1	EFFECTS OF GELATIN ORIGIN, BOVINE-HIDE AND TUNA-SKIN, ON THE
2	PROPERTIES OF COMPOUND GELATIN-CHITOSAN FILMS
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10	** This centre has implemented and maintains a Quality Management System in
11	compliance with ISO standard 9001:2000.
12	
13	ABSTRACT
14	With the purpose to improve the physico-chemical performance of plain gelatin
15	and chitosan films, compound gelatin-chitosan films were prepared. The effect
16	of the gelatin origin (commercial bovine-hide gelatin and laboratory-made tuna-
17	skin gelatin) on the physico-chemical properties of films was studied. The
18	dynamic viscoelastic properties (elastic modulus G', viscous modulus, G" and
19	phase angle) of the film forming solutions upon cooling and subsequent heating
20	revealed that the interactions between gelatin and chitosan were stronger in the
21	blends made with tuna-skin gelatin than in the blends made with bovine-hide

23 (~18% water solubility for tuna vs 30% for bovine) and more deformable (~68%

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gelatin. As a result, the fish gelatin-chitosan films were more water resistant

24 breaking deformation for tuna vs 11% for bovine) than the bovine gelatin-25 chitosan films. The breaking strength of gelatin-chitosan films, whatever the 26 gelatin origin, was higher than that of plain gelatin films. Bovine gelatin-chitosan 27 films showed a significant lower water vapor permeability (WVP) than the 28 corresponding plain films, whereas tuna gelatin-chitosan ones were only 29 significantly less permeable than plain chitosan film. In spite of gelatin-chitosan 30 interactions, all the chitosan-containing films exhibited antimicrobial activity 31 against S. aureus, a relevant food poisoning. Mixing gelatin and chitosan may 32 be a means to improve the physico-chemical performance of gelatin and 33 chitosan plain films, especially when using fish gelatin, without altering the 34 antimicrobial properties.

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Key words: bovine-hide gelatin, chitosan, fish gelatin, physico-chemical
properties, *S. aureus*, edible films

39 INTRODUCTION

40 As a consequence of the problems associated with disposal of packaging plastics, there is a growing interest concerning the development of 41 42 biodegradable materials. These newer materials can be obtained from several 43 sources, which include proteins (collagen/gelatin, soya, whey, wheat, etc), 44 polysaccharides (chitosan, starch, cellulose) and lipids (wax, fatty acids) 45 (Gennadios, Hanna & Kurth, 1997; Tharanathan, 2003). Although the entire 46 substitution of petrochemical polymers with bioplastics may not be possible due 47 to the worse physicochemical properties of the latter, it is necessary to research 48 on the improvement of such properties, as well as on the new applications, in 49 order that bioplastics corner the market and, therefore, the traditional synthetic 50 plastics are partially substituted.

51 Gelatin is a protein with a broad range of functional properties and applications, 52 including film-forming ability. Bovine and porcine wastes are the most frequent 53 sources to obtain gelatin of good quality. However other sources of gelatin are becoming increasingly relevant, such as fish bones and skins (Gomez-Guillen, 54 55 Turnay, Fernandez-Diaz, Ulmo, Lizarbe & Montero, 2002). Whatever the 56 species of origin, gelatin films fail in terms of mechanical properties and water 57 resistance, which may limit its field of application. Several strategies have been 58 used to improve the physical performance of gelatin films. These include 59 chemical or enzymatic treatments (Cao, Fu & He, 2007; Chiou et al., 2008; de Carvalho & Grosso, 2004; Spanneberg, Osswald, Kolesov, Anton, Radusch & 60 61 Glomb, 2010; Zhang et al., 2010)) and mixing with other polymers, as 62 composite films may be designed to take advantages of pure components 63 (Garcia, Pinotti, Martino & Zaritzky, 2004). For example mixing with apolar

64 components such fatty acids or oils reduces water vapour transmission rate 65 (Jongjareonrak, Benjakul, Visessanguan & Tanaka, 2006; Limpisophon, Tanaka 66 & Osako, 2010; Perez-Mateos, Montero & Gomez-Guillen, 2009). Furthermore 67 polymers can establish new bonds that may enhance the properties of the 68 resulting materials (Denavi, Perez-Mateos, Anon, Montero, Mauri & Gomez-69 Guillen, 2009; Sionkowska, Wisniewski, Skopinska, Kennedy & Wess, 2004). 70 Gelatin has been blended with casein (Chambi & Grosso, 2006), pectin (Liu, 71 Liu, Fishman & Hicks, 2007), chitosan (Arvanitoyannis, Nakayama & Aiba, 72 1998; Kolodziejska & Piotrowska, 2007; Kolodziejska, Piotrowska, Bulge & 73 Tylingo, 2006), starch (Arvanitoyannis et al., 1998) and soy protein (Denavi et 74 al., 2009), achieving in general an improvement of its physical performance.

75 Chitosan (poly b-(1,4)N-acetyl-D-glucosamine) polymer is industrially produced 76 by chemical deacetylation of the chitin found in arthropod exoskeletons. 77 Chitosan is largely utilized not only due to its film forming capability but also 78 because of its antimicrobial properties (Helander, Nurmiaho-Lassila, 79 Ahvenainen, Rhoades & Roller, 2001; Jung, Youn, Lee, No, Ha & 80 Prinyawiwatkul, 2010). For this reason, the use of chitosan as an edible coating 81 or film to extend the shelf-life of foods and inhibit pathogens is of growing 82 interest (Aider, 2010; Gomez-Estaca, Montero, Gimenez & Gomez-Guillen, 83 2007; Lopez-Caballero, Gomez-Guillen, Perez-Mateos & Montero, 2005). 84 Specifically, chitosan has been found to be active against S. aureus (Fernandes 85 et al., 2008). Some S. aureus strains are able to produce staphylococcal 86 enterotoxins and are the causative agents of staphylococcal food poisonings 87 (Le Loir, Baron & Gautier, 2003). The antimicrobial action of chitosan is 88 supposed to be derived from the positive charge that amino groups present at

89 acidic pH (below 6.5), which lead to cellular membrane depolarization and 90 microbial death. However the main drawback of chitosan under this condition is 91 its intrinsic water solubility, which limits the utilisation as self-standing packaging 92 material. Attempts to increase the water resistance of chitosan have been made 93 including crosslinking with glutaraldehyde, glyoxal or epichlorohydrin (Suto & Ui, 94 1996; Tual, Espuche, Escoubes & Domard, 2000; Zheng, Du, Yu & Xiao, 2000). 95 However, in a subsequent work (Tang, Du & Fan, 2003) it was confirmed that 96 the antimicrobial capacity of chitosan films diminishes with an increase in the 97 cross-linking. Mixing of chitosan with other biopolymers to obtain more insoluble 98 matrices has been proved as an effective means to improve the water 99 resistance of chitosan maintaining its antimicrobial properties (Fernández-Saiz 100 et al., 2008). Chitosan has also been blended with other biopolymers such as 101 methylcellulose and starch resulting in an improvement of its physico-chemical 102 properties (Garcia et al., 2004; Garcia, Pinotti & Zaritzky, 2006).

103 According to Taravel & Domard (1995), the interactions between gelatin and 104 chitosan are produced by both electrostatic and hydrogen bonding, with the 105 blends taking on new physical properties and thus becoming suited to potential 106 new applications (Sionkowska et al., 2004). Accordingly, combining these 107 biopolymers seems to be a promising way to enhance the physical properties of the resulting materials (Huang, Onyeri, Siewe, Moshfeghian & Madihally, 2005; 108 109 Mao, Zhao, Yin & Yao, 2003). There are some reports dealing with the physico-110 chemical properties of compound fish gelatin-chitosan films as well as its 111 improvement by adding different crosslinkers (Arvanitoyannis et al., 1998; 112 Kolodziejska et al., 2007; Kolodziejska et al., 2006). However there is no 113 previous report on the effect of the gelatin origin on the film forming ability and

the physico-chemical properties of the gelatin-chitosan admixtures. The objective of the present work has thus been to study the physico-chemical properties of compound gelatin-chitosan films as function of gelatin origin: a commercial bovine-hide gelatin and a laboratory-obtained tuna-skin gelatin. The antimicrobial activity of the resulting films over *S. aureus* was also evaluated.

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120 MATERIALS AND METHODS

121 Materials

Tuna-skin gelatin was extracted according to a patented method (GómezGuillén & Montero, 2001). A commercial type A bovine-hide gelatin (Bloom
200/220) was purchased from Sancho de Borja S.L. (Zaragoza, Spain).
Chitosan from shrimp shells (85 % deacetylated; 141,000 Da) was purchased
from Guinama (Valencia, Spain). Glycerol and sorbitol were from Panreac
(Barcelona, Spain).

For the microbial analyses, brain heart infusion (BHI) broth was purchased from Oxoid (Basingstoke, UK) and bacteriological agar were from Scharlau (Barcelona, Spain). *Staphylococcus aureus* (CECT 240) was obtained from the Spanish Type Culture Collection (Valencia, Spain).

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133 **Preparation of the film-forming solutions and films**

Seven different formulations were prepared: The batches were: B (bovine-hide gelatin), T (tuna-skin gelatin), Ch (chitosan), B-Ch 0.75% (bovine-hide gelatin plus 0.75% chitosan), T-Ch 0.75% (tuna-skin gelatin plus 0.75% chitosan), BCh 1.5% (bovine-hide gelatin plus 1.5% chitosan), T-Ch 1.5% (tuna-skin gelatin

138 plus 1.5% chitosan). The gelatin film-forming solutions (solutions B and T) were 139 prepared at a concentration of 4 g gelatin/100 ml of distilled water. The chitosan 140 film-forming solution (solution Ch) was prepared by dissolving chitosan in a 141 proportion of 3 g/100 mL in 0.15 M acetic acid. The gelatin-chitosan film-forming 142 solutions (solutions B-Ch 0.75%, B-Ch 1.5%, T-Ch 0.75%, T-Ch 1.5%) were 143 prepared by mixing a 4% bovine-hide or tuna-skin gelatin solution with a 1.5% 144 or 3% solution of chitosan in 0.15 M acetic acid, in a proportion of 1:1 (v/v), to 145 obtain a film-forming solution containing 2% gelatin and 0.75% or 1.5% 146 chitosan, respectively. A mixture of glycerol and sorbitol was added to the film-147 forming solutions as previously described (Thomazine, Carvalho & Sobral, 148 2005) at a concentration of 0.15 g each per gram of the total polymeric agent 149 (gelatin and/or chitosan). Plasticizing molecules of this kind reduce inter-chain 150 interactions, improving film flexibility and, consequently, film handling.

The film-forming solutions, which had a pH of 4.6 \pm 0.3 in all cases, were warmed and stirred at 45 °C to obtain a good blend. The films were made by casting an amount of 40 mL onto plexiglass plates (12 x 12 cm) and drying at 45 °C in a forced-air oven for 15 h to yield a uniform thickness of 100 µm \pm 11 in all cases except the 0.75% chitosan formulations, which was 80 µm \pm 7 thick. Prior to the determinations, the films were conditioned at 22 °C over a saturated solution of NaBr (58% RH) in desiccators for 3 d.

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159 Physical characterization of the film-forming solutions

Dynamic viscoelastic analysis of the film-forming solutions was carried out on a
Bohlin CSR-10 rheometer/rotary viscometer (Bohlin Instruments Ltd.,

Gloucestershire, UK) using a cone-plate geometry (cone angle = 4°, gap = 0.15 mm). Cooling and heating from 40 to 6 °C and back to 40 °C took place at a scan rate of 1 °C/min, a frequency of 1 Hz, and a target strain of 0.2 mm. The elastic modulus (G'; Pa), viscous modulus (G''; Pa) and phase angle (°) were determined as functions of temperature. Two determinations were performed for each sample, with an experimental error of less than 6% in all cases.

The film-forming solutions were poured into glasses 2.3 cm in diameter and 3.6 cm in height and left to mature in a refrigerator at 2 °C for 16-18 h. Gel strength at 9±1 °C was determined on an Instron model 4501 Universal Testing Machine (Instron Co., Canton, MA, USA) with a 100 N load cell, a crosshead speed of 1 mm/s, and a flat-faced cylindrical plunger 1.27 cm in diameter. The maximum force (g) reading was taken when the plunger had penetrated 4 mm into the gelatin gels.

175 **Physico-chemical characterization of the films**

176 Mechanical properties

177 A puncture test was performed to determine the maximum strength and 178 deformation of the films at the breaking point. The films were placed in a cell 5.6 179 cm in diameter and perforated to the breaking point using an Instron model 4501 180 Universal Testing Machine (Instron Co., Canton, MA, USA) with a round-ended 181 stainless-steel plunger 3 mm in diameter at a crosshead speed of 1 mm/s and a 182 100 N load-cell. Breaking strength was expressed in N and breaking deformation 183 in percent as previously described (Sobral, Menegalli, Hubinger & Roques, 184 2001). All determinations were the means of at least five measurements.

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186 Thermal properties

187 Calorimetric analysis was performed using a model TA-Q1000 differential 188 scanning calorimeter (TA Instruments, New Castle, DE, USA) previously 189 calibrated by running high purity indium (melting point 156.4 °C; enthalpy of melting 28.44 W/g). Sample amounts on the order of 10 mg ± 0.002 weighed 190 191 out using a Sartorious model ME235S electronic balance (Goettingen, 192 Germany) were tightly encapsulated in aluminum pans and scanned under dry 193 nitrogen purge (50 mL/min). Freshly conditioned films were rapidly cooled to 0 194 °C and scanned between 0 and 90 °C, at a rate of 10 °C/min. Glass transition 195 temperatures, Tg (°C), were determined on first heating scans consistently with 196 the other physical determinations carried out on the same original (after 197 conditioning at 58% RH) material. Tg was estimated as the midpoint of the line 198 drawn between the temperature at the intersection of the initial tangent with the 199 tangent through the inflection point of the trace and the temperature of the 200 intersection of the tangent through the inflection point with the final tangent. Tg 201 data were the mean values of at least three replications per film sample.

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203 Water solubility

Film portions measuring 4 cm² were placed in aluminum capsules with 15 mL of distilled water and shaken gently at 22 °C for 15 h. The solution was then filtered through Whatman no. 1 filter paper to recover the remaining undissolved film, which was desiccated at 105 °C for 24 h. Film water solubility was calculated using the equation *FS* (%) = $((W_o-W_f)/W_o) \cdot 100$, where W_o was the initial weight of the film expressed as dry matter and W_f was the weight of the

210 undissolved desiccated film residue. All determinations were carried out in211 triplicate.

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213 Water vapour permeability

214 Water vapour permeability (WVP) was determined by a gravimetric method. Films were attached over the openings of cells (permeation area = 15.9 cm^2) 215 216 containing silica gel, and the cells were placed in desiccators with distilled water 217 at 22 °C. The cells were weighed daily for 7 d. Water vapour permeability was 218 calculated using the equation $WVP = w \cdot x \cdot t^{-1} \cdot A^{-1} \cdot \Delta P^{-1}$, where w was weight gain 219 (g), x film thickness (mm), t elapsed time (h) for the weight gain, and ΔP the 220 partial vapour pressure difference between the dry atmosphere and pure water (2642 Pa at 22 °C). Results have been expressed as g·mm·h⁻¹·cm⁻²·Pa⁻¹. All 221 222 determinations were carried out in duplicate.

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224 Microbial analysis

225 For microbial analysis, prior to film casting, the pH of all the film-forming 226 solutions was adjusted to 6. This pH was considered suitable to enable any 227 potential in vitro antibacterial effects of the film-forming solutions and/or films to 228 be attributed to the chitosan rather than to the pH. The antimicrobial activity of 229 both the film-forming solutions and the films was determined against S. aureus. The bacteria were stored at -80 °C in BHI broth with 25 % glycerol until use. 230 231 Strain was grown in BHI (Oxoid) broth at 37 °C overnight to a final bacterial concentration of 107-108 cfu/mL. Spread plates of brain heart agar were 232 233 prepared. Sterile filter paper disks immersed in the film-forming solutions or

pieces of the films themselves were placed on the plate surfaces and incubated
at 37 °C for 24 h. The appearance of a clear area below or around the film or
filter paper disks was deemed to be positive for antimicrobial activity.

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238 Statistical analysis

Statistical tests were performed using the SPSS® computer program (SPSS Statistical Software, Inc., Chicago, IL, USA). One-way analysis of variance was carried out. Differences between pairs of means were compared using a Tukey test. The level of significance was set at $p \le 0.05$.

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245 RESULTS AND DISCUSSION

246 Film-forming solutions

247 The changes in the viscoelastic properties of the film-forming solutions during 248 cooling and subsequent heating have been plotted in Figure 1. Solution B had a 249 higher elastic modulus (G') value and had thermal transition (gelling and 250 melting) points at higher temperatures than solution T. These findings are 251 indicative both of bovine-hide gelatin's greater capacity to refold into triple-helix 252 chains as it cools and its higher thermostability. These findings were fully 253 expected and wholly consistent with the different origins of the gelatins, as 254 reported previously (Joly-Duhamel, Hellio, Ajdari & Djabourov, 2002; Joly-255 Duhamel, Hellio & Djabourov, 2002). Bovine-hide gelatin has a higher imino 256 acid (Pro+Hyp) content than tuna-skin gelatin (210 residues in bovine gelatin 257 vs. 185 residues in tuna gelatin) (Gomez-Estaca, Montero, Fernandez-Martin &

258 Gomez-Guillen, 2009), and this is well known to be related to enhanced 259 physical properties and higher thermostability of the resulting gelatin gels 260 (Ledward, 1986; Norland, 1990).

261 Adding chitosan modified the viscoelastic properties of both gelatins. The 262 maximum G' values were considerably lower than the values for solutions B and 263 T, and the thermal transition points also occurred at lower temperatures. This is 264 indicative of a pronounced loss of the gelatin's ability to refold into triple-helix chains in the presence of chitosan. A decrease in percentage renaturation of 265 266 food grade gelatin by incorporation of chitosan has been previously reported 267 (Arvanitoyannis et al., 1998). Adding chitosan at the higher concentration led to 268 a substantial increase in the viscous modulus (G") during cooling. Furthermore, 269 phase angle values at both concentrations were higher than the values for the 270 corresponding gelatin solutions at temperatures below 10 °C. Both these effects 271 also indicate that chitosan interferes with protein network formation as a result 272 of gelatin-chitosan interactions. These effects were considerably more marked 273 for the tuna-skin gelatin, especially at the higher chitosan concentration, as 274 indicated by the higher increase in G" and the not so steep slope of the phase 275 angle during cooling and subsequent heating. Thus, the chitosan interacted 276 more with the tuna-skin gelatin than with the bovine-hide gelatin. Interactions 277 between chitosan and collagen have previously been described (Sionkowska et al., 2004) and were also reflected by an increase in the viscosity values. In the 278 279 present experiment the collagen had previously been denatured to obtain 280 soluble gelatin, presumably heightening this interaction. The interactions 281 between gelatin and chitosan are produced by both electrostatic and hydrogen 282 bonding (Taravel & Domard, 1995). The former are a consequence of the

283 different charges of gelatin, an anionic biopolymer, and the cationic chitosan. 284 The latter occur extensively between the –COOH, -NH₂, and –OH groups on the 285 amino acids in the gelatin and the –OH and –NH₂ groups on the chitosan, and 286 are particularly notable at low temperatures. Regarding the contribution of 287 electrostatic interactions, it should be noted that both gelatins are type A, i.e., 288 they had been subjected to an acidic pre-treatment, therefore both are presumed to have similar isoelectric points, being net positively charged at pH 289 290 6. The most likely explanation for such a different interaction could be the 291 differing imino acid content (Pro+Hyp), which was considerably lower in the 292 tuna-skin gelatin. The abundance of pyrolidine iminoacid content (Pro, and 293 especially Hyp), is well known to be directly implicated in the gelling mechanism 294 promoting the formation of junction zones of triple collagen-like helices 295 (Ledward, 1986). In this connection, the lower iminoacid content in the tuna 296 gelatin would lead to a less extensive self-aggregation of the gelatin chains, 297 which might have contributed to an increment of the gelatin-chitosan 298 interactions.

299 The gel strength determinations (Figure 2) carried out on the cold-matured film-300 forming solutions revealed a decrease in the mechanical properties of the 301 gelatin gels when chitosan was added, which is ascribed not only to a dilution 302 effect of the gelatin in the film-forming solution but also to the gelatin-chitosan 303 interactions. The effect was more pronounced at the higher concentration of 304 added chitosan, indicating that chitosan not only interfered with nucleation point 305 formation but also with triple-helix chain growth during cold maturation. No 306 differences were found between both types of gelatin.

307

308 Films

309 Warm-water fish gelatins have been reported to have guite similar properties to 310 mammalian gelatins (Gilsenan & Ross-Murphy, 2000), and the tuna-skin gelatin 311 films showed Tg values that were very slightly lower (within experimental error) 312 than the bovine-hide gelatin films (Figure 3a, curves B and T; Table 1), in 313 agreement with previous reports (Gomez-Estaca et al., 2009). Films made from 314 various chitosans (different degrees of deacetylation, OH and amino groups, 315 crystallinity, etc.) in the presence of various amounts of different hydrophilic 316 plasticizers have been reported to exhibit very high (>>100°C) glass transition 317 temperatures (Suyatma, Tighzert & Copinet, 2005). The chitosan films tested 318 here, however, had a rather low glass transition temperature, even lower than 319 the values for the two gelatins (Table 1); as the ratio between the polymer and 320 the glycerol plus sorbitol equiproportional mixture remained constant in all 321 cases, the considerable higher water content in Ch (~22%) was presumably a 322 main responsible. The value seemed to be consistent with a phase diagram 323 previously published (Lazaridou & Biliaderis, 2002) for a system comprising 70 324 % chitosan and 30 % sorbitol at different moisture contents.

325 As discussed above, chitosan has been reported to interact molecularly with 326 collagen by forming a wide range of blends where new hydrogen bonding 327 networks appear, the triple helical structure of the collagen being replacing as 328 the chitosan level increases (Sionkowska et al., 2004). In other work, the 329 formation of gelatin-chitosan polyelectrolyte complexes has been shown to give 330 rise to a decrease of the crystallinity of the system (Yin, Yao, Cheng & Ma, 331 1999). Other factors affecting the properties of gelatin-chitosan films are the 332 evaporation temperature and the presence of plasticizers (water and polyols)

333 (Arvanitoyannis et al., 1998). Furthermore, the interactions between gelatin and 334 chitosan are supposed to be dependent on the physical and chemical properties 335 of the gelatin, which vary considerably with gelatin origin. As a general rule, 336 mammalian gelatins afford better physical properties and thermostability than 337 fish gelatins, and there are, moreover, appreciable differences among gelatins 338 from different fish species (Gomez-Guillen et al., 2002). The main factors 339 determining the physical properties of a gelatin and hence gelatin quality are the 340 amino acid composition and the molecular weight distribution (Gomez-Guillen et 341 al., 2002). Mammal's gelatins are typically richer in imino acids whereas the 342 average molecular weight highly depends on the extraction procedures. Both 343 these factors will presumably affect the interactions between the gelatin and 344 chitosan. Thus, the physico-chemical properties, and ultimately the potential 345 applications of the resulting materials will all be closely related to the properties 346 of the polymer admixture employed.

347 In the present work, the glass transition temperatures of compound films were 348 higher than those for the single components, indicating some level of 349 interaction/association. At each of the two chitosan levels, the Tg values for 350 both series of compound gelatin-chitosan blends were quite close and at the 351 lower Ch concentration were nearly 6 °C higher than for the gelatins alone 352 (Table 1). Augmenting the chitosan component resulted in a further increase of 353 around 3 °C in the Tg. Higher water contents may produce higher plasticization 354 effects in the blended films by an increase in the macromolecules mobility. 355 However, it appeared clearly overcame by the increase matrix density effect 356 caused by the strong associations between gelatins and chitosan, with the final 357 result of glass transition processes occurring at higher temperatures in the

358 blends than in the single films. Initially, the polyols mixture is highly 359 hygroscophic so that water may plasticize both the low-molecular-weight high-molecular-weight components. 360 mixture and the The water-alone 361 contribution to the plasticization of the different polymeric blends seemed hardly 362 difficult to being separated from the whole plasticizer system. Additionally the 363 humidity values for the B-Ch and T-Ch films were quite similar; small 364 differences in water content could be basically modulated by the glycerol plus 365 sorbitol plus water entire system, so that small variations in water are expected 366 not to play an essentially differential role in the plasticization process and Tg 367 values of the complex films. Therefore, the glass transition behaviour of the B-368 and T-type compounded films with Ch reasonably showed a paralleled 369 evolution.

370 All these findings were consistent with the thermal scan data obtained for the 371 corresponding gels by rheometry, as shown in Figure 1 (bottom, curves a and 372 b). At both Ch levels hysteresis of the sol-gel-sol transitions increased in the 373 film-forming solutions of both gelatin-chitosan blends relative to those in the 374 corresponding gelatins. Adding the larger amount of chitosan increased the 375 complexity of the gel-sol and sol-gel profiles of both systems, particularly T-376 Ch1.5, which could be interpreted as a certain level of phase separation in the 377 systems. Gelatin-rich phases (more frequent in the T systems than in the B 378 systems) have higher gel-sol and sol-gel transition temperatures and, 379 conversely, chitosan-rich phases have lower gel-sol and sol-gel transition 380 temperatures. Although this gelling behaviour is not necessarily transferred to 381 the respective films, DSC did not reveal any phase separation in the Ch1.5 382 systems.

383 Table 1 lists the glass transition temperature, mechanical properties (breaking 384 strength and breaking deformation), solubility, and water vapour permeability of 385 the different films. Breaking strength and water solubility values were similar for 386 both gelatin films (B and T), a result attributable to the presence of very high-387 molecular-weight aggregates (Gomez-Estaca et al., 2009). In comparison, the 388 chitosan film (Ch) was stronger than either of the gelatin films (B and T). 389 although it was also more soluble in water. Perhaps the most interesting feature 390 was that the breaking deformation value for film T was ~10 times higher than 391 the values for films B and Ch. Deformation differences among gelatin films have 392 been previously attributed to different imino acid contents, giving bovine-hide 393 gelatin a higher degree of molecular rigidity (Gomez-Estaca et al., 2009). The 394 water vapour permeability values showed all the films to be quite permeable, 395 entirely consistent with films prepared from biopolymers plasticized with polyols. 396 This has been reported to be also a result of increases in the free volume 397 between polymer chains caused by a reduction in intermolecular attractive 398 forces, making the polymer network less dense and thus more permeable (Cuq, 399 Gontard, Cug & Guilbert, 1997). However, some differences among the 400 formulations were found. Thus, the Ch film was more permeable to water 401 vapour than the B and T films. Gelatin origin was also a factor, T being less 402 permeable than B, again attributable to both the amino acid profile and the molecular weight distribution (Avena-Bustillos et al., 2006; Gomez-Estaca et al., 403 404 2009). Comparisons among different authors are somewhat difficult because of 405 the different film manufacture and measurement procedures. There is a study in 406 which the WVP of compound fish gelatine-chitosan films as well as that of the 407 pure components were studied (Kolodziejska et al., 2007). In this work neither

408 the gelatine and the chitosan pure films nor the compound ones were 409 significantly different. In other study (Arvanitoyannis et al., 1998) mammal 410 gelatine and chitosan were blended at a ratio 1:1, plasticized with different 411 amounts and type of plasticizers and dried at different temperatures, resulting in 412 WVPs ranging from 1.1 to 6.3×10^{-11} g m/m² s Pa, which are three orders of 413 magnitude lower than ours. In this case the single components were not 414 analyzed.

415 On the whole, the physico-chemical properties of the compound gelatin-416 chitosan films were more appropriated for practical purposes than the properties 417 of the films made from a single biopolymer only (films B, T, and Ch). As a 418 general rule, the compound films had breaking strengths similar to the values 419 for the Ch films ($p \le 0.05$), the strongest films, and significantly higher ($p \le 0.05$) 420 than the values for the gelatin films. Breaking deformation either stayed the 421 same (B-Ch blends) or decreased (T-Ch blends) significantly (p≤0.05) 422 compared with the respective gelatin films. In spite of the decrease of breaking 423 deformation in T-Ch films compared to T ones, these mixtures showed a higher 424 breaking deformation than B, Ch and B-Ch films. Solubility of the compound 425 films was significantly lower ($p \le 0.05$) compared with that of chitosan films 426 whatever the gelatin employed. Furthermore, in tuna-skin gelatin-chitosan 427 mixtures the solubility was significantly ($p \le 0.05$) lower than that of the tuna-skin 428 gelatin. Water vapour permeability of the compound films decreased 429 significantly ($p \le 0.05$) compared with the single biopolymer component films Ch 430 and B, and was similar to the values for T film, the one with the lower WVP 431 among plain films, in all cases. All the alterations in the physico-chemical 432 properties of the films can be attributed to the gelatin-chitosan interactions

433 previously reported by different researchers (Sionkowska et al., 2004; Taravel 434 et al., 1995; Yin et al., 1999) and confirmed in this study by the results for the 435 viscoelastic properties and gel strength of the film-forming solutions. The 436 interactions were observed to be stronger in the case of the tuna-skin gelatin, 437 hence the alterations in the physical and chemical properties upon blending with 438 chitosan were most discernible for the T-Ch formulations, especially regarding 439 the film breaking deformation and water solubility.

440 It was also evaluated the possible loss of antibacterial activity of chitosan upon 441 mixing with the bovine or the tuna gelatins. For this purpose, S. aureus was 442 selected as a model of gram-positive microorganism that is also a relevant food 443 poisoning microorganism. As expected, neither the B and T gelatin film-forming 444 solutions nor the B and T gelatin films employed as controls exhibited any 445 antibacterial effect (Figure 4). On the other hand, all the chitosan-containing 446 formulations displayed distinctly discernible antimicrobial effects against S. 447 aureus. For purposes of economy, only the photographs for the T films have 448 been included here. So, the good inhibitory effects of the chitosan against S. 449 aureus were maintained in spite of the gelatin-chitosan interactions observed. 450 Other authors (Fernandez-Saiz, Lagaron, Hemandez-Munoz & Ocio, 2008) also 451 found a gliadin-chitosan blend to effectively reduce the growth of S. aureus.

In conclusion, mixing gelatin and chitosan may be a means to improve the
physico-chemical performance of gelatin and chitosan plain films, especially
when using fish gelatin, without altering the antimicrobial properties.

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614 Figure captions

615

Figure 1. Dynamic viscoelastic properties of the film-forming solutions. B
(bovine-hide gelatin), T (tuna-skin gelatin), Ch (chitosan), B-Ch 0.75% (bovinehide gelatin plus 0.75% chitosan), T-Ch 0.75% (tuna-skin gelatin plus 0.75%
chitosan), B-Ch 1.5% (bovine-hide gelatin plus 1.5% chitosan), T-Ch 1.5%
(tuna-skin gelatin plus 1.5% chitosan).

Figure 2. Gel strength of the film-forming solutions after cold maturation at 2 °C.Batch designations as in Figure 1.

- Figure 3. Glass transition behaviour of one (a) and two (b) component films. (a)
- 624 One-component films: Bovine-hide gelatin, curve B; tuna-skin gelatin, curve T;
- 625 chitosan, curve Ch. (b) Two-component films: curves B-Ch 0.75%, B-Ch 1.5%,
- 626 T-Ch 0.75%, T-Ch 1.5%.
- 627 Figure 4. Antimicrobial activity of the tuna-skin gelatin (T) and the tuna-skin
- 628 gelatin plus 1.5% chitosan (T-Ch 1.5%) film-forming solutions (FFS) and films
- 629 against *S. aureus*. Arrows indicate the inhibition areas.

Table 1. Physico-chemical properties (glass transition temperature, breaking strength and breaking deformation by puncture test, solubility, and water vapour permeability) of the film batches [B (bovine-hide gelatin), T (tuna-skin gelatin), Ch (chitosan), B-Ch 0.75% (bovine-hide gelatin plus 0.75% chitosan), T-Ch 0.75% (tuna-skin gelatin plus 0.75% chitosan), B-Ch 1.5% (bovine-hide gelatin plus 1.5% chitosan), T-Ch 1.5% (tuna-skin gelatin plus 1.5% chitosan)].

В	Т	Ch	B-Ch 0.75%	B-Ch 1.5%	T-Ch 0.75%	T-Ch 1.5%
41.6±0.6 ^a	41.0±0.6 ^a	34.6±0.7 ^b	46.9±0.8 ^c	49.7±0.9 ^d	47.4±0.9 ^c	49.6±09 ^d
$10.7\pm2.2^{\text{a}}$	$8.5\pm1.6^{\text{a}}$	$23.0\pm2.9^{\text{b}}$	$18.4\pm3.6^{\text{b}}$	$18.8\pm3.3^{\text{b}}$	$13.2\pm2.2^{\text{a}}$	$20.0\pm3.5^{\text{b}}$
$14.1\pm5.0^{\text{a}}$	$154\pm36^{\text{b}}$	$19.2\pm4.7^{\text{a}}$	$11.0\pm3.1^{\text{a}}$	$19.9\pm4.3^{\text{a}}$	$68\pm\mathbf{6^c}$	$40\pm9^{\text{a}}$
$34.3\pm0.6^{\text{ab}}$	$39.9 \pm \mathbf{1.3^{b}}$	82 ± 11^{c}	$29.4\pm2.0^{\text{ab}}$	$33.1\pm1.8^{\text{ab}}$	$17.5\pm1.6^{\text{a}}$	$19.0\pm4.4^{\text{a}}$
$\textbf{2.20}\pm\textbf{0.11}^{a}$	$1.65\pm0.4^{\text{b}}$	$\textbf{2.43} \pm \textbf{0.26}^{c}$	$1.91\pm0.06^{\text{b}}$	$1.44\pm0.23^{\text{b}}$	$1.75\pm0.37^{\text{b}}$	$1.98\pm0.13^{\text{ab}}$
	$\begin{array}{c} & \\ 41.6 \pm 0.6^{a} \\ \\ 10.7 \pm 2.2^{a} \\ \\ 14.1 \pm 5.0^{a} \\ \\ 34.3 \pm 0.6^{ab} \end{array}$	41.6 ± 0.6^{a} 41.0 ± 0.6^{a} 10.7 ± 2.2^{a} 8.5 ± 1.6^{a} 14.1 ± 5.0^{a} 154 ± 36^{b} 34.3 ± 0.6^{ab} 39.9 ± 1.3^{b}	41.6 ± 0.6^{a} 41.0 ± 0.6^{a} 34.6 ± 0.7^{b} 10.7 ± 2.2^{a} 8.5 ± 1.6^{a} 23.0 ± 2.9^{b} 14.1 ± 5.0^{a} 154 ± 36^{b} 19.2 ± 4.7^{a} 34.3 ± 0.6^{ab} 39.9 ± 1.3^{b} 82 ± 11^{c}	41.6 ± 0.6^{a} 41.0 ± 0.6^{a} 34.6 ± 0.7^{b} 46.9 ± 0.8^{c} 10.7 ± 2.2^{a} 8.5 ± 1.6^{a} 23.0 ± 2.9^{b} 18.4 ± 3.6^{b} 14.1 ± 5.0^{a} 154 ± 36^{b} 19.2 ± 4.7^{a} 11.0 ± 3.1^{a} 34.3 ± 0.6^{ab} 39.9 ± 1.3^{b} 82 ± 11^{c} 29.4 ± 2.0^{ab}	41.6 ± 0.6^{a} 41.0 ± 0.6^{a} 34.6 ± 0.7^{b} 46.9 ± 0.8^{c} 49.7 ± 0.9^{d} 10.7 ± 2.2^{a} 8.5 ± 1.6^{a} 23.0 ± 2.9^{b} 18.4 ± 3.6^{b} 18.8 ± 3.3^{b} 14.1 ± 5.0^{a} 154 ± 36^{b} 19.2 ± 4.7^{a} 11.0 ± 3.1^{a} 19.9 ± 4.3^{a} 34.3 ± 0.6^{ab} 39.9 ± 1.3^{b} 82 ± 11^{c} 29.4 ± 2.0^{ab} 33.1 ± 1.8^{ab}	41.6 ± 0.6^{a} 41.0 ± 0.6^{a} 34.6 ± 0.7^{b} 46.9 ± 0.8^{c} 49.7 ± 0.9^{d} 47.4 ± 0.9^{c} 10.7 ± 2.2^{a} 8.5 ± 1.6^{a} 23.0 ± 2.9^{b} 18.4 ± 3.6^{b} 18.8 ± 3.3^{b} 13.2 ± 2.2^{a} 14.1 ± 5.0^{a} 154 ± 36^{b} 19.2 ± 4.7^{a} 11.0 ± 3.1^{a} 19.9 ± 4.3^{a} 68 ± 6^{c} 34.3 ± 0.6^{ab} 39.9 ± 1.3^{b} 82 ± 11^{c} 29.4 ± 2.0^{ab} 33.1 ± 1.8^{ab} 17.5 ± 1.6^{a}

Different letters (a, b, c) in the same row indicate significant differences ($p \le 0.05$).

Tg = Glass transition temperature

WVP = Water vapour permeability

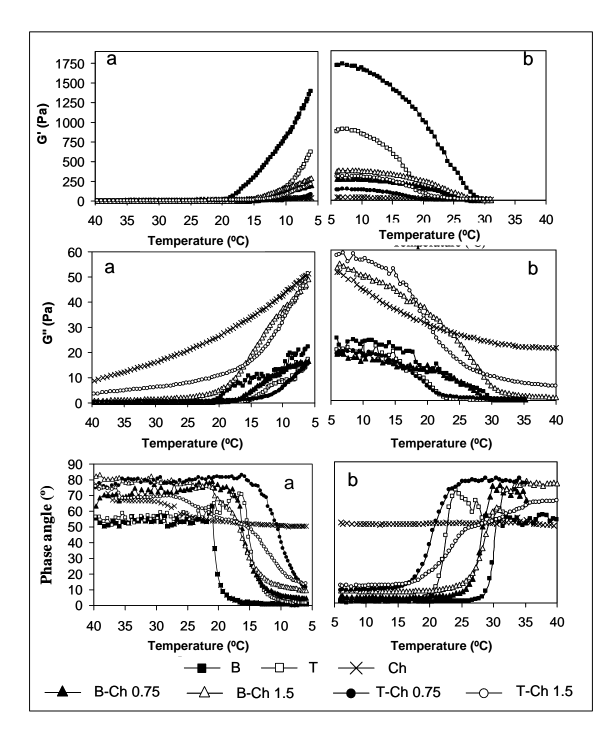


Figure 1

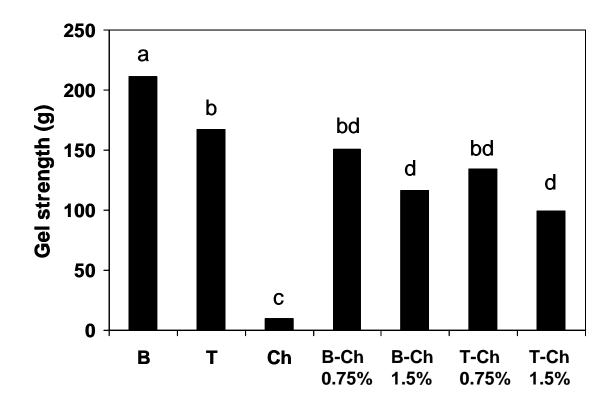


Figure 2

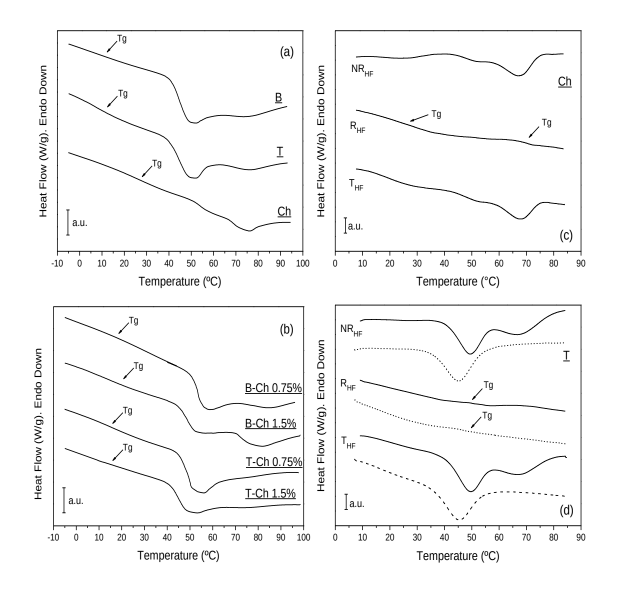


Figure 3

