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The Effect of Fermentation Temperature on the Microstructure, Physicochemical and Rheological Properties of Probiotic Buffalo Yoghurt

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16 **ABSTRACT**

17 The properties of buffalo and bovine milk differ and the procedures developed to make  
18 bovine yoghurt may require optimisation for the production of buffalo yoghurt. This study  
19 aimed to apply cryo scanning electron microscopy and confocal laser scanning microscopy to  
20 determine the optimal temperature for processing buffalo yoghurt. Milk was fermented at  
21 three different temperatures (37°C, 40°C and 43°C), stored for 28 days and the yoghurt  
22 microstructure, physicochemical and rheological properties assessed. Yoghurt fermented at  
23 37°C had a compact microstructure and the probiotic *Lactobacillus acidophilus* La-5 was  
24 more viable on storage. In contrast, yoghurt produced from a faster fermentation at 43°C was  
25 firmer with a more porous microstructure that exhibited a higher degree of syneresis. The  
26 rheological properties during storage including the thixotropy, consistency coefficient and  
27 flow behaviour index were not significantly affected by temperature nor was the  
28 concentration of lactose, ionic calcium or titratable acidity. This study shows how changes to  
29 processing can be used to alter the microstructure of buffalo products and suggests that a  
30 decrease in fermentation temperature could be used to improve the quality of buffalo yoghurt.

31 **KEY WORDS:** buffalo yoghurt, fermentation temperature, syneresis, microstructure,  
32 rheological properties

33

## 34 1. Introduction

35 Buffalo milk is significantly different to bovine milk in both chemical composition and  
36 physicochemical properties. These differences lead to advantages but also disadvantages  
37 during milk processing. Benefits include a higher yield of cheese, cream, ghee and butter  
38 (Menard et al. 2010), faster separation of cream, as well as easier churning and reduced fat  
39 loss during butter production (Sahai 1996). Yoghurt production is simpler without the need  
40 for fortification with milk powder or the addition of thickeners or stabilisers (Addeo et al.  
41 2007). Drawbacks include an acceleration of the Maillard browning reaction during  
42 pasteurisation or sterilisation (Sahai 1996) and a higher buffering capacity that results in  
43 slower acidification and longer fermentation during the production of cheese and yoghurt  
44 (Ahmad et al. 2008). The larger fat globules and the higher fat content in buffalo milk also  
45 lead to a more porous yoghurt microstructure that has a high degree of syneresis; a major  
46 defect that requires the optimisation of processing conditions (Nguyen et al. 2013).

47 Much is known about the factors that affect gel formation and syneresis for bovine yoghurt,  
48 although the effect of these variables on buffalo yoghurt is not well understood. The  
49 fermentation temperature, starter culture type, starter culture concentration and milk base used  
50 all affect yoghurt production from bovine milk (Abbasi et al. 2009; Folkenberg et al. 2004;  
51 Lucey et al. 1998b; Lucey et al. 1998a; McClements 2007; Sodini et al. 2004; Xu et al. 1992).  
52 Among these, fermentation temperature is considered most significant, due to the significant  
53 impact of temperature on gel formation and acidification rate (Lee and Lucey 2003; Sodini et  
54 al. 2004; Wu et al. 2009; Lee and Lucey 2004; Purwandari et al. 2007; Tamime and Robinson  
55 2007; Laligant et al. 2003). A fermentation temperature of 37°C - 45°C is typically selected  
56 for the production of bovine yoghurt to achieve optimal growth of the mixed bacterial starter  
57 cultures, such as *Streptococcus thermophilus* and *Lactobacillus bulgaricus* (Purwandari et al.  
58 2007; Sfakianakis and Tzia 2011; Paseephol et al. 2008). While faster fermentation at a higher  
59 fermentation temperature may be advantageous in industrial production, it can lead to several  
60 defects, such as an increase in whey separation (Lee and Lucey 2003; Purwandari et al. 2007;  
61 Lee and Lucey 2004), a decrease in gel firmness, viscosity and smoothness, a decrease in  
62 desirable sensory properties (Tamime and Robinson 2007; Wu et al. 2009), and a weaker  
63 protein network with a coarser microstructure (Lee and Lucey 2004; Lucey and Singh 1997).  
64 Conversely, the physical and sensory properties of bovine yoghurt can be improved when a

65 lower fermentation temperature of 32°C - 39°C is employed (Sodini et al. 2004; Martin et al.  
66 1999), although these improvements come at the cost of increased production time.

67 Research on buffalo yoghurt, especially on factors affecting the syneresis such as the  
68 fermentation temperature, is still limited. Buffalo yoghurt has long been produced in countries  
69 such as India and Pakistan using traditional processing technology (Ahmad et al. 2008) but  
70 the only readily accessible prior study is by Shiby and Mishra (2008), who examined the  
71 simultaneous effects of fermentation temperature, starter culture and milk total solids  
72 concentration on the rate of acidification, firmness and syneresis of buffalo yoghurt.  
73 Unfortunately, the use of a mesophilic lactic acid bacteria that necessitates overnight  
74 incubation limits the application of this study to commercial yoghurt production where short  
75 incubations are preferred (Chandan and O'Rell 2006; Tamime and Robinson 2007). Most  
76 other studies have focused on the fortification of buffalo yoghurt (Kumar and Mishra 2003;  
77 Ghadge 2008) or yoghurt production from a mixture of buffalo and other milk types (Bezerra  
78 et al. 2012). Further, buffalo yoghurt is usually prepared with unhomogenised and  
79 unstandardised milk (Tamime and Robinson 2007; Addeo et al. 2007), which is different to  
80 the normal case for bovine yoghurt. These attributes together with the markedly different  
81 chemical composition and properties of buffalo milk affect the processing and quality of the  
82 resulting dairy products. Consequently, the impact of fermentation temperature on buffalo  
83 yoghurt may be substantially different. Additionally, the fermentation required for buffalo  
84 yoghurt takes longer than for bovine yoghurt. For this reason, it is important to determine an  
85 optimum fermentation temperature for this product that will balance the extended production  
86 time and the quality of the yoghurt. This forms the focus of the present paper.

87 This study aims to investigate the effect of three fermentation temperatures (37°C, 40°C and  
88 43°C) on the properties of probiotic buffalo yoghurt during fermentation and cold storage.  
89 The second aim is to apply microscopy tools known to preserve the microstructure of  
90 hydrated gels to investigate the microstructure of buffalo yoghurt as a function of  
91 fermentation temperature.

## 92 **2. Materials and methods**

### 93 *2.1 Yoghurt preparation*

94 Buffalo yoghurt was produced from raw buffalo milk provided by a local dairy farmer in  
95 Shaw River (Victoria, Australia) with chemical composition of  $4.1 \pm 0.4$  (% w/w) protein, 7.9

96  $\pm 0.3$  (% w/w) fat,  $5.0 \pm 0.2$  (% w/w) lactose and  $17.1 \pm 0.4$  (% w/w) total solids. Four litres  
97 of the buffalo milk were batch-pasteurised (85°C, 30 min) using a water bath (Qualtex,  
98 Watson Victor Ltd., Australia). The pasteurised milk was cooled to three different  
99 fermentation temperatures of 37°C, 40°C or 43°C before inoculation with  $0.062 \text{ g.L}^{-1}$  freeze  
100 dried direct vat starter culture ABT-5 containing probiotic *Lactobacillus acidophilus La-5*,  
101 *Bifidobacterium lactis Bb-12* and *Streptococcus thermophilus* (CHR-Hansen, Bayswater,  
102 Victoria, Australia). The inoculated milk was distributed into several plastic containers and  
103 fermented at 37°C, 40°C or 43°C in three water baths containing water that had been tempered  
104 to the appropriate fermentation temperature. The fermentation was terminated when the milk  
105 reduced to a pH of 4.5. Two trials of yoghurt were produced for each fermentation  
106 temperature on different days.

## 107 2.2 Chemical analysis

### 108 2.2.1 Measurement of pH and ionic calcium concentration

109 The changes in pH during fermentation were measured using an electrode pH meter (Orion  
110 720A plus, Orion Pacific Pty Ltd., Victoria, Australia) while the changes in ionic calcium  
111 concentration were determined using an Orion 93 - 20 calcium half-cell electrode in  
112 conjunction with an Orion 90 - 02 Ag/AgCl double junction reference electrode (Orion  
113 Pacific, Victoria, Australia) as previously described (Nguyen et al. 2013). Three independent  
114 samples per trial were used for the measurement of pH and ionic calcium concentration at  
115 each time point during the fermentation. Two trials of yoghurt production were carried out for  
116 each fermentation temperature.

### 117 2.2.2 Determination of fat, protein, lactose, total solids and titratable acidity

118 The concentration of milk fat and protein was determined following the methods previously  
119 described by Atwood and Hartmann (1992), Pesce and Strand (1973) and modified by  
120 Nguyen et al. (2013) using a spectrophotometer (Fluostar Optima, BMG labtech, Ortenberg,  
121 Germany). Concentration of lactose was analysed following a previous method (Gosling et al.  
122 2009) using an HPLC Shimadzu Prominence system (NSW, Australia) equipped with a  
123 RID-10A refractive index detector and a 300 x 7.8 mm Rezex RCM-Monosaccharide  $\text{Ca}^{2+}$   
124 column (Phenomenex, NSW, Australia). Total solids and titratable acidity were determined  
125 using the methods of Association of Official Analytical Chemists (AOAC 2006). Three  
126 independent samples per trial were used for the analysis of lactose and titratable acidity at

127 each time point during the fermentation while fat, protein and total solids content of the milk  
128 used for yoghurt production were analysed in triplicates in each trial. Two trials of yoghurt  
129 production were carried out for each fermentation temperature.

### 130 *2.3. Syneresis determination*

131 Syneresis of yoghurt was determined following the method described previously (Purwandari  
132 et al. 2007) using a bench-top centrifuge (Eppendorf 5810R, VIC, Australia). Syneresis was  
133 expressed as a weight percentage of the whey separated from the gel over the initial weight of  
134 the gel. Three independent samples per trial were used for the determination of syneresis at  
135 each time point during storage. Two trials of yoghurt production were carried out for each  
136 fermentation temperature.

### 137 *2.4 Texture analysis*

138 The texture of yoghurt was analysed following a previously described method (Nguyen et al.  
139 2013) using a TA.XT-2 texture analyser (Stable Microsystems, Surrey, England) equipped  
140 with 2 kg load cell and a 10 mm diameter cylindrical probe. The contact area was set at 1  
141 mm<sup>2</sup> and the contact force set at 5 g. The instrument speed was set at 1 mm.s<sup>-1</sup>. The  
142 compression distance, the distance of penetration from the surface of the sample, was set at 20  
143 mm. Data were recorded at a rate of 200 points per second. Three independent samples per  
144 trial were used for the determination of texture at each time point during storage. Two trials of  
145 yoghurt production were carried out for each fermentation temperature.

### 146 *2.5 Rheological analysis*

#### 147 *2.5.1 Rheological properties (storage modulus G') during fermentation*

148 Rheological properties during fermentation were determined using a controlled strain  
149 rheometer (Advanced Rheometrics Expansion System, TA Instruments, New Castle, U.S.A.)  
150 equipped with a cup 34 mm in diameter and a six-blade vane fixture 32 mm in diameter and  
151 33 mm in height as previously described (Nguyen et al. 2013). A total of two rheological  
152 analyses were performed for each temperature treatment.

153 2.5.2 Rheological properties (thixotropy, flow behaviour index and consistency index)  
154 during storage

155 The rheological properties of yoghurt during storage were investigated following a previously  
156 described method (Purwandari and Vasiljevic 2009) using a controlled stress rheometer (AR-  
157 G2, TA instruments Ltd., New Castle, U.S.A.) fitted with a cone plate (40 mm diameter / 4°  
158 angle). Three independent samples per trial were used for the rheological analysis at each time  
159 point during storage. Two trials of yoghurt production were carried out for each fermentation  
160 temperature.

161 2.6 *Microstructural analysis using confocal laser scanning microscopy (CLSM) cryo*  
162 *scanning electron microscopy (cryo-SEM)*

163 The CLSM analysis was carried out using an inverted confocal scanning laser microscope  
164 (Leica TCS SP2; Leica Microsystems, Heidelberg, Baden-Wurttemberg, Germany) with  
165 sample preparation for CLSM analysis described in details in our previous work (Nguyen et  
166 al. 2013). The cryo-SEM analysis was performed using a field emission scanning electron  
167 microscope (Quanta, Fei Company, Hillsboro, Oregon, U.S.A.) as previously described by  
168 Ong and co-authors (2011). Two images were taken for each yoghurt sample in each trial.  
169 Two trials of yoghurt were carried for each fermentation temperature and hence, a total of  
170 four images were collected for each sample and a typical image is presented.

171 2.7 *Microbiological analysis*

172 The growth and viability of bacteria during fermentation and storage were assessed using the  
173 pour plate technique and different selective media as previously described (Nguyen et al.  
174 2013). Two plates with 25 - 250 colonies were selected for manual counting per trial for each  
175 yoghurt sample. Two trials of yoghurt production were carried out for each fermentation  
176 temperature.

177 2.8 *Statistical analysis*

178 Data were analysed using statistical Minitab software (V16, Minitab Inc., Stage College, PA,  
179 U.S.A.). Two way and one way analysis of variance (ANOVA) and Fisher's paired  
180 comparison were used to assess the differences between means, with a significance level of P  
181 = 0.05.

182 **3. Results and discussion**

183 The effect of fermentation temperature on the properties of buffalo yoghurt was assessed  
184 using pasteurised, unhomogenised, unstandardised milk and commercial starter cultures to  
185 reflect industrially relevant conditions used for yoghurt production in Australia.

### 186 *3.1 The effect of fermentation temperature on gel development*

187 During fermentation, the bacteria in the starter culture consume lactose and produce acid,  
188 leading to an increase in H<sup>+</sup> concentration and titratable acidity (Fig. 1a and 1b) and a  
189 decrease in lactose concentration (Fig. 1c). As the pH of the milk decreases towards the  
190 isoelectric point, the colloidal calcium phosphate present within the casein micelles  
191 dissociates and causes an increase in the concentration of ionic calcium (Fig. 1d). These  
192 biochemical changes were negligible during the first 150 min of the fermentation, with  
193 significant changes occurring later in the fermentation after the lag phase of bacterial growth.

194 The decreasing negative charge of the casein micelles also results in a decrease in the  
195 electrostatic repulsion and an increase in the hydrophobic interactions between casein  
196 molecules. These factors facilitate the formation of a casein network which in turn leads to an  
197 increase in the storage modulus G' (Fig. 1e). The increase in the storage modulus G' is  
198 minimal until it reaches the gelation point, defined as the time when the storage modulus G'  
199 exceeded 1 Pa (Lee and Lucey 2003) .

200 The fermentation time, defined as the time for milk samples to reach an H<sup>+</sup> concentration of ~  
201  $3.2 \times 10^{-5}$ M (equivalent to pH 4.5), increased from 360 to 420 and then 510 min, for samples  
202 incubated at 43°C, 40°C and 37°C respectively (Fig. 1a). The gelation time was also shortest  
203 for yoghurt formed at 43°C compared to 40°C or 37°C (inset of Fig. 1e). The shortest gelation  
204 and fermentation time observed for samples at 43°C are consistent with the fastest  
205 acidification rate under these conditions, as indicated by the steepest gradients in the plots of  
206 H<sup>+</sup> concentration versus fermentation time (Fig. 1a). This observation is similar to that  
207 reported in previous studies for bovine yoghurt (Purwandari et al. 2007; Lee and Lucey 2003;  
208 Laligant et al. 2003). For example, Lee and Lucey (2003) found a decrease in fermentation  
209 and gelation time from 790 and 389 min to 490 and 180 min when the fermentation  
210 temperature decreased from 46°C to 34°C.

211 While the rate of change in these parameters varies with the fermentation temperature, at the  
212 end of the fermentation time, no significant difference is observed in titratable acidity or the

213 concentration of lactose between treatments ( $P>0.05$ ). The ionic calcium concentration in  
214 samples fermented at 40°C or 37°C was slightly lower than at 43°C ( $P<0.05$ ) possibly due to  
215 the slower rate of fermentation. These results suggest there was no significant difference in  
216 the overall metabolism of the lactic bacteria as a function of temperature under the conditions  
217 used in this study.

218 At the end of the fermentation (pH~4.5, also indicated by the black arrows in Fig. 1e), the  $G'$   
219 value was lowest at the highest fermentation temperature (43°C). This result is consistent with  
220 the previous work of Lee and Lucey (2003) who reported a lower storage modulus  $G'$  when  
221 bovine yoghurt was fermented at a higher temperature. An increase in fermentation  
222 temperature results in a faster acidification rate but allows less time for the interaction  
223 between protein particles leading to the formation of a less branched network that decreases  
224 the storage modulus  $G'$  (Fig. 1e). Furthermore, according to Lee and Lucey (2004), the  
225 greater mobility of the protein molecules at the high fermentation temperature may also  
226 contribute to an increased protein network rearrangement. This possibly results in a less stable  
227 and weaker gel network, indicated by the lower storage modulus  $G'$ .

228 The concentration of lactose at the end of fermentation of buffalo yoghurt in our study was  
229 lower than in previous studies for buffalo yoghurt,  $3.9 \pm 0.2$  (% w/v) vs. 4.7 - 5.0 (% w/v)  
230 (Shiby and Mishra 2008) while the titratable acidity was higher,  $1.1 \pm 0.1$  (% lactic acid  
231 equivalents) vs. 0.7 - 0.9 (% lactic acid equivalents) (Shiby and Mishra 2008; Yadav et al.  
232 2007; Nahar et al. 2007). This is likely due to the lower pH used to define the end of  
233 fermentation in our study compared to others (pH 4.50 - 4.54 vs. pH 4.80 - 4.90) (Bezerra et  
234 al. 2012; Yadav et al. 2007), consistent with the requirement for yoghurt production in  
235 Australia (Australia and New Zealand Food Standards 2006).

### 236 *3.2 The effect of fermentation temperature on the firmness of buffalo yoghurt*

237 The yoghurt gel firmness was affected by fermentation temperature ( $P<0.05$ ) and was higher  
238 for yoghurt fermented at 43°C compared to 37°C as shown in Fig. 2a. Our result is in  
239 contradiction to a number of other studies using bovine milk who reported that a lower  
240 fermentation temperature leads to a stronger gel network (Lee and Lucey 2003, 2004; Anema  
241 2008). In these cases, it was argued that the lower temperature allows more interaction and  
242 cross-links within proteins in the gel leading to the formation of a network that is more  
243 branched and homogenous in structure.

244 The gel firmness of the buffalo yoghurt in this study increased significantly with storage time  
245 by 40 to 50 % for all treatments. This behaviour is consistent with that observed for bovine  
246 yoghurt (Saccaro et al. 2009; Salvador and Fiszman 2004), but not in other reports for buffalo  
247 yoghurt where the gel firmness decreased (Yadav et al. 2007). This inconsistency in previous  
248 studies is possibly due to the different starter cultures used. Starter cultures such as  
249 *Bifidobacterium lactis* have been observed to contribute to the increase in gel firmness in  
250 bovine yoghurt (Saccaro et al. 2009), possibly due to its capacity to produce  
251 exopolysaccharide (EPS) including L-rhamnopyranose, D-glucopyranose, D-galactopyranose  
252 and D-galactofuranose (Hidalgo-Cantabrana et al. 2013; Leivers et al. 2011) which may  
253 interact with the protein network, resulting in yoghurt with an improved texture (Zhang and  
254 Zhang 2012). This bacteria strain was also present in the starter culture used in our study and  
255 hence, the gel firmness was also expected to increase during cold storage.

### 256 3.3 *The effect of fermentation temperature on the syneresis of buffalo yoghurt*

257 Syneresis is a major physical defect in yoghurt and is determined by the amount of whey that  
258 separates from the yoghurt over time. The centrifugation method was used in this study to  
259 facilitate the collection and assessment of the whey expelled.

260 The fermentation temperature had a significant effect on the syneresis of buffalo yoghurt  
261 ( $P < 0.05$ ) (Fig. 2b). A significantly greater mass of whey was expelled from buffalo yoghurt  
262 samples fermented at 43°C compared to those fermented at 40°C or 37°C ( $P < 0.05$ ), while no  
263 difference was observed between buffalo yoghurt fermented at these two lower temperatures  
264 ( $P > 0.05$ ). Syneresis was also affected by storage time ( $P < 0.05$ ); prolonged storage time  
265 increased syneresis for samples fermented at 43°C. The greater expulsion of whey from  
266 buffalo yoghurt samples fermented at this temperature is possibly linked to the more rapid  
267 acidification occurring. A higher acidification rate may result in a less developed protein  
268 network with fewer protein cross-links leading to a weaker gel that is more susceptible to  
269 syneresis (Lee and Lucey 2003; Purwandari et al. 2007; Wu et al. 2009). Furthermore, the  
270 increased hydrophobic interactions at an increased fermentation temperature could also lead  
271 to the contraction of the protein strands resulting in a weaker network containing thinner  
272 protein strands as reported in the study of Lee and Lucey (2004).

273 While there was a significant decrease in syneresis when the fermentation temperature was  
274 lowered from 43°C to 40°C, there was no significant change when this was further decreased

275 to 37°C. This result is of practical importance as it indicates that fermentation at 40°C could  
276 allow for a relatively short production time while maintaining a level of syneresis similar to  
277 that at 37°C. However, the lower syneresis level observed here for buffalo yoghurt samples  
278 fermented at reduced fermentation temperature was still approximately 8 times higher than  
279 that observed for bovine yoghurt in our previous work (Nguyen et al. 2013). This result shows  
280 that more parameters other than the fermentation temperature need to be investigated to  
281 improve the quality of buffalo yoghurt.

### 282 3.4 *The effect of fermentation temperature on the rheological properties of buffalo yoghurt*

283 The storage time was found to have a significant effect ( $P < 0.05$ ) on the rheological properties  
284 of buffalo yoghurt, whereas the effect of fermentation temperature was minimal ( $P > 0.05$ )  
285 (Fig. 3). Three measures were used to assess the buffalo yoghurt rheological properties. These  
286 were (i) the thixotropy, which is defined as the difference in the energy required to recover to  
287 original structure after deformation, (ii) the consistency coefficient  $K$ , which indicates the  
288 viscosity of the fluids, and (iii) the flow behaviour index  $n$ , which measures the deviation  
289 degree from a Newtonian fluid.

290 During the 28 days of storage, the thixotropy and consistency coefficient  $K$  increased for all  
291 samples, while the flow behaviour index  $n$  decreased. These measures indicate that the  
292 buffalo yoghurt was more susceptible to structural breakdown under external force and less  
293 capacity to recover to its original structure after storage. No significant differences were  
294 observed in the thixotropy and consistency coefficient  $K$  within buffalo yoghurt fermented at  
295 different temperatures ( $P > 0.05$ ) (Fig. 3a and 3b). A subtle difference was observed in the flow  
296 behaviour index  $n$  of buffalo yoghurt fermented at different temperatures, but only on the first  
297 day of storage ( $P < 0.05$ ) (Fig. 3c).

298 Our results differ to the findings of previous studies of bovine yoghurt where rheological  
299 properties were significantly affected by the fermentation temperature. Haque and co-authors  
300 (2001) observed a considerable increase in the viscosity of bovine yoghurt fermented at  
301 higher temperatures (46°C vs. 37°C) as measured by funnel flow. Purwandari and co-authors  
302 (2007) found the highest thixotropy of bovine yoghurt at the intermediate temperature (37°C  
303 vs. 30°C or 42°C) after 30 days of storage. The differences in the response of the rheological  
304 properties of buffalo and bovine yoghurt to the changes in fermentation temperature is likely  
305 due to the differences in the properties of the two milk types. The minimal effect of

306 fermentation temperature on the rheological properties of buffalo yoghurt indicates that  
307 procedures that have long been established to produce bovine yoghurt might not be  
308 appropriate for buffalo yoghurt, and such protocols therefore require optimisation prior to  
309 application for buffalo yoghurt production.

310 It is also noted that the rheological properties observed here for buffalo yoghurt were  
311 considerably different to bovine yoghurt made with homogenised milk reported in our  
312 previous study (Nguyen et al. 2013). For example, the thixotropy values measured at the first  
313 day of storage for buffalo yoghurt fermented at 43°C, 40°C or 37°C were  $1686 \pm 517$  (Pa.s<sup>n</sup>),  
314  $1508 \pm 381$  (Pa.s<sup>n</sup>) and  $1767 \pm 207$  (Pa.s<sup>n</sup>) respectively, significantly higher than that of  $479 \pm$   
315  $35$  (Pa.s<sup>n</sup>) observed for the bovine yoghurt. The flow behaviour index  $n$  at the first day of  
316 storage of buffalo yoghurt fermented at 43°C, 40°C or 37°C were  $0.07 \pm 0.06$ ,  $0.16 \pm 0.07$ ,  
317  $0.16 \pm 0.06$  respectively, significantly lower compared to  $0.42 \pm 0.03$  for the bovine yoghurt.  
318 This comparison gave an indication that the use of homogenised milk for buffalo yoghurt  
319 production may improve the viscosity and reduce the thixotropy value, hence giving yoghurt  
320 structures that recover better upon deformation.

### 321 3.5 *The effect of fermentation temperature on the microstructure of buffalo yoghurt*

322 Both fermentation temperature and storage were found to significantly affect the  
323 microstructure of buffalo yoghurt as assessed by CLSM and cryo-SEM (Fig. 4). While large  
324 serum pores (black areas) could be observed in all yoghurt samples regardless of the  
325 fermentation temperature, these were more numerous within the gel network of yoghurt  
326 fermented at 43°C. This difference was particularly apparent after the yoghurt samples were  
327 stored for 28 days (Fig. 4i vs. Fig. 4a, e; Fig. 4k vs. Fig. 4c, g). The protein network of buffalo  
328 yoghurt fermented at 43°C appeared less dense compared to other treatments (Fig. 4j vs. Fig.  
329 4b, f; Fig. 4l vs. Fig. 4d, h). This observation correlates well with the observation that buffalo  
330 yoghurt fermented at a higher temperature or left in storage is more susceptible to whey  
331 separation (Fig. 2b), more sensitive to the external force indicated by the flow behaviour  
332 index  $n$  (Fig. 3c) and less able to recover to the original structure after deformation, as  
333 indicated by the thixotropy value (Fig. 3a). The protein strands at day 28 of the storage are  
334 denser than day 1 (Fig. 4d vs. 4b; Fig. 4h vs. 4f; Fig. 4l vs. 4j), probably due to the fusion of  
335 the protein aggregates or the further development of the network during the cold storage. This  
336 observation is consistent with the increase in gel firmness of buffalo yoghurt with storage  
337 (Fig. 2a)

338 These results are in agreement with the findings of Lee and Lucey (2003) who observed the  
339 larger pores within a less dense protein network in bovine yoghurt fermented at a higher  
340 temperature (46°C vs. 40°C). These authors suggest that hydrophobic interaction within the  
341 casein particles increase with temperature, leading to a decrease in the voluminosity and the  
342 contact area between casein particles within the protein network, resulting in a yoghurt  
343 microstructure with larger serum pores and thinner protein strands.

### 344 **3.6 The effect of fermentation temperature on the bacterial growth during** 345 **fermentation and viability of bacteria during storage**

346 Similarities were observed in the growth and viability of starter culture bacteria during  
347 fermentation and storage regardless of the fermentation temperature. During fermentation, the  
348 number of all three bacterial starter strains increased significantly ( $P < 0.05$ ) from a similar  
349 level of bacterial inoculation with the fastest growth rate observed for *Streptococcus*  
350 *thermophilus*, followed by *Lactobacillus acidophilus* La-5 and the lowest rate in  
351 *Bifidobacterium lactis* Bb-12 (Fig. 5). During storage, the number of viable probiotic  
352 *Lactobacillus acidophilus* La-5 and *Bifidobacterium lactis* Bb-12 bacteria reduced  
353 significantly ( $P < 0.05$ ) with the more profound decrease observed in La-5, while the viability  
354 of *Streptococcus thermophilus* remained unchanged ( $P > 0.05$ ).

355 The superior growth and viability of *Streptococcus thermophilus* and the slower growth and  
356 poorer viability of *Lactobacillus acidophilus* La-5 have been reported in previous studies of  
357 bovine yoghurt (Damin et al. 2008; Oliveira et al. 2001; Dave and Shah 1997). Briefly, this  
358 phenomenon is thought to arise due to the higher proteolytic activity of *Streptococcus*  
359 *thermophilic* compared to *Lactobacillus acidophilus* and the susceptibility of the latter strain  
360 to acidic and cold conditions during storage (Ozer and Kirmaci 2010; Marafon et al. 2011;  
361 Gilliland and Lara 1988).

362 The fermentation temperature did not significantly affect the growth of bacteria but altered the  
363 bacterial viability during storage ( $P > 0.05$ ). During storage, *Bifidobacterium lactis* Bb-12  
364 survived in greater number in the buffalo yoghurt fermented at 43°C and 40°C (Fig. 5b) while  
365 *Lactobacillus acidophilus* La-5 survived better in buffalo yoghurt fermented at 37°C (Fig. 5c).

366 Several studies reported on the optimum growth of *Lactobacillus acidophilus* at 37°C (Baati  
367 et al. 2004; Bozanic et al. 2008; Shafiee et al. 2010) while there are no studies to date on the

368 effect of fermentation temperature on the survival of this probiotic bacteria in yoghurt during  
369 cold storage. It is, however, interesting to note that a relationship exists between the  
370 fermentation temperature and the survival rate of *Lactobacillus acidophilus* La-5 during  
371 frozen stage or cryo-storage at -20°C that is often used for the preservation of the starter  
372 culture (Murga et al. 2000; Wang et al. 2005b; Wang et al. 2005a). Wang and co-authors  
373 (2005a) found that *Lactobacillus acidophilus* was more cryo-resistant when it was cultured at  
374 low temperatures between 30°C and 37°C compared to 42°C. *Lactobacillus acidophilus*  
375 survival after freeze-thawing also increased systematically from 14% to 67% when it was  
376 fermented at 25°C compared to 40°C. This improved resistance is thought to be due to the  
377 increase in the ratio of unsaturated fatty acids, the high concentration of 19-carbon  
378 cyclopropane membrane fatty acid (cyc C19:0) and the upregulation of specific proteins that  
379 help the bacterial cells to adapt to freezing (Murga et al. 2000; Wang et al. 2005b; Wang et al.  
380 2005a). These adaptations may also assist *Lactobacillus acidophilus* during cold storage,  
381 leading to the higher numbers of survival of this bacteria strain in the yoghurt samples  
382 fermented at 37°C observed here.

#### 383 **4. Conclusion**

384 The fermentation temperature used to produce buffalo yoghurt significantly affected the  
385 yoghurt quality, altering the physical appearance and yoghurt microstructure. The faster  
386 fermentation at a higher fermentation temperature of 43°C increased the ionic calcium  
387 concentration and the gel firmness of buffalo yoghurt but altered the protein network leading  
388 to larger and more numerous serum pores within the microstructure resulting in higher  
389 syneresis. The rheological properties, lactose and titratable acidity were not affected by  
390 fermentation temperature but the viability of the probiotic *Lactobacillus acidophilus* La-5  
391 bacteria was significantly reduced with increased fermentation temperature. In contrast,  
392 fermentation at the lower temperatures of 37°C or 40°C was longer, leading to a more  
393 consistent product and these temperatures are recommended to improve the microstructure  
394 and syneresis of buffalo yoghurt. The gel firmness, thixotropy and consistency coefficient  
395 increased while the flow behaviour index decreased on storage for all treatments. The high  
396 level of syneresis observed here relative to bovine yoghurt, even for those samples fermented  
397 at lower fermentation temperatures, suggests that further work is required to optimise the  
398 processing condition for buffalo yoghurt. In particular, while this yoghurt is typically made  
399 from unhomogenised milk, it may be useful to examine whether homogenisation might be

400 used to improve the yoghurt structure. This study also confirms that while there are many  
401 similarities between buffalo and bovine yoghurt, studies specific to buffalo products are  
402 required to optimise production.

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- 578

579 **List of Figures and Figure legends**

580

581 **Figure 1.** Changes in the concentration of dissociated H<sup>+</sup> ion (a), titratable acidity (b), lactose  
582 (c), ionic calcium (d) and storage modulus G' (e) during the fermentation of buffalo yoghurt  
583 fermented at 37°C (●), 40°C (●) or 43°C (○). Each data point is the average of six replicates  
584 (n=6) in Figure 1a-d and two replicates (n=2) in Figure 1e. The error bars are the standard  
585 deviation of the mean. The inset in (e) corresponds to the data between 140 - 260 min of  
586 fermentation time.

587 **Figure 2.** Changes in the gel firmness (a) and syneresis (b) during cold storage of buffalo  
588 yoghurt fermented at 37°C (●), 40°C (●) or 43°C (○). Each data point is the average of six  
589 replicates (n=6) and the error bars are the standard deviation of the mean.

590 **Figure 3.** Changes in the thixotropy (a), consistency coefficient (b) and flow behaviour index  
591 (c) during cold storage of buffalo yoghurt fermented at 37°C (●), 40°C (●) or 43°C (○). Each  
592 data point is the average of six replicates (n=6) and the error bars are the standard deviation of  
593 the mean.

594 **Figure 4.** Microstructure of buffalo yoghurt fermented at 37°C (a-d), 40°C (e-h) o 43°C (i-l) at  
595 day 1 (two left column images) and day 28 (two right column images) of storage as observed  
596 by CLSM and cryo-SEM. Nile Red stained fat appears red, FCF stained protein appears  
597 green, the black areas are serum pores and CLSM images were captured using a 63x objective  
598 using a 1x digital zoom (first and third column images). Cryo-SEM images were captured  
599 using a solid state detector at 16000x magnification (second and fourth column images). The  
600 scale bars are 10 μm (first and third column CLSM images) or 5 μm in length (second and  
601 fourth column cryo-SEM images).

602 **Figure 5.** Bacterial growth and viability during the fermentation and storage of buffalo  
603 yoghurt fermented at 37°C (●), 40°C (●) or 43°C (○). Each data point is the average of six  
604 replicates (n=6) and the error bars are the standard deviation of the mean. Storage commenced  
605 after 6 hours, 7 hours and 8.5 hours for buffalo yoghurt fermented at 37°C, 40°C and 43°C  
606 respectively.

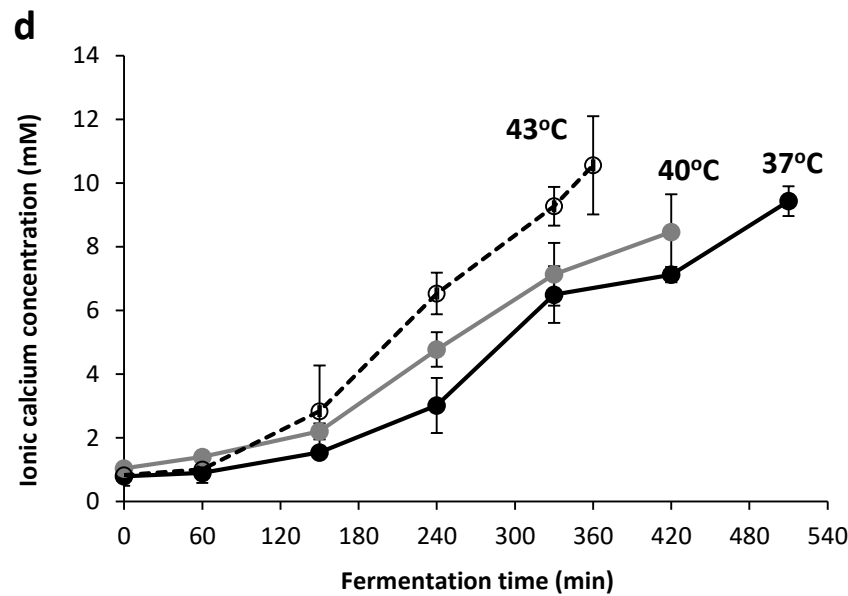
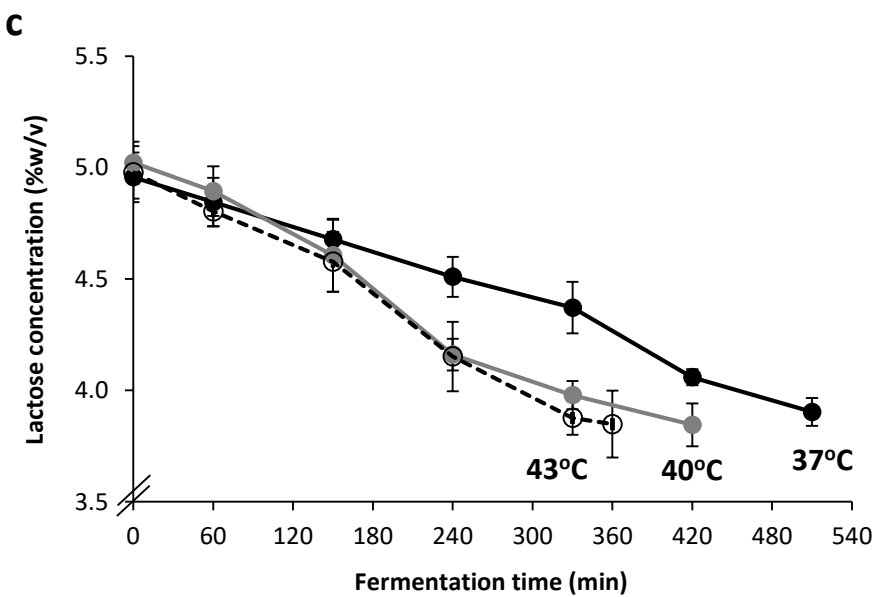
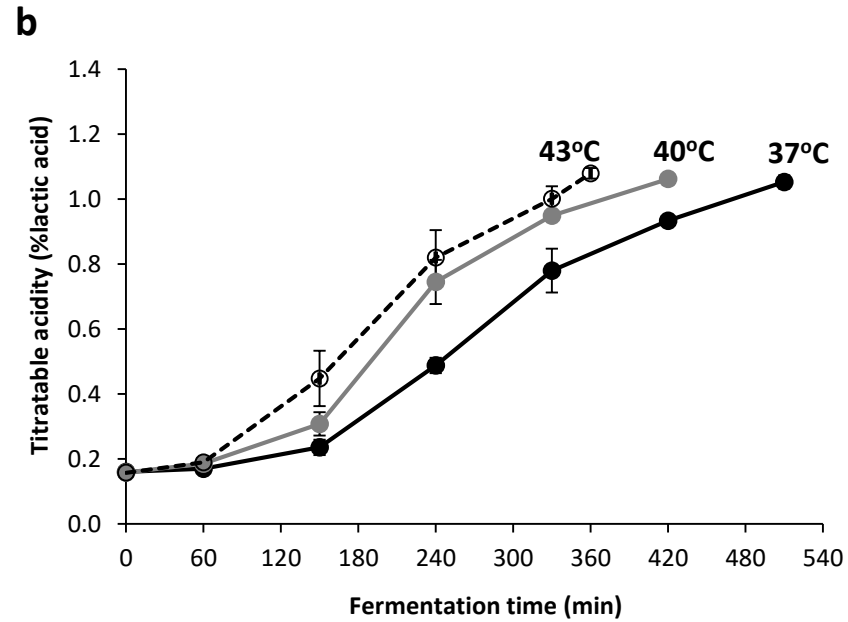
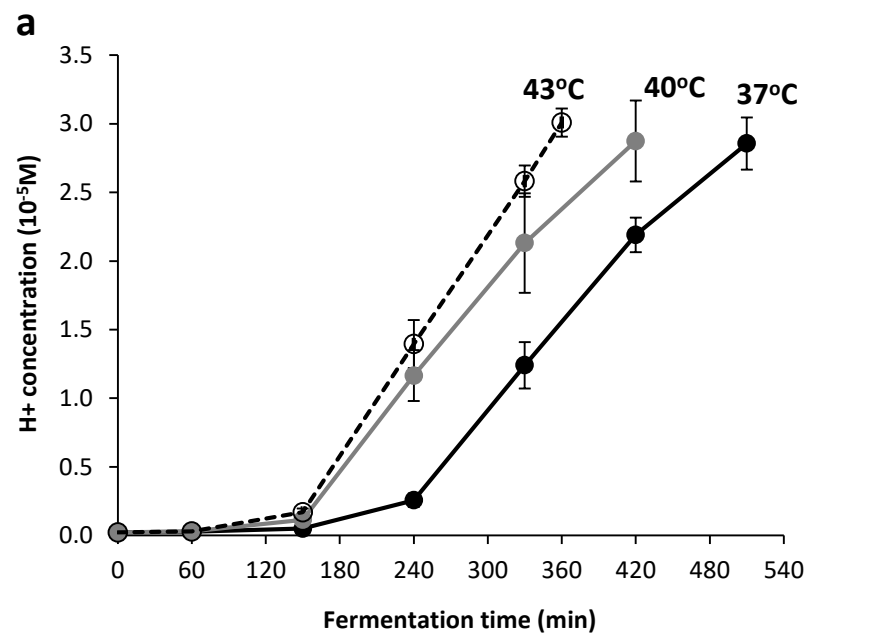


Figure 1

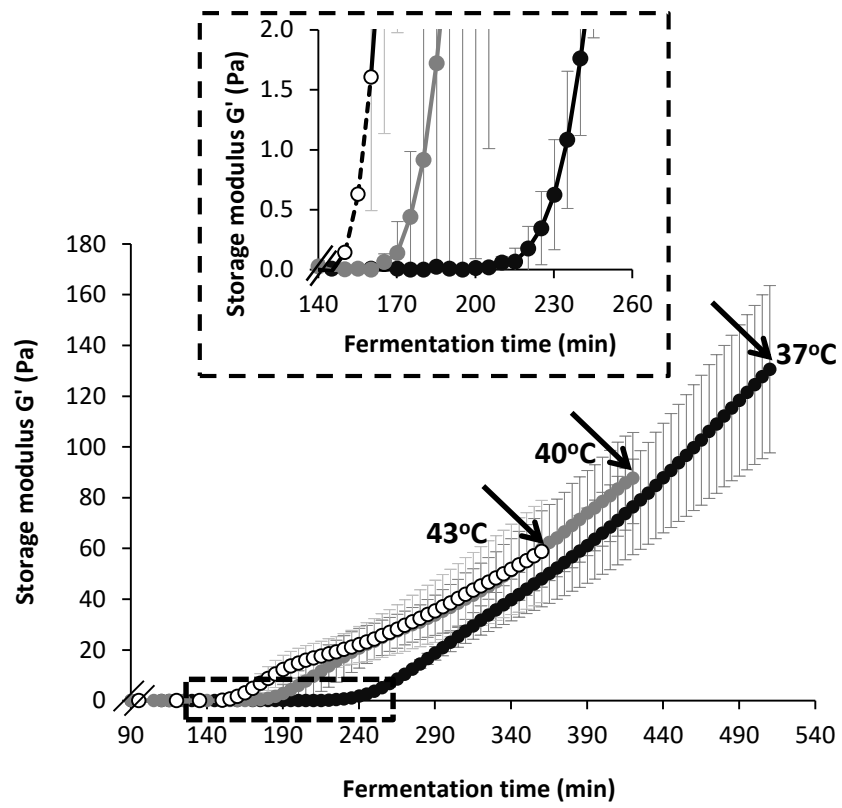


Figure 2

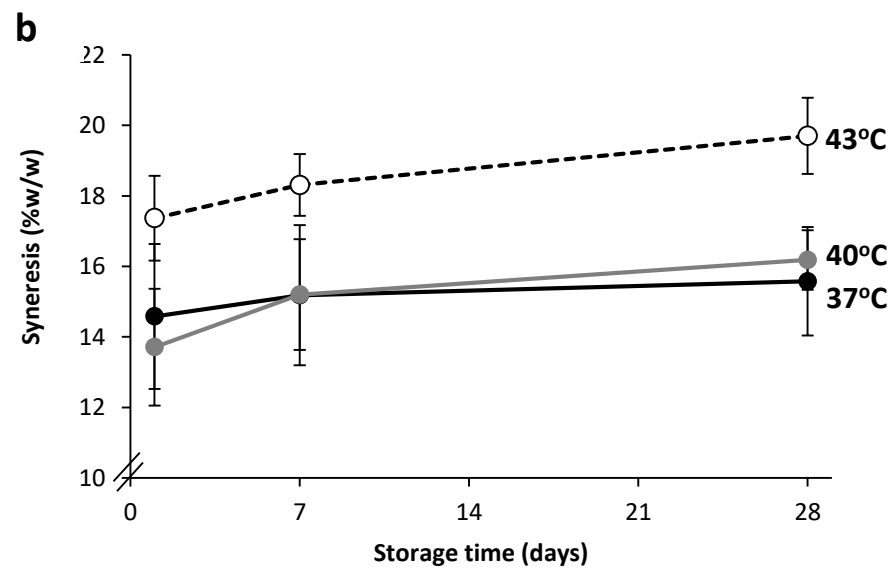
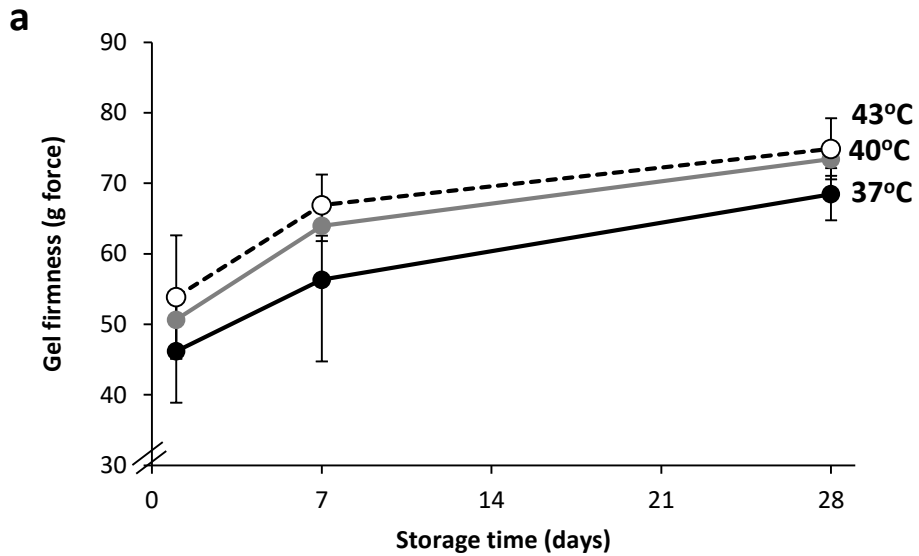


Figure 3

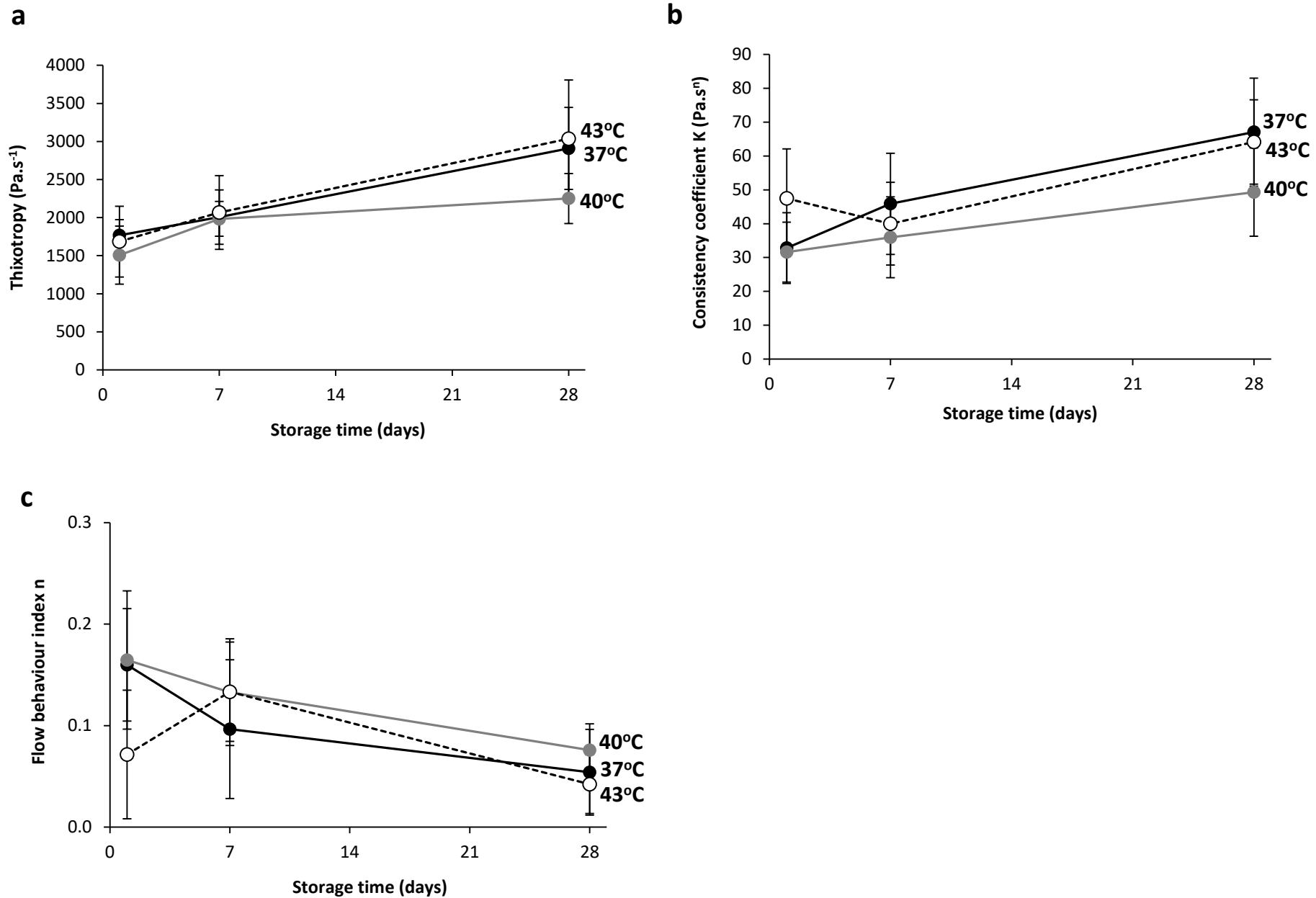
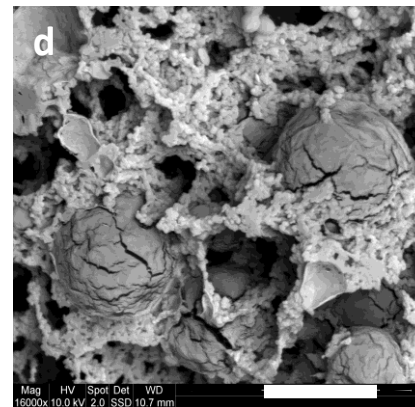
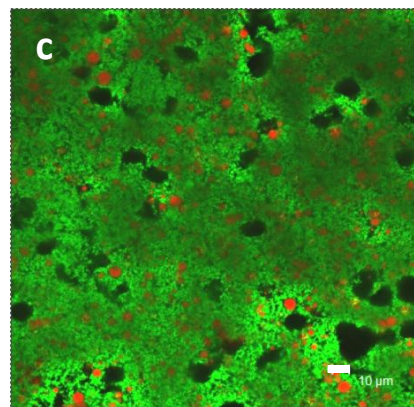
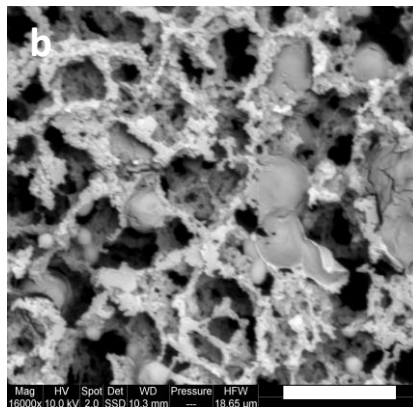
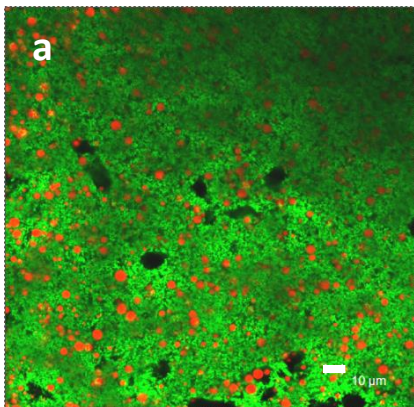


Figure 4

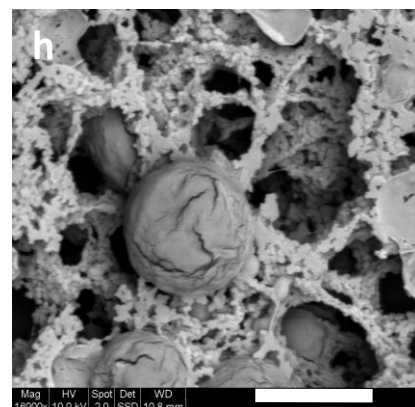
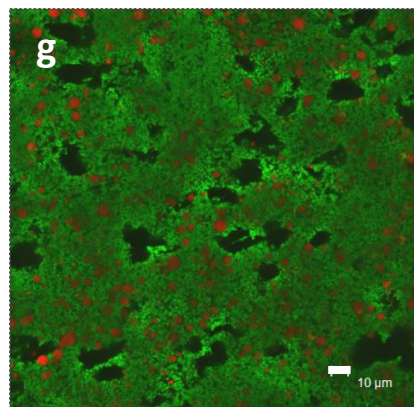
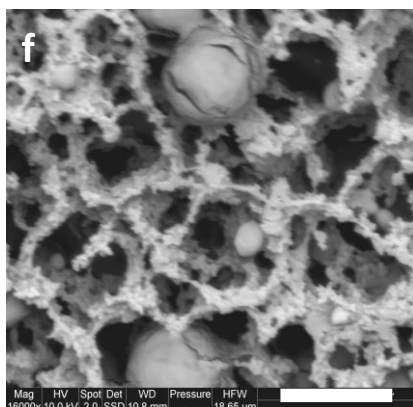
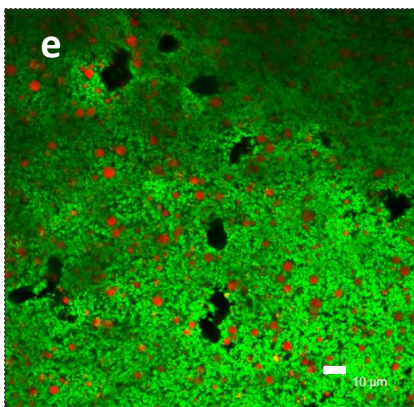
Day 1

Day 28

37°C



40°C



43°C

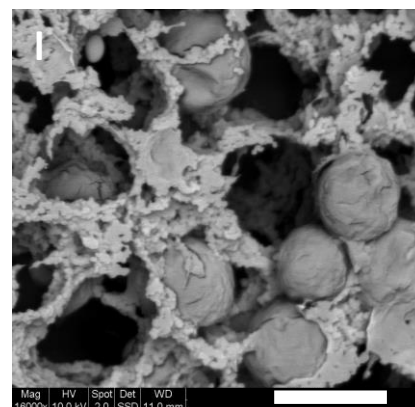
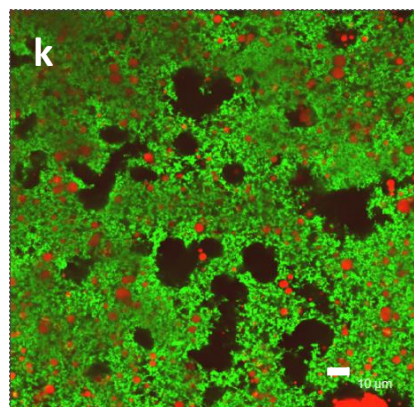
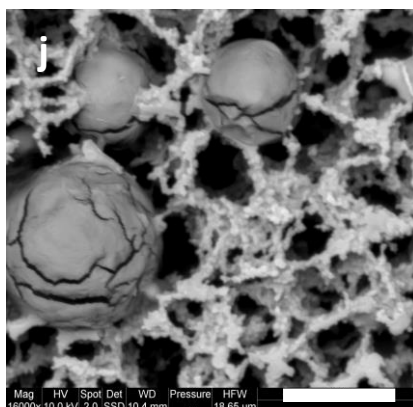
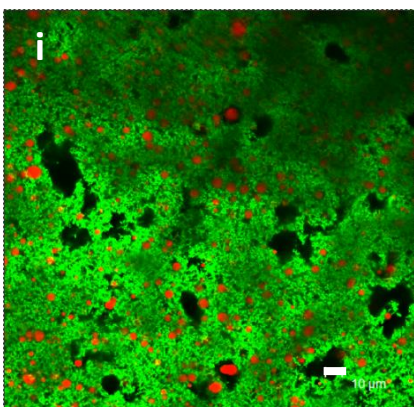


Figure 5

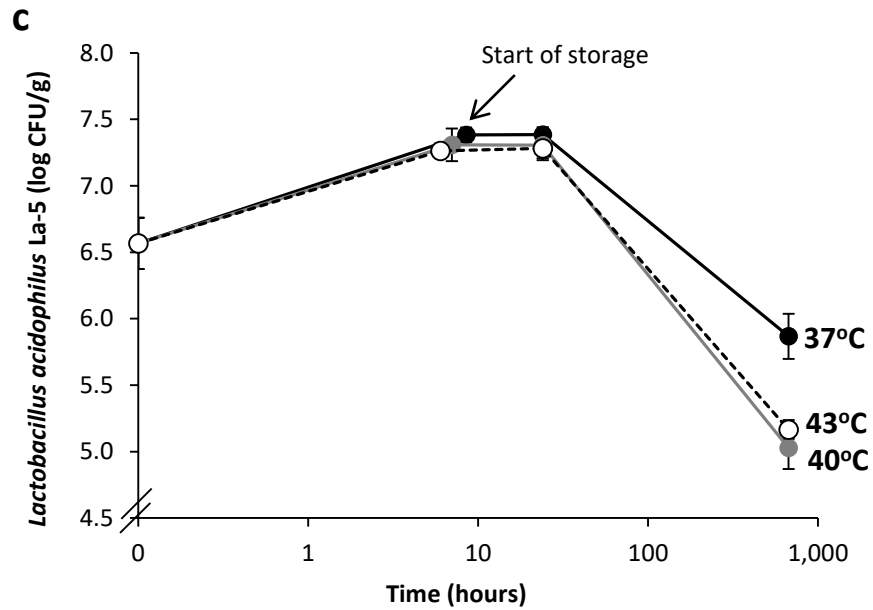
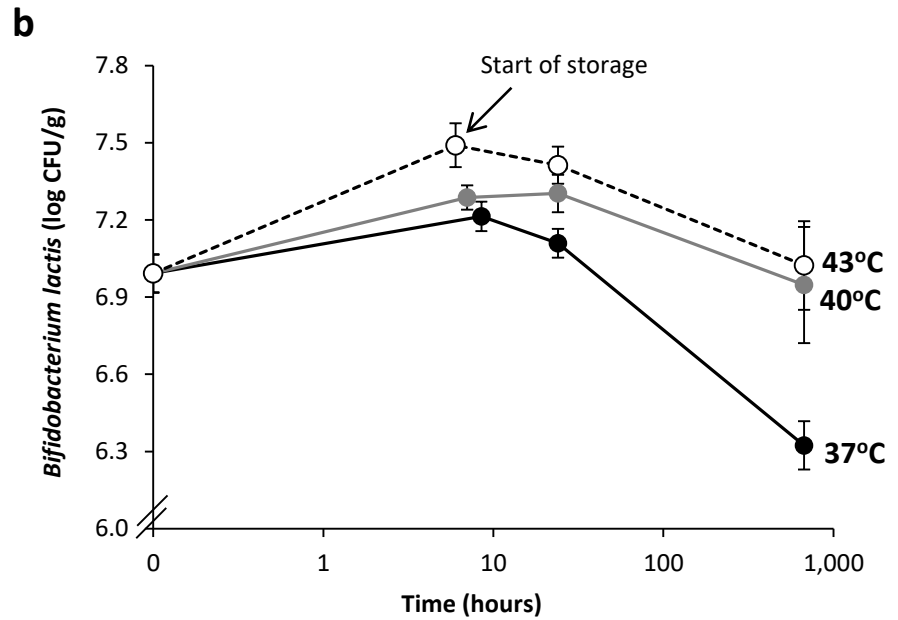
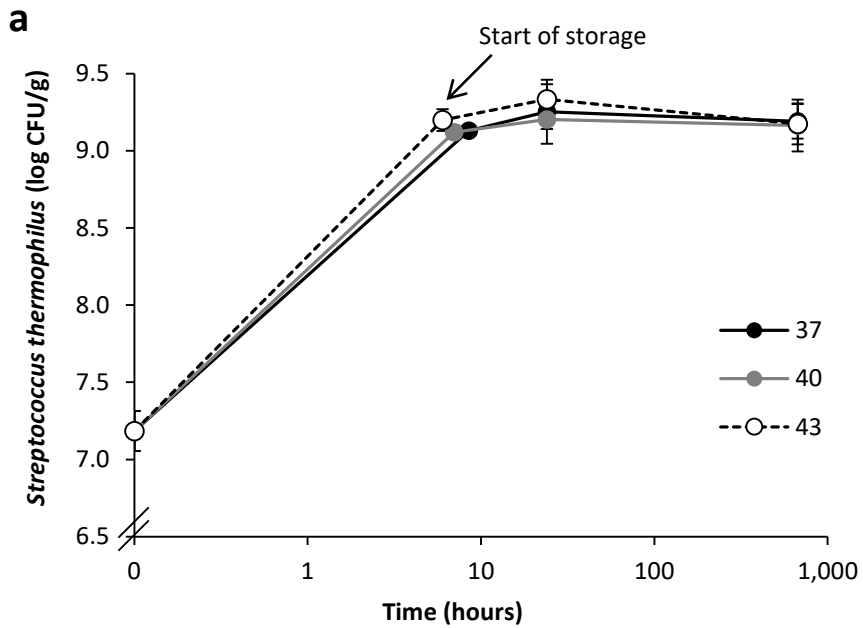


Figure 6