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Additional Information

1	USE OF DIFFERENT BIOPOLYMERS AS CARRIERS FOR PURPOSES OF OBTAINING A
2	FREEZE-DRIED ORANGE SNACK
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8	
9	Abstract
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10	In addition to colour, one of the most important qualities of a snack-type product is its crunchy
11	texture. A freeze-dried fruit snack is characterised by its low water content, which creates the
12	problem of a loss of crunchiness related to its low glass transition temperature (Tg). In this sense, a
13	common technique with which to increase the Tg of these types of products is to add different
14	biopolymers. However, these compounds can, at the same time, affect the colour and texture of the
15	product. In this study, different biopolymers have been tested in order to discover their similarities or
16	differences in terms of hygroscopicity, antiplasticising character, colour and impact on the
17	mechanical properties of a freeze-dried orange snack formulated from their different mixtures. Gum
18	Arabic, maltodextrin, starch modified with octenylsuccinic anhydride, pea fibre, bamboo fibre and
19	native corn starch have been selected as biopolymers. The impact of any of them on the studied
20	properties can be confirmed, without any of them being more or less effective than the others.
21	
22	Keywords: gum Arabic, maltodextrin, modified and native starch, pea and bamboo fibre,
22	crunchiness
<i>,</i> ¬	CHRISTORS

#### 1. Introduction

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The consumption of fruit is part of a healthy diet and it is recommended as a way of improving our health and well-being (WHO, 2018). With the intention of promoting the consumption of fruit among the population, there is continuous interest in offering consumers new products that stimulate this consumption (Telis & Martínez-Navarrete, 2012). To this end, there exists the possibility of obtaining a snack from freeze-dried orange puree.

The final water content of the freeze-dried fruit will condition its characteristics, especially its optical and textural properties, and the expected shelf life under given conditions (Telis & Martínez-Navarrete, 2012). During the processing of typical snacks, a low water content is determinant for their characteristic brittleness, crispness or crunchiness. The plasticising effect of water on the mechanical properties of the food systems is well known (Roos, 1995). The primary mechanical effect of a plasticizer is that of weakening or breaking intermolecular bonds, thus decreasing its mechanical resistance and increasing its deformability, gumminess or sogginess (Pittia & Sacchetti, 2008). The impact of water content change on the mechanical behaviour of a food has been related to its physical state (Roos, 1995). Dehydrated fruits obtained by freeze-drying are normally in an amorphous state. Above the glass transition temperature (Tg), the change from the glassy state, more stable, to the rubbery occurs. The much lower viscosity of the rubbery state determines the aforementioned product softening, so that the typical crunchiness is lost. For this reason, it is of great importance to ensure the glassy state of a crispy product. To this end, it should be processed and stored at temperatures below its Tg. Likewise, it is important to know, at a specified storage temperature, the critical water content (CWC) and the critical water activity (CWA) of the product in order to ensure the glassy state. As in the low water content range, a small increase in the food water content leads to a significant decrease in the corresponding Tg, the storage relative humidity (RH) has to be strictly controlled. The state diagram of a food is one tool with which to predict and control the phase transitions. According to different authors, it is the right tool for the purposes of finding out the relationships between Tg and the water content in order to design efficient processes to obtain high-quality products and also to optimize storage conditions (Fabra, Talens, Moraga, & Martínez-Navarrete, 2009; Rahman, 2006; Roos & Karel, 1991; Roos, 1995).

Freeze-dried fruit pulps, such as sugar-rich foods, present structural problems of stickiness, caking and collapse. The Tg of freeze-dried fruit products present very low values, between 25 °C and -38 °C for a water content of between 3.3 and 25% (Telis & Martínez-Navarrete, 2010). Therefore, it is fairly easy to find these products in a rubbery state under the usual storage conditions. One way to prevent this, is, for example, the addition of biopolymers of high molecular weight that contribute to increasing the value of Tg or that even play a steric role delaying the structural collapse. Biopolymers, such as gums (gum Arabic, xanthan), maltodextrins, proteins (whey protein concentrate), starches (octenyl succinic anhydride, waxy starch) and natural fibres (bamboo fibre), have been used as drying carriers to obtain stable dehydrated products (Agudelo, Igual, Camacho, & Martínez-Navarrete, 2017; Bhusari, Muzaffar, & Kumar, 2014; Cano-Chauca, Stringheta, Ramos, & Cal-Vidal, 2005; Da Silva et al., 2013; Fongin, Kawai, Harnkarnsujarit, & Hagura, 2017; González, García-Martínez, Camacho, & Martínez-Navarrete, 2019; Martínez-Navarrete, Salvador, Oliva, & Camacho, 2019; Telis & Martínez-Navarrete, 2009). However, the addition of biopolymers may cause unintended effects in other properties, such as changes in the colour or texture of the final product. Non-enzymatic browning reactions, such as Maillard reactions and caramelization, may stem from heating or occur during the long-term storage of foods containing carbohydrates, especially reducing sugars (BeMilller & Whistler, 1996; Telis & Martínez-Navarrete, 2012). Since the biopolymers differ as to their composition and nature, their interactions with the food matrix may vary, causing different water-solid interactions. According to Acevedo, Schebor, & Buera (2008), there is a correlation of the effects of water-solid interactions and water mobility on the nonenzymatic browning rates in freeze-dried potato. Some authors have studied the effect of different storage vapour pressure atmospheres on the stability of fruit powders containing biopolymers. They have found that the mechanical properties of the dried food are closely dependent on the aw (Pérez-Alonso, Beristain, Lobato-Calleros, Rodríguez-Huezo, & Vernon-Carter, 2006; Telis & Martínez-Navarrete, 2009).

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In this study, different biopolymers, gum Arabic, maltodextrin, starch modified with octenylsuccinic anhydride, pea fibre, bamboo fibre and native corn starch, have been tested in order to evaluate the

- 80 possible advantages or disadvantages of each one of them when added to an orange puree for the
- 81 purposes of freeze-drying and obtaining a snack.
- 82 2. Materials and Methods
- 83 2.1. Raw materials
- 84 2.1.1. Fruit
- The oranges (*Citrus x sinensis* var. Navel) used in this study were bought from a local supermarket
- in the city of Valencia (Spain). The fruit pieces were chosen by visual inspection based on the size,
- 87 homogeneity, colour and good physical integrity.
- 88 2.1.2. Biopolymers
- 89 The carriers used to obtain the dehydrated orange samples were gum Arabic (GA, Scharlab,
- 90 Sentmenat, Spain), maltodextrin (MD, Roquette, France), starch modified with octenylsuccinic
- 91 anhydride (OSA, Roquette, France), pea fibre (PF, Roquette, France), native corn starch (NC,
- 92 Roquette, France) and bamboo fibre (BF, VITACEL®, Germany). These biopolymers were selected
- 93 to avoid structural collapse of the dehydrated product, GA, MD and OSA for its ability to increase Tg
- and BF, NCS and PF because of its steric role avoiding the formation of interparticle bridges.
- 95 2.2. Samples preparation
- 96 In order to obtain the orange puree, the fruit was washed, peeled, cut and triturated in a bench top
- 97 electrical food processor for 40 s at speed 4 (2000 rpm) followed by 40 s at speed 9 (91000 rpm)
- 98 (Thermomix TM 21, Vorwerk, Spain). The orange puree was mixed (10 min at speed 3 (1000 rpm)
- 99 with the biopolymers (Table 1) to obtain five different formulated samples to be freeze-dried, in
- 100 addition to the orange puree with no biopolymers (O) that was also considered. The ratio orange
- puree:biopolymers was selected to ensure the physical stability of the dried product (Agudelo, Igual,
- 102 Camacho, & Martínez-Navarrete, 2017).
- For freeze-drying purposes, each of the samples was distributed on two aluminium plates, 10.5 x
- 104 7.8 cm, 0.5 cm thickness, and immediately frozen at -45 °C (Liebherr LGT 2325, Germany) and
- then dried (Telstar Lioalfa-6, Spain), at 0.05 mbar, -45 °C on the condenser and 40 °C on the

- 106 shelves for 20 hours, to obtain two cakes from each of the six different freeze-dried cakes (O,
- 107 GA+BF, MD+PF, MD+NCS, OSA+PF, OSA+NCS). One cake from each sample was crushed
- manually with a mortar to obtain the corresponding powder.
- 109 2.3. Sorption experiments
- 110 The freeze-dried cake and the powder from each of the six samples were placed at 20 °C in
- 111 hermetic chambers containing saturated salt solutions (BrLi, ClLi, CH<sub>3</sub>COOK, MgCl<sub>2</sub>, K<sub>2</sub>CO<sub>3</sub>,
- 112 Mg(NO<sub>3</sub>)<sub>2</sub>, the corresponding relative humidity (RH) ranging between 6% and 53% (Greenspan,
- 113 1977). The samples were weighed every week in order to determine the equilibrium condition with
- the surroundings (Δm<±0.001 g) (Kaymak-Ertekin & Gedik, 2004). In this moment, the water activity
- of each sample was assumed to be equal to the corresponding RH/100. The time needed to reach
- equilibrium was approximately two months.
- 117 2.4. Analytical determination
- 118 2.4.1. Water content (x<sub>w</sub>)
- 119 The mass fraction of water (x<sub>w</sub>) was obtained by drying the equilibrated powdered samples in a
- vacuum oven (Selecta®, Vaciotem-T, J.P. Selecta S.A., Spain) at 60°C ± 1°C under p < 100 mm Hg
- 121 until constant weight (AOAC, 1990). Three replicates were carried out on each of the six
- equilibrated freeze-dried samples and the mean value was considered as the corresponding x<sub>w</sub>.
- 123 2.4.2. Glass transition temperature (Tg)
- 124 The Tg of the powdered samples conditioned at different water contents was determined by
- 125 differential scanning calorimetry (DSC 220CU-SSC5200, Seiko instruments Inc., Japan).
- 126 Approximately 15 mg of each sample were placed into DSC pans (P/N SSC000C008, Seiko
- 127 Instruments Inc., Japan). The heating rate was 5 °C/min and the temperature range varied between
- 128 -80 °C and 80 °C, depending on the water activity of each sample. The onset, midpoint and
- endpoint of the glass transition were obtained from each thermogram.
- 130 2.4.3. Fitted models

131 In order to predict the water sorption behaviour of the samples, the linearized BET model (Eq. 1) 132 was used to predict water sorption up to a<sub>w</sub> ≈ 0.5 (Brunauer, Emmett & Teller, 1938).

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$$\frac{a_w}{(1-a_w)\,W_e} = \frac{1}{W_0\,C} + \frac{C-1}{W_0\,C}.\,a_w \tag{1}$$

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Where W<sub>e</sub> is the equilibrium water content (g water/g dry solute), W<sub>o</sub> is the monolayer water content value (g water/g dry solute), C is the energy constant related to the sorption heat and aw is the water activity.

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140 In order to predict glass transition temperatures, experimental Tg -  $x_{\text{w}}$  data were fitted to the 141 linearized Gordon & Taylor (1952) equation (Eq. 2) considering the onset, midpoint and end of the 142 transition.

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$$T_g = T_{g(s)} + k \frac{x_w \cdot (T_{g(w)} - T_g)}{(1 - x_w)}$$
 (2)

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Where Tg is the glass transition temperature ( ${}^{\circ}$ C), Tg<sub>(s)</sub> is the glass transition temperature of the anhydrous solids (°C), k is the Gordon y Taylor constant model, xw is the mass fraction of water (g water/g product) and  $Tg_{(w)}$  is the glass transition temperature of amorphous water: -135 °C (Roos, 1995).

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The relationship between Tg - a<sub>w</sub> is given by the linear regression proposed by Roos (1995) (Eq. 3), 152 where Y and Z are the constants of the model.

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$$T_g = Ya_w + Z \tag{3}$$

155 2.4.4. Mechanical properties

> The mechanical behaviour of the equilibrated samples was registered using a texture analyser TA-XT2i (Stable Micro Systems, UK). Portions of 20 x 20 mm of the freeze-dried cakes were

compressed using a cylindrical probe of 10 mm diameter applying a strain of 80% with a test speed of 1 mms<sup>-1</sup>. Five replicates were performed per sample. The parameters analysed in the test were the maximum force (F), expressed in Newtons.

#### 2.4.5. Colour measurements

CIE L\*a\*b\* colour space was selected as a uniform and objective method with which to specify the colour of equilibrated freeze-dried cake samples. The L\* coordinate denotes lightness on a 0–100 scale from black to white; a\*, (+) red or (-) green; b\*, (+) yellow or (-) blue. Colour coordinates (10° observer and D65 illuminant) were obtained from the reflectance spectrum using a spectrophotometer (Minolta, CM 3600D, Japan). The colour was measured at four different points of the whole cake and the mean value was considered. From the colour coordinates, the hue angle (h\*, Eq. 4), and chroma or saturation (C\*, Eq. 5) were obtained. Measurements were taken with the specular component excluded.

$$h^* = \operatorname{arctg}(b^*/a^*) \tag{4}$$

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$$C^* = (a^{*2} + b^{*2})^{0.5}$$
 (5)

## 3. Results and Discussion

# 3.1. Sorption behaviour

The obtained sorption isotherms predict the relationship between w<sub>e</sub> and a<sub>w</sub> of the different freezedried orange purees at 20 °C (Fig. 1). As can be observed, all the added biopolymers reduced the hygroscopicity of the orange cake over the whole aw range of, OSA seeming to be the most effective for this purpose. The experimental data of each sample were fitted to the BET model (Table 2). The BET monolayer water content (W<sub>o</sub>) was in the range of 0.0619 - 0.0742 g water/g dry solid, the highest value being that of the orange sample without biopolymers and the lowest that of the orange with OSA ones. The W<sub>o</sub> indicates the amount of water that is tightly adsorbed in specific sites on food surfaces,

which has been related with a security water content below which the product stability is guaranteed (Choudhury, Sahu, & Sharma, 2011; Telis & Martínez-Navarrete, 2010; Wan et al., 2018). The values of constant C, the other BET parameter, varied between 2.25 – 2.96, which allows these sorption isotherms to be classified as type II (Brunauer, Deming, Deming & Teller, 1940). In order to evaluate the significant differences in the sample's sorption behaviour, the BET equation fitted to each sample and that fitted to different groups of samples were statistically compared through the values of the statistic E, which was compared with tabulated F – Snedecor (Moraga et al., 2004). The obtained results permitted a twofold confirmation: that there were no differences between the formulated samples (P>0.05) and that all of them were different to the O sample (P<0.05). In this way, the capacity of any of the biopolymers used to reduce the hygroscopicity of the freeze-dried orange snack can be confirmed, without any of them being more or less effective than the others.

### 3.2. Glass transition temperature- water content-water activity relationships

The glass transition temperature of dried foods is extremely important as a means of predicting the conditions of a proper drying process and product storage (Roos & Karel, 1991). Nevertheless, glass transition is a state transition that is developed over a temperature range. That is why the onset (Tg°), mid-point (Tg<sup>m</sup>) and end-point (Tg°) of glass transition can be characterized (Fig. 2). The majority of studies published take Tg<sup>m</sup> as the characteristic Tg value (Goula, Karapantsios, Achilias, & Adamopoulos, 2008; Khallooufi & Ratti, 2003; Roos, 1995; Wu, Sun, & Liu, 2019). However, the change in the food properties associated with glass transition, could start from the moment in which the transition begins or might not become patent until the entire amorphous matrix has changed to the rubbery state, at the end of the Tg. For this reason, it may be interesting to consider Tg° and Tg° as being related to the physical properties (Rahman, 2006; Wan et al., 2018). The samples considered in this study showed a Tg amplitude of about 5-15 °C and, as expected, the Tg fell as the water content increased (Fig. 2). The orange sample without biopolymers presented lower Tg values than the samples with added biopolymers, especially in the lower water content range (approximately up to 0.07g water/g product). From this value, the added biopolymers do not seem to have so much influence on the Tg. The GA+BF sample seems to be the one that

most increases the Tg over the entire water content range. Each one of the 3 characteristic values of the Tg of each sample was related to the corresponding water content by fitting the Gordon and Taylor model (Eq. 2). Table 3 shows the corresponding k and Tg(s) parameters. The value of these parameters confirms the above comments. In this sense, the lowest  $Tg_{(s)}$  was that of the O sample and the highest that of the GA+BF sample, the amplitude of the corresponding Tg ranging between 7.6 and 14.7 °C, respectively. Again, the statistical comparison among the different Gordon and Taylor fittings, performed through comparing the values of statistic E with the tabulated F-Snedecor, showed that only the O sample was different as regards the water plasticization behaviour. No significant differences (P>0.05) were detected among the formulated samples. As shown in Fig. 1, Tg-a<sub>w</sub>-w<sub>e</sub> relationships may be plotted together to construct the stability map, which is a tool that permits an easy identification of the critical processing or storage conditions (temperature and relative humidity) for glass transition (Fabra et al., 2009). Table 4 shows the critical water content (CWC) and critical water activity (CWA) values that have to be exceeded for the glass transition to take place, both at 20 and 4 °C, common room and refrigeration temperatures. The freeze-dried orange cake with added biopolymers showed higher values of CWC and CWA than the O sample, related to the greater capacity of these samples to maintain the more stable glassy state. It is remarkable that, despite the great microbiological and chemical stability of the dehydrated products, decreasing the storage temperature appreciably increases the CWC and CWA of all the samples. In this regard, at 20 °C for instance, the glass transition of the GA+BF sample will take place, at a surrounding RH in the range of 19.0-30.2%, this increasing to 31.1-42.5% at 4 °C. This increase in the critical values may be related to the physical stability of the products in terms of aspects such as the texture or colour of the snack. On the other hand, in order to obtain a tool that makes it easy to relate both temperature and RH storage conditions for glass transition, Tg and a<sub>w</sub> were related using the linear relationship (Eq. 3) proposed by Roos (1995). Table 2 shows the results of this fit, where a very good and useful linear relationship can be observed.

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## 3.3. Colour and mechanical properties

The addition of biopolymers led to an increase in L\* and a decrease in a\* and b\*, so that the chroma of the samples decreased and the hue angle increased (Fig. 3). This colour change is related to the colour of the biopolymers themselves and to the dilution of the orange pigments in the formulated samples (Telis & Martínez-Navarrete, 2009). The whitish colour of the biopolymers contributes to the increase in lightness, the colour becoming yellower and less pure. The most significant colour change in every sample was observed at water activities between 0.328 and 0.432. At this a<sub>w</sub>, the amorphous matrices of all the samples are already in a totally rubbery state (Table 4), and the water content is enough for enzymatic and non-enzymatic oxidation and browning reactions to occur. From this water activity, the greater amount of water present in the samples exerts a dilution effect of the components responsible for the colour, which means that the change in colour is no longer so marked.

The variation in the mechanical properties of the studied snacks was evaluated from the force—deformation curves. Fig. 4 shows the typical shape of the force—distance curves obtained from the different studied samples with several  $a_w$ . For the purposes of clarity, only one of the replicates at each  $a_w$  was selected for the Figure. In every case, the increase in the water content dramatically affected the shape of the curve, so that a change from a jagged to a smooth and regular trend of the force—deformation relationship was observed. This change is related to the material transformation from hard and brittle (or crunchy and crispy) when dried to soft and ductile at a higher water content (Martínez-Navarrete et al., 2019). The loss of crispness in the orange sample without added biopolymers was evident at water activities between 0.113 and 0.225, while in the samples with added biopolymers it was in the range of 0.225-0.328. In every case, the number of fracture peaks observed at the lower  $a_w$  decreased when the water content rose. These results point to the fact that any of the studied biopolymers may be used to protect the mechanical properties of the snack over a wider water content range, although there is still a limit to the gain of water if the quality of the product is to be ensured.

As a measurement of the strength of the samples with different  $a_w$  and so different water contents, the maximum force achieved in the mechanical test (Fmax) was characterized from the force—distance curves (Fig. 5). Significant differences (p<0.05) were observed between samples and  $a_w$ .

In order to confirm the impact of the physical state of the snack matrix on the measured mechanical response, the CWA for the glass transition of every sample (Table 4) was related with Fmax. Fig. 5 shows CWA for the onset and end point of the Tg (CWA $^{\circ}$  and CWA $^{\circ}$ , respectively) of the O and GA+BF samples, and also those of the OSA+PF sample, whose values were genuinely close to the rest of the samples. As may be observed, the values of Fmax remain low below CWA $^{\circ}$ , while all the product matrix is in the glassy state. Fmax sharply increases while the transition from the glassy to the rubbery state occurs (between CWA $^{\circ}$  and CWA $^{\circ}$ ) and sharply decreases again when the rubbery state is fully achieved, above CWA $^{\circ}$ . The small differences observed in the Tg-x<sub>w</sub>-a<sub>w</sub> relationships of the different samples containing biopolymers, despite not being significant (P>0.05), were enough to ensure a significantly (P<0.05) higher Fmax value of the GA+BF sample at a<sub>w</sub> = 0.328 compared to the other samples, MD+PF having intermediate values.

The decrease in Fmax is related with the plasticizing effect of water in the fully rubbery state. In this state, the amount of water in the sample is enough to ensure all the primary interactions with the matrix and there is still water available to contribute to the softening of the sample by weakening intermolecular bonds. The increase in Fmax at the lowest  $a_w$ , also observed in other studies (Chang, Cheah, & Seow, 2000; Moraga, Talens, Moraga, & Martínez-Navarrete, 2011; Sogabe, Kawai, Kobayashi, Jothi, & Hagura, 2018), has been called the anti-plasticization effect of water and it has been related with an increase in the cohesiveness of the glassy food matrix that increases its rigidity and firmness (Pittia & Sacchetti, 2008). This offers a greater resistance upon force application and more energy is needed to compress the sample. For these authors, the anti-plasticization effect could be considered as a mere hardening or 'toughening' effect. This hardening limit the number of fractures when  $a_w$  increases over this low  $a_w$  range. In fact, it seems that the loss in the number of observed multiple fracture peaks occurs at the same  $a_w$  from which Fmax starts to decrease (Figs 4-c and 5), when the whole matrix is in the rubbery state.

The above results indicate the need to keep the orange snack in a completely glassy state to ensure a crispy texture and adequate colour. Despite the fact that the W<sub>o</sub> value has frequently been considered as a secure water content below which product stability is guaranteed (Choudhury et al., 2011; Telis & Martínez-Navarrete, 2010; Wan et al., 2018), in this study, the CWC that ensured the

glassy state of the studied snack was lower than W<sub>o</sub> (Tables 2 and 4). In this sense, regardless of whether W<sub>o</sub> may be considered an optimum value for the purposes of preventing processes such as oxidative deterioration (Goula et al., 2008), it is not only a<sub>w</sub> but also Tg data that may be considered as complementary tools with which to ensure the stability of some products.

#### 4. Conclusions

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Any of the biopolymers studied permit a reduction in hygroscopicity and an increase in the glass transition temperature of the freeze-dried orange snack, without any of them being more or less effective than the others. The typical crunchy characteristics of the snack are lost at the onset of glass transition, while the colour changes are evident at its end. This indicates the need to keep the snack at temperatures below its Tg, depending on its water content. In this sense, it is recommended that the orange puree be formulated with any of the biopolymers studied and, although unnecessary from the perspective of chemical and microbiological stability, stored in refrigeration.

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# 415 Figure captions

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- 416 Figure 1. Water content (w<sub>e</sub>) water activity (a<sub>w</sub>) onset glass transition temperature (Tg<sup>0</sup>)
- 417 relationships of freeze-dried orange puree (O) and that formulated with: starch modified with
- 418 octenylsuccinic anhydride and native corn starch (OSA+NCS) or pea fibre (OSA+PF); maltodextrin
- with native corn starch (MD+NCS) or pea fibre (MD+PF); gum Arabic with bamboo fibre (GA+BF).
- The continuous lines represent the  $Tg-a_w$  fitted model, the discontinuous lines predict  $x_w-a_w$  data
- 421 obtained from the BET fitted model and the points correspond to experimental data. The black
- dashed lines show, for the O sample, how to obtain the critical water activity and water content
- 423 (CWA and CWC, respectively) for glass transition at 20°C.

Figure 2. Mid-point glass transition temperature (Tg<sup>m</sup>)-water content (x<sub>w</sub>) relationship of freeze-dried orange puree (O) and that formulated with: starch modified with octenylsuccinic anhydride and native corn starch (OSA+NCS) or pea fibre (OSA+PF); maltodextrin with native corn starch (MD+NCS) or pea fibre (MD+PF); gum Arabic with bamboo fibre (GA+BF). The lines represent the Gordon and Taylor fitted model and the points correspond to experimental data. The inner graph shows an example of one of the DSC thermograms obtained with GA+BF sample with  $a_w = 0.06$ . The dashed lines and arrows indicate the way to obtain the temperature at the onset (Tg<sup>0</sup>), midpoint (Tg<sup>m</sup>) and end-point (Tg<sup>e</sup>) of glass transition. Figure 3. Values of a) luminosity (L\*), b) chroma (C\*) and c) hue angle (h\*) of freeze-dried snack obtained from orange puree (O) and that formulated with: starch modified with octenylsuccinic anhydride and native corn starch (OSA+NCS) or pea fibre (OSA+PF); maltodextrin with native corn starch (MD+NCS) or pea fibre (MD+PF); gum Arabic with bamboo fibre (GA+BF), conditioned at different water activity (a<sub>w</sub>). Figure 4. Force (F) - distance (d) curves obtained with freeze-dried snack obtained from orange puree (O) and that formulated with: starch modified with octenylsuccinic anhydride and native corn starch (OSA+NCS) or pea fibre (OSA+PF); maltodextrin with native corn starch (MD+NCS) or pea fibre (MD+PF); gum Arabic with bamboo fibre (GA+BF), conditioned at different water activity (a<sub>w</sub>). Figure 5. Values of maximum force (Fmax) obtained in the mechanical test carried out with the freeze-dried snack obtained from orange puree (O) and that formulated with: starch modified with octenylsuccinic anhydride and native corn starch (OSA+NCS) or pea fibre (OSA+PF); maltodextrin with native corn starch (MD+NCS) or pea fibre (MD+PF); gum Arabic with bamboo fibre (GA+BF),

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conditioned at different water activity (aw).

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Table 1. Sample code as a function of the corresponding formulation.

Biopolymers added and concentration					
Sample	5g/100 g orange puree	1g/100 g orange puree			
GA+BF	gum Arabic	bamboo fibre			
MD+PF	maltodextrin	pea fibre			
MD+NC	maltodextrin	native corn starch			
OSA+PF	starch modified with octenylsuccinic anhydride	pea fibre			
OSA+NC	starch modified with octenylsuccinic anhydride	native corn starch			

Table 2. Parameters of the BET ( $W_o$ , C) and linear (Y, Z) models fitted to experimental water content-water activity (Eq. 1) and mid-point glass transition temperature-water activity (Eq. 3) data, respectively. The sample codes can be identified in Table 1.

Sample C		W₀ (g water/g dry solids)	R <sup>2</sup>	Y	Z	R <sup>2</sup>
Ο	3,0	0,074	0,911	-124,4	36,8	0,981
OSA+PF	2,7	0,062	0,708	-137,8	51,9	0,969
OSA+NCS	2,8	0,062	0,837	-140,5	52,7	0,981
MD+PF	2,6	0,066	0,901	-130,4	49,3	0,982
MD+NCS	2,3	0,071	0,580	-136,2	50,9	0,985
GA+BF	2,8	0,066	0,827	-140,2	55,6	0,987

R<sup>2</sup>: adjusted determination coefficient

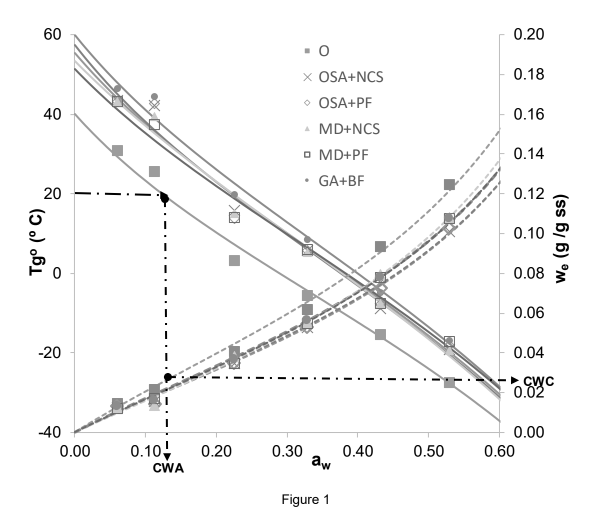
Table 3. Gordon and Taylor parameters (K and Tgs) obtained when fitting the experimental onset, midpoint and end point of the glass transition temperature (Tg<sup>0</sup>, Tg<sup>m</sup> and Tg<sup>e</sup>, respectively) - water content relationship (Eq. 2). The sample codes can be identified in Table 1.

		0	OSA+PF	OSA+NCS	MD+PF	MD+NCS	GA+BF
	К	5,69	6,93	7,24	6,16	6,52	6,62
$Tg^{o}$	Tgs	37,2	48,6	50,6	45,1	47,7	52,6
	$R^2$	0,985	0,972	0,981	0,970	0,984	0,985
	K	5,19	6,56	6,81	5,74	6,02	6,28
$Tg^{m}$	Tgs	40,0	55,4	57,5	51,4	53,4	60,0
	$R^2$	0,958	0,963	0,982	0,964	0,989	0,988
	K	4,86	6,22	6,44	5,36	5,56	5,97
Tg <sup>e</sup>	Tgs	44,8	62,3	64,4	57,8	59,2	67,3
	R <sup>2</sup>	0,953	0,951	0,981	0,957	0,985	0,991

R<sup>2</sup>: adjusted determination coefficient

Table 4. Values of the critical water content (CWC) and critical water activity (CWA) for the onset (°) and end (°) point of the glass transition at 20 and 4°C. The sample codes can be identified in Table 1.

		0	OSA+PF	MD+PF	OSA+NCS	MD+NCS	GA+BF
	CWA°	0,095	0,170	0,160	0,165	0,169	0,190
20 °C	CWA <sup>e</sup>	0,165	0,279	0,275	0,275	0,275	0,302
20 C	CWC°	0,019	0,026	0,026	0,026	0,026	0,031
	CWC <sup>e</sup>	0,032	0,042	0,044	0,042	0,043	0,049
	0)4/40	0.040	0.000	2 222	0.005	0.005	0.044
	CWA°	0,213	0,293	0,289	0,285	0,295	0,311
4 °C	CWA <sup>e</sup>	0,305	0,404	0,409	0,400	0,405	0,425
4 0	CWC°	0,040	0,044	0,046	0,044	0,046	0,050
	CWC <sup>e</sup>	0,057	0,063	0,067	0,063	0,067	0,071



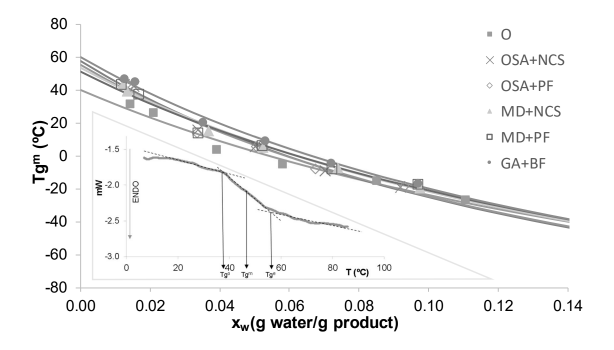
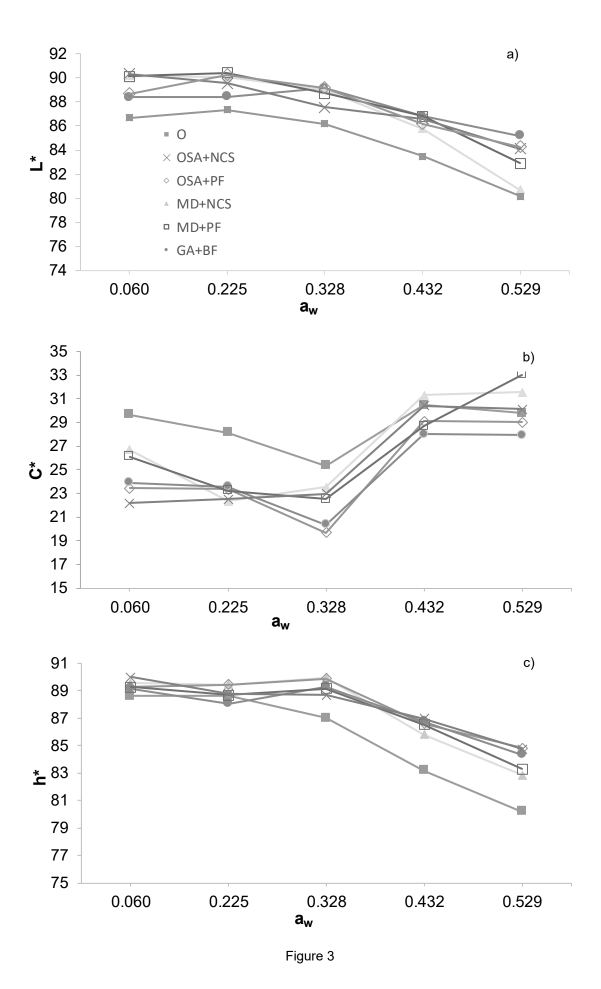


Figure 2



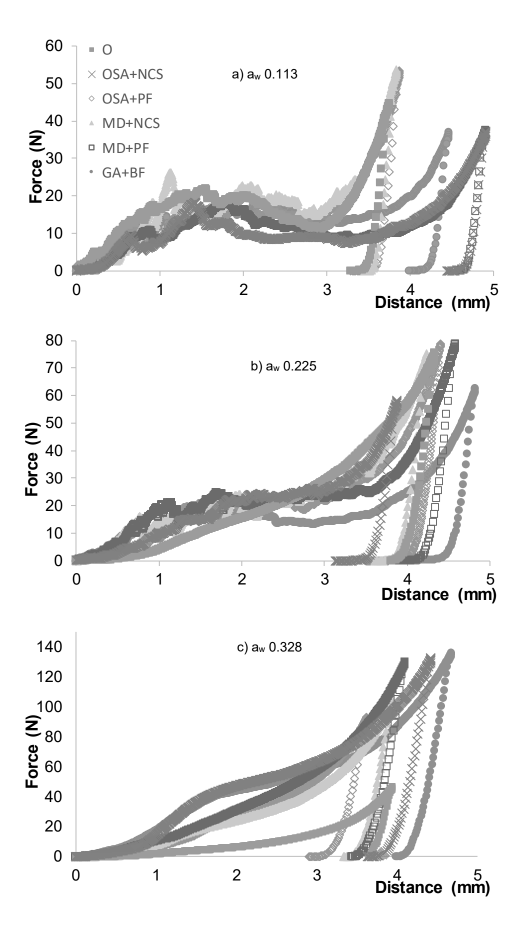


Figure 4

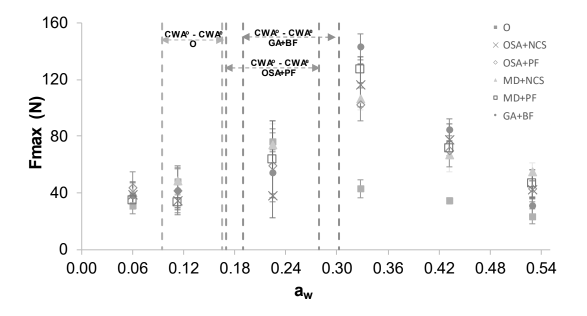


Figure 5

## **CREDIT AUTHOR STATEMENT**

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**Nuria Martínez-Navarrete**: Conceptualization, Formal analysis, Funding acquisition, Methodology, Project Administration, Supervision, Writing – original draft, Writing – review & editing.

\*Conflict of Interest Form

The authors have no competing interests to declare  $% \left( 1\right) =\left( 1\right) \left( 1\right) \left($