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Iodine uptake in brown seaweed exposed to radioactive liquid discharges from the reprocessing plant of ORANO La Hague

3

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13

Abstract

14 lodine-129 is present in controlled liquid radioactive waste routinely released in seawater by the 15 ORANO nuclear fuel reprocessing plant in La Hague (Normandy, France). Brown algae are known for their exceptional ability to concentrate iodine from seawater. They also potentially emit volatile iodine 16 17 compounds in response to various stresses, such as during emersion at low tide. For these reasons, 18 brown seaweed is routinely collected for radioactivity monitoring in the marine environment. Despite 19 the high concentration ratio the exact mechanism of iodine uptake is still unclear. Chemical imaging 20 by laser desorption/ionization mass spectrometry provided evidence that iodine is stored by kelps as 21 I. In this study we investigate in vivo iodine uptake in kelps (Laminaria digitata) with an emphasis on 22 seawater iodine chemical speciation. Our results showed that kelp plantlets were able to take up iodine 23 in the forms of both IO_3^- and I^- . We also observed transient net efflux of I^- back to seawater but no IO_3^- 24 efflux. Since the seaweed stores I⁻ but takes up both IO₃⁻ and I^{<math>-}, IO₃⁻ was likely to be converted into I^{<math>-} at</sup></sup></sup> 25 some point in the plantlet, or through the action of associated microorganisms. One major outcome 26 of our experiments was the direct observation of the kelp-based biogenic conversion of seawater IO₃⁻ 27 into Γ . On the basis of both IO_3^- and I^- uptakes by the seaweed, we propose new steps in the possible 28 iodine concentration mechanism used by kelp.

- 29 Keywords: radioactive iodine, uptake, kelp, chemical speciation
- 30

31 1 Introduction

32 Though a minor element in seawater, iodine is mainly present on Earth dissolved in the oceans 33 (Carpenter, 2003; Carpenter et al, 2021; Chance et al, 2014; Wong, 1991). In the terrestrial 34 environment, it is present at very low levels in soils, water and plants, but in vertebrates, it is involved 35 in the crucial functions of the thyroid gland as organically bound in thyroid hormones (Fuge & Johnson, 36 2015). lodine is also present as multiple volatile compounds involved in a complex geochemical cycle 37 between the oceans and the atmosphere. The high reactivity of iodine is considered to play a key role 38 in atmospheric chemistry and possibly the climate, at least in coastal environments (Carpenter et al, 2021; Carpenter et al, 2013; Kolb, 2002; Saiz-Lopez et al, 2012). In addition to stable ¹²⁷I, iodine has 36 39 radioactive isotopes, and with the exception of ¹²⁹I (half-life of 15.7 10⁶ years), most iodine 40 radioisotopes quickly decay (half-lives of a few days). Two main iodine radioisotopes are generated by 41

human activities. ¹³¹I is a short-lived (half-life of 8.5 days) fission by-product of ²³⁵U in nuclear electricity 42 production, and high activities are also used in medicine for cancer radiotherapy. ¹²⁹I is generated in 43 44 nuclear reactors, and because of its very slow decay rate, it is a major residual iodine radioisotope in the nuclear fuel cycle, which adds to naturally occurring ¹²⁹I (cosmic ray spallation of Xe in the 45 46 atmosphere and spontaneous fission of natural uranium). It is also present in fallout from past 47 atmospheric nuclear weapons testing and major powerplant accidents. ¹²⁹I is present in controlled 48 liquid radioactive waste routinely released in seawater by the ORANO nuclear fuel reprocessing plant 49 in La Hague (Normandy, France) and the annual amounts released are between 1 and 2 TBq.yr⁻¹ (Fiévet et al, 2020). Discharging liquid ¹²⁹I waste into seawater results in isotopic dilution in naturally-present 50 stable ¹²⁷I. The background isotopic ratio ¹²⁹I/¹²⁷I is in the range [10⁻¹³-10⁻¹²] and reaches [10⁻⁷-10⁻⁶] in 51 seawater in the English Channel (Hou et al, 2007). Considering a concentration of dissolved ¹²⁷I in 52 53 seawater of 0.5 µM, this means ¹²⁹I activities in the range [0.05-0.5] Bq.m⁻³. However, higher seawater 54 ¹²⁹I activities could be expected in the close vicinity of the ORANO plant outlet.

55 Brown algae are known for their exceptional ability to concentrate iodine from seawater. The ratio 56 of iodine content between kelps (i.e. Laminariales) and seawater reaches 10⁵, especially in young 57 plantlets (Leblanc et al, 2006). Moreover, brown algae are also known to release volatile iodine 58 compounds in response to various stresses, such as during emersion at low tide (Ball et al, 2010; 59 McFiggans et al, 2010; McFiggans et al, 2004; O'Dowd et al, 2002; Whitehead et al, 2009). Wherever brown seaweed is exposed to ¹²⁹I discharges in seawater, the radioisotope could return to the 60 61 terrestrial environment via an airborne route, namely via the wind and aerosols. For these reasons, 62 brown seaweed is routinely collected in the marine environment in the English Channel to monitor 63 radionuclide concentrations, and in particular that of ¹²⁹I (Fiévet et al, 2020). In a previous study on 64 radionuclide transfers between seawater and biota, we observed that the bioavailability of ¹²⁹I 65 appeared to decrease with distance from the source of input (the ORANO La Hague plant outlet). We 66 emphasized the role of the chemical speciation of dissolved iodine in seawater (Fiévet et al, 2021). 67 lodine uptake by Laminariales, and more generally brown algae, is hypothesized to be mediated by enzymes from the vanadium-haloperoxidase (VHPO) family, through the specific oxidation of iodide 68 69 (Küpper et al, 1998). Dissolved iodine is naturally present in seawater in the forms of iodide (I⁻) and 70 iodate (IO_3). Although iodate is the dominant chemical form because of pH and redox potential in 71 seawater, both chemical forms are present in various proportions (Carpenter et al, 2013; Hou et al, 72 2007; Wong, 1991). The conversion mechanisms between iodine species are still unclear because 73 iodate does not spontaneously reduce to iodide in natural seawater for thermodynamic reasons (Hou 74 et al, 2007). Nevertheless this reaction between iodate and iodide takes time to reach equilibrium 75 (Luther et al, 1995), around two weeks according to estimates (Carpenter, pers. Comm.). Now, 76 depending on the chemical form of ¹²⁹I discharged by the reprocessing plant, the question arises as to 77 bioavailability its for seaweed. The PUREX process 78 (https://www.cea.fr/Documents/monographies/Procédé-PUREX.pdf) operated at the ORANO reprocessing plant results in the liquid release of ¹²⁹I as iodide. As such, it is assumed to be a potential 79 substrate for VHPOs, being incorporated by kelps. But as conversion into ¹²⁹IO₃⁻ occurs in seawater, it 80 may become unavailable for the enzyme. In summary, two crucial questions should be answered to 81 better understand ¹²⁹I uptake by brown seaweed in the marine environment around the reprocessing 82 83 plant of ORANO La Hague: 1- Which form of iodine is bioavailable for uptake by brown seaweed? 2-84 Does it evolve with time and distance from the outlet once released in seawater by the plant? More 85 generally, the exact mechanism of iodine uptake is still unclear, and was challenged by chemical 86 imaging results confirming a major apoplastic storage location (Lebeau et al, 2021). Assessing which 87 chemical form of seawater iodine is taken up by kelps in the marine environment should contribute to the exploration of its bioaccumulation mechanism. 88

89 In this study, we investigate in vivo iodine uptake in L. digitata with an emphasis on chemical speciation between iodide vs iodate. We also use radioactive ¹³¹I⁻ as a tracer of unidirectional iodide 90 influx to estimate iodide turnover in seaweed as well as the relationship with seawater iodide 91 concentration. We also report an isotopic ratio ¹²⁹I/¹²⁷I measured in the marine environment in the 92 93 vicinity of the outlet of the ORANO reprocessing plant in La Hague. Iodine-129 activity and stable iodine 94 concentrations were determined in total dissolved iodine in seawater and as total iodine extracted 95 from two species of brown seaweed, L. digitata and Fucus serratus collected in Goury (6 Km from the 96 plant outlet). Finally, laboratory experiment results and data from the natural environment are put 97 into perspective to attempt to better understand radioactive iodine transfers between seawater and 98 brown seaweed.

99 2 Material and methods

100 2.1 Algal material

For iodine uptake experiments, plantlets of *Laminaria digitata* [0.05-3 g] were collected on the rocks in Goury (Normandy, France) at low tide (spring-tides) and transferred to the laboratory (LRC, IRSN, Cherbourg, France) for maintenance in an aquarium. The temperature was set to that of the natural environment upon collection (12.5 °C), the water was aerated and recirculated. About 200 g of total biomass (plantlets and young sporophytes, see below) was kept in a 60 L tank in natural light, and the seawater was replaced every week with pre-cooled fresh seawater. The iodine concentration in seawater (see 2.2.2) was checked to ensure no depletion occurred.

For ¹³¹I experiments, we used both whole plantlets and tissue disks. In the latter case, disks of about 0.04-0.05 g (diam. 14 mm) were punched out of the blade of young sporophytes (20-40 cm long; collected and maintained as described above) of *L. digitata* by die-cutting. Algal disks were then kept in filtered seawater (seawater collected in Goury, filtered at 0.45 μm; Millipore ref. HAWG0700) until iodine leakage stopped, as indicated by the monitoring of iodine concentration in seawater (with one disk in 10 mL of seawater: no iodine leakage was observed after 1 hour).

For the environmental monitoring of ¹²⁹I/¹²⁷I, adult sporophytes of *L. digitata* (1-1.5 m in length) and *Fucus serratus* (30-70 cm in length) were collected monthly at low tide at Goury for 37 months (see supplementary material S2-1 for sampling location).

117 2.2 Determination of I^- and IO_3^- in seawater

118 2.2.1 Separation of seawater I^{-} and IO_{3}^{-}

119 After sample filtration (0.45 μ m, Millipore ref. HAWG0700), seawater I⁻ and IO₃⁻ were separated by 120 anion exchange chromatography (AEC) using Bio-Rad AG1-X4 resin and mini-spin columns according 121 to (Hou et al, 2001; Zhang, 2015). Briefly, AG1-X4 resin (provided in Cl⁻ form) was conditioned into NO₃⁻ 122 form as follows: 1- a first wash in NaClO 5% removed any residual I⁻ from the fresh resin. 2- Cl⁻ was 123 replaced by NO₃⁻ in NaNO₃ 2M. 3- Excess NO₃⁻ was finally flushed with MilliQ water (Millipore). About 124 0.5 cc of conditioned resin (NO₃⁻ form) was added to mini Bio-spin columns (Bio-Rad #7326207) and 125 allowed to drain by gravity. Then 3 more 0.9 mL washes of MilliQ water were applied and left to drain prior to starting AEC on filtered (0.45 μm) seawater as follows: 1- two 0.9 mL seawater samples were 126 127 passed through the resin to replace the mobile phase. 2- The bottom of the column was closed with 128 the stopper tip, another 0.9 mL sample was added and the top was closed with the cap. The column 129 was flipped twice and agitated horizontally for 1 hour on a 3D-platform shaker (Polymax) at slow 130 speed. 3- The bottom stopper tip and the top cap were removed and the mobile phase was allowed to 131 drain in a 1.5 mL centrifuge tube. The flow-through contained IO_3^- anions whilst I⁻ anions stayed 132 trapped on the resin. Total dissolved iodine concentration was measured (see 2.2.2) before the AEC (I⁻ + IO_3^{-1} and in the column flow-through (IO_3^{-1}), I⁻ was calculated as the difference. We checked the accuracy of the separation with the standard addition method using KI, KIO₃ and both together.

135 2.2.2 Determination of iodine concentration

136 Total dissolved iodine (as I⁻ and/or IO₃⁻) concentration in seawater was determined by 137 spectrophotometry according to (Sandell & Kolthoff, 1937) in the range [0.04-0.30] µM. Seawater samples were filtered (0.45 µm, Millipore ref. HAWG0700) and diluted in MilliQ water to fit within that 138 139 range. The color reaction was prepared in a 1.5 mL centrifuge tube with 500 μ L of either seawater or 140 an AEC-treated seawater sample (diluted), 50 µL NaCl (10% g/g), 500 µL As₂O₃ (0.1 M) and transferred 141 to a water bath at 30°C. Then 125 µL of (S0₄)₄Ce(NH₄)₄, 2 H₂O, (0.02 M) was added and the reaction was 142 allowed to run for exactly 20 min at 30°C in the water bath. Absorbance at 420 nm was then measured 143 in 1 mL disposable cuvettes in a spectrophotometer. A calibration curve was carried out with standard solutions between 0.039 µM and 0.316 µM (dilutions of 1.31 g.L⁻¹ KI in MilliQ water solution). The 144 145 repeatability of the method showed that the uncertainty of the results was ± 5%. Total iodine 146 concentration before $(I^2 + IO_3)$ and after (IO_3) separation by AEC (see 2.2.1) was determined in order 147 to obtain the I⁻ concentration based on the difference. We checked the accuracy of the quantification 148 with the standard addition method using KI, KIO₃ and both together. The reliability of method was 149 further checked by comparing the results obtained by spectrophotometry with duplicates analyzed by 150 ICPMS measurements on some of our samples.

151

152 2.3 Measurement of net iodine flux between seaweed and seawater

153 Plantlets of L. digitata (pooled to about 2 g) were incubated in beakers with 50-75 mL of filtered 154 seawater (0.45 μ m, Millipore ref. HAWG0700) at the same temperature as in the host aquarium (12.5 155 \pm 1.0 °C), with bubbling. Seawater samples were collected over time (a picture of the incubation setup 156 is provided in the supplementary material S1-1). The sample volume was 0.5 mL for total dissolved 157 iodine ($I^{-} + IO_{3}^{-}$) and 3 mL for AEC separation. Beakers without seaweed plantlets served as controls for 158 changes in iodine concentrations into the filtered seawater used due to adsorption onto the beakers 159 surface. No changes in iodine concentrations were observed in those beakers. At some point in the 160 experiment, seawater was spiked either with KI, KIO₃ or both. Upon completion of each experiment, the plantlets were patted dry between absorbent paper sheets and weighed. Concentrations of both 161 162 I^{-} and IO_{3}^{-} in the water were plotted as a function of time.

163 2.4 Measurement of unidirectional ¹³¹I⁻ influx in seaweed

164 Iodine-131 as KI was purchased from CIS Bio International (I-131-S-2) and the activity information was provided by the manufacturer with the calibration date (Act.=1.11E+08 Bq; Vol.=0.67 cc; 165 ¹³¹/¹²⁷I=4.64E+08). Because of constraints in accurate nuclear metrology (geometry calibration), 166 unidirectional radioactive ¹³¹I influx was measured on tissue disks punched out of *L. digitata* blade as 167 previously published (Küpper et al, 1998; Shaw, 1959). Tissue disks were incubated in seawater spiked 168 169 with ¹³¹I in the form of KI. The disks were rapidly patted dry on absorbent paper, transferred to a plastic 170 container and counted for 60 sec on a hyper pure GeHP gamma detector. The measurement procedure 171 lasted 1.5 min and the disks were then returned to the incubation medium to carry on the uptake 172 kinetics experiment. Radioactive iodine disk content was then plotted as a function of time and it was found to be linear for 15 min. The activity in seawater was found to be constant for short-term 173 experiments (15 min). We investigated the relationship between the ¹³¹I uptake by the disks and iodine 174 175 concentration in seawater. For that purpose, seawater iodine concentration was increased by addition of KI to 1, 3, 10, 50, 500 µM and spiked with ¹³¹I (230 Bq.mL⁻¹). The slope of the 15 min linear ¹³¹I uptake 176 177 allowed the iodine permeability of the disks to be estimated (in nmol.min⁻¹.g⁻¹) since the exchange surface was constant. Plotting the iodine permeability as a function of seawater concentration allowed
 for the derivation of the values of Vmax and Km according to the Michaelis-Menten model as
 previously described by (Küpper et al, 1998). The Michaelis-Menten model was fitted using Microsoft
 Excel[™] Solver function.

182 A longer time intake experiment was carried out over 14 days with a 0.1 g plantlet kept in a 10 L tank to estimate iodine turnover by the algae. The procedure was the same as for tissue disks with the 183 plantlet patted dry and lying in a plastic dish for gamma counting. The activity of the seawater (30 184 Bq.mL⁻¹ seawater) was maintained constant by adding ¹³¹I to compensate for intake by the plantlet as 185 well as ¹³¹I decay. The very large volume of the tank compared to the small size of the plantlet ensured 186 187 that no total iodine depletion occurred in seawater. The biological half-life of iodine in L. digitata plantlet was estimated as $tb_{1/2} = Ln(2)/k$ by fitting the data to equation Act_t = Act₀.[1-exp^(-k.t)] (with 188 t=time; Act=¹³¹I activity at time t; k=time constant) with the Solver function of Microsoft Excel[™] (we 189 don't need to implement ¹³¹I decay since the radioisotope activity is artificially maintained constant in 190 191 seawater).

192 2.5 Measurement of iodine-129 activity in seawater

193 120 L-seawater samples were collected monthly at Goury for 37 months, and filtered at 0.45 μm 194 (Millipore, Durapore membrane ref. HVLP2932A). Total dissolved iodine concentration was 195 determined in an aliquot as described above (2.2.2). A known amount of stable iodine (0.500 g KIO_3) 196 was added to the 120 L as a carrier and to allow for the calculation of the extraction yield. Dissolved 197 iodine in the forms of I^{-} and IO_{3}^{-} were merged by the addition of 120 g Na₂SO₃ at pH 2 (HCl-36%) to reduce IO₃⁻ into I⁻. Total I⁻ was then precipitated by the addition of 6 g AgNO₃. The AgI precipitate was 198 199 recovered in 500 mL with MilliQ water and then solubilized by the addition of 35 g L+ascorbic acid 200 $(C_6H_8O_6)$ at pH 10 (NaOH 5N). The solution containing dissolved iodine was then transferred into a 201 separation funnel and covered with 100 mL 1,2,4-Trimethylbenzene (TMB). HNO₃ 65% was added until 202 pH 2 was reached, and NaNO₂ added whilst pH was maintained with HNO₃ to oxidize iodide into I_2 . The 203 funnel was vigorously shaken to trap l₂ in the organic phase and then left to decant. The aqueous phase 204 was recovered and another 50 mL TMB was added to further extract I₂. The two TMB organic phases 205 were pooled (150 mL) and washed twice with 100 mL MilliQ water (the water was discarded). Extracted 206 iodine was finally recovered from the organic phase by adding 20 mL of NaOH 0.5 N; Na₂SO₃ 0.2 M to 207 reduce I₂ into I⁻. The separation funnel was vigorously shaken, allowed to settle, and the 20 mL aqueous 208 phase was recovered in a plastic vial where traces of TMB adhered to the plastic wall. The aqueous 209 phase was finally transferred into a clean 20 cc geometry. The final 20 cc geometry contained initial + 210 added stable iodine and unknown iodine-129. Iodine-129 activity was determined by gamma 211 spectrometry in the 20 cc geometry previously calibrated with a standard solution (IL29-ELSF15 212 #4793/4, provided by CEA/DAMRI), using a hyper pure GeHP gamma detector (29,78 keV double ray, 213 emission rate 57,3%, detection limit = 0.05 Bq in the final sample with a counting time of 80 000 s, 0.03 214 Bq in the final sample with a counting time of 240 000 s). A quenching curve was obtained by 215 incrementing stable iodine concentration in the standard IL29-ELSF15 solution using the same 20 cc geometry: the quenching by stable iodine was lower than 15% (15% at 600 mg iodine, 5% at 200 mg 216 217 iodine). Stable iodine concentration in the 20 cc geometry was determined by potentiometric method 218 using an Orion 96-53 ionplus® Series Iodide Electrode according to the manufacturer's instructions. 219 The iodine concentration in the geometry was used to correct the results for the extraction yield and 220 for gamma counting quenching. Extraction yield was in the range [70%-80%]. Final results were expressed in Bq.m⁻³ for ¹²⁹I and in μ M for stable ¹²⁷I. The isotopic ratio ¹²⁹I/¹²⁷I could then be calculated 221 in seawater for total dissolved iodine ($I^2 + IO_3^2$; with 1 Bq ¹²⁹I = 1.186E-09 mol). 222

223 2.6 Iodine-129 activity measurement in seaweed

In parallel with seawater, iodine-129 activity was determined in *L. digitata* and *F. serratus* samples collected monthly (see **Erreur ! Source du renvoi introuvable.**), dried at 90°C and finely ground as described in (Bouisset et al, 1999; Lefevre et al, 2003). In both species, stable total iodine concentration was determined in parallel on aliquots by the CNRS/SCA (Service Central d'Analyses, Vernaison, France). Seaweed dry material was mineralized in Schöninger combustion flasks and the solutions were analyzed using ion exchange chromatography with electrochemical detection (the quantification limit

- 230 was 0.25 μ g.L⁻¹ iodine). Final results were expressed in Bq.Kg⁻¹ dry weight for ¹²⁹I and in mg.Kg⁻¹ dry for
- stable 127 I. The isotopic ratio 129 I/ 127 I could then be calculated in dry seaweed material).
- 232

233 **3 Results**

234 3.1 Which form of inorganic iodine in seawater is taken up by *Laminaria digitata*?

A first series of experiments consisted in the incubation of *L. digitata* plantlets in natural seawater collected in Goury at the same location as the seaweed with $[IO_3^-] = 0.35 \ \mu\text{M}$ and $[I^-] = 0.25 \ \mu\text{M}$. The changes in seawater IO_3^- and I^- concentrations showed that only IO_3^- appeared to decrease in the medium in the presence of the plantlet, and this was confirmed by spiking seawater with IO_3^- at time 90 min (Figure 1 left). However, spiking seawater with I^- instead of IO_3^- clearly demonstrated that both iodine forms decreased (Figure 1 right). These experiments were replicated 6 times for I^- and $IO_3^$ independently with the same results (supplementary materials S1-2 and S1-3).

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247 When replicating the experiment, the transfer of *L. digitata* plantlet to the incubation vial 248 sometimes resulted in transient iodine leaks from the seaweed. In those cases, only I⁻ increased in 249 seawater whilst IO_3^- continued to decline (Figure 2).





Figure 2: Transient iodine leak by the plantlet which only resulted in net I^- output whilst IO_{3^-} was still taken up (same symbols as in Figure 1, uncertainties ± 5 %).

Since both IO_3^- and I^- seem to be taken up by *L. digitata*, the question then arose as to why the $I^$ concentration didn't decrease in the experiment illustrated in Fig.1 left? A simple hypothesis is that the concentrations in seawater determine which form of iodine is pumped. We repeated the experiment with different concentrations of IO_3^- and I^- in seawater, starting with the same concentrations of both forms of iodine, and increased the levels up to 1.4 μ M for each iodine species (Figure 3).



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Figure 3: Incubating *Laminaria digitata* plantlets in seawater with increasing levels of IO_3^- and I^- at the same concentration (same symbols as in Figure 1, uncertainties ± 5 %).

When both IO_3^- and I^- were present at the same concentration, their concentrations decreased equally in the presence of the plantlet. So, in the left experiment in Figure 1, was there a minimum $I^$ concentration below which the uptake just balanced the output? Likewise, does a minimum $IO_3^$ concentration exist below which this iodine form uptake just balances the output? To address this 266 point, we extended the net flux experiment for 6 hours to attempt to see the "asymptotic" seawater

 IO_3^- and I⁻ concentrations when no more net flux were observed. At the end, the net iodine uptake was re-stimulated by a fresh spike of I⁻ and IO_3^- . The results of this long-term kinetics are shown in Figure 4.



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Figure 4: Long-term kinetics experiment on net iodine flux using plantlets of *Laminaria digitata* (same symbolsas in Figure 1).

This long-term kinetics experiment was replicated 7 times (supplementary materials S1-4) and every time, IO_3^- concentrations declined below I concentrations⁻, as in the example displayed in Figure 4. The average iodine concentrations (± SE) at time 0 were $[IO_3^-] = 0.35 \pm 0.01 \,\mu\text{M}$; $[I^-] = 0.31 \pm 0.06 \,\mu\text{M}$ and at time 360 min, they were $[IO_3^-] = 0.16 \pm 0.03 \,\mu\text{M}$; $[I^-] = 0.29 \pm 0.03 \,\mu\text{M}$, (N=7).

277 3.2 Tracing iodine uptake in *Laminaria digitata* with ¹³¹I

278 Since both IO_3^- and I^- seemed to be taken up equally by the seaweed (provided their concentrations were high enough), using radioactive ¹³¹I⁻ was considered to trace dissolved iodine uptake whatever 279 280 the chemical form. We investigated the relationship between the uptake and iodine concentration for L. digitata using disks punched out from the thallus blade as previously described (Küpper et al, 1998; 281 282 Shaw, 1959). The unidirectional flux J_{in} (nmol.min⁻¹.g⁻¹) was proportional to seaweed iodine permeability, since the disk surface was constant during the experiment. The results are presented in 283 Figure 5 fitted with Michaelis-Menten model kinetics with apparent Km = 59 μ M and Vmax = 216 284 nmol.min⁻¹.g⁻¹. From those results, we calculate an approximate unidirectional influx capacity of ¹³¹I of 285 c.a. 100 nmol.min⁻¹.g⁻¹. This unidirectional influx can be compared to results obtained in our net fluxes 286 experiments (Figures 1 to 4), in which we obtained influxes of around 1 nmol.min⁻¹.g⁻¹. Net fluxes result 287 from the balance between input and output fluxes. Therefore, the 2 orders of magnitude higher 288 unidirectional input fluxes measured in our ¹³¹I uptake experiment indicate that output fluxes must be 289 290 very high to result in a 100 fold reduction when considering net fluxes.



291

Figure 5: Relationship between unidirectional iodine influx and seawater iodine concentration for tissue disks of *Laminaria digitata*. Fitting the Michaelis-Menten model kinetics yielded Km = 59 μ M and Vmax = 216 nmol.min⁻¹.g⁻¹.

A two-week experiment was performed to estimate the biological half-life of iodine in *L. digitata* plantlets. The ¹³¹I uptake experiment displayed in Figure 6 yielded an estimated $tb_{1/2}$ value of c.a. 5 days.

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- 299
- Figure 6: ¹³¹I uptake over two weeks by *Laminaria digitata* plantlet leading to the estimation of the iodine
 turnover in the seaweed as 5 days.
- 3.3 ¹²⁹I transfer between seawater and brown seaweed exposed to discharges from the
 303 ORANO reprocessing plant in La Hague

304 3.3.1 Seasonal changes of iodine in seawater and in brown seaweed

Seawater samples were periodically collected in Goury and analyzed for iodine chemical speciation.
 Additional samples were collected in different locations around the Cap of La Hague and along the
 coast of Brittany (Bretagne, France). The data are presented in the supplementary material (S2-2).

Since brown seaweed iodine content may vary during the seasonal cycle, it was crucial to take these changes into account when looking at radioactive iodine transfer to seaweed. Total iodine in mg.Kg⁻¹ dry was determined in *L. digitata* and *F. serratus* samples were collected monthly in Goury. The data are displayed in the supplementary material (S2-3).

312 3.3.2 Iodine-129 activity in seawater and in brown seaweed

- 313 Iodine-129 activity determined in dissolved iodine in seawater $(IO_3^- + I^-)$ and in the two brown
- seaweed species *L. digitata* and *F. serratus* are presented in Figure 7.
- 315

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Figure 7: lodine-129 activity in seawater and in brown seaweed collected in Goury, near the outlet of the ORANO reprocessing plant in La Hague. Error bars account for metrological uncertainties (2 x sigma; may be masked by symbol).

Because iodine content varies in seaweed over the season, expressing the radioactivity data as the isotopic ratio ${}^{129}I/{}^{127}I$ was necessary to explore ${}^{129}I$ transfer between seawater and seaweed. The ratio ${}^{129}I/{}^{127}I$ was determined in dissolved iodine in seawater ($I^- + IO_3^-$) and in *L. digitata* and *F. serratus* dry material and the data are presented in Figure 8.



Figure 8: lodine-129 isotopic ratio in seawater and in brown seaweed collected in Goury, near the outlet of the ORANO reprocessing plant in La Hague.

327 4 Discussion

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The fate of ¹²⁹I liquid discharged by the ORANO reprocessing plant in La Hague (Normandy, France) 328 329 deserves special attention for several reasons: due to its very slow decay, this radionuclide is a major 330 residual iodine radioisotope in the nuclear fuel cycle and a tracer of the discharges by the ORANO 331 reprocessing plant in the environment. As part of the liquid waste released in seawater in the Alderney 332 Race (Bailly du Bois et al, 2020; Fiévet et al, 2020), it mixes with natural stable iodine so that isotopic 333 dilution minimizes its impact in terms of radioprotection. However, brown seaweed is known to be 334 exceptionally efficient at bioaccumulating iodine from seawater. The recommended value for the 335 iodine concentration factor (CF) in macroalgae is 10^4 (IAEA, 2004), and ratios of up to 4.3 10^4 were observed in kelps (Fiévet et al, 2021). Moreover, brown seaweed has the ability to transform 336 337 accumulated iodine into volatile compounds, such as I₂, released in response to various stresses, such as emersion at low tide (Ball et al, 2010; McFiggans et al, 2010; McFiggans et al, 2004; O'Dowd et al, 338 2002; Whitehead et al, 2009). This process might challenge the strategy to keep ¹²⁹I away from humans 339 340 because the isotope can return to the terrestrial environment via airborne routes via the wind and 341 aerosols. For these reasons, although the potential contribution of this atmospheric route is not 342 known, brown seaweed is focused on as part of radionuclide monitoring of the marine environment. 343 Understanding the transfer of radioactive iodine released by the nuclear industry (under control) in 344 the marine environment is not only essential for the sake of radioprotection, the fate of radioactive 345 iodine released into the English Channel also has a much larger scope than radioprotection as a potential tracer of the complex behavior of iodine in the marine environment. The different key steps 346 347 of iodine transfer from seawater to the bioaccumulation in brown seaweed are part of the bigger 348 puzzle of the biogeochemical cycle of iodine in the ocean and its consequences on the chemistry of 349 the atmosphere (Carpenter et al, 2021). The transfer of iodine from the ocean to the atmosphere is assumed to occur through the oxidation of iodide (I⁻) into diiodine (I₂) in the presence of atmospheric ozone under solar radiation, but also through biological oxidation processes (Carpenter et al, 2021). lodate (IO₃⁻) is the preferential chemical form of iodine in seawater, therefore its conversion into iodide (I⁻) is not spontaneous as regards thermodynamics (Hou et al, 2007; Wong, 1991). After decades of literature on the topic, the reduction of iodate into iodide still remains poorly understood, though microorganisms or marine algae clearly play a role (Carpenter et al, 2021; Wong, 1991) and this factor has recently been confirmed for some marine bacteria (Reyes-Umana et al, 2022).

357 In the present study, we provide new information on the uptake mechanism of iodine by kelp (L. 358 digitata). First, our results suggest that kelp plantlets were able to take up iodine in the forms of both 359 IO_3^- and I^- with the same efficiency (Figure 1). When both iodine species were present at the same 360 concentration, they declined at equivalent rates in seawater (Figure 3). It should be emphasized that 361 our experimental conditions magnified naturally occurring phenomena due to a change in the ratio 362 between the seaweed mass and the volume of its surrounding seawater compared to the natural 363 environment. This magnification allowed us to easily assess the net IO_3^- and I^- fluxes between the 364 plantlet and seawater by measuring the decrease in those iodine species in seawater, with our 365 measurements accounting for the balance between inputs and outputs. Plantlets of L. digitata are very 366 sensitive and required extreme care during transfers from the natural environment and between the 367 host aquarium and the flux experiment setup. Chemical imaging data provided information on a very 368 unequal compartmentalization of iodine in the tissue of L. digitata plantlet. Iodine appears to be 369 preferentially concentrated in the peripheral apoplastic tissue (Lebeau et al, 2021). As such, transient 370 iodine leakage might occur when manipulating a plantlet. We assumed that plantlets underwent 371 minimal stress because after each experiment, they were returned to the host aquarium and could be 372 re-used for more experiments, and the iodine leaks sometimes observed at the beginning of the 373 experiments were only transient and brief. Indeed, a transient iodine leak by the plantlets was 374 observed in 11 out of 42 net flux experiments. In all cases it should be emphasized that only I⁻ leaks 375 were ever detected in seawater whilst IO_3^- always continued to decline (Figure 2). This observation was 376 an *in vivo* evidence that iodine is stored in L. digitata tissue as iodide (I^{-}) , as shown by laser 377 desorption/ionization mass spectrometry (Lebeau et al, 2021). We could then interpret the results of 378 our net flux experiment on the basis of this information. When assessing IO_3^- and I^- net fluxes, we 379 expected those fluxes to stop when the inputs balanced the outputs, or when the seawater 380 concentrations were too low to meet the needs of the uptake process. In the long-term (6 hours) 381 experiment (Figure 4), a change in seawater iodine speciation was observed in the presence of L. 382 digitata plantlets. The IO₃⁻ concentration was initially above the I⁻ concentration, and this reversed 383 after 2 hours. IO_3^- declined down to around 0.15 μ M whilst I⁻ stayed at around 0.25 μ M. If the seaweed 384 only stores I but can take up both IO₃ and I, we conclude that IO₃ was likely to be converted into I at some point in the plantlet (apoplast), or through the action of associated microorganisms. After 5-6 385 386 hours of incubation, the stable IO_3^- and I^- concentrations in seawater therefore resulted from $IO_3^- + I^-$ 387 influxes and I⁻ outflux only. We assume that the IO₃⁻ stabilized in seawater because its concentration 388 was so low that the uptake mechanism was no longer able to pump this iodine form, "by exhaustion", 389 as indicated by the more than 200 fold difference between the K_M of 59 μ M and the final iodine 390 concentrations of 0.15-0.25 μ M we measured. Conversely, the extreme chemical gradient of I⁻ 391 between the seaweed and seawater and its permeability resulted in I⁻ outflux. I⁻ stabilization in 392 seawater corresponded to the balance between this I⁻ leak and the ability of the mechanism to take 393 up I⁻. So, when the IO₃⁻ is low enough in seawater to exhaust its uptake, we only observe I⁻ recycling. A 394 major conclusion of this laboratory experiment was that we directly observed the kelp-based biogenic 395 conversion of seawater IO₃⁻ into I⁻. Even if the quantitative contribution of the process was considerably

magnified with our biomass/seawater ratio, this provided direct evidence of this iodine conversion
 mediated by brown algae and associated microorganisms observed in the marine environment.

398 Second, we also concluded that *L. digitata* plantlets can take up either IO_3^{-} or I^{-} indifferently. lodine 399 bioaccumulation in kelps is assumed to be mediated by vanadium-dependent haloperoxidases (VHPO) 400 family (Küpper et al, 1998), and especially by vanadium-dependent iodoperoxidases (VIPO) 401 characterized in L. digitata, specific to iodide oxidation (Colin et al, 2005). The substrates of vHPOs are 402 halides and H_2O_2 (Colpas et al, 1996) so VIPO is assumed to oxidize I⁻ and not IO_3^- . The hypothetical 403 mechanism involving only VIPO-based iodide uptake in iodine bioaccumulation by kelp was recently 404 questioned by our data on the apoplastic localization of iodine storage in the plantlets (Lebeau et al, 405 2021; Verhaeghe et al, 2008). The parallel uptake of IO_3^- and I⁻ that we observed in our net flux 406 experiments suggests a need to further validate in vivo the biological function of VHPO and/or VIPO in 407 iodine bioaccumulation. However, we suggest the following speculative scheme: if iodide is the only 408 form of iodine in the storage site, as shown previously (Lebeau et al, 2021; Verhaeghe et al, 2008), 409 oxidized forms may have a favorable chemical gradient to enter the seaweed. This would rely on the 410 strong reduction capacity of the storage site in order to maintain this gradient. The question of this 411 reduction capacity inside the seaweed storage compartment is still pending, but in this scheme, the 412 driving force would be this reduction capacity of the apoplastic storage site and the uptake would only 413 rely on oxidized iodine accessing the site along a favorable chemical gradient. Natural IO₃⁻ in seawater 414 would be available for this, and VIPO would be there to oxidize I⁻ when present. It should be kept in 415 mind that a major role for the I⁻ bioaccumulated in the seaweed is to react with reactive oxygen species 416 to reduce oxidative stress and/or to release oxidized iodine in response to stress as a defense 417 mechanism (Küpper et al, 2008).

We used an ¹³¹I tracer to further characterize iodine uptake in kelp. This replicated a kinetic 418 419 experiment (3 hours) performed many decades ago (Shaw, 1959). The isotope was added in the form 420 of ¹³¹I⁻, but our net flux experiment showed that both IO_3^- and I^- uptakes exactly matched. This 421 suggested that ¹³¹I⁻ unidirectional influx data were relevant to trace iodine uptake. The dependence of 422 I⁻ uptake on concentration followed Michaelis-Menten kinetics, as previously published with disks of 423 L. digitata (Küpper et al, 1998). The Michaelis-Menten kinetics confirmed the enzymatic mediation of 424 I⁻ uptake but our values of Km and Vm parameters were quite different. Our Km value of 59 μM was 425 more than 7 times lower and our Vm value of 219 μ M was more than three time higher. We have no 426 explanation for this discrepancy except maybe the physiological condition of the disks punched out 427 from the blade of L. digitata. On the basis of our Km value of 59 µM, we may speculate that at 428 concentrations more than two orders of magnitude lower, like in our flux experiments (Figure 4), the 429 uptake mechanism reached a steady state. A biological half-life of 5 days for iodine was estimated based on the long-term ¹³¹I⁻ uptake experiment on *L. digitata* plantlets. Although the values are 430 consistent, this was below our previous estimate of 14 days in brown seaweed based on modeling 431 432 using ¹²⁹I environmental data (L. digitata and F. serratus) in the marine environment in the vicinity of 433 the ORANO reprocessing plant in La Hague (Fiévet et al, 2021). Our laboratory experiment lasted 15 days, but we can't rule out that we had not reached the plateau for ¹³¹I activity in the seaweed. Besides, 434 435 it was not surprising that the young plantlet of *L. digitata* showed small differences in iodine uptake 436 kinetics compared to adult and other brown seaweed used in our environmental modelling.

The apparent convergence of IO_3^- and I^- uptake by *L. digitata* plantlets observed in our net flux experiments was essential for interpreting our study of radioactive iodine transfer between seawater and seaweed in the marine environment under the influence of radioactive liquid discharges from the ORANO reprocessing plant in La Hague. ¹²⁹I should be measured in seawater based on total iodine (IO_3^- + I^-) to account for ¹²⁹I bioavailability at least for kelps. Seasonal changes in iodine concentrations in 442 seawater and in seaweed were also essential to take into account. In seawater, if we merge all total 443 $(IO_3^- + I^-)$ dissolved iodine measurements we took in seawater along the coast of the English Channel 444 and Brittany (Map in the supplementary material S2-1), the average concentration (\pm SE) was 0.54 \pm 445 0.05 μM (N=56). Our recent measurements around the Cape of La Hague (supplementary material S2-446 2) with IO_3^- and I^- partitioning showed relatively stable IO_3^- concentration whilst I^- seemed to fluctuate 447 over the year. Longer time-series measurements are required to confirm these chemical speciation 448 data, but a question already arose from these I⁻ data. In seawater samples collected from the shore, 449 where brown seaweed including kelp was present, our IO3⁻ concentration measurements of around 450 0.3 μ M were consistent with offshore values reported in the literature (Hou et al, 2007). But our I⁻ 451 concentrations onshore were around 0.2 μ M, whilst they were in the range [0.04-0.08] μ M in the 452 middle of the English Channel. Interestingly, iodide concentrations around 0.15 µM were reported at 453 coastal sampling locations from North Germany (Hou et al, 2007). This difference in iodide partition 454 between coastal and offshore waters is in agreement with previous reviews by (Wong, 1991; Wong, 455 2001; Wong & Cheng, 2001a; Wong & Cheng, 2001b). Did the presence of local brown algae fields 456 influence the iodine chemical partition ($IO_3^- vs I^-$) in our coastal seawater sample? This question is now under further investigation (Carrano et al, 2021). Nevertheless, total dissolved iodine concentration 457 458 data in seawater were used to calculate the isotopic ratio of 129 in total iodine (IO₃⁻ + I⁻) measurements 459 in Goury (Figure 7 and Figure 8). Iodine content in brown seaweed clearly cycled with season, with 460 higher levels in winter than in summer (supplementary material S2-3). Does a higher growth rate in 461 summer overwhelm the iodine uptake capacity of the seaweed? Does solar exposure at low tide or 462 other stresses (thallus grazing by marine organisms) result in more volatile iodine release and loss? 463 Regardless of the cause, it was critical to consider iodine seasonal fluctuations in seaweed (Ar Gall et 464 al, 2004) when investigating radioactive iodine transfer between seawater and seaweed. The isotopic ratio ¹²⁹I/¹²⁷I in seaweeds and in seawater gave us a better understanding of iodine transfer. Up to 10-465 fold higher isotopic ratios were found in seaweed vs seawater (Figure 8). Although the ¹²⁹I activities 466 were lower in *F. serratus* than in *L. digitata* (Figure 7), the ¹²⁹I/¹²⁷I ratio values in the former were above 467 those in the latter (Figure 8). The discrepancy between the isotopic ratio $^{129}I/^{127}I$ in seawater and brown 468 seaweed was puzzling. Our net laboratory IO3⁻ and I⁻ flux experiments in L. digitata plantlets showed 469 470 that both forms of iodine are equally taken up by the seaweed. Therefore, a conversion of ¹²⁹Idischarged by the reprocessing plant into ¹²⁹IO₃⁻ would not result in a change in ¹²⁹I⁻ bioavailability for 471 kelp, as we had however proposed in our recent review (Fiévet et al, 2021). This also invalidates this 472 hypothesis to explain why the apparent ¹²⁹I⁻ bioavailability declined with distance from the plant outlet, 473 as we noticed in seaweed with regards to hydrodynamic dispersion modelling (Fiévet et al, 2021). We 474 cannot currently explain the difference in the isotopic ratio ¹²⁹I/¹²⁷I between seawater and brown 475 476 seaweed. Although we cannot rule out isotopic fractionation in the uptake mechanism, it seems 477 unlikely with such a small relative difference in the nucleus mass of the two iodine isotopes. A likely explanation was that, when looking at IO3⁻ + I⁻ in seawater, we missed an essential ¹²⁹I compartment 478 479 (chemical form) in the marine environment to which brown seaweed are exposed, an iodine compartment that remains to be identified. The difference in the isotopic ratio ¹²⁹I/¹²⁷I between the 480 two seaweed species that we analyzed may provide a clue. F. serratus grows in the mid intertidal zone 481 482 whilst L. digitata grows at the very bottom. It means that the former emerges twice a day at low tide 483 and the latter only a few times a month during spring tides. More frequent contact with the seawater 484 surface and longer contact with the atmosphere occurs for Fucus than Laminaria. To test this 485 hypothesis in a preliminary experimental setup, we detached adult L. digitata from the rocks and artificially maintained the seaweed in a floating net for 1 month (photo in the supplementary material 486 S2-4). At the same location, we compared ¹²⁹I activities in natural specimens from the bottom with 487 488 those from the net and observed a 50% increase (not shown). The thallus tissue was not visibly 489 degraded, and an identical ⁴⁰K level was taken as an indication of regular mineral balance. This experiment was anecdotal and should be confirmed and further characterized. The question raised by
the different isotopic ratio ¹²⁹I/¹²⁷I between seawater and brown seaweed is still open, but the sea
surface and its interface with the atmosphere may be a focal point when investigating iodine transfer
to seaweed in the marine environment.

494 **5 Conclusion**

495 The present paper aimed to improve our understanding of iodine transfers between seawater and 496 brown seaweed in the context of radioactive liquid discharges from nuclear facilities in the marine 497 environment. We are left with a series of questions: 1- The iodine uptake mechanism in brown 498 seaweed, which yields an exceptional bioaccumulation level, unknown in any other living organism, is 499 still mysterious. The difference of five orders of magnitude in concentration with respect to seawater 500 is even a low estimate, since iodine storage compartments are probably only peripheral in the 501 apoplast, as suggested by previous results in *L. digitata* (Lebeau et al, 2021; Verhaeghe et al, 2008). 502 However, our net IO_3^- and I^- flux experiments demonstrated that both iodine chemical forms are taken 503 up by L. digitata plantlets. The role of iodoperoxidases identified in this species and hypothesized to 504 be key to specific iodide uptake (Colin et al, 2005) remains to be further validated in vivo. In this new 505 context, a reduction process is likely to be a key step for completing the picture. 2- No step in the 506 radioactive iodine biological cycle between seawater and brown seaweed is expected to change the 507 isotopic ratio, except unlikely isotopic fractionation. However, our environmental data on the isotopic 508 ratio ¹²⁹I/¹²⁷I in seawater and brown seaweed exposed to liquid discharges from the ORANO 509 reprocessing plant in La Hague showed an unexpected discrepancy. This means that this in situ monitoring certainly missed a key compartment involved in ¹²⁹I transfer between seawater and brown 510 511 seaweed. The question raised in our previous review on radioactive transfers in the marine 512 environment around the reprocessing plant (Fiévet et al, 2021) regarding the non-conservative hydrodynamic dispersion of ¹²⁹I also remains open. Although important questions remain to be 513 addressed, our laboratory experiments provided a direct observation of the ability of marine brown 514 515 algae to modify iodine chemical speciation in seawater and to convert seawater IO_{3}^{-} into I⁻. The 516 endogenous extracellular pathways through which seaweed modify iodine chemistry in seawater 517 remain to be further investigated. This conversion is a key step in the Earth's biogeochemical iodine 518 cycle between the Ocean and the Atmosphere (Carpenter et al, 2021).

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529 8 Supplementary material

530 Additional data are provided in two files: Supplementary_Material_1.pdf and 531 Supplementary_Material_2.pdf.

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