

1 **Physico-chemical properties of tuna-skin and bovine-hide gelatin films**
2 **with added aqueous oregano and rosemary extract**

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11 **ABSTRACT**

12 The properties of edible, gelatin-based films with added oregano or rosemary
13 extract (two different concentrations) were studied. Gelatins from different
14 sources (bovine hides and tuna skins) were employed with a view to elucidating
15 how inherent gelatin characteristics may affect interaction of the gelatin with the
16 polyphenols in the added extract and hence the properties of the resulting films.
17 The bovine-hide gelatin reacted only slightly with the polyphenols in the extracts
18 as shown by the electrophoretic profile and analysis of the dynamic viscoelastic
19 properties, and consequently the attributes (mechanical properties, water
20 solubility, water vapour permeability) of the films were practically unchanged
21 compared with the film made without any added plant extract. The tuna-skin
22 gelatin did evidence some interactions with the polyphenols in both the oregano
23 and the rosemary extracts, especially for the more concentrated of the two
24 extracts tested, thereby altering the attributes of the corresponding films,
25 namely, a higher glass transition temperature, decreased deformability, and, in

26 particular, increased water solubility. Opacity increased irrespective of gelatin
27 origin and plant extract type and concentration.

28

29 **Key words:** bovine-hide gelatin, tuna-skin gelatin, polyphenols, films, physico-
30 chemical properties

31

32 **INTRODUCTION**

33 Considerable efforts are being expended lately to develop new biodegradable
34 packaging materials from natural polymers because of environmental concerns
35 relating to synthetic plastic packaging waste. Gelatin is a protein with a broad
36 range of functional properties and applications, including film-forming ability,
37 and is obtained by hydrolyzing collagen. Collagen is composed of three α -
38 chains (two α_1 and one α_2) intertwined in the collagen triple helix (Gómez-
39 Guillén, Turnay, Fernández-Díaz, Ulmo, Lizarbe, & Montero, 2002). Extraction
40 yields a pool of α -chains (both α_1 and α_2), β components (two covalently linked
41 α -chains), and γ components (three covalently linked α -chains), along with high-
42 molecular-weight aggregates (different covalently linked α -chains, β
43 components, and γ components) and peptide fractions with molecular weights
44 of <100 kDa. The properties and film-forming ability of gelatins are directly
45 related to the molecular weight, i.e., the higher the average molecular weight,
46 the better the quality of the gelatin. The molecular weight distribution depends
47 mainly on the degree of collagen cross-linking and the extraction procedure.
48 However, the physical properties of gelatins are related not only to the
49 molecular weight distribution but also to the amino acid composition.
50 Mammalian gelatins commonly have better physical properties and

51 thermostability than most fish gelatins (Ledward, 1986), and this has been
52 related mainly to their higher imino acid content (Norland, 1990), which
53 promotes refolding into the triple-helix configuration at low temperature (Gómez-
54 Guillén et al., 2002). Various studies have dealt with the physical and chemical
55 properties of mammalian gelatin films (Menegalli, Sobral, Roques, & Laurent,
56 1999; Sobral, Menegalli, Hubinger, & Roques, 2001; Simon-Lukasik &
57 Ludescher, 2004; Bertan, Tanada-Palmu, Siani, & Grosso, 2005). Recently
58 attention has focused on the properties of films made from fish gelatins
59 (Muyonga, Cole, & Duodu, 2004; Thomazine, Carvalho, & Sobral, 2005; Avena-
60 Bustillos et al., 2006; Jongjareonrak, Benjakul, Visessanguan & Tanaka, 2006a,
61 2006b; Gómez-Guillén, Ihl, Bifani, Silva, & Montero, 2007; Pérez-Mateos,
62 Montero, & Gómez-Guillén, 2008; Carvalho et al., 2008).

63

64 Film coatings improve storage, mainly by acting as barriers to water, oxygen,
65 and light (Gennadios, Hanna, & Kurth, 1997). Furthermore, active substances,
66 e.g., plant extracts, have also been included in formulations of edible films to
67 afford enhanced antioxidant and/or antimicrobial properties (Zivanovic, Chi, &
68 Draughon, 2005; Kim et al., 2006; Seydim & Sarikus, 2006; Gómez-Guillén et
69 al., 2007) and thus improve the quality and stability of foods during storage
70 (Oussalah, Caillet, Salmiéri, Saucier, & Lacroix, 2004; Gómez-Estaca, Montero,
71 Giménez, & Gómez-Guillén, 2007). However, adding such extracts may alter
72 the mechanical and barrier properties of the gelatin films (Gómez-Guillén et al.,
73 2007) and affect the release of active components (Gómez-Estaca, Bravo,
74 Gómez-Guillén, Alemán, & Montero, 2009) as a consequence of polyphenol-
75 protein interactions, but little information is available in this regard to date.

76 Because of its intrinsic open structure, gelatin is generally agreed to be more
77 prone to interact with polyphenols than globular proteins (Frazier,
78 Papadopoulou, Mueller-Harvey, Kisson, & Green, 2003; Naczk, Grant,
79 Zadernowski, & Barre, 2006). Nevertheless, exactly how the intrinsic
80 characteristics of gelatins from different sources may act on these interactions
81 is unknown. Frazier et al. (2003) used isothermal titration microcalorimetry to
82 investigate protein-tannin interactions and found that tannins bound to gelatin
83 by a two-stage mechanism, namely, a first stage of cooperative binding of
84 tannins to the protein, followed by a second stage of gradual saturation of
85 binding sites. Naczk et al. (2006) studied the protein precipitating capacity of
86 phenolics from wild blueberry leaves and fruits and found that the tannin-rich
87 fractions of the extracts were more effective gelatin precipitants than the
88 fractions comprising low-molecular-weight compounds (monomers, dimers, and
89 trimers) only.

90

91 The object of this experiment was therefore to evaluate changes in the physical
92 properties of gelatin films on adding aqueous extracts of oregano and rosemary,
93 taking into consideration differences related to gelatin source, i.e., either tuna
94 skins or bovine hides.

95

96 **MATERIALS AND METHODS**

97

98 **Preparation of the antioxidant extracts**

99 Freeze-dried oregano (*Origanum vulgare*) and rosemary (*Rosmarinus*
100 *officinalis*) leaves were purchased at a local market. Quantities of 5 and 20 g,

101 respectively, were mixed with 100 mL of distilled water that had been pre-
102 warmed to 45 °C, the mixture was then extracted by continuous stirring in a
103 warm water bath at 45 °C for 10 min. The aqueous extract obtained was filtered
104 through Whatman no. 1 filter paper. The total phenolic content of the aqueous
105 extracts as determined by the Folin-Ciocalteu method according to Montreau
106 (1972) was $2\,080 \pm 23$ µg of caffeic acid/mL for the oregano extract and $665 \pm$
107 11 µg of caffeic acid/mL for the rosemary extract.

108

109 **Formulation of the film-forming solutions (FFSs) and film formation**

110 The FFSs were prepared using gelatin made from tuna skins [obtained
111 according to the method described by Gómez-Guillén & Montero (2001)] or
112 bovine hides (Bloom 200/220 from Sancho de Borja S.L., Saragossa, Spain) at
113 a concentration of 4 g/100 mL of distilled water. Based on the results of
114 Thomazine et al. (2005), a mixture of sorbitol (0.15 g/g gelatin) and glycerol
115 (0.15 g/g gelatin) was employed as plasticizer. The oregano (OE) and rosemary
116 (RE) extracts were added to the FFSs in the proportion of 6.25 mL OE/100 mL
117 FFS (batch O-L, theoretical phenol content of 130 µg caffeic acid/mL FFS);
118 25 mL OE/100 mL FFS (batch O-H, theoretical phenol content of 520 µg caffeic
119 acid/mL FFS); 12.5 mL RE/100 mL FFS (batch R-L, theoretical phenol content
120 of 83 µg caffeic acid/mL FFS); and 100 mL RE/100 mL FFS (batch R-H,
121 theoretical phenol content of 665 µg caffeic acid/mL FFS). Distilled water was
122 employed to prepare the extract/FFS dilutions, except for batch R-H, which was
123 not diluted. For comparative purposes an FFS with no added extract was also
124 made up (batch C). All mixtures were warmed and stirred at 40 °C for 15 min to
125 obtain a good blend, and the films were prepared by casting an amount of 40 ml

126 on 12 cm x 12 cm-square plates and drying in a forced-air oven at 45 °C for
127 15 h to obtain a uniform thickness (100 µm; $p < 0.05$) in all cases. Before
128 performing the determinations, the films were conditioned in desiccators over a
129 saturated solution of NaBr at 22 °C for 2 days.

130

131 **Electrophoretic profile of the gelatins dissolved in the aqueous extracts**

132 The molecular weight distributions of the solutions of both the bovine-hide and
133 tuna-skin gelatins with the oregano and rosemary extracts at the two
134 concentrations employed were determined by polyacrylamide gel
135 electrophoresis in the presence of SDS, and the SDS-PAGE profiles of were
136 examined to assess the interactions among the aqueous oregano and rosemary
137 extract components and the gelatins. The gelatin solutions were mixed with
138 loading buffer (2 % SDS, 5 % mercaptoethanol, and 0.002 % bromophenol
139 blue) in a proportion of 1:4, resulting in a final protein concentration of 5 mg/mL.
140 Samples were heat-denatured at 90 °C for 5 min and analysed according to
141 Laemmli (1970) using 4% stacking gels and 6% resolving gels in a Mini Protean
142 II unit (Bio-Rad Laboratories, Hercules, CA, USA) at 25 mA/gel. Loading volume
143 was 15 µL in all lanes. Protein bands were stained with Coomassie brilliant Blue
144 R250. Type I collagen from fetal calf was used as a marker for α -chain and β -
145 component mobilities. A molecular weight standard composed of myosin
146 (212 kDa), α_2 -macroglobulin (170 kDa), β -galactosidase (116 kDa), transferrin
147 (76 kDa) and glutamic dehydrogenase (53 kDa) (Amersham Pharmacia
148 Biotech, Buckinghamshire, UK) was also employed.

149

150 **FFS analysis**

151 *Dynamic viscoelastic properties*

152 Dynamic viscoelastic analysis of the film-forming solutions was carried out using
153 a Bohlin CSR-10 rheometer rotary viscometer (Bohlin Instruments Ltd.,
154 Gloucestershire, UK) using a cone-plate geometry (cone angle=4°, gap= 0.15
155 mm) as described in Gómez-Guillén et al. (2007). Cooling and heating from 40
156 to 6 °C and back to 40 °C took place at a scan rate of 1 °C/min, a frequency of 1
157 Hz, and a target strain of 0.2 mm. The elastic modulus (G' ; Pa), viscous
158 modulus (G'' ; Pa) and phase angle (°) were determined as functions of
159 temperature. Two determinations were performed for each sample, with an
160 experimental error of less than 6% in all cases.

161

162 *Gel strength*

163 The film-forming solutions were poured into glasses 2.3 cm in diameter by 3.6
164 cm in height and left to mature in a refrigerator at 2 °C for 16-18 h. Gel strength
165 at 8-9 °C was determined on an Instron model 4501 Universal Testing Machine
166 (Instron Co., Canton, MA, USA) with a 100 N load cell, a cross-head speed of
167 60 mm/min, and a flat-faced cylindrical plunger 1.27 cm in diameter. The
168 maximum force (g) was determined when the plunger had penetrated 4 mm into
169 the gelatin gels.

170

171 **Film analysis**

172 *Mechanical properties*

173 A puncture test was performed to determine the breaking force and breaking
174 deformation of films. Films were placed in a cell 5.6 cm in diameter and
175 perforated to the breaking point using an Instron model 4501 Universal Testing

176 Machine (Instron Co., Canton, MA, USA) with a rounded stainless-steel plunger
177 3 mm in diameter at a cross-head speed of 60 mm/min and a 100-N load cell.
178 The puncture force was expressed in N and breaking deformation in percent, as
179 per Sobral et al. (2001). All determinations are the means of at least five
180 measurements.

181

182 *Thermal properties*

183 Calorimetric analysis was performed using a model TA-Q1000 differential
184 scanning calorimeter (DSC) [TA Instruments, New Castle, DE, USA] previously
185 calibrated by running high purity indium (melting point: 156.4 °C; enthalpy of
186 melting: 28.44 W/g). Samples of approximately 10 mg (± 0.002 mg) were
187 weighed out using a model ME235S electronic balance (Sartorius, Goettingen,
188 Germany) and were tightly encapsulated in aluminium pans and scanned under
189 dry nitrogen purge (50 mL/min). Freshly conditioned films were rapidly cooled to
190 0 °C and scanned at between 0 and 90 °C at a heating rate of 10 °C/min. Glass
191 transition temperatures, T_g (°C), were determined only on the first heating
192 scans, the values obtained on the second scans being deemed insufficiently
193 reliable because of the virtual impossibility, in practice, of reproducing the
194 original film conditioning. The glass transition temperature was estimated as the
195 midpoint of the line between the temperature at the intersection of the initial
196 tangent with the tangent through the inflection point of the trace and the
197 temperature of the intersection of the tangent through the inflection point with
198 the final tangent. T_g data have been reported as the mean values of at least
199 duplicate samples of each film, usually within ± 1 °C.

200

201 *Water solubility*

202 Film portions of 4 cm² were placed in aluminium capsules with 15 ml of distilled
203 water and shaken gently at 22 °C for 15 h. The solution was then filtered
204 through Whatman no. 1 filter paper to recover the remaining undissolved film,
205 which was desiccated at 105 °C for 24 h. Film solubility was calculated using
206 the equation $FS (\%) = [(W_o - W_f) / W_o] \cdot 100$, where W_o was the initial weight of the
207 film expressed as dry matter and W_f was the weight of the undissolved
208 desiccated film residue. All tests were carried out in triplicate.

209

210 *Water vapour permeability*

211 Water vapour permeability was determined according to the gravimetric method
212 described by Sobral et al. (2001). Films were attached over the openings of
213 cells (permeation area = 15.9 cm²) containing desiccated silica gel, and the
214 cells were placed in desiccators with distilled water at 22 °C. The cells were
215 weighed daily for 7 d. Water vapour permeability was calculated using the
216 equation $WVP = w \cdot x \cdot t^{-1} \cdot A^{-1} \cdot \Delta P^{-1}$, where w was weight gain (g), x film thickness
217 (mm), t elapsed time at weight gain (h), A permeation area, and ΔP partial
218 vapour pressure difference between the dry atmosphere and pure water
219 (2 642 Pa at 22 °C). Results have been expressed as g·mm·h⁻¹·cm⁻²·Pa⁻¹. All
220 tests were carried out in duplicate.

221

222 *Opacity*

223 Film portions of 0.8 cm x 4 cm were placed in a spectrophotometer test cell, and
224 absorbance was measured at 600 nm using a UV-1601 spectrophotometer
225 (Model CPS-240, Shimadzu, Kyoto, Japan). The opacity index (O) was

226 calculated as the quotient of the absorbance value at 600 nm divided by film
227 thickness in mm.

228

229 **Statistical analysis**

230 Statistical tests were performed using the SPSS® computer program (SPSS
231 Statistical Software, Inc., Chicago, IL, USA). One-way and two-way analyses of
232 variance were carried out. Differences between pairs of means were resolved by
233 means of confidence intervals using a Tukey-b test. The level of significance was
234 set for $p < 0.05$.

235

236 **RESULTS AND DISCUSION**

237 *Electrophoretic profile of the gelatins dissolved in the aqueous extracts*

238 The molecular weight distributions of the blends of the bovine-hide and tuna-
239 skin gelatins with the aqueous oregano and rosemary extracts at the different
240 concentrations studied are presented in Figure 1. The presence of aqueous
241 oregano and rosemary extracts in the bovine-hide gelatin solution had hardly
242 any effect on the electrophoretic profiles as compared to the control gelatin
243 solution (batch C), irrespective of extract concentration. In contrast, the tuna-
244 skin gelatin with both concentrations of added oregano extract registered
245 appreciable decreases in β -components and high-molecular-weight aggregates
246 (HMW-a) and a certain increase of polypeptides with molecular weights below
247 100 kDa. The increase in hydrolyzed fractions was more intense in the batch
248 with the more concentrated oregano extract (batch O-H). These results
249 suggested a high degree of interaction between the phenolic substances in the
250 oregano extract and the fish-gelatin polypeptides, giving rise to cleavage or

251 disruption of the covalently associated α -chains (β -components and HMW-a).
252 Adding rosemary extract to the tuna-skin gelatin also brought about a decrease
253 in the amount of HMW-a present, but in this case without any appreciable rise in
254 the hydrolyzed low-molecular-weight fractions, although the new protein
255 fractions could be located in the >200 kDa region. This difference with respect
256 to the samples with the added oregano extract could be attributed to qualitative
257 differences in the two extracts as previously observed using confocal laser
258 scanning microscopy and HPLC analysis (Gómez-Estaca et al., 2009). In that
259 study, rosmarinic acid was found to be the most abundant compound in both
260 extracts, but the oregano extract also contained gallic acid and protocatechuic
261 acid while the rosemary extract contained chlorogenic acid. There were also
262 appreciable amounts of a series of other compounds that could not be
263 identified, though based on their absorption spectra they may have been
264 hydroxybenzoic acid derivatives, caffeic acid derivatives, and various flavonoids
265 (primarily flavone derivatives). The stronger polyphenol-protein interaction
266 observed for the tuna-skin gelatin compared with the bovine-hide gelatin can be
267 ascribed to differences in gelatin characteristics. According to Frazier et al.
268 (2003) and Naczki et al. (2006), proteins with an open structure are more prone
269 to interact with polyphenols. Although gelatin is a clear example of a protein
270 with an open structure subsequent to collagen denaturation, there are certain
271 differences that could affect the degree of interaction depending on gelatin
272 origin. Mammalian gelatins are well known to have higher imino acid (Pro+Hyp)
273 contents than fish gelatins (Norland, 1990; Avena-Bustillos et al., 2006), which
274 is conducive to intra and interchain interactions. In addition, the collagenous
275 material in fish skins has low levels of intra and interchain covalent cross-

276 linking, mainly involving lysine and hydroxylysine residues and aldehyde
277 derivatives (Montero, Borderías, Turnay, & Leyzarbe, 1990). Thus, the
278 presumably higher protein-protein interactions in the bovine-hide gelatin may
279 interfere with the ability of the polyphenols to interact with the protein chains
280 compared with the tuna-skin gelatin.

281

282 *FFS analysis*

283 Possible interactions between the extract components and the gelatin
284 molecules were also investigated by following the changes in the dynamic
285 viscoelastic properties of the different FFSs upon cooling and subsequent
286 heating (Figure 2 for bovine-hide FFS, Figure 3 for tuna-skin FFS). Addition of
287 the antioxidant extracts did not substantially alter the G' curves or the thermal
288 transition points for either gelatin type, indicative of minor interference with
289 triple helical structure formation. However, the viscous modulus (G'') value
290 increased in tuna-skin gelatin batches O-H and R-H, suggesting the presence
291 of certain peptide fractions that did not participate directly in the protein gel
292 matrix upon cold renaturation, probably because of interactions with the
293 phenolic compounds. This would be consistent with the electrophoretic profiles
294 of the phenolic-containing fish-gelatin samples, which exhibited an
295 accumulation of hydrolyzed peptide fractions resulting from cleavage of
296 covalently linked peptide chains. Similarly, interference with gelatin polypeptide
297 chains by polyphenolics upon cold renaturation and subsequent melting was
298 also reported in tuna-fish gelatin film-forming solutions with added *murta*
299 extracts (Gómez-Guillén et al., 2007). In that study the increase in the G''
300 values was also proportional to the amount of polyphenolics added to the FFSs.

301

302 The gel strength of the FFSs was also determined (Figure 4) to shed further
303 light on the possible interactions or interference between the polyphenolic
304 compounds and the protein matrix during cold maturation of the gels, during
305 which triple helix growth takes place by association of different α -chains via
306 hydrogen bonding. The addition of aqueous oregano or rosemary extract
307 brought about an increase in gel strength irrespective of gelatin origin, with
308 minor variations depending on extract type and concentration in the FFSs. This
309 gel strengthening effect could be attributable to non-covalent polyphenol-protein
310 interactions during cold maturation of the gelatins overnight, which may be
311 conducive to the formation of side-by-side associations of gelatin chains without
312 disturbing triple helix formation and growth. According to Oh, Hoff, Armstrong &
313 Haff (1980) and Haslam (1996), the polyphenol-protein interaction initially
314 results from hydrophobic interactions and is subsequently augmented by the
315 possible formation of hydrogen bonds between the polyphenol –OH groups and
316 the protein –COOH groups. Frazier et al. (2003) found that multidentate tannin
317 ligands formed intermolecular cross-links between binding sites on adjacent
318 gelatin molecules, contributing to protein precipitation. In this regard, Naczki et
319 al. (2006) reported that gelatin precipitation due to phenolic compounds from
320 blueberry leaves and fruits was due mainly to the condensed tannin fraction.
321 The aqueous oregano and rosemary extracts employed in the present
322 experiment did not contain these high-molecular-weight polyphenol complexes
323 (Gómez-Estaca et al., 2009), and therefore no protein precipitation was
324 apparent. However, the low-molecular-weight polyphenols present exerted a

325 strengthening effect, most probably by promoting interactions among adjacent
326 gelatin chains.

327

328 *Film analysis*

329 The breaking force and breaking deformation values for the resulting films are
330 set out in Table 1. Breaking force values for the control films were similar
331 ($p > 0.05$), but addition of the polyphenolic extracts resulted in differences
332 according to gelatin type. Still, the added plant extracts did not significantly
333 ($p > 0.05$) alter the breaking force for any of the gelatin admixtures compared to
334 the control batches. Similarly, no differences were observed for breaking
335 deformation on adding the plant extracts, except for the tuna-skin gelatin film
336 with the more concentrated rosemary extract (batch R-H), which had a
337 significantly lower breaking deformation value. Other workers have reported
338 lower breaking strength and breaking deformation on adding antioxidant
339 extracts to films made either from soya or from gelatin (Kim et al., 2006;
340 Gómez-Guillén et al., 2007). This behaviour was explained by Orliac, Rouilly,
341 Silvestre, & Rigal (2002) by a weakening of the interactions that stabilize the
342 protein matrix on adding polyphenolic antioxidants, especially higher-molecular-
343 weight polyphenols. The general absence of significant differences in the
344 mechanical property values of the films studied here when the plant extracts
345 were added, in contrast to other published work, is largely ascribed to the
346 characteristics of the phenolic composition of the aqueous oregano and
347 rosemary extracts, which had a predominance of low-molecular-weight
348 compounds (Gómez-Estaca et al., 2009).

349

350 Figure 5 plots the DSC traces. Despite encapsulation in non-hermetically sealed
351 pans, samples shrank only by ~1-2% by weight after DSC scanning, indicating
352 good water retaining capacity of the films. However, this produced a certain
353 bending in the trace from water vaporization, which distorted the DSC profiles
354 somewhat at the glass transition exit. Additionally, devitrification overlapped
355 completely with a relaxation effect that introduced considerable uncertainty into
356 the Tg determinations. The films did not exhibit any melting event attributable to
357 crystallization of the gelatins. The glycerol plus sorbitol (Sobral et al., 2001) and
358 the low water contents used yielded wholly amorphous films. It is generally
359 accepted that, in addition to their plasticizing effects, polyols present at low
360 moisture levels may inhibit crystalline structures by constraining molecular
361 mobility (Cheng, Karim, & Seow, 2006). The tuna-skin gelatin batches exhibited
362 Tg values slightly lower than but similar to the corresponding bovine-hide
363 gelatin batches (Table 2). Adding the herb extracts raised the glass transition
364 temperatures in both the bovine-hide and tuna-skin gelatin films. The increase
365 was relatively higher as the amount of added extract rose, with the bovine-hide
366 gelatin being slightly more sensitive than the tuna-skin gelatin. Furthermore, the
367 rosemary extract seemed to bring about a larger increase in the Tg than the
368 oregano extract, despite the similar phenolic contents. However, compared with
369 the respective tuna-skin gelatin batches, the bovine-hide gelatin films displayed
370 a saturation effect with rosemary extract concentration, in that batch R-H
371 resulted in a considerably lower increase than would at first be expected based
372 on the previous effect of batch R-L. Thus, the binding sites for polyphenols in
373 the bovine-hide gelatin appeared to be saturated, and the rosemary extract

374 appeared to be incorporated into the bovine-film matrix partly in the manner of a
375 filler.

376

377 Addition of the antioxidant extracts to the tuna-skin gelatin brought about a
378 pronounced increase in film solubility (Table 3) in the case of batches O-H and
379 R-H, as well as batch O-L. Electrophoretic analysis indicated that the higher
380 solubility could be due to the cleavage or degradation of covalently linked α -
381 chains (β -components) or even HMW-a induced by the phenolic compounds
382 present. This effect observed in the electrophoretic profile was more apparent in
383 the films containing the more concentrated oregano extract, but adding the
384 more concentrated rosemary extract also exerted an influence, reducing the
385 quantities of HMW-a present to some extent. The differences in water solubility
386 observed between batches R-L and O-L could be due to qualitative differences
387 in the polyphenolic composition, as indicated in a previous paper (Gómez-
388 Estaca et al., 2009). Kim et al. (2006) also reported increased film solubility on
389 adding green tea extract to soya-protein films. However, the presence of the
390 phenolic extracts did not produce any significant differences in the solubility of
391 the bovine-hide gelatin films. This finding was in consonance with the
392 corresponding electrophoretic profiles, which likewise were barely affected by
393 the different blends.

394

395 Table 3 also sets out the water vapour permeability (WVP) of the films. All the
396 tuna-skin gelatin films had significantly lower WVP values than the bovine-hide
397 gelatin films. According to Avena-Bustillos et al. (2006), this could be related to
398 higher levels of hydrophobicity resulting from the lower amounts of proline and

399 hydroxyproline present in gelatins from cold-water fish species as compared to
400 mammalian gelatins. Adding the plant extracts did not significantly ($p > 0.05$)
401 alter the water vapour permeability in either the bovine-hide or the tuna-skin
402 gelatin films. Gómez-Guillén et al. (2007) recorded a decrease in the WVP on
403 adding murta extract to tuna-fish gelatin films when the extract from the murta
404 ecotype with the higher phenolic content was added.

405

406 To the unaided eye all the films were quite transparent, though tinted by the
407 natural colour of the oregano and rosemary extracts. Adding the plant extracts
408 increased the O values of both types of gelatin, especially for the more highly
409 concentrated oregano extract (Table 4). The higher film opacity on adding plant
410 extracts is directly ascribed to enrichment of the films with polyphenols and to
411 some extent probably also to polyphenol-protein interactions. Similar results
412 were reported for tuna-fish gelatin films with added murta extract (Gómez-
413 Guillén et al., 2007).

414

415 **CONCLUSIONS**

416 The physical and chemical properties of the bovine-hide and tuna-skin gelatins
417 determine their reactivity with the polyphenols in added plant extracts and
418 hence the properties of the corresponding composite films. The bovine-hide
419 gelatin thus did not react with the polyphenols to the same extent as the tuna-
420 skin gelatin. In consequence, the properties of bovine-hide gelatin films with
421 added oregano or rosemary extract were similar to the properties of the control
422 film with no added extract. In contrast, the tuna-skin gelatin interacted more
423 readily with the polyphenols, appreciably increasing the water solubility of the

424 films. Opacity increased irrespective of gelatin origin and added plant extract
425 type and concentration.

426

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558 **Figure captions**

559 Figure 1. Electrophoretic profiles of the bovine-hide and the tuna-skin gelatins
560 with added oregano or rosemary extract. C: no added extract (control batch); O-
561 L: a low concentration of added oregano extract; O-H: a high concentration of
562 added oregano extract; R-L: a low concentration of added rosemary extract; R-
563 H: a high concentration of added rosemary extract. HMW-a: high-molecular-
564 weight aggregates.

565

566 Figure 2. Dynamic viscoelastic properties of the FFSs prepared from bovine-
567 hide gelatin with added oregano or rosemary extract (batch designations as in
568 Figure 1) during cooling (a) and subsequent heating (b) ramps.

569

570 Figure 3. Dynamic viscoelastic properties of FFSs prepared from tuna-skin
571 gelatin with added oregano or rosemary extract (batch designations as in Figure
572 1) during cooling (a) and subsequent heating (b) ramps.

573

574 Figure 4. Gel strength of FFSs (batch designations as in Figure 1) after cold
575 maturation. Different letters (a, b, c) indicate significant differences between the
576 formulations prepared using the different gelatin types. Different letters (x, y)
577 indicate significant differences between the two gelatin types for each
578 formulation.

579

580 Figure 5. DSC traces for the bovine-hide (a) and tuna-skin (b) gelatin films
581 (batch designations as in Figure 1) at the Tg point.

582

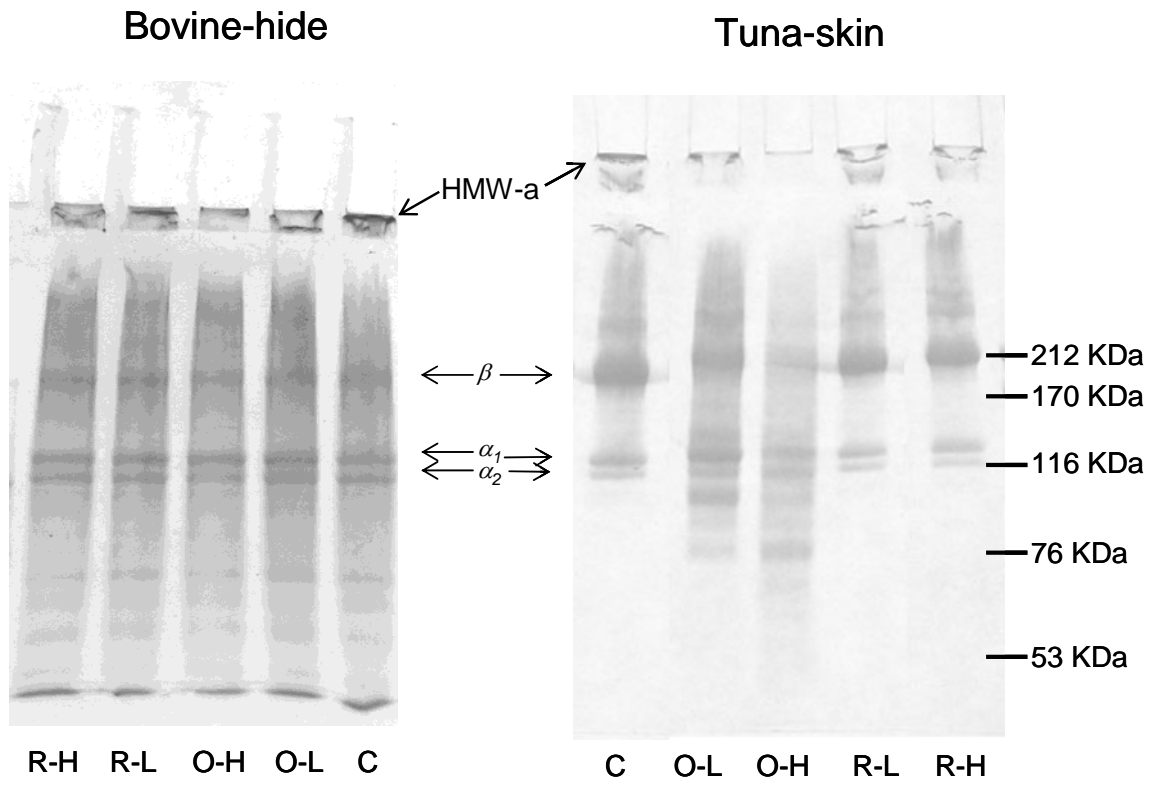


Figure 1

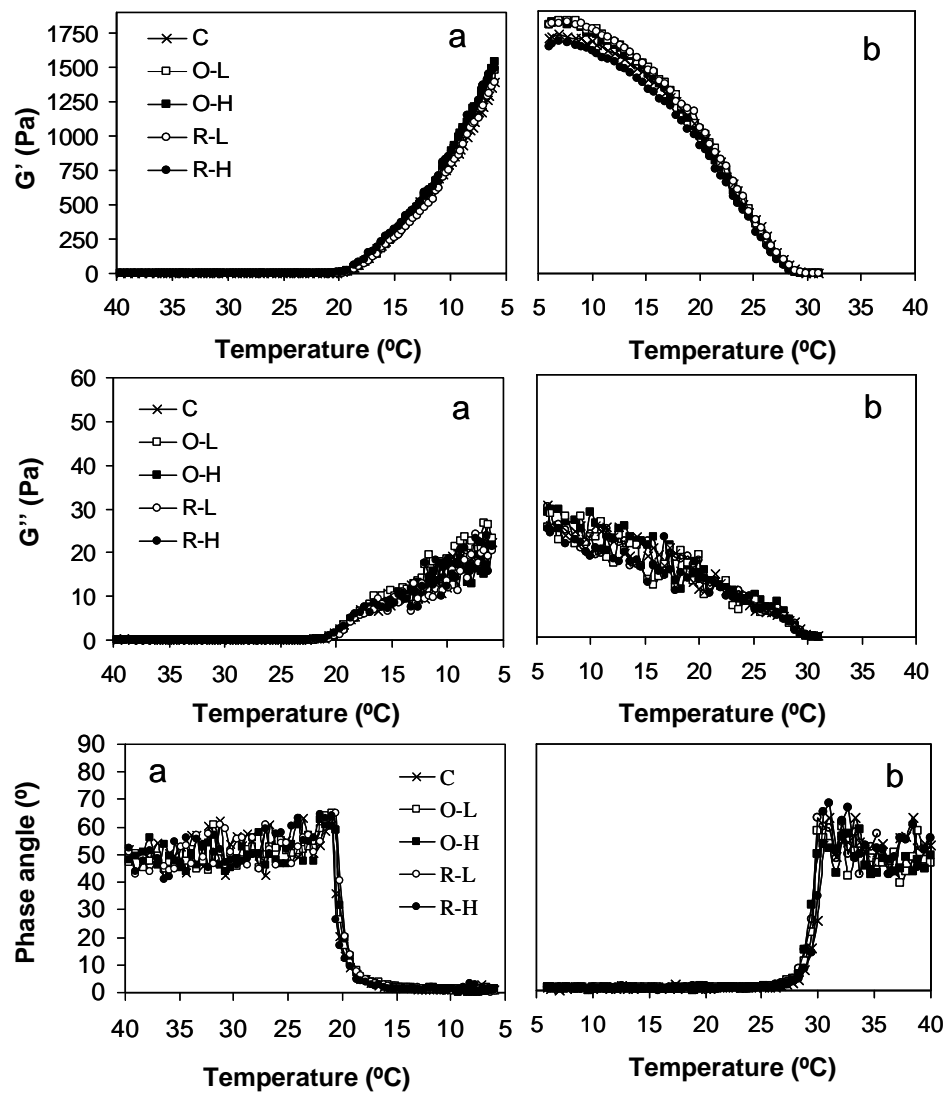


Figure 2

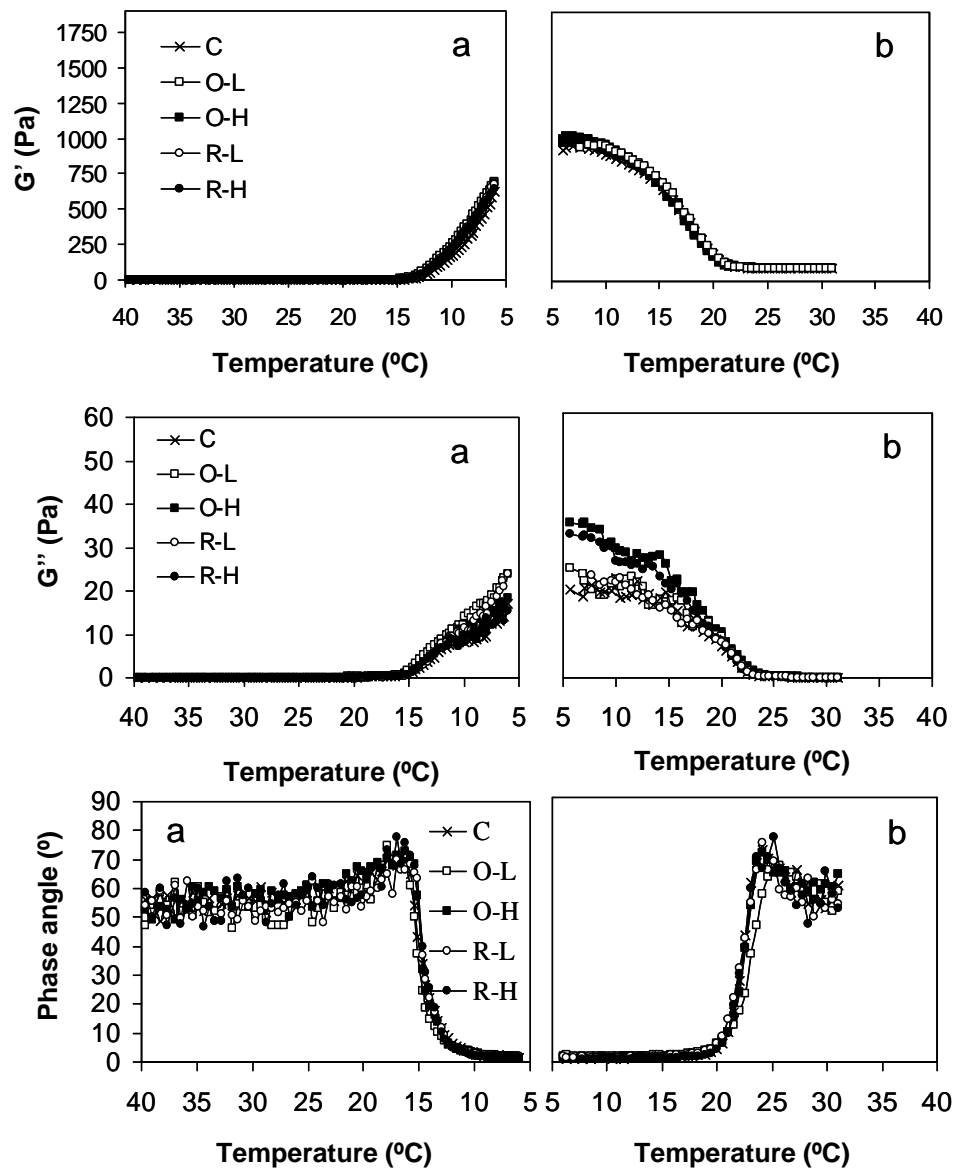


Figure 3

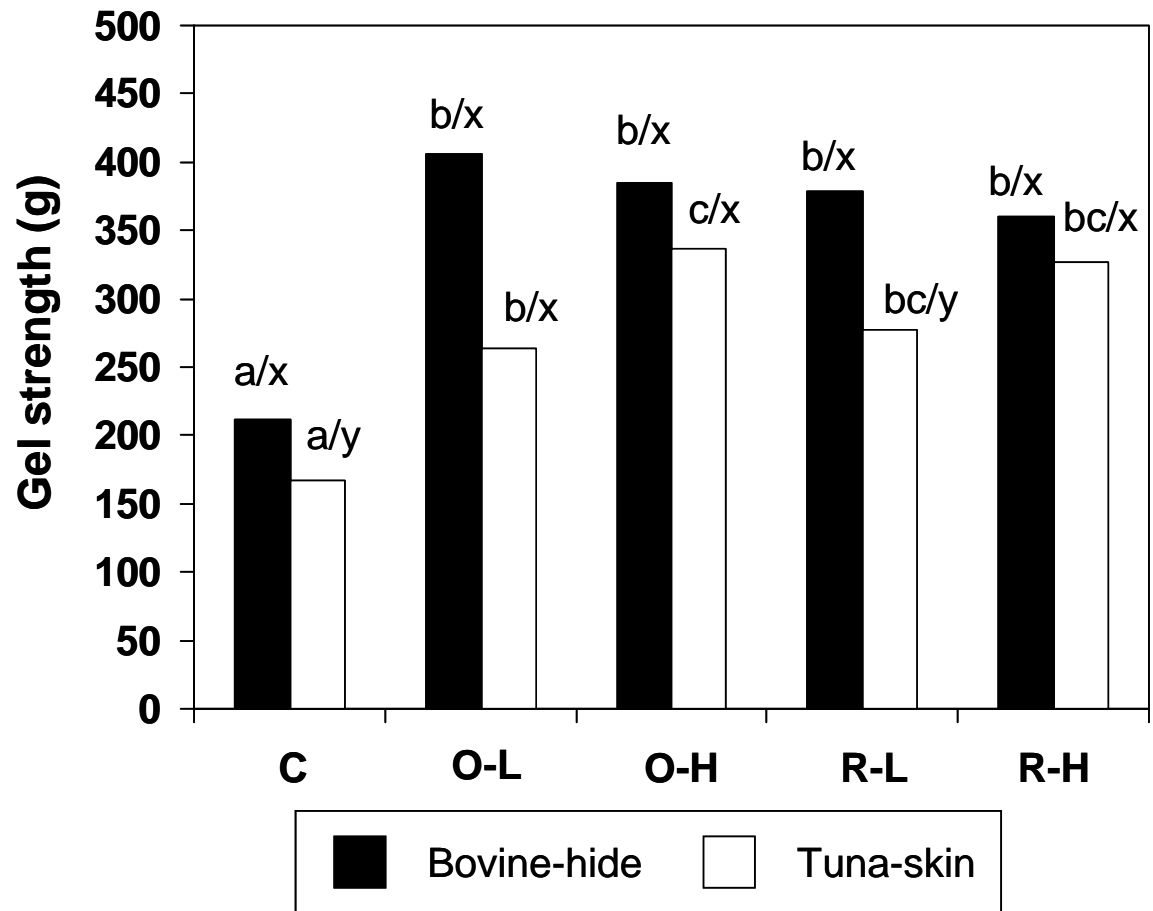


Figure 4

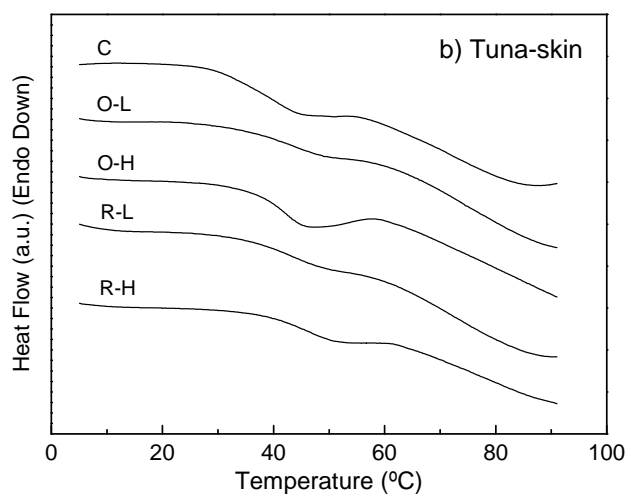
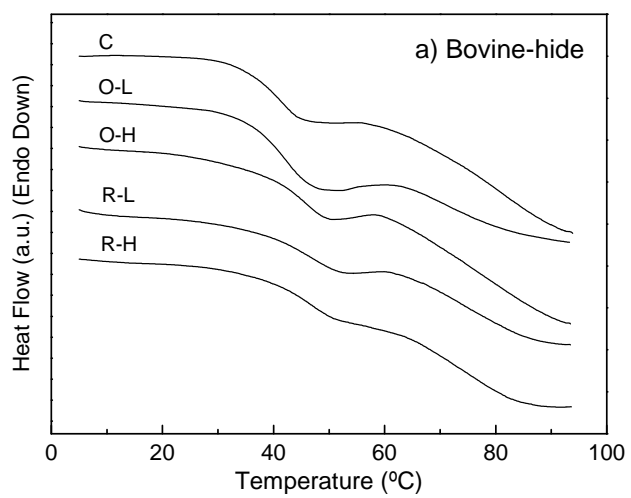


Figure 5

Table 1. Mechanical properties of the films (batch designations as in Figure 1). Results have been expressed as the mean value \pm standard deviation. Different letters (a, b, c) within the columns indicate significant differences between formulations with/without extract. Different letters (x, y) within the rows indicate significant differences between gelatine types.

Batch	Puncture force (N)		Breaking deformation (%)	
	Bovine-hide	Tuna-skin	Bovine-hide	Tuna-skin
C	10.7 \pm 2.2 ax	8.5 \pm 1.6 ax	14.1 \pm 5.0 ax	154 \pm 35 cy
O-L	10.2 \pm 1.3 ax	5.2 \pm 1.7 ay	14.1 \pm 4.7 ax	116 \pm 26 cy
O-H	8.8 \pm 0.8 ax	6.1 \pm 1.0 ay	19.4 \pm 4.2 ax	132 \pm 14 bcy
R-L	9.9 \pm 1.3 ax	6.2 \pm 1.3 ax	14.9 \pm 6.9 ax	147 \pm 38 cy
R-H	12.4 \pm 0.7 ax	5.6 \pm 1.0 ay	11.6 \pm 0.9 ax	87 \pm 9 aby

Table 2. Tg values (°C) of the films (batch designations as in Figure 1). Results are the mean values of at least two samples of each film, usually within ± 1 °C.

Batch	Tg (°C)	
	Bovine-hide	Tuna-skin
C	41.5	40.7
O-L	42.6	41.2
O-H	44.3	42.5
R-L	45.3	42.9
R-H	47.4	47.0

Table 3. Water solubility and water vapour permeability of the films (batch designations as in Figure 1). Results have been expressed as the mean value \pm standard deviation. Different letters (a, b, c) within the columns indicate significant differences between formulations with/without extract. Different letters (x, y) within the rows indicate significant differences between gelatine types.

Batch	Water solubility (%)		Water vapour permeability ($10^{-8} \cdot \text{g} \cdot \text{mm} \cdot \text{h}^{-1} \cdot \text{cm}^{-2} \cdot \text{Pa}^{-1}$)	
	Bovine-hide	Tuna-skin	Bovine-hide	Tuna-skin
C	34.3 \pm 0.6 ax	39.9 \pm 1.3 ax	2.20 \pm 0.11 bcx	1.65 \pm 0.39 ay
O-L	35.7 \pm 1.5 ax	76.6 \pm 0.1 by	2.13 \pm 0.08 abcx	1.59 \pm 0.08 ay
O-H	38.6 \pm 1.4 ax	83.7 \pm 3.6 by	2.40 \pm 0.03 cx	1.35 \pm 0.06 ay
R-L	34.8 \pm 1.6 ax	36.6 \pm 0.6 ax	2.17 \pm 0.08 bcx	1.30 \pm 0.59 ay
R-H	34.4 \pm 0.7 ax	82.6 \pm 8.1 by	1.89 \pm 0.07 abx	1.42 \pm 0.26 ay

Table 4. Opacity index values for the films (batch designations as in Figure 1).

Batch	Opacity index	
	Bovine-hide	Tuna-skin
C	0.461 ± 0.001	0.377 ± 0.000
O-L	0.542 ± 0.000	0.546 ± 0.001
O-H	0.725 ± 0.002	0.655 ± 0.003
R-L	0.530 ± 0.001	0.411 ± 0.001
R-H	0.684 ± 0.002	0.557 ± 0.003