

23 **Highlights:**

- 24
- Oxidation of iodide to iodate by marine nitrifying bacteria demonstrated for first time
- 25
- Laboratory cultures of ammonium oxidising bacteria produced iodate from iodide substrate
- 26
- Nitrification used to parameterise iodide sink in global marine iodine cycling model
- 27
- Changes in nitrification may increase sea surface iodide, impacting atmospheric chemistry

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31

32 **Abstract**

33 Reaction with iodide (I) at the sea surface is an important sink for atmospheric ozone, and causes
 34 sea-air emission of reactive iodine which in turn drives further ozone destruction. To incorporate this
 35 process into chemical transport models, improved understanding of the factors controlling marine
 36 iodine speciation, and especially sea-surface iodide concentrations, is needed. The oxidation of I to
 37 iodate (IO₃⁻) is the main sink for oceanic I, but the mechanism for this remains unknown. We
 38 demonstrate for the first time that marine nitrifying bacteria mediate I oxidation to IO₃⁻. A significant
 39 increase in IO₃⁻ concentrations compared to media-only controls was observed in cultures of the
 40 ammonia-oxidising bacteria *Nitrosomonas* sp. (Nm51) and *Nitrosococcus oceani* (Nc10) supplied
 41 with 9-10 mM I, indicating I oxidation to IO₃⁻. Cell-normalised production rates were 15.69 (±4.71)
 42 fmol IO₃⁻ cell⁻¹ d⁻¹ for *Nitrosomonas* sp., and 11.96 (±6.96) fmol IO₃⁻ cell⁻¹ d⁻¹ for *Nitrosococcus oceani*,
 43 and molar ratios of iodate-to-nitrite production were 9.2±4.1 and 1.88±0.91 respectively. Preliminary
 44 experiments on nitrite-oxidising bacteria showed no evidence of I to IO₃⁻ oxidation. If the link
 45 between ammonia and I oxidation observed here is representative, our ocean iodine cycling model
 46 predicts that future changes in marine nitrification could alter global sea surface I fields with
 47 potential implications for atmospheric chemistry and air quality.

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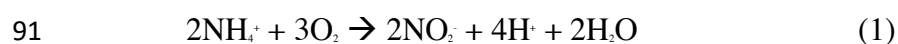
51 **Introduction**

52 Iodine plays an important role in catalytic ozone destruction and new particle formation in the
 53 troposphere, thereby impacting the oxidative capacity of the atmosphere (Sherwen *et al.*, 2016) and
 54 the Earth's radiation balance (O'Dowd *et al.*, 2002). Sea-to-air iodine transfer is known to be the
 55 main source of iodine to the atmosphere (Carpenter, 2003; Sherwen *et al.*, 2016). Reactive inorganic
 56 iodine (I₂, HOI) emissions resulting from the reaction of gas-phase ozone with sea surface iodide (I⁻)
 57 is now thought to be the dominant mechanism mediating sea-air iodine emissions (Carpenter *et al.*,
 58 2013). The strength of the surface reactive iodine flux is related to sea surface I concentrations
 59 (Carpenter *et al.*, 2013) so knowledge of ocean I distributions is required in order to estimate the
 60 significance of this process. Furthermore, a detailed understanding of the processes controlling
 61 inorganic iodine speciation is needed to allow us to develop predictive capacity regarding sea surface
 62 I₂, ozone-deposition rates and sea-air emission of reactive iodine.

63 Total inorganic iodine is found at 400-500 nM in seawater and predominantly exists as iodate (IO₃⁻)
 64 and I⁻ (Chance *et al.*, 2014) with inter-conversion between these two species alongside physical
 65 mixing being the main causes of spatial and temporal variability in sea surface I. Iodate is the
 66 thermodynamically stable form and the dominant form in the deep ocean. The existence of relatively
 67 higher levels of I⁻ in the euphotic zone (reviewed by Chance *et al.*, 2014) has led to the suggestion
 68 that IO₃⁻ reduction to I⁻ is linked to primary productivity. This theory has been supported by
 69 observations of I⁻ production in cultures of a wide range of marine phytoplankton (e.g. Chance *et al.*,
 70 2007; Bluhm *et al.*, 2010) and some field studies (Chance *et al.*, 2010). Proposed mechanisms for IO₃⁻
 71 reduction to I⁻ by marine phytoplankton include nitrate reductase enzymes (Hung *et al.*, 2005) and
 72 reactions of iodate with reduced sulphur species exuded from cells during senescence (Bluhm *et al.*,
 73 2010), but neither has yet been confirmed as the dominant route of conversion. I⁻ oxidation to IO₃⁻ is
 74 also known to occur with rate estimates ranging from ~4 to 670 nM yr⁻¹ (reviewed in Chance *et al.*,
 75 2014). Abiotic oxidation of I⁻ back to IO₃⁻ in the ocean (e.g. by oxygen, hydroxyl radicals, hydrogen

76 peroxide and ozone) is thought to occur so slowly as to be insignificant (e.g. Wong, 1991), and so I
 77 oxidation to IO₃ is also thought to be associated with marine microbiological activity. The rates and
 78 processes involved in I to IO₃ oxidation are associated with large uncertainty (Truesdale *et al.*, 2001;
 79 Amachi *et al.*, 2008), and the mechanisms involved remain undefined. This uncertainty has been
 80 suggested to be one of the factors hindering the development of mathematical models of iodine
 81 transformations in the global oceans (Truesdale *et al.*, 2001).

82 I oxidation to I₂ has been observed in bacterial isolates obtained from a range of environments
 83 including seawater aquaria (Gozlan *et al.*, 1968), natural gas brines (Iino *et al.*, 2016) and
 84 seawater/marine mud (Fuse *et al.*, 2003). Additionally, based on field observations, a number of
 85 studies (Truesdale *et al.*, 2001; Žic *et al.*, 2013) have proposed that I oxidation to IO₃ is linked to
 86 nitrification in marine systems. Nitrification is the two-stage biological transformation of ammonia
 87 (NH₃) to nitrate (NO₃) (Equations 1 and 2; Koops & Pommerening-Röser, 2001) mediated by
 88 chemoautotrophic ammonia-oxidising bacteria (AOB), and nitrite-oxidising bacteria (NOB).
 89 Previously thought to only occur outside of the euphotic zone, nitrification is now known to occur
 90 throughout the oceanic water-column (reviewed by Yool *et al.*, 2007).



93 A link between I oxidation/ IO₃ production and nitrification is yet to be confirmed but, if established,
 94 would suggest that I oxidation to IO₃ is widespread throughout the world's oceans (Yool *et al.*,
 95 2007).

96

97 The primary aim of this study was to establish whether I oxidation to IO₃ is associated with marine
 98 nitrification. Our objectives were to determine if IO₃ production occurs in cultures of marine

99 ammonia- and nitrite-oxidising bacteria supplied with I, determine the relative rates of IO₃
 100 production and nitrification and explore the possible implications of the findings.

101

102 **Methods**

103 ***Cultures***

104 Two AOB cultures (*Nitrosomonas* sp. [Nm51] and *Nitrosococcus oceani* [Nc10]) were investigated
 105 for IO₃ production in the presence of I as the only iodine source. Cultures were grown in the dark in
 106 a water bath at 25 °C in autoclaved ESAW artificial seawater mixture (Berges *et al.*, 2001) made up
 107 using distilled water. The ESAW media was supplemented with 7-8 mM ammonium chloride and
 108 potassium phosphate. We also conducted preliminary tests on three active marine NOB (*Nitrospira*
 109 *marina* [295], *Nitrospina gracilis* [3/211], *Nitrococcus mobilis* [231]) but saw no evidence of IO₃
 110 production in any of the cultures studied. These results are not discussed further. Handling of
 111 cultures was done at all times in a biosafety cabinet using sterile equipment.

112

113 ***Experimental Set Up***

114 For the AOB experiments triplicate cultures were incubated alongside triplicate media-only controls
 115 for periods of 8-12 days. The experiments were kept as short as possible to avoid significant changes
 116 in pH in the bulk media which would impact inorganic iodine speciation. Hence experiments were
 117 only run until an increase in nitrite across two time-points was observed. Samples were taken at
 118 regular intervals of between 1 to 6 days for pH measurement, cell counts and determination of NO₂⁻,
 119 IO₃⁻, I and NH₄⁺/NH₃ concentrations. In all cases, I (Aristar) was added to be at similar concentrations
 120 with the NH₄⁺ required in the growth media. The levels of I are much higher than those encountered
 121 in the oceans (global ocean median=77 nM I [interquartile range 28-140 nM], Chance *et al.*, 2014)
 122 but were chosen to be similar to the levels of NH₄⁺. This is because in the marine environment

123 nitrifiers would be exposed to similar ratio of NH_4^+ and I. For example, Rees *et al.* (2006) show that
 124 $\text{NH}_4^+/\text{NH}_3$ occurs at concentrations ranging from 60-300 nM in the Atlantic between 60°N to 50°S.

125

126 ***pH***

127 A spectrophotometric method using a Lambda 25 UV/Vis spectrophotometer (Perkin-Elmer) and m-
 128 cresol purple dye (Dickson *et al.*, 2007) with measurements at 730, 578 and 434 nm was used to
 129 determine pH in the cultures and media-only controls. Salinity, needed for the pH calculation, was
 130 calculated from conductivity measured using a calibrated Hanna Instruments hand-held probe.

131

132 ***Cell counts***

133 Immediately after sampling, 4 mL of the culture was fixed with 15 μL of 50% glutaraldehyde (Alfa
 134 Aesar), flash frozen in liquid nitrogen and placed in a -80 °C freezer for later determination of cell
 135 density. Cell counts were made using a Beckman Coulter Cytoflex S flow cytometer (flow rate of 10
 136 $\mu\text{L min}^{-1}$) within 2 months of collection. DAPI (Sigma; 2 $\mu\text{g mL}^{-1}$) stained samples were excited by a
 137 laser at 405 nm and the emitted fluorescence detected using an avalanche photodiode detector with a
 138 reflective band pass filter 450/45. The flow cytometer thresholds were set using the 405 nm laser
 139 side scatter and the DAPI fluorescence signals.

140

141 ***Nitrite concentration***

142 NO_2^- was measured in 0.45 μm (Millex) filtered samples using a spectrophotometric method
 143 (Lambda 25 UV/Vis spectrophotometer, Perkin-Elmer) developed by Norwitz & Keliher (1984). The
 144 method involves diazotizing nitrite with sulfanilamide (Fisher, analytical reagent grade) and coupling
 145 with N-1-naphthylethylenediamine dihydrochloride (Fisher, analytical reagent grade) to form a

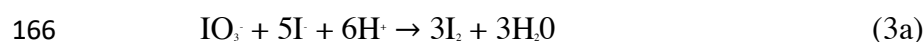
146 coloured azo dye which is measured spectrophotometrically at 540 nm. The method was calibrated
 147 using NaNO₂ standards (Fisher, analytical reagent grade) prepared in the ESAW-based media.

148

149 ***Iodate Concentration***

150 IO₃⁻ concentrations were measured in 0.45 μm (Millex) filtered samples using a manual version of the
 151 spectrophotometric (Lambda 25 UV/Vis spectrophotometer) method detailed in Truesdale &
 152 Spencer, 1974 and Jickells *et al.*, 1988. Absorbance was measured at 350 nm. Strictly, this method
 153 determines all oxidised (0 to +5 oxidation state) forms of inorganic iodine, but in seawater derived
 154 media this is predominantly IO₃⁻, and so will be referred to as IO₃⁻ iodate hereafter. The method was
 155 calibrated using potassium iodate (Aristar) standard solutions made up in ESAW.

156 Some validation and modification to the method was required due to the nature of our experimental
 157 set-up. Chapman & Liss (1977) show that NO₂⁻ can interfere with spectrophotometric IO₃⁻
 158 measurements (using sulfamic acid) at ambient seawater concentrations with a 15% error. Clearly
 159 significant interference would be an issue for our experiments where NO₂⁻ was being produced so we
 160 ran tests. We found that the presence of NO₂⁻ up to 10 μM had negligible impact on IO₃⁻ measurements
 161 (between 0.1-50 μM). We did however identify that the high starting concentration of I⁻ (~10 μM) in
 162 the culture media was problematic. The iodate analysis method comprises two steps: the first
 163 involves an initial absorbance reading after the addition of sulfamic acid; the second involves the
 164 addition of excess I⁻. Under acidic conditions I⁻ reacts with IO₃⁻ to form I₂ (equation 3a) which reacts
 165 with excess I⁻ to form the coloured ion I₃⁻ (equation 3b) that can be measured spectrophotometrically.



168 The difference between the first and second absorbance readings is then used to calibrate the method.
 169 In the case of our experiments the media already contained excess I so the formation of I₂ and I₃⁻ was
 170 initiated as soon as the acid was added in the first step. Hence we calibrated the method based on a
 171 single absorbance reading obtained after acid and then additional I was added. Calibrations and
 172 standard checks revealed this approach did not have any impact on the quality of the data.

173

174 *Ammonium Concentration*

175 NH₄⁺ concentrations were measured in 0.45 μm (Millex) filtered samples with a Seal Analytical
 176 Autoanalyser 3 according to method G-109-93 rev. 10 (Seal Analytical) using sodium salicylate,
 177 dichloro-isocyanuric acid and citrate buffer. The method was calibrated using standards ranging from
 178 0-2 mg/L prepared from dilutions of a 1000 mg/L ammonium standard solution (Merck).

179

180 *Iodide Concentration*

181 I concentrations were determined using a Dionex ICS-2000 ion chromatograph equipped with an
 182 EGC III KOH elugen cartridge, AG18 (2 x 50 mm) guard column, AS18 (2 x 250 mm) analytical
 183 column, ASRS 300 (2 mm) suppressor, DS6 heated conductivity cell and AS40 autosampler.
 184 Samples were diluted 100-fold with 18 MΩ deionised water for analysis and 5 μL was injected onto
 185 the ion chromatograph. Aqueous potassium hydroxide was used as the eluent at a flow rate of 0.25
 186 mL min⁻¹ with a gradient program starting from an initial concentration of 2 mM hydroxide (hold 1
 187 min) to 20 mM at 18 min then to 41 mM at 19 min (hold 2 min) before returning to 2 mM. The I
 188 retention time was 19 min. The instrument was calibrated with matrix-matched standards ranging
 189 from 0-100 nM (I⁻), prepared from dilutions of a 1000 mg/L iodide standard solution (Fisher
 190 Scientific) with 18 MΩ deionised water and containing a final concentration of 1% ESAW.

191

192 **Data Analysis**

193 As in Guerrero and Jones (1996), the NH_4^+ oxidation rate is defined here as the rate of increase in
 194 NO_2^- . Similarly, we define the rate of I oxidation as the rate of increase in IO_3^- . This is appropriate as
 195 no other iodine species were supplied to the cultures and conversion between I and IO_3^- is known to
 196 be the main cause of variability in inorganic iodine speciation (Bluhm *et al.*, 2010; Chance *et al.*,
 197 2014). Average NO_2^- and IO_3^- production rates were calculated for each replicate culture using
 198 Equation 4.

199
$$\text{Production Rate (nM day}^{-1}\text{)} = \frac{(C_{\text{end}} - C_0)}{t} \quad (4)$$

200 where C_0 and C_{end} are the NO_2^- or IO_3^- concentrations observed at the start and end of the experiment
 201 and t is the experimental duration in days. Cell-normalised rates were calculated by dividing these
 202 rates by the final cell density observed in each AOB culture and are hence likely to be minimum
 203 values.

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205

206 **Results**

207 **Cell counts and pH**

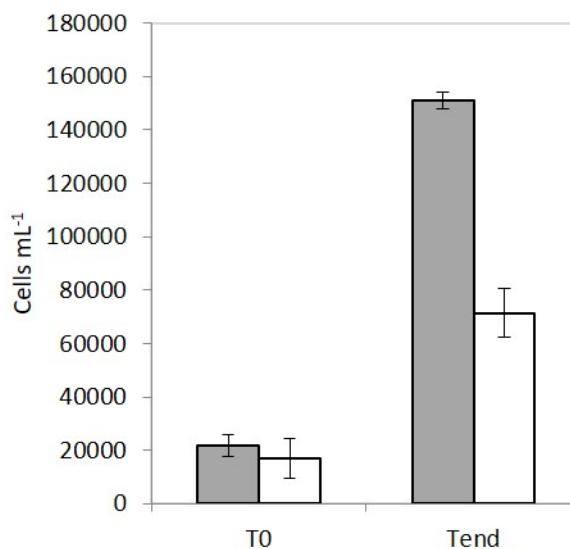
208 Increases in cell density were observed in all replicates of *Nitrosomonas* sp. and *Nitrosococcus*
 209 *oceanii* between the start and end of the experiment indicating growth (Figure 1). Average initial cell
 210 density in the *Nitrosomonas* sp. cultures was 21,767 ($\pm 4,046$) cells mL^{-1} and this increased to 150,983
 211 ($\pm 7,585$) cells mL^{-1} by the end of the experiment (8 days). For *Nitrosococcus oceanii* start and end (12
 212 days) cell densities were 16,947 ($\pm 3,098$) and 71,430 ($\pm 9,062$) cells mL^{-1} , respectively. Average pH
 213 levels in the culture experiments calculated from measurements at each time point (data not shown)
 214 were 7.69 (± 0.07) for *Nitrosomonas* sp. and 7.41 (± 0.12) for *Nitrosococcus* sp. These pH levels are

215 consistent with those found in the media-only controls (7.64 ± 0.07 for *Nitrosomonas* sp; 7.64 ± 0.15
 216 for *Nitrosococcus oceani*).

217

218

219



220

221 **Figure 1.** Average cell number in the *Nitrosomonas* sp. (grey bars) and *Nitrosococcus oceani* (white
 222 bars) cultures used in this study at the start (T₀) and end (T_{end}; 8 days for *Nitrosomonas* sp. and 12
 223 days for *Nitrosococcus oceani*) of each experiment. Error bars are standard deviations from three
 224 replicate cultures.

225

226 *Iodine and nitrogen speciation*

227 Figure 2 shows that significant increases in the concentrations of IO₃⁻ (compared to media-only

228 controls) were observed alongside NO₂⁻ production in both AOB cultures studied. In *Nitrosomonas*

229 sp. (Figure 2ai and 2bi) there was a steady increase in IO₃⁻ concentrations throughout the experiment

230 reaching a maximum of 19,921 (±4,754) nM by the end of the experiment (day 8). In contrast NO₂⁻

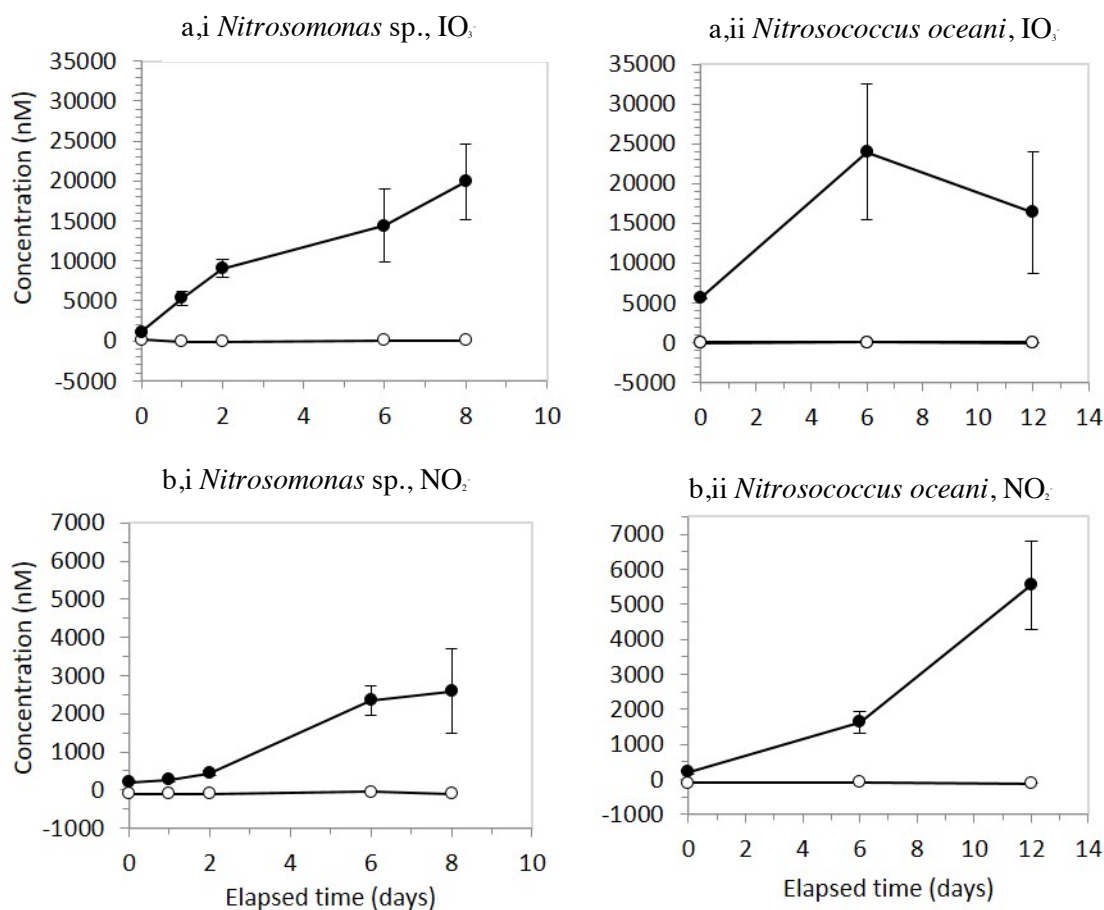
231 concentrations reached a maximum of 2,360 (±386) nM by day 6 and remained at around that level

232 until the end of the experiment. In *Nitrosococcus oceani* (Figure 2aii and 2bii) IO₃⁻ concentrations

233 increased rapidly during the initial stages of the experiment reaching 23,943 (±8,568) nM by day 6.

234 IO₃⁻ concentrations at the end of the experiment (day 12) were 16,365 (±7,603) nM. NO₂⁻

235 concentrations increased gradually throughout the experiment reaching 5,547 ($\pm 1,251$) nM by day
 236 12. There was larger variability in IO_3^- concentrations between replicates for *Nitrosococcus oceani*
 237 but despite this a clear increase in all replicates was observed.



238

239 **Figure 2.** Changes in iodate (a) and nitrite (b) concentrations in cultures (closed symbols) and
 240 media-only controls (open symbols) for two cultures of ammonia-oxidising bacteria: i) *Nitrosomonas*
 241 sp.; and, ii) *Nitrosococcus oceani* supplied with 9-10 mM iodide and 7-8 mM NH_4^+ . Error bars show
 242 the standard deviation of three replicate cultures.

243

244 Average production rates of IO_3^- and NO_2^- are presented in Table 1. In *Nitrosomonas sp.* average

245 rates (\pm standard deviation) were 2,348 (± 593) nM IO_3^- day $^{-1}$ and 298 (± 141) nM NO_2^- day $^{-1}$. In

246 *Nitrosococcus oceani* averages rates were 897 (± 640) nM IO_3^- day $^{-1}$ and 445 (± 99) nM NO_2^- day $^{-1}$.

247 Minimum cell-normalised rates (based on the final cell density observed in each culture) were 15.69

248 (± 4.71) fmol IO_3^- cell $^{-1}$ day $^{-1}$ and 1.96 (± 0.88) fmol NO_2^- cell $^{-1}$ day $^{-1}$ for *Nitrosomonas sp.*, and 11.96

249 (± 6.96) $\text{fmol IO}_3^- \text{ cell}^{-1} \text{ day}^{-1}$ and $6.19 (\pm 0.56) \text{ fmol NO}_2^- \text{ cell}^{-1} \text{ day}^{-1}$ for *Nitrosococcus oceani*. Molar
 250 ratios of iodate-to-nitrite production were 9.2 ± 4.0 for *Nitrosomonas* sp. and 1.88 ± 0.91 for
 251 *Nitrosococcus oceani*.

252

253 **Table 1.** Nitrite and iodate production rates (\pm standard deviations) observed in cultures of the
 254 ammonia-oxidising bacteria *Nitrosomonas* sp. and *Nitrosococcus oceani*. Cell-normalised values are
 255 a minimum as they are calculated using maximum cell densities.

256

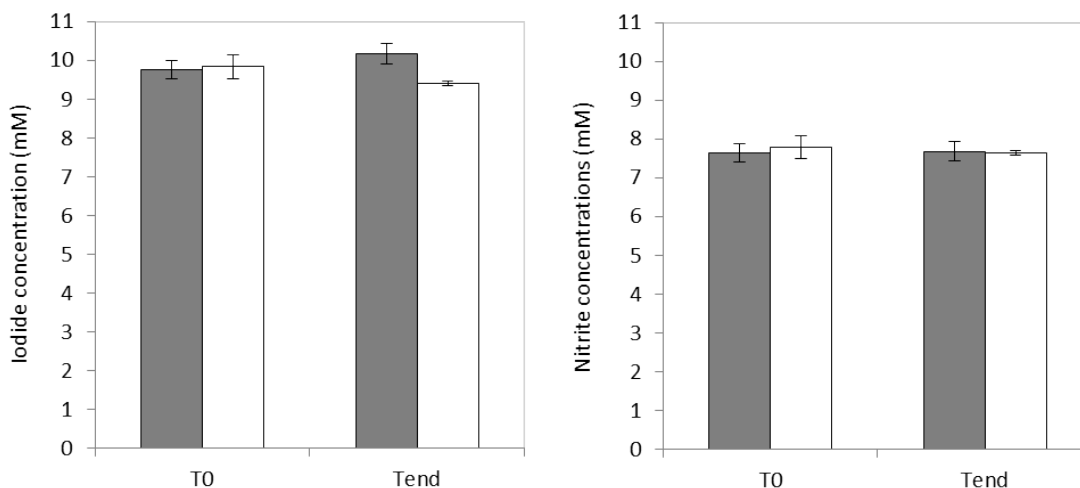
Culture	Nitrite		Iodate	
	nM day ⁻¹	fmol cell ⁻¹ day ⁻¹	nM day ⁻¹	fmol cell ⁻¹ day ⁻¹
<i>Nitrosomonas</i> sp.	298 (± 141)	1.96 (± 0.88)	2,348 (± 593)	15.69 (± 4.71)
<i>Nitrosococcus oceani</i>	445 (± 99)	6.19 (± 0.56)	897 (± 640)	11.96 (± 6.96)

257

258

259 Figure 3 shows that, within error, a decline in I or NH_4^+ concentrations was not observed during
 260 either of the AOB experiments. Average starting I or NH_4^+ concentrations in *Nitrosomonas* sp. were
 261 $9.8 (\pm 0.2) \text{ mM}$ and $7.6 (\pm 0.1) \text{ mM}$ respectively. At the end of the experiment these values were 10.2
 262 $(\pm 0.3) \text{ mM}$ I and $7.7 (\pm 0.1) \text{ mM}$ NH_4^+ . For *Nitrosococcus oceani* the start and end concentrations
 263 were $9.8 (\pm 0.3)$ and $9.4 (\pm 0.1) \text{ mM}$ for I and $7.8 (\pm 0.1)$ and $7.7 (\pm 0.1) \text{ mM}$ for NH_4^+ . This result was
 264 expected as the average standard deviations associated with the observed concentrations of I or NH_4^+
 265 (i.e. 0.1 to 0.3 mM) are at least an order of magnitude higher than the maximum levels of IO_3^- and
 266 NO_2^- observed in the culture experiments, i.e. very little of the initial stock of NO_2^- or NH_4^+ was
 267 oxidised during the experiments.

268



269

270 **Figure 3.** Start and end concentrations of a) iodide and b) ammonia in cultures of *Nitrosomonas* sp.
 271 (grey bars) and *Nitrosococcus oceani* (white bars). Error bars show the standard deviation of three
 272 replicate cultures.
 273

274

275

276

277 **Discussion**

278 ***Iodate production by ammonia-oxidising bacteria***

279 Our results confirm that IO_3^- production occurs in cultures of the ammonia-oxidising bacteria

280 *Nitrosomonas* sp. and *Nitrosococcus oceani* supplied with I, but not in cultures of nitrite oxidising

281 bacteria. Coincident increases in NO_2^- (Figure 2) show that both cultures were actively oxidising

282 ammonia throughout the experiments at rates of $1.96 \pm 0.08 \text{ fmol NO}_2^- \text{ cell}^{-1} \text{ day}^{-1}$ for *Nitrosomonas*

283 sp. and $6.19 \pm 0.56 \text{ fmol NO}_2^- \text{ cell}^{-1} \text{ day}^{-1}$ for *Nitrosococcus oceani*. Whilst these cell-normalised

284 oxidation rates are of the same order as those reported in the literature (e.g. 6-20 $\text{fmol NO}_2^- \text{ cell}^{-1} \text{ day}^{-1}$;

285 Ward *et al.*, 1987; 1989) they are at the lower end. This is consistent with the approach taken here to

286 calculate the rates by normalising to the final (highest) cell densities. It is also worth noting that the

287 cultures were at an early stage of growth and had relatively low cell densities during the experiment.

288 This was done to avoid significant changes in pH in the bulk media which would impact inorganic

289 iodine speciation (Section 3.2). The observation of an increase in IO_3^- concentrations alongside active

290 biological ammonia oxidation supports previous studies (e.g. Truesdale *et al.*, 2001; Zic *et al.*, 2013)
 291 which have shown that high aqueous concentrations of IO_3^- are found in regions of enhanced
 292 nitrification, and provides the first direct confirmation of a biological basis for at least one
 293 mechanism of iodide oxidation

294

295 Whilst we did not set out to establish the mechanism for I^- to IO_3^- oxidation by marine nitrifiers, some
 296 speculations can be made. As I^- oxidation to IO_3^- requires the transfer of six electrons, it may occur in
 297 a series of one- or two- electron transfer steps. Initially, I^- may be oxidised to molecular iodine ($\text{I}^- \rightarrow$
 298 I_2), a reaction which is thermodynamically unfavourable at the pH of seawater (Luther *et al.*, 1995).
 299 Further oxidation to IO_3^- by disproportionation ($\text{I}_2 \rightarrow \text{HOI} \rightarrow \text{IO}_3^-$) can occur spontaneously, but in
 300 seawater is subject to competition with reduction of I_2 by organic matter (Truesdale & Moore, 1992;
 301 Truesdale *et al.*, 1995). It is not known whether the ammonia-oxidisers mediate just the first stage of
 302 I^- oxidation, with the observed IO_3^- production due to subsequent spontaneous reactions in the culture
 303 media, or if they are involved in driving the complete conversion of I^- to IO_3^- . However, bacteria
 304 which just oxidise I^- to I_2 have been isolated from seawater aquaria (Gozlan, 1968), I-rich natural gas
 305 brine waters (Amachi *et al.*, 2005) and marine environmental samples (Fuse *et al.*, 2003; Amachi *et*
 306 *al.*, 2005).

307

308 The observed IO_3^- production is either linked to the nitrification process itself or associated with other
 309 metabolic activities of the AOB studied. Truesdale *et al.* (2001) has proposed that I^- oxidation to IO_3^-
 310 would be energetically advantageous for chemoautotrophic AOB. In that case the key enzymes used
 311 to obtain energy during the oxidation of NH_4^+ to NO_2^- (ammonia monooxygenase [AMO] and
 312 hydroxylamine oxidoreductase [HAO]) could also have the potential to use I^- as a substrate. The
 313 observed IO_3^- -to- NO_2^- molar production rates (9.2 ± 4.0 for *Nitrosomonas* sp. and 2.3 ± 1.1 for
 314 *Nitrosococcus oceani*) are intriguing. If AMO/HAO are involved, this suggests that the enzymes

315 have higher affinities for I⁻ than NH₄⁺/NH₂OH given the similar concentrations of I⁻ and NH₄⁺ used in
 316 the experiments. Other enzymes that have been implicated in I⁻ oxidation include the
 317 chloroperoxidases (Thomas & Hager, 1968) but we do not know if they occur in AOB. The exact
 318 metabolic pathway driving the observed IO₃⁻ production and its controls (i.e. substrate concentrations,
 319 light intensity) will need to be determined in future work. To establish if such further
 320 experimentation is warranted we need to explore whether the link between nitrification and I⁻
 321 oxidation is likely to be an important part of inorganic iodine cycling in seawater.

322

323 ***Implications for inorganic iodine speciation in the oceans***

324 Our culture studies suggest that the molar rate of I⁻ oxidation (IO₃⁻ production) is ~2-9 times higher
 325 than that for ammonia oxidation (nitrification). Note that although ammonium and iodide
 326 concentrations were much higher in the experimental media than in the oceans, the concentration
 327 ratio of these species was comparable to that found naturally. Ammonia oxidation rates in seawater
 328 range from below detection to 10³ nM day⁻¹ (Table 2). Literature estimates of the rate of I⁻ oxidation
 329 in the marine environment range from ~4 to 670 nM year⁻¹ or 0.01 to 1.84 nM day⁻¹ (reviewed in
 330 Chance *et al.*, 2014). If the oxidation molar ratios observed in this study (~2-9) are representative,
 331 predicted rates of I⁻ oxidation are in-line (i.e. 2-9 times higher) with the lower end of observed
 332 ammonia oxidation rates (Table 2).

333

334 **Table 2.** Ammonia-oxidation rates measured in a range of ocean regions.

335

Study	Location	Rate (nM day ⁻¹)
Newell <i>et al.</i> (2011)	Arabian Sea, Indian Ocean	undetected to 21.6
Smith <i>et al.</i> (2015)	Northeast Pacific	< 0.01 to 90
Peng <i>et al.</i> (2015)	Eastern tropical north Pacific	< 1 to 8.6
Newell <i>et al.</i> (2013)	Subtropical Atlantic, Sargasso Sea (BATS)	< 2
Lam <i>et al.</i> (2007)	Black Sea	7-24
Beman <i>et al.</i> (2012)	Gulf of California, eastern tropical north Pacific	0-348

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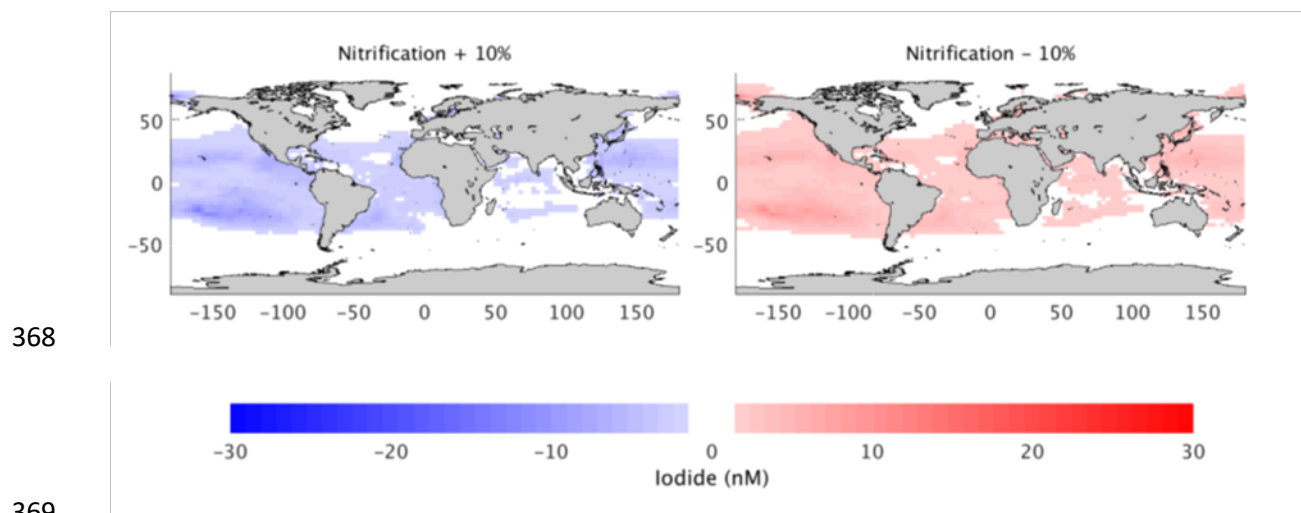
338 Truesdale *et al.* (2001) derive likely I⁻ oxidation (or IO₃⁻ production) rates for the near surface Black
 339 Sea using an iodine budget and this allows us to examine the potential importance of the link
 340 between nitrification and I⁻ oxidation on a local scale. They predict a minimum I⁻ oxidation flux of
 341 $3.89 \times 10^{-4} \text{ mol I m}^{-2} \text{ year}^{-1}$ which is an average of 0.02 nM day^{-1} at a mixed-layer depth (MLD) of 50 m
 342 or 0.11 nM day^{-1} at an MLD of 10 m. Lam *et al.* (2007) report an AOB abundance of $\leq 1,400 \text{ cells}$
 343 mL^{-1} in the Black Sea. If we apply a cell density of $1,400 \text{ AOB cells mL}^{-1}$ to the average cell-
 344 normalised rates of IO₃⁻ production observed in this study (Table 1) we derive I⁻ oxidation rates of
 345 $\sim 20 \text{ nM d}^{-1}$. This is clearly much higher than the rates suggested in Truesdale *et al.* (2001). This
 346 discrepancy could be explained in a number of ways. Firstly, Lam *et al.* (2007) state that net
 347 nitrification only takes place within a narrow depth range of the Black Sea water column (i.e.
 348 between 71 and 81 m) and, the I⁻ oxidation values derived in Truesdale *et al.* (2001) are minimum
 349 values. It is also possible that the AOB studied here have a higher capacity for I⁻ oxidation (per unit
 350 ammonia-oxidised) than other ammonia-oxidisers or that our culture conditions (e.g. substrate
 351 availability) promoted higher I⁻ oxidation rates than would be observed in marine systems. For
 352 example, ammonia-oxidising Archaea (AOA), which can outnumber known bacterial ammonia
 353 oxidisers by orders of magnitudes in environments such as the marine water-column (reviewed by
 354 Schleper & Nicol, 2010), may have a very different capacity for I⁻ oxidation compared to the AOB
 355 studied here. Further studies are needed to establish the relationship between ammonia- and I⁻
 356 oxidation in the marine environment.

357

358 ***Potential implications for future oceanic inorganic iodine distributions***

359 Environmental factors which are known to be currently undergoing change in the oceans (e.g.
 360 oxygen, light, pH, temperature) have all been found to impact rates and patterns of marine
 361 nitrification (reviewed by Pajores and Ramos, 2019). Whilst there remains some uncertainty about
 362 the future magnitude and, in some cases, sign of the response, some of the expected future changes in

363 marine nitrification are large. For example, whilst some studies have seen no impact on specific
 364 marine nitrifiers (e.g. Qin et al., 2014), Beman et al. (2011) suggest that expected rates of
 365 acidification could cause a decline in ammonia oxidation by up to 44% within the next few decades.
 366 It is hence worth exploring how possible future changes in marine nitrification could impact ocean
 367 iodine cycling.



370 **Figure 4.** Modelled changes in surface I concentration (nM) resulting from a) +10%, b) -10%,
 371 changes in the rates of nitrification. Negative percent values indicate a decline in the rate of
 372 nitrification and *vice-versa*. Negative values on the scale bar indicate a decrease in I concentrations
 373 and *vice versa*.
 374

375 In order to explore the possible impact of future changes in marine nitrification rates on sea surface
 376 iodine fields we used the ocean cycling model described in Wadley et al. (2020). Within the model
 377 iodide production is driven by primary productivity, and I oxidation to IO_3^- linked to nitrification in
 378 the mixed layer. Nitrogen fluxes and the spatial distribution of mixed layer ammonia oxidation are
 379 derived from a global biogeochemical cycling model (Yool *et al.*, 2007). I is oxidised to IO_3^- in
 380 association with the ammonia oxidation, with the same I:N:C ratio as associated with iodide
 381 production (Truesdale *et al.*, 2001; Long *et al.*, 2015). The model does not use any of the rates
 382 derived in the current study as these are based on results from only 2 AOB species cultured at high
 383 substrate concentrations. Model outputs (Figure 4) show that even with small (+/- 10%) changes in

384 ammonia oxidation there is a clear alteration to sea surface I fields. Sea surface I concentrations
 385 increase as ammonium oxidation rates decrease and *vice-versa*. For example, the ocean cycling
 386 model suggests there could be an average global increase of 0.13 nM I per 1% decrease in
 387 nitrification. The outputs suggest that the change in the iodine fields is spatially variable and will
 388 increase as the perturbation to ammonia oxidation increases. For example, at the 44% decline in
 389 nitrification predicted by Beman et al. (2011) the model predicts there will be a 25% increase (+30
 390 nM) in sea surface I in the sub-tropical gyres. Carpenter *et al.* (2013) show that I₂ emissions due to
 391 ozone deposition increase near linearly with I concentration. Hence, the predicted changes to sea
 392 surface I fields under future ocean acidification could have a major impact on ozone deposition to
 393 the sea surface, atmospheric chemistry and resulting sea-air iodine emissions.

394

395

396 **5.3.Conclusions**

397 This study has shown that I oxidation to IO₃ occurs in cultures of ammonia oxidising (nitrifying)
 398 bacteria, but not nitrite oxidising bacteria. Our calculations suggest that I oxidation by AOB could be
 399 an important control on inorganic iodine speciation in seawater, but to confirm this further study is
 400 needed on a wider range of ammonia-oxidisers including ammonia oxidising archaea (AOA).

401 Simulations from our iodine cycling model suggest that changes in nitrification rate, such as those
 402 predicted to occur under acidification (Beman *et al.*, 2011), could have an important impact on sea
 403 surface I fields. A future change in marine nitrification could alter sea surface I fields. In turn, this
 404 could lead to a change in ozone deposition to the sea surface and sea-air iodine emissions with
 405 potentially major implications for atmospheric chemistry and air quality.

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409

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413

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426

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433

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