

Interactions of bile salts with a dietary fibre, methylcellulose, and impact on lipolysis

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1 Interactions of bile salts with a dietary fibre, methylcellulose, and impact

2 on lipolysis

- 3 <u>Olivia Pabois^{a, b}, Amandine Antoine-Michard^a, Xi Zhao^b, Jasmin Omar^b, Faizah Ahmed^b, Florian</u>
- 4 Alexis^a, Richard D. Harvey^c, Isabelle Grillo^a, Yuri Gerelli^a, Myriam M.-L. Grundy^d, Balazs Bajka^e,
- 5 Peter J. Wilde^f, Cécile A. Dreiss^{b*}
- 6 ^a Institut Laue-Langevin, Grenoble 38000, France
- ^b Institute of Pharmaceutical Science, King's College London, London SE1 9NH, United Kingdom
- 9 ^c Institut für Pharmazie, Martin-Luther-Universität Halle-Wittenberg, Halle (Saale) 06099,

10 Germany

- ^d School of Agriculture, Policy and Development, University of Reading, Reading RG6 6AR,
 United-Kingdom
- 13 ^e Department of Nutritional Sciences, King's College London, London SE1 9NH, United-
- 14 Kingdom
- ¹⁵ ^f Quadram Institute Bioscience, Norwich Research Park, Norwich NR4 7UA, United-Kingdom
- 16 E-mail addresses:

17	<u>olivia.pabois@kcl.ac.uk</u> ;	amandine-03@h	<u>otmail.fr</u> ;	<u>xi.zhao@kcl.ac.uk</u> ;
18	jasmin.1.omar@kcl.ac.uk;	faizah.ahmed@kc	:l.ac.uk;	floalexis0@gmail.com;
19	richard.harvey@pharmazie.uni-h	nalle.de;	<u>grillo@ill.fr</u> ;	gerelli@ill.fr;
20	m.m.grundy@reading.ac.uk;	balazs.bajka@kcl.a	<u>c.uk; pete</u>	r.wilde@quadram.ac.uk;

- 21 cecile.dreiss@kcl.ac.uk
- 22 Corresponding author:
- 23 Cécile A. Dreiss:
- 24 King's College London
- 25 Institute of Pharmaceutical Science
- 26 Franklin-Wilkins Building

- 27 150 Stamford Street
- 28 SE1 9NH London, UK
- 29 Tel: +44 (0)207 848 3766

30 Highlights

- BS, NaTC and NaTDC, impact the rheological properties and gelation of MC.
- NaTDC has a greater impact on the viscoelasticity of MC compared to NaTC.
- NaTDC desorbs from a MC-stabilised interface at lower concentrations than NaTC.
- Upon digestion, NaTDC destabilises more readily MC-stabilised emulsion droplets.
- During MC-stabilised emulsion digestion, NaTDC generates less FFA than NaTC.

36 Abstract

37 Methylcellulose (MC) has a demonstrated capacity to reduce fat absorption, hypothetically 38 through bile salt (BS) activity inhibition. We investigated MC cholesterol-lowering mechanism, 39 and compared the influence of two BS, sodium taurocholate (NaTC) and sodium 40 taurodeoxycholate (NaTDC), which differ slightly by their architecture and exhibit contrasting 41 functions during lipolysis.

42 BS/MC bulk interactions were investigated by rheology, and BS behaviour at the MC/water 43 interface studied with surface pressure and ellipsometry measurements. *In vitro* lipolysis 44 studies were performed to evaluate the effect of BS on MC-stabilised emulsion droplets 45 microstructure, with confocal microscopy, and free fatty acids release, with the pH-stat 46 method.

Our results demonstrate that BS structure dictates their interactions with MC, which, in turn, impact lipolysis. Compared to NaTC, NaTDC alters MC viscoelasticity more significantly, which may correlate with its weaker ability to promote lipolysis, and desorbs from the interface at lower concentrations, which may explain its higher propensity to destabilise emulsions.

51 Keywords

52 Methylcellulose; bile salts; rheology; surface pressure measurements; *in vitro* duodenal 53 lipolysis

55 Graphical abstract



Oil droplets stabilised by methylcellulose



57 **1. Introduction**

58 Obesity and associated health risks (such as chronic cardiovascular diseases and type-59 2-diabetes mellitus) have become increasingly prevalent worldwide. In 2016, 39% of the 60 world's adult population were classified as overweight, and 13% as obese (World Health 61 Organization, 2019). Controlling the digestion of dietary lipids (fats) and optimising their 62 absorption are therefore crucial to addressing this ongoing health crisis (McClements & Li, 63 2010b; Mei, Lindqvist, Krabisch, Rehfeld, & Erlanson-Albertsson, 2006). With their 64 demonstrated capability to reduce food intake and aid weight loss, dietary fibres have shown 65 great potential against obesity (Slavin, 2005). Nonetheless, a better understanding of the 66 processes responsible for their ability to regulate calorie uptake still needs to be provided. 67 Due to its approved (Younes et al., 2018) and wide (The Dow Chemical Company, 2002) use in 68 the food industry, as well as its proven capacity to diminish blood cholesterol levels (without 69 inducing any adverse effect) (Agostoni et al., 2010), methylcellulose (MC) is an appropriate 70 model of dietary fibre for elucidating the mechanism by which dietary fibres reduce 71 hyperlipidaemia.

72 MC is a non-ionic polysaccharide belonging to the large family of cellulose ethers and 73 containing repeating anhydroglucose units, with methyl (hydrophobic) moieties substituting 74 hydroxyl (hydrophilic) groups (Nasatto et al., 2015b) (Figure 1). The capacity of this dietary 75 fibre to hinder lipolysis has been mainly attributed to its ability to induce loss of bile salts (BS) 76 and cholesterol in faeces by (i) increasing the viscosity of the small intestine content (Christos 77 Reppas, Meyer, Sirois, & Dressman, 1991), which slows down fat digestion and reduces 78 nutrients absorption (Bartley et al., 2010; Carr, Gallaher, Yang, & Hassel, 1996; Maki et al., 79 2009; C. Reppas, Swidan, Tobey, Turowski, & Dressman, 2009; van der Gronde, Hartog, van 80 Hees, Pellikaan, & Pieters, 2016), and/or by (ii) trapping BS and/or cholesterol molecules in its 81 network, via hydrophobic interactions occurring both in the bulk aqueous phase and at the 82 fat droplet interface (Pilosof, 2017; Pizones Ruiz-Henestrosa, Bellesi, Camino, & Pilosof, 2017; 83 Torcello-Gómez et al., 2015; Torcello-Gómez & Foster, 2014). BS are biosurfactants produced 84 in the liver and released into the small intestine (duodenum) (Hofmann & Mysels, 1987), 85 which play key roles in lipid digestion and absorption (Maldonado-Valderrama, Wilde, 86 Macierzanka, & Mackie, 2011; Wilde & Chu, 2011): on the one hand, they facilitate enzyme 87 adsorption to fat droplet interfaces, thus promoting enzyme-catalysed lipolysis (Borgström,

88 Erlanson-Albertsson, & Wieloch, 1979; Bourbon Freie, Ferrato, Carrière, & Lowe, 2006; 89 Erlanson-Albertsson, 1983; Labourdenne, Brass, Ivanova, Cagna, & Verger, 1997); on the 90 other, they remove the enzyme-inhibiting insoluble lipolysis products (diacylglycerols (DAG), 91 monoacylglycerols (MAG) and free fatty acids (FFA)) present at the interface, carrying them 92 to the gut mucosa for absorption (Hofmann & Mysels, 1987). In this work, we are focusing on 93 the interactions between MC and BS, which have been hypothesised to explain (i) MC 94 cholesterol-lowering effect, due to the reduction in BS re-absorption in the ileum and the 95 subsequent increased production of BS by the liver from cholesterol, and (ii) the early 96 signalling of satiation and lengthening of satiety feeling, by the accumulation of undigested 97 materials in the duodenum, due to BS being entrapped and prevented from fulfilling their 98 functions during lipolysis (Gunness & Gidley, 2010). Recent studies have demonstrated BS 99 inhibitory effect on MC thermally-induced structuring using microcalorimetry and rheology 100 (Torcello-Gómez et al., 2015; Torcello-Gómez & Foster, 2014), and the competition of BS with 101 MC for adsorption at the lipid droplet/water interface with tensiometry (Torcello-Gómez & 102 Foster, 2014). However, there is little structural evidence for the hypothesis of entrapment of 103 BS by MC, and a mechanistic understanding of the competitive processes leading to enzyme 104 inhibition, delayed fat digestion and the associated health benefits, is still lacking. Therefore, 105 further studies are required to clarify how MC interacts with BS during lipid digestion and how 106 this, in turn, correlates to BS molecular structure and their contrasting roles.

107 The work presented here increases our understanding of the mechanisms underlying 108 MC capacity to regulate fat digestion in the small intestine, with a particular focus on its ability 109 to compete with BS for adsorption at the lipid droplet/water interface. More specifically, by 110 combining bulk and interfacial experiments with in vitro lipolysis studies, we examined the 111 interactions between MC and BS in bulk water, at the MC/water interface, and at the oil/water 112 interface of fat droplets mimicking food colloids. It has been hypothesised that BS structural 113 diversity is responsible for the different functions they carry out in fat digestion; to explore 114 this postulate, two BS, sodium taurocholate (NaTC) and sodium taurodeoxycholate (NaTDC) 115 (Figure 2), were selected, as they display contrasting adsorption/desorption dynamics, which 116 are thought to reflect their different roles in the gut (Pabois et al., 2019; Parker, Rigby, Ridout, 117 Gunning, & Wilde, 2014). Since BS are expected to interact with MC both in the bulk aqueous 118 phase and at the surface of MC-stabilised emulsion droplets, we assessed the impact of BS on 119 MC rheological properties, using oscillatory shear rheology, and BS/MC interfacial behaviour

- at the air/water interface, through surface pressure measurements in a Langmuir trough set up and ellipsometry. We then investigated how these interactions affect the lipolysis of an
 MC-stabilised emulsion, by monitoring the structure of emulsion droplets after addition of BS
 and enzymes, with different optical microscopy techniques, and by measuring the amount of
 FFA released throughout *in vitro* lipid digestion, with the pH-stat method.

126 **2. Experimental section**

127 2.1 Materials

128 Methocel[™] SG A7C (solution viscosity: 700 mPa.s at 2% w/w at 20°C; methoxyl degree of 129 substitution: 1.8; molecular weight: 400 - 500 kDa) (Figure 1) was kindly supplied by Dow Wolff 130 Cellulosics GmbH (Bomlitz, Germany). Chloroform (CHCl₃) was purchased from Fisher 131 Scientific (Loughborough, UK). NaTC (P97.0% TLC) (Figure 2A), NaTDC (P95.0% TLC) (Figure 132 2B), paraffin oil, ethanol (EtOH, P99.8% GC), orlistat (P98.0%), Nile red, fluorescent brightener 133 28 (calcofluor), dimethyl sulfoxide anhydrous (P99.9%), sunflower seed oil from Helianthus 134 annuus, pancreatin from porcine pancreas (or pancreatic lipase/co-lipase; activity: 40 U/mg 135 of solid, based on lipase activity using tributyrin as a substrate), sodium phosphate monobasic 136 dihydrate (NaH₂PO₄, P99.0% T), sodium phosphate dibasic dihydrate (Na₂HPO₄, P99.0% T), 137 sodium chloride (NaCl, P99.8%), calcium chloride dihydrate (CaCl₂, P99.0%) and sodium 138 hydroxide (NaOH, 0.1 M) were all obtained from Sigma-Aldrich (Gillingham, UK). Ultrapure 139 water, or MilliQ-grade water (H₂O, 18.2 MΩ·cm, Merck Millipore, Molsheim, France), was 140 used in all experiments. Phosphate buffer (10 mM, pH = 7.04 at 21°C) was prepared by mixing 141 0.01% wt NaH₂PO₄ with 0.01% wt Na₂HPO₄, in ultrapure water. All reagents were used as 142 supplied.



143



Figure 1: Structure of methylcellulose (MC)



145



Figure 2: Structures of sodium taurocholate (NaTC) (A) and sodium taurodeoxycholate (NaTDC) (B)

147 **2.2 Methods**

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2.2.1 Bulk and interfacial studies

149

2.2.1.1 Preparation of MC and MC/BS aqueous solutions

MC aqueous solution was prepared using the "hot/cold" method (The Dow Chemical Company, 2002, 2013). Solid MC was first dispersed into one third of the required mass of ultrapure water heated to 80°C (for around 15 minutes), until complete wetting of particles; then, the dispersion was transferred into an ice bath, and the remaining two thirds of cold ultrapure water (4°C) were added progressively into the stirred solution, which was finally left to stir overnight at 4°C, to ensure complete solubilisation. MC/BS solutions were prepared simply by mixing both components together at the required concentrations.

157

2.2.1.2 Rheology measurements

158 Rheology experiments were performed with a strain-controlled rheometer (ARES, TA 159 instruments, Inc, Borehamwood, UK), fitted with a 25 mm diameter titanium parallel plate 160 and equipped with a temperature-controlled Peltier system (with a \pm 0.1°C temperature 161 stability at thermal equilibrium). Each sample was loaded onto the lower plate, and the upper 162 plate was adjusted to a gap size of 0.8 \pm 0.3 mm. A thin layer of low viscosity paraffin oil was 163 deposited around the edges of the sample exposed to air to prevent sample drying and 164 evaporation throughout the measurement.

165 Dynamic temperature sweeps were performed at a fixed angular frequency of 6.28 rad/s and 166 strain of 1%, from 20°C to 80°C, with a heating rate of 2°C/min, to measure the evolution of 167 the storage (G') and loss (G'') moduli as a function of temperature, in the absence and 168 presence of BS. Dynamic frequency sweeps were performed over an angular frequency range 169 of 0.1 - 100 rad/s, at a fixed strain of 1%, and a fixed temperature of 60°C (above MC transition 170 temperature (T_t), which is the point where a break in the slope of G' is detected in the dynamic 171 temperature sweep curves). The strain of 1% was chosen within the linear viscoelastic regime, 172 which was established by performing dynamic strain amplitude sweeps on MC and MC/BS 173 solutions, over a strain range of 0.01 - 100%, at a constant angular frequency of 6.28 rad/s and 174 a temperature of 60°C. Each test was repeated at least twice to confirm reproducibility; 175 representative curves (rather than averages) are shown in the manuscript.

176

2.2.1.3 Langmuir trough measurements

177 Interfacial tension measurements were performed in a 50 mm diameter perfluoroalkoxy Petri 178 dish (19.6 cm² surface area and 20 mL volume of subphase), to study the adsorption of MC 179 and its interaction with BS at the air/water interface. All experiments were carried out under 180 constant stirring, at a fixed area, and at a temperature of 23 ± 2°C (room temperature). The 181 surface pressure (π) was measured by a Wilhelmy plate made of chromatographic paper 182 (Whatman International Ltd, Maidstone, UK) of 2.3 x 1.0 cm (length x width) and attached to 183 a calibrated Nima PS4 microbalance (Nima Technology Ltd, Coventry, UK). Prior to any 184 measurement, the trough was thoroughly cleaned with EtOH and CHCl₃ to remove organic 185 impurities, and then filled with ultrapure water (subphase). Surface-active contaminants, dust 186 and bubbles were all removed from the subphase by suction with a pump, and the subphase 187 was considered as clean when changes in surface pressure did not exceed ± 0.2 mN/m over 188 approximately two minutes.

189 **MC adsorption at the air/water interface.** Using a 1 mL syringe (Becton Dickinson, Madrid, 190 Spain) fitted with a 19 G x 1 ½ in. needle (Becton Dickinson, Madrid, Spain), a specific amount 191 of pure MC solution in ultrapure water was injected into the subphase, under constant stirring. 192 Surface pressure (π) was measured over time until it reached a plateau. Each experiment was 193 repeated at least twice; either a representative curve or an average measurement is shown.

BS interaction with a MC layer at the air/water interface. A MC layer was first formed at the air/water interface, by addition of a specific amount of MC aqueous solution into the clean and stirred water ($\pi_{MC} = 21 \pm 1 \text{ mN/m}$ with 0.5‰ w/w, and $\pi_{MC} = 18 \pm 2 \text{ mN/m}$ with 0.5×10⁻ 2 ‰ w/w). After film equilibration (ca. 1-2 hours), a specific amount of pure BS aqueous solution was injected beneath the MC layer. The corresponding changes in surface pressure (π) were recorded over time. Each experiment was repeated at least twice; either a representative curve or the average measurement is shown.

201

2.2.1.4 Ellipsometry

MC adsorption and interaction with BS at the air/water interface was further investigated by ellipsometry (Beaglehole Instruments, Wellington, New Zealand). Time-dependent measurements were performed with a 632.8 nm-wavelength laser hitting the surface at an incident angle of 50°. In this configuration, changes in the polarisation of light reflected by the 206 interface are measured over the 1 mm² area and ~1 μ m depth probed by the laser beam; 207 these changes can be correlated to the amount of material adsorbed at this interface over 208 time. The polarisation state of the incident light is composed of an *s*- and *p*-component (where 209 the *s*-component is oscillating parallel to the sample surface, and the *p*-one parallel to the 210 plane of incidence). The ratio of the reflectivity of these two components (r_s for the *s*-211 component and r_p for the *p*-component) characterises the polarisation change and is 212 expressed by the following equation:

213

$$\frac{r_p}{r_s}$$
 = tan(Ψ). e^{i Δ} (1)

214 where Ψ is the amplitude change and Δ the phase shift. In the thin film limit at the air/water 215 interface (i.e., film thickness << laser wavelength), Δ is found to be much more sensitive to 216 changes in the amount adsorbed at the interface than Ψ (Motschmann & Teppner, 2001). 217 Therefore, time-dependent changes in phase shift ($\Delta\Delta$) were measured, with $\Delta\Delta(t) = \Delta(t)$ -218 $\Delta(t_0)$, where $\Delta(t_0)$ is the phase shift at the beginning of a given experiment, namely, the phase 219 shift of the bare air/water interface (Δ_0) for MC adsorption and interaction with BS, at the 220 air/water interface. Changes in the phase shift are directly proportional to the amount of 221 material adsorbed at the interface (Motschmann & Teppner, 2001). In order to measure 222 simultaneously the surface pressure and phase shift for the same surface, the instrument was 223 mounted on top of the Petri dish, used as a Langmuir trough. Data were acquired at a rate of 224 0.2 Hz, using the Igor Pro software.

225

2.2.2 In vitro lipolysis studies

226

2.2.2.1 Preparation of MC-stabilised emulsion

227 MC (0.5% w/w) was dispersed into sunflower oil (15% w/w). Cold phosphate buffer (84.5% 228 w/w, at T < $T_{dissolution}$ = 10°C) was added to the oil phase and the mixture stirred for a few 229 minutes. The dispersion was then pre-emulsified at 11,000 rpm for 1 minute, using a high-230 shear mixer (T25 digital Ultra-Turrax, IKA[®]-Werke GmbH & Co. KG, Staufen, Germany). This 231 pre-emulsion was transferred into a 10 mL volume beaker in an ice bath and was sonicated at 232 a frequency of 20 kHz and amplitude of 70% for 5 minutes with a tip sonicator (SONOPULS HD 233 3100 ultrasonic homogeniser, microtip model: MS 73, BANDELIN electronic GmbH & Co. KG, 234 Berlin, Germany).

235

2.2.2.2 Simulation of the duodenal lipolysis environment

236 For each *in vitro* lipolysis experiment, the following model (Grundy, Wilde, Butterworth, Gray, 237 & Ellis, 2015) was employed to simulate the duodenum (small intestine) environment: 19 mL 238 of MC-stabilised emulsion was added to a thermostatically-controlled and mechanically-239 stirred reaction vessel at 37°C, followed by 15 mL of a BS aqueous solution (NaTC, NaTDC; 2.5, 240 25, 125 mM, in phosphate buffer). Then, 1 mL of NaCl (4.9 M, in ultrapure water) and 1 mL of 241 CaCl₂ (0.37 M, in ultrapure water) were added to the mixture, under continuous stirring. 242 Finally, 1.5 mL of either phosphate buffer (for the blank assay, used as a control) or freshly 243 prepared pancreatic lipase/co-lipase suspension (17 mg/mL, in phosphate buffer) (for the 244 lipolysis assay) were added. The final system was made up of 7.6% w/w lipid, 1, 10, or 50 mM 245 BS, 130 mM NaCl, 10 mM CaCl₂, and 0.68 mg/mL pancreatic lipase/co-lipase.

246

2.2.2.3 Optical microscopy

247 The structural changes induced on an MC-stabilised emulsion upon duodenal digestion, were 248 monitored over time by brightfield optical (Olympus BX61 microscope, Olympus France S.A.S., 249 Rungis, France) and confocal (Leica TCS SP2, DMIRE2 inverted, Leica Microsystems UK Ltd, 250 Milton Keynes, UK) microscopy. Prior to in vitro lipolysis studies, the pure emulsion was 251 characterised; then, the mixture modelling the duodenal environment was added to the 252 emulsion and samples measured at different time points (t = 5, 15, 30 and 60 min), to analyse 253 the evolution of emulsion droplet microstructure from the beginning to the end of duodenal 254 lipolysis. The influence of each component (NaTC, NaTDC, NaCl and CaCl₂) used individually 255 and together was assessed to better understand their impact on duodenal lipolysis. A blank 256 assay was also measured as a control to monitor changes over time in the absence of enzymes.

For confocal microscopy, prior to visualisation, samples were mixed with 1 mg/mL orlistat (prepared in dimethyl sulfoxide) to stop lipolysis, and then stained with 10 μ g/mL Nile red (prepared in dimethyl sulfoxide) and 20 μ g/mL calcofluor (prepared in ultrapure water), to detect lipids (red fluorescence) and MC (blue fluorescence), respectively. Samples were excited at 488 nm (for Nile red) and 405 nm (for calcofluor), and the fluorescence emitted by the samples was detected between 510 - 650 nm (for Nile red) and 410 - 480 nm (for calcofluor). Images were captured using objective lenses of 10×, 20× or 63×, and micrographs were compiled with the Olympus image analysis software (for optical microscopy, Olympus
France S.A.S., Rungis, France) and Fiji software ("Fiji," 2019) (for confocal microscopy).

266

2.2.2.4 pH-stat measurements

267 The rate and extent of lipolysis were evaluated by titrating the amount of FFA released from 268 an MC-stabilised emulsion with 0.1 M NaOH, at 37°C and pH 7.0, in conditions mimicking the 269 duodenal (small intestine) environment. Each assay was carried out over 1 hour of digestion, 270 using a pH-stat titration unit (848 Titrino plus, Metrohm AG, Herisau, Switzerland). The blank 271 experiment was performed as a control, to measure pH fluctuation in the absence of enzymes; 272 the volume of NaOH released during this assay was then subtracted from the data recorded 273 in the presence of pancreatic lipase/co-lipase (lipolysis assay). Each blank and lipolysis 274 experiment was repeated at least six times.

The volume of NaOH released during MC-stabilised emulsion digestion was converted into the
 percentage of FFA produced, using this equation:

277

%FFA (t) = 100 ×
$$\frac{V_{\text{NaOH}} (t) \cdot [\text{NaOH}] \cdot M_{\text{Lipid}}}{2 \cdot m_{\text{Lipid}}}$$
 (2)

where V_{NaOH} is the volume of NaOH required to neutralise the FFA produced over time, [NaOH] the concentration of the NaOH solution used, M_{Lipid} the molecular weight of the oil employed in this experiment (in our case, $M_{Sunflower oil} = 876$ g/mol (Sánchez, Maceiras, Cancela, & Rodríguez, 2012)), and m_{Lipid} the mass of triacylglycerol (TAG) initially present in the digestion vessel. This equation has been established considering the ideal case where the hydrolysis of one molecule of TAG leads to the formation of one molecule of MAG and two molecules of FFA. The results are shown as the proportion of FFA release as a function of time.

The pH-stat data were analysed with the GraphPad Prism software ("GraphPad Prism," 2019); statistical analysis was carried out using the two-way analysis of variance (ANOVA), followed by the Tukey post-test, with a 95% confidence level, meaning that differences were considered as statistically significant when P < 0.05.

3. Results

3.1 BS interaction with MC in the bulk

291 **MC viscoelastic behaviour.** The temperature-dependence of MC rheological properties was 292 investigated by performing dynamic temperature sweep measurements on MC solutions 293 prepared at concentrations ranging between 0.1 and 2.0% w/w (Figure S1).

294 At all the MC concentrations studied, a relatively flat region is observed for the storage 295 modulus (G') in the lower temperature range (ca. 20 - 40° C), followed by a steep increase 296 beyond a transition temperature (T_t) and a final plateau at high temperatures. As MC 297 concentration increases, the transition temperature from which G' starts to level off shifts 298 towards lower values (from 55°C at 0.1% w/w, to 37°C at 2.0% w/w). Below and above T_t, MC 299 behaves as a predominantly solid-like material over the whole range of temperatures studied 300 (G' dominates over G'' over the range of frequencies measured), and above T_t , both moduli 301 increase and are still independent of frequency (data not shown) (Funami et al., 2007; L. Li et 302 al., 2001; Lin Li, 2002); the transition temperature thereby corresponds to a weak-to-strong 303 gel transition. The increase in MC concentration also induces a relatively weak change in MC 304 elastic properties (G') at low temperatures, and a much more significant one in the high 305 temperature region, in agreement with previous studies (Nasatto et al., 2015a).

306 The gelation of MC – whose chains are arranged as 'bundles' at room temperature (or packed 307 'strands' held together by packing of unsubstituted regions and the hydrophobically-driven 308 aggregation of methyl groups in regions of denser substitution) - has been postulated to 309 follow two steps (Haque & Morris, 1993; Sarkar, 1995; Desbrières, Hirrien, & Rinaudo, 1998; 310 Hirrien, Chevillard, Desbrières, Axelos, & Rinaudo, 1998; Kobayashi, Huang, & Lodge, 1999; L. 311 Li et al., 2001, 2002; Lin Li, 2002; Lin Li, Wang, & Xu, 2003; Funami et al., 2007; Torcello-Gómez 312 & Foster, 2014; Torcello-Gómez et al., 2015; Nasatto et al., 2015a; Isa Ziembowicz et al., 2019): 313 upon heating, MC strands separate, allowing intermolecular associations to form between MC 314 hydrophobic (methyl) groups, therefore inducing the formation of a strong, physical gel 315 network; at low temperatures, these hydrophobic polymer-polymer interactions take place to 316 a much lower extent because of water molecules surrounding MC methyl moieties (via 317 hydrogen bonds), thus resulting in the swelling of 'bundles' and the formation of a softer, 318 weaker gel. The effect of MC concentration on its rheological properties is therefore attributed to the increase in the number of methyl groups in solution, resulting in a largernumber of hydrophobic interactions from lower temperatures.

Effect of BS on MC viscoelastic behaviour. The impact of the two BS on MC rheological properties was assessed by following the dynamic moduli (G', G'') of a 1.0% w/w MC solution over a range of temperatures and frequencies (Figure 3). The evolution of the transition temperature (T_t , from which the increase in G' becomes steeper) and of both dynamic moduli (G', G'') are shown as a function of BS concentration in Figures 4A and 4B, respectively.



326

Figure 3: (A, B) Temperature-dependent evolution of the storage modulus (G') obtained from dynamic temperature sweeps, and (C, D) angular frequency-dependent evolution of the dynamic moduli: (•) G', the storage modulus, (○) G", the loss modulus, obtained from dynamic frequency sweeps performed at a constant temperature of 60°C, on a 1.0% w/w MC aqueous solution containing increasing amounts (1, 10, 25, 50, 100, 200, 500 mmol/kg) of BS: (A, C) NaTC, (B, D) NaTDC. The curves obtained in the absence of BS are also shown for comparison.



Figure 4: Evolution of MC transition temperature (T_t) (A) and dynamic moduli: (•) G', the storage modulus, (o) G'', the loss modulus, obtained at an angular frequency of 1 rad/s (B), as a function of the concentration in BS: NaTC, NaTDC. The transition temperature (T_t) is the temperature from which G' starts changing. These data are extracted from, respectively, (A) dynamic temperature sweeps performed over a temperature range of 20 - 80°C (Figures 3, A, B), and (B) dynamic frequency sweeps performed over an angular frequency range of 0.1 - 100 rad/s, at a constant temperature of 60°C (Figures 3, C, D).

332

339 In the presence of BS, the dynamic temperature sweeps of MC solutions show a similar profile 340 as the pure MC solution, namely, a moderate increase in G' followed by a sharp rise (Figures 341 3, A, B). However, BS have a significant impact on MC rheological properties, leading to a 342 notable, and gradual increase in the transition temperature from around 50°C, in the absence 343 of BS, to 53°C with 500 mmol/kg NaTC and 58°C with 500 mmol/kg NaTDC (Figure 4A). In 344 addition, both BS (from the lowest concentration studied of 1 mmol/kg) induce a drop in MC 345 viscoelasticity (G') at all temperatures studied, most visibly at high temperatures (Figures 3, A, B). At physiological temperature $(37^{\circ}C)$, a decrease from G' = 10 Pa in the absence of BS, to 346 347 G' = 5 and 2 Pa in the presence of 1 mmol/kg of, respectively, NaTC and NaTDC, is observed. 348 Dynamic frequency sweeps performed at 60°C, where MC forms a strong gel and changes 349 caused by BS are most visible (Figures 3, C, D), reveal a 10-fold decrease in G', from ca. 280 Pa 350 in the absence of BS, to ca. 20 Pa with the highest concentration of BS studied, at a frequency 351 of 1 rad/s (Figure 4B). In addition, G' shows an increasing dependence on frequency with the 352 addition of BS, more notably so with NaTDC. Overall therefore, the presence of the BS 353 converts MC gel into a less solid-like material. Comparing the two BS, it is clear that NaTDC 354 has a much stronger impact; for instance, only 10 mmol/kg of NaTDC are needed to significantly reduce the value of the storage modulus (G') (Figures 3D and 4B), while 25 355 356 mmol/kg of NaTC are required to induce the same effect (Figures 3C and 4B). Similar 357 observations have been reported elsewhere (Torcello-Gómez et al., 2015).

Overall, over the whole temperature range studied, MC behaves as a gel whose strength increases with temperature. The addition of BS induces a transition to a softer material (lower elastic modulus (G')), both above and below MC transition temperature (T_t); in addition, this transition occurs at lower concentrations of NaTDC, compared to NaTC.

362 3.2 BS interfacial properties in the presence of MC

363 **MC adsorption dynamics at the air/water interface.** MC behaviour at the bare air/water 364 interface was studied using both a Langmuir trough and ellipsometer, by monitoring the time-365 dependent evolution of the surface pressure (π) and phase shift ($\Delta\Delta$), respectively, upon 366 injection into the water subphase of either successive quantities of MC (0.5×10⁻¹, 0.25 and 367 0.5‰ w/w (Figure S2); 0.5×10⁻², 0.25×10⁻¹ and 0.5×10⁻¹‰ w/w (Figure S3)), or fixed amounts 368 over a longer period of time (0.5×10⁻³, 0.5×10⁻², 0.5×10⁻¹ or 0.5‰ w/w) (Figure S4).

369 Upon addition of 0.5×10⁻¹‰ w/w MC into the aqueous subphase, the surface pressure 370 increases until reaching a near-plateau at π = 19 ± 1 mN/m, which stays relatively constant 371 with following injections ($\pi = 19 \pm 1 \text{ mN/m}$ at 0.25% w/w, and $\pi = 18 \pm 3 \text{ mN/m}$ at 0.5% w/w) 372 (Figure S2A). With the same injection sequence, the ellipsometry phase shift, which is 373 measured at the same time as the surface pressure and relates to the amount of material 374 adsorbed at the interface (Motschmann & Teppner, 2001), exhibits the same trend as the 375 surface pressure (Figure S2B): it reaches a value of $\Delta\Delta = 0.033^\circ$, which then slightly increases 376 to $\Delta\Delta$ = 0.035° at 0.25‰ w/w and $\Delta\Delta$ = 0.036 at 0.5‰ w/w. Both measurements thus show 377 that MC adsorbs at the air/water interface up to a saturation point, independently of its 378 concentration in the bulk. The two experiments differ, nevertheless, by the presence of peaks 379 of surface pressure visible straight after MC injection, not detected in the phase shifts, which 380 could be explained by an initial strong adsorption, followed by a relaxation process as the 381 polymer rearranges at the air/water interface, changing conformation (Graham & Phillips, 382 1979). These transient surface pressure peaks were also observed in a previous study with BS injected under the air/water interface (Pabois et al., 2019). The trends in surface pressure 383 384 (Figure S3A) and phase shift (Figure S3B) are reproduced with lower amounts of MC (0.5×10⁻ 385 2 , 0.25×10⁻¹, and 0.5×10⁻¹‰ w/w) injected into water.

386 In order to study the kinetics of adsorption of MC molecules at the air/water interface, surface 387 pressure measurements were performed over longer periods of time (Figure S4). Results show 388 that, above 0.5×10^{-2} % w/w, the same equilibrium surface pressure ($\pi = 17 \pm 1 \text{ mN/m}$) is

389 always reached, irrespective of MC concentration, whereas a much lower value is obtained at 390 the lowest concentration studied of 0.5×10^{-3} % w/w ($\pi = 10 \pm 0.4$ mN/m). Arboleya and Wilde 391 (Arboleya & Wilde, 2005) also observed a saturation point from a similar MC concentration 392 (i.e., 1×10⁻²‰ w/w), and obtained comparable interfacial tension values. Furthermore, as MC concentration decreases, the surface pressure rises at a slower rate: a change in surface 393 394 pressure is immediately observed after injection of both 0.5×10^{-1} and 0.5% w/w, while a lag 395 period of about 3 and 40 min is seen with solutions containing 0.5×10^{-2} and 0.5×10^{-3} % w/w 396 MC, respectively. The amount injected into the aqueous subphase thus affects MC adsorption 397 rate and extent, such that the lower the concentration, the slower the adsorption process and 398 the lower the quantity of material adsorbed, therefore indicating a diffusion-controlled 399 adsorption mechanism, as already observed elsewhere with hydroxypropyl MC (Avranas & 400 Tasopoulos, 2000; Camino, Pérez, Sanchez, Rodriguez Patino, & Pilosof, 2009; Pérez, Sánchez, 401 Pilosof, & Rodríguez Patino, 2008; Wollenweber, Makievski, Miller, & Daniels, 2000). In the 402 literature, MC adsorption has been suggested to occur in three stages: MC first slowly diffuses 403 from the bulk phase to the sub-surface region and then adsorbs at the air/water interface, 404 while undergoing conformational changes (Arboleya & Wilde, 2005).

405 All these results are consistent with data reported elsewhere (Nasatto et al., 2014; Pizones
406 Ruiz-Henestrosa et al., 2017).

407 BS interaction with a MC layer at the air/water interface. The interfacial behaviour of the 408 two selected BS (NaTC and NaTDC) in the presence of a MC film at the air/water interface was 409 then evaluated, by injecting BS below the polysaccharide layer. Measurements were carried 410 out either by adding increasing amounts of BS every hour (2, 4, 6, 8 and 10 mM) (Figures 5 411 and S5) or by injecting fixed concentrations and measuring over longer times (1, 5 or 10 mM) 412 (Figures 6 and S6). These BS concentrations were selected to be below, around, and above 413 their critical micelle concentration (CMC), which is 4 – 7 mM for NaTC (gradual micellisation 414 process) and 2 mM for NaTDC in ultrapure water (data not shown) (Matsuoka, Maeda, & 415 Moroi, 2003). Prior to BS injection, a saturated film of MC at the interface was formed by 416 injecting it into the water subphase, at either 0.5‰ w/w (Figures 5, 6 and S6) or $0.5.10^{-2}$ ‰ 417 w/w (Figure S5).



418

Figure 5: Time-dependent evolution of (A, B) the surface pressure (π) measured in a Langmuir trough, and (C, D) phase shift ($\Delta\Delta(t) = \Delta(t) - \Delta_0$) measured by ellipsometry, upon successive injections of either (A, C) NaTC or (B, D) NaTDC into the aqueous subphase (at 23 ± 2°C). The first increase in surface pressure corresponds to the adsorption of MC at the air/water interface, which was added into water at a concentration of 0.5‰ w/w ($\pi_{MC} = 21 \pm 1 \text{ mN/m}$, $\Delta\Delta_{MC} = 0.039 \pm 0.005^\circ$). Each addition of BS is shown by an arrow, together with the corresponding BS concentration achieved in the subphase. Each experiment was reproduced twice, and a representative measurement was selected for each experiment.

425 The evolution of the surface pressure is quite different for the two BS (Figures 5, A, B): while 426 the successive injections of NaTC lead to a continuous increase in surface pressure (up to π = 427 23 ± 0.5 mN/m at 10 mM) (Figure 5A), with NaTDC, a steep rise to π = 25 ± 1 mN/m (at 2 mM), 428 followed by a gradual drop to π = 22 ± 1 mN/m (at 10 mM), is observed (Figure 5B). These 429 trends are also obtained with a lower amount of MC at the air/water interface (Figure S5). The 430 ellipsometry phase shift obtained in parallel follows the same trends (Figures 5, C, D): with 431 NaTC, it gradually increases up to $\Delta\Delta = 0.045 \pm 0.003^{\circ}$ upon successive additions of BS into the 432 subphase (Figure 5C); instead, the injection of 2 mM NaTDC into the water induces a sharp 433 increase to $\Delta\Delta$ = 0.047 ± 0.003°, followed by a decrease to $\Delta\Delta$ = 0.042 ± 0.001° from 4 mM 434 (Figure 5D). As observed with successive injections of MC, temporary surface pressure peaks 435 are also present after each addition of BS; here again, these peaks could be attributed to MC

436 film compression and subsequent relaxation, induced by BS adsorption (Graham & Phillips,

437 1979).





Figure 6: Evolution of the surface pressure ($\Delta \pi = \pi_{Equilibrium} - \pi_{MC}$) as a function of BS concentration, measured in a Langmuir trough, upon injection of fixed concentrations (1, 5, 10 mM) of BS (NaTC, NaTDC) into the aqueous subphase (at 23 ± 2°C). 0.5‰ w/w MC were injected into water to form a layer at the air/water interface, at $\pi_{MC} = 21 \pm 1$ mN/m. These data were extracted from individual BS injections measurements (Figure S6). Each experiment was reproduced at least twice, and the average measurement was selected for each BS at each concentration.

444 Upon injection of fixed BS concentrations, the surface pressure increases sharply over time 445 until reaching a plateau value, independently of the BS type and concentration (Figure S6). 446 The surface pressure values achieved at equilibrium are summarised in Figure 6, showing $\Delta \pi$ 447 = $\pi_{Equilibrium}$ - π_{MC} , where π_{MC} is the initial MC layer surface pressure (π_{MC} = 21 ± 1 mN/m). The 448 surface pressure changes induced by the two BS are relatively small, in agreement with 449 previous studies performed on the interaction of a hydroxypropyl MC layer with bile extract 450 (Pizones Ruiz-Henestrosa et al., 2017). At 1 and 5 mM, NaTDC induces a higher increase in 451 surface pressure ($\Delta \pi$ = 5 ± 0.4 mN/m at 1 mM, and $\Delta \pi$ = 6 ± 0.2 mN/m at 5 mM), compared to 452 NaTC ($\Delta \pi$ = 3 ± 0.3 mN/m at 1 mM, and $\Delta \pi$ = 4 ± 1 mN/m at 5 mM); at high BS concentration 453 (10 mM), the opposite trend is observed ($\Delta \pi = 6 \pm 0.4$ mN/m for NaTC, and $\Delta \pi = 4 \pm 0.3$ mN/m 454 for NaTDC).

455 **3.3 Effect of BS structure and concentration on the duodenal digestion of an MC-stabilised** 456 emulsion

A range of *in vitro* duodenal lipolysis studies was carried out on a sunflower oil emulsion stabilised by MC. Before reaching the small intestine, ingested fat droplets pass through simulated oral and gastric digestion, where their physicochemical and structural properties are significantly affected; however, because our main aim is to understand BS roles during lipolysis, the work performed here focuses on the duodenum part of the lipolysis process, where BS are acting.

Evolution of emulsion droplets microstructure. The structure of the pure MC-stabilised emulsion droplets was first characterised using both optical and confocal microscopy (Figure S7). Optical microscopy demonstrates that emulsion droplets are uniformly dispersed with a size ranging between 2 and 5 μ m, and with a small number of larger droplets around 10 μ m (Figure S7A). Confocal microscopy highlights the presence of a MC network (stained in blue with calcofluor) in the bulk and at the interface of emulsion droplets (stained in red with Nile red) (Figure S7B).

470 In vitro lipolysis studies were performed on the emulsion by adding the digestive medium and 471 monitoring the structural changes of the emulsion droplets by microscopy (Figures 7, S8 and 472 S9). Using brightfield optical microscopy, the influence of both BS type and concentration on 473 the structure of MC-stabilised emulsion droplets was assessed in control assays (no enzyme), 474 as well as the effect of enzymes (lipolysis assays) (Figure 7). In the absence of enzymes (blank 475 assays), the emulsion droplets microstructure is affected by the digestive fluid, as revealed by 476 the occurrence of droplets flocculation, and some - limited - coalescence, which is more 477 visible with NaTDC, and particularly evident for both BS at high concentration (10 mM). Upon 478 the addition of enzymes (lipolysis assays), flocculation occurs to a higher extent, and droplet 479 coalescence (size increase) is observed in all samples, to a larger extent, again, with NaTDC. 480 To further elucidate the mechanism of digestion of an MC-stabilised emulsion, the influence 481 of the different components of the digestive fluid (NaCl, CaCl₂ and BS) on droplet stability was 482 also evaluated (Figures S8 and S9). Brightfield optical micrographs show that extensive 483 flocculation occurs when both BS and salts are present, which suggests that the association of 484 BS with the different salts (NaCl, CaCl₂) is responsible for the droplet aggregation observed in 485 Figure 7.



486

Figure 7: Time-dependent evolution of the microstructure of MC-stabilised emulsion droplets in the presence of BS: NaTC,
 NaTDC, used at 1 and 10 mM, under duodenal digestion conditions (at 37°C). MC-stabilised emulsion was made up of 0.5%
 MC and 15% sunflower oil. Both blank (without enzymes) and lipolysis (with enzymes) assays were performed to assess,
 respectively, the effect of BS type and concentration on the droplets stability, and of enzymes on the droplets
 microstructure. Microscopy observations were made at t = 5 and 60 minutes. The scale bar is 200 μm.

492 This in vitro lipolysis study was complemented with micro-structural assessment of the 493 emulsion droplets with confocal microscopy, to determine the localisation of MC throughout 494 the emulsion and its evolution during lipid digestion (Figures 8 and 9). Based on our pH-stat 495 results (see the following section), 50 mM BS was used here, as it shows the higher extent of 496 FFA release. The images obtained suggest that the addition of digestive fluid not only breaks 497 down the network of MC, but also displaces it from the lipid/water interface (Figure 8); 498 interestingly, MC bulk network is disrupted to a higher extent in the presence of NaTDC, 499 compared to NaTC. Additionally, the lipid droplets become non-spherical with "rough" 500 surfaces, compared to the initial emulsion (Figures 8 and 9). This demonstrates coalescence

and may be an indication of fats being digested by enzymes; in particular, small oil droplets
were seen to flocculate or coalesce onto the surface of larger droplets (Figure 9A) and areas
with an undefined oil/water interface suggest the presence of digestion products (Figure 9B).



Figure 8: Time-dependent evolution of the microstructure of MC-stabilised emulsion droplets in the presence of 50 mM
 BS: NaTC, NaTDC, under duodenal digestion conditions (at 37°C). MC-stabilised emulsion was made up of 0.5% MC and
 15% sunflower oil. The lipid droplets are stained in red (with Nile red), while MC is stained in blue (with calcofluor).
 Microscopy observations were made at t = 5, 15, 30 and 60 minutes, to compare the structural changes occurring during
 digestion; at each time point, orlistat was used to inhibit lipolysis. The scale bar is 150 μm.





- 510
- Figure 9: (A) Cross-section confocal image of MC-stabilised emulsion droplets in the presence of 50 mM NaTC, under duodenal digestion conditions (at 37°C). The microscopy observation was made at t = 15 minutes. (B) MC-stabilised emulsion droplets in the presence of 50 mM NaTDC, under duodenal digestion conditions (at 37°C). Insoluble lipolysis products seem to be presumably present at the fat droplet interface (see the arrow). MC-stabilised emulsion was made up of 0.5% MC and 15% sunflower oil. The lipid droplets are stained in red (with Nile red), while MC is stained in blue (with calcofluor). The scale bar is 10 μm.

Quantification of FFA release from the MC-stabilised emulsion. The ability of NaTC and NaTDC to promote or inhibit the duodenal digestion of an MC-stabilised emulsion was compared by monitoring the release of FFA (*%FFA*) over time with the pH-stat method (Y. Li, Hu, & McClements, 2011) (Figure 10). The effect of BS concentration on the rate of lipolysis and its extent was also evaluated using the two BS at both 10 and 50 mM.





Figure 10: (A) Proportion of FFA released (%*FFA*) over time from an MC-stabilised emulsion, using two different BS: NaTC, NaTDC, at two different concentrations: (\Box) 10 and (\blacksquare) 50 mM, under duodenal digestion conditions (at 37°C). (B) Proportion of FFA released (%*FFA*) after 1 hour of digestion of an MC-stabilised emulsion, using the two BS, at 10 and 50 mM, under duodenal digestion conditions (at 37°C). Statistical significance was determined using the two-way ANOVA, followed by the Tukey post-test (**** indicates P < 0.0001, i.e., differences are extremely significant). MC-stabilised emulsion was made up of 0.5% MC and 15% sunflower oil.

529 Independently of the BS type and concentration, the proportion of FFA generated during 530 lipolysis increases steeply after the addition of enzymes (Figure 10A); this rapid initial rate of 531 lipolysis, already observed elsewhere (Bellesi, Martinez, Pizones Ruiz-Henestrosa, & Pilosof, 532 2016; McClements & Li, 2010a), can be attributed to the immediate adsorption of lipase/co-533 lipase onto fat droplets surfaces, which then triggers TAG break-down and thus lipid digestion. 534 After a certain time (t = 0.07 h with both BS at 10 mM, t = 0.18 h with 50 mM NaTDC and t = 535 0.24 h with 50 mM NaTC), the release of FFA starts slowing down, until it reaches a near-536 plateau. This decrease in the rate of lipolysis can be explained by the accumulation of lipolysis 537 products at the oil/water interface during the process of fat digestion (Patton & Carey, 1979; 538 P. Reis, Holmberg, Watzke, Leser, & Miller, 2009; P. Reis et al., 2008; P. M. Reis et al., 2008), 539 which then leads to the inhibition of enzymes binding to the substrate, as previously 540 demonstrated (Bellesi et al., 2016; Borel et al., 1994). Increasing BS concentration from 10 to 541 50 mM leads to a significant increase in the percentage of FFA produced, for both BS (P_{NaTC} <

- 542 0.0001 and P_{NaTDC} < 0.0001) (Figure 10B): more specifically, a 14% and 9% increase is obtained 543 with, respectively, NaTC and NaTDC. This can be attributed to the larger amount of BS 544 micelles, which can solubilise a larger amount of FFA released, thereby preventing droplets 545 surface saturation by these products (Wilde & Chu, 2011). While no significant differences are 546 observed between the two BS at the lowest concentration (10 mM) (%FFA t = 1h = 6 ± 1 % for 547 both NaTC and NaTDC; P_{10 mM} = 0.4), a significant difference is seen at high concentration (50 548 mM), with NaTC inducing a higher extent of lipolysis (%FFA t = 1h = 20 ± 1 % and 14 ± 1 % for,
- 549 respectively, NaTC and NaTDC; $P_{50 \text{ mM}} < 0.0001$).

550 **4. Discussion**

551 The objective of this study was to investigate the interactions of MC with BS, in 552 particular the ability of MC to inhibit BS activity, and thus to shed light on the mechanism of 553 lipid digestion regulation by MC – a dietary fibre with a proven potential to lower cholesterol 554 levels (Agostoni et al., 2010). Bulk (rheology) and interfacial (surface pressure measurements 555 and ellipsometry) studies were carried out to characterise the interactions between these two 556 components in the bulk and at the interface, while in vitro lipolysis (microscopy, pH-stat) 557 experiments were performed to link these interactions to the lipid digestion of an MC-558 stabilised emulsion. The two BS, which differ by the presence (NaTC) or absence (NaTDC) of a 559 hydroxyl group on their steroid skeleton (Figure 2) and constitute 20% of human bile (Staggers, 560 Hernell, Stafford, & Carey, 1990), were chosen for this study, as they have been reported to 561 exhibit different interfacial behaviours, hypothesised to explain the contrasting roles they play 562 during the process of lipolysis (Pabois et al., 2019; Parker et al., 2014).

563 **4.1 Interaction between MC and BS in the bulk and at the interface**

564 The impact of BS on MC rheological properties was investigated to explore the 565 interaction of BS with MC in the bulk, where MC is present in excess. Increasing the amount 566 of BS in solution led to a notable shift in the transition temperature (T_t) to higher values, as 567 well as a gradual drop in viscoelastic properties, which were more substantial with NaTDC 568 (Figures 3 and 4). In particular, MC – which presents predominantly solid-like properties in the 569 absence of BS – turned into a softer gel above a threshold concentration of BS (25 mmol/kg 570 for NaTC vs. 10 mmol/kg for NaTDC, at 60°C) (Figures 3, C, D and 4B). MC gelation occurs via 571 the association of the hydrophobic (methyl) moieties (Haque & Morris, 1993; Sarkar, 1995; 572 Desbrières, Hirrien, & Rinaudo, 1998; Hirrien, Chevillard, Desbrières, Axelos, & Rinaudo, 1998; 573 Kobayashi, Huang, & Lodge, 1999; L. Li et al., 2001, 2002; Lin Li, 2002; Lin Li, Wang, & Xu, 2003; 574 Funami et al., 2007; Torcello-Gómez & Foster, 2014; Torcello-Gómez et al., 2015; Nasatto et 575 al., 2015a; Isa Ziembowicz et al., 2019). The presence of BS and its association with MC may 576 thus prevent hydrophobic groups from assembling with each other, thus weakening the gels 577 or hindering gelation altogether. The stronger effect observed with NaTDC may be attributed 578 to its higher hydrophobicity (Armstrong & Carey, 1982), which may result in a more efficient 579 connection between BS and MC hydrophobic regions (Torcello-Gómez et al., 2015). Overall, these rheological measurements reveal the presence of strong interactions between BS and the dietary fibre, which have a substantial impact on MC viscosity; the presence (NaTC) or absence (NaTDC) of a hydroxyl group on BS steroid backbone impacts this behaviour considerably.

The interfacial properties of BS in the presence of a MC layer formed at the air/water 584 585 interface were then studied to determine the interactions occurring when a BS molecule 586 approaches a fat droplet stabilised by MC. Studies with a Langmuir trough set-up (Figures 5, 587 A, B, S5, 6 and S6) combined to ellipsometry (Figures 5, C, D) demonstrate that the two BS 588 behave quite differently when injected beneath an almost-saturated MC film: NaTC was 589 shown to gradually adsorb at the interface with increasing concentration, whereas NaTDC first 590 adsorbed at low concentrations (up to 2 - 3 mM) and then desorbed above 4 - 5 mM. This 591 contrasting interfacial behaviour correlates with their micellisation behaviour, which occurs 592 over 4 – 7 mM for NaTC and at 2 mM for NaTDC. Similar differences have been observed when 593 BS were injected below a phospholipid monolayer (Pabois et al., 2019). Nevertheless, BS were 594 found to adsorb and/or desorb to a much lower extent in the presence of a MC film, compared 595 to the phospholipid monolayer (surface pressures changes as high as 30 mN/m were 596 monitored in the presence of the lipid film, whereas an increase of up to 10 mN/m was 597 observed with the MC layer). This may in part be explained by the likely presence of MC excess 598 in the bulk, which could interact with BS and therefore limit their adsorption at the interface.

599 **4.2 Impact of BS/MC interactions on fat digestion**

600 Next, we performed *in vitro* lipolysis studies by following the evolution of the structure 601 of an MC-stabilised emulsion with optical and confocal microscopy, to compare the effect of 602 the two BS on the droplets (Figures S7A and 7) and shed light on the behaviour of MC during 603 emulsion digestion (Figures S7B, 8 and 9). The characterisation of the MC-stabilised emulsion 604 by confocal microscopy clearly demonstrates that fat droplets are entrapped in a network of 605 MC present in excess in the bulk, which may be responsible for the stabilisation of the 606 emulsion against droplets flocculation or coalescence (Figure S7B). Optical microscopy images 607 (Figure 7) demonstrate that, even in the absence of digestive enzymes, the presence of both 608 BS destabilises the emulsion, inducing some flocculation; upon the addition of lipases, 609 droplets destabilisation (namely, flocculation and coalescence) was found to occur to a large

610 extent, and more markedly with NaTDC, compared to NaTC. Confocal microscopy images 611 (Figure 8) suggest that flocculation and coalescence observed during lipolysis are due to the 612 MC network being broken down and removed from the lipid/water interface. The better 613 ability of NaTDC to induce coalescence could therefore be explained by its higher capacity to 614 disturb MC bulk network (as observed by confocal microscopy observations), which, in turn, 615 could be attributed to its stronger interactions with MC (as seen from rheology 616 measurements) and higher propensity to desorb from the interface at lower concentrations 617 (as detected by interfacial measurements). While the displacement of MC from the interface 618 by BS may facilitate the access of BS and enzymes to the lipid droplets surface, the network of 619 MC remaining in the bulk may also trap BS (via hydrophobic interactions) and thus prevent 620 them from removing insoluble lipolysis products, which could explain how MC hinders lipase 621 activity. Emulsion droplets coalescence (and thus the decrease in droplets surface area), which 622 occurs under duodenal digestion conditions, could also explain the slowing down of lipolysis.

The capacity of the two BS to promote or inhibit MC-stabilised emulsion digestion was then explored with the pH-stat method; results revealed that NaTC favoured FFA release to a higher extent than NaTDC (at 50 mM) (Figure 10). The lower proportion of FFA release obtained with NaTDC can be explained by its higher efficiency at binding to MC network (as suggested by rheology measurements), which may result in this BS becoming trapped in the bulk and therefore not contributing to the lipolysis process.

629 **5. Conclusion**

The demonstrated potential of MC, a dietary fibre, to regulate lipolysis is thought to be due to its ability to reduce BS activity by sequestration; the objective of this work was to compare the interactions of two structurally different BS, NaTC and NaTDC, with MC, and to determine their impact on the digestion of an emulsion stabilised by this polysaccharide. These findings are key to establishing a molecular-level, mechanistic understanding of the ability of MC to lower fat absorption.

636 Both BS were found to decrease the elasticity of MC gels, and to shift the transition 637 temperature (T_t) to stiffer gels to higher temperatures, to a higher extent with NaTDC. When 638 injected below a MC film at the air/water interface, NaTC remained adsorbed at the interface 639 over a wider concentration range, compared to NaTDC, which desorbed at a lower 640 concentration, correlating with the onset of micellisation in the bulk (between 4 - 7 mM for 641 NaTC and at 2 mM for NaTDC). The small difference in the two BS molecular structure, 642 specifically their bile acid portion, is responsible for their contrasting behaviour, and explains 643 the different results obtained during *in vitro* lipid digestion: (i) NaTDC has a higher propensity 644 to disrupt MC network in the bulk and interfacial layer, and thus induces more extensive 645 emulsion destabilisation (as seen from optical and confocal microscopy); (ii) the release of FFA 646 is lower with NaTDC, which can be linked to its higher capacity to bind MC in the bulk, resulting 647 in BS being unable to access the oil/water interface. Overall, it is clear that BS architectural 648 diversity - whose importance is often neglected - plays a key role in their functionalities 649 during fat digestion.

This work is a first step towards unlocking the mechanism of lipid digestion regulation by MC. Additional structural studies, in particular with techniques such as small-angle neutron scattering and neutron reflectometry, should bring significant knowledge to the area, in particular to examine the structure of MC in the presence of BS and the evolution of the fat droplet interface during digestion; this is the focus of current work. Building upon these results, the next challenge will be to engineer MC-stabilised lipid emulsions with appetitesuppressing or satiety-enhancing properties and evaluate their effect on cholesterol levels.

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670 **Declarations of interest**

671 None

673 **References**

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