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Influence of pH on foaming and rheological properties of aerated high sugar system with egg white protein and hydroxypropylmethylcellulose

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- 1 Influence of pH on foaming and rheological properties of aerated high sugar system with egg
- 2 white protein and hydroxypropylmethylcellulose

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Abstract

The objective of this study was to evaluate the effects of the total biopolymer (egg white protein - EW 3 and hydroxypropylmethylcellulose - HPMC) concentration (1.4 to 5.6 g/100 g of sugar) and 4 EW/HPMC ratio (2/1 to 18/1 g/g) on the apparent viscosity before whipping, foaming capacity 5 (density and overrun) and foam rheological properties (G', G" and δ) of sugar/EW/HPMC mixtures 6 using a central composite rotatable design (CCRD). The conditions to obtain intermediate apparent 7 viscosity, high foaming capacity, elastic and solid behaviour were total biopolymer concentration 5.0 8 g/100 g of sugar and EW/HPMC ratio 14/1 (g/g). Under these conditions, experiments were carried 9 10 out to evaluate the effect of interactions between EW and HPMC at pH 3.0, 4.5 and 6.0 on the foaming and rheological properties. The greatest foaming capacity, elastic and solid behaviour, with 11 no liquid drainage, were obtained at pH 3.0. At pH 4.5, foams possessed monodisperse bubble size 12 distribution and viscoelastic behaviour, leading to better stability with respect to disproportionation 13 14 and coalescence compared to foams at pH 3.0. At pH 6.0, foam showed the poorest foaming properties and viscous behaviour. The interactions between EW and HPMC in aerated confectionery 15 at different pH affect foaming and rheological properties. 16

17

18 Keywords: foam, biopolymer interaction, elastic behaviour, semi solid, stability

19 **1. Introduction**

Food foam is a dispersion of air bubbles in a continuous liquid phase or solid phase, stabilized by surface-active ingredients (Damodaran, 2008). It is a thermodynamically unstable system where drainage, coalescence and disproportionation are the factors that affect its stability.

Liquid drainage from thin film lamella due to gravity leads to coalescence of adjacent bubbles via rupture of the lamella film between them. Disproportionation is the diffusion of gas from small to large bubble or to atmosphere. Even in the absence of liquid drainage and coalescence, disproportionation is difficult to prevent because the pressure in a small bubble is greater than in larger ones (Damodaran, 2005; Murray & Ettelaie, 2004; Walstra & van Vliet, 2008).

Many foods such as bakery products, beverages, mousses, ice cream and confectionery 28 items are foams. The aeration process results in changes in the texture and rheology providing a 29 30 different mouthfeel and appearance (Campbell & Mougeot, 1999). Aerated confectionery such as marshmallows and nougats are manufactured using high-boiled sugar syrup and surface-active 31 32 agents such as proteins, which can be combined with polysaccharides (Lees & Jackson, 1992). In 33 confectionery products, to prevent microbial growth at ambient temperature, the product has to be 34 higher than 76 g of sugar/100 g. At this level of sugar, to avoid crystal formation, part of the sucrose should be replaced by others sugars such as glucose syrup and/or invert sugar to increase the 35 36 system solubility (Stansell, 1995).

37 Sugars, proteins and polysaccharide may interact with each other, affecting foaming capacity, foam stability and rheological properties. Sugars influence the functional properties of proteins such 38 as adsorption and gelation. Interaction with sucrose decreases ovalbumin surface activity at pH 7.0, 39 whereas for sodium caseinate there is an increase in the protein surface activity (Antipova, 40 Semenova, & Belyakova, 1999). Sucrose concentration influences the gelation rate of whey proteins 41 (Bryant & Mcclements, 2000) and the adsorption rate of bovine serum albumin (BSA) to air-aqueous 42 interfaces. The difference in adsorption rate of BSA depends on the type and concentration of sugar. 43 The process of adsorption may be attributed to an increase in aqueous phase viscosity and in 44 45 protein surface hydrophilicity or to the preferential interactions of protein with solvent components (Guzey, Mcclements, & Weiss, 2003). High sugar concentration (> 60 g of sugar/100 g) improves the 46 47 stability of aerated confectionery by decreasing the drainage rate by the increasing the liquid continuous phase viscosity, but decreases the foam overrun (Lau & Dickinson, 2005; Raikos, 48 49 Campbell, & Euston, 2007).

In order to perform as a good foaming agent, proteins should be able to adsorb rapidly at the air-water interface, undergo rapid conformational change and rearrangement at the interface and form a cohesive viscoelastic film via intermolecular interactions (Damodaran, 2008; Dickinson, 2011; Mine, 1995). Egg white (EW) protein is used as surface-active ingredient to produce marshmallow and nougat. Its excellent foaming properties are due to the interaction between its protein components. Globulins contribute to foamability, ovomucoid prevents foam drainage by imparting high viscosity, and lysozyme forms complexes with other proteins enhancing film strength and foam
stability (Dickinson, 2011; Mine, 1995).

Polysaccharides act as thickening, water-holding or gelling agents and their use 58 can increase foam stability by either increasing the viscosity of the continuous phase or via forming a 59 dimensional network (Dickinson, 2003; Walsh, Russell, & 60 three Fitzgerald, 2008). Hydroxypropylmethylcellulose (HPMC) is a polysaccharide that has some surface activity due to 61 presence of the methyl (hydrophobic) group and the hydroxypropyl (hydrophilic) group (Perez, 62 Carrera Sanchez, Rodrigues Patino, & Pilosof, 2007). The functionality of EW in bulk aqueous 63 medium related to foaming properties could be improved by using HPMC and it depends on pH 64 65 (Berg, Jara & Pilosof, 2015; Sadahira et al, 2015).

The objective of this study was to evaluate the effect of total biopolymer concentration (g/100 g of sugar) and EW/HPMC ratio (g/g) in a high sugar content system on the foaming and rheological properties of the systems. The effect of pH (3.0, 4.5, and 6.0) on foaming properties was also evaluated.

70

71 2. Materials and methods

72 2.1. Materials

Sucrose (Tate & Lyle, London, UK) was purchased from local market. Glucose syrup (40 73 74 D.E.) and invert sugar syrup (80 g of sugar/100 g) were donated by Brenntag UK & Ireland (Leeds, UK) and by British Sugar (Peterborough, UK), respectively. Dried egg white protein (EW) and 75 hydroxypropylmethylcellulose (HPMC), METHOCEL F50 (methyl content 27.00 - 30.00 g/100 g, 76 hydroxypropyl content 4.00 - 7.75 g/100 g, 0.05 Pa.s viscosity in 2 g/100 g of solution, according to 77 78 manufacturer) were provided by Saltos Alimentos LTDA (Salto, Brazil) and Down S.A. (Midland, 79 USA), respectively. EW presented, in wet basis, 79.9 ±1.2 g of protein/100 g, 10.20 ± 0.02 g of 80 moisture/100 g and 5.64 ± 0.22 g of ash/100 g, determined according to methodologies described by AOAC (2010). SDS-PAGE analysis of EW (Laemmli, 1970) presented an eletrophoretic profile with 81 bands of 77.7, 44.5 and 14.3 kDa that correspond to conalbumin, ovalbumin and lysozyme, 82 respectively. Other reagents used were analytical grade and Milli-Q water was used in all 83 84 experiments.

85

86 2.2. Preparation of solutions and foams

The sugar mixture used as a model system to evaluate the foaming and rheological properties in aerated products was composed of sucrose (42.5 g of sugar/100 g), glucose syrup (42.5 g of sugar/100 g) and invert sugar (15 g of sugar/100 g). This composition is adequate to

obtain foams with density between 0.25 g/mL and 0.50 g/mL and water activity from 0.665 to 0.778,
which are characteristics of aerated confectionery such as marshmallow (Jackson, 1995; Wills,
1998).

In order to reach 80 g of sugar/100 g of solution, the sugar mixture was heated on hot plate stirrer and cooled to the whipping temperature, 70 °C. According to Table 1, the biopolymers were hydrated together in 36 g of water under magnetic stirring for 1 h at room temperature. The pH was adjusted to 3.0, 4.5 and 6,0 using 4 mol L⁻¹ citric acid.

97 For the foam preparation, the sugar mixture (500 g) and hydrated EW/HPMC blends were 98 mixed using a Kitchen Aid 5KPM5 stand mixer (Havant, UK) at speed setting 4, for 1 min and 99 equipped with a flat beater. The foams were then produced using a whisk beater, operating at speed 100 setting 10 under atmospheric pressure and whipping time of 6 min (Sadahira, Rodrigues, Akhtar, 101 Murray, & Netto, 2016).

102

103 2.3. Foaming properties

104 2.3.1. Foaming capacity: density and overrun

Foam samples were carefully filled into cylindrical containers $(35.43 \pm 0.21 \text{ mL})$ and to obtain constant volume the top of the container was leveled with a metal spatula to achieve a uniform and plane surface. The foam weight was recorded and then the foam density/overrun was determined according to Equation 1 (Lau & Dickinson, 2004).

109

110 Overrun (%) = $100(m_i - m_f)/m_f$

(1)

where m_i is the mass of the initial solution (before whipping) and m_f is the mass of the resulting foam with the same volume of m_i .

113 The density was determined by Equation 2:

114

115 Density (g/mL) = mf/volume of cylindrical container (2)

116

117 2.3.2. Liquid drainage

Foam samples were placed into plastic cylindrical containers (25 mL) and stored at 25 °C. Liquid drainage was followed for 20 or 30 days by visual observation and recorded via digital photography. 121

122 2.3.3. Bubble size distribution

Microscopy images of the foam samples were carried out using a Leica Confocal Scanning 123 Laser Microscope, CLSM, (model TCS SP2, Heidelberg, Germany) equipped with an Ar/HeNe laser 124 and 10x objective lens (HC PL APO CS 20 × 0.7 DRY). Rhodamine B (tetraethylrhodamine; with 125 purity of 95%, purchased from Aldrich (Dorset, UK), was used as the labeling dye at a level of 0.1 mL 126 of 0.1 (g/100 mL). The fluorescence dye Rhodamine B (Aldrich, UK), was excited at 50% of 127 maximum absorption at 488 nm, and the detection bandwidth was set from 500 to 600 nm. Images 128 were recorded at low magnification and analyzed via Image J software (Rasband, 1997-2016). A 129 130 fresh foam sample was placed into a welled slide (18 mm inner diameter x 3 mm depth) and the dye 131 was then added. The well was covered with a cover slide, pressed down to maintain a flat surface 132 over the well and the images were recorded after 24h.

Foam bubble size distributions were measured by analyzing the CLSM images via Image J software: 1000 bubbles were measured for each sample. According to Nicorescu et al. (2011) and Labbafi, Thakur, Vial & Djelveh (2007) sample size between 500 and 600 bubbles is sufficient for statistical analysis, bubble size distribution and Sauter Diameter (d₃₂).

137 Mean bubble size was characterized using Feret diameter. In order to calculate Sauter mean 138 diameter, a spreadsheet was built with number of bubble (frequency) within the range size bubble 139 (block). From the mid-point of each range/block we calculated the area and volume mean diameter 140 for each block. For each block the volume fraction (vol %) was calculated and then the bubble size 141 distribution was built.

142 d₃₂ was calculated using the Equation 3:

143
$$d_{32} = \frac{\sum_{n}^{1} \text{volume}}{\sum_{n}^{1} \text{surface}} = d_{32} = \sum_{i} di^{3} / \sum_{i} di^{2}$$
 (3)

144

145 2.4. Rheological properties

A stress-controlled rheometer (Kinexus, Malvern Instruments Limited, Worcestershire, UK) 146 147 equipped with parallel-plate geometry (65 mm flat plate) was used to measure the rheological properties at 25 °C. Apparent viscosity of sugar/EW/HPMC mixture before whipping was measured 148 as a function of shear rate (0.1 to 100 s⁻¹), using a 1 mm gap, according to previous studies with 149 glucose syrup and honey (Schellart, 2011). The increasing apparent viscosity of continuous phase 150 151 enhances the foam stability related to liquid drainage. In order to evaluate the viscosity of sugar syrup and drainage of liquid, the shear rate close to 10 s⁻¹ was used for CCRD because it is the 152 typical shear rate range for materials that presents drainage induced by gravity and during food 153

154 consumption (Barnes, Hutton, & Walters, 1989). The dynamic viscoelastic moduli (elastic modulus G', viscous modulus G") of the foams were determined at a maximum low shear strain amplitude of 155 0.02%. and a gap of 3 mm, which was selected to avoid crushing or destroying of the gas bubbles 156 157 (Zmudzinski et al., 2014). To determine the linear viscoelastic region in oscillatory shear, stress sweep tests were carried out at 1 Hz. Samples were also subjected to a frequency sweep from 0.1 158 to 10 Hz at constant strain amplitude (0.02%) within the linear viscoelastic region of each sample. 159 The rheological measurements were carried out in 3 repetitions for fresh foams and foams aged for 160 24 h. 161

- 162
- 163

2.4 Central Composite Rotatable Design (CCRD)

A Central Composite Rotatable Design CCRD (2² factorial design with 4 trials under the axial 164 conditions and 3 repetitions at the central point) totaling 11 trials (Table 1) (Rodrigues & lemma, 165 2015) was carried out. The effect of total biopolymer concentration (g/100 g of sugar) and EW/HPMC 166 ratio (g/g) on the apparent viscosity of the sugar/biopolymer mixture before whipping at 10 s^{-1} , 167 foaming capacity (density and overrun) of the fresh foam and the rheological properties (G', G" and 168 δ at 1 Hz) of the fresh foam and foam aged for 24 h were evaluated. High frequency corresponds to 169 170 short time while low frequency corresponds to long time ($\omega = 1/t$; ω : frequency, t: time). G' and G" 171 were used at 1 Hz for CCRD analysis in order to relate the elastic and viscous behavior, 172 respectively, to the texture of samples and their longer term stability. Data were analyzed via 173 Protimiza Experiment Design Software (http://experimental-design.protimiza.com.br). Second-order models were obtained and analyzed statistically by analysis of variance (ANOVA). 174

In order to evaluate the effect of pH on the foaming and rheological properties of the 175 sugar/EW/HPMC mixtures, experiments were carried out at pH 3.0, 4.5 and 6.0 under the conditions 176 used for the model validation (total biopolymer concentration 5.0 g/100 g of sugar, EW/HPMC ratio 177 g/g 14/1, 80 g of sugar/100 g of solution and 70 °C). The results were analyzed for differences 178 among means via Tukey's test (p < 0.05). 179

180

3. Results and discussion 181

182

3.1. Apparent viscosity, foaming and rheological properties of high sugar system/EW/HPMC mixtures

183 A CCRD was carried out with total biopolymer concentration and EW/HPMC ratio as independent variables to evaluate the effect of these variables on the apparent viscosity of 184 sugar/EW/HPMC mixture before whipping, the foaming capacity and rheological properties of 185 186 aerated samples. The experimental conditions as well as the results are shown in Table 1.

187 Mathematical models were built for the responses: apparent viscosity of sugar/ EW/HPMC 188 mixture before whipping at 10 s⁻¹, foaming capacity (density and overrun) and rheological properties 189 (G', G" and δ at 1 Hz) for fresh and aged for 24 h foams. On the basis of ANOVA, the adequacy of 190 the fitted model was evaluated (Table 2).

191

192 Table 1.

193

194 According to Table 2, R^2 and calculated F indicated that are adequate to obtain the second-195 order model (Equations y₁, y₂, y₃, y₄, y₆, y₇ and y₉) for the responses apparent viscosity, density, 196 overrun, G', and δ , within the range studied.

197

198 Table 2.

199

The equations from Table 2 were used to generate the contour curves for the dependent variables: apparent viscosity of sugar/EW/HPMC mixture before whipping (y_1) , foaming capacity (density (y_2) and overrun (y_3)) of fresh foam, rheological properties of fresh foam G' (y_4) , and δ (y_6) and foam aged for 24 h G' (y_7) and δ (y_9) (Fig. 1). According to Fig. 1, G' values of foams aged for 24 h are lower than G' values of fresh foams, indicating that the fresh foams were not completely stable. After 24 h, the microstructure changed, leading to a less elastic behaviour.

206 The apparent viscosity of the sugar/EW/HPMC mixtures before whipping increases with increasing total biopolymer concentration and decreasing EW/HPMC ratio (Fig. 1). In the regions of 207 low density and high overrun, G' values are higher and δ values are lower, for fresh foams and foams 208 209 aged for 24 h foams. The incorporation of air bubbles into liquids modifies food texture, which then 210 exhibit more semi-solid behaviour (Thakur, Vial & Djelveh, 2008). G' and G" represent the elastic and viscous behaviour of a material, respectively. When G' is higher than G", the material can be 211 said to be more solid-like, whereas when G" is higher than G', it can be said to be more liquid-like 212 213 (Rao, 1999). The loss factor is defined by tan δ (G"/G') or by the phase angle δ value. Tan $\delta = 0$ (phase angle $\delta = 0$) and tan $\delta = \infty$ ($\delta = 90^{\circ}$) characterize an ideal solid and viscous behaviour, 214 respectively (Steffe, 1996). Therefore increasing air incorporation improves the foam elastic and 215 solid behaviour, in accordance with previous work (Goff et al., 1995; Thakur, Vial, & Djelveh, 2008). 216

Foam presented low density, high overrun, high G' and low δ for fresh sample and sample aged for 24 h at total biopolymer concentration above 5 g/100 g of sugar and EW/HPMC ratio above 10/1.

220

221 Fig. 1.

222

The apparent viscosity of sugar/EW/HPMC mixtures before whipping were measured in order 223 to evaluate its influence on foaming capacity and foam rheological properties. Above 15 Pa.s., 224 225 increasing density and decreasing overrun values were observed, possibly due to the difficulty of 226 incorporating air bubbles. Low apparent viscosity of sugar/EW/HPMC mixtures led to greater liquid 227 drainage. Foams from trials 1, 3 and 5 (Table 1), which were prepared with mixtures with apparent 228 viscosity below 8 Pa.s, showed liquid drainage after one week of storage at 25 °C (Figure 2). Foams prepared from sugar/EW/HPMC mixtures with viscosity between 8 to 17 Pa.s did not present drained 229 liquid after 20 days at 25 °C (data not shown). On the other hand, mixtures with high apparent 230 viscosity such as the one from trial 7 (21.48 Pa.s) resulted in foam with high density value (0.72 231 232 g/mL) and low overrun value (46.9%). The high apparent viscosity possibly hampered the 233 incorporation of air bubbles during whipping and also influenced molecular diffusion - decreasing the 234 adsorption rate of proteins (Yang & Foegeding, 2010). Due to the lower foaming capacity, the foam presented low G' (363.1 Pa) and high δ (61.4°). These values indicate that this foam did not behave 235 236 as a solid, leading to creaming and liquid drainage after 20 days of storage at 25 °C (Figure 2).

237

238 Fig. 2.

239

The contour curves (Fig. 1) were jointly analyzed to determine the conditions to obtain high foaming capacity, elastic and solid behaviour, which characterize good foam properties. Thus total biopolymer concentration 5 g/100 g of sugar and EW/HPMC ratio 14/1 were the conditions to obtain low density, high overrun and G', small δ and intermediate apparent viscosity values (9 a 12 Pa.s). Foam obtained at these conditions showed density and δ of 0.35 g/mL and 20 °C, respectively, which are found in products such as marshmallows, chocolate mousse, whipped cream and dairy toppings (Jackson, 1995; Thakur et al., 2008).

247 Model validation was carried out under the previous established conditions (total biopolymer 248 concentration 5 g/100 g of sugar, 14/1 EW/HPMC (g/g ratio). The relative error between the experimental tests and predicted values by the coded model for apparent viscosity, density, overrun, G' (fresh foam), δ (fresh foam), G' (foam aged for 24 h) and δ (foam aged for 24 h) were -2.5, 3.1, 5.5, 9.8, 14.6, -1.2 and 16.5%, respectively. In general, the experimental results were close to the predicted values (Table 3). The exceptions were the experimental δ (fresh foam and foam aged for 24 h). In spite of this deviation, the results from validation experiments were satisfactory.

254

255 3.2. Effect of pH on foaming and rheological properties

Thermodynamic incompatibility of proteins and polysaccharides in solution (Grinberg & Tolstoguzov, 1997) and the effect of sucrose on the thermodynamic properties (protein hydrophilicity and surface activity) of proteins depend on the pH (Antipova et al., 1999). Thus, in order to study the influence of pH on foaming properties in a high sugar content system with EW and HPMC, experiments were carried out under the model validation conditions (total biopolymer concentration 5 g/100 g of sugar, 14/1 EW/HPMC ratio, 80 g of sugar/100 g of solution and 70 °C) at pH 3.0, 4.5 and 6.0. The results are presented in Table 3.

- 263
- 264 Table 3.
- 265

According to Table 3, the pH did not significantly affect the apparent viscosity of 266 sugar/EW/HPMC mixtures before whipping. However, the foams obtained at pH 3.0, 4.5 and 6.0 267 showed differences (p<0.05) in density, overrun and δ . The highest foaming capacity (density and 268 overrun) was obtained at pH 3.0. At this pH, the foam showed G' and δ values which characterized 269 elastic and solid behaviour for the fresh foam and foam aged for 24 h. At pH 3.0 and 4.5, G' of the 270 271 foams aged for 24 h did not differ (p>0.05) while at pH 4.5, G" values were higher than at pH 3.0 (p < 0.05). At pH 6.0 the lowest foaming capacity was obtained and G" value higher than G' for fresh 272 273 foam and foam aged for 24 h (Table 3), indicating viscous behaviour.

The highest foaming capacity being obtained at pH 3.0 is possibly due to the thermodynamic compatibility between EW and HPMC (Sadahira et. al., 2015). At pH 4.5 the foaming capacity is lower than at pH 3.0 possibly because pH 4.5 is close to protein pl (isoeletric point), which favours aggregation of ovalbumin. In addition, in the presence of sucrose, due to strengthening of the protein-protein net attractive interactions, significant aggregation of protein occurs leading to decrease of ovalbumin surface activity (Antipova et al., 1999).

At pH 6.0, the lowest foaming capacity and the highest foam instability (Fig. 3i) were possibly due to the interaction between ovalbumin and sucrose which leads to increase protein hydrophilicity in the bulk medium and decrease the protein surface activity (Antipova et al., 1999). Moreover,

thermodynamic incompatibility between biopolymers takes place at pH values higher than protein pl
(Grinberg & Tolstoguzov, 1997; Rodríguez Patino & Pilosof, 2011). Thermodynamic incompatibility at
the interface affects foam stability (Damodaran & Razumovsky, 2003).

The bubble size distribution of foams aged for 24 h obtained at pH 3.0, 4.5 and 6.0 are 286 287 presented in Fig. 3. At pH 3.0, foams had the smallest average bubble diameter (d₃₂) and a bimodal bubble size distribution (Fig. 3b). The splitting of the bubble size distribution suggests that the 288 smaller bubbles may be evolving into the larger ones due to gas diffusion from smaller bubble to 289 290 larger bubble (disproportionation). After 30 days of storage, foam at pH 3.0 did not present drainage (Fig 3c). At pH 4.5 d₃₂ was larger than at pH 3.0 and the bubble size distribution was monodisperse. 291 292 At this pH, the foam did not show drainage which led to greater stability related to disproportionation 293 and coalescence (Fig. 3f). Foam stability increases at pH values near the pI due to lower repulsion of 294 proteins that increase the interactions at interface air-water and a more stable and firm protein film is 295 created (Kuropatwa, Tolkach, A., & Kulozik, 2009). The foam prepared at pH 6.0 showed the largest bubble d_{32} (56.5 µm) and the widest bubble size distribution (Fig. 3h). These factors led to larger 296 297 foam instability mechanism such as creaming and liquid drainage after 30 days of storage at 25 °C 298 (Fig. 3i).

In order to analyze the degree of frequency dependence of the storage modulus (G') and 299 300 phase angle (δ), a power law model was fitted to the results from Fig. 4, i.e., G' = $a\omega^{n'}$ and $\delta = c\omega^{e'}$. 301 The fitted power law parameters are shown in Fig. 4. The coefficients a and c represent the 302 magnitude of the intercepts at frequency 1 Hz, whereas the n' and e' values represent the slopes of 303 G' and δ as a function of frequency (ω), respectively. According to Hatami et al. (2014) and Smith, 304 Goff, & Kakuda (2000) a and c are related to strength (elastic structure) and flexibility (rigid or 305 viscoelastic) of a sample. High frequency corresponds to short time while low frequency corresponds to long time ($\omega = 1/t$; ω : frequency, t: time) (Tadros, 2004). A n' value close to zero is characteristic of 306 307 a truly solid-like material, i.e., G' is independent of frequency and does not change with time. For n' value = 1 the system behaves as a viscous material (Hatami, Nejatian, Mohammadifar, & Pourmand, 308 2014). Thus, for 0 < n' < 1 the frequency dependence of G' is characteristic of a viscoelastic 309 310 structure (Smith, Goff, & Kakuda, 2000). The *n*' values and δ are lower at pH 3.0 than at pH 4.5, indicating that foam at pH 3.0 is more solid than the foam at pH 4.5. Moreover e' values were 311 constant (= 0.15) for foams at pH 3.0, whereas e' decreased from 0.12 to 0.05 for foams at pH 4.5 312 after 24 h. The lower e' values indicate that the stability of foam is related its viscoeslaticity, since δ 313 does not change over time. Foam with viscoelasticity characteristic is more able to resist the 314 destabilization processes (Smith, Goff, & Kakuda, 2000). Therefore, foams at pH 4.5 were more 315

stable than foams at pH 3.0. At pH 6.0, foams showed viscous behaviour, i.e., G' > G' value and $\delta > 0.45$, leading to the highest instability.

318

319 Fig. 3.

320 Fig. 4.

321

322 4. Conclusions

323 Total biopolymer concentration (egg white protein - EW and hydroxypropylmethylcellulose -HPMC), EW/HPMC ratio and pH influenced on foaming and rheological properties of aerated high 324 sugar system. At pH 3.0, systems had the highest foaming capacity, elastic and solid-like behaviour, 325 with little drainage, whereas systems prepared at pH 4.5 showed lower foaming capacity, but with 326 better stability to disproportionation and coalescence than foams prepared at pH 3.0 because of the 327 328 viscoelastic behaviour of the foams at pH 4.5. At pH 6.0, foams showed the lowest foaming capacity, 329 the highest instability and more liquid-like behaviour. The evaluation of the frequency degree 330 dependence of the storage modulus (G') and phase angle (δ) indicates the foam rheological behaviour (solid-like, viscoelastic and liquid-like) in order to evaluate the foam stability. HPMC may 331 be considered to increase the stability of aerated confectionery at pH 4.5 but not at pH 6.0. 332

333

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Table 1. Design matrix of the Central Composite Rotatable Design (CCRD) with independent variables total biopolymer concentration (g/100 g of sugar) and egg white protein (EW)/hydropropylmethylcellulose (HPMC) ratio (g/g), and the results for responses: apparent viscosity of sugar/ egg white protein (EW)/hydropropylmethylcellulose (HPMC) mixture before whipping at 10 s⁻¹, foaming capacity (density and overrun) and rheological properties (elastic modulus G', viscous modulus G' and phase angle δ at 1 Hz) for fresh foam and foam aged for 24 h, at pH 3.0.

Trial	Total Biopol.	EW/HPMC	η	ρ	Overrun		Fresh foa	m	Foam aged for 24 h		
	conc. (g/100	ratio	(Pa.s)	(g/mL)	(%)	G' (Pa)	G" (Pa)		G' (Pa)	Ğ" (Pa)	δ (°)
	g of sugar) x ₁	(g/g) x ₂	y ₁	y ₂	y 3	У4	y 5	y ₆	У7	У 8	y 9
1	-1 (2.00)	-1 (4:1)	7	0.56	102.2	1202	1317	48	523	811	57
2	1 (5.00)	-1 (4:1)	17	0.48	140.5	2485	1808	36	1024	834	39
3	-1 (2.00)	1 (16:1)	3	0.42	180.0	2411	1216	28	928	773	40
4	1 (5,00)	1 (16:1)	11	0.39	212.8	4700	1909	22	1434	879	31
5	-1.41 (1.40)	0 (10:1)	3	0.45	169.4	1893	1036	30	562	693	51
6	1.41 (5.60)	0 (10:1)	10	0.39	198.0	4697	1788	21	1232	724	30
7	0 (3.50)	-1.41 (2:1)	21	0.72	46.9	363	664	61	269	511	62
8	0 (3.50)	1.41 (18:1)	8	0.40	198.9	3938	1603	22	1163	767	33
9	0 (3.50)	0 (10:1)	10	0.41	187.6	4079	1738	25	1145	851	37
10	0 (3.50)	0 (10:1)	8	0.42	181.4	2769	1214	25	908	596	33
11	0 (3.50)	0 (10:1)	10	0.41	190.5	3962	1644	23	989	648	33

Coded values and () true values of the independent variables; Total biopolymer conc.: total biopolymer concentration; η: apparent viscosity; ρ: density; G': elastic modulus; G'': viscous modulus; δ: phase angle.

Table 2. Analysis of variance (ANOVA) (Percentage of explained variance (R^2), $F_{calculated}$ value and $F_{tabulated}$) for the responses: apparent viscosity of sugar/egg white protein (EW)/hydropropylmethylcellulose (HPMC) mixture before whipping at 10 s⁻¹, foaming capacity (density and overrun) and rheological properties (elastic modulus G', viscous modulus G' and phase angle δ at 1 Hz) for fresh and aged for 24 h foams.

Response		R² (%)	Fcalculated	F* _{tabulated}	Equation	
Apparent Viscosity (η) Pa.s		9 <mark>2</mark> .1	17.4	4.53	$y_1 = 9.33 + 3.5x_1 \cdot 1.7x_1^2 - 3.5x_2 + 2.3x_2^2$	
Density (ρ) (g/mL)		91. <mark>4</mark>	25.0	4.35	$y_2 = 0.41 - 0.02x_1 - 0.09x_2 + 0.07x_2^2$	
Overr (%)	un	97.0	73.8	4.35	$y_3 = 186.5 + 13.9x_1 + 45.7x_2 - 30.4x_2^2$	
	G' (Pa)	91. <mark>3</mark>	24. <mark>6</mark>	4.35	$y_4 = 3452.8 + 942.2x_1 + 1060.0x_2 - 685.2x_2^2$	
Fresh foam	G" (Pa)	63.9	1.8	5.12	It was not possible to establish a model	
Ľ	δ (°)	95. <mark>5</mark>	49.5	4.35	$y_6 = 24.9 - 3.8x_1 - 11.1x_2 + 8.4x_2^2$	
24h	G' (Pa)	90. <mark>4</mark>	22,1	4.35	$y_7 = 999.1 + 244.3x_1 + 259.9x_2 - 101.7x_2^2$	
Foam aged for 24h	G" (Pa)	25.0			No regression coefficient was statistically significant (p > 0.10)	
Foam	δ (°)	91.1	23.8	4.35	$y_9 = 36.7 - 7.1x_1 - 8.2x_2 + 5.3x_2^2$	

x, x: coded independent variables for total biopolymer concentration and EW/HPMC ratio, respectively. ---: there is no regression coefficient.

Table 3. Results of experimental validation conditions of sugar/egg white protein (EW)/hydropropylmethylcellulose (HPMC) mixture (biopolymer concentration 5 g/100 g of sugar, 14/1 egg white protein (EW)/hydropropylmethylcellulose (HPMC) ratio, 80 g of sugar/100 g of solution), for responses apparent viscosity (before whipping at 10 s⁻¹), foam density (ρ), overrun, rheological properties of fresh foam and foam aged for 24 h (elastic modulus G', viscous modulus G" and phase angle δ at 1 Hz) obtained at pH 3.0, 4.5 and 6.0.

		рН						
		3.0 (CCR	D pH)	4.5	6.0			
		Experimental	Predicted					
η	(Pa.s)	10 <u>+</u> 0.9 ^a	10	11 <u>+</u> 0.6 ^a	9 <u>+</u> 2 ^a			
Density	(g/mL)	0.38 <u>+</u> 0.00 ^a	0.37	0.42 <u>+</u> 0.01 ^b	0.51 <u>+</u> 0.02 ^c			
Overrun	(%)	206 <u>+</u> 11 ^a	218	168 <u>+</u> 4 ^ь	140 <u>+</u> 15 [°]			
	G' (Pa)	5326 <u>+</u> 227 ^a	4803	3538 <u>+</u> 721 ^ь	903 <u>+</u> 226°			
Fresh foam	G" (Pa)	1924 <u>+</u> 57 ^a	÷	1837 <u>+</u> 266 ^a	1380 <u>+</u> 114 ^ь			
	δ (°)	20 <u>+</u> 0.5 ^a	17	28 <u>+</u> 0.9 ^b	61 <u>+</u> 2 ^c			
	G' (Pa)	1360 <u>+</u> 115 ^a	1376	1234 <u>+</u> 76 ^a	466 <u>+</u> 104 ^b			
Foam aged for 24h	G" (Pa)	850 <u>+</u> 36 ^a		1230 <u>+</u> 32 ^b	525 <u>+</u> 72 [°]			
	δ (°)	32 <u>+</u> 2 ^a	27	45 <u>+</u> 2 ^b	54 <u>+</u> 2 [°]			

Values are mean \pm SD of triplicates, except G' and δ fresh sample that are mean \pm SD of duplicates. For the same response, mean with different small letters in the same row differ significantly (p <0.05) by Tukey's test; density (p), *overrun*, rheological properties of fresh sample

and sample aged for 24 hours (elastic modulus G', viscous modulus G" and $\delta)$. ---: there is no predicted value.

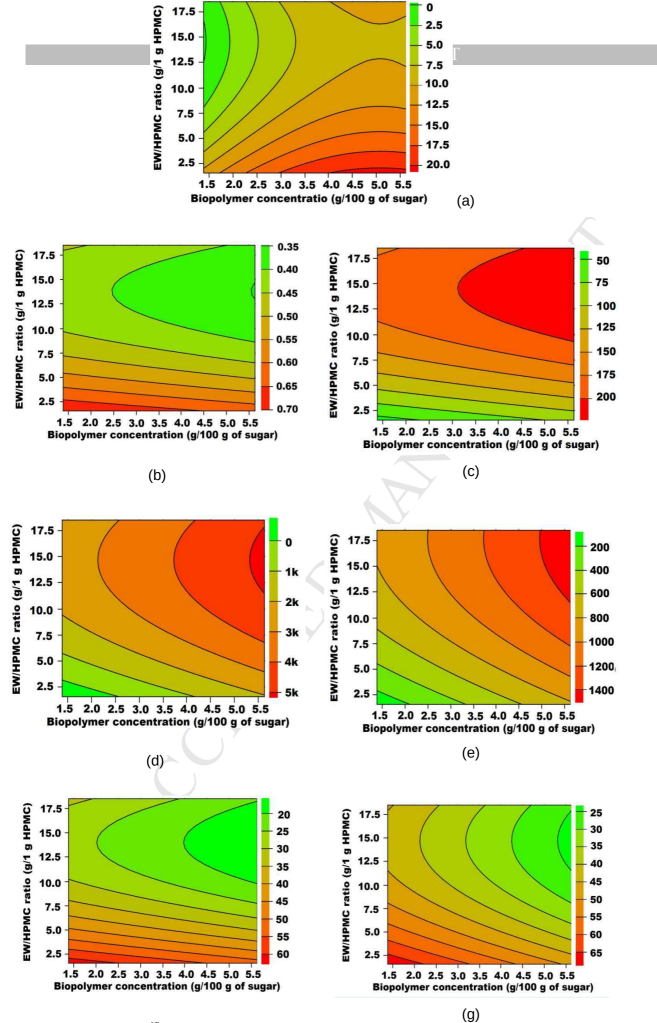


Fig. 1. Contour curves from de Central Composite Rotatable Design CCRD for the dependent variables apparent viscosity (η) of sugar/egg white protein-EW/hydropropylmethylcellulose-HPMC mixtures: (y_1) before whipping (a), foaming capacity of fresh foam density (y_2) (b) and *overrun* (y_3) (c), rheological properties of fresh foam G' (y_4) (d) and δ (y_6) (f) and foam aged for 24 h G' (y_7) (e) and δ (y_9) (g).



Fig. 2. Liquid drainage of foams obtained from de Central Composite Rotatable Design CCRD under the conditions of Trial 1, 3 and 5 (pH 3.0; 70 °C) after 1 week of storage at 25 °C; drainage and creaming of Trial 7 after 20 days of storage at 25° C. η : apparent viscosity of sugar/egg white protein (EW)/hydropropylmethylcellulose (HPMC) mixture before whipping; ρ : foam density. Bio. conc.: biopolymer concentration.

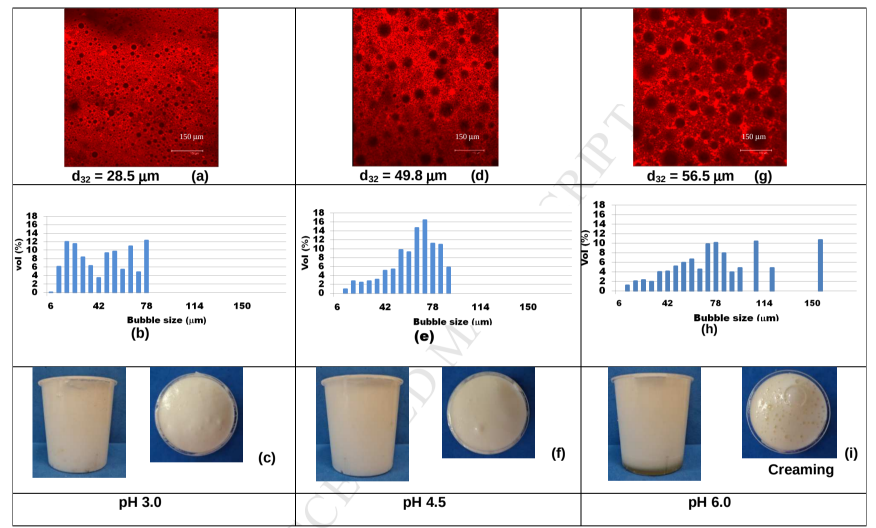


Fig. 3. Confocal microscopy (after 24h of storage at 25 °C) (a, d, g), bubble size distribution (b, e, h) and photographs (after 30 days of storage at 25 °C) (c, f, i) of aerated samples containing 5 g biopolymer/100 g of sugar and egg white protein (EW)/hydropropylmethylcellulose (HPMC) ratio 14/1 (g/g) at pH 3.0, pH 4.5 and pH 6.0. Average bubble diameter: d₃₂

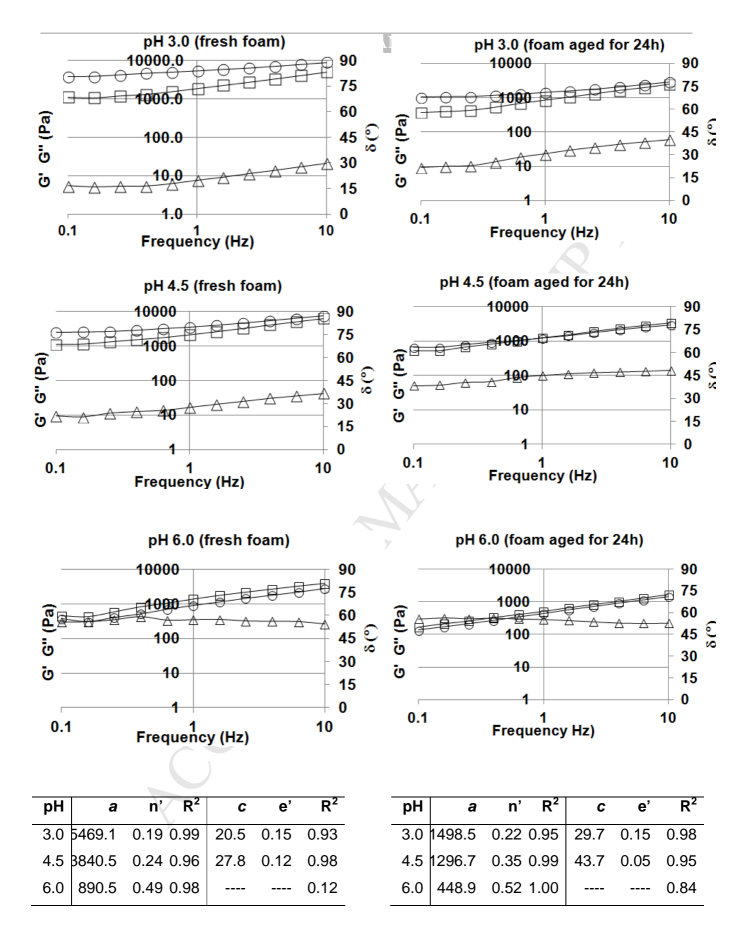


Fig.4. Dynamic frequency sweep of aerated samples containing 5 g biopolymer/100 g of sugar and egg white protein (EW)/hydropropylmethylcellulose (HPMC) ratio 14/1 (g/g) at pH 3.0, pH 4.5 and pH 6.0. Power law parameters for storage modulus G' (G' = a $\omega^{n'}$) and phase angle δ (δ = c $\omega^{e'}$) where The

coefficients *a* and *c* represent the magnitude of the intercepts at frequency 1 Hz and the n' value and e' value represent the slope of G' and δ in function of frequency (ω), respectively. ----: there is no R², explained percentage of variation. G' (O); G'' (\Box); δ (Δ)

Highlights

High sugar foam stability depends on pH and rheological properties of the mixtures.

High sugar foam with viscoelasticity characteristic shows higher stability.

Hydroxypropylmethylcellulose increases high sugar foam stability at pH 4.5.

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