

1 **TITLE: Effect of alkalis on konjac glucomannan gels for use as potential gelling**
2 **agents in restructured seafood products**

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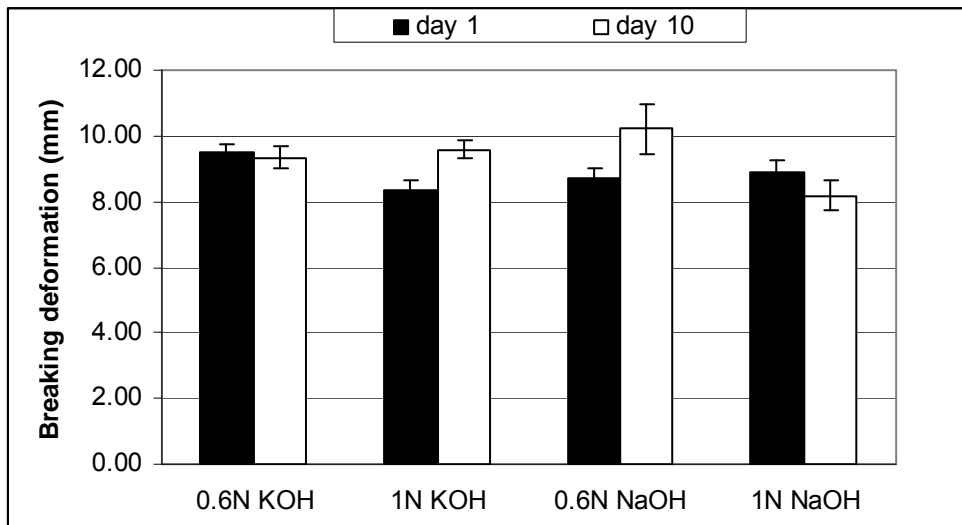
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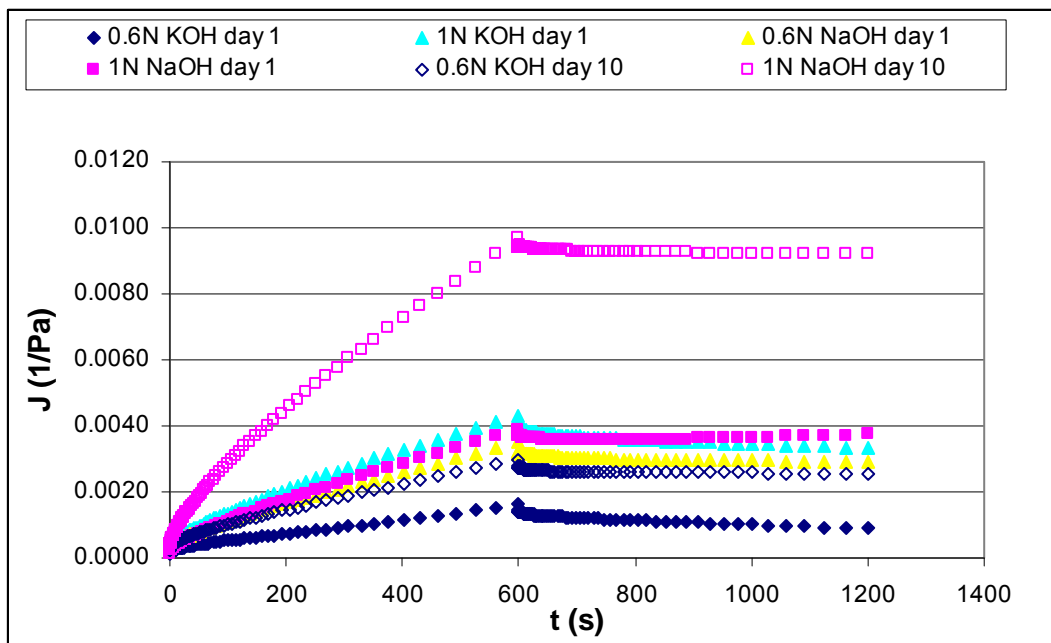
13 **Graphical abstract**

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17 Breaking deformation in gels deacetylated with different alkalis at 0.6N and 1N after 1 and 10
18 days of refrigeration.

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23 Influence of chilled storage time at 5 °C on creep and recovery compliance data of
24 glucomannan gels deacetylated with KOH 0.6N and NaOH 1N.

25 **Abstract**

26

27 Four dispersions of 3% glucomannan in water, deacetylated with 5% 0.6N and 1N
28 KOH (lots L1 and L2) and 0.6N and 1N NaOH (lots L3 and L4) as gelling agents, were
29 evaluated for use in raw restructured seafood products. Several properties (pH,
30 moisture content, water binding capacity, cooking loss and lightness) together with
31 puncture data (breaking force and breaking deformation) were determined after 1 and
32 10 days of chilled storage at 5°C. All these data were analysed together with different
33 viscoelastic parameters obtained at small amplitude oscillatory strain (SAOS) after 1
34 day of chilled storage, showing that L1 and L4 samples were the most suitable gels for
35 incorporation in raw restructured fish products. In both cases the highest stress (σ_{max})
36 and strain (γ_{max}) amplitude values were found in the linear viscoelastic (LVE) range;
37 however, L1 showed both high strain amplitude and breaking deformation values.
38 Moreover, creep and recovery (transient) data showed that L1 was the most-time
39 stable gel with the highest elasticity and the lowest relaxation exponent (n). L4 gel
40 showed strong rigidity, i.e. the highest values of breaking force and storage moduli (G')
41 and the highest n value, making it less gel-like. Both L1 and L4 gels became
42 significantly less gel-like over 10 days of chilled storage due to the loss of gel strength
43 (S) and a noticeable increase of n . These chilled storage effects were more intense in
44 L4 than in L1.

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48 **Keywords:** Konjac glucomannan; Breaking deformation; Restructured seafood
49 product; Alkaline coagulant; Deacetylation

1. Introduction

Restructured seafood products are processed from minced and/or chopped muscle, usually with added ingredients, to make products with a new appearance and texture. The last 30 years have seen the development of a new generation of seafood products called analogues or substitutes, most of which mimic seafood or other high-value products and are formulated essentially from surimi, or in some cases mince. The processing is the result of thermal gelation or in some cases cold gelation with the help of ingredients like alginates or transglutaminase (Moreno, Carballo, & Borderias, 2008). However, if muscle protein functionality has been lost in processing, for example heating, gel formation to produce structures is not possible. One possibility in that case would be to find a substance, neutral in colour and flavour, that after being mixed with minced muscle could form a thermo-stable gel. Such a substance could be Konjac glucomannan (KGM) a neutral polysaccharide derived from the tuber of *Amorphophallus konjac* C. Koch (Nishinari, Williams, & Phillips, 1992) that has the property of making thermo-stable gels in certain conditions (Nishinari, 1987; Nishinari et al., 1992; Yoshimura & Nishinari, 1999; Zhang et al., 2001) and is generally recognized as safe (GRAS). It consists of a linear backbone of β -1—4-linked D-glucopyranose and D-mannopyranose sugars in a random order in a molecular ratio of 1:1.6 (Kato & Matsuda, 1969) and possesses between 5-10% of acetyl substituted residues (Dea et al., 1977; Maekaji, 1978a), which makes it soluble in aqueous solution and improves chain flexibility. Eliminating acetyl groups produces deacetylated konjac glucomannan, which is able to build junction zones through hydrogen bonding, Coulombic (between solvated (Na^+ , K^+) cations and KGM deacetylated anions ($\text{R-CH}_2\text{O}^-$) being R the framework of carbohydrate, dipole-dipole (among water molecules and OH groups of deacetylated KGM), van der Waals, charge transfer and hydrophobic interactions (Lapasin & Prici, 1999). However, it is not yet fully understood how this mechanism works under different conditions yet (Yin, Hongbin, Huang, & Nishinari, 2008; Yoshimura & Nishinari, 1999). The type of binding in the junction zone and the amount of molecules forming the junction zones are the most important factors determining the rheological and thermal properties of a gel (Williams & Phillips, 2004). Acidic deacetylation is possible, but alkaline reactions are preferred to limit hydrolysis (Imeson, 2010). The importance of KGM gels in rheological terms is firstly its capacity to extensively modify the rheology of aqueous media to which they are added, even at fairly low concentrations, and that is the basis of their functional properties as thickening and gelling agents. Secondly they are also involved in many other types of

37 applications, such as encapsulation, controlled release, etc. (Lefebvre & Doublier,
38 2005).

39 Therefore, small- and large-deformation mechanical tests need to be monitored in
40 various different alkaline conditions to obtain information about the structure and
41 mechanisms of gelation in different gels at a constant polysaccharide concentration.

42 The rate of gel formation is dependent on various factors: concentration and
43 molecular weight of glucomannan, processing temperature, degree of acetylation,
44 alkaline concentration (Nishinari & Zhang, 2004), and also pH (Huang, Takahashi,
45 Kobayashi, Kawase, & Nishinari, 2002). In general, at a fixed alkaline concentration the
46 gelation process accelerates with increasing molecular weight, KGM concentration or
47 heating temperature (Yoshimura & Nishinari, 1999; Huang et al., 2002) but it is delayed
48 with an increasing degree of acetylation (Huang et al., 2002). As regards pH, Kohyama
49 and Nishinari (1990) reported that the gelation of KGM occurs from pH 11.3 to 12.6,
50 suggesting that a change in molecular structure is necessary for gelation. However,
51 Thomas (1997) indicated that the pH range necessary for gelation is 9 to 10. It seems
52 that the gelation rate also depends on the alkaline concentration and is lower at higher
53 concentrations, but there is a critical alkaline concentration below which gelation does
54 not occur (Huang et al., 2002). In a previous work, Maekaji (1978a) suggested that the
55 deacetylation produced by the addition of alkalis is governed by the concentration of
56 hydroxide ion, irrespective of the kind of alkali, and the deacetylation ratio—i.e. acetyl
57 groups removed/total acetyl groups—was practically independent of gelling
58 temperature and increased with decreasing KGM concentration.

59 Very little scientific literature is available reporting studies of mixes of KGM and
60 fish. Park (1996) reported that KGM has the ability to reinforce gel hardness 8-10 fold
61 in both whiting and pollock surimi. Thomas (1997) also reported the use of KGM at a
62 concentration of 1 % in surimi, but he did not elaborate. Iglesias-Otero, Borderias, &
63 Tovar (2010) added glucomannan to reinforce squid surimi gelation.

64 Also a recent paper verified the good cryoprotective effect of KGM on protein
65 from grass carp (*Ctenophryngodon idella*) during frozen storage (Xiong et al., 2009). The
66 present authors have studied different KGM solubilization conditions, different KGM
67 concentrations and types of alkali at different concentrations with a view to producing
68 homogeneous gels with a texture similar to that of fish fillets. As regards the type of
69 alkali, the literature reports the use of various different alkali coagulants such as
70 phosphate buffers (Penroj, Mitchell, Hill, & Ganjanagunchorn, 2005), Na_2CO_3 (Hata,
71 Ono, & Toda, 1951; Huang et al., 2002; Huang & Lin, 2004), K_2CO_3 (Case, Knopp,
72 Hamman, & Schwartz, 1992; $\text{Ca}(\text{OH})_2$ (Hata et al., 1951; Jiménez-Colmenero et al.,
73 2010; Na_3PO_4 (Nishinari et al., 1992), NaOH or KOH (Maekaji, 1978b). Using this

74 information, in preliminary studies for the work (not included in this paper), various
75 types of alkaline solutions (KOH, NaOH, Ca(OH)₂, Na₂CO₃, K₂CO₃, Na₃PO₄ and
76 K₃PO₄) at different concentrations (0.2, 0.6 and 1N) were added to make gels from
77 KGM aqueous solutions. In these experiments gels firm enough to hold the mince as a
78 filler and with a suitable colour only were obtained using KOH and NaOH, the alkaline
79 agents used in the present work, at concentrations of 0.6 and 1N. These levels
80 translate as 0.073- 0.196 g/kg, which is very little for standard food products, and do
81 not pose problems as regards flavour or health properties.

82 This work is the first step of a study whose objective is to produce restructured
83 seafood products in which KGM aqueous solutions and non-functional minced fish
84 muscle are mixed, in such a way that the glucomannan forms a gel structure and the
85 mince acts as a filler, with a view to forming structures of various kinds fibres,
86 myotomes, etc.). The aim of this paper is to study the influence of the type and
87 concentration of the alkali on the viscoelastic, mechanical and water retention
88 properties of KGM gels in order to choose the most suitable thermo-stable gel for
89 making restructured seafood products with different textures.

90

91 **2. Materials and methods**

92 *2.1. Preparation of glucomannan gels*

93 Aqueous solutions (3% (w/v) of glucomannan from konjac (glucomannan purity
94 100%, Guinama, Valencia, Spain) were prepared by continuous stirring for 30 min at
95 low speed in a homogenizer (Stephan UM5, Stephan u. Söhne GmbH & Co., Hameln,
96 Germany) at 60° C. Then 5% of NaOH or KOH (Panreac Química, S. A., Barcelona,
97 Spain) was added at a concentration of 0.6 or 1N of alkali coagulant, mixing for 1
98 minute at 50 rpm to induce gel formation. After that, Petri dishes were filled with these
99 mixtures and immediately vacuum packed into plastic bags (Barrier bag®, Cryovac Air
100 Corporation, Barcelona, Spain) to compact the samples. Subsequently the samples
101 were set by heating at 30°C for 1 hour and then at 5°C for 4 hours to obtain heat-stable
102 gels. The last step was to reduce the high pH values by taking the gels out of the Petri
103 dishes and placing them in a 0.2M citrate-phosphate buffer at pH 5 (gel:buffer
104 proportion 1:10). After 20 hours at 5°C, the pH of thermostable gels changed to neutral.
105 They were then kept refrigerated (5°±1°C) for 10 days. Analyses were performed at day
106 1 after gel preparation and after 10 days of chilled storage.

107 The different lots were named L1, L2, for gels made with 0.6N and 1N of KOH
108 respectively, and L3 and L4 for the gels made with 0.6N and 1N of NaOH respectively.

109

110 *2.2. Analyses*

111 2.2.1. Physicochemical analysis

112 The pH was measured using a model 9165BNWP pH probe (Analítica
113 Instrumental, S.A., Barcelona) inserted in the gel. The pHmeter was an Orion model
114 720A (Analítica Instrumental, S.A., Barcelona).

115 Water content was determined by drying the sample to constant weight at 110°C
116 and the results are expressed as a percentage (AOAC, 2000). Samples were analysed
117 in triplicate on days 1 and 10 after gel preparation.

118 The pH was measured in triplicate on days 0, 1, 5 and 10 of chilled storage; day 0
119 was the day immediately prior to putting the samples into the pH 5 buffer.

120

121 2.2.2. Water binding capacity (WBC)

122 Gels were cut into small pieces (2 g) and placed in a centrifuge tube (diameter 10
123 mm) with enough filter paper (2 filters Whatman nº 1, diameter 90 mm). Then the
124 samples were centrifuged in a Jouan MR1812 centrifuge (Saint Nazaire, France) for 10
125 min at 3000g at room temperature. WBC was expressed as per cent water retained per
126 100 g water present in the sample prior to centrifuging. Measurements were carried out
127 in triplicate on days 1 and 10 of refrigeration storage.

128

129 2.2.3. Cooking loss determination

130 A sample (40 g) was cut into small pieces and placed in a plastic bag where small
131 holes had been made to drain the drip. Then, this bag with the sample inside was put
132 inside another bag, hung with the holes at the bottom and cooked in that position in an
133 oven (Rational Combi-Master CM6) for 20 min at 100°C. The sample was then cooled
134 and weighed. Cooking loss was expressed as g/100 g by weight difference between
135 uncooked and cooked samples.

136

137 2.2.4. Colour measurements

138 Lightness (L*), was analysed using a CIELab scale. Measurement was analysed
139 using a colorimeter (Minolta Chroma Meter Cr-200, Japan). The colour coordinates
140 were measured five times on the surface of the gel at three different analysis times (on
141 days 0, 1 and 10 of chilled storage). Before use, the colorimeter was standardized
142 using a white calibration plate.

143

144 2.2.5. Puncture tests

145 Cylindrical samples (diameter 3 cm x height 3.5 cm) were filled after gel
146 preparation. After neutralization in the buffer probes they were removed from the
147 cylindrical cells. Before the analyses, probes were tempered at room temperature

148 (25°C). They were then taken out of the bags and analysed. Gels were pierced to
149 breaking point using a TA-XTplus Texture Analyser (Stable Micro System Ltd., Surrey,
150 UK) with a 5 mm–diameter round–ended metal probe. Crosshead speed was 1 mm/s,
151 and a 5 kg load cell was used. The load (as breaking force) and the depth of
152 depression (as deformation) when the gel sample lost its strength and ruptured were
153 recorded. All determinations were carried out at least in sextuplicate.

154

155 2.2.6. Dynamic rheometry measurements

156 Small deformation shear oscillatory testing was performed using a Bohlin CVO
157 controlled stress rheometer (Bohlin Instruments, Inc. Cranbury, NJ). The
158 measurements were carried out using parallel-plate geometry (20 mm in diameter and
159 1 mm gap). Definitive gels were cut into disk-shaped slices 20 mm in diameter and 1
160 mm thick on a 570 S.T.E slicer (Germany). Any excess sample protruding beyond the
161 upper plate was carefully removed. Samples were allowed to rest for 15 min before
162 analysis to ensure both thermal and mechanical equilibrium at the time of
163 measurement. Samples were covered with a thin film of Vaseline oil (Codex
164 purissimum) to limit evaporation. No evidence of specimen slippage at the bottom plate
165 was detected in any case (disk-shaped slices remained intact at the same initial
166 position). The temperature was controlled to within 0.1°C by a Peltier element in the
167 lower plate and was kept at 25.0 °C.

168 *Stress sweep tests*

169 To determine the linear viscoelastic (LVE) region, stress amplitude sweeps
170 were run at 6.28 rad/s, and 25 °C. The amplitude sweeps were conducted by varying
171 the shear stress (σ) of the input signal from 0.24 to 1000 Pa. 300 points in the
172 continuous mode were used in all instances. Changes in storage modulus (G'), loss
173 modulus (G'') and complex modulus (G^*) were recorded. The critical (maximum) values
174 of the amplitude sweeps—shear stress (σ_{max}) and shear strain (γ_{max}) at which the G^*
175 values are just beginning to show a noticeable deviation from the previously constant
176 values—were determined from these data. The range of tolerable deviation ($\pm 10\%$)
177 was corroborated using creep and recovery tests (Mezger, 2006).

178 *Creep and recovery tests*

179 An instantaneous stress σ_0 (30 Pa) was applied for 600 s to each sample in the
180 creep tests and the resulting change in strain over time was monitored. When the
181 stress was released, some recovery was also observed for 600s. Creep measurements
182 were made over the linear viscoelastic range on each sample (σ_0 corresponding to
183 0.5% shear strain). The creep and recovery results are described in terms of the shear

184 compliance function, $J(t) = \gamma(t)/\sigma_0$. Compliance curves generated at different linear
185 stress levels overlap, making it possible to examine and compare the structural
186 properties of the different food gels on larger time scales (Steffe, 1996).

187 From $J(t)$ data we obtained the relaxation modulus $G(t)$, which was used to find
188 the gel strength (S) and relaxation exponent (n) (Ferry, 1980).

189 All measurements were made at 25 °C.

190 *Mechanical spectra*

191 Samples were subjected to stress that varied harmonically with time at a variable
192 frequency. The shear strain amplitude was fixed at 0.5 %; oscillatory frequency sweeps
193 were run from 10 to 0.1 Hz, and measurements were made at 25 °C. The complex
194 modulus (G^*), storage modulus (G'), loss modulus (G''), and loss factor, $\tan \delta$, were
195 determined as functions of frequency. Data were obtained in such a way as to ensure
196 that the resulting σ in the sample would always fall within the linear viscoelastic range.

197

198 2.2.7. Statistical analyses

199 At least five independent batches were tested for each experiment and data are
200 presented as averages. Statistical analysis was carried out using Microsoft Excel
201 software. Trends were considered significant when means of compared sets differed at
202 $p < 0.05$ (Student's t-test).

203 Statistical correlations between the textural and viscoelastic parameters were
204 determined by multiple regression with confidence intervals of 95% ($p < 0.05$) using the
205 SPSS Statistics 17.0 software.

206

207 **3. Results and discussion**

208

209 *3.1. Evolution of pH*

210 At Table 1 shows, the evolution of pH was the same in all the gels. The values
211 were around 12 just after addition of 5% alkali and gel setting (day 0). These values
212 dropped drastically to around 6-7 after the samples had been kept for 20 hours in
213 citrate-phosphate buffer 0.2M pH 5 and remained constant over 1, 5 and 10 days of
214 chilled storage. There were no significant differences ($p < 0.05$) in the average values
215 in the different lots.

216

217 *3.2. Moisture content, Water Binding Capacity (WBC) and cooking loss*

218 Table 2 shows the moisture content, water binding capacity (WBC) and cooking
219 loss of samples L1- L4.

220 All the samples showed very high moisture content (96-97%). There were no
221 significant differences ($p < 0.05$) among the different lots at days 1 and 10. Teramoto &
222 Fuchigami (2000) reported that konjac glucomannan gel as a food had high water
223 content (approximately 97%) but he did not indicate the percentage of glucomannan
224 and the type of alkali (sodium carbonate or calcium hydroxide) used in the formation of
225 these gels. Some authors (Kök, Abdelhameed, Ang, Morris, & Harding, 2009; Shinzato,
226 Broussalis, & Ferraro, 1996) have reported that KGM forms highly viscous solutions
227 when dissolved in water, suggesting that KGM has the highest viscosity at lowest
228 concentration of any known dietary fibre (Shinzato et al., 1996; Yassen, Herald,
229 Aramouni, & Alavi, 2005) so that it can take up to 200 times its weight in water.

230 WBC values were between 64.8%-78.5%. At day 1 values seemed to tend to be
231 higher in samples made with NaOH (L3 and L4), although none of the samples showed
232 significant differences ($p < 0.05$). These WBC values were stable after 10 days of
233 chilled storage. L3 (NaOH 0.6N) showed the highest value ($p < 0.05$) but was only
234 significant respect L2 (KOH 1N). Extraordinarily high water binding capacity has been
235 reported in KGM although no data were given (Kök et al., 2009). The values for
236 cooking loss were slightly but significantly ($p < 0.05$) higher in samples treated with
237 KOH (L1 and L2) than in samples with NaOH added (L3 and L4), both at day 1, just
238 after making the gel, and after 10 days. Cooking loss values were higher ($p < 0.05$) in
239 the sample with the lower alkaline concentration (0.6N KOH) (L1) than in the one with
240 the higher concentration (1N KOH) L2. Water retention values of about 86% have been
241 reported in restructured fish muscle products made with minced fresh horse mackerel
242 muscle, and about 75% in frozen hake muscle (Sánchez-Alonso, Haji-Maleki, &
243 Borderias, 2007).

244 In general, all these data seem to indicate that glucomannan could reinforce the
245 ability of the final product to capture moisture during cooking and retain its texture.
246 There also seem to be some small and non significant differences depending on the
247 alkali used for WBC and slightly higher for cooking drip retention when Na^+ ions are
248 added as NaOH than when K^+ ions are added as KOH. One possible explanation is the
249 difference in the radius of Na^+ and K^+ ions, which would contribute to the degree of
250 hydration of the structures. Large ions, as in the case of K^+ , possess a lower hydration
251 number (N_w), i.e. they are surrounded by a larger number of water molecules (not
252 cation-linked), which retain their translational degrees of freedom. K^+ ions show a
253 smaller average hydration number ($N_w = 2$) compared to Na^+ ions ($N_w = 3$). The later
254 one possess a higher charge density since the same positive charge is located on the

255 smaller Na⁺ ion. Therefore it polarizes the negative electronic clouds of water
256 molecules more effectively (Moore, 1978). Hence NaOH can bind the water more
257 easily (Fennema, 1976). Moreover, Kragh (1977) reported that in the ordered
258 Lipotropic Series of Hofmeister, a cation series from higher to lower hydration ability,
259 the Na⁺ is classified with more hydration ability. Kragh (1977) reported that the
260 mechanism in this series is not clear but is related to the polarity and size of the ion. As
261 noted earlier, these could be the reasons why in general WBC and cooking drip
262 retention are higher when Na⁺ ions are added, since this causes greater hydration.

263

264 3.3. Colour

265 All the gels obtained were translucent, showing high values of lightness (L*). At
266 day 1, L* was higher in gels made with NaOH than those made with KOH (p < 0.05)
267 (Table 3), but after 10 days of chilled storage, L* did not differ significantly in any of the
268 samples except L2 (KOH 1N). The other two colour parameters (a* and b*) were not
269 affected by the type and/or concentration of alkali (data not shown). The lightness of
270 whiting and pollock surimi with added glucomannan was reported by Park (1996), who
271 found that L* was increased by the addition of glucomannan-rich konjac flour. We have
272 found no more colour data for konjac gels in the literature.

273 In general, when gelation is performed at low ionic strength and in acidic or
274 alkaline conditions, it produces fine-stranded gels which are translucent and have high
275 water binding capacity (Rao, 2007). For that reason, in the light of their colour and the
276 WBC data (Table 2) the four samples may be classified among the fine-stranded type
277 physical gel networks.

278

279 3.4. Puncture test

280 Figure 1 shows the Breaking force (1a) and Breaking deformation (1b) of samples
281 L1-L4 at 0 and 10 days of chilled storage. There are significant differences (p < 0.05) in
282 breaking force (1a) between L4 (1N NaOH) and the other samples. After 10 days, L4
283 still had the highest values for strength and L1 the lowest.

284 Although there was an increase in breaking force between day 1 and day 10 in all
285 samples, this was much smaller and not significant in L1 than in the other samples,
286 which could mean that the gel was more stable over chilled storage.

287 On the other hand, breaking deformation was very similar in all samples. The
288 higher values were found in L1 and L4 at day 1, and L1 differed significantly (p < 0.05)
289 from L2 and L3 (Figure 1b). As in the case of breaking force, L1 showed practically no
290 change between day 1 and day 10. L4 likewise showed no significant differences (p <
291 0.05) over the storage time, although these differences were slightly greater than in L1.

292 Therefore again it seems that the L1 network was the most time-stable of all the
293 samples.

294 In short, these large-deformation rheological measurements made it possible to
295 distinguish L1 and L4 samples, since when NaOH 1N was used as an alkaline agent,
296 the resulting gel possessed similar breaking deformation but significantly higher
297 breaking force, making the gel significantly more rigid. This mechanical behaviour is
298 consistent with the fact that viscoelastic moduli values from mechanical spectra were
299 higher [for the respective samples](#), as will be discussed in the next paragraph. The
300 trend was sustained after 10 days of chilled storage (Figures 1a and 1b).

301 Although large-strain deformation mechanical tests, provide information that
302 correlates more with the sensory perception and handling properties of food products
303 (Bollaín, Angioloni, & Collar, 2005; van Vliet, 1995), small deformation oscillatory
304 measurements have been considered a necessary and useful tool to study the network
305 structure of foods, especially in food gels that present complex viscoelastic behaviour
306 (Romero et al., 2009). Therefore, we considered it necessary to conduct a thorough
307 rheological study to identify the properties of these gel networks.

308

309 *3.5. Rheological measurements at small deformation. Influence of chilled storage*

310

311 3.5.1. Overview of the small amplitude oscillatory (SAOS) results

312 The viscoelastic behaviour of the four samples was characterized at initial time in
313 terms of several critical parameters which determine the linear viscoelastic range (LVE) ,
314 together with their mechanical spectra. The information from the oscillatory tests was
315 then supplemented by the results of transient tests, also at initial time, to discriminate
316 which samples had better gel properties. Thereafter, we examined the influence of 10
317 days of chilled storage on the viscoelastic magnitudes, but only in the samples with
318 better gel characteristics ([higher stress and strain amplitudes from stress sweeps, low
319 frequency-dependence on G', and lower viscous moduli from mechanical spectra](#)).

320

321 3.5.2. Stress sweeps at initial time

322 The first step was to investigate the LVE range for all the polysaccharide gels
323 when a different kind and concentration of alkali was used. Within this range the
324 viscoelastic moduli are independent of the stress. As the applied stress increases, the
325 bonds holding the network together begin to rupture. At critical values of stress (σ_{max})
326 and strain (γ_{max}), the network structure breaks down leading to a sharp decrease in the

327 moduli. These critical values, which may serve as a measure of the stability of
328 viscoelastic materials, were obtained from an automatic analysis (Campo-Deaño,
329 Tovar, Pombo, Solas, & Borderias, 2009) and were corroborated using creep and
330 recovery tests, since compliance curves generated at different stress levels overlap
331 when data are collected in the LVE range (Steffe, 1996).

332 Table 4 shows the more representative magnitudes which characterize the
333 amplitude of this LVE region. $\tan\delta=G''/G'$ values were similar (≈ 0.2), and significantly
334 lower than 0.5 in the four samples, meaning that samples behave like a viscoelastic gel
335 since G' is larger than G'' indicating the presence of a network structure (Mezger,
336 2006).

337 There was no statistically appreciable difference in the overall rigidity of their networks
338 (G^*) (Table 4), however L1 and L4 samples showed higher stress and strain
339 amplitudes than L2 and L3, which is consistent with the fact that they showed the
340 highest breaking force and breaking deformation at initial time, as confirmed by large
341 deformation measurements (Figure 1a and b).

342

343 3.5.3. Frequency sweeps at initial time

344 The frequency dependence of G' and G'' can provide valuable information about
345 the structure of a gel. Mechanical spectra for selected alkali/concentration values of
346 glucomannan gels are presented in Figure 2. The mechanical spectra show little
347 variation of G' and G'' over the entire frequency range studied. The G' values can be
348 fitted to the power law (equation 1), but the G'' moduli remain practically constant from
349 high to low frequencies.

$$350 \quad G' = G_0' \cdot \omega^{n'} \quad (1)$$

351 Table 5 shows the power law parameters for G_0' and n' together with the mean
352 viscous modulus between 0.63 and 63 rad/s. In L1-L4 samples, G' is practically
353 frequency-independent over this time scale ($n' < 0.1$). In addition $G_0' \gg G''$ so samples,
354 can be classified as true gel systems (Kaur, Singh, Singh, & McCarthy, 2008). There
355 was significant ($p < 0.01$) negative correlation of n' (-0.998) and $\tan\delta$ (-0.970) with
356 WBC , confirming that when a network gel approximates to a true gel (low n' and $\tan\delta$),
357 the gel functionality is better and hence water binding capacity noticeably increases.

358 This rheological behaviour is also quantifiable in terms of quality factor Q
359 (Campo-Deaño, Tovar, & Borderias, 2010), a dimensionless quantity which represents
360 the degree of damping of an oscillator. The Q factor is 2π times the ratio between the
361 energy stored and the average energy loss per period (Arya, 1990). On the basis of the
362 oscillatory character of frequency sweeps and the peculiar mechanical spectra data

363 (Figure 2 and Table 5), Q can be calculated from G' (eq. 1) and the mean viscous
364 modulus in a sinusoidal strain (Table 5):

$$365 \quad Q = 2\pi \frac{G_0'}{G''} \omega^{n'} \quad (2)$$

366 Note that except for L2 gel, all the rest possess high Q values, which is a
367 measure of structural stability in their networks on short time scales. The fact that the
368 mean viscous moduli are lowest in L1 gel in particular indicates that the relevant polar
369 interactions to cross-links are junction zones with finite energy, which act more
370 cooperatively to ensure gel stability at short times. A more permanent three-
371 dimensional network was formed, so that although the gels contained around 97%
372 solvent (Table 2), they are macroscopically connective, generating a superstructure
373 which can store more energy. Thus, a minimum amount of energy is required for the
374 internal relative motion between molecular segments (low molecular friction) and hence
375 Q increases, indicating higher viscoelastic stability in physical gels over short
376 experimental times.

377

378 3.5.4. Creep and recovery tests at initial time

379 Creep analysis is a transient test which was done at constant stress within the
380 LVE range. The results of these experiments were used to compare the different
381 structural responses of the four gels. This type of analysis produces data on creep
382 compliance, $J(t)$, the ratio of strain to stress over time. Thus, the time dependent
383 properties connected with the viscoelastic characteristics of physical gels can be
384 studied on longer time scales than those associated with oscillatory tests (Mezger,
385 2006). The rheological characterization was completed with transient experiments,
386 which provide a means of trying regimes of $t > 100$ s and can help to distinguish among
387 noncovalent crosslinked gels. Long-term behaviour may be associated with the re-
388 orientation of chain segments, and probably with the movement of whole molecules
389 relative to one another, causing relatively strong crosslinks to break (Lapasin and Prici,
390 1999). Transient tests, then, can cause the breakage of short-range interactions.
391 Therefore information can be obtained about the relative long-range properties of these
392 physical networks (Steffe, 1996). From J_{max} and J_{min} on the creep and recovery curves
393 respectively, it is possible to quantify the percentage of elasticity in the networks
394 according to equation 3:

$$395 \quad Elasticity(\%) = \left(\frac{J_{max} - J_{min}}{J_{max}} \right) \cdot 100 \quad (3)$$

396 Figure 3 shows creep-recovery compliances for glucomannan gels. L1 sample
397 presents the lowest compliance data over the entire time interval. When the load was
398 applied at time $t=0$, there was an instantaneous deformation, from which this sample
399 showed the most complete recovery upon removal of the load, indicating that it was the
400 most elastic (Table 6). Moreover, the fact that it presented the least permanent
401 deformation indicated that for longer loading times there was less structural collapse
402 and less irreversible breakage of interactions (Deman & Beers, 1987).

403 By means of an experiment based on a different physical principle, we can
404 corroborate the viscoelastic stability of the physical network of L1 sample. This stability
405 effect was deduced from stress sweeps (higher strain amplitude) and from mechanical
406 spectra (lowest loss modulus) (Tables 4 and 5 respectively). We should note that the
407 evaluation of the viscous component of gels from frequency sweeps presented the
408 same trend as the irreversible compliance data obtained from the residual strain at the
409 end of the recovery process. We can therefore state that the polymer network in L1
410 sample possesses similar connectivity to the covalently crosslinked networks, as was
411 also reported by Case et al. (1992) for konjac gels.

412 For their part, the L2-L4 gels exhibited higher creep and recovery compliance
413 values, with a significantly low degree of elasticity (Table 6). During creep time, break a
414 lot of hydrogen bonds which originated certain degree of the irreversibly structural
415 damage, thereafter when the load was removed the higher proportion of the structure
416 collapsed gave $J(t)$ on the recovery increased and elasticity diminished. This result is
417 consistent with the higher viscous modulus values of L2-L4 samples (Table 5);
418 therefore, if residual strain increases at the end of the recovery period, as in L2-L4
419 gels, can be attributed to rearrangement of the gel network (Williams & Phillips, 2004),
420 then there would be more broken interactions, which would explain their higher G''
421 values.

422 L4 gel was made with the alkali with the lowest cation diameter (Na^+) at maximum
423 concentration, so that there were more Na^+ ions to link a larger number of water
424 molecules, producing [more layers of associated water molecules](#) and hence a larger
425 number of ion-dipole interactions. This could originate a *physical principle* based on the
426 predominance of strong cation-dipole interaction, added to the corresponding anion-
427 dipole (deacetylated KGM-water), which is less important since the anion is larger. This
428 causes first an average preferential direction among molecular domains, which may
429 generate a particular dipolar chain orientation (local order), and second a noticeable
430 decrease of the available water, which cannot act as a lubricant between anionic
431 polymer chains; this could explain the fact that the sample's physical network was more

432 brittle (high breaking force and low breaking deformation) than L1 (Figure 1a and b), as
433 discussed in section 3.4.

434 Moreover, the $J(t)$ from the creep values gives us the relaxation modulus $G(t)$, since if
435 we plot $\log J(t)$ versus $\log t$ over the entire time interval, the slope of the function $m \ll 1$
436 (data not shown here), $G(t)$ becomes the reciprocal of $J(t)$ (Ferry, 1980). Thus, the
437 equation of Winter and Chambon (te Nijenhuis, 1997) can be used to calculate other
438 parameters related to *gel strength* (S), and also to the *relaxation exponent* (n), by:

$$439 \quad G(t) = S \cdot t^{-n} \quad (4)$$

440 Where S ($\text{Pa} \cdot \text{s}^n$) depends on the strength of the zone junctions between
441 molecular domains and n is related to the density of these zone junctions, i.e. the
442 degree of connectivity in the gel (Gabriele, de Cindio, & D'Antona, 2001). Table 6
443 shows that L1 gel presented the highest S and the lowest n parameters. The lower the
444 n values, the higher is the density of physical crosslinks, which increases the extension
445 of the junction zones in noncovalently crosslinked networks (Lapasin & Prich, 1999). L4
446 gel possessed a similar S value to L1, but n was the highest as a result of less
447 connectivity and may reflect more levels of heterogeneity than in the other samples.
448 Also, there was a significant ($p < 0.01$) positive correlation between S and the critical
449 stress ($r^2 = 0.997$) and strain ($r^2 = 0.996$) (from stress sweeps). This is because strong
450 gels may remain in the LVE range at greater strains level, reflecting higher gel strength
451 values (Steffe, 1996).

452 To summarize, from large and SAOS measurements we can deduce that sample
453 L1, followed by L4, was the most stable, with the most ordered three dimensional
454 network. Both had the highest maximum stress (σ_{max}) and strain (γ_{max}) values, as
455 reflected by their high values of breaking force and breaking deformation respectively.
456 However, L1 was the most elastic, presenting the lowest viscous moduli (G''), and
457 recovery compliance (J) together with the lowest *relaxation exponent* (n), indicating a
458 better-organized and better-ordered network as explained above. In addition, L1
459 showed the least variation in breaking force and breaking deformation over 10 days,
460 which means that this gel was very stable, an important characteristic in a restructured
461 fish product intended for chilled storage. All this means that L1 (0.6N KOH) offers the
462 best properties for use as a gelling agent in restructured fish products. L4 (1N NaOH)
463 could also be used although its network is less well-ordered and time-stable than L1's.
464 It was therefore decided to study the influence of 10 days of chilled storage on
465 viscoelastic properties of L1 and L4 gels.

466 The last step of processing is neutralization with a citrate-phosphate buffer. Although
467 this should theoretically stimulate hydrogen bonding, the gel texture remains

468 unchanged as unpublished previous analyses showed. As most of the bonds probably
469 might already be formed at the time of deacetylation and therefore contribute to a
470 certain thermodynamical stability rearrangements of bonds should be limited.

471 3.5.5. Effect of chilled storage time on the LVE interval and mechanical spectra

472 First, stress sweeps were used to evaluate the influence of 10 days of chilled
473 storage on the σ_{max} and γ_{max} values in both L1 and L4 samples. There were practically
474 no differences with respect to the corresponding values at initial time (Table 4): σ_{max}
475 values after 10 days were: 90 ± 9 and 70 ± 7 Pa for L1 and L4 respectively. Although
476 γ_{max} values increased after the 10 days, 2.19 ± 0.59 and 1.62 ± 0.33 for L1 and L4
477 respectively, there were no significant differences ($p < 0.05$) with respect to the
478 corresponding γ_{max} values at day 1 (Table 4).

479 However, the frequency sweeps showed considerable changes in the viscoelastic
480 properties when compared to day 1. There was thus a general decrease in the gel-like
481 character of both samples after 10 days: on the one hand the G_o' parameter (equation
482 1) decreased significantly in both samples, more so in L4 (29 % with respect to initial
483 value) than in L1 (18%) (Fig. 4a). There was also a greater increase in frequency-
484 dependence, similar in L1 and L4 (Fig. 4c), while G'' increased considerably in L1 (40%
485 with respect to initial value) and decreased in L4 (Figure 4b), making for a smaller gap
486 between G' and G'' . This meant a sharp decline in the elasticity of the networks.

487 These results show that in this kind of transient networks, chilled storage breaks
488 the junction zones stabilized by thermolabile polar interactions such as ion-dipole and
489 hydrogen bonds between polymer chains and with water. This effect was more
490 pronounced in L1 sample given that the quality factor Q decreased by 39% while in L4
491 it decreased by 10% with respect to the corresponding values at day 1 (Table 5). This
492 trend reflects a higher degree of connectivity by means of physical interactions in L4
493 sample, since Na^+ has more hydration ability (Kragh, 1977), and hence the loss of
494 solidity was smaller than in L1 on short time scales

495

496 3.5.6. Effect of chilled storage time on the creep and recovery data

497 Figure 5 shows the influence of 10 days of chilled storage on the gel strength and
498 relaxation exponent parameters from equation 4. In both L1 and L4 samples, these
499 transient data confirm that chilled storage time caused some structural damage. This
500 effect can be seen in the noticeable loss of the gel strength (Figure 5a) and the
501 significant increase of the relaxation exponent (Figure 5b), which is associated with a
502 decrease of polymer molecular weight due to the rupturing of a sequence of physical
503 crosslinks (Lapasin & Prici, 1999). After 10 days this damage was somewhat greater in

504 L4, which presented a slightly greater decrease in extent (17%) and strength of
505 connectivity (45%) than L1 (12 % and 39% respectively). In general, the effect of
506 chilled storage was slightly greater in gels made with NaOH than those made with
507 KOH; in the former the number and distribution of anionic sites of KGM is altered by
508 higher number of big hydrated ions that could generate, more dipolar fluctuations within
509 the network (number and position of all these noncovalent cross-links), and hence the
510 effect of experimental and storage time will be greater. On the other hand, as noted
511 earlier, L1 (with minor quantity of small hydrated ions) was more time-stable in terms of
512 breaking force and breaking deformation (Fig. 1), with hardly any difference between
513 day 1 and day 10 for either parameter.

514

515 **4. Conclusions**

516 In this study various physicochemical measurements and various rheological
517 techniques based on different physical principles such as large and SAOS
518 deformations on different time scales (oscillatory and transient tests), were used to
519 determine the influence of several alkalis at different concentrations on deacetylated
520 glucomannan gels. Of the samples analyzed aqueous dispersions of 3% glucomannan
521 deacetylated with 0.6N KOH and with 1N NaOH seem to possess well-structured gel
522 properties and to be moderately stable over time, making them suitable for use in raw
523 restructured seafood products that have to be stored chilled.

524 Of the samples assayed, the one with 0.6 N KOH performed best in terms of
525 elasticity and time- and structural stability. Tests done over longer time scales, such as
526 creep-recovery experiments, show low residual strain values and greater connectivity,
527 which is consistent with results from experiments performed over shorter times, such
528 as low viscous moduli values from mechanical spectra and high breaking deformation
529 (from large deformations).

530 In order to be able to use them as gelling agents in restructured fish products, a
531 thermal rheological study of these two gels (0.6N KOH and 1N NaOH) is required to
532 demonstrate their thermo-stability.

533

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688 **Captions for Figures**

689

690 **Figure 1.** The change of the breaking force (a) and breaking deformation (b) of
691 glucomannan gels deacetylated with different alkali at 0.6N and 1N after 1 and 10 days
692 of chilled storage.

693

694 **Figure 2.** Mechanical spectra data of glucomannan gels deacetylated with different
695 alkali at 0.6N and 1N after 1 day of chilled storage. Closed symbols G' , open symbols
696 G'' . $T= 25^{\circ}\text{C}$

697

698 **Figure 3.** Creep and recovery compliance $J(t)$ data of glucomannan gels deacetylated
699 with different alkali at 0.6N and 1N at day 1 of elaboration. $T=25^{\circ}\text{C}$.

700

701 **Figure 4.** Effect of refrigerated time at 5°C on power law parameters from equation 1
702 G_0' , n' , and viscous modulus (G'') from frequency sweeps of L1 and L4 . $T= 25^{\circ}\text{C}$.

703

704 **Figure 5.** Influence of refrigerated time at 5°C of L1 and L4 gels, on gel strength (S)
705 and relaxation exponent (n) from equation 4. $T= 25^{\circ}\text{C}$.

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