, ST3 from the
103	Chr Hansen culture collection (Chr Hansen A/S) were used alone (pure culture) or in
104	combination (co-culture) in seven different starter cultures. From manufacturer information,
105	strain ST1 possess a cell wall protease (PrtS+) and could play a role as acidifying strain. ST2
106	or ST3 were texturing strains without cell wall protease (PrtS-) and were chosen to provide
107	different textural types of milk gels: firm or viscous, respectively.
108	Cultures ST1, ST2 and ST3 were composed of a single S. thermophilus strain
109	corresponding to the name of the culture. The four co-cultures ST1+2 (1/1.7), ST1+2 (5/1),
110	ST1+3 (1/1.7)-and ST1+3 (5/1) consisted of the combination of acidifying strain ST1 with a

112	corresponding to the name of the culture. All strains were stored at -80 °C. Strains were
113	thawed and mixed then 2 g of the final culture were added to 100 mL of UHT skim milk. Ten
114	millilitres of this pre-inoculated milk were immediately added to 1 L of pasteurised milk to
115	start the lactic fermentation. The final inoculation rate for the seven starter cultures was $10^6$
116	cfu mL <sup>-1</sup> pasteurised milk. Sampling, thawing and formulation were performed in the
117	laboratory under sterile conditions.
118	
119	2.3. Acid milk gel manufacture
120	
121	The milk-based preparation was carried out as described by Nguyen et al. (2018b).
122	The milk formulation contained 11.8% (w/w) dry matter, $4.0\%$ (w/w) proteins and $6.7\%$
123	(w/w) lactose. The skim milk powder was dissolved in ultra-pure water during 1h at room
124	temperature and then kept for 12 h at 4 °C before the pasteurisation at 95 °C for 6 min. After
125	the heating step, milk suspensions were cooled to 43 °C and immediately inoculated with the
126	starter cultures. Fermentation procedures, pH monitoring and gel structuring have previously
127	been reported by Nguyen et al. (2018b). In brief, the fermentation was carried out at 43 $^{\circ}C$
128	with pH control until a final pH of 4.65 was reached. In parallel, gel formation was monitored
129	using a Physica MCR 300 rheometer (Anton Paar, St Albans, UK) equipped with concentric
130	cylinder geometry (CC27) that operated in continuous oscillation mode at a frequency of 1
131	Hz and 1% strain. The gel point was defined as the moment when the storage modulus (G')
132	first exceeds the loss modulus (G'') (Winter, 1987). After acidification, the gels were stored
133	undisturbed in plastic cups at 4 °C for 7 days before determination of texture properties.

135 2.4. Gel texture properties

The textural parameters of set-type acid milk gel at day 7 including the firmness, the
viscosity, the spontaneous syneresis and the grain defect were characterised as described by
Nguyen et al. (2018a).

141 The firmness of acid milk gel was determined immediately after removing out of the 142 cold room (4 °C) using a texture analyser equipped with an acrylic cylinder compression 143 probe (diameter = 40 mm, thickness = 5 mm). The compression test was carried out with a 144 speed of 5 mm s<sup>-1</sup>, the trigger force of 5 g and a penetration depth of 15 mm. The gel firmness 145 (N) corresponding to the maximum compression force, i.e., the rupture point of the gel, was 146 calculated with the Exponent software (Stable Micro Systems, Godalming, UK).

For the viscosity measurement, the set-type acid milk gels were stirred 20 times with a spoon and 20 g of sample was placed in the rheometer cup. The viscosity of the stirred gel ( $\eta$ , Pa s) at a shear rate of 300 s<sup>-1</sup> (13 °C) was determined using Physica MCR 300 rheometer equipped with concentric cylinder geometry and the shear rates  $\gamma$  (s<sup>-1</sup>) increased with a linear ramp from 0.271 s<sup>-1</sup> to 300 s<sup>-1</sup> in 210 s.

The spontaneous syneresis (% w/w) was the percentage of whey on the surface of the
undisturbed gel to the total gel weight and was measured according to the siphon method
(Amatayakul, Sherkat, & Shah, 2006).

155The grain defect was determined by image analysis. After stirring 5 times with a156spoon, 1 g of acid gel was dispersed in 100 mL of distilled water by stirring at 400 rpm for1572 min. 10 mL of diluted sample was poured into a plastic Petri dish and visualised with a158digital camera in black and white mode. The image analysis was performed using ImageJ1591.49v software (National Institutes of Health, USA) with Fiji plug-in. Grains with a diameter160> 1 mm were enumerated (Remeuf et al., 2003). The grains were classified into different size

161	groups by their diameter: 1–1.5 mm, 1.5–2.0 mm, 2.0–2.5 mm, 2.5–3 mm or > 3 mm. The
162	results were expressed as the number of grains of each size per gram of gel.
163	
164	2.5. EPS quantification
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166	EPS quantification was performed using the protocol 1 described by Nguyen et al.
167	(2018b). The EPS were extracted in hot-acidic medium and dialysed to obtain a high
168	extraction yield. The total sugar content was determined by the phenol-sulfuric acid method
169	using glucose as the standard (range of concentration from 0 to 200 mg L <sup>-1</sup> ). The acidified
170	milk with glucono- $\delta$ -lactone (GDL) was used as blank and the residual milk sugars after
171	dialysis were subtracted from all data.
172	
173	2.6. EPS characterisation
174	
175	EPS were extracted without a heating step using the protocol 3 described by Nguyen
176	et al. (2018b) to avoid EPS denaturation. The extraction process included a step of pH
177	correction to 7, the protein precipitations in an acidic medium, followed by the precipitations
178	of EPS by ethanol before dialysis and lyophilisation. The structural properties of extracted
179	EPS were characterised using high-performance size exclusion chromatography coupled with
180	multi-angle laser light scattering detector (HPSEC-MALS). The material and operating
181	conditions were reported in Nguyen et al. (2018b). The weight-average molar mass $(M_w)$ and
182	the intrinsic viscosity ([ $\eta$ ]) were determined using the software ASTRA 6.1.2 (Wyatt

- 183 Technologies, Santa Barbara, CA). The  $M_w$  was calculated using the Berry's model (2<sup>nd</sup>
- 184 order). The data were analysed using a refractive index increment (dn/dc) of 0.145 mL  $g^{-1}$
- 185 according to the literature (Lambo-Fodje et al., 2007).

## 187 2.7. Statistical analysis

189		All analyses were conducted at least in triplicate and the values were expressed as the
190	mean	± standard deviation. The experimental data were analysed statistically using the
191	Fische	r's and Student's tests with 95% confidence level and using Pearson's correlation
192	coeffic	cient functions for the correlation between means.
193		
194	3.	Results and discussion
195		
196	3.1.	Characterisation of starter cultures
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198	3.1.1.	Acidification kinetics
199		The kinetics of milk acidification induced by lactic fermentation were monitored by
200	pH ev	olution between pH 6.5 and 4.6 (Fig. 1). The parameters evaluated were the total
201	acidifi	cation time and the maximum acidification rate $(V_{max})$ (Table 1). As an acidifying
202	strain,	ST1 presented the highest $V_{\text{max}}$ value of all pure cultures and the shortest acidification
203	time (·	~300 min) compared with ST2 and ST3. The acidification curve profiles of strains ST2
204	and ST	Γ3 (Fig. 1) were very different from that of strain ST1. ST2 and ST3 had significantly
205	slower	acidification rates during the first 160 min and also a stronger slowdown in
206	acidifi	cation after the shoulder observed around pH 6.25. This specific profile corresponded
207	to the	strains' urease activity, which produced ammonia from milk urea and induced a slight
208	alkalis	ation of the medium. Furthermore, the final count of bacterial cells at the end of the
209	fermer	ntation was ~ $10^9$ , $10^7$ or $10^6$ cfu g <sup>-1</sup> for ST1, ST2 or ST3, respectively, and was
210	consis	tent with the kinetics observed. These differences in acidification kinetics and cell

growth were related to the proteolytic activity of these 3 strains. Indeed, strain ST1 has a cell envelope-associated proteinase (PrtS+) that hydrolyses caseins into peptides and essential amino acids to ensure high cell density growth and thereby generating high acidification activity (Galia, Perrin, Genay, & Dary, 2009). When this enzymatic activity is lacking, such as for ST2 and ST3 (PrtS-), the growth of *S. thermophilus* strains in milk is less optimal depending on the specific intracellular peptidases and amino acids requirements of the strain (Galia et al., 2009).

218 The addition of ST1 to ST2 or ST3 in four co-cultures resulted in a considerable 219 decrease in acidification time and a significant increase in  $V_{\text{max}}$  (Fig. 1, Table 1). For 220 example, the total acidification time for cultures of ST2 was reduced from 1351 min (pure 221 culture) to 528 min or 304 min when ST1 was added with a ST1/ST2 ratio of 1/1.7 or 5/1, 222 respectively. Furthermore, while pure culture of ST3 had the slowest acidification kinetics of 223 all cultures tested, the ST1+3 co-cultures had better acidification performance as compared 224 with ST1+2 co-cultures for the same ST1 ratio. These results suggested that the proteinase 225 activity of ST1 produced peptides and essential amino acids that stimulated growth of both 226 texturing strains. However, the amino acid requirements seemed to be different between ST2 and ST3, as indicated by their differing rates of acidification. ST1 metabolites may be more 227 228 adapted to the needs of ST3 than to those of ST2, which allowed for a better symbiotic 229 relation between ST1 and ST3 than between ST1 and ST2.

- 230
- 231

#### 3.1.2. EPS production and characterisation

The EPS produced by the seven starter cultures were extracted for quantification (Fig. 2) and for macromolecular characterisation (molar mass distribution ( $M_w$ ) and intrinsic viscosity ( $[\eta]$ )) (Fig. 3). Although the strain ST1 was not considered as textural strain, it was able to produce a substantial amount of EPS of ~55 ± 4 mg glucose kg<sup>-1</sup> of gel in a short time

of acidification (~300 min) (Fig. 2). However, the EPS produced have low  $M_w$  of ~9.2×10<sup>4</sup> g 236 237 mol<sup>-1</sup> and low  $[\eta]$  of ~42 mL g<sup>-1</sup>, which could explain the poor texturing property of this strain. These results also confirmed that macromolecular properties are a more critical 238 239 characteristic for the functional properties of EPS than the total amount of polymers (Mende, 240 Rohm, & Jaros, 2016). Even after an acidification time that was 4–5 times longer than for 241 culture ST1, cultures ST2 and ST3 did not produce a greater quantity of EPS (~30 mg glucose kg<sup>-1</sup> and ~52 mg glucose kg<sup>-1</sup>, respectively), particularly for ST2 (Fig. 2). However, 242 the EPS produced by these two texturing strains had very different macromolecular properties 243 244 from each other and from those of the ST1 culture. Based on elution profiles of HPSEC-245 MALS analysis, EPS produced by ST2 or ST3, could be divided into two groups of low and 246 high M<sub>w</sub> as also observed in the literature (Petry et al., 2003; Vaningelgem et al., 2004). The 247 EPS produced by the strain ST3 has a M<sub>w</sub> and intrinsic viscosity significantly higher than 248 those produced by cultures ST2 or ST1 (Fig. 3). Furthermore, EPS from ST3 consisted of 249 80% high M<sub>w</sub> polymers in number, in contrast to EPS from ST2, which contained only 35% 250 high M<sub>w</sub> molecules.

251 Regarding the strain ST2 and its co-cultures with ST1, the increase in the proportion of ST1 from 0 to 37% [culture ST1+2 (1/1.7)] and to 83 % [culture ST1+2 (5/1)] induced a 252 253 significant increase in the amount of EPS produced from 30 to ~52 and to ~67 mg glucose kg<sup>-</sup> 254 <sup>1</sup>, respectively (Fig. 2), while the time of acidification was significantly decreased (Table 1). 255 The same trend was also observed when ST1 was added to strain ST3. Furthermore, the 256 macromolecular characteristics of the EPS produced by ST2 and its co-cultures were broadly similar (Fig. 3A,D). For the strain ST3 and its co-cultures ST1+3, a significant decrease of 257 EPS M<sub>w</sub> was observed when the acidification time decreased (Pearson's r = 0.951) due to the 258 259 increase in ST1/ST3 ratio. These results suggested a chain length extension of EPS from ST3 during longer acidification time. Laws, Leivers, Chacon-Romero, and Chadha (2009) 260

reported a similar trend in which EPS M<sub>w</sub> produced by *Lactobacillus acidophilus* increased
3.7 times from 6 h to 24 h of fermentation in skim milk supplemented with glucose at 42 °C
and at a controlled pH of 5.8. In association with cell counting, the authors observed that EPS
synthesis and chain length extension continued after the end of the exponential phase, using
sugar nucleotides already produced during in the latter phase.

266 When looking in more details at the results of EPS M<sub>w</sub> distribution (in mass or in 267 number, Fig. 3B,C), different trends could be observed. Firstly, the presence of the acidifying 268 strain ST1 at a level of 37% in culture ST1+2 (1/1.7) and ST1+3 (1/1.7) did not change the 269 M<sub>w</sub> distribution characteristics as compared with pure strains (Fig. 3B,C), suggesting that in 270 these conditions ST1 produced a negligible quantity of EPS, but had an enhancing effect on 271 the EPS production of texture strains. In contrast, a significant change in the M<sub>w</sub> distribution 272 of EPS was observed with 83% ST1 in cultures ST1+2 (5/1) and ST1+3 (5/1), in that the low 273 M<sub>w</sub> EPS then became predominant. These findings could be related to the short acidification 274 time observed with these co-cultures and/or to the prevalence of the acidifying strain over the 275 texturing strain. Furthermore, we observed that the high M<sub>w</sub>EPS group from culture ST1+3 276 (5/1) presented a lower average M<sub>w</sub>, but a higher [ $\eta$ ] than for ST1+2 (5/1). These findings 277 suggested that the structure of EPS produced by ST2 could be more flexible and/or branched 278 than those produced by ST3. It has been shown that for two EPS with identical structure (i.e. 279 repeating units branching, stiffness of side chain), a higher  $M_w$  could lead to a higher [n] and 280 therefore to a higher gel viscosity (Faber, Zoon, Kamerling, & Vliegenthart, 1998; 281 Looijesteijn, van Casteren, Tuinier, Doeswijk-Voragen, & Hugenholtz, 2000; Mende et al., 2016). However, other structural factors such as side chain stiffness and branching also made 282 283 a considerable contribution to the  $[\eta]$  of EPS. Generally, for a given M<sub>w</sub>, a rigid linear 284 polysaccharide will tend to extend under shearing, as opposed to a flexible branched

molecules which will tend to compact, and will then contribute to increase the viscosity of thesolution (Whistler & BeMiller, 1997).

287 Overall, the results presented in this section showed that the acidification kinetics, the 288 production and the macromolecular characteristics of EPS depended not only on the strains 289 but also on the interaction between textural and acidifying strains. The combination between 290 ST1 and ST3 seemed to be more efficient than the combination between ST1 and ST2. 291 Furthermore, all pure cultures could produce EPS, but these EPS differed in terms of 292 quantity, distribution in mass or in number and intrinsic viscosity. In the same way, the 293 overall distribution of low and high M<sub>w</sub> EPS is strongly correlated with the interaction 294 between acidifying and texturing strains. It is quite difficult to compare these findings with 295 other studies because very few data are reported on the EPS production of mixtures of S. 296 thermophilus. Folkenberg et al. (2006) observed different trends of EPS production in 297 fermented milk by 5 S. thermophilus strains in pure or in mix cultures of two strains. Their 298 results suggested that the interaction between two S. thermophilus strains could be 299 compatible or not, depending on the strain used. However, they did not report acidification 300 time or molecular characteristics of EPS, which made difficult the comparison with our 301 results.

302

303 3.2. Influence of starter cultures on milk gelation and textural properties of gels

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305 3.2.1. Gelation kinetics

The kinetics of gelation were monitored by the evolution of the storage modulus (G') and the loss modulus (G''), characterised by time and pH of the gel point. As presented in Table 1, the time and pH of gelation depended on the culture used. The pH of the gel point was approximately 5.2 for the cultures ST1, ST2, ST1+2 (1/1.7) and ST1+2 (5/1), and was

310 not influenced by the rate of acidification. The same pH value was reported for the gel point 311 of heated milk acidified chemically using glucono-delta-lactone (Lucey, Tamehana, Singh, & 312 Munro, 1998; Tamime et al., 2001). In the case of ST3, the pH of the gel point was found to 313 be significantly higher at 5.57 and decreased to 5.4 and 5.2 when strain ST1 was added at 37 314 and 83%, respectively (Table 1). The occurrence of the gel point at a higher pH could be 315 related to the presence of EPS and their interactions with milk proteins (Mende et al., 2013b). 316 In the case of strain ST3 and its co-cultures, a positive correlation (Pearson's r =317 0.992) was observed between the pH of gelation and EPS macromolecular properties (M<sub>w</sub>, 318  $[\eta]$ ), but not with the EPS quantity. The decrease in the pH of the gel point could be related 319 to a decrease of the average M<sub>w</sub> of EPS and/or of the proportion of high M<sub>w</sub> EPS (Fig. 3). On 320 the other hand, for all cultures, a predominant proportion in number of low Mw EPS 321 corresponded to a gel point occurring at a lower pH of ~5.2 (Fig. 3C). Mende et al. (2013a,b) 322 also reported a similar trend. The authors observed early gelation due to the depletion effect 323 between casein micelles and added bacterial EPS or commercial dextran (Tuinier, Grotenhuis, Holt, Timmins, & de Kruif, 1999). These two neutral polysaccharides, bacterial 324 EPS and dextran had different  $M_w$  of 2.6  $\times 10^6$  Da and 5 $\times 10^5$  Da, respectively. Compared with 325 bacterial EPS, addition of 15 - 80 times more dextran was required to achieve a similar effect 326 327 on gelation point.

328

329 *3.2.2. Gel texture* 

Gel texture was characterised by four parameters: (i) firmness, (ii) viscosity, (iii) spontaneous syneresis (Table 2) and (iv) graininess (Fig. 4). Gels from the ST3 culture are not mentioned in this section because after 24 h of acidification, the gels obtained at a final pH of ~4.8 were heterogeneous with an upper gel-like part and a lower liquid-like part.

335 *Firmness.* Gel firmness is the main texture property of set-type acid milk gel and this 336 parameter is mainly governed by the protein network and not by bacterial EPS (Folkenberg et 337 al., 2006; van Marle & Zoon, 1995). For the monocultures used in the present study, ST1 338 provided the lowest gel firmness while ST2 produced the highest gel firmness, which could 339 be related to its long acidification time of nearly 24 h. Moreover, it was observed for strain 340 ST2 and its co-cultures ST1+2 that the higher the proportion of ST1, the faster the 341 acidification and the lower the gel firmness (Pearson's r = 0.998). According to the literature, 342 slow protein aggregation could result in continuous network formation with more connected 343 proteins, reducing the rearrangement of particles during gel formation and contributing to an 344 increase in gel rigidity during fermentation at low temperatures (Lucey & Singh, 1997; 345 Sodini et al., 2004).

In the present study, it is interesting to note that although the cultures ST1, ST1+2 (5/1) and ST1+3 (5/1) induced similar acidification times (Table 1), the resulting gel firmness was significantly different in the order: ST1+2 (5/1) > ST1+3 (5/1) > ST1, consistent with the order of the average M<sub>w</sub> of EPS produced. In other words, the M<sub>w</sub> of EPS from ST1+2 (5/1) was higher than from those of ST1+3 (5/1) and ST1 (Pearson's r = 0.987). These findings suggested that the presence of high M<sub>w</sub> EPS was also an important determining parameter for gel firmness, though not as much as the acidification rate.

When comparing the gels of co-cultures with the same ST1 ratio, the gels containing ST2 had a higher firmness than those containing ST3 (Table 2). Other studies have also reported a decreased gel firmness when using a ropy culture which is defined as a high EPS producing culture with high [η] such as the culture ST3 (Hassan et al., 2003; Hess, Roberts, Ziegler, 1997; Mende et al., 2012). Using microscopic analyses, Hassan et al. (2003) observed that the EPS of ropy cultures were mainly located and segregated in the pores of the gel rather than attached to the protein network. Hence, the gel structure from a ropy culture was less dense than that of non-ropy culture. Mende et al. (2012) and Rohm and Kovac
(1994) also suggested that the large amount of uncharged EPS produced by ropy culture
could have partially prevented the development of protein-protein bonds during gel formation
and thus reduced the rigidity of acid milk gel.

364

365 Stirred gel viscosity. Stirred gel viscosity is correlated with a thick mouthfeel and with 366 the smoothness of fermented milk products (Folkenberg et al., 2006). In this study, the stirred 367 gel viscosity varied depending on the culture and the macromolecular properties of the 368 produced EPS. ST1 produced only low  $M_w$  EPS with low  $[\eta]$  and formed gels presenting the 369 lowest viscosity. When ST2 was used, even if a low quantity of EPS was produced, the 370 viscosity of the stirred gels formed was higher than for strain ST1. This result could be 371 explained by the high percentage (80%) in mass of EPS with high Mw produced by culture 372 ST2.

373 Concerning the co-cultures of ST2 or ST3, increasing the ratio of the acidifying strain 374 decreased the stirred gel viscosity from 193 to 164 mPa s for ST1+2(1/1.7) and (5/1), 375 respectively, or from 301 to 239 mPa s for ST1+3 (1/1.7) and (5/1), respectively (Table 2). 376 These results could be directly linked to the macromolecular properties of the EPS produced 377 (Fig. 3) and suggested a positive correlation between the quantity of high M<sub>w</sub> EPS and the 378 stirred gel viscosity (Pearson's r = 0.890). Petry et al. (2003) also reported a striking 379 correlation between an elevated proportion of the high M<sub>w</sub> EPS fraction and viscosity of acid 380 milk gel. In the present study, we also observed that even though the amount and molar mass 381 distribution of EPS were similar, gels from culture ST1+3 (1/1.7) were more viscous than 382 those from culture ST1+2 (1/1.7). This result could be partially explained by the higher  $[\eta]$  of 383 the EPS produced by ST3 and probably by a different molecular rigidity as already mentioned in section 3.1.2. Also when comparing ST1+2 (5/1) and ST1+3 (5/1), despite both 384

385 co-cultures exhibiting a low proportion of high  $M_w$  EPS and similar [ $\eta$ ] of EPS, the viscosity 386 of stirred gels was significantly different, suggesting that EPS were not the only factor 387 contributing to this textural property. In a previous study based on gel microstructure 388 observations, Nguyen et al. (2018a) suggested that in parallel to EPS production, the 389 morphology of bacterial cell chain length should also be taken into account to understand the 390 texture results.

391

392 Spontaneous syneresis defect. Spontaneous syneresis is the contraction of a gel 393 without exerting external forces and is related to an instability of the gel network that results 394 in the loss of ability to entrap the entire serum phase (Lucey & Singh, 1997). Gels obtained 395 with pure culture of ST1 presented a medium level of syneresis while those from ST2 had the 396 highest syneresis, possibly related to the long acidification time. For co-cultures of ST2, this 397 texture defect decreased from 2.7 to 1.4% when the ratio of ST1/ST2 changed from 1/1.7 to 398 5/1. In this case, a positive correlation was observed between syneresis, gel firmness and time 399 of acidification (Pearson's r = 0.999) suggesting that gel contraction could occur during the 400 slow fermentation, thereby hardening the gel and expelling the serum. In contrast, low 401 syneresis levels were observed for co-culture of ST1+3. Decreased gel syneresis has been 402 reported with the use of ropy strains that produce a high quantity of high M<sub>w</sub> EPS 403 (Amatayakul et al., 2006; Kristo, Miao, & Corredig, 2011). It can be assumed that the ropy 404 EPS located in the gel pores could bind the serum and thus reduce the syneresis.

405

406 *Graininess defect.* Graininess or lumpiness is considered an undesirable defect
407 because consumers usually expect a smooth product (Lucey, 2004). Remeuf et al. (2003)
408 measured graininess by the number of grains with a diameter above 1 mm in 1 g of acid milk
409 gel. In the present study, the graininess defect was determined by the number of grains, but

also by their diameter from 1 mm to above 3 mm. Larger grains were more damaging for
texture perception even in low number (Fig. 4). The graininess was high when using pure
cultures ST1 and ST2. In the mixtures, regardless of the strain ratio, co-cultures of ST3
presented a low graininess while the co-cultures of ST2 showed a medium granular texture
(Fig. 4).

415 According to the literature, grains are large aggregates of proteins that appear under 416 certain conditions such as a high fermentation temperature, excessive whey protein to casein 417 ratio or with the use of certain types of starter culture (Lucey, 2004). However, only few 418 studies have focused on the impact of starter culture on the appearance of grain. In agreement 419 to our result, Küçükçetin et al. (2009) reported that the number of grains and their size were 420 significantly reduced when using a high-level EPS-producing starter culture, rather than a 421 medium- or low-level EPS-producing culture. Results from the present study provided 422 additional data on this phenomenon. Indeed, it is worth noting that cultures ST1+2 (5/1) and 423 ST1+3 (5/1) had a similar kinetics of acidification and gelation (Table 1) and similar 424 production of EPS (Fig. 2) that presented identical intrinsic viscosity and EPS molar mass 425 distribution (Fig. 3 B,C,D). However, despite these similarities, the texture of the resulting 426 gels from ST1+2 (5/1) presented more major defects (syneresis and graininess) as compared 427 with gels from culture ST1+3 (5/1). These new data support the inhibitory effect of strain 428 ST3 against appearance of texture defects, specifically graininess, which has been observed 429 in a previous study (Nguyen et al., 2018a). Therefore, we suggested that this interesting 430 technological property of culture ST3 could be related to the bacterial chains morphology 431 and/or the structural properties of the EPS produced and that it did not depend on the acidification kinetics or final EPS content. 432

#### 434 **4.** Conclusion

435

It is known that the symbiosis between texturing and acidifying strains of *S*. *thermophilus* reduces the acidification time and has an impact on the final acid milk gel structure. In this work, we selected one acidifying strain (ST1) and two texturing strains (ST2, ST3) strains, and used them singly or in co-cultures (ST1+2 or ST1+3) at two different ratios (1/1.7 or 5/1) in pasteurised skim milk to assess and understand the influence of their interaction on the acidification kinetics, EPS production, gel structure and appearance of texture defects.

In pure culture, due to the presence of proteinases in the cell wall, the acidifying strain provided rapid acidification and produced a medium amount of EPS. However, these EPS had a low  $M_w$  and intrinsic viscosity [ $\eta$ ] hence the gel obtained had the lowest firmness and viscosity, and the most graininess defect. Both texturing strains did not have the proteinase activity and showed a slow acidification. They both produced EPS with high molar mass but with different macromolecular properties.

449 In co-culture, the strains ST2 and ST3 had positive symbiotic relations with ST1, as 450 showed by the reduction in acidification time and the increase in EPS production. 451 Furthermore, the viscosity of gels increased significantly, probably due to the production of 452 high  $M_w$  EPS by the texturing strains. Texture defects such as syneresis and graininess 453 disappeared with use of the co-cultures of ST1+3. Moreover, co-cultures ST1+3 (5/1) and 454 ST1+2 (5/1) had some similarities in terms of (i) acidification, (ii) gelation kinetics, (iii) EPS 455 production and molar mass distribution and (iv) gel firmness. However, despite these 456 similarities, gels from co-culture ST1+2 (5/1) showed worse texture defects as compared with 457 ST1+3 (5/1).

458	These results highlight the fact that the inhibitory effect of a strain against the
459	appearance of texture defects is not correlated to the acidification rate or EPS content, but
460	that it could instead be explained by differences in the structural properties of EPS and/or in
461	the morphology of bacterial chains.
462	
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471	References
472	
473	Amatayakul, T., Sherkat, F., & Shah, N. P. (2006). Syneresis in set yogurt as affected by EPS
474	starter cultures and levels of solids. International Journal of Dairy Technology, 59, 216-
475	221.
476	Faber, E. J., Zoon, P., Kamerling, J. P., & Vliegenthart, J. F. G. (1998). The
477	exopolysaccharides produced by Streptococcus thermophilus Rs and Sts have the same
478	repeating unit but differ in viscosity of their milk cultures. Carbohydrate Research, 310,
479	269–276.
480	Folkenberg, D. M., Dejmek, P., Skriver, A., & Ipsen, R. (2006). Interactions between EPS-
481	producing Streptococcus thermophilus strains in mixed yoghurt cultures. Journal of
482	Dairy Research, 73, 385–393.

- 483 Galia, W., Perrin, C., Genay, M., & Dary, A. (2009). Variability and molecular typing of
- 484 *Streptococcus thermophilus* strains displaying different proteolytic and acidifying
  485 properties. *International Dairy Journal*, *19*, 89–95.
- 486 Hassan, A. N., Ipsen, R., Janzen, T., & Qvist, K. B. (2003). Microstructure and rheology of
- 487 yogurt made with cultures differing only in their ability to produce exopolysaccharides.
- 488 *Journal of Dairy Science*, 86, 1632–1638.
- 489 Hess, S. J., Roberts, R. F., & Ziegler, G. R. (1997). Rheological properties of nonfat yogurt
- 490 stabilized using *Lactobacillus delbrueckii* ssp. *bulgaricus* producing exopolysaccharide
- 491 or using commercial stabilizer systems. *Journal of Dairy Science*, 80, 252–263.
- 492 Kristo, E., Miao, Z., & Corredig, M. (2011). The role of exopolysaccharide produced by
- 493 *Lactococcus lactis* subsp. *cremoris* in structure formation and recovery of acid milk gels.
  494 *International Dairy Journal*, *21*, 656–662.
- Küçükçetin, A., Weidendorfer, K., & Hinrichs, J. (2009). Graininess and roughness of stirred
  yoghurt as influenced by processing. *International Dairy Journal*, *19*, 50–55.
- 497 Lambo-Fodje, A. M., Leeman, M., Wahlund, K. G., Nyman, M., Öste, R., & Larsson, H.
- 498 (2007). Molar mass and rheological characterisation of an exopolysaccharide from
- 499 *Pediococcus damnosus* 2.6. *Carbohydrate Polymers*, 68, 577–586.
- 500 Laws, A. P., Leivers, S., Chacon-Romero, M., & Chadha, M. J. (2009). Variation in the
- 501 molecular mass of exopolysaccharides during the time course of extended fermentations
- 502 of skimmed milk by lactic acid bacteria. *International Dairy Journal*, *19*, 768–771.
- 503 Looijesteijn, P. J., van Casteren, W. H., Tuinier, R., Doeswijk-Voragen, C. H., &
- 504 Hugenholtz, J. (2000). Influence of different substrate limitations on the yield,
- 505 composition and molecular mass of exopolysaccharides produced by *Lactococcus lactis*
- subsp. *cremoris* in continuous cultures. *Journal of Applied Microbiology*, 89, 116–122.
- 507 Lucey, J. A. (2004). Cultured dairy products: an overview of their gelation and texture

- 508 properties. *International Journal of Dairy Technology*, 57, 77–84.
- Lucey, J. A., & Singh, H. (1997). Formation and physical properties of acid milk gels: A
  review. *Food Research International*, *30*, 529–542.
- 511 Lucey, J. A., Tamehana, M., Singh, H., & Munro, P. A. (1998). Effect of interactions
- between denatured whey proteins and casein micelles on the formation and rheological
- 513 properties of acid skim milk gels. *Journal of Dairy Research*, 65, 555–567.
- 514 Marshall, V. M., & Rawson, H. L. (1999). Effects of exopolysaccharide-producing strains of
- thermophilic lactic acid bacteria on the texture of stirred yoghurt. *International Journal of Food Science & Technology*, *34*, 137–143.
- 517 Mende, S., Mentner, C., Thomas, S., Rohm, H., & Jaros, D. (2012). Exopolysaccharide
- 518 production by three different strains of *Streptococcus thermophilus* and its effect on
- 519 physical properties of acidified milk. *Engineering in Life Sciences*, *12*, 466–474.
- 520 Mende, S., Peter, M., Bartels, K., Dong, T., Rohm, H., & Jaros, D. (2013a). Concentration
- dependent effects of dextran on the physical properties of acid milk gels. *Carbohydrate Polymers*, 98, 1389–1396.
- 523 Mende, S., Peter, M., Bartels, K., Rohm, H., & Jaros, D. (2013b). Addition of purified
- exopolysaccharide isolates from *Streptococcus thermophilus* to milk and their impact on
  the rheology of acid gels. *Food Hydrocolloids*, *32*, 178–185.
- 526 Mende, S., Rohm, H., & Jaros, D. (2016). Influence of exopolysaccharides on the structure,
- 527 texture, stability and sensory properties of yoghurt and related products. *International*528 *Dairy Journal*, 52, 57–71.
- 529 Nguyen, A. T.-B., Nigen, M., Jimenez, L., Ait-Abderahim, H., Cunault, C., Marchesseau, S.,
- 530 et al. (2018a). A multi-scale approach to identify the role of heat treatment, milk protein
- 531 composition and starter culture on the gel formation and the texture defects of acid milk
- 532 gel. *Food Hydrocolloids*, 85, 299–310.

533	Nguyen, A. TB., Nigen, M., Jimenez, L., Ait-Abderrahim, H., Marchesseau, S., & Picart-
534	Palmade, L. (2018b). Performances of different protocols for exocellular
535	polysaccharides extraction from milk acid gels: Application to yogurt. Food Chemistry,
536	239, 742–750.
537	Petry, S., Furlan, S., Waghorne, E., Saulnier, L., Cerning, J., & Maguin, E. (2003).

- 538 Comparison of the thickening properties of four *Lactobacillus delbrueckii* subsp.
- 539 *bulgaricus* strains and physicochemical characterization of their exopolysaccharides.

540 FEMS Microbiology Letters, 221, 285–291.

- 541 Remeuf, F., Mohammed, S., Sodini, I., & Tissier, J. P. (2003). Preliminary observations on
- 542 the effects of milk fortification and heating on microstructure and physical properties of
- 543 stirred yogurt. *International Dairy Journal*, *13*, 773–782.
- Rohm, H., & Kovac, A. (1994). Effects of starter cultures on linear viscoelastic and physical
  properties of yogurt gels. *Journal of Texture Studies*, 25, 311–329.
- 546 Sodini, I., Remeuf, F., Haddad, S., & Corrieu, G. (2004). The relative effect of milk base,
- starter, and process on yogurt texture: A Review. *Critical Reviews in Food Science and Nutrition*, 44, 113–137.
- 549 Tamime, A. Y., Robinson, R. K., & Latrille, E. (2001). Yoghurt and other fermented milks.
- 550 In A. Y. Tamime & B. A. Law (Eds.), *Mechanisation and automation in dairy*
- 551 *technology* (pp. 152–203). Sheffield, UK: Sheffield Academic Press.
- 552 Tuinier, R., Grotenhuis, E. T., Holt, C., Timmins, P. A., & de Kruif, C. G. (1999). Depletion
- interaction of casein micelles and an exocellular polysaccharide. *Physical Review E*, 60,
  848–856.
- van Marle, M. E., & Zoon, P. (1995). Permeability and rheological properties of microbially
  and chemically acidified skim-milk gels. *Netherlands Milk and Dairy Journal*, 49, 47–
- 557 65

- 558 Vaningelgem, F., Zamfir, M., Mozzi, F., Adriany, T., Vancanneyt, M., Swings, J., et al.
- 559 (2004). Biodiversity of exopolysaccharides produced by *Streptococcus thermophilus*
- 560 strains is reflected in their production and their molecular and functional characteristics.
- 561 *Applied and Environmental Microbiology*, 70, 900–912.
- 562 Whistler, R. L., & BeMiller, J. N. (1997). Carbohydrate chemistry for food scientists. St.
- 563 Paul, MI, USA: Eagan Press.
- 564 Winter, H. H. (1987). Can the gel point of a cross-linking polymer be detected by the G' G''
- 565 crossover? *Polymer Engineering & Science*, 27, 1698–1702.

#### **Figure legends**

Fig. 1. Kinetics of acidification monitored during skim milk fermentation by cultures ST1 (- - -), ST2 (----), ST1+2 (1/1.7) (----), ST1+2 (5/1) (----), ST3 (-----), ST1+3 (1/1.7)
(-----), ST1+3 (5/1) (----). Skim milk suspensions were pasteurised at 95 °C for 6 min.

**Fig. 2.** Quantity of EPS (mg glucose kg<sup>-1</sup>) produced by the 7 cultures/co-cultures (ST1, ST2, ST1+2 (1/1.7), ST1+2 (5/1), ST3, ST1+3 (1/1.7), ST1+3 (5/1)) during fermentation of skim milk pasteurised at 95 °C for 6 min. Mean values  $\pm$  standard deviation (n = 3). Values affected with different letters were significantly different for p = 0.05.

**Fig. 3.** Macromolecular properties of extracted EPS from the 7 cultures/co-cultures (ST1, ST2, ST1+2 (1/1.7), ST1+2 (5/1), ST3, ST1+3 (1/1.7), ST1+3 (5/1): A, average molar mass; B, molar mass distribution in mass; C, molar mass distribution in number of particles; D, intrinsic viscosity. The EPS were classified in high molar mass ( $\Box$ ) or low molar mass ( $\Box$ ).

**Fig. 4.** Graininess defect of acid milk gels made by 6 different cultures [ST1, ST2, ST1+2 (1/1.7), ST1+2 (5/1), ST1+3 (1/1.7), ST1+3 (5/1)]. Grains were classified by their diameters: 1–1.5 mm ( $\Box$ ), 1.5–2 mm ( $\Box$ ), 2–2.5 mm ( $\blacksquare$ ), 2.5–3 mm ( $\boxdot$ ) and > 3 mm ( $\Box$ ). Mean values ± standard deviation (n = 9).



Figure 1



Figure 2





Figure 4

#### Table 1

Acidification parameters of skim acid milk gel obtained using 3 pure strain cultures (ST1,

ST2, ST3) and 4 co-cultures [ST1+2 (1/1.7	, ST1+2 (5/1), ST1+3 (1/	1.7), ST1+3 (5/1)]. <sup>2</sup>
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P	arameters	ST1	ST2	ST1+ST2		ST3	ST1+ST3	
-			~	1/1.7	5/1		1/1.7	5/1
А	cidification							
	Acidification time (min)	$348 \pm 49^{ad}$	$1351 \pm 109^{b}$	$528 \pm 33^{\circ}$	$304 \pm 24^{a}$	ND	$395 \pm 27^{d}$	$287 \pm 11^{a}$
	$V_{\rm max}$ (10 <sup>-3</sup> pH unit min <sup>-1</sup> )	$17.1 \pm 1.7^{a}$	$14.0 \pm 0.3^{b}$	$16.9 \pm 1.3^{a}$	$20.0\pm0.3^{\rm c}$	$6.5 \pm 0.1^{d}$	$18.1 \pm 1.7^{ac}$	$20.0 \pm 0.1^{\circ}$
G	elation							
	Time of gel point (min)	$206 \pm 13^{a}$	388 ±36°	$208 \pm 10^{a}$	$157 \pm 11^{b}$	371 ±18°	179 ± 11 <sup>b</sup>	$164 \pm 10^{b}$
	pH of gel point	$5.16 \pm 0.05^{a}$	5.15 ±0.05 <sup>a</sup>	$5.23 \pm 0.01^{a}$	$5.21 \pm 0.05^{a}$	$5.57 \pm 0.01^{\circ}$	$5.39 \pm 0.06^{b}$	$5.22 \pm 0.01^{a}$

<sup>a</sup> Reconstituted skim milk was pasteurised at 95 °C, 6 min.  $V_{\text{max}}$  is the maximum acidification rate. Data were obtained by monitoring the pH evolution (acidification) and rheological parameters (gelation) during fermentation at 43 °C; values of each characteristic affected with different letters were significantly different for p = 0.05 (ND, not determined).

#### Table 2

Cultures	ST1	ST2	ST1+ST2		ST1+ST3	
			1/1.7	5/1	1/1.7	5/1
Firmness (N)	$3.36 \pm 0.18^{a}$	$6.44 \pm 0.15^{\text{ d}}$	$4.78 \pm 0.08$ °	4.14 ± 0.13 <sup>b</sup>	4.43 ± 0.25 <sup>b</sup>	$3.85 \pm 0.11^{\text{ e}}$
Viscosity (mPa s) at $\gamma = 300 \text{ s}^{-1}$	$138 \pm 12^{a}$	$186 \pm 4^{\circ}$	$193 \pm 3^{\circ}$	$164 \pm 2^{b}$	$301 \pm 10^{e}$	$239 \pm 15^{d}$
Spontaneous syneresis (%, w/w)	$1.8 \pm 0.3^{a}$	$6.3 \pm 0.3^{\circ}$	$2.7 \pm 0.4^{b}$	$1.4 \pm 0.4^{a}$	$0.4 \pm 0.2^{e}$	0 <sup>d</sup>

Textural parameters of pasteurised acid milk gel using 6 different cultures <sup>a</sup>

<sup>a</sup> Reconstituted skim milk was pasteurised at 95 °C for6 min; values of each characteristic affected with different letters were significantly different at p = 0.05.