

Enzyme-assisted extraction of k/I-hybrid carrageenan from Mastocarpus stellatus for 1 2 obtaining bioactive ingredients and their application for edible active films development 3 N. BLANCO-PASCUAL, A. ALEMÁN, M.C. GÓMEZ-GUILLÉN & M.P. MONTERO* 4 Instituto de Ciencia y Tecnología de Alimentos y Nutrición (ICTAN-CSIC)**, 5 6 C/ José Antonio Nováis, 28040 – Madrid (Spain) 7 8 * Corresponding author. Tel: +34-31-5492300; fax: +34-91-5493627. 9 E-mail address: mpmontero@ictan.csic.es 10 ** This centre has implemented and maintains a Quality Management System which fulfils the 11 requirements of the ISO standard 9001:2000. 12 13

14 Abstract

15 Two hydrolysates were obtained from dried Mastocarpus stellatus using alcalase. Phenolic 16 content was partially removed from one of them. The phenolic-partially-removed hydrolysate (H) 17 was found to be a potent Angiotensin-I converting enzyme (ACE) inhibitor. However, the 18 phenolic-containing hydrolysate (Hp), showed a higher Folin-reactive substance content and 19 antioxidant capacity (reducing power and radical scavenging capacity). Hp was therefore 20 selected for the development of antioxidant Mastocarpus carrageenan-based films. F-Hp0 (without hydrolysate), F-Hp15 (with 15% hydrolysate) and F-Hp30 (with 30% hydrolysate) films 21 22 were developed. K/I-hybrid carrageenan was the main film constituent and hydrolysate addition 23 resulted in an increased sulfated proportion, higher protein content and higher number of 24 hydrogen bonds. Therefore interactions between carrageenan helices, plasticizer and peptides 25 in the film-forming solution were enhanced, especially in F-Hp15, and consequently the water 26 vapour permeability (WVP) of the resulting film decreased. Nevertheless, F-Hp30 considerably 27 improved transparency, UV/Vis light barrier, water resistance and elongation at break (EAB). Hp presence increased both puncture force (F) and puncture elongation (E), but not tensile strength 28 29 (TS) or Young's modulus (Y). The addition of an increased concentration of hydrolysate to the 30 films led to a considerable increase in the Folin-reactive substance content and the antioxidant 31 activity, especially the radical scavenging capacity.

32 Keywords: Seaweeds, Mastocarpus, films, hydrolysate, antioxidant, antihypertensive

34 **1**. Introduction

Seaweeds have recently been included in Western diets as food and also as components of functional products because of their richness in polysaccharides, proteins,¹ minerals and vitamins. Moreover, seaweeds are an excellent source of bioactive substances such as sulfated polysaccharides, peptides and polyphenols with biological activities, including antioxidant and antihypertensive properties.²⁻⁵

In the last decade, new marine bioprocess technologies have allowed the isolation of substances with antioxidant properties or bioactive peptides by enzymatic hydrolysis.⁶ Seaweeds have proved to be a good source of peptides and polyphenols.⁷⁻⁹ Red algae (Rhodophyta) are known to have a high protein content, mainly composed of bioactive phycobiliproteins¹⁰ and other wall proteins that might be more efficiently extracted by an enzyme-assisted treatment.^{11,12}

46 *Mastocarpus stellatus* is one of the few carrageenophyte species on the Atlantic coast currently
 47 harvested for phycocolloid industry purposes, but it is still underutilized.¹³

Commercial carrageenan is commonly extracted at alkaline conditions (pH 7–9) at temperatures
near boiling point (80–110 °C) for 3–4 h, providing yields of 20–40%.^{13,14} However, high
molecular weight carrageenan can also be extracted at mild temperatures (50 °C) for 1–5 h.¹⁵

51 κ/ι-hybrid carrageenan has been reported to be the main biopolymer structure extracted from *M.*52 stellatus,¹⁶ although other components, such as proteins, minerals and polyphenols, are also
53 present in significant amounts.¹⁷ Mastocarpus enzymatic hydrolysis could produce both
54 antihypertensive and antioxidant extracts, as previously reported with another species of the
55 Rhodophyta phylum.^{18,19} Protein hydrolysates from different origins have been incorporated in
56 the formulation of protein-based films to improve or confer bioactivity.^{20,21}

57 Mastocarpus extraction could be maximized by first performing an enzymatic hydrolysis at mild 58 temperatures and alkaline conditions followed by carrageenan precipitation and bioactive 59 compound isolation. Antioxidants have been widely used as food additives both to improve lipid oxidation stability and to extend the product shelf life. The potential use of films as antioxidant releasing packages, capable of improving food preservation, would be a rather interesting application.

The aims of the present study were: (i) to obtain two different potentially bioactive hydrolysates with antioxidant and/or antihypertensive capacities from dried *Mastocarpus stellatus*, and (ii) to develop antioxidant Mastocarpus carrageenan-based films as an active edible food packaging material.

67 2. Materials and Methods

68 2.1. Seaweed sampling

69 Samples of *Mastocarpus stellatus* (M), kindly supplied by Porto-Muiños (Cerceda, A Coruña, 70 Spain), were washed several times with running tap water and air-dried at 50 °C for 24–48 h in 71 a ventilated oven. Seaweed samples were stored in sealed plastic bags at 2–4 °C for 1 week 72 until use.

73 **2.2. Unrefined biopolymer extraction**

74 Dried seaweed was homogenized using an Osterizer blender (Oster, Aravaca, Madrid, Spain) 75 with water in a 1:15 (w:v) proportion and kept for 12 h at 3 ± 2 °C. The seaweed was then 76 filtered and subjected to two consecutive extractions in water at a 1:30 (w:v) proportion, at 91 °C 77 for 2 h during the first step and 1.5 h during the second one. Each extract was centrifuged at 78 3000 rpm for 5 min (Sorvall Evolution RC Centrifuge, Thermo Fisher Scientific Inc., Landsmeer, 79 The Netherlands) and blended. The supernatant was dried in an oven (FD 240 Binder, 80 Tuttlingen, Germany) at 65.0 ± 0.8 °C and this constituted the Mastocarpus biopolymer extract, 81 which was stored at room temperature.

82 2.3. Seaweed hydrolysis

Dried seaweed was mixed with 4% distilled water (w/v) and subjected to enzymatic hydrolysis for 3 h, using alcalase 2.4L (EC 3.4.21.14, 2.64 AU/g, Sigma-Aldrich Inc., St. Louis, MO, USA) in optimal conditions for enzymatic activity (pH 8, 50 °C). The enzyme-substrate (seaweed) ratio 86 was 1:20 (w:w) and the pH of the reaction was kept constant by addition of 1 N NaOH solution 87 to the reaction medium using a pH-stat (TIM 856, Radiometer Analytical, Villeurbanne Cedex, 88 France). The enzyme was inactivated by heating at 90 °C for 10 min. The hydrolysate was 89 centrifuged at 7000 g for 15 min. The supernatant was subjected to two consecutive 90 carrageenan extractions by precipitation with ethanol 1:3 (v/v) at 4 °C for 2 h. The precipitated 91 carrageenan was dried at 65 ± 0.8 °C and weighed in order to evaluate extraction yields. The 92 carrageenan-free liquid phase was centrifuged at 13000 g for 5 min. The supernatant was 93 concentrated by rotary evaporation and was subsequently subjected to five organic extractions 94 with ethyl acetate 1:5 (v/v), to remove most of the polyphenols and other compounds such as 95 pigments. After decanting, the successive aqueous phases were concentrated by rotary 96 evaporation. The concentrate was lyophilized, and this constituted the phenolic-partially-97 removed hydrolysate (H). The phenolic-containing hydrolysate (Hp) was obtained under the 98 same conditions as described above, with the exception of the removal of polyphenol compounds with ethyl acetate. The Hp hydrolysate was selected for active film development. 99

100 **2.3.1. Amino acid analysis of hydrolysates**

101 The amino acid composition of the hydrolysates (H, Hp) was determined using a Biochrom 20 102 amino acid analyser (Pharmacia, Barcelona, Spain) according to the method described by 103 Alemán *et al.*²² The results were expressed as number of amino acid residues per 1000 104 residues.

105 2.3.2. Angiotensin-converting enzyme (ACE) inhibition of hydrolysates

106 Reversed-phase high performance liquid chromatography (RP-HPLC) was used to determine 107 ACE-inhibitory capacity of the hydrolysates (H, Hp), according to the method described by 108 Alemán *et al.*²² The IC₅₀ value was defined as the concentration of hydrolysate (μ g/mL) required 109 to inhibit 50% of ACE activity.

110 2.4. Film preparation

111 Three film-forming solutions were prepared to obtain the following films: F-Hp0 (without the 112 addition of hydrolysate), F-Hp15 (with 15% hydrolysate) and F-Hp30 (with 30% hydrolysate). 113 Film-forming solutions (FS) (2% w/v) were prepared from Mastocarpus biopolymer extract by 114 adding hot distilled water (90 °C) and homogenizing with a T25 basic Ultra-Turrax (IKA-Werke 115 GmbH & Co. KG, D-79219 Staufen, Germany) at 17500–21500 rpm for 5 min. A portable pH-116 meter series 3 Star Orion with an electrode pH ROSS (Thermo Fisher Scientific Inc., 117 Landsmeer, Netherlands) was used for pH measurements (6.4-6.8). Glycerol (Panreac 118 Química S.A., Barcelona, Spain) was added at 10% (w/w) in relation to the seaweed extract 119 content. The film-forming solutions were centrifuged at 3000 rpm for 3 min to remove air 120 bubbles. Hp was then added at 15 and 30% (w/w) in relation to the seaweed extract content, 121 and was magnetically stirred for 5 minutes. The film-forming solutions were cast into petri dishes 122 and dried in an oven (FD 240 Binder, Tuttlingen, Germany) at 35.0 ± 0.8 °C for 21 h. All the 123 films were conditioned at 58.0 ± 0.2% RH and 22 ± 1 °C for 4 days prior to analysis.

124 **2.5.** Viscoelastic properties of film-forming solutions (FS)

A dynamic viscoelastic study of the film-forming solutions was carried out on a Bohlin CVO-100 rheometer (Bohlin Instruments Ltd., Gloucestershire, UK) using a cone-plate geometry (cone angle 4°, gap 0.15 mm). A dynamic frequency sweep from 0.1 to 10 Hz took place at auto stress, at a temperature of 10 °C and a target strain of 0.005%. The elastic modulus (*G'*; Pa) and viscous modulus (*G'*; Pa) were plotted as functions of the frequency ramp. To characterize the frequency dependence of *G'* over the limited frequency range, the following power law was used:

132 $G' = G_0 \omega^n$

where G_0 is the energy stored and recovered per cycle of sinusoidal shear deformation at an angular frequency of 1 Hz, ω is the angular frequency and *n* is the power law exponent, which should exhibit an ideal elastic behaviour near zero in gels. At least two determinations were performed for each sample. The experimental error was less than 6% in all cases.

137 **2.6. Viscosity**

A viscosity test for film-forming solutions was performed at 25 °C in the cone-plate cell (cone angle 4°, gap = 150 mm) of the Bohlin rheometer at a constant shear rate of 0.5 s^{-1} . The results are averages of eight determinations and are expressed as Pa·s.

141 **2.7. Thermal properties**

142 Calorimetric analyses of extracts and films were performed using a differential scanning 143 calorimeter (DSC) model TA-Q1000 (TA Instruments, New Castle, DE, USA) previously 144 calibrated by running high-purity indium (melting point, 156.4 °C; melting enthalpy, 28.44 J/g). 145 Samples of around 10-15 mg were tightly encapsulated in aluminium hermetic pans. They were 146 scanned under dry nitrogen purge (50 mL/min) between 5 and 180 °C at a heating rate of 10 147 °C/min. Peak temperatures (T_{peak} , °C) and enthalpies of conformational changes (ΔH) were measured at least in triplicate, the latter data being normalized to dry matter content (J/g_{dm}) after 148 149 desiccation of each particular capsule.

150 2.8. ATR-FTIR spectroscopy

Extract and film infrared spectra between 4000 and 650 cm⁻¹ were recorded at least in triplicate
using a Perkin Elmer Spectrum 400 Infrared Spectrometer (Perkin–Elmer Inc., Waltham, MA,
USA), as described by Ojagh *et al.*²³

154 **2.9. Antioxidant activities of hydrolysates and films**

ABTS radical [2,2'-azino-bis-(3-ethylbenzothiazoline-6-sulfonic acid)] scavenging capacity and 155 FRAP (ferric reducing ability of plasma) were used to measure the antioxidant activity of the 156 hydrolysates (H, Hp) and films (F-Hp0, F-Hp15, F-Hp30). Both hydrolysates and films were 157 158 dissolved in distilled water and shaken until they were totally homogeneous. The film solutions were filtered through Whatman No. 1 paper. The method used for the FRAP and ABTS assays 159 was previously described by Alemán et al.22 Results were expressed as µmoles Fe2+ 160 161 equivalents/g for FRAP and mg Vitamin C Equivalent Antioxidant Capacity (VCEAC)/g) for ABTS, based on standard curves of FeSO₄7H₂O and vitamin C, respectively. All determinations 162 163 were performed at least in triplicate.

164 **2.10.** Folin-reactive substances content of hydrolysates and films

165 Total Folin-reactive substances content was determined according to a modified method by 166 Slinkard and Singleton with the Folin–Ciocalteu reagent.²⁴ An aliquot of 10 μ L of sample was 167 mixed with 750 μ L of distilled water and oxidized with 50 μ L of Folin–Ciocalteu reagent. The reaction was neutralized with 150 µL of sodium carbonate solution and incubated for 2 h at room temperature. The absorbance of the resulting blue colour was measured at 765 nm (UV-1601, model CPS-240, Shimadzu, Kyoto, Japan). Results were expressed as mg gallic acid (GA) equivalent/g of sample. All determinations were performed at least in triplicate.

172 2.11. Film determinations

173 **2.11.1. Thickness**

The thickness was measured using a micrometer (MDC-25M, Mitutoyo, Kanagawa, Japan),
averaging the values of 6–8 random locations in 15 films for each treatment as described by
Pérez-Mateos *et al.*²⁵

177 2.11.2. Moisture content

The moisture content was determined at least in triplicate by drying samples of around 0.5 g at
105°C for 24 h, according to A.O.A.C. (Association of Official Analytical Chemists, 1995).²⁶
Water content was expressed as a percentage of the total weight.

181 **2.11.3. Protein content**

The protein content was determined by a LECO FP-2000 nitrogen/protein analyser (Leco Corp.,
St. Joseph, MI, USA), according to Dumas (A.O.A.C., 2005)²⁷ and using a nitrogen-to-protein
conversion factor of 6.25.

185 **2.11.4. Light absorption and transparency**

186 The light barrier properties and transparency of the films were calculated at least in triplicate 187 using a UV-1601 spectrophotometer (model CPS-240, Shimadzu, Kyoto, Japan) at selected 188 wavelengths from 250 to 800 nm following the method described by Pérez-Mateos *et al.*²⁵

189 The films were cut into a rectangle piece and directly placed in the spectrophotometer test cell,

190 using an empty test cell as the reference. Transparency was calculated by the equation

191 Transparency = $-\log(T_{600}/x)$, where T600 is the light transmission (*T*) at 600 nm, and *x* is the

192 film thickness (mm).

193 **2.11.5. Colour**

194 The colour parameters lightness (L^*), redness (a^*), and yellowness (b^*) were measured 195 following the method described by Blanco-Pascual *et al.*²⁸8

196 2.11.6. Water vapour permeability (WVP)

197 The water vapour permeability (WVP) was determined at least in triplicate following the method described by Sobral et al.²⁹ A round portion of film was cut out and mounted on a plastic cup 198 (permeation area = 15.90 cm^2) containing silica gel, and the cups were then placed in desiccators 199 200 with distilled water. Weights were taken every hour for seven hours at 22°C. Water vapor permeability was calculated from the equation WVP= $w.x.t^{1}.\Delta P^{1}$, where w was weight gain 201 202 (g), x was film thickness (mm), t was time of gain (h), A was permeation area, and ΔP was the change in the partial atmospheric vapor pressure with the silica gel and pure water (2642 Pa at 203 22°C). Results were expressed as g.mm. h⁻¹.cm⁻².Pa⁻¹. All measurements were carried out in 204 205 triplicate.

206 2.11.7. Water solubility

Film circumferences of 40 mm in diameter were placed in plastic containers with 50 mL distilled water and placed at 22 °C for 24 h. The solution was then filtered through Whatman # 1 filter paper to recover the remaining undissolved film, which was desiccated at 105 °C for 24 h. Film solubility *FS* (%) was calculated using the expression $[(W_o - W_f)/W_o] \times 100$, where W_o was the initial weight of the film expressed as dry matter and W_f was the weight of the undissolved desiccated film residue. All tests were carried out at least in triplicate.

213 2.11.8. Water resistance

Films were fixed onto the opening of calibrated cells (area 15.90 cm²) and the cells placed in desiccators at 22°C and exposed over distilled water. Distilled water (5 mL) was poured over the film surface. The film deformation due to the water effect, the time when the water started to leak and the time when the film broke were annotated. All tests were carried out at least in triplicate.

219 2.11.9. Mechanical properties

220 Tensile and puncture tests were run using a texture analyzer TA.XT plus TA-XT2 (Texture 221 Technologies Corp., Scarsdale, NY, USA) (58% RH and 22°C) controlled by the Texture 222 Exponent Software (Texture Technologies and by Stable Micro Systems, Ltd., Scarsdale, NY, 223 USA), using a 5 kg load cell. Tensile test: At least three probes were cut rectangular (100 mm x 224 20 mm), leaving initial grips separation (I_0) of 60mm and using cross-head speed of 100 225 mm/min. The tensile strength (TS, MPa) (break force/initial cross-sectional area) and elongation at break $[(I_{break} - I_0)/I_0] \times 100$, (EAB, %), were determined from the stress vs strain curves at the 226 227 breaking point, and the elastic modulus or Young's modulus (Y, MPa) calculated as the slope of the linear initial portion (elastic response zone) of the curve $(I_{break} - I_0)/I_0$. Puncture test: Films of 228 229 100X100 mm were fixed in a 35 mm diameter cell and punctured to breaking point with a round-230 ended stainless steel plunger (5 mm) at a cross-head speed of 100 mm/min, for breaking force (F, N), and breaking deformation (D, %) data according to Sobral et al.²⁹ which were carried out 231 232 at least in triplicate at room temperature and keeping the samples at 58% RH until the text 233 performance.

234 2.12. Statistical analysis

Statistical tests were performed using the SPSS computer programme (SPSS Statistical Software Inc., Chicago, Illinois, USA) for one-way analysis of variance. The variance homogeneity was evaluated using the Levene test, or the Brown-Forsythe when variance conditions were not fulfilled. Paired comparisons were made using the Bonferroni test or the Tamhane test (depending on variance homogeneity), with the significance of the difference set at $P \le 0.05$.

241 3. Results and Discussion

242 3.1. Extraction yield of seaweed hydrolysis

Carrageenan extraction yield was 28.65% (dry weight basis) and hydrolysate yields were 19.04% for H and 39.17% for Hp (dry weight basis); therefore total seaweed extraction yield by H and Hp hydrolysis was 47.69 and 67.82%, respectively. While H extraction resulted in a similar yield to another previously reported alcalase red seaweed hydrolysis, Hp was much
higher than almost all the protease extracts tested.¹⁸

Although carrageenan extraction is normally performed at high temperatures (80 °C),¹⁴ good extraction yields were obtained in the present using milder temperatures (50 °C), thereby demonstrating that these conditions could be suitable for extraction. Montolalu *et al.* indicated that the carrageenan obtained in extractions at 50 °C for long times (5 h) showed good gelling properties, being better than those obtained with shorter times.¹⁵ Therefore, enzymatic hydrolysis would allow concomitant extraction of bioactive compounds (hydrolysate) and carrageenan, improving the total yield and adding value to the seaweed extraction.

255 3.2. Mastocarpus protein hydrolysates

256 **3.2.1. Protein content and amino acid composition**

The protein content of marine algae varies greatly within species. Reports have shown that, in general, red seaweeds contain high levels of proteins.³⁰ The dried *Mastocarpus stellatus* contained $15.02 \pm 0.53\%$ of protein. Protein content was concentrated in the hydrolysates up to $37.86 \pm 1.07\%$ for H and $31.32 \pm 0.96\%$ for Hp.

The amino acid composition of H and Hp, expressed as residues per 1000 total amino acid 261 residues, is shown in Table 1. As expected, a similar amino acid profile was observed in both 262 263 hydrolysates. Both H and Hp showed high contents of Ser, Gly, Ala, Asp and Glu, and relatively 264 high contents of Leu, Thr, Val, Pro and Phe. The sum of the aspartic and glutamic acid contents 265 was 192 residues/1000 residues and 198 residues/1000 residues for H and Hp, respectively. The high acidic amino acid content is typical of red seaweeds.¹⁹ Nevertheless, some differences 266 between the hydrolysates were noteworthy. Some amino acids (Ser, Thr, Arg, His) were 267 268 concentrated in the more purified hydrolysate (H), owing to the removal of other amino acids, mainly hydrophobic residues (Ala, Val, Ile, Leu, Pro, Met). These hydrophobic amino acids 269 270 might have been extracted during the ethyl acetate extraction, suggesting that some of them 271 could be linked to the polyphenols extracted.

272 **3.2.2. ACE-inhibitory capacity**

Angiotensin-I converting enzyme (ACE) plays an important role in the regulation of blood pressure and hypertension, because it catalyses the conversion of inactive angiotensin-I into angiotensin-II, a potent vasoconstrictor, and inactivates bradykinin, a potent vasodilator.³¹ The amount of Mastocarpus hydrolysate required to inhibit 50% of the ACE activity (IC₅₀) is shown in Table 2.

278 Both H and Hp showed a high ACE-inhibitory capacity. The phenolic-partially-removed hydrolysate (H) showed considerably higher ACE-inhibitory capacity (IC₅₀ of 91 µg/mL) than the 279 280 phenolic-containing hydrolysate (Hp), probably owing to H's higher peptide concentration (7%). 281 Furthermore, some peptides of Hp might be interacting with polyphenols, therefore being less available for ACE binding. However, Jeon reported that, among seven flavourzyme enzymatic 282 283 digestions of brown seaweed, the hydrolysate with the highest polyphenol content showed the highest ACE-inhibitory capacity.³² Moreover, some polyphenolic compounds have been shown 284 to exert ACE-inhibitory activity.³³ ACE-inhibitory capacity might also be influenced by small 285 286 differences in the amino acid compositions of the hydrolysates. Peptide ACE-inhibitory activity 287 could be strongly influenced by the presence of hydrophobic (aromatic or branched side chains) amino acid residues at the C-terminal positions.^{31,34} The hydrophilic-hydrophobic partitioning in 288 the sequence is also a critical factor in the inhibitory activity.35 ACE inhibition is also highly 289 290 dependent on the molecular weight of peptides, those that are very short and have low molecular weight being more active.³⁶ 291

The IC₅₀ of the ACE-inhibitory capacity of the H hydrolysate was 17.5 times lower than the IC₅₀ value of the alcalase hydrolysate derived from red algae *Porphyra yezoensis*.³⁷ The ACEinhibitory capacity of algae hydrolysates has been reported in other works.^{34,38,39} Although, the use of a different method and its associated modifications to test ACE-inhibitory capacity makes direct comparison of IC₅₀ values difficult,²² H could be considered as a potent ACE-inhibitory hydrolysate.

298 **3.2.3. Folin-reactive substances and antioxidant activity**

Folin-reactive substances, ferric reducing power and ABTS radical scavenging ability of the hydrolysates are shown in Table 3. Both hydrolysates presented a noticeable amount of Folin301 reactive substances. As was expected, considering the method of hydrolysate preparation, the 302 Folin-reactive substances content was higher in Hp (phenolic-containing hydrolysate) than in H 303 (phenolic-partially-removed hydrolysate). Some polyphenols, however, might not have been 304 fully separated with ethyl acetate in the H hydrolysate. Moreover, although the Folin–Ciocalteu 305 assay is a widely used method to determine total phenolic content, additional substances can 306 react with the Folin reagent, including sugars and proteins, and should be taken into account.⁴⁰

The hydrolysis process would allow an improved extraction of phenolic compounds as well as the release of low molecular weight peptides,¹⁸ which contribute to enhance the antioxidant properties. Hp showed higher antioxidant activity than H (1.4 times higher for reducing power and 2.7 times higher for ABTS radical scavenging), probably owing to a greater presence of phenolic compounds in Hp. The positive correlation between the polyphenolic content of algae and their antioxidant activity has been well documented.⁴¹⁻⁴⁵

313 On the other hand, the hydrolysate peptide fraction can also contribute to antioxidant activity. It 314 is well known that biological activities of protein hydrolysates are related to the amino acid composition, sequence, molecular weight and peptide configuration. For example, 315 phosphorylated serine and threonine are known to bind metals,⁴⁶ being more hydrophilic and 316 317 reactive because of their hydroxyl group. Amino acids with non-polar aliphatic groups, such as 318 alanine, leucine or proline, have high reactivity to hydrophobic PUFA radicals, while hydrogen 319 donors such as aspartic and glutamic acids are able to quench unpaired electrons or radicals by 320 supporting protons.⁴⁷ The abundance of these amino acids in the peptide sequences of 321 hydrolysates could also be responsible for their antioxidant activity. As previously mentioned, the hydrophobic amino acid content was higher in Hp than in H, which might also have 322 323 contributed to the higher antioxidant capacity (ABTS and FRAP) of Hp compared with H.

Various studies have been carried out to evaluate the antioxidant potential of marine algae hydrolysates.^{7,8,18,48-50} However, to our knowledge, no reference has been made in previous studies to the antioxidant or ACE-inhibitory activity of Mastocarpus hydrolysates.

Given the Folin-reactive substances content and antioxidant activity results of the hydrolysates,
both hydrolysates could be considered potential antioxidants. However, because of its greater
potential, Hp was selected to develop active Mastocarpus films with antioxidant activity.

330 **3.3. Development of active films**

Increasing concentrations of hydrolysate (Hp) were added to Mastocarpus biopolymer filmforming solutions. Concentrations above 30% produced sticky, unmanageable films. For this reason, 30% was chosen as the maximum hydrolysate concentration that could be used for film development. A concentration of 15% hydrolysate was also chosen in order to maintain a balance between the film's physico-chemical properties and the active properties that could be provided by the hydrolysates.

337 3.3.1 FTIR-ATR

338 Infrared spectra of F-Hp0, F-Hp15 and F-Hp30 films and Hp freeze-dried Mastocarpus stellatus 339 hydrolysate were measured (Figure 1). Hp was analysed to assess its possible contribution to film structure. All film spectra showed a band at approximately 845 cm⁻¹ (C-O-S vibration), 340 341 assigned to D-galactose-4-sulfate (present in both K- and I-carrageenan), and a strong band at 342 924 cm⁻¹, which indicated the presence of 3,6-anhydro-D-galactose, a typical feature of κcarrageenan.^{51,52} The concomitant presence of κ - and ι -carrageenan features strongly 343 344 suggested a greater extraction of κ/i -hybrid carrageenan, as previously reported with M. stellatus.^{14,53} F-Hp0 and F-Hp15 spectra had similar band intensities, while F-Hp30 had the 345 346 above-mentioned bands considerably reduced, attributed to the reduced carrageenan amount in this film formulation in comparison with Hp, which had a much lower IR intensity at these wave 347 348 numbers. Although high temperature (80-90 °C) is adequate for suitable carrageenan 349 extraction, a certain amount of biopolymer might have been extracted during the hydrolysis 350 carried out at 50 °C. In this regard, Montolalu et al. reported an appreciable extraction yield of high molecular weight carrageenan at 50 °C in Kappaphycus alvarezii.¹⁵ The strong band at 351 1037 cm⁻¹ in Hp confirmed the predominantly polysaccharide nature of the hydrolysate. 352 353 Moreover, the greater absorption in the Hp IR-spectrum at wave numbers between 1100 and 354 1150 cm⁻¹ as compared to the films also suggested an increased proportion of shorter polysaccharide chains.⁵⁴ A small band at approximately 803 cm⁻¹ in the film spectra indicated 355 the presence of two sulfate ester groups on the anhydro-D-galactose residues (sulfation on C2), 356 characteristic and distinctive of I-carrageenan.^{51,55} This feature, which was not found in the 357 358 hydrolysate, was most prominent in F-Hp0, and became smaller with an increasing Hp amount in the film formulation (F-Hp0>F-Hp15>F-Hp30). Despite the presence of I-carrageenan, the *M.* stellatus film spectra were quite similar to κ -carrageenan standards, as previously shown by Gómez-Ordóñez and Rupérez.⁵² No evidence of a broad band between 820 and 830 cm⁻¹ was found, indicating the absence of highly sulfated λ -carrageenan. The second derivative spectra of the films revealed trace evidence at 871 cm⁻¹ of μ -carrageenan (κ -carrageenan precursor) and v-carrageenan (I-carrageenan precursor), which was not observable in Hp (data not shown).

The strong absorption bands at ~1216–1217 cm^{-1} (S = O), assigned to the presence of ester 366 sulfate groups, were noticeably lower in F-Hp30, coinciding with the highest proportion of added 367 hydrolysate. The 1217/924 cm⁻¹ ratio, as a measure of relative total sulfate groups with respect 368 to κ-carrageenan content, was much higher in Hp (2.6) than in any of the films studied (0.88 in 369 370 F-Hp0, 0.91 in F-Hp15 and 0.94 in F-Hp30), suggesting an additional source of sulfated compounds in Hp, probably phycobiliproteins with sulfur-containing amino acids.⁵⁶ In this 371 372 respect, Dumay et al. observed that enzymatic digestion was an effective treatment for phycoerythrin extraction.⁵⁷ Accordingly, the 1217/924 cm⁻¹ ratio became higher as a result of 373 374 the increase in the hydrolysate amount in the film formulation.

The IR-spectrum of Hp also revealed strong bands at 3277 cm⁻¹, 2929 cm⁻¹ and 1600 cm⁻¹ and 375 376 a small shoulder at 1518 cm⁻¹, which could be assigned, respectively, to amide A, amide B, 377 amide I and amide II of constituent proteins, most likely phycoerythrin and phycocyanin.58 378 Comparison of the films showed that as the added hydrolysate percentage increased the amide I amplitude became more evident, and it exhibited a wave number down-shift to 1631 cm⁻¹ in F-379 Hp30 as compared to 1639 cm⁻¹ in F-Hp0 and F-Hp15, which denoted more hydrogen bonding 380 381 in F-Hp30, attributed to the higher proportion of shorter peptides. In addition, the reduced band 382 intensity of the ester sulfate group in F-Hp30 and the slight frequency up-shift from 1216.1 cm⁻¹ 383 in F-Hp0 and F-Hp15 to 1217.5 in F-Hp30 could be indicative of appreciable carrageenan-384 peptide interactions in the film with the highest amount of added hydrolysate.

385 3.3.2. DSC

386 DSC thermograms of the freeze-dried Mastocarpus stellatus hydrolysate, Hp, and the F-Hp0, F-387 Hp15 and F-Hp30 films are shown in Figure 2. The hydrolysate showed two main endothermic peak temperatures, T_{peak} (°C), at 86.62 ± 6.22 and 130.16 ± 0.63, and corresponding ΔH (J/g_{dm}) 388 values of 3.51 ± 0.72 and 0.48 ± 0.05 , respectively, which might correspond to phycoerythrin 389 and phycocyanin fragments.⁵⁹ Temperatures were higher than those where protein normally 390 391 features, probably because hydrolysis might shift maximal peak temperature towards higher temperatures.⁶⁰⁻⁶² Low enthalpies also suggested the presence of hydrolysis products stabilized 392 393 by different amounts of hydrogen bonds and hydrophobic interactions. Another endothermic 394 transition in Hp with T_{peak} (°C) at 50.75 ± 0.66 and ΔH (J/g_{dm}) 0.39 ± 0.05 was evidence of the κ carrageenan constituent.60 395

396 DSC thermograms of F-Hp0, F-Hp15 and F-Hp30 are shown in Figure 2B. Slight endothermic 397 transitions were hardly observable, with T_{peak} values (°C) of 57.73 ± 0.20 in F-Hp0, 61.07 ± 2.14 398 in F-Hp15 and 64.65 \pm 1.86 in F-Hp30, and ΔH (J/g_{dm}) of 0.41 \pm 0.02, 0.19 \pm 0.01 and 0.11 \pm 399 0.04 respectively, indicative of the helix-to-coil transition suffered by the κ /i-carrageenan as a result of the breakage of weak physical cross-links.⁶³ Thermal transitions, however, were not as 400 sharp as those in a pure carrageenan curve.^{64,65} The increased protein content in films with 401 added hydrolysate might promote carrageenan-peptide interactions,⁶⁶ which could explain the 402 increase in T_{peak} temperatures. However, the enthalpy reduction suggested a hydrolysate-403 404 induced plasticizing effect in the films as a result of increasing the free water and chain mobility.

405 3.3.3. Rheology

406 Figure 3 shows the mechanical spectra of the film-forming solutions at 10 °C, in terms of elastic 407 modulus (G') and viscous modulus (G'') as a function of angular frequency. The F-Hp0 solution, 408 which had the lowest G' values, was the only one with a crossover point where G' = G''. At 409 frequencies below 2 Hz, the F-Hp0 solution was characterized by a dominant viscous behaviour 410 (G' < G'), which turned into a gel-like behaviour at higher frequencies; thus it could be classified 411 as a concentrated solution constituting an entanglement network. In the absence of KCI, 412 solutions of k-carrageenan cooled down to 9 °C have been shown to adopt helical structures, which did not aggregate to form self-supporting gels.⁶⁷ 413

414 In contrast, the F-Hp15 and F-Hp30 solutions showed a typical gel-like behaviour denoted by G' 415 > G" values within the whole frequency range, as previously reported in other studies on carrageenan.⁶⁸ The G' values were successfully modelled according to the power law ($r^2 \sim 0.99$) 416 417 in all three cases. The hydrolysate addition to the film-forming solution at 15% concentration 418 caused a remarkable increase in G', much higher than with 30%. The rheological behaviour of 419 the F-Hp15 solution showed lower frequency dependence of G' than the F-Hp30 and F-Hp0 420 solutions, as deduced from the lowest power law exponent value (n') (0.63 in F-Hp0, 0.40 in F-421 Hp15, 0.52 in F-Hp30). Hp concentrations higher than 30% conferred sticky, unmanageable 422 properties to the films. All these findings suggest that the hydrolysate added at the appropriate 423 concentration led to stronger gels with increased structural stability, probably due to hydrolysate 424 components (mainly peptides and phenolic compounds) favouring aggregation of carrageenan 425 helices to form a three-dimensional network. Interactions between carrageenan and proteins have previously been shown to produce much stronger gels than single carrageenan gels.⁶⁶ 426 427 Similarly, the ability of polyphenols to interact with polysaccharides forming complexes has been well documented.⁶⁹ Nevertheless, the higher hydrolysate amount in the F-Hp30 solution 428 429 considerably reduced the gel-forming capacity with respect to the F-Hp15 solution, with the 430 helical aggregates probably having more difficulty in being created as a result of a carrageenandilution effect.70 431

The apparent viscosity of the film-forming solutions, measured at 25 °C and shear rate of 0.5 s^- ¹, was considerably higher in the F-Hp15 solution (14.89 ± 0.53 Pa·s) than in the F-Hp0 and F-Hp30 solutions (3.47 ± 0.01 Pa·s and 4.72 ± 0.15 Pa·s, respectively), strongly suggesting effective interactions at the right concentration between carrageenan and other compounds naturally present in the hydrolysate, presumably peptides and phenolic compounds.

437 3.3.4. Light barrier properties

Colour parameters, L^* (lightness), a^* (reddish/greenish) and b^* (yellowish/bluish), are shown in Table 4. All the films were quite similar, having low lightness (28–29) and slightly greenish and yellowish tendencies. The F-Hp30 film exhibited the highest (P≤0.05) lightness and greenish colouration, and lowest (P≤0.05) yellowish tendency. Changes in $L^*a^*b^*$ values, however, did not correlate with increasing amounts of added hydrolysate in the film, which could be due to a different degree of interactions between protein pigments and carrageenan. Comparing these results with previously developed commercial κ -carrageenan films, the present *M. stellatus* films presented considerably lower lightness and more red tendency, owing to the concomitant extraction of non-carrageenan compounds.⁷¹⁻⁷³

447 In general, the films exhibited low light transmission in the UV range (250-300 nm) (0-1.12%) (Figure 4), as compared to commercial κ -carrageenan films,⁷⁴ with F-Hp0 providing the least 448 efficient UV barrier. Two absorption peaks were defined in all the films in the ranges 400-450 449 450 nm and 600-700 nm, which might be associated with the presence of pigments, such as carotenoids and chlorophyll, which absorb at 400-450 (violet-blue-green colours), and 451 phycoerythrin and phycocyanin at 600 nm (red colour).⁷⁵ In the visible range, the light 452 transmission was significantly (P≤0.05) lower in F-Hp30, especially in the wavelength range 453 454 between 350 and 700 nm, which might be largely due to the increase in thickness associated 455 with the hydrolysate addition, as Table 5 shows. The hydrolysate contained small molecules 456 (mainly peptides and oligosaccharides) that might have interfered in carrageenan helix 457 aggregation during the film drying process. This interference might have caused a plasticizing 458 effect with an increase in free volume that would have resulted in thicker films.

459 3.3.5. Physico-chemical properties

Slight variations in moisture content were observed among the three film formulations (Table 5), with F-Hp30 showing slightly higher values, which could be related to its increased thickness. The protein content in the films increased significantly ($P \le 0.05$) with the addition of increasing amounts of hydrolysate (Table 5).

464 **3.3.5.1. Water barrier**

No significant differences in film water solubility were found in *M. stellatus* films with either 15 or 30% added hydrolysate (Table 5). A similar finding was reported earlier in gelatin films incorporating different percentages of gelatin hydrolysate.⁷⁶ Although the solubility values were not high, the films totally lost their original structure, becoming a very viscous solution with gelling tendency at low temperatures. Solubility was similar to previous results obtained in commercial carrageenan films.⁷³ No significant (P \leq 0.05) differences were found between F-Hp0 and F-Hp30 water vapour permeability (*WVP*) (Table 5). In contrast, F-Hp15 had the lowest permeability, probably owing to effective carrageenan-protein interactions, as previously commented. Despite the greater thickness of F-Hp30, *WVP* was not reduced by adding 30% hydrolysate. The extra protein and plasticizer effect caused by Hp addition may have resulted in a less dense network.⁷⁷ The present films were more water vapour permeable than previously reported commercial carrageenan films.^{71-73,78,79}

Water resistance test results are shown in Figure 5. Noticeable differences among samples were observed after 10 minutes. Although every film elongated up to 2 cm until breakage, F-Hp0 showed a faster elongation speed (3.2 cm/h) than F-Hp15 and F-Hp30 (~2.9 cm/h). The hydrolysate addition led to a significantly higher breakage resistance in F-Hp30 in comparison with F-Hp0 and F-Hp15, probably related to the higher carrageenan peptide interactions. No water leaking before film breakage could be detected in any sample.

484 3.3.5.2. Mechanical properties

485 F-Hp0 had the significantly highest (P \leq 0.05) tensile strength (TS) (Table 5), which was lowest in 486 F-Hp30. The opposite behaviour was found regarding the elongation at break (EAB) values. 487 confirming the hydrolysate-induced plasticizing effect in the film. The TS and EAB values in the 488 three *M. stellatus* films studied were, respectively, higher and lower than the results reported with commercial k-carrageenan or I-carrageenan films,^{73,79} suggesting a reinforcement effect 489 490 caused by the presence of other non-carrageenan components. As far as Young's modulus (Y) is concerned (Table 5), the highest stiffness also corresponded to F-Hp0 (P≤0.05), decreasing 491 492 with increasing amount of Hp. The small molecules (mainly peptides and oligosaccharides) that 493 form part of the hydrolysate have been proved to act as film plasticizers by preventing 494 carrageenan helix associations and increasing the molecular mobility of polymer chains, which 495 in the case of F-Hp30 was favoured by the increased water plasticizing effect. Similarly, 496 Salgado et al. also observed a reduction in TS and Y and an increase in EAB in protein films 497 with added hydrolysate, which, in view of the lack of film moisture increase, was attributed to interferences in protein cross-linking.²¹ 498

499 There were no significant differences in puncture force (F) between F-Hp0 and F-Hp15 (Table 500 5), whereas F-Hp30 had higher values ($P \le 0.05$). Puncture deformation (D) was also significantly 501 higher (P≤0.05) in F-Hp30, with no differences between F-Hp0 and F-Hp15. It is worth noting 502 the different information provided by both type of mechanical tests. The high plasticizing effect 503 exerted by the hydrolysate in F-Hp30, mostly by preventing carrageenan helix associations, was 504 the main factor determining the film tensile properties; however, this film showed higher 505 resistance to perforation, which could be attributed to some carrageenan-protein interactions. 506 Tensile strength gives more general information about the film resistance to traction, while 507 puncture test focuses on the resistance to perforation in a specific matrix area.

508 **3.3.6.** Antioxidant activity and Folin-reactive substances of the films

Folin reactive substances, ferric reducing power and ABTS radical scavenging capacity of the films are shown in Table 3. Films without algae hydrolysates (F-Hp0) contained Folin reactive substances and exhibited some antioxidant activity measured by both FRAP and ABTS assays. The incorporation of increasing concentrations of Hp significantly increased the Folin-reactive substances content (3-fold increase in F-Hp30), as well as FRAP and ABTS values (3- and 9fold increase, respectively, in F-Hp30).

515 The antioxidant properties of squid gelatin films and sunflower films were also improved by the 516 addition of hydrolysates from squid gelatin and bovine plasma, respectively, but the antioxidant 517 activity increase reported was much lower than in the present work.^{21,76}

518 4. CONCLUSION

519 Enzyme-assisted Mastocarpus stellatus hydrolysis could be a complementary way to extract 520 bioactive components in addition to carrageenan. The more purified peptide hydrolysate 521 (phenolic content partially removed) showed the highest ACE-inhibitory capacity; while keeping 522 peptides and polyphenols together resulted in a more antioxidant hydrolysate, which was 523 suitable as active ingredient for antioxidant film development. The addition of the hydrolysate 524 resulted in a more plasticized film by reducing the tensile strength and increasing the elongation 525 at break, however, the film was more resistant to perforation. In view of the results obtained, 526 these films might be used as a complementary material for active packaging development.

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669	Table	1. Amino	acid	composition	of Mastocar	pus hy	/drolysates	(H,	Hp)
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Amino acid	Number of residues / 1000 residues		
	Н	Нр	
Asp	102	105	
Thr	73	66	
Ser	153	92	
Glu	90	93	
Gly	112	116	
Ala	104	120	
Cys	5	7	
Val	47	65	
Met	16	19	
lle	34	40	
Leu	72	88	
Tyr	28	28	
Phe	43	43	
His	12	6	
Lys	29	26	
Arg	27	23	
Pro	48	57	
Нур	0	0	
Hyl	5	5	

672 Table 2. ACE-inhibitory capacity of Mastocarpus hydrolysates

		C72
		673 IC₅₀* (μg/mL) 674
	н	91.62 ± 2.44 a 75
	Нр	148.32 ± 3.16 6 76
677	Different	t letters (a, b) indicate significant differences (p≤0.05).
678	* IC ₅₀ : co	oncentration (µg/mL) required to inhibit 50% of ACE ad

679

* IC₅₀: concentration (μ g/mL) required to inhibit 50% of ACE activity.

680 **Table 3.** Antioxidant activity and Folin reactive substances of hydrolysates and films

Sample	ABTS (mg Vit C eq/g)	FRAP (µmol Fe/g)	Folin reactive substances (mg/g)
н	35.95 ± 1.59 a	84.52 ± 1.38 a	36.02 ± 3.26 a
Нр	93.26 ± 2.55 b	106.19 ± 1.05 b	75.61 ± 0.56 b
F-Hp0	3.07 ± 0.18 a	4.54 ± 0.08 a	7.33 ± 0.34 a
F-Hp15	17.56 ± 0.90 b	11.77 ± 0.38 b	15.97 ± 1.46 b
F-Hp30	27.51 ± 0.83 c	13.75 ± 0.06 c	22.17 ± 0.36 c

681 Results are the mean ± standard deviation. One-way ANOVA: Different letters indicate

682 significant differences among the different hydrolysates (H) or different films (F) (P≤0.05).

684 **Table 4.** L^* , a^* , b^* and Transparency ($-\log(T_{600}/x)$) of F-Hp0, F-Hp15 and F-Hp30

Film	L*	a*	b*	Transparency
F-Hp0	28.65 ± 0. 29 a	-0.57 ± 0.03 a	4.57 ± 0. 11 a	7.14 ± 0.24 a
F-Hp15	27.99 ± 0.08 b	-0.30 ± 0.05 b	4.72 ± 0. 04 b	6.61 ± 0.30 ab
F-Hp30	29.25 ± 0.05 c	-0.70 ± 0.03 c	4.36 ± 0.04 c	6.12 ± 0.12 b

685 Results are the mean ± standard deviation. One-way ANOVA: Different letters indicate

686 significant differences among the different films ($P \le 0.05$).

Table 5. Thickness, moisture, protein content, film solubility, water vapour permeability (*WVP*),

689	tensile strength (TS), elongation at break (EAB), Young's modulus (Y), puncture force (F) and
690	puncture deformation (D) of F-Hp0, F-Hp15 and F-Hp30

	F-Hp0	F-Hp15	F-Hp30
Thickness (µm)	51.82 ± 3.34 a	56.05 ± 3.65 a	68.36 ± 4.04 b
Moisture (%)	13.46 ± 0.28 a	12. 13 ± 0. 14 b	14.59 ± 2.75 ab
Protein content (%)	8.90 ± 0.06 a	10.98 ± 0.21 b	13.05 ± 0.24 c
Film solubility (%)	20.97 ± 4.5 a	25.77 ± 3.64 a	22.16 ± 2.95 a
WVP (x10 ⁻⁸ g m ⁻¹ s ⁻¹ Pa ⁻¹)	3. 78 ± 0. 17 a	3. 20 ± 0. 12 b	4.04 ± 0. 30 a
<i>TS</i> (MPa)	59.94 ± 2.27 a	51.37 ± 3.75 b	41.63 ± 2.95 c
EAB (%)	0.95 ± 0.11 a	1.59 ± 0.09 b	2.47 ± 0.24 c
Y (MPa)	1797 ± 61 a	1347 ± 74 b	1054 ± 45 c
<i>F</i> (N)	23.47 ± 1.08 a	26.36 ± 2.14 b	30.38 ± 2.27 b
D (%)	7.73 ± 0.47 a	8.61 ± 0.53 a	12.24 ± 1 b

691 Results are the mean ± standard deviation. One-way ANOVA: Different letters indicate

692 significant differences among the different films (P≤0.05).

693







Fig. 2



Fig. 3



Fig. 4



