

**Supplementation with low amount of seaweed improves iodine status in iodine-insufficient British women**

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17

18 **Abstract**

19 Iodine-insufficiency is now a sustained issue in the UK and other European countries, due to  
20 low intakes of dairy and seafoods (especially where iodine fortification is not in place). Here,  
21 we tested commercially-available encapsulated edible seaweed (Napiers Hebridean  
22 Seagreens® *Ascophyllum nodosum* species - NaHS) for its acceptability to consumers, iodine  
23 bioavailability and the impact of a 2-week long daily supplementation on iodine levels and  
24 thyroid function. Healthy non-pregnant women of childbearing age, self-reporting low dairy  
25 and seafood consumptions, with no history of thyroid or gastro-intestinal disease were  
26 recruited. Seaweed iodine (712 µg, in 1g seaweed) was modestly bioavailable at 31-46% of  
27 the ingested iodine dose (n=22). After supplementation (2 weeks, 0.5g seaweed daily, n=42),  
28 urinary iodine excretion increased from 78 (IQR 47) to 140 µg/L (IQR 92), p<0.001. Thyroid  
29 stimulating hormone increased from 1.5 (IQR 1) to 2.1 mUI/L (IQR 1.6) (p<0.001) with two  
30 subjects exceeding the normal range after supplementation (but normal free thyroxine). There  
31 was no change in other thyroid hormones levels after supplementation. The seaweed was  
32 palatable and acceptable to consumers as a whole food or as an ingredient, and effective as a  
33 source of iodine in an insufficient population. Incorporation in staple foods would provide an  
34 alternative to fortification of salt or other foods with potassium iodine.

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37 **Introduction**

38 Iodine is essential for the synthesis of the thyroid hormones triiodothyronine (T<sub>3</sub>) and  
39 thyroxine (T<sub>4</sub>) which play a key roles in metabolism, and are vital for a growing fetus, for  
40 normal growth and brain development <sup>46</sup>. While hypothyroidism complicates some  
41 pregnancies <sup>1</sup>, it does not preclude hypothyroid women to become pregnant <sup>32</sup>, and iodine  
42 intake is crucial during the period surrounding child-bearing. When the iodine intake is  
43 below the recommended intake (140 µg/day) <sup>10</sup>, adequate secretion of the thyroid hormones  
44 may still be achieved by physiological adaptation. Modifications of thyroid and pituitary  
45 activities increases thyroid stimulating hormone (TSH) secretion, which enhances production  
46 of T<sub>3</sub> relative to T<sub>4</sub> and rapid iodine turnover <sup>9</sup>, but fetal supply and placental transfer remain  
47 low. For epidemiological purposes, iodine insufficiency is defined as a population, or  
48 subgroup, with a median urinary excretion (UIC) less than 100 µg/l for non-pregnant adults,  
49 and below 150 µg/L for groups of pregnant women <sup>47</sup>. While iodine fortification of common  
50 foods is widespread, it is not provided in all countries. There is no requirement for iodine  
51 fortification of foods in UK, and iodine fortification is unusual. There is growing concern that  
52 subclinical iodine deficiency may be emerging in post-industrial countries previously  
53 assumed to be iodine sufficient and there is currently very little evidence about the need for  
54 specific dietary advice, or for iodine fortification / supplementation targeted towards these  
55 two key vulnerable groups: young women and their infants.

56 With dairy and seafoods as main dietary source of iodine <sup>20</sup>, the UK has been considered  
57 iodine replete. Areas with historical endemic goitre ('Derbyshire neck') no longer see clinical  
58 dietary hypothyroidism, in what was hailed an accidental public health success, following  
59 change to farming practice and supplementation of dairy herds <sup>36</sup>. However, a recent survey  
60 of British schoolgirls has highlighted mild iodine deficiency with median urinary iodine

61 concentrations of 80 µg/L <sup>44</sup>. Similar results were found in a Scottish survey of women of  
62 childbearing age <sup>25</sup>. Although few people have frank iodine deficiency and hypothyroidism, a  
63 low or marginal intake presents a potential hazard in pregnancy due to the increased demand  
64 placed on maternal thyroid function <sup>16</sup>. This level of iodine insufficiency in the population is  
65 sufficient to impair intellectual development of future generations. Bath *et al.* showed that  
66 low maternal iodine status in pregnancy (individual iodine-to-creatinine ratios below 150 µg/g  
67 in spot samples) was associated with decreased cognitive functions in the ALSPAC cohort of  
68 1040 children from the south of England <sup>4</sup>. While there is no lack of availability of dietary  
69 iodine in these regions <sup>29</sup>, the explanation may be that many of the young female population  
70 commonly exclude fish and/or dairy products from their diets, for social or other reasons,  
71 leading to either low or marginal iodine intakes <sup>34</sup>.

72 Seaweeds used to feature as cheap and natural traditional foods in the British diet <sup>23</sup> until  
73 more recently when proper standards have come in to ensure suitability as human food  
74 seaweed. Despite this, it is still rather neglected in the UK, and data on its consumption are  
75 lacking, despite the fact that it is a rich source of iodine, with wide variation between species  
76 (from 16 to 8165 µg/g) <sup>43</sup>.

77 This study aimed to investigate the potential of seaweed as a safe and acceptable option for  
78 dietary iodine supplementation, specifically answering the following research questions:

- 79 1) What is the bioavailability of iodine from an encapsulated edible seaweed  
80 (Seagreens® *Ascophyllum nodosum* species), in a group of asymptomatic non-  
81 pregnant women reporting to consume low amounts of iodine-rich foods?
- 82 2) What is the impact of daily consumption of the encapsulated seaweed on iodine levels  
83 and thyroid function, in the same group of women?
- 84 3) Is the encapsulated seaweed acceptable for consumers (taste / use)?

85 **Material and Methods**

86 **Seaweed supplement**

87 Each capsule contained 0.5g Seagreens *Ascophyllum nodosum* (Napiers Hebridean Seagreens  
88 Capsules - NaHS), equivalent to 356 µg iodine (suppliers information based on  
89 measurements from independent UKAS accredited laboratories). NaHS is a dried and milled  
90 seaweed, sourced in Scotland and produced to distinct human food seaweed™ standards  
91 (patents pending) ensuring the safety, quality, sustainability and consistency of the products.  
92 All products are rigorously monitored during harvesting, drying and milling, and analyzed  
93 independently by UKAS accredited laboratories for nutritional composition, contaminants  
94 and heavy metals.

95 **In vitro iodine bioavailability assays**

96 The *in vitro* determination of the bioavailability of iodine in seaweed is based on the simple  
97 simulation of gastric and intestinal digestion according to the method developed by Romaris  
98 Hortas *et al.*<sup>37</sup>.

99 Digestion was carried out in triplicate. In brief, powdered NaHS (0.5 g) was added to distilled  
100 water (20mL) and the pH was adjusted to 2.0 with a 6M hydrochloric acid. Fresh gastric  
101 solution (0.15 g, pepsin 6.0% (w/v) dissolved in 6.0M HCl) was added to the flask, prior to  
102 incubation (37°C in a shaking bath at 150 rpm for 120 minutes). Digestate aliquots (0.5 mL)  
103 were transferred to -20°C prior to iodine determination. The digestate pH was neutralized  
104 with NaOH (pH 7.5). Dialysis bags filled with 0.15N PIPES (20 mL) were placed inside each  
105 flask, along with intestinal digestion solution (pancreatin 4.0% (m/v) and bile salts 2.5%  
106 (m/v) dissolved in 0.1M sodium hydrogen carbonate, 5mL). The flasks were incubated at  
107 37°C in a shaking water bath at 150 rpm for 120 min. The enzymatic reaction was stopped by

108 immersing the flasks in an ice water bath. The dialysis bags were removed and residual or  
109 non-dialyzable fraction (remaining slurries in the flasks) were transferred to polyethylene  
110 vials and separately weighed. Aliquots (1.5 mL) from the dialysate (20 mL) and non-  
111 dialysate fractions (25 mL) were transferred to - 20°C prior iodine determination.

112 Colonic fermentation was carried out as described by Edwards <sup>12</sup>. Briefly, faecal samples  
113 (16g) from three healthy volunteers were homogenized with a blender (30 s) in fermentation  
114 buffer (50 mL) to make a 32% faecal slurry. An aliquot (5 mL) of the non-dialyzable fraction  
115 of the intestinal digestate was added to faecal slurries (50 mL). The bottle was purged with  
116 OFN (1 min) and sealed and incubated in a shaking water bath at 37°C and 60 stroke/min.  
117 Samples were taken at t= 0h, 2h, 4h, 6h and 24h to measure pH and were immediately stored  
118 at -20°C prior to iodine determination.

### 119 **Human iodine bioavailability experimental design**

120 The study was approved by the University of Glasgow Medical Veterinary and Life Sciences  
121 College Ethics committee. All participants provided written informed consent.

122 Healthy women aged 18-46, self-reporting as low-iodine consumers, were recruited locally  
123 using via posters and word-of-mouth, to take part in cross-over iodine bioavailability study.  
124 Those with existing thyroid or gastro-intestinal conditions, taking medication other than the  
125 contraceptive pill or smoking were excluded, as well as pregnant or lactating women and  
126 those planning to conceive. Those taking dietary supplements containing iodine were also  
127 excluded.

128 Height, weight, waist circumference and blood pressure were measured after recruitment.  
129 Usual dietary intake was determined using an iodine-specific food frequency questionnaire <sup>8</sup>.  
130 Participants were allocated at random to treatment order (potassium iodine (KI) or seaweed

131 first) and were asked to avoid all iodine-rich foods (dairy and seafood) for the duration of the  
132 study. Prospective food dietary were filled for the duration of the study, and the iodine  
133 content of participants diet was determined using Windiets 2005 (Robert Gordon University).  
134 A 7-day wash out period between each leg of the cross-over intervention. Participants were  
135 asked to replicate their diet during the second leg of the study.

136 All urine passed on Day 1 (baseline 24h urines) was collected. On Day 2, participants  
137 received either a seaweed supplement (NaHS, 1 g) or potassium iodide (KI) supplement  
138 (equivalent iodine content; 712 µg) to be taken fasted with a breakfast of white toast and a  
139 glass of water. Urine was collected for 24 hours, in fractions for the periods 0-2h, 2-5h, 5-8h,  
140 8-20h and 20-24h.

#### 141 **Seaweed supplementation study – experimental design**

142 Healthy women aged 18-50, self-reporting as low-iodine consumers, were recruited locally  
143 using via posters and word-of-mouth, to take part in cross-over seaweed supplementation  
144 study. Those with existing thyroid or gastro-intestinal conditions, or taking medication other  
145 than the contraceptive pill were excluded, as well as those taking iodised dietary  
146 supplements. None had taken part in the bioavailability study. The supplementation study  
147 was approved by the University of Glasgow Medical Veterinary and Life Sciences College  
148 Ethics committee. All participants provided written informed consent. The a priori sample  
149 size was calculated in G Power (Kiel University, Germany) using UIC as a primary outcome  
150 for mean difference between two groups using the Wilcoxon signed-Rank test for matched  
151 pairs, assuming a logistic parent distribution. A sample size of n=42 was calculated, to detect  
152 (or not) an increase from the current population UIC for the target group (median 75µg/L,  
153 calculated mean 94 µg/L, standard deviation 80 µg/L<sup>25</sup>) to a sufficient UIC (100 µg/mL),  
154 equivalent to a ~14% increase in UIC, and an effect size of 0.47, with  $\alpha=0.05$ ,  $\beta=0.80$ ).

155 Participants' height, weight, waist circumference and blood pressure were measured at the  
156 beginning and end of the supplementation period. Usual dietary intake was determined using  
157 an iodine-specific food frequency questionnaire <sup>8</sup>. During the run-in period, participants were  
158 asked to keep a 4-day weighed food diary. Urine was collected for 24 hours on Day 4. On day  
159 5, participants were supplied with a stock of supplements, and instructed to consume one  
160 capsule of NaHS daily (0.5 g per day, equivalent to an intake of 356 µg/d) for 14 days, while  
161 following their usual diet. A fasted venous blood sample was collected, and the total volume  
162 of the urine collection measured. At the end of the supplementation period, participants  
163 replicated the diet recorded on the 4-day weighed diary (Days 16-19), and collected 24-hour  
164 urine on the last day of supplementation (Day 19). A final fasted venous blood sample was  
165 collected (Day 20). All urine and plasma samples were aliquoted and stored at -80°C until  
166 analysis. Compliance was checked by counting the number of capsules remaining in the  
167 container supplied to volunteers.

#### 168 **Urinary iodine measurements**

169 Urinary iodine and iodine concentration in digestates were analysed using the colorimetric  
170 Sandell-Kolthoff reaction adapted for the 96-well microtiter plate, as described by Ohashi *et*  
171 *al.* <sup>33</sup>, using a custom-made sealing cassette. Sample were measured in triplicates (CV%  
172 <10%).

#### 173 **Thyroid function tests**

174 Thyroid stimulating hormone (TSH), thyroglobulin (Tg), triiodothyronine (T<sub>3</sub> and fT<sub>3</sub>) and  
175 thyroxine (T<sub>4</sub> and fT<sub>4</sub>) were measured in plasma in duplicates using immunoassays (ELISA  
176 assays, Astra biotech GmbH, Luckenwalde, Germany).



177 **Acceptability of the supplement**

178 Participants filled a self-administered questionnaire focusing on habitual frequency of  
179 consumption of seaweed products (6-point Likert scale, “daily” to “never”), opinions on taste  
180 (3 statements, 5-point Likert scales, “strongly agree” to “strongly disagree”), after-taste (1  
181 statement, 5-point Likert scales, “strongly agree” to “strongly disagree”) and overall  
182 acceptability of seaweed as a food or ingredient (3 statements, 5-point Likert scales, “strongly  
183 agree” to “strongly disagree”). Open questions were used to gather information on taste, after  
184 taste, and views on seaweed as an ingredient in foods.

185 **Statistical analyses**

186 Data were expressed as mean  $\pm$  SD or as median and inter-quartile range (IQR) depending on  
187 normality, which was checked using the Shapiro-Wilks test. Categorical data (Likert scale)  
188 was described using the mode and IQR. Significance was implied at  $p < 0.05$ . Wilcoxon  
189 signed-Rank test for matched pairs or paired t-test was used to assess the difference between  
190 paired groups depending on their data distribution, while the Mann-Witney U-test or  
191 independent t-test was used to compare unrelated samples. Analysis was carried out using  
192 SPSS 18.0 (SPSS Inc., Chicago, IL, USA).

193

194 **Results**

195 **In vivo bioavailability study**

196 Healthy females (n=22), median age 24.5 (IQR 14.3) were recruited and completed the  
197 bioavailability study. Socio-demographic and anthropometric details for the group are  
198 summarized in Table 1.

199 Dietary iodine intake was low (below 55 µg/day) throughout the bioavailability study period,  
200 for each study arm (Table 2). The baseline median UIC, for the 24 hours preceding the study,  
201 was 40 µg/L (IQR 42) for the seaweed arm and 31 µg/L (IQR 52) for KI arm. Correcting for  
202 total urine volumes, this was equivalent to 50 µg/24h (IQR 43) preceding seaweed intake,  
203 and 50 µg/24h (IQR 54) preceding KI intake.

204 Urinary iodine output, in µg.L<sup>-1</sup>.h<sup>-1</sup> is presented in Figure 1, with cumulated iodine excretion  
205 in µg presented in Figure 2. The peak iodine excretion time occurred earlier for KI (0-2h)  
206 compared to the seaweed (2-5h). The amount of iodine excreted over the 24h period  
207 following ingestion was greater (p<0.001) following KI intake (421 µg, IQR 199) compared  
208 to seaweed intake (239 µg, IQR 153).

209 Participants were grouped according to habitual iodine intake, as either sufficient (n=7) or  
210 insufficient (n=13). The dose of iodine excreted in urine was calculated based on the iodine  
211 load of the NaHS capsule / KI plus the dietary iodine intake of day 3 (Table 2). The dose of  
212 iodine excreted was significantly higher following KI intake than seaweed intake (p<0.001).  
213 This was true for both subgroups (p=0.009 and p=0.017 for insufficient and sufficient group,  
214 respectively). However, while the dose of iodine excreted after KI was higher in the sufficient  
215 group (73% vs. 46%, p=0.036), there was no difference between groups after seaweed  
216 ingestion (46% vs 31%) (Table 3).

### 217 **In vitro bioavailability assays**

218 After digestion in the simulated gastric compartment, only 9.9±0.1% of the iodine present in  
219 the sample was available and in solution. After digestion in the simulated intestinal  
220 compartment, 4.9±0.1% of the initial iodine dose present was recovered in the dialysis bag,  
221 with a further 5.0±0.0% in the non-dialysable fraction. This indicates that approximately 90%  
222 of the iodine was still trapped in the seaweed matrix at that point and consistent with the

223 cumulated dose excretion in urine during the in vivo bioavailability study, which was  
224 approximately 12% of the dose ingested (IQR 8%). After faecal fermentation of an aliquot of  
225 the non-dialysable fraction, 51.2±10.4% of the iodine present was available, and in solution.

## 226 **Impact of seaweed supplementation on urinary iodine**

227 A total of 42 healthy females of childbearing age took part in the 2-week supplementation  
228 study. The demographic, anthropometric and dietary profiles of participants are presented in  
229 Table 4.

230 At baseline, median UIC was well below the cut-off for sufficiency (100 µg/L) at 78 µg/L  
231 (IQR 47). The group average iodine intake was 110 µg (IQR67), with 31 participants with an  
232 intake below the recommended intake of 140 µg/day. Subsequently, individuals were  
233 classified as having iodine-sufficient (>140 µg) or insufficient intake (<140 µg) based on  
234 their habitual iodine consumption as estimated by the FFQ. There was no difference in  
235 weight, BMI, waist circumference between the subgroups with sufficient or insufficient  
236 iodine intake at baseline.

237 After supplementation, median UIC increased significantly to 140 µg/L (IQR 92) (p<0.001).  
238 This increase in UIC differed between sufficient and insufficient group (+23 µg/L, IQR49 for  
239 the sufficient group, +97 µg/L, IQR75 for the insufficient group; p=0.041) and was only  
240 statistically significant in participants with insufficient habitual iodine intake (p<0.001). The  
241 total amount of iodine excreted over 24 hours was however significantly increased for both  
242 insufficient (from 93, IQR48 to 262, IQR 103 µg/day, p<0.001) and sufficient groups (from  
243 138, IQR 84 to 214, IQR 268 µg/day, p<0.041). Neither weights nor waist circumferences  
244 changed during the supplementation study.

245 **Impact of seaweed supplementation on thyroid function**

246 The thyroid function tests are presented in Table 5. At baseline, Tg and fT3 levels were  
247 different between iodine sufficient and insufficient subgroups ( $p=0.047$  and  $p=0.048$ ,  
248 respectively). Tg values were within the Tg reference range in healthy adults (3 - 40  $\mu\text{g/L}$ )  
249 but higher than the proposed cut-off for iodine sufficiency (10  $\mu\text{g/L}$ ).

250 TSH levels were within the normal range (0.4 – 4.5 mUI/L)<sup>3</sup> for all but one participant, who  
251 had a borderline TSH level of 5.72 (but normal fT4 levels).

252 There was no significant change in the thyroid hormones T3, T4, fT3, fT4 following  
253 supplementation, or Tg (with values remaining over 10  $\mu\text{g/L}$ )<sup>45</sup>. There was however a  
254 significant increase in TSH, from a median 1.5 mUI/L (IQR 1) to 2.1 mUI/L (IQR 1.6)  
255 ( $p<0.001$ ). This increase was significant in both insufficient and sufficient groups ( $p=0.027$   
256 and  $p=0.006$ , respectively), but more marked in those with sufficient habitual iodine intake  
257 ( $p=0.044$ ). Serum TSH did exceed the normal range for two participants (7.3 and 8.0 mUI/L)  
258 with fT4 still within the normal range. While fT3 levels did not significantly change for the  
259 whole group, those in the insufficient group had a decrease after supplementation ( $p=0.048$ ).

260 **Seaweed consumption and acceptability of the supplement**

261 Participants in the bioavailability and supplementation studies answered a side questionnaire  
262 on seaweed consumption (combined  $n=63$ ). They had very rarely been exposed to seaweed as  
263 a foodstuff, with 19% never having consumed it knowingly; 60% of participants had  
264 consumed it as sushi, on a monthly basis (18%) or less often (37%). Less than half (40%) of  
265 participants had consumed whole seaweed (less than twice a year). Most had never consumed  
266 lava bread (90%), nor seaweed as a tablet (92%) or a capsule (87%). The main reasons for the  
267 low consumption was lack of opportunity (mentioned by 64% of participants), and lack of  
268 appeal (54%).

269 Participants agreed that the taste of the supplement was acceptable when swallowed as a  
270 capsule (mode 5, median 4, IQR2) and disagreed that there was an unpleasant after-taste  
271 (mode 2, median 2, IQR2) or that the capsule were difficult to swallow (mode 1, median 2,  
272 IQR1). Supplementation study participants who had added the seaweed to foods (n=24)  
273 neither agreed nor disagreed on the acceptability of its taste as an ingredient (mode 3, median  
274 3, IQR0) or its ease of use for cooking (mode 3, median 3, IQR1).

275 Participants agreed that encapsulated seaweed is a good way to include seaweed in the diet  
276 (mode 4, median 4, IQR1). Preferred ways to consume seaweed included encapsulated  
277 (71%), as an ingredient in food (33%) or as a whole food (19%). Most (67%) saw the  
278 potential use of seaweed as a food ingredient as a positive. The main reasons were assumed  
279 health benefits and extra nutrients (35%) and flavour enhancement (24%). A minority (7%)  
280 held negative view on seaweed as an ingredient, with taste the main concern (75%). The rest  
281 were either unsure or with no opinion.

282

## 283 **Discussion**

284 This study showed that asymptomatic young women in the UK with diets low in seafoods  
285 and dairy products do indeed display biochemical evidence of quite marked iodine  
286 deficiency. It then shows how an acceptable/palatable commercially available seaweed  
287 product can boost the iodine intake of a group of mostly iodine-insufficient women, without  
288 deleterious impact on thyroid function.

289 Daily intake of an encapsulated seaweed (NaHS) was effective at raising the UIC of a group  
290 of females after a two-week supplementation period with a slight increase in the TSH levels  
291 after seaweed supplementation. Our results are in agreement with Teas *et al.* who

292 supplemented iodine-replete healthy post-menopausal women with *Alaria esculenta* capsules  
293 for 7 weeks (475 µg iodine/day)<sup>41</sup> and Clark *et al.* (kelp, 1 g iodine/day for 6 weeks)<sup>6</sup>. The  
294 TSH levels remained within the normal range for all but two participants, with no change  
295 observed for the thyroid hormones, whereas Clark *et al.* observed a decrease in total T3 after  
296 supplementation. Tg values remained higher than the proposed 10 µg/L cut-off for iodine  
297 insufficiency<sup>45</sup>, even after the supplementation, which might be indicative of a lag period for  
298 Tg values to fall within iodine sufficiency range after achieving iodine sufficient status.

299 The iodine contained in NaHS was bioavailable, although to a lesser extent (30%) than  
300 previously reported by Aquaron (90-100% for iodine-sufficient women, and 62-85% for  
301 iodine-insufficient women over 48-hours)<sup>2</sup> or Teas (60% for iodine-sufficient women over  
302 48-hours)<sup>41</sup>. This may be directly related to our shorter (24-hour) urine collection, and the  
303 type of seaweed used in the other studies (*Gracillaria verrucosa*, *Laminaria hyperborea* and  
304 *Alaria esculenta*). Incomplete collections are also a possible explanation. We showed a  
305 difference in excretion between those with either sufficient or insufficient iodine intake as  
306 previously described<sup>2</sup>. *In vitro* digestion confirmed limited release of the iodine from the  
307 seaweed matrix in the first gastric and intestinal phases of simulated digestion. We showed  
308 that colonic fermentation of seaweed is important to free iodine from the seaweed matrix,  
309 with mechanism relying on fermentation of the polysaccharide matrix<sup>30</sup> or metabolism of  
310 organic iodine<sup>37</sup>. Therefore, the seaweed matrix may delay iodine absorption (compared to  
311 KI), with iodine released from the food over a longer period. Impact of further processing  
312 such as cooking needs to be taken in consideration if seaweed is used as an ingredient, as it  
313 would lead to partial loss via evaporation<sup>27; 43</sup>.

314 Several studies reported that iodine insufficient populations were diagnosed with iodine-  
315 induced hyper- or hypothyroidism following high iodine intake<sup>39; 5; 14; 26</sup>, however, a two-

316 week iodine supplementation with up to 500  $\mu\text{g}/\text{d}$  had no impact on thyroid function tests in  
317 euthyroid subjects<sup>35</sup>. Upper tolerable limit of iodine intake in healthy individuals have been  
318 defined as 1.1 mg/d in the United States and 600  $\mu\text{g}/\text{d}$  in the European Union<sup>15; 17</sup>. While  
319 epidemiological evidence has linked high daily seaweed/iodine intake with higher thyroid  
320 cancer risk in Japan<sup>31</sup>, this observation is not supported by experimental studies in rats with  
321 chronic high iodine intake (up to 1g/L in drinking water)<sup>40</sup>. The thyroid gland can adapt to  
322 excessive iodine intake after initial diminution in the excretion of thyroid hormone due to the  
323 Wolff-Chaikoff effect. This effect was demonstrated to have a longer lasting suppression of  
324 the thyroid gland in those ingesting excess seaweed<sup>28</sup>. Restricting the seaweed intake was  
325 able to reverse iodine-induced goiter and transient hypothyroidism<sup>48</sup>.

326 Reports of widespread iodine insufficiency in Britain and other European countries, the  
327 renewed interest in iodine nutrition and the lack of iodine prophylaxis in the UK represent an  
328 opportunity for seaweed as a foodstuff. Iodine insufficiency results from low intake of dairy  
329 (especially milk, which consumption has been steadily decreasing since 1975<sup>13</sup>), and seafood  
330 (which consumption is low in the UK population at 37g/day<sup>11</sup>). Iodised salt is the main  
331 method of iodine prophylaxis worldwide but its implementation would be in direct conflict  
332 with the efforts to reduce salt consumption in relation to the prevention of chronic diseases.  
333 With table salt usage falling following successful public health campaigns, it may be  
334 contradictory to portray salt as a vehicle for iodine. A more viable option to increase iodine  
335 status includes fortification of staple foods with seaweed, which was previously successfully  
336 incorporated in a nutritionally-balanced pizza, designed in the context of health-by-stealth  
337 improvement of ready meals. Seaweed addition enabled to reduce the sodium content of the  
338 product, while improving nutritional content, without compromising the taste or appearance<sup>7</sup>.  
339 Given that iodine is extensively stored in the thyroid, it can safely be consumed

340 intermittently, which makes seaweed use in a range of foods attractive, and occasional  
341 seaweed intake enough to ensure iodine sufficiency.

342 Seaweed consumption in most Western cultures has been low, due to low availability and  
343 poor consumer awareness regarding potential health benefits <sup>22</sup>. The benefits of incorporating  
344 seaweed isolates into the habitual diet goes further than addressing iodine deficiency, with  
345 impact of seaweed consumption on serum oestradiol, reduction of the glycaemic response to a  
346 carbohydrate load, and increased satiety via lowered gastric emptying. These aspects may be  
347 relevant to the development of functional foods for weight management <sup>24; 18; 42; 19; 21</sup>.  
348 Incorporation in bread had no impact on taste or appearance <sup>22</sup>. With an average trade price of  
349 £8 per kg, the additional cost per loaf would be minimal considering that seaweed is iodine-  
350 rich and that little would be required.

351 The contaminants and heavy metal content of seaweed is sometimes a concern, especially in  
352 retailed products with poor traceability and limited compositional analysis, as consumption  
353 may expose the consumer to heavy metals such as organic / inorganic arsenic <sup>38</sup>. Water  
354 quality is important for seaweed quality, and France is the only European country with  
355 specific regulations for the use of seaweeds as vegetables <sup>27</sup>. The seaweed used in this study  
356 (NaHS) was grown in Scottish Grade A Pristine water (SEPA/SNH evaluation) and produced  
357 to Human Food Seaweed™ standards (patents pending). Compositional analysis, carried out  
358 on every batch, showed no contaminants and heavy metals below threshold levels. This is  
359 important if seaweed will become a more commonly used ingredient in processed foods.

360 In conclusion the answers to the research questions behind this study are:

361 1) Iodine bioavailability from the encapsulated seaweed was low in the group of women  
362 studied. The seaweed matrix may be a key factor for this low bioavailability.



363 2) Daily consumption of 0.5g of NaHS increased urinary iodine level to 140 µg/L for the  
364 group. TSH increased slightly, within the normal range for all but two participants,  
365 with no change to thyroid hormones levels.

366 3) Participants indicated that the encapsulated seaweed had an acceptable taste, was easy  
367 to use, and were positive about seaweed use as an ingredient.

368

369 The study conclusions would have been strengthened with a randomised controlled crossover  
370 study design, longer exposure time and reassessment of iodine status and thyroid function  
371 after the end of the intervention, but that would demand an impractical duration of high  
372 tolerance from volunteers. It would be of value to repeat the biochemical aspects in different  
373 subject groups. The influence of the seaweed matrix on bioavailability will be an important  
374 factor to consider if seaweed is incorporated in cooked and uncooked staple foods. A large-  
375 scale survey needs to take place to properly investigate attitudes to seaweed utilisation in  
376 processed foods and cuisine in general.

377

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489

490 **Figure legends**

491

492 Figure 1: Urinary iodine excretion in  $\mu\text{g/L/h}$  over 24h, after ingestion of a dose of  $712\mu\text{g}$  iodine,  
493 from KI (■) or NaHS (○).

494 Figure 2: Cumulated iodine output in  $\mu\text{g}$  over 24h, after ingestion of a dose of  $712\mu\text{g}$  iodine, from  
495 KI (■) or NaHS (○).

496

497 **Table 1: Characteristics of the bioavailability study participants (n=22)**

		<b>Median</b>	<b>IQR</b>
<b>Demographic &amp; anthropometric details</b>	Age (yrs)	24.5	14.3
	Height (cm)	165.3	4.7
	Weight (kg)	59.6	14.8
	Waist (cm)	71.0	12.5
	BMI (kg/m <sup>2</sup> )	22.0	4.9
<b>Usual diet</b>	Milk (mg/day)	131.1	144.3
	Other dairy (mg/day)	114.9	90.2
	Seafood (mg/day)	23.6	15.7
	Daily iodine intake (µg/day)	126.8	54.8
		<b>Count (n)</b>	<b>(%)</b>
<b>Ethnicity</b>	White British	6	27%
	White Europeans	4	18%
	Other ethnicities	12	55%
<b>Body composition</b>	Overweight (BMI>25)	3	14%
	Obese (BMI>30)	1	5%
<b>Iodine intake</b>	Daily iodine intake >140 µg/day	7	33%
	Daily iodine intake <140 µg/day	14	67%

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501 **Table 2: Daily dietary iodine intake (µg) according to study arm**

<b>Study arm</b>	<b>Day 1</b>		<b>Day 2</b>		<b>Day 3</b>	
	median	IQR	median	IQR	median	IQR
<b>NaHS - KI</b>	54	52	45	36	39	36
<b>KI - NaHS</b>	53	25	48	65	38	40

502

503 **Table 3: Percentage iodine dose excreted, according to habitual iodine intake (sufficient &**  
 504 **insufficient)**

	<b>Seaweed</b>		<b>KI</b>	
	median	IQR	median	IQR
<b>insufficient (n=13)</b>	31% <sup>a</sup>	13%	46% <sup>b</sup>	28%
<b>sufficient (n=7)</b>	46% <sup>a</sup>	16%	73% <sup>b</sup>	13%
<b>All (n=22)</b>	33% <sup>a</sup>	18%	57% <sup>b</sup>	28%

505 <sup>a,b</sup> significantly different change (Δ pre-post supplementation) between groups at p<0.05

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512 **Table 4: Characteristics of the participants in the 2-week supplementation study (n=42)**

		<b>Median</b>	<b>IQR</b>
<b>Anthropometric and demographic information</b>	Age (yrs)	27.0	15.0
	Height (cm)	164.3	6.8
	Weight (kg)	61.6	14.1
	Waist (cm)	72.1	14.9
	BMI (kg/m <sup>2</sup> )	22.6	4.8
<b>Usual diet</b>	Milk (mg/day)	180.3	169.8
	Other dairy (mg/day)	70.6	124.0
	Seafood (mg/day)	19.6	31.4
	Daily iodine intake (µg/day)	109.7	67.4
		<b>Count (n)</b>	<b>(%)</b>
<b>Ethnicity</b>	White British	25	60%
	White Europeans	9	21%
	Other ethnicities	8	19%
<b>Body composition</b>	Overweight (BMI>25)	10	24%
	Obese (BMI>30)	4	10%
<b>Iodine intake</b>	Daily iodine intake >140 µg/day	11	26%
	Daily iodine intake <140 µg/day	31	74%

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514

**Table 5: Iodine status and thyroid function pre and post supplementation in participants meeting the daily iodine recommendation (n=11) or not (n=31). Data are presented as median (IQR).**

	All (n=42)			Insufficient (n=31)			Sufficient (n=11)		
	Pre	Post	Δ	Pre	Post	Δ	Pre	Post	Δ
UIC (μg/L)	78.0 (74.8)	140.0 (91.8) ***	72.5 (105.2)	50.1 (61.2)	148.9 (89.2) ***	97.4 (75) <sup>a</sup>	103.7 (36.4)	139.0 (94.3)	23.5 (49.1) <sup>b</sup>
UIC (μg/24h)	94.1 (81.5)	248.2 (128.2) ***	147.4 (108.5)	93.0 (48.3)	262.3 (103.3) ***	149.1 (93.2)	137.8 (83.9)	214.3 (268.8) *	76.5 (142.4)
TSH (mUI/L)	1.5 (1)	2.1 (1.6) ***	0.5 (1.1)	1.4 (1.1)	1.9 (1.6) *	0.4 (0.9) <sup>a</sup>	1.7 (0.8)	2.7 (0.9) **	0.8 (0.7) <sup>b</sup>
Tg (μg/L)	21.8 (15.3)	20.6 (13.1)	-1.0 (6.1)	26.6 (17.7)	24.0 (14.1)	-1.7 (6.8)	17.2 (10.9)	15.8 (5.5)	-0.4 (3.5)
T3 (nmol/L)	1.9 (0.5)	1.9 (0.5)	-0.1 (0.3)	1.9 (0.6)	2.0 (0.4)	-0.1 (0.4)	1.9 (0.2)	1.9 (0.6)	-0.1 (0.2)
T4 (nmol/L)	86.9 (21.8)	86.0 (26.2)	2.3 (14.9)	89.9 (23.2)	86.9 (35.8)	-0.3 (12.7)	80.8 (12.1)	83.8 (18.9) *	2.9 (13.6)
fT3 (pmol/L)	5.5 (4.5)	4.4 (3.8)	-0.2 (1.6)	4.1 (3.9)	3.3 * (3.7)	-0.3 (1.4)	6.8 (2.5)	6.8 (2.5)	0.0 (1.5)
fT4 (pmol/L)	13.8 (3.2)	14.4 (3.5)	0.4 (1.7)	13.9 (3.6)	14.5 (3.2)	0.4 (1.4)	13.5 (2.5)	14.3 (3.7)	0.2 (2.8)

Δ difference between parameters measured pre and post supplementation

\* p<0.05, \*\* <p<0.01, \*\*\* p<0.001 pre vs post supplementation

<sup>a,b</sup> significantly different change (Δ pre-post supplementation) between groups at p<0.05

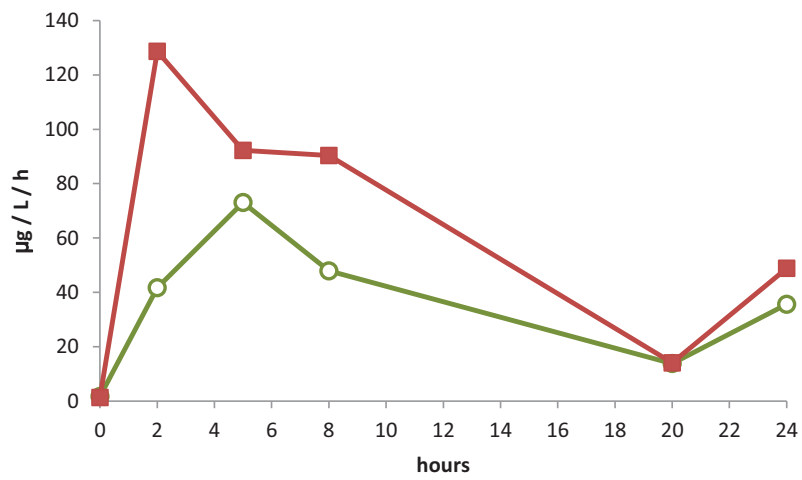


Figure 1: Urinary iodine excretion in  $\mu\text{g/L/h}$  over 24h, after ingestion of a dose of 712  $\mu\text{g}$  iodine, from KI (■) or NaHS (○).



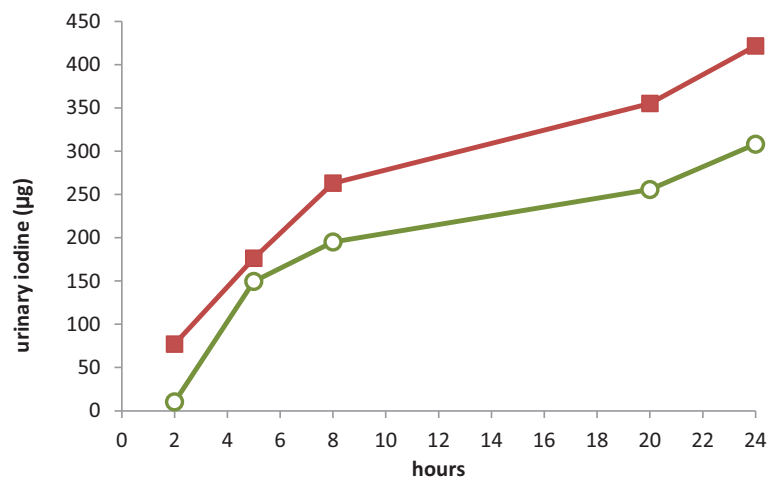


Figure 2: Cumulated iodine output in µg over 24h, after ingestion of a dose of 712ug iodine, from KI (■) or NaHS (○).