

1 **Gastrointestinal modifications and bioavailability of brown seaweed phlorotannins and**
2 **effects on inflammatory markers.**

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4 Giulia Corona^{1,2*}, Yang Ji², Prapaporn Aneboonlap², Sarah Hotchkiss³, Chris Gill⁴, Parveen
5 Yaqoob², Jeremy P.E. Spencer² and Ian Rowland².

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8 ¹Health Sciences Research Centre, University of Roehampton, London SW15 4JD, UK.

9 ²Department of Food and Nutritional Sciences, University of Reading, Reading RG6 6AP, UK,

10 ³CyberColloids Ltd., Carrigaline Industrial Estate, Carrigaline, County Cork, Ireland

11 ⁴Northern Ireland Centre for Food & Health, University of Ulster, Coleraine, BT52 1AA

12

13

14 *Correspondence to: Dr Giulia Corona, Health Science Research centre, Life Sciences
15 Department, Whitelands College, University of Roehampton, Holybourne Avenue, London,
16 SW15 4JD

17 Tel: +44 (0)20 8392 3622, fax +44 –(0)208392 3610, email giulia.corona@roehampton.ac.uk

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25 **Abstract:**

26 Brown seaweeds such as *Ascophyllum nodosum* are a rich source of phlorotannins (oligomers
27 and polymers of phloroglucinol units), a class of polyphenols that are unique to **Phaeophyceae**.
28 At present there is no information on the bioavailability of seaweed polyphenols and limited
29 evidence on their bioactivity *in vivo*. Consequently we investigated the gastrointestinal
30 modifications *in vitro* of seaweed phlorotannins from *Ascophyllum nodosum* and their
31 bioavailability and effect on inflammatory markers in healthy participants. *In vitro*, some
32 phlorotannin oligomers were identified after digestion and colonic fermentation. In addition 7
33 metabolites corresponding to *in vitro* absorbed metabolites were identified. Urine and plasma
34 samples contained a variety of metabolites attributed to both un-conjugated and conjugated
35 metabolites (glucuronides and/or sulfates). In both urine and plasma, the majority of the
36 metabolites were found in samples collected at late time points (6-24 h), suggesting colonic
37 metabolism of high molecular weight phlorotannins, with three phlorotannin oligomers
38 (hydroxytrifuhalol A, 7-hydroxyeckol, C-O-C dimer of phloroglucinol) identified in urine
39 samples. A significant increase of the cytokine IL-8 was also observed. Our study shows for
40 the first time that seaweed phlorotannins are metabolized and absorbed, predominantly in the
41 large intestine, and there is a large inter-individual variation in their metabolic profile. Three
42 phlorotannin oligomers present in the capsule are excreted in urine. Our study is the first
43 investigation of the metabolism and bioavailability of seaweed phlorotannins and the role of
44 colonic biotransformation. In addition IL-8 is a possible target for phlorotannin bioactivity.

45

46 **Introduction:** There has been increasing interest in the past few years on the bioactive
47 compounds present in seaweeds ^(1; 2; 3). Traditionally, seaweeds are consumed as a food product
48 in Asian countries and are increasingly used worldwide as **ingredients** for industrial
49 applications. **In Japan, over 20 species of red, green, and brown algae (seaweed) are included**
50 **in meals ⁽⁴⁾, and daily seaweed consumption per person has remained relatively consistent over**
51 **the last 40 years, in the range of 1.50 to 3.65 kg/person/year as reported by a range of studies**
52 **^(5; 6; 7). Seaweeds are a rich source of polyphenolic compounds ⁽⁸⁾, and polyphenols extracted**
53 **from algae ^(9; 10) show some similarities to those found in land plants ^(9; 10; 11). Thus the main**
54 **polyphenols found in brown seaweeds are phlorotannins ^(12; 13; 14; 15), a type of phenolic**
55 **compound only found in brown seaweeds ⁽¹⁶⁾. Brown seaweed phlorotannins are oligomers**
56 **and polymers of phloroglucinol units, and their oligomer and polymer molecular weights can**
57 **greatly vary, from 126 Da to 650 kDa ⁽³⁾, comprising up to 15 % of the plant dried weight ⁽¹¹⁾.**
58 **It has been reported that the consumption of brown algae is on average 1.342 kg/person/year,**
59 **containing 66.652 g of phlorotannins/person/year and 183 mg/person/day ^{(4) (4)}. Phlorotannins**
60 **are classified according to the type of linkages between phloroglucinol units into four main**
61 **groups: eckols (with dibenzodioxin linkages), fucols (with a phenyl linkage), fuhalols and**
62 **phloroethols (with ether linkages), and fucophloroethols (with ether and phenyl linkages) ⁽¹⁶⁾.**
63 **Phlorotannins are being increasingly investigated for their vast array of bioactivities ^(10; 17; 18)**
64 **such as antioxidant ^(19; 20; 21; 22; 23; 24), anti-inflammatory ^(20; 25; 26), antibacterial ^(27; 28), anticancer**
65 **^(29; 30; 31; 32; 33), and antidiabetic ^(29; 34; 35), showing promising potential to further develop**
66 **seaweed-derived products rich in bioactive components with commercial potential for food and**
67 **pharma applications ⁽³⁶⁾.**
68 **Bioavailability is a critical factor influencing *in vivo* biological activity of polyphenols and we**
69 **have reasonable understanding of the bioavailability of polyphenols from fruits and vegetables,**
70 **and some of the mechanisms by which they exert beneficial effects *in vivo* have been**
71 **determined ⁽³⁷⁾. Their ability to act as effective bioactive molecules *in vivo* is dependent on the**
72 **extent of their biotransformation ⁽²⁴⁾ and conjugation during absorption from the**
73 **gastrointestinal (GI) tract, in the liver and finally in cells ⁽³⁷⁾. Consequently, consideration must**
74 **be given to the way polyphenols are absorbed and metabolised during gastrointestinal digestion**
75 **and colonic fermentation and how this may impact on bioactivity ⁽³⁸⁾. It is noteworthy that there**
76 **is no information on the bioavailability of seaweed phlorotannins and this is a limitation to**
77 **understanding their bioactivity and mechanism of action *in vivo*. In the absence of specific data**
78 **regarding phlorotannin absorption and bioavailability, it is useful to consider the absorption**
79 **and metabolism of other polyphenols as a guide ⁽³⁷⁾. In general, after ingestion of a polyphenol-**

80 rich diet, their protective effects *in vivo* are determined by measuring a range of suitable
81 biomarkers, and correlate with the absorption of polyphenols from the gut and their
82 circulation/excretion ⁽³⁸⁾. Polyphenols can be extensively conjugated to form glucuronide,
83 sulphate and methyl group in the gut mucosa and inner tissues ^(37; 39), and absorption occurs in
84 the small intestine ⁽³⁷⁾. Polyphenols unabsorbed in the upper gastrointestinal tract or re-
85 excreted in the bile, are extensively metabolised by colonic microflora into a wide range of low
86 molecular weight phenolic acids ⁽⁴⁰⁾. The aim of this study was to elucidate the gastrointestinal
87 modifications of seaweed phlorotannins, and the effects on their metabolism and
88 bioavailability. A food grade seaweed polyphenol extract (SPE) rich in phlorotannins (from
89 the brown seaweed *Ascophyllum nodosum*) was subjected to *in vitro* gastrointestinal digestion
90 and fermentation to examine the gastrointestinal modifications occurring in the upper- and
91 lower- GI tract. Furthermore, the absorption and metabolism of polyphenols in healthy subjects
92 was investigated, after oral ingestion of a SPE capsule containing 101.89 mg of polyphenols.
93 This amount represents an intake lower than the average daily intake of seaweed polyphenols
94 in the Asian diet, and it is not expected to exert any cytotoxic effect ⁽⁴¹⁾. The impact of
95 absorption and gastrointestinal modifications on phlorotannins anti-inflammatory potential is
96 explored.
97

98 **Experimental methods**

99

100 **Seaweed material.** Fresh *Ascophyllum nodosum* was supplied by The Hebridean Seaweed
101 Company, Isle of Lewis, Scotland in March 2011. **The seaweed biomass was harvested by**
102 **hand, cleaned and then** shipped refrigerated to the processing facility in France where it was
103 immediately chopped and frozen.

104

105 **Preparation of food-grade SPE and capsule.** **A novel SPE from *Ascophyllum nodosum* was**
106 **produced by CEVA (France) using a solvent based extraction system that was specifically**
107 **developed for this study and for use with either fresh or frozen *Ascophyllum nodosum*. The**
108 **solvent used was a 60:40 ethanol:water mixture which allowed for the water content of the**
109 **seaweed itself. The extraction was carried out over 5 hours using constant stirring and at all**
110 **times protected from light. A solvent:seaweed ratio of 3:1 was used. The mixture was filtered**
111 **to remove the supernatant and subsequently the alcohol was removed using a rotary evaporator.**
112 **A hydrometer was used to check that all the alcohol had been removed. The final extract was**
113 **recovered by centrifugation and further filtration before freeze drying.**

114 Approximately half of the produced extract (**basic extract**) was then fractionated using
115 tangential flow ultra filtration to produce further extracts of varying molecular weight range
116 and with varying polyphenol content. A blended **SPE** was formulated (Table 1) using **175 mg**
117 of basic extract and **50 mg** of **high molecular weight (HMW) fraction** (>10 kDa cut off) for use
118 in the current study. Maltodextrin (**175 mg**) was added to the capsule formulation as an
119 excipient. This was done in order to maximise the polyphenol content (>**100 mg** per day) but
120 also to minimise the level of iodine to within accepted regulatory guidelines (< **500 µg** per day).
121 Blending was carried out at the food grade CEVA facilities in France. **400 mg** doses of the
122 **SPE *Ascophyllum* blend** were packed into white, opaque, vegetarian capsules by Irish
123 Seaweeds, Belfast, UK and used for the clinical study. The food grade seaweed capsule was
124 characterized by NP-HPLC and LC-MS analysis. Phlorotannins were quantified using the
125 Folin-Ciocalteu Method ⁽⁴²⁾ using phloroglucinol as the standard ⁽⁸⁾.

126

127 **Simulated Gastro-Intestinal Digestion and Fermentation:** The Gastro-Intestinal Digestion
128 procedure was adapted from Mills *et al.* (2008) and McDougall *et al.* (2005). This method
129 consists of two sequential stages: gastric digestion and small intestinal digestion followed by
130 dialysis. 10 g of **SPE** was dissolved in 30 ml acidified water (pH=2) and pepsin (320 U/ml)
131 was added. Samples were incubated at 37 °C for 2 h on a shaker covered with foil to protected

132 from light. 5 ml aliquots (G) were removed. The pH was adjusted to 7.5 by adding few drops
133 of 6 M NaOH, and pancreatin (4 mg/ml) and bile extracts (25 mg/ml) were added. The samples
134 were incubated at 37 °C for 2 h on a shaker. 5 ml aliquots (SI) were removed. Samples were
135 transferred into the dialysis tubing (100-500 Da, cut-off, 1.8 ml/cm, Spectra/Por, Biotech) and
136 dialysed overnight at 4 °C against water (4 L) to remove low molecular weight digests. 5 ml
137 aliquots of dialysis solution (D1) were removed. The dialysis fluid was changed and dialysis
138 continued for additional 2 h. 5 ml aliquots of second dialysis solution (D2) were removed.
139 Samples (SI+D) were freeze-dried and subjected to colonic fermentation (Batch Culture): The
140 method was adapted from Tzounis *et al.* (2008). Batch-culture fermentation vessels (300 ml;
141 one vessel per treatment) were autoclaved and filled with 135 ml sterilized basal medium.
142 Medium was stirred and gassed overnight with O₂-free N₂. Before addition of SI+D digested
143 extracts equivalent to 1.5 g of undigested extracts, the temperature inside the vessels was set to
144 37 °C by a circulating water bath and the pH was controlled at 6.8 by an Electrolab pH
145 controller, in order to mimic conditions in the distal region of the human large intestine
146 (anaerobic; 37 °C; pH 6.8). Vessels were inoculated with 15 ml faecal slurry (1:10, w/v) and
147 batch cultures were run for 24 h 7 ml samples were collected at five time points (0, 2, 4, 8 and
148 24 h), centrifuged at 13,000 rpm at 4 °C for 10 min and the supernatants were kept. All the
149 samples collected during the digestion and fermentation procedure were stored at -80 °C until
150 LC-MS analysis.

151

152 **Study design:** This study was conducted according to the guidelines laid down in the
153 Declaration of Helsinki and all procedures involving human subjects/patients were approved
154 by the University of Reading Ethics Committee before initiation of the study. Written informed
155 consent was obtained from all participants. Exclusion criteria for subjects were: smokers, BMI
156 < 18 or > 30, abnormal liver function and haematology, alcohol intake of > 21 units/wk,
157 gastrointestinal disease or chronic gastrointestinal disorders, consumption of antibiotics in
158 previous 3 months before study, females who were pregnant or intending to become pregnant.
159 Potentially suitable participants underwent a screening process and individuals with blood
160 pressure > 150/90 mmHg; Hb > 125 g/l for men and > 110 g/l for women; g-glutamyl
161 transferase > 1.3 mkat/l or cholesterol > 6.5 mmol/l were excluded from the study. In total 24
162 volunteers were recruited - 12 females (6 aged 18-30 years and 6 aged 30-65 years) and 12
163 males (6 aged 18-30 years and 6 aged 30-65 years). Participants were asked to follow a low
164 phenolic diet for 1 day prior to the study day (devoid of tea, coffee, fruit, vegetables, alcoholic
165 beverages, cocoa, whole- grain and seaweed-containing products). On the study day, the

166 subjects were cannulated and a baseline blood sample was taken. Participants were asked to
167 consume one **SPE** capsule (**400 mg**) containing **101.89 mg** of polyphenols. Blood samples were
168 collected at 0, 1, 2, 3, 4, 6, 8 and 24 hours after ingestion of the SPE capsule, and urine samples
169 were collected at baseline, 0-8 and 8-24 h after the ingestion. During the day, participants were
170 provided with a lunch and dinner of low phenolic content. The study is registered at
171 clinicaltrials.gov (study ID: NCT02496806).

172

173 **Sample collection and storage:** One aliquot of blood was collected in heparin tubes and
174 cultured immediately (whole blood culture for cytokine analysis). One aliquot of blood was
175 collected in EDTA tubes and centrifuged at 3000 rpm for 15 min at 4°C. The plasma was
176 separated and 1 mg/ml ascorbic acid was added as preservative. Aliquots were stored at -80 °C
177 until analysis. Total volume of collected urine was recorded and aliquots were stored at -80 °C
178 until analysis.

179

180 **Plasma sample processing for metabolite analysis:** Plasma samples were prepared
181 following a procedure similar to the one described by Ottaviani *et al.* ⁽⁴³⁾. **10 µl** of internal
182 standard solution (resorcinol **200 µg/ml**) were added to **450 µl** of plasma, then **50 µl** of **1.2 M**
183 acetic acid were added and samples were mixed. Samples were analysed with and without
184 enzymatic treatment (**37 °C**, 40 min) in the presence of 1500 IU of β-glucuronidase and 50 IU
185 of sulfatases from *Helix pomatia* (Type H-1). **1 ml** of **100 %** methanol acidified with **0.5 %**
186 acetic acid was added and samples were centrifuged for **15 min** at **16,100 x g** at **4 °C** and
187 supernatants were collected. This step was repeated 3 times (last time with **50 %** methanol
188 acidified with **0.5 %** acetic acid) and the supernatants were dried using a Speedvac. The pellets
189 were dissolved with **125 µl** of mobile phase and transferred to vials for RP-HPLC analysis.

190

191 **Urine sample processing for metabolite analysis:** Urine samples were prepared following a
192 procedure similar to the one described by Ottaviani *et al.* ⁽⁴³⁾. Briefly, **10 µl** of internal standard
193 solution (resorcinol **200 µg/ml**) were added to **250 µl** urine. Samples were analysed with and
194 without enzymatic treatment (**37 °C**, 40 min), in the presence of 1500 IU of β-glucuronidase
195 and 50 IU of sulfatases from *H. pomatia* (Type H-1). **1 ml** of **100 %** methanol acidified with
196 **0.5 %** acetic acid was added, samples were mixed and centrifuged for **15 min** at **16.100 g** (rcf)
197 at **4 °C**, supernatants were transferred to a new tube and dried on a Speedvac. Dried samples
198 were re-suspended on **125 µl** of mobile phase, completely dissolved, centrifuged and
199 transferred to vials for RP-HPLC analysis and LC-MS analysis.

200

201 **NP-HPLC analysis:** The phlorotannins in the food-grade **SPE** used to produce the capsule
202 were analyzed by normal phase HPLC analysis ⁽⁸⁾ using an HPLC 1100 series equipped with
203 LiChrospher Si60-5 column (250 mm × 4.0 mm ID, 5 µm particle size from HICHROM
204 (LISP60-5-250AF), fitted with a guard column LiChrospher Si60-5 from HICHROM (LISP60-
205 5-10C5). The mobile phase contained A: **82 %** dichloromethane + **2 %** methanol + **2 %** acetic
206 acid in water and B: **96 %** methanol + **2 %** acetic acid in water and was pumped through the
207 column at 1 ml/min. 10 µl of samples were injected and analyzed by the gradient program
208 which were (min/%B): 0/0, 30/17.6,45/30.7, 50/87.8, 60/87.8, 80/0, 105/0 for detection of all
209 compounds. The compounds were detected at a wavelength of 268 nm. All data were analyzed
210 by ChemStation software. The phloroglucinol standard was injected at 0.1-100 µg/ml and
211 phlorotannins in the capsules were analysed as phloroglucinol equivalents.

212

213 **RP-HPLC analysis:** The analysis of plasma and urine samples was carried out with a Hewlett-
214 Packard 1100 series liquid chromatograph (Hewlett-Packard, Palo Alto, CA) as previously
215 described ⁽⁴⁴⁾. Samples were analyzed by reverse-phase HPLC using a Nova-Pak C18 column
216 (4.6 x **250 mm**) with 4 µm particle size. The temperature of the column was maintained at 30
217 °C. The mobile phases consisted of a mixture of aqueous methanol 5 % in 0.1 % hydrochloric
218 acid **5 M** (A) and a mixture of aqueous acetonitrile 50 in 0.1 % hydrochloric acid **5 M** (B) and
219 were pumped through the column at 0.7 ml/min. The following gradient system was used (min
220 / % B): 0/5, 5/5, 40/50, 55/100, 59,9/100, 60/5, with 10 min post-run for both compound and
221 metabolite detections. The eluent was monitored by photodiode array detection at **280 nm** and
222 spectra of products obtained over the 200–600 nm range. Peaks were characterized by their
223 retention time and spectra characteristics. A calibration curve of phloroglucinol was
224 constructed using authentic standards (0.1–100 µg/ml) and in each case were found to be linear
225 with correlation coefficients of **0.995**. Metabolites were quantified as phloroglucinol
226 equivalents.

227

228 **LC-MS analysis:** LC-MS analysis was conducted to analyse the food grade seaweed capsule,
229 the urine samples and the digested materials, and was carried out in the negative ion mode
230 using LC-MS/MS utilizing electrospray ionisation (ESI) as previously described ⁽⁴⁵⁾.
231 Characterization was achieved using LC-MS/MS utilizing electrospray ionisation (ESI). This
232 consisted of an Agilent 1200 HPLC system equipped with a binary pump, degasser, auto-
233 sampler, thermostat, column heater, photodiode array detector and an Agilent 1100 Series

234 LC/MSD Mass Trap Spectrometer. Separation of samples was achieved using a Zorbax SB
235 C18 column (2.1 x 100 mm; 1.8 μ m) (Agilent) and HPLC conditions were as follows: injection
236 volume: 1 μ L; column temperature: 25 $^{\circ}$ C; binary mobile system: (A) 0.1 % aqueous formic
237 acid and (B) 0.1 % of formic acid in acetonitrile; flow rate: 0.2 mL/min. A series of linear
238 gradients were used for separation (min/%B): 0/10, 3/10, 15/40, 40/70, 50/70, 65/10. Mass
239 spectrometry was performed in the negative ion mode (scan range, m/z 100-1000 Da; source
240 temperature, 350 $^{\circ}$ C). All solvents used were LC-MS grade.

241

242 **Cytokine production:** Blood samples collected during the clinical intervention (baseline, 1 h,
243 2 h, 4 h, 6 h, 8 h) into heparin tubes, were immediately cultured as follows: 500 μ l blood
244 aliquots were mixed with 500 μ l of RPMI media containing antibiotics on a 24 well plate and
245 LPS (1 μ g/ml) or vehicle (control group) were added before incubation at 37 $^{\circ}$ C for 24 h. At
246 the end of the culture period, samples were centrifuged at 2000 x g for 5min and the
247 supernatants were collected and kept at -20 $^{\circ}$ C until analysis for inflammatory cytokine levels.
248 The supernatants were collected and stored at -20 $^{\circ}$ C. Cytokines (IL-1 β , IL-6, IL-8, IL-10 and
249 TNF- α) in the supernatants were measured with Luminex xMAP Technology using
250 commercially available Fluorokine MAP kits (R&D systems) and data analyzed on the
251 xPONENT software. Final data are presented as the difference between LPS-treated and un-
252 stimulated control.

253

254 **Statistical analysis:** The statistical evaluation of the results was performed by one-way
255 analysis of variance (ANOVA) followed by a Bonferroni post-hoc t-test using GraphPad InStat
256 version 5 (GraphPad Software, San Diego, CA, USA). Significant changes are indicated as P
257 < 0.05.

258

259 **Results**

260

261 **SPE characterization:** The chromatogram (Figure 1) illustrates the trace obtained by NP-
262 HPLC with diode array detection after injecting a water solution of the SPE. The chromatogram
263 shows a number of peaks (20-70 min) representing different high molecular weight
264 phlorotannins, the characteristic phenolics in brown seaweeds. Longer phlorotannin polymers,
265 which consisted of more hydroxy groups, resulted in tighter attachment to the column material.
266 Consequently, shorter compounds were released earlier than longer compounds. Owing to a
267 lack of phlorotannin standards and the complexity of the oligomeric and polymeric forms, the
268 calibration curve of phloroglucinol was used to quantify all the phlorotannins contained in the
269 SPE as phloroglucinol equivalents. The SPE comprised a wide range of molecular weight of
270 phloroglucinol derivatives with a total phlorotannin concentration of 312µg/mg quantified as
271 phloroglucinol equivalents. Further characterization of the SPE was achieved with LC-MS/MS
272 (Figure 2) utilizing electrospray ionisation (ESI). The data were collected in a non-targeted
273 fashion, by acquiring full spectrum data in negative ion mode from m/z 100 to 1000. The data
274 were then analysed by searching for the theoretical mono-isotopic masses corresponding to
275 possible phlorotannin oligomers and the presence of ions (1-6) which could correspond to
276 phlorotannins (Figure 2). The ion 1 with [M-H]⁻ at m/z 405 corresponded to the trimer
277 hydroxytrifuhalol A, whereas the MS² fragment -387 corresponded to the loss of one molecule
278 of water (-18), a characteristic pattern of phlorotannin fragmentation. Compound 2 ([M-H]⁻ at
279 m/z 497) was considered to be a phlorotannin tetramer composed of 4 phloroglucinol units,
280 such as tetrafucol or fucodiphlorethol and also in this case the fragment -479 corresponds to
281 the loss of a molecule of water (-18), whereas the fragment -353 corresponds to the loss of
282 water (-18) and a phloroglucinol unit (-126), in accordance with analytical profiles recently
283 described in positive ion mode by Wang *et al.* ⁽⁴⁶⁾ and by Ferreres *et al.* ⁽⁴⁷⁾. The ion 3 has a
284 [M-H]⁻ at m/z 247 corresponding to a C-O-C dimer of phloroglucinol as previously indicated
285 by Nwosu *et al.* ⁽²⁹⁾. The ion 4 (387) corresponds to the trimer 7-hydroxyeckol, and we observe
286 the presence of a fragment at -369 deriving from the loss of one molecule of water. Isomers 5
287 and 6 with [M-H]⁻ at m/z 249 were also observed, which can correspond to the dimers
288 diphlorethol and difucol.

289 **In vitro digestion and characterisation:** Due to the lack of commercially available standards
290 for phlorotannins and the complexity of the oligomers and polymers in the extract, the analysis
291 of phlorotannins and their metabolites is challenging and requires a combination of approaches.
292 Similarly to other polyphenol classes, phlorotannins may undergo extensive modification by

293 phase I and phase II enzymes and the colonic microbiota during their transit through the
294 gastrointestinal tract⁽³⁷⁾, and the implication of such metabolic modifications on the bioactivity
295 of phlorotannins has not been investigated yet. Consequently we subjected the **SPE** to *in vitro*
296 digestive and fermentative processes. An *in vitro* gastric and ileal digestion and colon microbial
297 fermentation of the **SPE** was initially conducted, followed by dialysis to simulate absorption
298 into the circulation. The MS spectra and fragmentations of the compounds detected in the
299 samples were studied (Figure 3). After *in vitro* digestion and fermentation procedures, the
300 samples were analysed by LC-MS/MS utilizing electrospray ionisation (ESI) in negative ion
301 mode as previously described, searching for the theoretical mono-isotopic masses
302 corresponding to the low molecular weight phlorotannins previously identified in the capsule
303 (Figure 2). We were able to identify molecular ions and fragments corresponding to
304 hydroxytrifuhalol A (405), the C-O-C dimer of phloroglucinol (247), the dimer diphlorethol/
305 difucol (249) and 7-hydroxyeckol (387), also found after colonic fermentation. In addition, in
306 digested and fermented samples subjected to dialysis to mimic absorption into the circulation,
307 we reported the presence of 7 compounds (DM1 to DM7) corresponding to *in vitro* absorbed
308 metabolites.

309

310 **Plasma and urine analysis:** The food grade SPE was given to healthy subjects (Figure 4) in
311 the form of a capsule (400 mg) containing 101.89 mg of polyphenols, and blood and urines
312 were analysed for phlorotannin metabolites. HPLC-DAD analysis of the plasma (Figure 5) and
313 urine (Figure 6) samples with and without glucuronidase/sulfatase treatment showed the
314 presence of a variety of metabolites absent in the baselines (before the seaweed capsule
315 ingestion) in samples from 15 volunteers out of 24. **In plasma, the total level of phlorotannins
316 and their metabolites ranges from 0.011 to 7.757 µg/ml, and in urine from 0.15 to 33.52 are
317 excreted.** Some metabolite peaks were present in samples with and without enzymatic
318 treatment, and therefore could be assigned to un-conjugated metabolites. Some other
319 metabolite peaks were present only in samples without enzymatic treatment or were only
320 appearing in samples enzymatically treated, and were attributed to conjugated forms
321 (glucuronides and/or sulfates) and their enzymatically released un-conjugated forms. In urine,
322 some metabolites were found in samples collected at 0-8 h after capsule ingestion, but the
323 majority of the metabolites were found in samples collected at 8-24 h. Some metabolites, such
324 as UM6 and UM7, show similar UV spectra characteristics and might therefore be structurally
325 related (Figure 6). In plasma (Figure 5) some metabolites were found in samples collected at
326 2, 3 and 4 h after capsule ingestion, but the majority of the metabolites were found in samples

327 collected at later time-points (6-24 h). This could be the result of poor absorption of the high
328 molecular weight phlorotannins in the upper gastrointestinal tract resulting in them reaching
329 the colon and undergoing fermentation to lower molecular weight derivatives by the colonic
330 microbiota. In addition, urine samples were subjected to LC-MS/MS (Figure 7) utilizing
331 electrospray ionisation (ESI) as previously described, searching for the theoretical mono-
332 isotopic masses corresponding to the low molecular weight phlorotannins previously identified
333 in the capsule (Figure 2). We were able to identify molecular ions and fragments corresponding
334 to hydroxytrifuhalol A, 7-hydroxyeckol and the C-O-C dimer of phloroglucinol, which
335 corresponded to the HPLC metabolite UM3. In addition, we reported the presence of 2 ions
336 (289 and 377) corresponding to metabolites that we characterized in samples from SPE
337 subjected to *in vitro* gastrointestinal digestion and fermentation (DM4 and DM7, figure 3) as
338 previously detailed.

339

340 ***Ex-vivo* Cytokine production:**

341 The ex-vivo production of cytokines (IL-1 β , IL-6, IL-8, IL-10 and TNF- α) relative to baseline
342 levels in cultured blood collected at various time-points (0, 1, 2, 3, 4, 6 and 8 h) during the
343 intervention study (LPS treated – unstimulated controls) is reported in Fig 8. The amounts of
344 TNF- α and IL-10 remained quite stable over time, as well as the amount of all cytokines at 1h
345 and 2h. IL-6 levels were observed to decrease at later time-points (4 h to 8 h) without reaching
346 statistical significance ($p > 0.05$). The levels of both IL-1 β and IL-8 were observed to increase
347 from 3 h to 8 h after the intervention; however the statistical analysis revealed that the only
348 significant change from baseline levels was the increase of IL-8 at 8 h.

349

350 Discussion

351 Polyphenols are ubiquitously found in plants and comprise a major part of a daily human diet.
352 Over the last 20 years, significant data have emerged with regard to the potential beneficial
353 effects of several classes of phenolic compounds against a number of chronic diseases. In
354 addition, a reasonable understanding has been gained of the bioavailability of many polyphenol
355 classes and this will be important for understanding the mechanisms by which they exert such
356 benefits *in vivo*. The interest in marine sources of phenolic compounds is recent and knowledge
357 on phlorotannin bioavailability is still lacking. The analysis of phlorotannins is challenging due
358 to high range of molecular weight present, and their characterisation is complicated further by
359 the lack of commercially available standards.

360 Chromatography techniques coupled to diode array and MS detection have been applied to the
361 analysis of phlorotannins and the advantages/disadvantages of their use described by Steevenz
362 *et al* ⁽³⁴⁾. Reversed phase High Performance Liquid Chromatography (RP-HPLC) is a
363 separation mode commonly used for polyphenols separation, however the very high polarity
364 of phlorotannins would causes them to elute with little or no retention due to the lack of
365 interaction with the non-polar stationary phase ^(34; 48). Normal phase liquid chromatography
366 (NP-HPLC) is more advantageous for retaining compounds with very high polarity, and the
367 NP-HPLC methodology developed by Koivikko was more suitable than RP-HPLC for the
368 separation of phlorotannins from the brow algae *Fucus vesiculosus* ^(34; 48). Thus we initially
369 analysed the phlorotannins in our SPE by Normal-Phase HPLC using a method adapted from
370 Koivikko *et al* ⁽⁸⁾. As expected, the SPE comprised a wide range of molecular weights (20-70
371 min), with abundance of very high molecular weight phlorotannins eluting at later retention
372 time (50-70min) in our Normal-Phase method. MS detection can provide higher sensitivity
373 and be advantageous to identify specific phlorotannins in the extract without commercially
374 available standards, however NP-HPLC solvents such as dichloromethane are not suitable for
375 MS analysis ⁽⁴⁹⁾, because they would result in poor ionisation and therefore significantly reduce
376 sensitivity ^(48; 50).

377 Nwosu *et al.* characterized a phlorotannin extract from *Ascophyllum nodosum* by RP-HPLC
378 using a C18 column the bound sample eluted in a unresolved set of peaks, and with MS
379 detection in negative ion mode they were able to assign the detected m/z spectra to a series of
380 phlorotannin structures ⁽²⁹⁾. Ferreres *et al.* identified 22 different phlorotannins belonging to
381 the eckol and fucophloroethol groups in four seaweed species belonging to the order Fucales
382 (genus *Cystoseira* and *Fucus*) with RP-HPLC separation combined with DAD-ESI-MSn
383 detection ⁽⁴⁷⁾. Using an equivalent RP-HPLC separation method coupled to ESI-MS analysis

384 in negative ion mode we were able to identify some phlorotannin oligomers such as
385 hydroxytrifuhalol A, tetrafucol, fucodiphlorethol, C-O-C dimer of phloroglucinol, 7-
386 hydroxyeckol, diphlorethol and difucol. The fragmentation patterns of the oligomers we
387 identified are in agreement with some recent published data in the field ^(46; 47). Recently,
388 Steevensz and his research group characterized the phlorotannins of five brown algae species
389 by ultrahigh-pressure liquid chromatography operating in “mixed-mode” (hydrophilic
390 interaction liquid chromatography mode) combined with high resolution mass spectrometry
391 ⁽³⁴⁾. The methodology proposed by this research group proved to be accurate for profiling
392 phlorotannins based on their degree of polymerization, and was demonstrated to be an effective
393 separation mode for low molecular weight phlorotannins, up to **6 KDa** ⁽³⁴⁾.

394 Phlorotannin characterization is a challenging and complex task, complicated by the lack of
395 commercially available standard compounds, thus chromatography separation coupled to MS
396 detection can help to elucidate phlorotannin complexity and its application to the analysis of
397 clinical samples from feeding trials, as well as the use of simplified *in vitro* digestion systems,
398 can help elucidate their beneficial health properties and the bioactive circulating forms. The
399 **SPE** was subjected to *in vitro* simulated gastrointestinal digestion and fermentation, followed
400 by dialysis to simulate as close as possible their absorption and biotransformation. The obtained
401 materials were analysed by LC-MS for a comparative characterisation of the phlorotannin
402 metabolites. LC-MA analysis of the digested and fermented **SPE** have indicated the presence
403 of some oligomeric phlorotannins also present in the undigested **SPE** (hydroxytrifuhalol A,
404 diphlorethol/difucol, 7-hydroxyeckol, C-O-Cdimer of phloroglucinol), in addition to a range of
405 newly formed metabolites (DM1 to DM7). *In vitro* conditions are indeed a great tool, allowing
406 a simpler and more convenient analysis, and our *in vitro* system predicted the formation of
407 metabolites subsequently identified in urine.

408 Intervention studies have investigated the fate of polyphenols from land plants in the human
409 body by measuring plasma concentrations and/or urinary excretion following intake from a
410 food source. Many studies performed to investigate polyphenol bioavailability are based on
411 the measurement of their excretion in urine and plasma by extraction, concentration and
412 chromatographic separation/analysis, and focused on the detection of polyphenols and their
413 metabolites in samples subjected or not to enzymatic treatment to release conjugate moieties
414 such as glucuronic acid and sulphate groups ^(43; 51). For example, following ingestion of a
415 polyphenol-rich meal, levels of phenolic compounds and conjugated metabolites can increase
416 rapidly achieving a peak concentration at approximately **1-2 h** in plasma and urine, indicating
417 small intestinal absorption, or peak at later time-points (**4-8 h** in plasma, **8-24 h** in urine),

418 indicating large intestinal metabolism and subsequent absorption ⁽⁵²⁾. In our study, the majority
419 of phlorotannin metabolites were found in samples collected at late time-points (6-24 h),
420 indicating limited small intestinal absorption followed by gut microbial metabolism, of the high
421 molecular weight phlorotannins in the large intestine.

422 In the upper gastrointestinal tract, dietary polyphenols act as substrates for a number of
423 enzymes and are subjected to extensive metabolism by glucosidase enzymes, phase I enzymes
424 (hydrolysing and oxidizing), such as cytochrome P450, and phase II enzymes (conjugating and
425 detoxifying) found both in the small intestine and the liver ⁽³⁷⁾. In particular there is strong
426 evidence for the extensive phase II metabolism (by UDP-glucuronosyltransferases,
427 sulphotransferases) to yield glucuronides and sulphate derivatives. Indeed, there is evidence of
428 efficient glucuronidation and sulfation of all classes of polyphenols to differing extents ⁽⁵³⁾.
429 Indeed, our results indicate that phlorotannin intake results in the formation of phase II
430 conjugate metabolites (glucuronides, sulfates).

431 Further transformations can occur in the colon, where the enzymes of the gut microbiota act to
432 breakdown complex polyphenolic structures to smaller units, which may also be absorbed and
433 further metabolized in the liver. Bacterial enzymes may catalyse many reactions including
434 hydrolysis, dehydroxylation, demethylation, ring cleavage and decarboxylation as well as rapid
435 de-conjugation ⁽⁵⁴⁾.

436 As predicted by the high molecular weight range of phlorotannins in our **SPE**, high colonic
437 metabolism seems to have occurred, following fermentation of high molecular weight
438 phlorotannins in the large intestine. By LC-MS analysis, we were able for the first time to
439 confirm the presence in urine of some phlorotannin oligomers, more specifically
440 hydroxytrifuhalol A, 7-hydroxyeckol and the C-O-C dimer of phloroglucinol. Interestingly, 2
441 of the urine metabolites (m/z 289 and 377) were present in the *in vitro* digested samples (DM4
442 and DM7).

443 There were substantial differences between volunteers in the pattern of phlorotannin
444 metabolites. Such inter-individual differences have been observed for other polyphenols and
445 have been attributed to differences in gut microbiota composition and the expression of
446 metabolizing enzymes ^(3; 55).

447 A secondary aim of our work has been to determine whether the **SPE** could modulate
448 inflammatory events in the blood, following the absorption of phlorotannin metabolites and
449 due to their presence into the circulation.

450 Polyphenols can exert numerous antioxidant and non-antioxidant functions of relevance in
451 chronic diseases development, and many of them have an important inflammatory component

452 ⁽⁵⁰⁾. In the present study, we observed an altered *ex vivo* production of IL-8, a low-molecular-
453 weight cytokine produced by mononuclear phagocytes and other cell types, with significant
454 increased levels of the cytokine after 8h compared to baseline.

455 IL-8 is an important inflammatory factor of the CXC chemokine family, involved in the
456 amplification of inflammatory signals ⁽⁵⁶⁾. IL-8 secretion is induced by TNF-alfa through a
457 transcriptional mechanism primarily regulated by nuclear factor-kB (NF-kB) ⁽⁵⁷⁾. Redox
458 signalling mechanisms are known to play a role in the regulation of such events, with reactive
459 oxygen species being able to promote IL-8 production and secretion ^(18; 23), whereas oxygen
460 radical scavengers are proven to inhibit IL-8 production in LPS-stimulated human whole blood
461 ⁽²²⁾. Dietary polyphenols such as catechins ⁽³³⁾ and curcumin ⁽²⁸⁾ have also been shown to
462 specifically interfere with IL-8 gene expression through inhibition of NF-kB activation ⁽⁵⁸⁾. In
463 consequence, we would have expected circulating seaweed polyphenol metabolites to
464 potentially be able to inhibit IL-8 secretion. Our results have given a preliminary indication
465 that the cytokine IL-8 is a possible target for phlorotannin metabolites. However a significant
466 increase on IL-8 levels at 8h after the intervention was observed, in parallel with the presence
467 of phlorotannin metabolites in plasma and urine samples. A recent study from our group
468 investigated the influence of a polyphenol-rich intervention on inflammation as primary
469 outcome. A randomised, double-blind, placebo-controlled, cross-over acute intervention was
470 conducted, and cytokine levels (IL-8) were measured with the same *ex-vivo* experimental
471 protocol. The results showed a time-dependent increase in IL-8 release compared to baseline,
472 in accordance with our findings. Thus the post-prandial *ex vivo* IL-8 production was
473 significantly attenuated by the intervention compared to the control, with a parallel appearance
474 in the circulation of polyphenol metabolites. Our trial was a single group interventional study
475 primarily designed to investigate the bioavailability of seaweed phlorotannins: however, on the
476 basis of this preliminary indication on their anti-inflammatory potential, not sufficient to draw
477 any conclusion, a chronic placebo-controlled intervention has been conducted to investigate
478 the anti-inflammatory effect in **deeper** detail.

479 The main limitations of this study arise from the phlorotannin complexity and lack of
480 commercially available analytical standards, potentially leading to possible quantification
481 inaccuracy as phloroglucinol equivalents. The lack of analytical standards also implies a
482 limited capability for method development, especially for the analysis of plasma, urine and
483 digested materials. In future, the availability of standards could allow a higher degree of
484 method optimization and the development of specific solid phase ion procedures for sample
485 clean-up and concentration.

486 The development of more recently explored analytical applications to phlorotannins, such as
487 HILIC ⁽³²⁾ and NMR ⁽³⁵⁾ could facilitate the development of more suitable protocols that
488 could lead to full identification of metabolites and improvements in phlorotannin **metabolites**
489 **quantification. In addition, The bioaccessibility of polyphenols in the form of a capsule/extract**
490 **might differ greatly from the bioaccessibility of the same compounds in a food matrix ⁽⁵⁹⁾. Future**
491 **work will be needed to determine the potential effects on bioavailability of different food**
492 **matrices and also any effects of cooking and/or processing.**

493 Nevertheless, the present work has for the first time started to shed light on the role of colonic
494 biotransformation on phlorotannin bioavailability, and its implication for their health benefits
495 *in vivo*. Our results provide a basis for further investigating the seaweed-derived bioactive
496 components in the body after ingestion, information which is necessary to understand their
497 mechanism of action *in vivo*.

498

499 **List of tables:**

500 **Table 1.**

Table 1: Key components of polyphenol rich basic extract, High Molecular Weight (HMW) fraction, and blend (capsule) showing crucial concentrations of polyphenols and iodine.

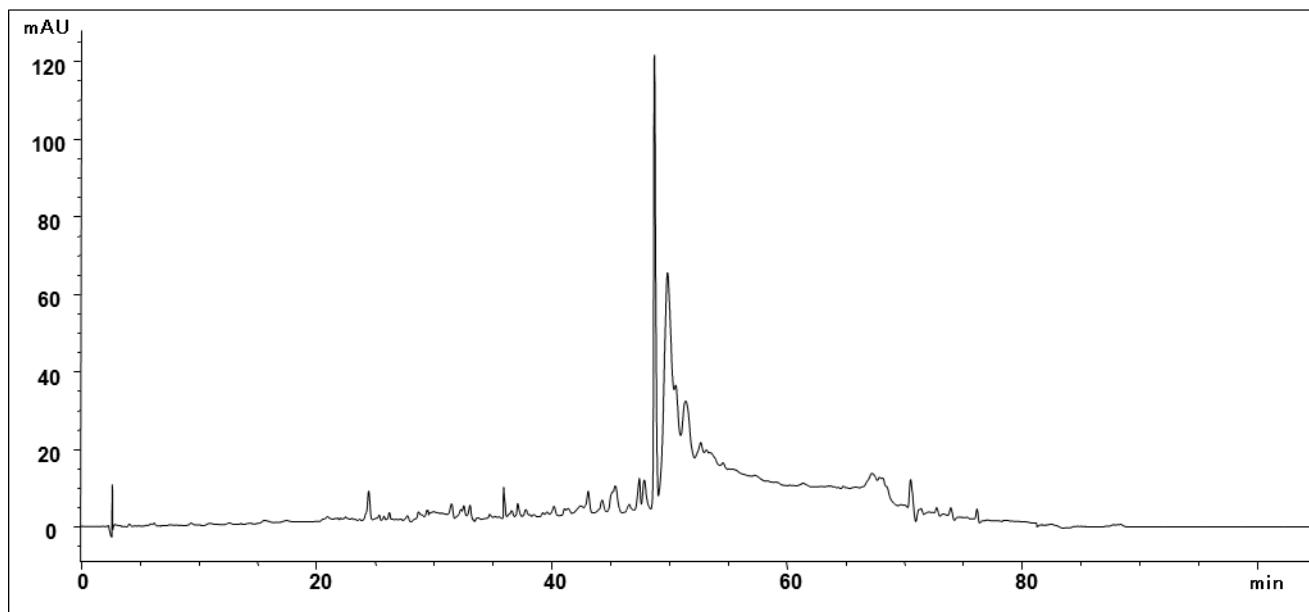
Extract component	basic extract mg/175mg of extract	HMW fraction mg/50mg of extract	blend (capsule) mg/400mg capsule
Polyphenols	58.74	43.15	101.89
Iodine	0.46	0.02	0.48
Maltodextrin*			175
Minerals	37.77	1.22	38.99
Fucoxanthin	<0.001	0.004	0.004
Laminarin as glucose	10.24	1.64	11.88
Fuoidan as fucose	<0.001	0.23	0.23
Mannitol	28.03	5.53	33.56
Inorganic arsenic	<0.001	<0.001	<0.001
Cadmium (LD 0.15mg/kg)	<LD	<LD	<LD
Mercury (LD 0.016mg.kg)	<LD	<LD	<LD
Lead (LD 1.1mg/kg)	<LD	<LD	<LD
Tin (LD 1.7mg/kg)	<LD	<LD	<LD

*maltodextrin was added to the capsule formulation as an excipient.

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502

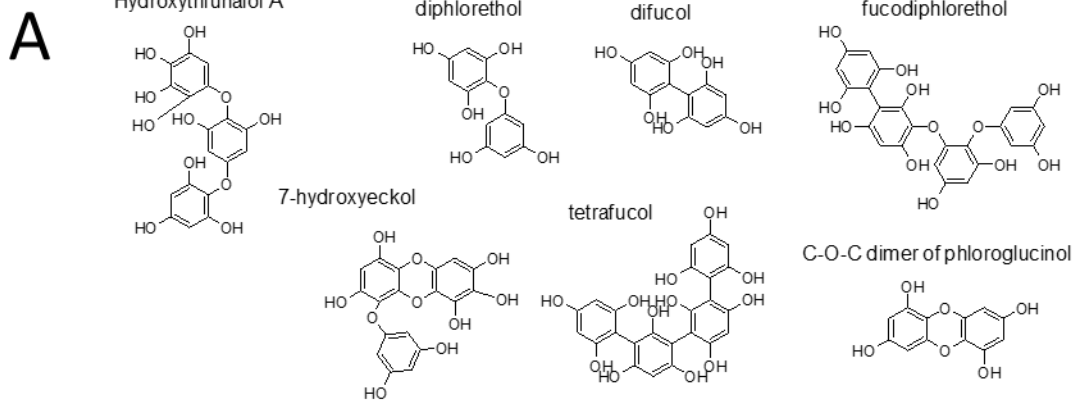
503 **Figure 1**



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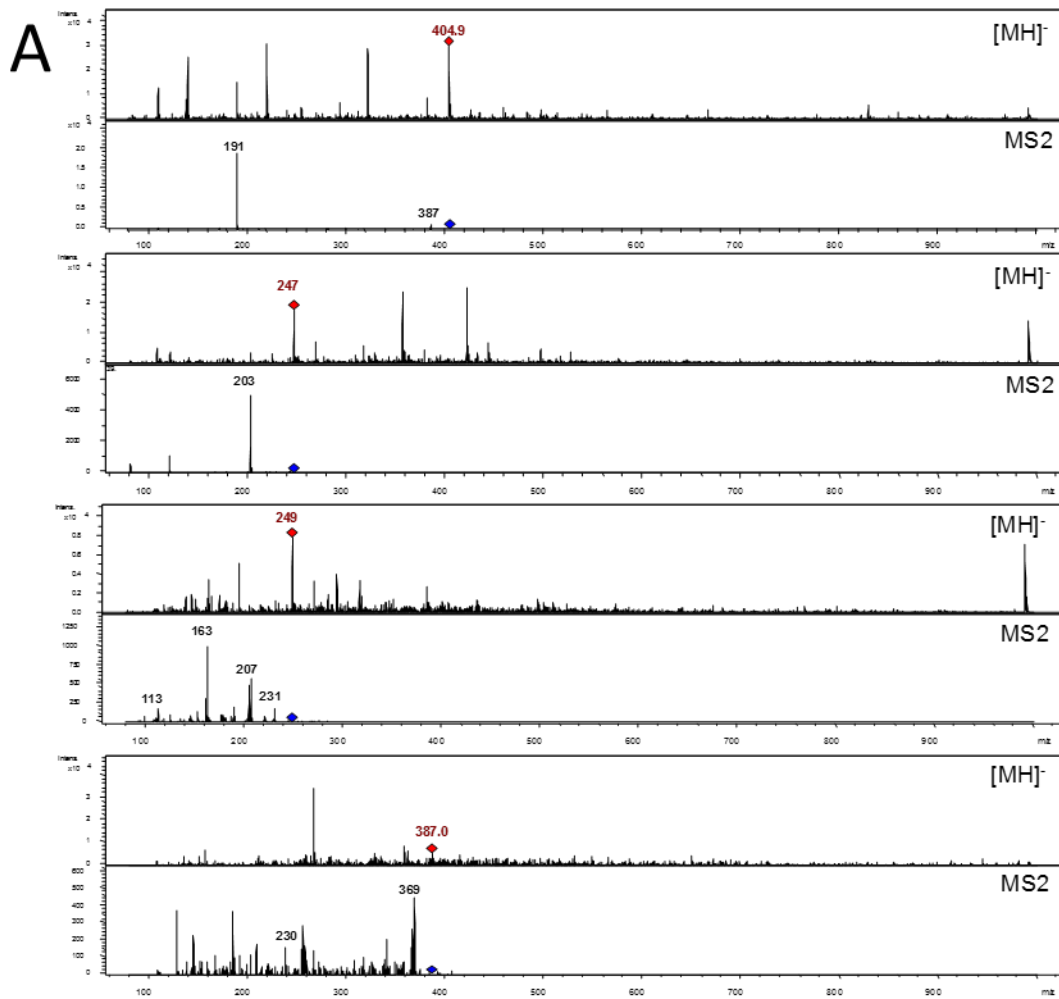
B

LC-MS analysis in negative ion mode of the seaweed extract phlorotannins

Peak N	RT (min)	[M-H] ⁻ m/z	MS ² m/z	tentative identification
1	2.7	405	387 191	hydroxytrifufahalol A
2	3.3	497	479 353 205	tetrafucol, fucodiphlorethol
3	5.6	247	203	C-O-C dimer of phloroglucinol
4	10.6	387	369 230	7-hydroxyeckol
5	43.7	249	181 113	diphlorethol / difucol
6	46.2	249	181 113	diphlorethol / difucol

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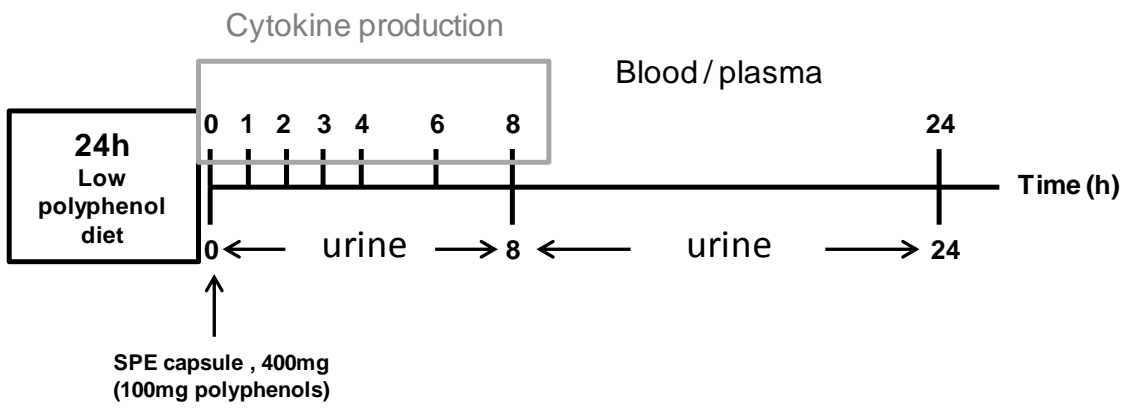
B LC-MS analysis in negative ion mode of the in vitro digested seaweed extracts

Metabolite	RT (min)	[M-H] ⁻ (m/z)	MS2 (m/z)			tentative identification	
<i>seaweed extract after small intestinal digestion</i>							
SIM1	2.70	405	387	191		hydroxytrifusalol A	
SIM2	5.00	247	203			C-O-C dimer of phloroglucinol	
SIM3	5.70	249	231	207	163	113	diphlorethol / difucol
SIM4	9.30	387	369	230			7-hydroxyeckol
<i>seaweed extract after colonic fermentation</i>							
FM1	9.40	387	369	230			7-hydroxyeckol
<i>dialysed seaweed extract (after digestion and fermentation).</i>							
DM1	1.50	377	311	243	175		Unknown
DM2	1.60	277	211	151	93		Unknown
DM3	1.65	249	181	113			Unknown
DM4	1.70	289	261	221	175		Unknown
DM5	1.75	317	249	225	181	113	Unknown
DM6	1.80	327	269	211	93		Unknown
DM7	1.85	377	341	281	221	179	Unknown

511

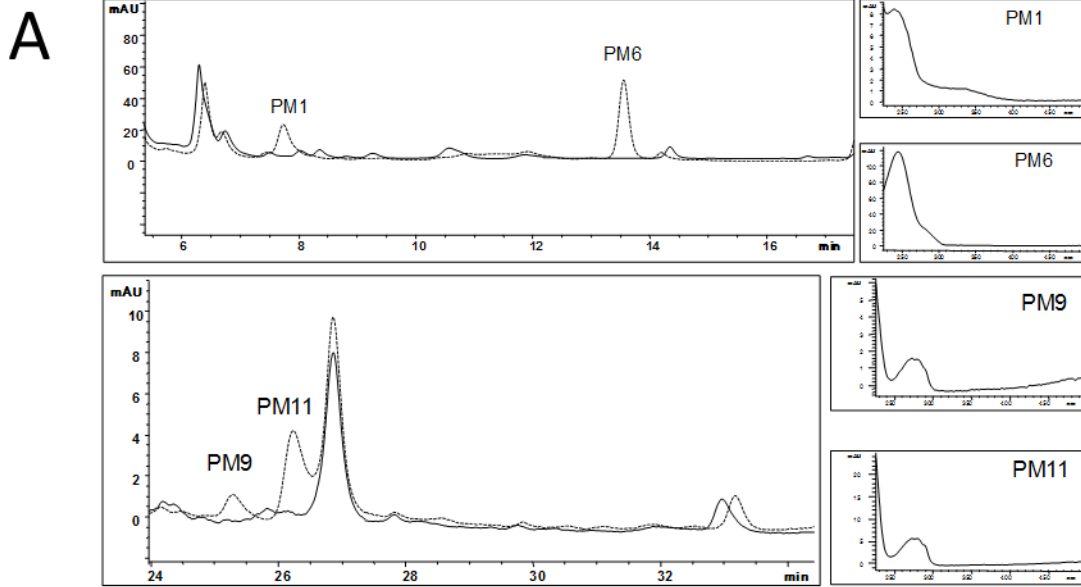
512

513 **Figure 4**



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B

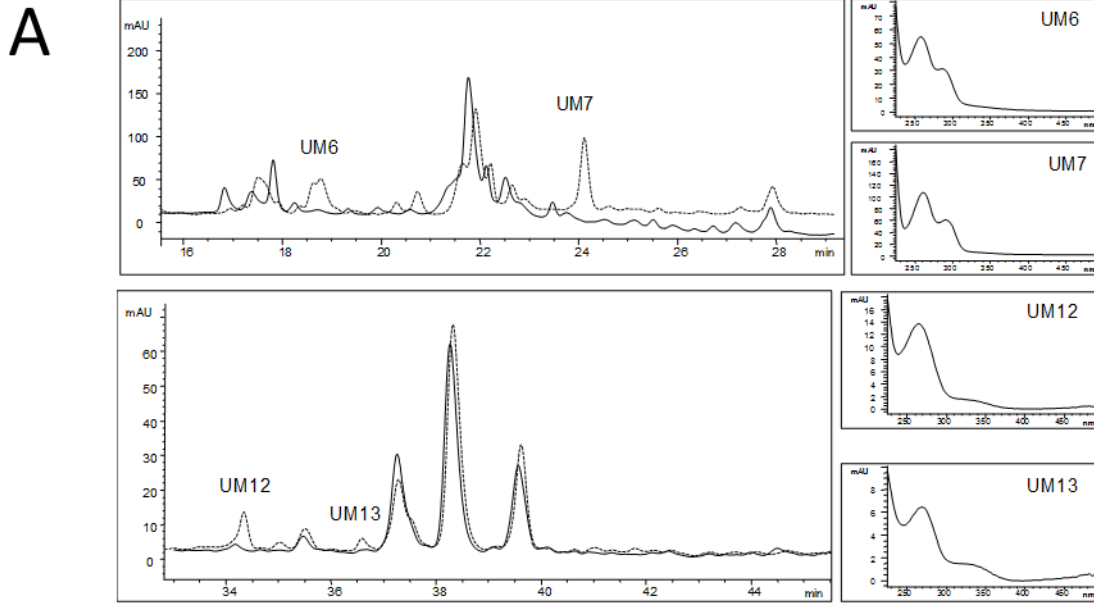
RP-HPLC analysis of seaweed metabolites in plasma samples

RT (min)	metabolite	mean (µg/ml)	stdev	N volunteers	time (h)	Enz -	Enz +
6.5	PM1	2.16	2.01	2	4; 24	no	yes
7.7	PM2	0.95		1	24	yes	no
9.2	PM3	0.07		1	8	yes	no
9.3	PM4	0.07	0.03	2	2; 3	yes	no
9.4	PM5	0.05	0.03	7	3; 4; 6; 8; 24	yes	no
13.5	PM6	2.82	1.85	2	24	yes	yes
20.4	PM7	0.03		1	3	no	yes
23.2	PM8	0.03		1	24	no	yes
25.2	PM9	0.06	0.01	2	8; 24	yes	no
25.5	PM10	0.01	0.00	2	24	yes	yes
26.2	PM11	0.22		1	24	yes	no
30.3	PM12	0.41		1	24	yes	no
32.7	PM13	0.05	0.04	3	24	yes	yes
33.2	PM14	0.04		1	4	yes	no
36.1	PM15	0.01		1	2	yes	no
38.0	PM16	0.02	0.01	3	2; 3	yes	no
42.4	PM17	0.04		1	4	yes	no
44.7	PM18	0.03	0.03	5	6; 24	yes	yes

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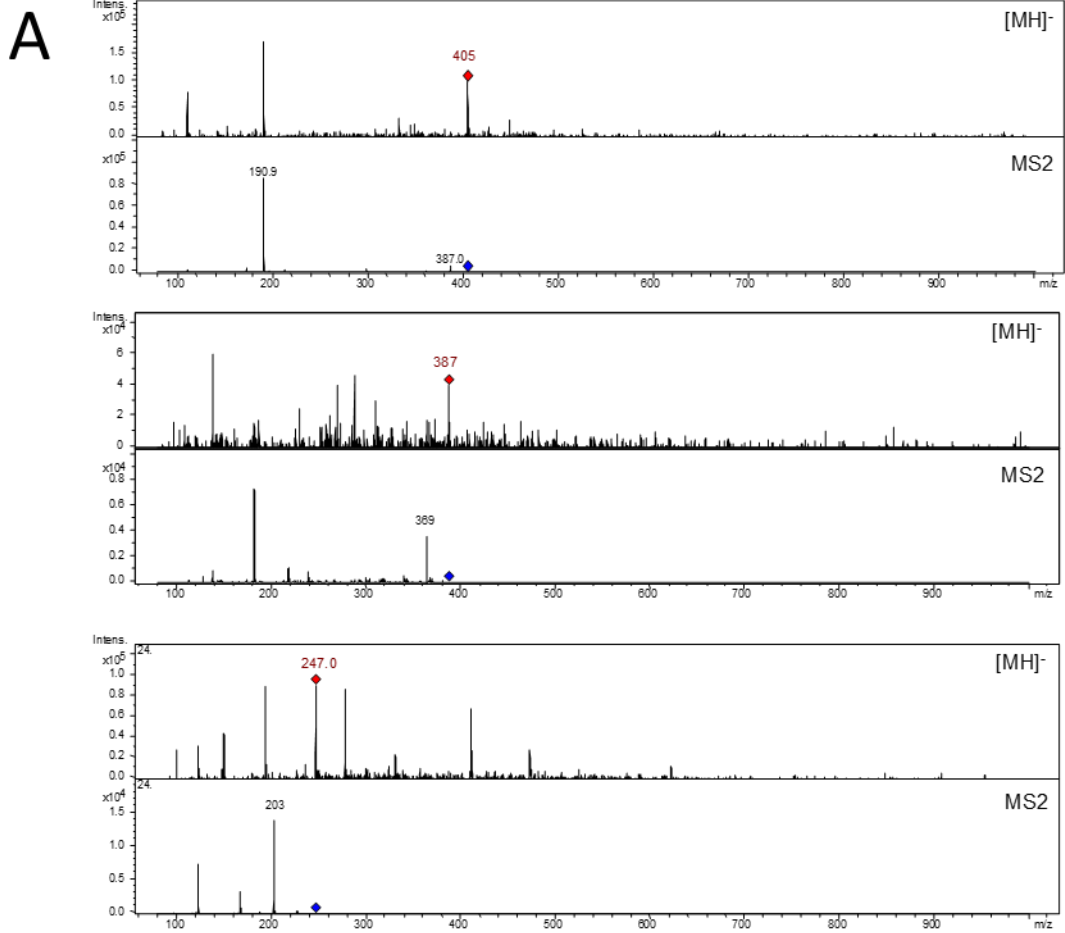
B

RP-HPLC analysis of seaweed metabolites in urine samples

RT (min)	metabolite	mean (mg)	stdev	N volunteers	time (h)	Enz -	Enz +
7.9	UM1	15.25	1.39	2	0 -- 8; 8 -- 24	yes	yes
10.4	UM2	10.77	4.29	7	8 -- 24	yes	yes
14.2	UM3	16.66	0.89	2	8 -- 24	yes	yes
16.7	UM4	2.94	1.82	3	8 -- 24	yes	yes
17.7	UM5	0.67		1	8 -- 24	yes	no
18.7	UM6	5.58	0.17	2	8 -- 24	yes	yes
24.1	UM7	9.85		1	8 -- 24	no	yes
29.6	UM8	0.30		1	8 -- 24	yes	no
30.8	UM9	0.51		1	8 -- 24	yes	no
33.0	UM10	1.26		1	8 -- 24	no	yes
33.1	UM11	1.28	0.09	2	8 -- 24	yes	yes
34.3	UM12	0.63	0.15	2	8 -- 24	yes	yes
36.5	UM13	0.61	0.55	6	8 -- 24	yes	yes
41.8	UM14	1.45	0.86	8	0 -- 8; 8 -- 24	no	yes
43.7	UM15	0.25		1	8 -- 24	no	yes
45.5	UM16	0.69		1	8 -- 24	yes	no
46.1	UM17	0.25		1	8 -- 24	no	yes
46.7	UM18	0.70		1	8 -- 24	no	yes

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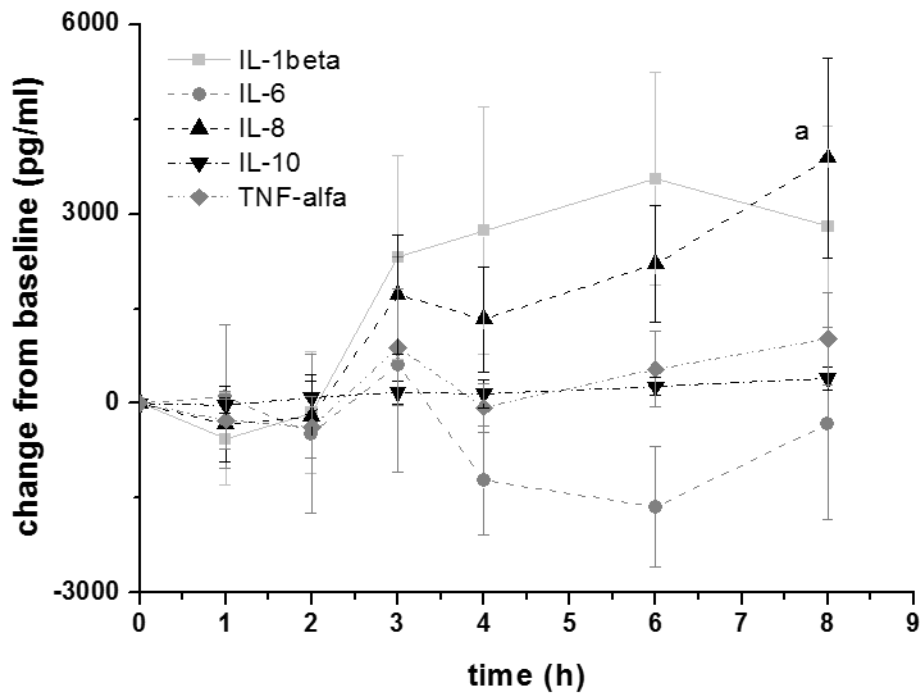
B **LC-MS analysis in negative ion mode of the urine samples**

RT (min)	[M-H] ⁻ (m/z)	MS2 (m/z)	tentative identification	
2.20	405	387	191	hydroxytrifuhald A
4.10	387	369		7-hydroxyeckol
10.40	247	203		C-O-C dimer of phloroglucinol , UMA3
1.70	289	221	175	Unknown, DM4
1.85	377	341	161	Unknown, DM7

524

525

526 **Figure 8**



527

528

529

530 **FIGURE LEGENDS**

531

532 **Figure 1.** Chromatographic separation of phlorotannins contained in the seaweed extract by
533 Normal-Phase HPLC with diode array detection (268 nm).

534

535 **Figure 2.** Characterisation of phlorotannins in the seaweed extract. A: Structures of
536 phlorotannins identified in the seaweed extract: B: Phlorotannins in the seaweed extracts
537 identified by LC-MS analysis in negative ion mode

538

539 **Figure 3.** LC-MS analysis in negative ion mode of the seaweed extract subjected to *in vitro*
540 gastrointestinal digestion, colonic fermentation and dialysis to mimic absorption. A: LC-MS
541 spectra and fragmentation of *in vitro* digested materials. B: Summary of LC-MS analysis of
542 the *in vitro* digested materials

543

544 **Figure 4.** Schematic illustration of the clinical intervention set up.

545

546 **Figure 5.** HPLC analysis of plasma samples for seaweed metabolites. A: HPLC
547 chromatograms (268 nm) and UV spectras showing examples of metabolites in plasma. B:
548 summary of seaweed metabolites present in plasma samples.

549

550 **Figure 6.** HPLC analysis of urine samples for seaweed metabolites. A: HPLC chromatograms
551 (268 nm) and UV spectras showing examples of metabolites in urine. B: summary of seaweed
552 metabolites present in urine samples.

553

554 **Figure 7.** LC-MS analysis in negative ion mode of urine samples. A: LC-MS spectra and
555 fragmentation of phlorotannins found in urine samples. B: Summary of LC-MS analysis of the
556 urine samples.

557

558 **Figure 8.** Cytokine production by whole blood cultures in cultured blood collected a various
559 time-points (0, 1, 2, 3, 4, 6 and 8 h) during the intervention study (LPS treated – unstimulated
560 controls). a = P < 0.05 vs. baseline.

561

562 **Financial support**

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564 *under the Capacities Programme (FP7)*

565

566 **Conflict of interest**

567 None

568

569 **Authorship**

570 G.C., J.P.E.S, P.Y. and I.R. designed the research. G.C. organized and coordinated all parts of
571 the clinical trial and analytical work. S.H. overviewed the preparation of seaweed materials,
572 extracts and capsules. P.A. aided running the clinical trial. Y.J. aided with the analysis of
573 metabolites in plasma and urines. G.C. analysed and summarised all the data. G.C. drafted the
574 manuscript, C.G., I.R, S.H. and P.Y. revised the manuscript. I.R. had primary responsibility for
575 final content.

576

577

578 **LIST OF ABBREVIATIONS**

579 SPE Seaweed Polyphenol Extract

580 DM Digestion Metabolite

581 UM Urine Metabolite

582 PM Plasma Metabolite

583 ESI electrospray ionization

584 RP-HPLC Reverse Phase High Performance Liquid Chromatography

585 NP-HPLC Normal Phase Liquid Chromatography

586

587 **References**

588

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