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# Influence of the Sea Surface Microlayer on Oceanic Iodine Emissions

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**ABSTRACT:** The influence of organic compounds on iodine  $(I_2)$  emissions from the  $O_3 + I^-$  reaction at the sea surface was investigated in laboratory and modeling studies using artificial solutions, natural subsurface seawater (SSW), and, for the first time, samples of the surface microlayer (SML). Gas-phase  $I_2$ was measured directly above the surface of liquid samples using broadband cavity enhanced absorption spectroscopy.  $I_2$  emissions were consistently lower for artificial seawater (AS) than buffered potassium iodide (KI) solutions. Natural seawater samples showed the strongest reduction of  $I_2$ emissions compared to artificial solutions with equivalent  $[I^-]$ , and the reduction was more pronounced over SML than SSW. Emissions of volatile organic iodine (VOI) were highest from SML samples but remained a negligible fraction (<1%) of the total iodine flux. Therefore, reduced iodine emissions from natural seawater cannot be explained by chemical losses of  $I_2$ 



or hypoiodous acid (HOI), leading to VOI. An interfacial model explains this reduction by increased solubility of the  $I_2$  product in the organic-rich interfacial layer of seawater. Our results highlight the importance of using environmentally representative concentrations in studies of the  $O_3 + I^-$  reaction and demonstrate the influence the SML exerts on emissions of iodine and potentially other volatile species.

# INTRODUCTION

Tropospheric iodine is attracting increasing research interest as insights are gained into its large influence on local and global tropospheric and stratospheric chemistry.<sup>1–8</sup> Reactive iodine species, such as IO radicals, induce cycles of catalytic ozone destruction,<sup>5,9,10</sup> change the oxidative capacity of the troposphere through their perturbation of the HOx and NOx cycles,<sup>1,2,11,12</sup> and are linked to particle nucleation.<sup>13,14</sup> Tropospheric iodine levels have tripled since the mid-20th century in certain regions,<sup>15,16</sup> thus a robust understanding of iodine sources into the atmosphere is crucial.

The main source of atmospheric iodine is oceanic emissions.<sup>22</sup> Although biogenic sources contribute to iodine emissions in coastal areas,<sup>17,18</sup> around 80% of atmospheric iodine is believed to arise from abiotic sea-air emissions of inorganic iodine in the form of molecular iodine (I<sub>2</sub>) and hypoiodous acid (HOI).<sup>2,19,20</sup> These emissions result from the reaction of ozone with iodide (I<sup>-</sup>), which, along with iodate (IO<sub>3</sub><sup>-</sup>),<sup>21,22</sup> comprise the main form of oceanic iodine at the sea surface (RR1–RR6)<sup>19,23,24</sup>

$$I_{(aq)}^{-} + O_{3(g \text{ or } aq)} \to IOOO^{-}$$
(R1)

$$IOOO^- \to IO^-_{(aq)} + O_2 \tag{R2}$$

$$IO_{(aq)}^{-} + H^{+} \rightarrow HOI_{(aq)}$$
 (R3)

$$HOI_{(aq)} + I^{-} + H^{+} \rightleftharpoons I_{2(aq)} + H_2O$$
(R4)

$$I_{2(aq)} \rightleftharpoons I_{2(g)} \tag{R5}$$

$$\text{HOI}_{(aq)} \rightleftharpoons \text{HOI}_{(q)}$$
 (R6)

On average, the global sea surface iodide concentration (upper 20 m) is estimated at  $9.5 \times 10^{-8}$  M.<sup>25</sup> Typical iodine sea-air fluxes calculated for the clean marine boundary layer lie in the range of 100–250 nmol m<sup>-2</sup> d<sup>-1</sup> for HOI and 2–10 nmol m<sup>-2</sup> d<sup>-1</sup> for I<sub>2</sub>.<sup>19,26</sup>

Iodide reacts very rapidly with ozone (RR1), much faster than the equivalent reactions of Cl<sup>-</sup> and Br<sup>-</sup> ( $k_{\rm I-} = 2 \times 10^9$  M<sup>-1</sup> s<sup>-1</sup>,  $k_{\rm Br-} = 1-2 \times 10^3$  M<sup>-1</sup> s<sup>-1</sup>,  $k_{\rm Cl-} \sim 3 \times 10^{-3}$  M<sup>-1</sup> s<sup>-1</sup>), explaining the major influence iodide has on the dry deposition of ozone, despite its much smaller concentration in seawater ([Cl<sup>-</sup>] =  $5.6 \times 10^{-1}$  M; [Br<sup>-</sup>] =  $8.6 \times 10^{-4}$  M).<sup>27-30</sup> The fast reactivity of iodide with O<sub>3</sub> and its enhancement at the air– water interface<sup>31,32</sup> suggests that heterogeneous surface reactions would be promoted. However, at low iodide

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The influence of organic compounds on the ozone + iodide reaction remains unclear. Due to the presence of other ions and virtually unknown quantities of various dissolved organics, the sea surface is a chemically complex, but dilute, system. The surface microlayer (SML), the uppermost 1 to 1000  $\mu$ m of the sea surface, represents a less dilute environment where surface-active organics can become significantly enriched.<sup>34,35</sup> By its nature, the SML constitutes the interface between the air and water, and its influence on air-sea exchange has been demonstrated for trace gases, e.g., CO<sub>2</sub><sup>36</sup> and N<sub>2</sub>O.<sup>37</sup> However, the underlying mechanisms remain largely unknown and it is not clear to what extent the presence of natural surfactants modifies oceanic gaseous emissions.<sup>38,39</sup>

Organics influence iodine emissions in different ways via several mechanisms, as summarized in Table S1. A suppression of I<sub>2</sub> emissions was observed in the presence of a monolayer of octanol,40 whereas short-chain carboxylic and fulvic acids enhanced I<sub>2</sub> emissions.<sup>41</sup> Chemical competition for O<sub>3</sub> by phenolate ions at the surface also suppresses I<sub>2</sub> emissions. The addition of a complex organic matrix, dissolved organic carbon (DOC) extracted from natural seawater, to buffered solutions of iodide has been found to lead to a strong reduction of  $I_{2(g)}$  emissions.<sup>43</sup> This reduction could not be explained by the reactivity of DOC toward  $O_3$  and  $I_2$ /HOI, and instead a decrease in the net transfer rate of I2 from the aqueous to gas phase was suggested,<sup>43</sup> as previously observed for octanol.<sup>40</sup> Conversely, ozonolysis of coastal seawater samples can generate certain halocarbons (CH<sub>2</sub>I<sub>2</sub>, CHI<sub>3</sub>, and  $CHClI_2$ ),<sup>44</sup> implying that reactions of the I<sub>2</sub> (or HOI) product in solution can yield organic iodine species. However, no direct link has been demonstrated between the emission of halocarbons and the presence of dissolved organics or reduced emission of inorganic iodine. Overall, these studies show that introducing a single organic component can alter iodine emissions through chemical enhancement, suppression, and/or physical hindrance.

Here, we compare iodine  $(I_{2(g)})$  emissions from the dark reaction of ozone with iodide in buffered potassium iodide solutions and artificial seawater (AS), against natural subsurface seawater (SSW) and, for the first time, SML samples. Importantly, these experiments were performed for ozone mixing ratios (20–150 ppbv) and iodide concentrations (1  $\times$  $10^{-7}$  to  $1.6 \times 10^{-5}$  M), which include ambient conditions. The dependence on ozone and iodide concentrations is investigated and the influence of organic materials is discussed. In separate experiments, we explore halocarbon production from the ozone + iodide reaction, comparing halocarbon emissions from artificial seawater, SSW and SML samples, as functions of ozone and iodide concentrations. The implications of these first I<sub>2</sub> and organic emission measurements using natural SML samples are explored using an adaptation of the aqueous interfacial layer model of Carpenter et al.<sup>19</sup>

## MATERIALS AND METHODS

**Chemicals.** Buffered solutions of iodide were prepared by adding concentrated KI stock solutions to a phosphate buffer at pH 8. Artificial seawater (AS) solutions were made by

dissolving KCl and KBr in a phosphate buffer and then adding aliquots of the KI stock solutions. Full details are in the Supporting Information (SI).

**Sampling and Analysis of Seawater.** The samples of natural seawater were obtained from the North Sea, 5 km off-shore from Bridlington (U.K.), and filtered through GF/F ashed quartz filters. Iodide in the samples was measured using cathodic stripping voltammetry. DOC was determined using a total organic carbon analyzer. Surface tension was measured using the DuNoüy ring method. Details about the sampling locations, dates, procedures, and methods can be found in SI, Sections S1.2–S1.5 and Table S2.

In Situ I<sub>2</sub> Measurements. Figure S1 shows the apparatus for in situ measurements of molecular iodine by broadband cavity enhanced absorption spectroscopy (BBCEAS). A 250 cm<sup>3</sup> min<sup>-1</sup> flow of synthetic air (BTCA-178, BOC special gases) passed through an ozone generator and mixed with 3500 cm<sup>3</sup> min<sup>-1</sup> of synthetic air before entering the custombuilt glass reaction vessel. The vessel contained 500 cm<sup>3</sup> of aqueous solution with a surface area of approx. 380 cm<sup>2</sup>, leaving a headspace of 4800 cm<sup>3</sup> where the BBCEAS light beam was integrated. Iodine concentrations were measured approximately 3.5 cm above the liquid's surface. Two additional air flows, 200 cm<sup>3</sup> min<sup>-1</sup> in total, were used to purge the cavity mirrors; thus, the total flow through the vessel was 3950 cm<sup>3</sup> min<sup>-1</sup>. The vessel was thermostatted and covered in aluminum foil to avoid photolytic losses of I<sub>2</sub> (or the production of IO radicals). The solution was actively stirred at the same rate for all experiments by means of a central magnetic stirrer, and solutions were brought to temperature before introduction into the vessel. Experiments recorded I<sub>2</sub> emissions versus increasing iodide concentrations by adding aliquots ( $\sim 1 \text{ cm}^3$ ) of concentrated potassium iodide solutions  $(1 \times 10^{-4} \text{ or } 1 \times 10^{-3} \text{ M})$  to the sample solutions through a lid at the top of the vessel.

Retrieving I<sub>2</sub> concentrations followed a similar procedure to previous BBCEAS measurements.<sup>45</sup> Further details appear in the SI. The errors reported include the statistical uncertainty of the spectral fit, dominant at small concentrations, and the systematic errors of the measurement (typically totaling 16% for I<sub>2</sub>, see the SI). The limit of detection for iodine (LoD) was 4 pptv (1  $\sigma$  in 60 s), which corresponds to a minimum detectable I<sub>2</sub> flux of 1.5 × 10<sup>7</sup> molecules cm<sup>-2</sup> s<sup>-1</sup>. All iodine data was corrected for losses in the reaction vessel due to the gas-phase reaction of I<sub>2</sub> + O<sub>3</sub> using the rate constant  $k_{(I_2 + O_3)} = 2.25 \times 10^3 \text{ M}^{-1} \text{ s}^{-1}$ .<sup>46</sup> These losses proved to be negligible (<1% of the I<sub>2</sub> emissions) for the low reactant concentrations of our experiments and the relatively short residence time of gas inside the reactor (73 s).

Ozone, measured using commercial UV absorbance ozone monitors, was monitored upstream and downstream of the vessel by switching a three-way valve.  $O_3$  measurements are detailed in the SI. All results presented here were obtained for solutions and seawater samples at 17 °C.

Halocarbon Measurements. The production of halocarbons was examined using the setup depicted in Figure S2. The system was designed to flow 500 cm<sup>3</sup> min<sup>-1</sup> of dry hydrocarbon-free air through a mass flow controller, with or without ozone, into the reaction vessel (500 cm<sup>3</sup> round-bottom glass flask). There, it passed over 250 cm<sup>3</sup> of degassed artificial seawater or natural samples (surface area of 105.7 cm<sup>2</sup>) before sampling. The entire reaction vessel and tubing



**Figure 1.** Panel A: BBCEAS measurements of  $I_2$  emissions as a function of  $[I^-]$  over buffered KI solutions (black points) for ozone concentrations of 13.6 ppbv (squares), 22.7 ppbv (circles), and 66.6 ppbv (triangles) at 17 °C. The red symbols are the  $I_2$  emissions over artificial seawater for  $[O_3] = 18$  ppbv (squares), 34.7 ppbv (circles), and 126.7 ppbv (triangles). Panel B: BBCEAS measurements of  $I_2$  emitted from the natural samples of subsurface seawater (blue, SSW), surface microlayer (green, SML), and a mixture of 20% SML + 80% SSW (gold) for ozone concentrations of 38.6, 38.5, and 36 ppbv, respectively (circles). The triangular symbols show  $I_2$  recorded at higher ozone concentrations over SSW ( $[O_3] = 128.1$  ppbv) and SML ( $[O_3] = 134.6$  ppbv). All measurements at 17 °C. The dotted lines are the straight segments between the points, meant to guide the eye. The error bars reflect the overall uncertainty on the measurements, including the uncertainty on the spectral fit, averaging and systematic errors.

were thermostatted and covered in aluminum foil to prevent halocarbon losses due to wall losses and photolysis. During all experiments, the solution was gently stirred using a magnetic stirrer to avoid depletion at the surface and to mimic the dynamics at the sea surface. The halocarbon products were trapped using an air server coupled to a thermal desorption unit (CIA-8, Unity-2, Markes, U.K.) and then analyzed using gas chromatography coupled to a mass spectrometer (GC– MS, Agilent 6890, 5975C). Further details are in the SI.

**Modeled Iodine Emissions: The Sea Surface Model.** The interfacial model described in Carpenter et al.<sup>19</sup> was used, with some modifications, to estimate I<sub>2</sub> (and HOI) emissions from this study's experiments. Full details can be found in the SI. Briefly, we assumed that the ozone uptake coefficient  $\gamma_{I}^{-}$  is controlled by the aqueous-phase O<sub>3</sub> + I<sup>-</sup> reaction and is equivalent to  $\gamma_{ag,I}$ , with

$$\frac{1}{\gamma_{\rm aq}^{\rm I}} = \frac{1}{\alpha_{\rm aq}^{\rm I}} + \frac{1}{\Gamma_{\rm aq}^{\rm I}}$$
(1)

where  $\alpha_{aq}^{I-}$  is the mass accommodation coefficient and  $\Gamma_{aq}^{I-}$  is the conductance of the aqueous-phase reaction, given by

$$\Gamma_{\mathrm{aq}}^{\mathrm{I}^{-}} = \frac{4s\sqrt{k^{\mathrm{I}^{-}} \times a_{\mathrm{I}^{-}} \times D_{\mathrm{aq}}}}{\omega}$$
(2)

In eq 2, *s* is the ozone solubility in nondimensional units (aqueous molarity/gas molarity),  $k^{I-}$  is the rate constant for the aqueous-phase reaction  $O_3 + I^-$ ,  $a_{I^-}$  is the activity of iodide, and  $D_{aq}$  is the diffusion coefficient of aqueous ozone. The values of *s* and  $D_{aq}$  were calculated according to the salt content of the water.

At the higher iodide conditions of our experiments ( $[I^-] > \sim 1 \times 10^{-5}$  M), surface reactions may add an appreciable extra component to O<sub>3</sub> uptake (e.g., refs 33, 50). In the SI, we describe a sensitivity study where we included total surface and bulk phase O<sub>3</sub> uptake in the model,<sup>50</sup> without any changes to the iodine emissions scheme. Noting that the model is

designed to simulate environmental conditions where the aqueous reaction dominates ( $[I^-] < 1 \times 10^{-5}$  M), and that those are the experimental conditions used here, we did not include surface reactivity for the remainder of this work.

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Rapid production of  $I_{2(aq)}$  follows the reaction of iodide at the aqueous surface with  $O_3$  deposited from the gas phase (RR1–RR4). The aqueous iodine reaction scheme used here was the same as in Carpenter et al.<sup>19</sup> except for a modification to reflect that  $I_{2(g)}$  emissions observed from artificial seawater (AS) were only around 50% of those from buffered potassium iodide solutions. The reasons for this are unknown, but a potential explanation could be a competing oxidation of HOI to iodate (IO3-) by HOCl/OCl- or HOBr/OBr,51 formed through heterogeneous reactions of Cl<sup>-</sup> and Br<sup>-</sup> with O<sub>3</sub>.<sup>52,53</sup> It is beyond the scope of this study to attempt to model such chemistry explicitly. Rather, we included the reactions HOI +  $HOCl/OCl^{-} \rightarrow IO_{3}^{-}$  as a proxy for the reduction of iodine emissions observed in the presence of Cl<sup>-</sup> or Br<sup>-</sup>. An assumed total of 2 mM of HOCl/OCl- (for seawater concentrations of  $Cl^{-}$  and  $Br^{-}$ ; 54% of deprotonated HOCl at pH = 8) was sufficient to dampen modeled  $I_2$  emissions by ~50% in artificial seawater compared to equivalent conditions over KI solutions. We included this HOCl/OCl- reaction in all simulations of natural or artificial seawater.

Concentrations of  $[I^-]$ ,  $[H^+]$ , and  $[OH^-]$  were fixed for each model run. For modeling iodine emissions from SSW and SML, we included (as in ref 19) pseudo-first-order rate constants for "O<sub>3</sub> + DOC" interfacial reactions of 100 s<sup>-1</sup> and for "I<sub>2</sub>/HOI + DOC" of  $7 \times 10^{-3}$  s<sup>-1</sup>.<sup>6,54</sup> We also utilized the latter reaction to explore the potential for volatile organic iodine production.

## EXPERIMENTAL RESULTS AND DISCUSSION

**Molecular lodine**  $(I_2)$  **Emissions.** The influence of organics in solution on gaseous inorganic iodine emissions was investigated using BBCEAS to monitor  $I_2$  emitted from the ozonolysis of buffered solutions of KI, artificial seawater (AS),

natural subsurface seawater (SSW), and sea surface microlayer (SML) samples.

Artificial Solutions. Figure 1, panel A, shows that iodine emissions were readily detected from KI solutions, even for low ozone and the lowest iodide concentrations tested  $(3 \times 10^{-7} \text{ M})$ . Increasing iodide concentrations led to higher concentrations of gas-phase I<sub>2</sub> under all experimental conditions. Emissions increased almost linearly with increasing ozone and increasing [I<sup>-</sup>], although some roll-off in linearity was observed for the highest iodide concentrations ( $\geq 8 \times 10^{-6} \text{ M}$ ).

I<sub>2</sub> production over artificial seawater (AS, Figure 1A) shows a very similar trend, but with overall lower I<sub>2</sub> flux rates than for KI solutions. Over AS, the lowest iodide concentration  $[I^-] =$  $1.5 \times 10^{-7}$  M and lowest ozone concentration (17 ppbv) did not produce I<sub>2</sub> emissions above the detection limit of the BBCEAS system. For all other ozone concentrations (35–127 ppbv) and iodide concentrations, I<sub>2</sub> was detected above the LoD, showing a generally linear increase with  $[O_3]$ .

The fluxes obtained in our experiments with KI solutions correspond well to previous observations. Carpenter et al.<sup>19</sup> reported a flux I<sub>2 emitted</sub> =  $4 \times 10^{10}$  molecules cm<sup>-2</sup> s<sup>-1</sup> for a buffered iodide solution with [I<sup>-</sup>] =  $1.5 \times 10^{-5}$  M and [O<sub>3</sub>] = 35 ppbv at 20°C. Under similar conditions, [I<sup>-</sup>] =  $1.6 \times 10^{-5}$  M and [O<sub>3</sub>] = 37.2 ppbv at 17 °C, the flux observed in this study is slightly lower, I<sub>2 emitted</sub> =  $3.3 \times 10^{10}$  molecules cm<sup>-2</sup> s<sup>-1</sup>. The observed I<sub>2</sub> fluxes also agree well with observations reported in MacDonald et al., from solutions without chloride.<sup>55</sup> The flux observed here, for [I<sup>-</sup>] =  $1.2 \times 10^{-6}$  M and [O<sub>3</sub>] = 66.6 ppbv, is I<sub>2 emitted</sub> =  $4.6 \times 10^9$  molecules cm<sup>-2</sup> s<sup>-1</sup>. This compares well with their I<sub>2 emitted</sub> =  $4.9 \times 10^9$  molecules cm<sup>-2</sup> s<sup>-1</sup> for a buffered solution of [I<sup>-</sup>] =  $1 \times 10^{-6}$  M and [O<sub>3</sub>] = 78 ppbv.<sup>55</sup>

When comparing I2 fluxes over artificial seawater, our observations are about 3 times smaller than in MacDonald et al.<sup>55</sup> using a similar chloride concentration (0.5 M) for  $[I^-] = 1$  $\times$  10<sup>-6</sup> M. MacDonald et al.<sup>55</sup> report an I<sub>2</sub> flux of 12  $\times$  10<sup>9</sup> molecules  $cm^{-2} s^{-1}$ , whereas the flux calculated by extrapolating our AS data to the same ozone concentration (222 ppbv) is around  $3.9 \times 10^9$  molecules cm<sup>-2</sup> s<sup>-1</sup>. However, there are some important differences that may explain this discrepancy. First, MacDonald et al. did not stir the liquid phase for their experiments, a condition known to lead to higher emissions due to reduced downmixing of products formed, making unstirred conditions less representative of the turbulent surface layer of the ocean.<sup>19</sup> Second, the AS used here contains bromide, whereas the MacDonald study used only chloride. Although the reaction of bromide with ozone is slow as stated in the introduction, bromide reacts quickly with HOI to form BrI (k =  $4.1 \times 10^{12} \text{ M}^{-1} \text{ s}^{-1}$ ), which could further contribute to the lower emissions observed here.<sup>56</sup>

The I<sub>2</sub> emissions observed over buffered KI solutions showed a near linear increase with increasing ozone concentrations (Figure 2, black squares). Although nonlinear behavior of I<sub>2</sub> emissions as a function of ozone has been observed under high iodide ( $5 \times 10^{-3}$  M) and high ozone conditions (over ~100 ppmv), linear behavior in ozone is expected for our iodide and/or ozone concentrations or lower.<sup>19,24</sup> At conditions representative for the open ocean's surface (low ozone, low iodide), I<sub>2</sub> emissions can thus be expected to scale linearly with [O<sub>3</sub>] as also predicted by the interfacial model (see section: Interfacial Model Results and Discussion). Figure 2 shows that lower I<sub>2</sub> emissions were observed when using artificial seawater solutions compared to



Figure 2. Measured I<sub>2</sub> emissions at 17 °C as a function of ozone concentration for total  $[I^-] = 1.2 \times 10^{-6}$  M (filled symbols) and total  $[I^-] = 7.9 \times 10^{-7}$  M (open symbols) over buffered KI solutions (black squares), artificial seawater (red circles), subsurface seawater (blue triangles), and surface microlayer sample (green diamonds). The I2 measurements from SSW and SML used samples collected on 04/05/ 2018, except measurements at  $[O_3] = 38.6$  ppbv, which were done with the samples collected on 15/08/2018, as indicated by the star symbol on the graph (see Table S2 for more details); variability in the organic content of the natural SSW and SML samples might explain why the I2 emissions recorded at 38.6 ppbv O3 lie below the trend of the data points at other O3 concentrations. The dotted lines are straight segments that join the data points, meant to guide the eye. The heavy dashed black lines through the buffered KI data points are linear regressions (y = 8.36 + 0.69x and  $R^2 = 0.94$  for  $[I^-] = 1.2 \times 10^{-6}$ M; y = 8.09 + 0.68x and  $R^2 = 0.93$  for  $[I^-] = 7.9 \times 10^{-7}$  M). The error bars reflect the overall uncertainty on the measurements, including the uncertainty on the spectral fit, averaging and systematic errors.

buffered KI solutions at all ozone concentrations. Since the only change between the experiments with the buffered KI solutions and artificial seawater is the addition of potassium chloride and bromide, this change in salinity seems to provoke the observed change in emissions.

Several factors could contribute in explaining the reduction in I<sub>2</sub> emissions from AS compared to buffered KI solutions. Magi et al.<sup>27</sup> estimated that  $O_3$  diffusivity decreases by 12% in a 3 M sodium iodide solution compared to pure water, but the resulting effect on the uptake of ozone (<6%) is negligible compared to the 3× differences we observed between KI and AS at our much lower salt concentrations. Based on these results and because diffusivity is difficult to predict, changes in diffusivity are generally ruled out as an important factor in the uptake of ozone.33 However, it is well documented that increased salinity almost always decreases the solubility of gases through the so-called "salting-out effect".<sup>57</sup> We calculated the solubility of ozone under our experimental conditions at 17 °C for KI and AS solutions following the approach in Moreno et al.<sup>33</sup> For the highest iodide concentration used here  $([I^-] = 1.6 \times 10^{-5} \text{ M})$ , the calculated solubility of ozone is  $1.15 \times 10^{-7}$  M atm<sup>-1</sup> in a solution of KI and  $1.01 \times 10^{-7}$  M  $atm^{-1}$  in AS, representing a decrease of 12%. This higher solubility of ozone in a solution of KI compared to AS alone cannot explain our observed differences in I<sub>2</sub> emissions. Additional reactions of ozone with Br<sup>-</sup> and Cl<sup>-</sup> could become important at ozone concentrations substantially above what

our study used, although the interfacial model does not predict this nonlinear behavior for  $I_2$  emissions. Other reasons for the reduced  $I_2$  emissions from AS compared with buffered KI solutions will be explored in detail in the model result section.

Natural Seawater Samples. Substantial reductions in La emissions were observed over subsurface seawater and surface microlayer samples compared to AS and KI solutions. This confirms the reduction of iodine emissions in the presence of organics reported in previous studies of the reaction of ozone with iodide<sup>19,40,43</sup> and shows, for the first time, a further reduction over SML samples. Figure 2 compares emissions observed over all four types of solution as a function of  $[I^-]$ . Two different samples were used for these experiments, as indicated by the star symbol in Figure 2 (further details in Table S2). For all ozone concentrations (20–145 ppbv),  $I_2$  was below the BBCEAS detection limit over SSW or SML (containing natural  $[I^-]$  of 1.04 to  $1.53 \times 10^{-7}$  M, see Table S2). However, I2 was detected from SSW and SML after the addition of relatively small amounts of iodide  $([I^-]_{total} \ge 2.98)$  $\times 10^{-7}$  M, i.e., approximately double the naturally occurring  $[I^-]$ ), even at 38 ppbv ozone (typical of mid-ocean ambient  $O_3$ concentrations).

Figures 2 and 1B both clearly show that I<sub>2</sub> emissions over SML samples are lower than those over SSW samples by an average of  $65 \pm 4\%$  (and by up to a maximum of 83%), and the reduction is similar for both sampling dates (Figure S3). A further experiment at  $[O_3] = 36$  ppbv with a mixture of SML/ SSW (20/80 by volume; gold symbols in Figure 1B) showed I<sub>2</sub> emission intermediate between the "pure" SSW and SML results. Interestingly, the I<sub>2</sub> fluxes from this mixed sample were 38% lower (averaged over all [I<sup>-</sup>] data) than the emissions expected from a simple 20:80 weighted average of the emissions from pure SML and pure SSW, which could indicate that organics from the minor SML component preferentially partitioned to the air-liquid interface where I<sub>2</sub> emissions are more efficiently suppressed. As discussed later, we attribute the substantially decreased emissions from the SML compared to the SSW to the enrichment of organics in the SML. However, I<sub>2</sub> emissions from different sets of the SML/SSW samples did not necessarily show the expected relationships with the presence of organics. For example, the SML and SSW samples from 15/08/18 (Table S2) had lower [DOC] and higher surface tension than the SML sample from 04/05/18, yet showed approximately 63% lower emission over SML compared to SSW collected on the same day. Detailed chemical analysis of a large number of SML and SSW samples (preferably collected from different geographical locations), which is beyond the scope of this present study, would be required to identify groups or individual compounds most involved in this reduction.

Similar to the artificial solutions, increasing ozone concentrations over natural seawater samples led to higher  $I_2$  emissions (Figure 2) in a generally linear trend. The emissions from SSW with  $[O_3] = 38.6$  ppbv seem to be lower than the general trend observed across the other ozone concentrations, but this sample was collected on a different date (15/08/18) than the samples used to determine  $I_2$  emissions from other ozone concentrations (04/05/18), which might explain the difference observed. More observations over natural samples are needed to further disentangle the relation between particular types of DOC, surface tension, and their effects to reduce  $I_2$  emissions.

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Halocarbon Emissions over Artificial and Natural Seawater. A separate set of experiments monitored emissions of halocarbons from the reaction of ozone at the surface of artificial solutions (buffered KI solution and AS) and natural samples (SSW, SML). Although no organic material was added, some production of halocarbons was observed upon the ozonolysis of artificial solutions, despite having purged the solution with N<sub>2</sub>. Without ozone, the emissions were close to or below the LoD, and therefore these zero ozone experiments were used as blanks. For the natural seawater samples, halocarbon emissions without ozone were mostly below the LoD, and where they were above, they were an order of magnitude smaller than with ozone.

The most abundant halocarbon produced from exposing natural and artificial samples to ozone was methyl iodide (CH<sub>3</sub>I); this was the only volatile organic iodine (VOI) compound that was consistently emitted. Other halocarbons observed above their LoDs were CH2ClI, C2H5I, 1-C3H7I, 2-C<sub>3</sub>H<sub>7</sub>I, CHBr<sub>2</sub>Cl, and CH<sub>2</sub>Br<sub>2</sub>, although the latter two were only observed for the highest ozone concentrations (1 ppmv) over natural samples. The summed total of these compounds represents less than 10% of the total VOI flux; the other >90% is CH<sub>3</sub>I. The 11 different volatile organic iodine compounds monitored (see Section S1.8 and Table S3 in the SI) were not all emitted from all 4 types of solutions, and different compounds showed different trends for KI, AS, SSW, and SML. But overall, the highest VOI emissions were seen when ozone reacted with the SML samples and increased with increasing ozone. CH<sub>2</sub>I<sub>2</sub> and CH<sub>2</sub>BrI were not observed above their LoDs and, due to a high background, CHBr3 was not significantly observed either.

Due to the complexity of the product distribution and the small flux for each compound individually, we focus only on the summed total of the VOI. Figure 3 shows the VOI emissions measured over the four different types of solutions for two ozone concentrations, both substantially above



**Figure 3.** Measured total volatile organic iodine emissions as a function of iodide concentration over buffered KI solution (black squares), artificial seawater (red circles), subsurface seawater (blue triangles), and surface microlayer (green diamonds) samples at 20 °C for (A)  $[O_3]$ = 400 ppbv (filled symbols) and (B)  $[O_3]$ = 1 ppmv. The dotted lines are straight segments between the points, meant to guide the eye. The error bars reflect the uncertainty on the quantification of the halocarbons.

ambient  $[O_3]$ , but at iodide concentrations relevant for ambient seawater. The VOI emissions show no clear trend with increasing iodide concentrations. The emissions after exposure to 400 ppbv of ozone (Figure 3A) are rather similar for the different solutions. However, VOI emissions are clearly higher for the 1 ppmv ozone experiments over the natural samples, particularly SML (Figure 3B). A maximum flux of VOI <sub>emitted</sub> =  $5.7 \times 10^7$  molecules cm<sup>-2</sup> s<sup>-1</sup> was measured over the SML sample exposed to 1 ppmv of ozone with  $[I^-] = 1.2 \times$  $10^{-7}$  M (i.e., without adding further iodide). Nevertheless, this peak VOI flux is still small compared to the inorganic I<sub>2</sub> fluxes reported in the previous sections, where much lower ozone concentrations were used. For example, a comparable flux of iodine of I <sub>emitted</sub> =  $3.0 \times 10^7$  atoms cm<sup>-2</sup> s<sup>-1</sup> (due to I<sub>2 emitted</sub> =  $1.5 \times 10^7$  molecules cm<sup>-2</sup> s<sup>-1</sup>) was observed over a surface microlayer sample with  $[I^-] = 4.5 \times 10^{-7}$  M exposed to only 38.5 ppbv of ozone (Figure 1). Clearly, the VOI flux will represent only a small fraction of the total iodine flux at environmentally relevant ozone concentrations. Using the interfacial model to estimate emission fluxes at  $[O_3] = 400$ ppbv over SML with  $[I^-] = 1$  to  $4.3 \times 10^{-7}$  M, we calculate  $I_{2 \text{ emitted}} = 50 \text{ to } 193 \times 10^7 \text{ molecules cm}^{-2} \text{ s}^{-1} \text{ and HOI}_{\text{emitted}} = 2.2 \text{ to } 8.3 \times 10^9 \text{ molecules cm}^{-2} \text{ s}^{-1}$ . Correspondingly, the observed VOI fluxes from SML for 400 ppbv  $O_3$  (1.5 to 2.6 ×  $10^7$  atoms cm<sup>-2</sup> s<sup>-1</sup> in Figure 3A) represent between 2.5 and 0.4% of the emissions of iodine atoms from  $I_2$  and VOI and only 0.1 to 0.8% of the total iodine flux (VOI +  $2 \times I_2$  + HOI). Thus, we conclude that VOI emissions make a negligible contribution to the total iodine flux.

Interfacial Model Results and Discussion. Figure 4 shows a comparison of the model results and  $I_2$  emissions observed from the four different types of solutions used in this study at atmospherically relevant ozone concentrations.

Artificial Solutions. The interfacial model was first used to predict iodine emissions over buffered KI solutions, and the



Figure 4. Observations (symbols) and modeling (lines) of  $I_2$  emissions at 17 °C as a function of I<sup>-</sup> concentration from buffered KI solutions (gray), artificial seawater (red), subsurface seawater (green), and surface microlayer (blue) for ozone concentrations of 22.7, 34.7, 38.6, and 38.5 ppbv, respectively. The error bars reflect the overall uncertainty on the measurements. The plot extends the calculation of the modeled  $I_2$  emissions back to an iodide concentration of  $1.5 \times 10^{-7}$  M, typical of natural oceanic surface iodide concentrations.

full dataset is shown in Figure S4A. The model captures the trends of  $I_2$  emissions with  $O_3$  and with  $I^-$  well, although it tends to underestimate the iodine flux at low  $[O_3]$ .

Modeled emissions over artificial seawater are compared to the observations in Figure S4B. Note that, as discussed in the methods section, a completing oxidation reaction of HOI (to iodate) by HOCl/OCl<sup>-</sup> was incorporated into the model to account for the ~50% decrease in I<sub>2</sub> emissions observed in AS compared to equivalent KI solutions. Although the model somewhat underpredicts I<sub>2</sub> emitted at low O<sub>3</sub>, overall the model shows skill in matching the observations. The experimental emissions for the lowest [I<sup>-</sup>] (black points, Figure S4) are close to the LoD, yet the modeled I<sub>2</sub> flux falls within the observational uncertainty.

Natural Seawater Samples. As detailed in the experimental results section, a substantial reduction in iodine emissions was observed from natural samples compared to artificial seawater and KI solutions. Previous studies on the reaction of gas-phase ozone with iodide solutions have established that  $I_2$  emissions are reduced in the presence of organics.<sup>19,40,43</sup> All of these studies attributed the reduction to a suppression of the liquid–gas transfer rate of  $I_2$ . Shaw and Carpenter<sup>43</sup> found that emissions were increasingly suppressed by DOC, by up to a factor of two, at ratios of [DOC]:[iodide] representative of their ambient reactivities to  $O_3(g)$ . Qualitatively, this is consistent with the reduction we observed in the SSW samples.

Using the same (but unmodified) interfacial model as we use in this study, Shaw and Carpenter<sup>43</sup> showed that neither DOC competing with I<sup>-</sup> to react with interfacial O<sub>3</sub> nor direct loss of I2 and/or HOI through reaction with DOC could fully explain the reduction of I2 emissions from SSW. Instead, they proposed a reduction in the net liquid-gas transfer rate of I<sub>2</sub> in SSW. Nevertheless, the reduction of the I2 liquid-gas transfer rate is a hypothesis that has hitherto not been explored in detail. Iodine  $(I_2)$  is a nonpolar molecule and is many times more soluble in organic solution than in water; for example, iodine has an octanol-water partition coefficient K<sub>OW</sub> of 309.58 An estimate of the octanol-air partition coefficient  $(K_{OA})$  of I<sub>2</sub> can be made by assuming  $K_{OA} = K_{OW}/K_{AW}$ , where  $K_{AW}$  is the air-water partition coefficient for I<sub>2</sub>.<sup>59</sup> Translated into the equations for mass transfer of I<sub>2</sub> under our laboratory conditions, the liquid-air mass transfer  $K_{\rm T}$  of I<sub>2</sub> from a pure octanol monolayer would be reduced by a factor of 99.3 compared to that from a purely aqueous solution at room temperature. We found that reducing the model's aqueous-air mass transfer term of I<sub>2</sub> at 17 °C from  $1.04 \times 10^{-6}$  to  $4 \times 10^{-7}$  $s^{-1}$  (i.e., ~40% of the pure water transfer term) produced a good agreement between the model and the SSW observations (see Figure S5A and further details in section 1.9 of the SI). Thus, assuming that the reduction in  $I_2$  emissions was entirely due to its increased solubility in the more organic-rich seawater than in pure water, this equates to an enhancement of I<sub>2</sub> solubility in seawater of about a factor of 6 (i.e., to  $\sim 0.2$  g/kg at 20 °C) compared to its value in pure water (0.03 g kg<sup>-1</sup> at 20 °C). Note that, while changes in solubility can explain the mass transfer of iodine under the still, laboratory conditions of the experiments presented here, additional factors caused by surfactants, such as, e.g., physical suppression of near surface mixing, might influence emissions under real-world conditions.

To explore the role of chemistry in reducing the  $I_2$  emissions, we modeled the loss of  $I_2$  and HOI through their reactions with DOC, as described in Materials and Methods. Including such chemistry had a negligible (<2%) impact on the

 $I_2$  emissions from SSW. This result strengthens our conclusion that  $I_2$  emissions are reduced in seawater compared to artificial seawater due to the enhanced solubility of  $I_2$ , rather than by its chemical loss.

I<sub>2</sub> emissions from SML samples were typically a factor of 3– 4 times lower than from subsurface seawater. The SML I<sub>2</sub> emissions were modeled satisfactorily, as shown in Figure S5B, by further reducing the aqueous–air mass transfer term for I<sub>2</sub> to  $1 \times 10^{-7}$  s<sup>-1</sup> (i.e., now only 10% of the pure water transfer term). This corresponds to the solubility of I<sub>2</sub> in the SML being around 5 times higher than in SSW, at around 1 g kg<sup>-1</sup> at 20 °C.

Figure 4 shows how the model performs well to predict iodine emissions from the natural samples over iodide concentrations up to  $4 \times 10^{-6}$  M for ambient ozone conditions. As the concentration of iodide in the open seawater generally ranges between 10 and  $150 \times 10^{-9}$  M,<sup>25</sup> this interfacial model can be a useful tool for predicting marine iodine emissions.

Volatile Organic lodine Emissions. I<sub>2</sub> and HOI reactions with DOC were included in the model to explore whether such chemistry could broadly explain the VOI emissions from the SML following the  $O_{3(g)}$  +  $I^-_{(aq)}$  reaction. We assumed a pseudo-first-order rate constant for the reaction of I2 and HOI with DOC of  $7 \times 10^{-3} \text{ s}^{-1}$  (see Materials and Methods), a 100% yield of VOI products (initially), and that VOI that is mixed downward out of the reacto-diffusive depth layer into the bulk is irreversibly lost, equivalently to I<sub>2</sub> and HOI. This gives a lower limit to the potential VOI emissions, since, unlike I<sub>2</sub> and HOI that react rapidly away in the bulk waters, some fraction of VOI molecules mixed down from the surface will persist long enough to be re-emitted. Nevertheless, this simple scenario produced VOI emissions an order of magnitude greater than we observed. However, it is known that reduction of I<sub>2</sub> and HOI emissions by DOC also leads to the formation of dissolved organic iodine (DOI), which was not monitored in our experiments, and reforms I<sup>-</sup> (e.g., ref 54, 60). We found that setting the VOI yield (from reaction of I<sub>2</sub> and HOI with DOC) to 5-10% gave the correct order of magnitude for the VOI emissions (1 to  $4 \times 10^7$  molecules cm<sup>-2</sup> s<sup>-1</sup> total VOI for  $[I^-]$  between 1 and 4 × 10<sup>-7</sup> M, Figure S6) and VOI fluxes scaled with the gaseous  $O_3$  concentration, as found experimentally (previous section, Figure 3). However, the model predicted an increase in VOI emissions as [I<sup>-</sup>] increased from 1 to  $4 \times 10^{-7}$  M, whereas the observed VOI emissions from the SML in Figure 3 actually declined; modeled VOI emissions only decline above  $[I^-] > \sim 1 \times$ 10<sup>-5</sup> M (Figure S6). Modeled VOI emissions as a fraction of the total iodine emissions (VOI + 2  $\times$  I<sub>2</sub> + HOI) decreased strongly with increasing  $[I^-]$ , which is likely due to the  $I_2 + I^$ reaction competing with  $I_2$  + DOC as  $[I^-]$  increases.

**Environmental Implications.** Our experiments show a clear reduction of molecular iodine emissions from the  $O_{3(g)} + I_{(aq)}^{-}$  reaction in seawater (compared to iodide solutions containing no added organics) over a broad range of iodide and ozone concentrations, confirming previous results.<sup>23,43,55,61</sup> For the first time, this reduction in I<sub>2</sub> is demonstrated to be larger for surface microlayer samples than for subsurface seawater samples. Unfortunately, there are very few observations of ambient open-ocean I<sub>2</sub> with which to compare our results. Lawler et al.<sup>61</sup> inferred an I<sub>2</sub> flux around 2.0 × 10<sup>7</sup> molecules cm<sup>-2</sup> s<sup>-1</sup> from measurements of night-time I<sub>2</sub> at Cape Verde ([O<sub>3</sub>] = 25 to 45 ppbv); however, the same paper

invoked range of I<sub>2</sub> fluxes  $7 \times 10^6$  molecules cm<sup>-2</sup> s<sup>-1</sup> to  $8.7 \times 10^7$  molecules cm<sup>-2</sup> s<sup>-1</sup> to model the diurnal cycles observed for I<sub>2</sub> and IO. Under similar conditions ([O<sub>3</sub>] = 38.5 ppbv, assuming oceanic [I<sup>-</sup>] =  $1.5 \times 10^{-7}$  M), the interfacial model constrained by our present measurements gives I<sub>2</sub> fluxes of 5.5  $\times 10^6$  molecules cm<sup>-2</sup> s<sup>-1</sup> for SSW and  $1.4 \times 10^6$  molecules cm<sup>-2</sup> s<sup>-1</sup> for SSW and  $1.4 \times 10^6$  molecules cm<sup>-2</sup> s<sup>-1</sup> for SML (extrapolated lines in Figure 4). Although our SML result is clearly lower, our SSW result is close to the lowest I<sub>2</sub> fluxes considered by Lawler et al. Interestingly, our SML result agrees well with the I<sub>2</sub> fluxes (1.4 to  $2.5 \times 10^6$  molecules cm<sup>-2</sup> s<sup>-1</sup>) reported from the early laboratory study of Garland and Curtis<sup>23</sup> on natural seawater "from the Dorset coast, U.K." (assumed [I<sup>-</sup>] =  $12.5 \ \mu g \ dm^{-3} = 1.0 \times 10^{-7} \ M$ , [O<sub>3</sub>] = 35 ppbv).

The presence of organics in natural seawater resulted in a very small flux of halocarbons, mainly CH<sub>3</sub>I, formed from chemistry subsequent to the surface reaction of ozone. The interfacial model predicts that the VOI flux makes its biggest relative contribution to the total iodine flux at low iodide concentrations (red line Figure S6), nevertheless the VOI emissions remain a negligible fraction (<5%) of the total iodine (VOI + 2 × I<sub>2</sub> + HOI) emissions for iodide concentrations relevant to environmental conditions.

We show that the observed reduction of  $I_2$  emissions is likely to be due to the increased solubility of I2 in the organicenriched seawater (compared to artificial seawater or buffered KI solutions). Our results are consistent with the solubility of  $I_2$  being a factor of 6 higher in SSW and  $\times 30$  higher in SML, compared to pure water. We calculated enrichment factors (EF), defined as the ratio of the SML over the SSW, based on the concentration of [DOC] (EF<sub>[DOC]</sub>) or on the surface pressure  $(EF_{\pi})$ . These can be used as an indication of the enrichment at the surface (see sections 1.4 & 1.5 of the SI and Table S2). The  $\text{EF}_{[\text{DOC}]}$  = 0.9 to 2.3 does not reflect the inferred factor of 5 difference in I<sub>2</sub> solubility between the SML and SSW. However, this difference in solubility does fall in the range of  $EF_{\pi}$  = 4.3 to 8.1 and hence seems more related to changes in the surface tension. More data are needed to confirm this relationship. This solubility effect may not result in reduced HOI emissions and may even lead to enhanced HOI emissions from organic-enriched seawater because HOI is very water-soluble. Experiments are highly desirable to confirm or otherwise this hypothesis.

# ASSOCIATED CONTENT

## **Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.est.0c02736.

Material and methods; literature overview of the influence of organics on iodine  $(I_2)$  emissions (Table S1); overview of sample locations and times for the natural seawater samples (Table S2); quantifier, qualifier, and retention times used in the GC analysis of halocarbons (Table S3); experimental setup for the molecular iodine  $(I_2)$  measurements by broadband cavity enhanced absorption (BBCEAS) (Figure S1); experimental setup for the halocarbon measurements (Figure S2); comparison of observed iodine fluxes for samples of different dates (Figure S3); observed and modelled iodine fluxes over natural

samples (Figure S5); relative contribution of simulated VOI emissions (Figure S6) (PDF)

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# Notes

The authors declare no competing financial interest.

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