1 Short-term iodine dynamics in soil solution

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8 ABSTRACT

Assessing the reactions of iodine (I) in soil is critical to evaluating radioiodine exposure and 9 10 understanding soil-to-crop transfer rates. Our mechanistic understanding has been constrained by 11 method limitations in assessing the dynamic interactions of iodine between soil solution and soil solid phase over short periods (hours). We use microdialysis to passively extract soil solution 12 spiked with radioiodine ($^{129}I^-$ and $^{129}IO_3^-$), to monitor short-term (≤ 40 hours) *in-situ* fixation and 13 speciation changes. We observed greater instantaneous adsorption of ¹²⁹IO₃⁻ compared to ¹²⁹I⁻ in 14 all soils and the complete reduction of ¹²⁹IO₃⁻ to ¹²⁹I⁻ within 5 hours of addition. Loss of ¹²⁹I from 15 solution was extremely rapid; the average half-life of ¹²⁹I⁻ and ¹²⁹IO₃⁻ in soil solution was 4.06 and 16 10.03 hours, respectively. We detected the presence of soluble organically bound iodine (org-¹²⁹I) 17 with a low molecular weight (MW) range (0.5-5 kDa) in all soils and a slower (20 to 40 hours) 18 time-dependent formation of larger MW org-I compounds (12-18 kDa) in some samples. This 19

study highlights the very short window of immediate availability in which I from rainfall or irrigation remains in soil solution and available to crops, thus presenting significant challenges to phytofortification strategies in soil-based production systems.

23 Introduction

Iodine (I) is an essential micronutrient for mammalian life, required for the synthesis of thyroid 24 25 hormones, approximately 1.9 billion people worldwide are estimated to be at risk of developing an iodine deficiency disorder.^{1, 2} Radioiodine (129 I; $t_{1/2} = 1.57 \times 10^7$ yr and 131 I; $t_{1/2} \otimes 0.02 \text{ d}$) released 26 as a consequence of anthropogenic activities poses a major ecological and health concern due to 27 its biophilic properties and relatively high environmental mobility.³ Once ¹²⁹I enters the 28 environment, it behaves similarly to ¹²⁷I, capable of bioaccumulating through the food chain and 29 subsequently stored in the thyroid where it can induce tumors.⁴⁻⁶ Understanding the 30 biogeochemical processes and mechanisms that affect I cycling in the terrestrial environment is 31 critical to evaluating the cause of I deficiency and potentially harmful exposure to radioiodine 32 33 isotopes.

Soil properties including pH, redox potential, concentration of soil organic matter (SOM) and 34 Fe/Al/Mn hydrous oxides affect I fixation, mobility and speciation.^{5, 7-11} Following natural and 35 anthropogenic I addition events (e.g. rainfall, irrigation, nuclear weapons testing), soluble 36 inorganic-I species, iodide (I⁻) and iodate (IO₃⁻), can be converted to soluble organic compounds 37 (org-I) and/or incorporated into the solid soil phase, bound to SOM or adsorbed to metal oxides.⁵⁻ 38 ^{7, 9, 12} It was demonstrated previously that IO₃⁻ and org-I are less mobile and have a greater affinity 39 for solid-soil components compared to I^{-.12-14} Metal hydrous oxides can influence iodine dynamics 40 in soils. Whilst ferric and aluminum oxides adsorb IO_3^- more strongly than I^{-} , ¹³ the presence of 41

42 manganese oxide birnessite (δ-MnO₂) can oxidize I⁻ to IO_3^{-} .¹⁵ However, in the presence of organic 43 matter (OM) the oxidation of I⁻ to IO_3^{-} is limited as the intermediate product, I₂, is incorporated 44 into OM (pH <7).¹⁶ Whilst the pH and redox potential of soils affects I speciation and mobility, 45 SOM appears to be the dominant factor controlling I retention in soils.^{12, 17, 18}

Microdialysis is an established method for neuroscience and pharmacokinetics and has been 46 used to investigate the mobility and bioavailability of macro-nutrients and metal ions in soil¹⁹⁻²²; 47 its applicability for following I reaction dynamics was previously evaluated by the authors.²³ The 48 49 passive sampling conditions of microdialysis enable the extraction of the soil solution phase with high temporal resolution and minimal disturbance to the soil structure. Unlike conventional soil 50 51 solution sampling methods which are destructive (e.g. centrifugation, high-pressure squeezing) or require soil to be close to saturation (e.g. Rhizon[™] samplers), microdialysis can perform 52 continuous extractions of the soil solution at a wide range of water contents (as low as ~50% water 53 54 holding capacity (WHC)). Evaluating soluble I without disturbing the ambient soil conditions, which could result in changes in I speciation, is vital for a more detailed understanding of I 55 biogeochemistry in soil. 56

In this study, we investigated short-term soil soluble I dynamics in three soils with contrasting physicochemical properties. Microdialysis was used to sample an isotopic I tracer (¹²⁹I) under ambient soil conditions to monitor changes in concentration and speciation following an addition event at 2.5 hour (hr) intervals over a 40 hr period. Iodine in the sampled dialysate was analyzed and speciated to determine: (i) the rate at which inorganic-I is removed from soil solution and adsorbed to the soil solid phase; and, (ii) the molecular weight (MW) range of soluble org-I compounds which are formed.

65 Materials and Methods

66 Soil sampling and physiochemical characteristics

Three topsoils (0 - 10 cm), from adjoining arable, grassland and woodland sites, were collected 67 from a location in Nottinghamshire, UK. The soils were sieved to <2 mm, homogenized and air-68 dried. Soil pH was determined using an Orion pH meter after equilibrating 5 g of soil in 12.5 mL 69 of 0.01 M CaCl₂ for 30 min. Loss-on-ignition (LOI), as an estimate of SOM content, was 70 determined after heating in a muffle furnace at 450 °C for 16 hr, after an initial drying period. The 71 water holding capacity (WHC) of the soil was determined gravimetrically in triplicate according 72 to Grace, et al.²⁴. Total soil ¹²⁷I was extracted with 5% tetramethylammonium hydroxide (TMAH) 73 at 70 °C for 3 hr (with shaking after 1.5 hr); once cooled, 5 mL of Milli-Q water was added and 74 the bottles were centrifuged (20 min at 3500 rpm) and the supernatant solutions retained for 75 analysis by ICP-QQQ.²⁵ To determine the concentrations of reactive iron, aluminum, and 76 77 manganese hydrous oxides, soils were extracted with dithionite-citrate-bicarbonate (DCB) solution, centrifuged (20 min at 2500 rpm), filtered (<0.22 µm) and the supernatant solutions 78 retained for analysis by ICP-QQQ.²⁶ Particle size analysis was determined using the method 79 described in Rawlins, et al. ²⁷, using a laser diffraction particle size analyzer. 80

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82 Soil incubation and microdialysis sampling

The radioiodine tracer (¹²⁹I) was obtained from the American National Institute of Standards as NaI (NIST, Gaithersburg, Maryland, USA; CRM 4949C, 0.004 mol L⁻¹ Na¹²⁹I, 3451 Bq mL⁻¹). The primary stock (¹²⁹I⁻) was made up to 100 mL with 0.01 M NaOH, as recommended by the suppliers. Iodate (¹²⁹IO₃⁻) was prepared from the ¹²⁹I⁻ stock by oxidation, using a method adapted from Yntema and Fleming ²⁸. To 50 mL of the ¹²⁹I⁻ stock, 5 mL of 0.1 M HCl was added in an

initial neutralization step, followed immediately by 5 mL of 0.2 M sodium chlorite for oxidation. 88 Successful oxidation to ¹²⁹IO₃⁻ was confirmed by anion exchange LC-ICP-OOO. The water 89 content of the air-dried soils (~90 g dry weight (dw)) was raised to 50% of WHC with Milli-Q 90 water (18.2 M Ω cm; Millipore) and allowed to equilibrate for 10 days at 20 ± 0.5 °C in the dark. 91 The moist soils were then homogenized with equivalent volumes of ¹²⁹I⁻ or ¹²⁹IO₃⁻ in solution to 92 bring the soils to 80% WHC with a final ¹²⁹I concentration of 0.2 mg kg⁻¹ (dw basis). The soils 93 were then distributed between triplicate 50 mL Corning® polypropylene centrifuge tubes (~30 g 94 95 dw of soil per replicate).

The microdialysis system consisted of a syringe pump (KD Scientific Legato 200 Series), 96 97 equipped with four plastic syringes (BD Plastipak; 20 mL) used to deliver the perfusate solution. Syringes were attached to microdialysis probes CMA 20 (10 mm length, 500 µm outer and 400 98 µm inner diameter) with a polyethersulfone (PES) membrane (100 kDa molecular weight cut-off 99 (MWCO). Within two minutes of ¹²⁹I addition to soil, incision holes were made, using an 100 introducing needle, and the microdialysis probes were inserted ~ 1.5 cm beneath the soil surface. 101 Probes were perfused with Milli-Q water at a flow rate of 5 µL min⁻¹ and dialysate solutions 102 collected at 2.5 hr intervals in glass microvials over a 40 hr extraction period. Samples were 103 initially stored in the refrigerated micro-fraction collector (6 °C; CMA 470) before freezing at -20 104 °C until defrosting immediately before analysis. Throughout the extraction, soils were kept in the 105 dark at a constant temperature of 20 ± 0.5 °C. The probes and micro-fraction collector were from 106 CMA Microdialysis AB (Kista, Sweden). In order to assess the concentration (µg L⁻¹) of ¹²⁹I in 107 the soil solution a correction factor was applied to the measured dialysate solutions as Csoil solution 108 $= C_{\text{dialvsate}}/(1 - e^{-PS/F})$. Where $C_{\text{soil solution}}$ and $C_{\text{dialvsate}}$ are the concentrations of the target analyte in 109

the external soil solution and the dialysate solution respectively, *PS* in the surface area of the microdialysis membrane and *F* is the perfusion flow rate, adapted from Clough ²⁹.

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113 Total iodine analysis

Total ¹²⁷I concentrations were determined in single MS, no-gas mode (Agilent 8900 ICP-QQQ, 114 Agilent Technologies). All ¹²⁹I concentrations were measured using an ICP-OOO (Agilent 8900, 115 Agilent Technologies). Analysis was conducted using both quadrupoles and the reaction cell, to 116 remove isobaric and polyatomic interferences, a full description of ¹²⁹I analysis can be found in 117 Humphrey, et al.³⁰. Due to the small sample volume, we used an Agilent 1260 Infinity Bio-inert 118 Quaternary LC pump and high performance autosampler to act as a low volume autosampler: vials 119 were also equipped with flat bottomed glass inserts to further reduce the sample volume 120 requirement. Throughout the analytical run, an in-house quality control sample with a known 121 concentration of 5 μ g L^{-1 129}I was measured with an average recovery of 101% (n = 28) and a limit 122 of detection (LOD) (3 x SD blanks) of 0.0075 μ g L⁻¹ for ¹²⁹I. 123

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125 Iodine speciation analysis

Inorganic-I (¹²⁹I⁻ and ¹²⁹IO₃⁻) concentrations were measured by ICP-QQQ following on-line chromatographic separation in isocratic mode using a Hamilton PRP X-100, anion exchange column (250 mm × 4.6 mm, 5 µm) and a samples injection volume of 25 µL. The mobile phase was 100 mM NH₄NO₃, adjusted to pH 9.5 with TMAH (25%), at a flow rate of 1.5 mL min⁻¹. Quality control samples with a known I concentration of 5 µg L⁻¹, ¹²⁹I⁻ and ¹²⁹IO₃⁻, were measured with an average recovery of 97% and 96% (n = 22), respectively. The LOD for ¹²⁹I⁻ and ¹²⁹IO₃⁻ was 0.007 and 0.006 µg L⁻¹, respectively.

The MW distribution of soluble org-I was determined using size exclusion 133 chromatography (SEC) hyphenated with UV absorbance detection and ICP-QQQ. An Agilent 134 1260 Infinity Bio-inert Quaternary LC pump and high performance autosampler in isocratic mode 135 with an AdvanceBio SEC 130 Å (2.7 µm, 4.6 x 300 mm column, MW range 100-120,000 Da, 136 Agilent) was used as the sample introduction system. Samples (10 µL) were injected onto the 137 column with a mobile phase (flow rate of 0.35 mL min⁻¹) consisting of 150 mM sodium phosphate 138 buffer, pH 7.0. Platinum tip sampler and skimmer cones were used on the ICP-QQQ due to the 139 140 high sodium concentration in the mobile phase. The UV detector scanned at 220 nm to measure the calibration protein standards, and 254 nm to detect dissolved organic carbon.^{9, 31-35} Protein 141 142 standards of known MWs: Ovalbumin (45 kDa); Myoglobin (17 kDa); Aprotinin (6.7 kDa); Neurotensin (1.7 kDa); and Angiotensin II (1 kDa) (AdvanceBio SEC 130Å protein standard, 143 Agilent) were used to calibrate the column. 144

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146 **Results and Discussion**

147 *Soil characteristics*

The soil physiochemical characteristics in Table 1 reflect their land use. Compared to the arable and grassland soils the acidic woodland soil had a much greater LOI, extractable Fe/Al/Mn hydrous oxide content and total I concentration. Given that all three soils were collected with 2.5 km of each other the greater I concentration in the woodland soil may reflect a greater ability to retain I from precipitation and the absence of annual crop removal.

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Table 1 Summary of soil physiochemical characteristics. Values expressed as mean ± standard
error (SE) (n=3)

	Units	Arable	Grassland	Woodland
Location (long, lat decimal degrees)		52.9001, -1.0884	52.8890, -1.0877	52.8986, -1.0744
Elevation	m	95	83	121
pH (CaCl ₂)		6.9	5.9	3.6
LOI	%	5.1	7.0	50.6
WHC	%	36.8	41.5	70.8
¹²⁷ I	mg kg ⁻¹	3.38 ± 0.05	3.63 ± 0.05	12.9 ± 0.10
¹²⁹ I	µg kg ⁻¹	<dl< td=""><td><dl< td=""><td><dl< td=""></dl<></td></dl<></td></dl<>	<dl< td=""><td><dl< td=""></dl<></td></dl<>	<dl< td=""></dl<>
Fe ₂ O ₃	%	0.180 ± 0.02	0.0131 ± 0.001	0.696 ± 0.06
Al(OH)3	%	0.0379 ± 0.003	0.0523 ± 0.007	0.925 ± 0.05
MnO ₂	%	0.0594 ± 0.005	0.0226 ± 0.003	0.622 ± 0.07
Texture (clay/silt/sand)	% mass	41/41/18	17/19/64	24/35/41
Texture classification		Clay	Sandy Loam	Clay Loam

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157 Total ¹²⁹I dynamics in soil solution

Total concentrations of soluble ¹²⁹I sampled from the soils spiked with ¹²⁹I⁻ or ¹²⁹IO₃⁻ are shown

159 in Figure 1.





Figure 1 Soluble ¹²⁹I (µg L⁻¹) sampled using microdialysis from arable, grassland and woodland 163 soils at 2.5 hour intervals over 40 hours. Symbols represent mean values obtained in triplicate, 164 error bars indicate \pm S.E. An asterisk at the top left of each chart denotes the initial spike 165 concentration; white circles (\circ) represent soils spiked with ¹²⁹I⁻; black triangles (\blacktriangle) represent soils 166 spiked with ¹²⁹IO₃⁻. Note the log scale on the Y-axis. The dashed line indicates the detection limit. 167 Each microdialysis dialysate sample was sampled at 5 μ L min⁻¹ for 2.5 hours. 168

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Losses of ¹²⁹I from soil solution were observed in all the soils within 2.5 hr, irrespective of the 170 initial spike speciation (¹²⁹I⁻ or ¹²⁹IO₃⁻). The concentration of ¹²⁹I in the first dialysate samples (0-171 2.5 hr) were significantly higher in soils treated with ¹²⁹I⁻ compared to ¹²⁹IO₃⁻. The significantly 172 larger initial loss of ¹²⁹IO₃⁻ from solution compared to ¹²⁹I⁻ observed in this study may be a 173 combination of adsorption on inorganic soil phases or, rapid immobilization within SOM.⁵ 174

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of SOM⁶ and the likelihood of it meaningfully contributing to total loss rates is low.³⁶ Therefore, 176 adsorption onto inorganic soil phases and rapid immobilization by SOM are far more likely reasons 177 for the losses of ¹²⁹I from soil solution; both of these processes are pH-dependent. Prior to the 178 incorporation of IO3⁻ with OM studies have shown that a reduction to an electrophilic intermediate 179 species, such as HOI or I₂ occurs, this reduction occurs much faster under acidic conditions (pH< 180 6).^{16, 37-40} Clays, hydrous oxides and SOM can absorb I⁻, where sorption generally increases with 181 decreasing pH, as observed in this study.⁴¹⁻⁴³ In previous studies it has been shown that when soil 182 pH >6.0 I⁻ sorption is predominantly influenced by OM, however, under more acidic conditions 183 the role of specific adsorption to iron and aluminum oxides becomes increasingly important.^{44, 45} 184 The faster immediate sorption of ¹²⁹IO₃⁻ compared to ¹²⁹I⁻ highlights the different adsorption 185 mechanisms of inorganic-I in soils. 186

Volatilization has previously been reported to occur from soils, but is suppressed by the presence

The significant proportion of apparent instantaneous loss of ¹²⁹IO₃⁻ from solution in all 187 experimental soil and the highest loss rates of ¹²⁹I⁻ from soil solution is consistent with previous 188 results.⁵ First-order kinetic equations were used to assess the rate at which total ¹²⁹I was removed 189 from soil solution as: $C_t = C_0 e^{-kt}$, where C_t is the ¹²⁹I concentration at time t (hr) (mg L⁻¹), C_0 is 190 the initial concentration of 129 I in soil solution (mg L⁻¹), and k is the first-order rate constant (hr⁻¹). 191 The half-life in soil solution was then calculated from $\ln(2)/k$. The average half-life of soils spiked 192 with ${}^{129}I^-$ and ${}^{129}IO_3^-$ was 4.06 \pm 0.96 and 10.03 \pm 0.78 hr, with r^2 values of 0.77 and 0.45, 193 respectively. Shetaya, et al.⁵ investigated I dynamics in soils and reported that the loss of I⁻ from 194 solution was extremely rapid, reaching completion over minutes-hours, whereas IO3⁻ loss from 195 solution was slower, typically occurring over hours-days, which is consistent with the results in 196 the present study. However, in the current study the use microdialysis has enabled far greater 197

198 sampling resolution compared to previous studies at time intervals within a similar range to the 199 reaction times reported by Shetaya, et al. ⁵. The results presented in this paper indicate that iodine 200 adsorption occurs more rapidly in soils under realistic conditions than previously observed using 201 batch sorption experiments and chemical extraction approaches.

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203 ¹²⁹I speciation in soil solution

The concentrations of soluble inorganic ($^{129}I^- + {}^{129}IO_3^-$) and organic (total ^{129}I minus the sum of inorganic species) ^{129}I in the three soils spiked with either $^{129}I^-$ or $^{129}IO_3^-$ are shown in Figure 2. A combined inorganic fraction was calculated due to the concentrations of $^{129}IO_3^-$ present in the soils treated with either $^{129}I^-$ or $^{129}IO_3^-$ falling below the LOD (shown in Figure S1).

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Figure 2 Soluble ¹²⁹I speciation of (a) iodate and (b) iodide spiked arable, grassland and woodland soils over 40 hours. Symbols represent the mean values obtained in triplicate, bars indicate \pm SE. White circles (\circ) represent inorganic-¹²⁹I concentrations; black triangles (\blacktriangle) represent soluble

org-¹²⁹I concentrations. Note the log scale on the primary Y-axis. Samples below the detection limit were omitted from the plot. Microdialysis dialysate was sampled at 5 μ L min⁻¹ for 2.5 hours.

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The proportion of inorganic-¹²⁹I present in soil solution was greater compared to org-¹²⁹I in soils spiked with I⁻, however, this trend was not observed consistently in IO₃⁻ spiked soils. In the arable soil spiked with ¹²⁹IO₃⁻, where the org-¹²⁹I became the dominant fraction in samples after 25 hr, when inorganic-¹²⁹I concentrations were below the LOD. In contrast, in woodland soil spiked with ¹²⁹IO₃⁻ the soluble org-¹²⁹I fraction was below the LOD 10 hr after addition, and only inorganic species were detected.

The speciation of stable and radioiodine present in soil systems has been the focus of several 222 studies^{9, 46-48} with I⁻ and org-I identified as the dominant species present. When ¹²⁹IO₃⁻ was added 223 to the soils we observed an immediate reduction to ¹²⁹I⁻ in soil solution, with very low 224 concentrations of ¹²⁹IO₃⁻ remaining in soil solution beyond 2.5 hrs (Figure S1, d). Reduction of 225 IO₃⁻ to I⁻ in soils has previously been observed^{49, 50} and may arise from biotic processes. However, 226 it seems more likely that reduction is caused by abiotic interactions with solid or dissolved OM as 227 the reaction rates appear to be too rapid for biological processes.⁵ The rapid reduction reported in 228 this study is much faster compared to previous work, where the reduction of IO3⁻ occurred over 229 days.⁵¹ Possible reasons for the discrepancies between the studies include the significantly lower 230 concentrations of ¹²⁹I used in this study and the lower soil pH used in this study as the reduction 231 of IO₃⁻ to the reactive iodine species is thermodynamically more favorable at lower pH. 232

In contrast, when soils were spiked with $^{129}I^-$, a very low concentration of $^{129}IO_3^-$ was detected in the arable and grassland soils within 7.5 hr, indicating that limited $^{129}I^-$ oxidation to $^{129}IO_3^-$ had occurred (Figure S1). We only observed I⁻ oxidation in the arable and grassland soils; there was

no evidence for oxidation in the organic-rich acidic woodland soil (Figure S1). Factors controlling 236 ¹²⁹I⁻ oxidation in soil remain unclear.⁵² The conversion of I⁻ to org-I requires I⁻ to be initially 237 oxidized to an intermediate such as I2 or HOI.^{16, 37} Iodide-oxidizing bacteria in soils have 238 previously been observed to influence I⁻ sorption/incorporation into SOM and oxidation to IO₃⁻ 239 and to produce volatile org-I compounds such as CH₃I.^{53, 54} In addition, metal oxides (Al, Fe, Mn) 240 and humic substances can act as oxidizing agents. In batch kinetic experiments pH and metal oxide 241 concentrations significantly influence reaction rates, with faster oxidation occurring at lower pH 242 and greater metal oxide concentrations.^{55, 56} However, in our *in-situ* experiments, we did not 243 observe I⁻ oxidation in the acidic, Fe/Al/Mn rich woodland soil. Figure 2 shows that the greatest 244 245 initial loss of I⁻ occurred in the woodland soil, which had the largest metal oxide concentrations, an order of magnitude higher than the arable and grassland soils, highest organic matter and the 246 lowest pH. The rapid loss of I⁻ from solution in organic-rich soils has been described as a first-247 order reaction.⁵⁷ Previous studies have demonstrated that I⁻ can easily be remobilized as it is only 248 bound by weak electrostatic attraction to solid soil media,⁵⁷ therefore, further studies assessing the 249 speciation of iodine during dynamic processes such as rainfall events or plant uptake should be 250 investigated to assess iodine mobility in soils. 251

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253 Molecular weight distribution of dissolved organic ¹²⁹I

The presence of org-¹²⁹I was confirmed using SEC-ICP-QQQ; the presence of multiple peaks indicates that soluble org-I exists at a range of MWs. Time-dependent changes in speciation and molecular weight of ¹²⁹I in the soil solution phase are shown in Figure 3.





Figure 3 Molecular weight distribution of soluble isotopically labeled ¹²⁹I in arable, grassland and woodland soils spiked with ¹²⁹IO₃⁻ or ¹²⁹I⁻ sampled over 40 hours, separated by size exclusion chromatography. Note the log Y-axis. High molecular weight organically bound I (HMW org-I), low molecular weight organically bound I (LMW org-I) and ¹²⁹I⁻ retention times labeled.

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Within all three experimental soils, for addition of both ¹²⁹I⁻ and ¹²⁹IO₃⁻, chromatographic peaks were observed for ¹²⁹I⁻ between 15 and 18 min that decreased over the 40 hr extraction period. The intensity of the ¹²⁹I⁻ peak was considerably less in the ¹²⁹IO₃⁻ treated soils due to the higher rates of instantaneous adsorption on inorganic soil phases and rapid immobilization by SOM. All soils spiked with ¹²⁹I⁻, had a broad peak representing LMW org-¹²⁹I between 9 and 15 min; the overall intensity of the org-I peak decreased over the 40 hr sampling period. There was a shift in the MW distribution in the arable soil spiked with ¹²⁹I⁻ between 20 and 40 hr when the signal intensity of

the LMW org-I peak decreased and a HMW org-I peak, with a retention time between 5 and 10 271 min emerged. In contrast, within the grassland soil spiked with ¹²⁹I⁻, dissolved org-¹²⁹I had a lower 272 MW distribution; no org-I compounds >5 kDa were present throughout the sampling period. The 273 emergence and rapid decline of LMW org-¹²⁹I between 0.5-5 kDa in the woodland soil further 274 illustrates how quickly initially soluble I is removed from solution in organic rich soils. However, 275 within the arable and grassland soils, the MW distribution of the org-I underwent various changes 276 within the 40 hr sampling period. The UV absorbance results confirmed that soluble organic 277 compounds had a broad MW range (Figure S2). 278

The transport behavior of inorganic-I in soils may be significantly impacted by the fixation of 279 dissolved I into macromolecular humic substances, however, many of the mechanisms controlling 280 the rate of fixation remain unknown.^{33, 38, 58, 59} Xu, et al.³ demonstrated that higher proportions of 281 125 I⁻ and 125 IO₃⁻ were associated with organic compounds <3 kDa, 16 and 20% compared to 6 and 282 283 3% in the >3 kDa fraction, respectively. In addition, they found that binding sites became saturated and less available when the treatment concentration increased.³ In resuspension experiments, Xu, 284 et al.¹⁸ found that mobile ¹²⁹I was associated with amphiphilic organic compounds with an average 285 MW between 13.5 and 15 kDa. Figure 3 shows that higher proportions of soluble inorganic-I were 286 incorporated into organic compounds with a low MW (<5 kDa). However, there was evidence of 287 larger dissolved org-I compounds with a MW between 12 and 18 kDa 20 hr after ¹²⁹I addition, in 288 agreement with Xu, et al.³. 289

The time-dependent formation of these larger dissolved org-I compounds could be due to smaller dissolved compounds binding together via weak chemical forces or the influence of microbial activity. The iodination of SOM in acidic conditions was shown to be predominantly an abiotic process, however, in less acidic conditions (pH \geq 5), microbial assisted iodination of SOM was

observed.³ Whilst microbial activity is generally not deemed essential for the transformation of IO₃⁻ into org-I (which seems to be primarily controlled by abiotic processes⁶), microbial laccases have been shown to enhance I⁻ sorption in soils.⁵³ Microbiological activities have previously been reported to incorporate I⁻ with humic acids and soils with a relatively HMW and low mobility.³³ Heumann, et al. ⁶⁰ identified that in the presence of microorganisms the production HMW org-I compounds significantly increased over an 8 week period; in contrast Figure 3 demonstrates that formation of relatively HMW compounds occurred over 20-40 hrs.

301 It has previously been suggested that the rapid rate of reaction of I with SOM could be beneficial, as humus may act as a natural barrier, accumulating any radioiodine released from nuclear waste 302 repositories as immobile org-I compounds.¹⁸ Following a radioiodine contamination event limited 303 soil-to-crop transfer would be beneficial for reducing the risk and duration of potentially harmful 304 exposure.^{18, 61, 62} The short-term experiments presented in this paper show that a relatively small 305 proportion of inorganic-¹²⁹I is rapidly incorporated into LMW organic compounds (<2.5 hr) which, 306 over time, can be sorbed to the solid soil phase or bind together to form HWM organic compounds 307 (~30 hr). 308

Evaluating the short-term transport, speciation and fate of I in soil is critical to assess the 309 environmental mobility and plant availability, as ¹²⁷I is an essential micronutrient for which dietary 310 intake often depends on the transfer of I from soil-to-crops.⁶³ Soil I concentrations are dependent 311 on the deposition of volatized I compounds from seawater.¹¹ Specific rainfall characteristics 312 influence the concentration of I in rainfall and soil properties affect the volume of I retained by 313 soils that would be accessible for plant uptake.⁵ Passive uptake is the predominant pathway for I 314 absorption in plants,³⁰ as such, plant uptake is highly dependent upon the concentration in the soil 315 solution phase. By using ¹²⁹I as a proxy for ¹²⁷I, it is possible to predict how much soluble I is 316

317 available for plant uptake following an addition event. We demonstrate that I only remains in the soil solution phase for a very limited period, before being sorbed on to inorganic soil phases or 318 immobilized by SOM. Consequently, there is a severely limited window of opportunity for plants 319 to absorb I from the soil solution immediately following rainfall or irrigation and limits the transfer 320 of I to animal and human diets. However, this study did not investigate the impact of rewetting 321 and remobilization of I bound to the solid soil phase, and this is should be the subject of further 322 research. Considering that there are an estimated 1.9 billion people at risk of I deficiency 323 worldwide,⁶⁴ the findings in this paper may contribute to better understanding of the efficacy of 324 phytofortification strategies. 325

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327 Supporting Information

328 Additional information on analytical methods and supplementary results.

329 TOC GRAPHIC FOR MANUSCRIPT



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- 340 The manuscript was written through contributions from all authors. All authors have given
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342 **Conflict of Interest**

343 The authors declare no competing financial interest.

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