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A laboratory characterisation of inorganic iodine emissions from the sea surface: dependence on oceanic variables and parameterisation for global modelling

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Abstract. Reactive iodine compounds play a significant role in the atmospheric chemistry of the oceanic boundary layer by influencing the oxidising capacity through catalytically removing O₃ and altering the HO_x and NO_x balance. The sea-to-air flux of iodine over the open ocean is therefore an important quantity in assessing these impacts on a global scale. This paper examines the effect of a number of relevant environmental parameters, including water temperature, salinity and organic compounds, on the magnitude of the HOI and I_2 fluxes produced from the uptake of O_3 and its reaction with iodide ions in aqueous solution. The results of these laboratory experiments and those reported previously (Carpenter et al., 2013), along with sea surface iodide concentrations measured or inferred from measurements of dissolved total iodine and iodate reported in the literature, were then used to produce parameterised expressions for the HOI and I₂ fluxes as a function of wind speed, sea-surface temperature and O₃. These expressions were used in the Tropospheric HAlogen chemistry MOdel (THAMO) to compare with MAX-DOAS measurements of iodine monoxide (IO) performed during the HaloCAST-P cruise in the eastern Pacific ocean (Mahajan et al., 2012). The modelled IO agrees reasonably with the field observations, although significant discrepancies are found during a period of low wind speeds $(< 3 \text{ m s}^{-1})$, when the model overpredicts IO by up to a factor of 3. The inorganic iodine flux contributions to IO are found to be comparable to, or even greater than, the contribution of organo-iodine compounds and therefore its inclusion in atmospheric models is important to improve predictions of the influence of halogen chemistry in the marine boundary layer.

1 Introduction

Reactive iodine compounds play an important role in the chemistry of the marine boundary layer (MBL) through their influence on ozone depletion and the oxidising capacity via repartitioning of HO_x and NO_x (Saiz-Lopez et al., 2012). Iodine oxides also form new particles spontaneously in coastal (O'Dowd et al., 2002) and polar regions (Atkinson et al., 2012), potentially leading to the production of cloud condensation nuclei. In coastal regions, emissions of molecular iodine and to a lesser extent of halocarbons from macroalgae are found to be the dominant source of reactive iodine to the MBL (O'Dowd et al., 2002). The source of reactive iodine and particles observed over the Antarctic sea ice remains to be determined (Atkinson et al., 2012).

Recent ground-, ship- and aircraft-based measurements have found that IO is ubiquitous over the open oceans, with MBL-averaged mixing ratios of around 0.5 pptv, and surface mixing ratios of up to 3 pptv (Allan et al., 2000; Mahajan et al., 2010, 2012; Dix et al., 2013; Gómez Martín et al., 2013; Großmann et al., 2013). Modelling studies have shown that these levels of IO cannot be sustained by the measured iodocarbon fluxes and that an additional source of reactive iodine from the open ocean, equivalent to > 50 % of the total surface iodine emission, is required to match the observations (Jones et al., 2010; Mahajan et al., 2010). Correlation studies of ground- and ship-based IO and reactive iodine $(IO_x = IO + I)$ measurements with oceanic variables have shown that there is a negative correlation with Chl a and CDOM (colored dissolved organic matter), suggesting that the additional iodine production over the oceans is not biological and could be inhibited by the presence of increased biological activity or organic matter (Mahajan et al., 2012; Gómez Martín et al., 2013; Großmann et al., 2013). This provides evidence for the widespread abiotic iodine source proposed by Garland and Curtis (1981): the sea surface oxidation of I⁻ by O₃ to yield HOI and I₂, which are then either released directly to the atmosphere or react with dissolved organic matter (Garland and Curtis, 1981; Martino et al., 2009; Carpenter et al., 2013). In addition, the correlation analysis showed significant correlations of IO_x with sea surface temperature (SST) and salinity (SSS), which suggests that this abiotic mechanism will be influenced by oceanic variables.

Previously we provided the first experimental evidence that both I₂ and HOI are emitted from the reaction of $I^- + O_3$ in the interfacial layer, and that these emissions can account for the missing source of reactive iodine over the tropical Atlantic Ocean (Carpenter et al., 2013). Very recently the first measurements of I₂ at a remote marine site (Cape Verde) have been reported (Lawler et al., 2014), with night time mixing ratios of up to 1.7 pptv. These mixing ratios are not enough to sustain the levels of IO observed at Cape Verde (Mahajan et al., 2010), which concurs with our laboratory observations of HOI being the major species emitted from the ocean surface. In this paper we investigate the dependence of the flux of inorganic iodine on relevant environmental parameters such as water temperature, salinity and the presence of organics. From these results and those previously reported (Carpenter et al., 2013), and using a kinetic model of the sea-air interface, parameterised expressions for the flux of both HOI and I₂ are derived. These expressions are then used in a 1-D atmospheric chemistry model for comparison with the field observations of reactive iodine species (IO and IO_x) recorded under a wide range of oceanic conditions during the HaloCAST-P cruise in the eastern Pacific Ocean (Mahajan et al., 2012).

2 Experimental

The experimental setup used has been described in detail elsewhere (Carpenter et al., 2013, Supplement) and involves detection of I₂ and HOI released from an ozonised iodide solution through selective photolysis, reaction of the resulting I radicals with an excess of O₃ to form iodine oxide particles (IOPs) (Saunders et al., 2010), and subsequent detection of these particles using a nano-differential mobility analyser (nano-DMA) (Fig. 1). For the selective photolysis, two methods were used: a xenon or tungsten lamp along with suitable

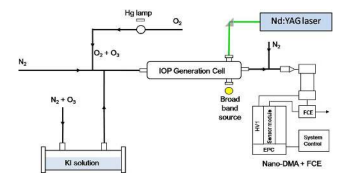


Figure 1. Experimental setup showing the two different types of light sources (W/Xe lamp or Nd:YAG laser) used to photolyse and distinguish between I_2 and HOI.

10 nm bandwidth interference filters, or photolysis at either 532 nm (I₂) (Saiz-Lopez et al., 2004) or 355 nm (HOI) (Rowley et al., 1999) using a frequency doubled or tripled Nd: YAG laser (Continuum Surelite). The laser energy was continuously monitored using a powermeter (Molectron Powermax 500A), so that the data could be normalised to the same pulse energy. The tungsten lamp showed high sensitivity for I₂ but did not allow measurements of HOI. The xenon lamp was used for measurements of HOI - however, the sensitivity and selectivity using this light source was poor. The laser was employed to measure I2 and HOI independently and enabled high sensitivity for both species and back-to-back operation. The experimental setup was calibrated using a known flow of I₂ vapour, produced by passing N₂ through a glass trap containing solid I₂ (Sigma Aldrich) in equilibrium with its vapour phase, for each of the light sources used. At the low I₂ concentrations and photolysis rates of the experiments the IOP mass is a linear function of the I2 concentration and accurate calibration coefficients can be obtained, typically with an uncertainty of ~ 8 % encompassing the uncertainty of the iodine vapour pressure (Baxter et al., 1907) and the statistical uncertainty from the linear regression. The detection limits using laser photolysis are typically ~ 2 pptv for I₂ and ~ 50 pptv for HOI. In the I⁻+O₃ experiments, the requirement to work at close to environmentally relevant concentrations of iodide and O_3 (within a factor of 10) results in a range of IOP numbers close to the detection limit of the particle counter, which may reduce the precision of the measurements.

For the temperature dependence experiments a doublewalled glass cell was used to allow cooled water to be flowed around the iodide solution. Temperatures ranging from 276 to 298 K were monitored using a thermocouple inserted into the solution in the cell. The iodide solutions were made up to the relevant concentrations using potassium iodide (KI) (Alfa Aesar, A12704) dissolved in deionised water buffered to pH 8 using 0.1 M sodium phosphate (Sigma Aldrich). For the salinity dependence experiments, reagent grade NaCl (Fisher Scientific S/3160/53) was added to 1×10^{-6} M KI solutions to give concentrations ranging from 0 to $0.5 \,\mathrm{M \, Cl^{-}}$. To estimate the trace iodide present as a potential contaminant in the NaCl, 0.104 M solutions in water (LC-MS Chromasolv - Fluka 39253) containing variable concentrations of KI (concentrations ranging from 0 to 99.4×10^{-9} M, acquisitions performed in duplicate) were prepared and directly infused into a Bruker MaXis Impact Q-TOF mass spectrometer in negative electrospray ionisation mode. Full profile data were acquired with a 1Hz acquisition rate for 3 minutes. The spectra were summed over the length of the acquisition and the relative intensity of the iodide peak (theoretical m/z 126.90522) relative to the cluster ion (NaCl)₂Cl⁻ (theoretical m/z 150.886645) was calculated. The spectra were calibrated using the $(NaCl)_n Cl^-$ clusters as internal references. The observed mass of the iodide ion peak was within 3 ppm of the theoretical value in all cases.

For the experiments involving organics, in the case of humic acid (Sigma Aldrich) a small amount was added to a prepared iodide solution which was then stirred overnight and any remaining solid filtered off. The concentration of humic acid was determined by UV/Vis spectroscopy to be around 2 mg dm^{-3} , using a specific absorbance of $5 \text{ L mg}^{-1} \text{ m}^{-1}$ at 254 nm (Weishaar et al., 2003). Sodium dodecyl sulfate (SDS, Sigma Aldrich) was added to a 1×10^{-7} M KI solution at a concentration of 6.2×10^{-3} M. Finally, for the experiments involving phenol, 1×10^{-7} M KI solutions containing phenol (Acros Organics) at concentrations ranging from 1×10^{-8} to 1×10^{-3} M were prepared.

3 Results

3.1 Salinity

The results of the salinity dependence experiments, using pH 8 buffered iodide solutions, are shown in Fig. 2. There is a clear increase in the I_2 flux with increasing salinity, with the flux around 2.5 times higher at 0.5 M NaCl (roughly equivalent to a salinity of 32 %) compared to a solution containing no chloride.

The quantitative mass spectrometric analysis of the NaCl employed in these experiments allows us to rule out a significant contribution of an iodide impurity to the increasing I₂ flux. The relative intensity of the iodide mass peak observed in the 0.104 M NaCl solution was less than that observed in the sample spiked with 9.9×10^{-9} M KI. Therefore, the 0.5 M NaCl solution contained less than 50×10^{-9} M I⁻ impurity, a factor of 20 smaller than the 1×10^{-6} M I⁻ concentration employed in the salinity experiments. The equivalent in terms of I₂ flux would be $< 6 \times 10^8$ molecule cm⁻² s⁻¹ at 0.5 M NaCl, i.e. well within experimental error ($\pm 4 \times 10^9$ molec. cm⁻² s⁻¹).

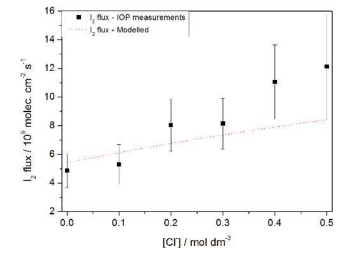


Figure 2. I_2 flux from solution as a function of chloride concentration, showing a clear increase in both model (red dashed line) and measurements (black squares).

3.2 Organics

A number of organic species were added to the KI aqueous solutions to investigate the effect of these compounds on the resulting I₂ flux. The first of these was humic acid (Sigma Aldrich), which was used as a proxy for dissolved organic matter found in seawater. Humic acid consists of a mixture of organic compounds formed from the degradation of organic matter. The I₂ flux was unaffected by the addition of relevant seawater concentrations ($\sim 2 \text{ mg dm}^{-3}$) of humic acid.

A further organic compound investigated was sodium dodecyl sulfate (SDS), which is a surfactant compound commonly found in a number of detergents and present at around $10 \,\mu g \, dm^{-3}$ in seawater (Ćosović et al., 1985). Surfactants are known to cause a reduction in the sea–air transfer of a number of atmospheric species by forming a barrier to emission. SDS was added to a 1×10^{-7} M KI solution at sufficient concentration to form a monolayer at the solution surface (6.2×10^{-3} M) (Hore et al., 2005). However, this amount of SDS was found to have no observable effect on the resulting IOP mass.

Previous studies have shown that phenol can have a significant effect on the I₂ flux in the I⁻+O₃ reaction (Hayase et al., 2010). It is commonly found in seawater, mainly from anthropogenic sources at concentrations of around 1×10^{-8} to 1×10^{-7} M. In our experiments, using realistic seawater concentrations of both iodide (1×10^{-7} M) and phenol (1×10^{-8} – 1×10^{-7} M) there was no difference observed in the measured I₂ flux within the 50% uncertainty of the observed IOP mass. The phenol concentration had to be increased up to 1×10^{-3} M before a 50% decrease in the I₂ flux was observed.

3.3 Temperature dependence

The temperature dependence of HOI and I2 emissions from ozonised iodide solutions was investigated using both broadband light sources (xenon and tungsten lamps) and the Nd:YAG laser source to photolyse I2 and HOI and produce detectable IOPs. The differing detection limits of the various systems necessitated different concentrations of I⁻ or O₃ to be able to measure both I2 and HOI. The iodide concentration used in these experiments ranged from 1×10^{-6} – 5×10^{-6} M and the ozone ranged from 222 to 3600 ppbv. In order to account for changes in the observed I₂ and HOI fluxes caused by different ozone and iodide concentrations, as well as for pH variation with temperature, the HOI and I₂ fluxes were normalised by dividing through by the O₃, I⁻ and H⁺ concentrations according to the individual experiments (we have shown previously (Carpenter et al., 2013) that the HOI and I₂ fluxes are proportional to the concentrations of these species over the ranges employed here). The Arrhenius plots for the resulting normalised fluxes are shown in Fig. 3, along with the 95 % confidence limits for the activation energies. The resulting activation energy for the I₂ flux is -7 ± 18 kJ mol⁻¹, and $17 \pm 50 \text{ kJ mol}^{-1}$ for the HOI flux (using conservative 95 % confidence limits rather than parameter errors). Even though the requirement of working under environmentally relevant conditions implies I₂ and HOI concentrations not far from the detection limit, the reproducibility of the room temperature data shows that the precision of the detection method is not a matter of concern. The major problem is the reproducibility of conditions of the experiment itself, i.e. in the cell containing the KI solution, and in particular in the experiments at low temperature, as shown by the error bars in Fig. 3. A possible explanation is uptake and release of iodine species on the inner walls and the outlet of the temperature stabilised cell.

4 Discussion

4.1 Salinity

There do not appear to have been any previous studies investigating the effect of chloride concentration on the rate of the $I^- + O_3$ reaction. The kinetic interfacial model described in Carpenter et al. (2013) satisfactorily predicts the observed positive dependence of the I₂ flux on the Cl⁻ concentration (Fig. 2). The reason for this increase is due to conversion of a small amount of HOI into I₂ via ICI:

$$Cl^- + HOI + H^+ \rightleftharpoons ICl + H_2O$$

 $ICl + I^- \rightleftarrows I_2Cl^-$

 $I_2Cl^-\rightleftarrows I_2+Cl^-.$

Note that this has very little effect on the HOI emission due to its great excess in the interfacial layer (Carpenter et al.,

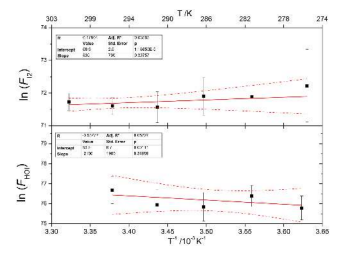


Figure 3. Arrhenius plots for I_2 and HOI produced from the combined data using each light source normalised for $[O_3]$, $[I^-]$ and $[H^+]$. Dashed lines indicate the 95 % confidence limits of the linear fits.

2013). As shown in Fig. 2, the observations exhibit a slightly larger chloride dependence, with a greater I_2 flux observed at 0.5 M Cl⁻ than the model predicts. This probably indicates the additional effect of the surface enhancement of I⁻ ions, which has been reported in a number of previous experimental and theoretical studies (Ghosal et al., 2005; Caleman et al., 2011; Gladich et al., 2011). In these studies I⁻ is shown to have a greater preference for the surface of an aqueous solution than Br⁻ or Cl⁻, which has been explained by the more polarisable I⁻ anions having a lower excess surface free energy compared to either Br⁻ or Cl⁻ (Jungwirth and Tobias, 2005; Gladich et al., 2011).

Although there is a clear dependence of the I_2 flux on the Cl⁻ concentration, in seawater this effect is unlikely to be important due to the very small changes in SSS. Over the range of SSS commonly observed in the oceans (32–37 %*c*), the resulting change in the I_2 flux based on the experimental results would be only around 1 %.

4.2 Organics

The results of our humic acid experiments are in agreement with those of Hayase et al. (2012) who also found no effect on I₂ emission from the addition of humic acid. However, they did observe an increase in I₂ emission when fulvic acid was added which they attributed to greater efficiency of proton supply from the carboxylic acid group compared to water. Hayase et al. (2010) investigated the effect that certain organic compounds have on the I⁻ + O₃ reaction. They found that phenol and other species containing the phenol functional group had a marked effect on the iodine emitted to the gas phase. Working at concentrations which were significantly higher than ambient levels (5 × 10⁻³ M NaI and 1 × 10⁻³ M phenol, at pH 6, so most phenol was phenolate), the I₂ and IO produced was around half that of a pure NaI solution. This was attributed to the fast reaction of the phenolate ion with O₃ $(1.4 \times 10^9 \text{ M}^{-1} \text{ s}^{-1})$, which competes with that of I⁻. Our results agree with that earlier study. However, the concentrations of phenol are orders of magnitude higher than that found in seawater, so phenolic compounds should not play an important role in regulating the emission of I₂ from the ocean.

Reeser and Donaldson (2011) showed that addition of an aqueous monolayer of octanol to ozonised KI solutions resulted in a decrease in the observed I₂ emission by a factor of ~ 2 , due to changes in gas-solution partitioning. Rouviere and Ammann (2010) found that certain surfactant species form a barrier to ozone uptake from an iodide solution, which implies inhibition of the release of iodine to the gas phase. Their work suggests, however, that chain lengths of > C15 may be necessary for a significant reduction in the O₃ loss. This is most likely due to the structure of the monolayer formed and how densely the monomers are packed. SDS has a carbon chain length of C11 which may explain the lack of an inhibiting effect observed in our experiments. Recently, Shaw and Carpenter (2013) have also reported a suppression of a factor of ~ 2 of the I₂ emission when dissolved organic carbon (DOC) of coastal seawater origin is added to ozonised iodide solutions, under conditions of O₃ reactivity with iodide and DOC representative of the open ocean, which they have also attributed to a surface physical effect.

4.3 Temperature

Magi et al. (1997) used the droplet train technique to measure the disappearance of O₃ following uptake in I⁻ solution droplets (0.5–3 M NaI). The O₃ concentrations used were not stated in the paper. They reported a large positive temperature dependence for the I⁻+O₃ rate coefficient, with an activation energy of 73 ± 30 kJ mol⁻¹. However, the Arrhenius expression they reported ($k = 1.44 \times 10^{22} \exp(-73080/RT)$ M⁻¹ s⁻¹) contains a pre-exponential factor which is roughly 10 orders of magnitude greater than the diffusion limited reaction rate, and therefore appears to be unphysical.

Hu et al. (1995), using a similar droplet train technique with I⁻ concentrations from 0.5–3 M and O₃ in the range 5×10^{12} to 1×10^{14} cm⁻³, measured $k(I^- + O_3) = 4 \times 10^9$ M⁻¹ s⁻¹ at 277 K. Although they do not report a room temperature measurement, the room temperature value for $k(I^- + O_3)$ appears to be reasonably well established from a number of previous studies (Garland et al., 1980; Hoigné et al., 1985; Magi et al., 1997; Liu et al., 2001), lying in the range $(1.2-2.4) \times 10^9$ M⁻¹ s⁻¹. Thus the Hu et al. (1995) result implies a slightly negative temperature dependence, although those authors state that within their experimental accuracy the reaction could have been temperature independent. We therefore assumed the temperature dependence for $k(I^- + O_3)$ to be zero for modelling the I₂ and HOI fluxes using the interfacial kinetic model (Carpenter et al., 2013) and included the various temperature dependences of parameters such as Henry's law constants, diffusion constants, mass transfer velocities, etc. This gave an activation energy for the modelled I₂ emission between -6 and 2 kJ mol^{-1} over the concentration ranges of I⁻ and O₃ used in the present study, in agreement with the experimental results (Fig. 3). For HOI, the modelled activation energy ranged from 20 to 30 kJ mol^{-1} , again within experimental error (Fig. 3).

4.4 Modelling reactive iodine field measurements

Recent field observations in the eastern Pacific (Mahajan et al., 2012) and Galápagos Islands (Gómez Martín et al., 2013) have shown a probable link between IO_x and SST on both temporal and spatial scales (Fig. 4). In contrast, there is either an insignificant or a negative correlation of IO_x with CDOM and Chl a, indicating that an inorganic ocean source is the most likely explanation of the missing contribution to IO_x . It is difficult to compare the available IO_x measurements as a consequence of the different experimental approaches adopted, which sampled different regions of the marine boundary layer. In addition, there is a lack of concurrent O₃ measurements, making the conversion from IO to IO_x more unreliable. In general, however, integrated over a 1 km boundary layer, the average daytime IO_x is around 0.5 pptv, with the IO_x at the surface around 2 pptv. Larger IO_x mixing ratios are generally observed towards the equator, which gradually decrease at higher latitudes (Fig. 4).

Sea surface iodide (SSI) data, measured (Tsunogai and Henmi, 1971) or derived from measurements of iodate and total iodine (Truesdale et al., 2000), also show a clear latitudinal variation with an increase in iodide towards the equator. According to Truesdale et al. (2000), the polewards increase in IO_3^- (decrease in I⁻) corresponds with the overall increase in vertical mixing in the water column, caused by the polewards decrease of SST. Thus, in tropical regions stratification appears to allow IO_3^- to be reduced to I⁻ in the sea surface either by blocking IO_3^- replenishment from below, or by facilitating undetermined reduction processes by prolonged isolation of surface water. Note that despite the pre-eminent role attributed to marine biota in the reduction of IO_3^- to I^- , IO_3^- does not correlate well with primary productivity (Truesdale et al., 2000).

From the discussion above, it follows that the surface iodide–ozone emission mechanism and the link between SSI and SST are a likely explanation for the observed correlations between SST and IO_x measurements. We have therefore combined the measurements of SSI and SST from several cruises in the Atlantic and Pacific oceans (Tsunogai and Henmi, 1971; Elderfield and Truesdale, 1980; McTaggart et al., 1994; Truesdale et al., 2000; Huang et al., 2005) into an Arrhenius-type plot in Fig. 5. This exhibits a sensibly linear

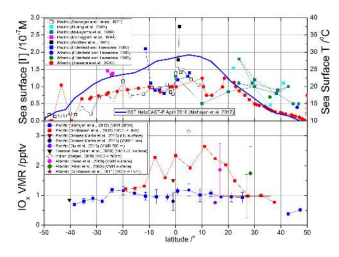


Figure 4. The top panel shows measured sea surface iodide from a number of ship campaigns in the Pacific and Atlantic oceans (Tsunogai and Henmi, 1971; Elderfield and Truesdale, 1980; Nakayama et al., 1989; Woittiez et al., 1991; McTaggart et al., 1994; Truesdale et al., 2000; Huang et al., 2005) along with SST measurements in the Pacific Ocean in March-April 2010 (Mahajan et al., 2012). The bottom panel shows IO_x volume mixing ratios from campaigns in the open ocean at different latitudes (Allan et al., 2000; Read et al., 2008; Oetjen, 2009; Großmann et al., 2011, 2013; Mahajan et al., 2012; Dix et al., 2013; Gómez Martín et al., 2013). In the legend, VMR refers to volume mixing ratio (layer height stated for MAX-DOAS measurements), VCD refers to vertical column density converted to mixing ratio using the indicated column height, and UL means upper limit. Scaled VCDs are used instead of mixing ratios in some cases where the uncertainty range from radiative transfer calculations obscures the latitudinal variability.

dependence of $\ln [I^-]$ with SST⁻¹ (apart from a small number of outliers marked in red in Fig. 5). The resulting Arrhenius expression

$$[I_{(aq)}^{-}] = 1.46 \times 10^{6} \times \exp\left(\frac{-9134}{\text{SST}}\right)$$
(1)

describes the iodide concentration (in mol dm⁻³) at the sea surface as a function of SST (in K). From the 95 % prediction bands plotted in Fig. 5, the uncertainty in [I⁻] from Eq. (1) is estimated to be ~ 50 % of the predicted value. This expression can then be incorporated into the flux expressions derived previously (Carpenter et al., 2013):

$$F_{\text{HOI}} = [O_{3_{(g)}}] \times \left(4.15 \times 10^5 \times \left(\frac{\sqrt{[I_{(aq)}]}}{\text{ws}} \right) - \left(\frac{20.6}{\text{ws}} \right) -23\,600 \times \sqrt{[I_{(aq)}]} \right)$$
(2)

$$F_{I_2} = [O_{3_{(g)}}] \times [I_{(aq)}^-]^{1.3} \times (1.74 \times 10^9 - (6.54 \times 10^8 \times \ln ws)),$$
(3)

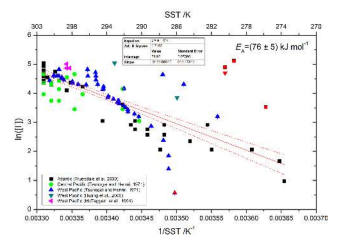


Figure 5. Arrhenius plot of SSI vs. SST field measurements in the Pacific and Atlantic oceans (Tsunogai and Henmi, 1971; McTaggart et al., 1994; Truesdale et al., 2000; Huang et al., 2005). Red points are outliers removed from the fit (most of them correspond to a high iodide episode at low latitude during the Atlantic cruise). The activation energy obtained from the straight line fit is (76 ± 5) kJ mol⁻¹ ($R^2 = 0.71$). Dashed lines: confidence bands (red) and prediction bands (blue) at 95 %.

where the flux is in nmol $m^{-2} d^{-1}$, $[O_{3_{(g)}}]$ is in ppbv (or nmol mol⁻¹), $[I_{(aq)}^{-}]$ in mol dm⁻³ and wind speed (ws) in m s⁻¹.

Equations (2) and (3) are room temperature expressions, since Carpenter et al. (2013) did not consider the potential temperature dependence of any processes in the interfacial layer. As discussed above, a large temperature dependence of the I^-+O_3 reaction can be ruled out, but the modest experimental precision does not tightly constrain the activation energies of the iodine fluxes (see Fig. 3). The interfacial layer model predicts activations energies for I2 and HOI emission of $\sim -2 \text{ kJ mol}^{-1}$ and 25 kJ mol^{-1} , respectively, which are within the experimental uncertainty of our measurements. Table 1 presents a summary of the relative error $(F_X(T) - F_X(293 \text{ K}))/F_X(293 \text{ K})$ made when using the 293 K expressions (2) and (3) at 283 and 303 K using different values for the activation energies. For this exercise, an additional Arrhenius-type exponential factor is assumed in (2) and (3), and the experimental uncertainty limits for the activation energies (E_{up} and E_{low}) and the activation energies derived from the interface kinetic model (E_{model}) are considered. The uncertainties in the HOI flux arising from the uncertainty in the activation energy are larger than for the I₂ flux, and since HOI this is the major inorganic species emitted, this must be kept in mind when modelling field data at SST other than 293 K.

Equations (1)–(3) provide a convenient parameterisation of the inorganic iodine flux for modelling purposes, since SST measurements are much more widely available than SSI measurements. Note that the fluxes are predicted to be

	I ₂			HOI		
	E _{low}	E _{up}	E _{model}	$E_{\rm low}$	E _{up}	E _{model}
	$-25 \mathrm{kJ} \mathrm{mol}^{-1}$	$11 \text{kJ} \text{mol}^{-1}$	$-2 \mathrm{kJ}\mathrm{mol}^{-1}$	$-33\mathrm{kJmol^{-1}}$	$67 \mathrm{kJ} \mathrm{mol}^{-1}$	$25 \mathrm{kJ} \mathrm{mol}^{-1}$
303 K	29 %	-16 %	3 %	37 %	-153 %	-41 %
283 K	-45%	15 %	-3%	-63%	63 %	31 %

Table 1. Relative differences of the inorganic iodine fluxes at 283 K and 303 K with respect to the 293 K values, for the experimental confidence limits and the modelled values of the activation energies.

highest under conditions of high $[O_{3(g)}]$, high $[I_{(aq)}]$, and/or low wind speed. The wind speed relationship arises from the assumption that wind shear (only) drives turbulent mixing of the interfacial layer with bulk seawater, thus reducing the proportion of I₂ and HOI evading into the atmosphere (Carpenter et al., 2013).

To test these expressions, daytime average measurements of IO_x recorded during the HaloCAST-P cruise in the eastern Pacific were compared with model output from the 1-D model THAMO (Saiz-Lopez et al., 2008). The HaloCAST-P cruise (Mahajan et al., 2012) was a 4-week cruise from 27 March to 26 April 2010, starting at Punta Arenas at the southern tip of Chile and continuing northwards to Seattle, USA. Measurements of IO were performed using a Multi-Axis Differential Optical Absorption Spectroscopy (MAX-DOAS) instrument, as well as measurements of CH₃I in seawater and air using gas chromatography-mass spectrometry (GC-MS) along with ancillary data such as SST, ws, Chl a, CDOM and SSS. IO_x mixing ratios were derived from the observed IO and CAM-Chem modelled O₃, assuming low NO_x concentrations (O_3 was not measured during the cruise), for which the IO_x partitioning is controlled by IO photolysis and the $I + O_3$ reaction, resulting in an IO/I ratio of ~ $0.12*[O_3] \text{ ppbv}^{-1}$ (for SZA < 60°).

Using the observed SST and ws from the HaloCAST-P cruise, and modelled O₃ from CAM-Chem (Fig. 6) (Mahajan et al., 2012), the predicted HOI and I₂ fluxes were calculated using Eqs. (1)–(3). The sum of these is plotted against latitude in Fig. 6a and against the measured IO and derived IO_x in Fig. 7. The three points highlighted in red in Fig. 7 were excluded from the linear regression fits. These excluded points correspond to measurements at the equatorial front in the Pacific Ocean (15-25° N, see Fig. 6d) , where the modelled O₃ concentrations are considerably more uncertain than earlier in the campaign. In this region the O₃ concentration is highly dependent on the water vapour concentration, which in turn will depend on the season and year due to fluctuations in temperature. As a result, the climatological O₃ output from CAM-Chem may not be representative of the actual O_3 during the Thompson cruise. The R^2 correlation coefficients with the IO_x and IO measurements (after excluding these points) were only 0.10 and 0.15 (p = 0.09), respectively. An interesting point to note is the significant intercept observed in both plots; 0.66 pptv for IO_x and 0.48 pptv for IO. This may be indicative of the contribution of iodocarbons to the total IO_x . The modelling work performed by Mahajan et al. (2012) suggested that around 30% of the IO_x could be explained by the observed emissions of CH₃I, and there are likely other iodocarbon species which will also contribute. As such, the 1-D model THAMO was initiated using only halocarbon fluxes with no contribution from the additional inorganic iodine emissions so that the resulting IO and IO_x matched the observed intercepts. These values were then fixed for subsequent model runs.

THAMO is a 1-D chemistry transport model (Saiz-Lopez et al., 2008) consisting of 200 stacked boxes of 5 m resolution giving a total height of 1 km. K_{zz} profiles were constructed for each measurement point based on the measured ws, assuming convective conditions and a capped boundary layer of 1 km. A number of species were constrained in the model using typical measured values over the open ocean: $[NO_x] = 25 \text{ pptv}$, [CO] = 110 ppbv, $[DMS] = 30 \text{ pptv}, [CH_4] = 1820 \text{ ppbv}, [ethane] = 925 \text{ pptv},$ $[CH_3CHO] = 970 \text{ pptv},$ [HCHO] = 500 pptv,[isoprene] = 10 pptv, [propane] = 60 pptv, [propene] = 20 pptv(Carpenter et al. (2013), Supplement). Equations (1)-(3) were included in THAMO to calculate the HOI and I₂ fluxes, driven by the measured ws and SST, and the CAM-Chem modelled O₃ mixing ratios at each point on the Thompson cruise track. Because HOI and I₂ can build up in the MBL close to the ocean surface, and therefore suppress the emission from the sea-air interface, the Henry's law constants and transfer coefficients $(K_{\rm T})$ for HOI and I₂ were calculated directly in the model (as described in Carpenter et al. (2013), Supplement).

The IO and IO_x predicted by THAMO, along with the measured IO and derived IO_x , were then plotted against latitude (Fig. 6d) and against the total inorganic iodine flux (Fig. 8). Note that the model IO and IO_x was computed by averaging over the first 200 m of the model vertical profiles (first 40 boxes) using only daylight data. This was done because the measured IO and IO_x mixing ratios were calculated from the MAX-DOAS IO slant column densities (dSCDs) measured with a 1° elevation angle, which should only be sensitive to the first 200 m in the boundary layer. The IO mixing ratios were then calculated using the O₄ method

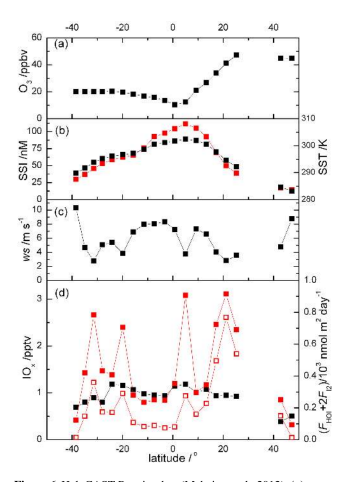


Figure 6. HaloCAST-P cruise data (Mahajan et al., 2012): (**a**) ozone (modelled using CAM-Chem, see text); (**b**) sea surface temperature (black, right axis) and iodide (red, left axis) derived from SST using Eq. (1); (**c**) wind speed; and (**d**) IO_x (left axis) and inorganic iodine flux (right axis). (**d**) shows IO_x derived from measurements (black) and modelled in this work using THAMO employing the shown inorganic iodine source function (empty red squares).

(Mahajan et al., 2012), and for quality control only daylight data with SZA $\leq 60^{\circ}$ were used.

Although there is reasonable agreement for the points with lower computed inorganic iodine flux, the predicted IO and IO_x is about a factor of 2 larger than measured at relatively high HOI and I_2 fluxes. Figure 6d also shows how the modelled IO_x around 20° N does not scale with the enhanced calculated flux at this latitude (equatorial front). The model sensitivity to the three key parameters in Eqs. (1)–(3) is described below, leading to a discussion of the possible reasons for the discrepancies observed.

4.5 Model sensitivity to ozone, SST (iodide) and wind speed

Sensitivity studies were conducted to investigate the effect of changing O_3 , SST and wind speed on the inorganic iodine fluxes predicted by Eqs. (1)–(3). The results of these inves-

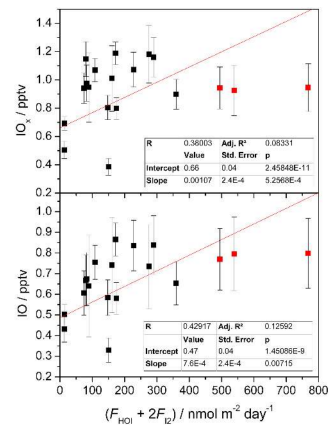


Figure 7. Measured IO_x (top panel) and IO (bottom panel) from the HaloCAST-P cruise against predicted inorganic iodine flux from the parameterised flux expressions for each data point. Points in red were excluded from the linear fit.

tigations are shown in Fig. 9. There is a weak sensitivity of IO to increasing O₃ (~ 0.01 pptv ppbv⁻¹ at noon), while IO_x decreases from 5 to 25 ppbv ozone and then converges with IO. This is because at higher O_3 concentrations most of IO_x is in the IO form, and a larger fraction of IO undergoes selfreaction and ends up as I_xO_y (there is around 10 times more $I_x O_y$ produced at 50 ppbv compared to 5 ppbv) which acts as a sink for IO_x in the model, forming IOPs. At low O₃ concentration (5 ppbv) a significant part of the reactive iodine remains as less reactive I atoms rather than being converted to IO, and therefore the IO_x remains high. In summary, the IO second-order losses counterbalance the significant increase in inorganic iodine flux for a 10-fold increase in ozone and may explain why the large increase of calculated inorganic flux around the equatorial front (Fig. 6d) does not reflect in the modelled IO_x mixing ratio.

An interesting result of this exercise is that it shows that IO is also produced at night (see Fig. 9). This is due to the presence of NO₃ and its reaction with the freshly emitted I_2 (Saiz-Lopez and Plane, 2004) This reaction produces I atoms which can further react with O₃ to produce IO. This reaction would not be important under clean conditions due to the

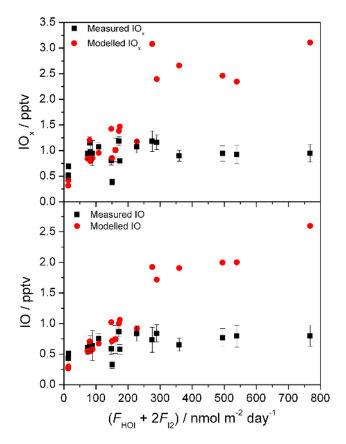


Figure 8. Measured (black squares) and THAMO modelled (red circles) IO_x (top panel) and IO (bottom panel) for all HaloCAST-P cruise data against predicted inorganic iodine flux from the parameterised expressions.

very low NO_x levels and indeed, the modelled IO is well below the detection limit of typical DOAS instruments. This may, however, be an important reaction for polluted conditions with much higher NO_x levels and there is the possibility that HOI will also react with NO_3 to produce IO.

The increase in IO and IO_x with SST is explained by the increase in SST leading to a higher predicted iodide concentration in the surface layer (Eq. 1). This will in turn lead to an increased predicted inorganic iodine flux from the flux expressions above. At a fixed ozone concentration of 25 pptv, the modelled IO_x shows a sensitivity to SST of ~ 0.05 pptv K⁻¹, which compares well with the 0.03 pptv K⁻¹ observed in the HaloCAST-P campaign.

Wind speed has a dramatic effect on the predicted IO and IO_x , with an increase observed at lower wind speeds. It is clear from inspection of Eq. (2) that the HOI flux will be heavily dependent on the wind speed due to the ws⁻¹ relationship. This means that Eq. (2) will become invalid at low wind speeds (< 3 m s⁻¹). A comparison of the predicted HOI and I₂ fluxes using Eqs. (1)–(3) with the output of the kinetic interfacial model (Carpenter et al., 2013) shows that there is increasing deviation at wind speeds less than 3 m s⁻¹. For

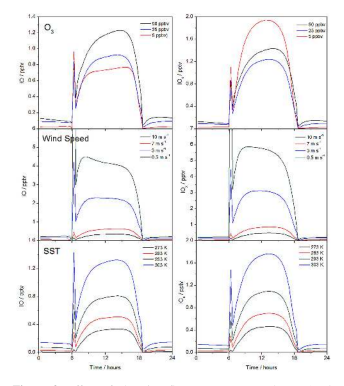


Figure 9. Effect of changing flux parameters on the THAMO modelled mixing ratios of IO and IO_x averaged over the first 200 m. For each sensitivity test, the two fixed parameters are set to the average of the HaloCAST-P campaign: $[O_3] = 25$ ppbv, ws = 6 m s⁻¹ and SST = 296 K. Top panels: modelled diurnal IO and IO_x with varying O₃ (black line = 50 ppbv, blue line = 25 ppbv, red line = 5 ppbv). Middle panels: modelled diurnal IO and IO_x with varying wind speed (black line = 10 m s⁻¹, red line = 7 m s⁻¹, blue line = 3 m s⁻¹, turquoise line = 0.5 m s⁻¹). Bottom panels: modelled diurnal IO and IO_x with varying SST (black line = 273 K, red line = 283 K, turquoise line = 293 K, blue line = 303 K).

instance, Eqs. (1)–(3) overpredict the combined HOI and I_2 fluxes by a factor of two at a wind speed of 0.5 m s⁻¹.

4.6 Discussion of discrepancies between observed and predicted IO

The high predicted inorganic iodine flux around 20° N is generated by high O₃ (from CAM-Chem) and low measured wind speeds (see Fig. 6a, c and d). As mentioned above, the high O₃ from CAM-Chem may not accurately represent the O₃ during the Thompson cruise. In order to investigate this possibility, the model was run for the conditions of the three highest flux points in Fig. 8, varying the O₃ concentration until the resulting IO matched the measured mixing ratios (the IO was used to match the observations, as the IO_x measurements were derived from the O₃ mixing ratios predicted by CAM-Chem). The O₃ had to be reduced to 2 ppbv before the observations could be matched. In fact, open ocean ozone mixing ratios below the detection limit of commercial instruments (~2 ppbv) have been reported (Gómez Martín et al., 2013; Großmann et al., 2013). However, we have also shown above that an enhanced iodine inorganic flux due to high ozone mixing ratios does not result in a large increase of IO_x mixing ratios due the loss to IOPs via IO self-reaction. This is clearly illustrated in Fig. 6 by similar modelled IO_x at 5° N and 20° N for similar wind speed and a difference of a factor of ~4 in the climatological ozone mixing ratios.

The model overpredictions could also be attributed to deviations of the iodide concentration from the parameterisation based on SST measurements. As shown in Fig. 5, the general trend of SSI is to follow the changes in SST. However, there are a number of localised phenomena which can alter this relationship, such as upwelling, lateral advection and rainfall (Nakayama et al., 1989; Truesdale et al., 2000). If the iodide at some points of the cruise track was significantly lower than indicated by the SST measurements this would lead to a reduced inorganic iodine flux, and therefore lower IO and IO_x. It is conceivable that SSI could show seasonal variability around the equatorial front which is not captured by the SST parameterisation. Note, however, that a related parameter such as SSS (Mahajan et al., 2012) shows a smooth behaviour throughout the HaloCAST-P cruise, which does not suggest strong localised phenomena affecting SSI. Regarding a potential depletion of iodide in the top layer of the ocean, at the highest calculated inorganic iodine flux ($\sim 800 \text{ nmol m}^{-2} \text{ d}^{-1}$), over the course of 24 h the iodide would be depleted in a layer at the surface of around 8 mm. Using the diffusion coefficient for iodide of $2 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ (Friedman and Kennedy, 1955), it would take around 2.5 h for the iodide in this layer to be replenished from below. Hence this explanation does not appear likely.

The sensitivity analysis in the previous section shows that the model is extremely sensitive to ws. In fact, a comparison of Fig. 6c and d shows that a correlation exists between deviation of simulated mixing ratios from observed data and wind speeds equal to or lower than 3 m s^{-1} . Below, we examine several reasons which could lead to an overestimation of IO_x by the model as a result of insufficient detail in the treatment of wind-related phenomena.

First, it is plausible that the interfacial model overpredicts I₂ and HOI emissions at very low wind speeds. The interfacial model only includes wind shear-driven turbulent mixing of the interfacial layer with bulk seawater. However, vertical transport from the surface to the bulk mixed layer is governed by a combination of processes, including convection generated by cooling at the surface, Langmuir turbulence, rainfall, etc. Convection is important for mixing when wind is in the low to intermediate speed regime (Rutgersson et al., 2011), thus it is likely that mixing of interfacial iodine to bulk waters is underestimated under low wind speed conditions. Second, it can be hypothesised that reduced wind speeds over the ocean enable the formation of organic films at the surface (Frew et al., 2004), inhibiting the sea-to-air exchange of I₂ and HOI (Reeser and Donaldson, 2011; Shaw and Carpenter, 2013).

A further possibility is that low wind speeds result in much less vertical mixing above the surface of the ocean, causing an accumulation of I₂ and HOI immediately above the ocean surface, which would suppress further emission due to equilibration. High mixing ratios of I₂ and HOI above the surface during daylight hours could lead to sufficient IO concentrations for IOP formation to occur (this is a highly non-linear process, as shown by Mahajan et al. 2010), reducing the overall IO mixing ratio over the first 200 m. To test this possibility, the K_{77} profile for the three highest predicted IO concentrations was adjusted to be an order of magnitude lower than originally calculated in the lowest 5 m box. However, this was found to have a limited effect on the modelled daytime average IO, with a reduction of only around 1 %. Note also that reduced atmospheric vertical mixing at low ws would retain the IO close to the ocean surface for a number of hours, creating a much steeper vertical gradient. The MAX-DOAS daytime average could possibly be insensitive to the IO that is very close to the ocean surface (the scattering altitude in the visible even for an elevation angle of 1° is tens of metres). If low ws conditions are sustained during the day the boundary layer would not develop as efficiently and so for the same scattering altitude, but lower mixing rates, a daytime averaged MAX-DOAS measurement could be lower under low ws, compared to high ws, for a very reactive species.

5 Conclusions

The experimental results reported show that over the large range of conditions considered in the laboratory, parameters such as salinity can have a large effect on the resulting total inorganic iodine flux, although these effects are likely to be minimal under real seawater conditions where the variations in salinity tend to be relatively small. None of the organics used in the reported experiments had any significant effect on the resulting inorganic iodine fluxes, but we cannot rule out the possibility that surface organic films may play a role in limiting the transfer of HOI and I_2 to the gas phase in real seawater.

SST was found to be a good proxy for SSI concentrations reported in the literature, and due to the wide availability of SST measurements, the derived relationship provides a useful tool for atmospheric modelling purposes. Modelled and measured IO and IO_x were found to be in reasonable agreement when the predicted I2 and HOI fluxes were low, although under low wind speed conditions there is a substantial over-prediction in the model. A limiting factor may need to be introduced for modelling IO and IO_x under very calm ocean conditions. Based on the comparison with measurements in this paper, a simple approach would be to set a lower limit of ws to 3 m s^{-1} when implementing Eqs. (1)–(3) in a model. The modelling results indicate that consideration of the inorganic iodine flux from reaction of O₃ and I⁻ is necessary for reproducing measurements of iodine oxides over the open ocean, although further experimental work is required to refine the parameterised flux expressions. These should be also tested against other long term/wide spatial coverage observations of reactive iodine recently reported.

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