

1 A novel technique for the humidity dependent calibration of 2 hypiodous acid (HOI) and iodine (I₂)

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12
13 **Abstract.** Hypiodous acid (HOI) and molecular iodine (I₂) are important precursors of reactive gaseous iodine, which plays
14 an important role in the oxidative capacity of the atmosphere and in aerosol formation in the marine boundary layer. HOI and
15 I₂ are emitted from the ocean surface and recycled on atmospheric aerosol via heterogeneous chemistry. Measurements of
16 these molecules, which are typically present in the marine boundary layer at the low-to-sub part per trillion (ppt) level, are
17 sparse, in part due to difficulties in quantification with a lack of appropriate instrumentation and calibration techniques. A
18 novel calibration technique is developed for HOI via generation from I₂ hydrolysis and then 1:1 conversion of HOI back to I₂
19 through a NaI trap, allowing the sensitivity of HOI to be calculated relative to I₂, which is readily calibrated using a permeation
20 tube system. Using this calibration method, we describe the use of a reduced pressure high resolution chemical ionisation mass
21 spectrometer (CIMS) to characterise the sensitivities of HOI and I₂ over a range of humidities representative of the marine
22 boundary layer and to measure these molecules in the field. At humidities of over 50% RH, the CIMS sensitivity of I₂ is
23 humidity independent whereas HOI exhibits a slight negative humidity dependence. The effect of inlet interactions on HOI
24 and I₂ signals is investigated, with HOI observed to convert to I₂. The implications of these inlet interactions and humidity
25 sensitivities for future ambient measurement configurations are discussed.

26 1 Introduction

27 Iodine species play an important role in atmospheric chemistry and climate. They influence the oxidative capacity of the
28 atmosphere through the destruction of ozone (O₃) (Chameides and Davis, 1980; Solomon et al., 1994; Read et al., 2008; Saiz-
29 Lopez et al., 2014; Sherwen et al., 2016a) and alteration of hydrogen oxides (HO_x) and nitrogen oxides (NO_x) cycles (Simpson

30 et al., 2015; Saiz-Lopez and Von Glasow, 2012). They can also oxidize elemental mercury (Calvert and Lindberg, 2004;
31 Auzmendi-Murua et al., 2014; Lee et al., 2024) and are involved in new particle formation, with iodine oxide radicals (IO),
32 produced during the reaction between O₃ and iodine, self-reacting and further ozonising to form I₂O₅ which can hydrolyse to
33 form HIO₃ (O'dowd et al., 2002; Sipilä et al., 2016; Finkenzeller et al., 2023).

34
35 Of the halogens, iodine has the most profound impact on tropospheric O₃ cycling and significantly modifies the atmospheric
36 response to anthropogenic perturbations of ozone precursor emissions. Global model simulations show that atmospheric iodine
37 chemistry lowers the global tropospheric O₃ burden by 6–20% and reduces O₃ concentrations by several parts per billion (ppb),
38 with significant effects over polluted and populated regions (Saiz-Lopez et al., 2014; Sherwen et al., 2016a; Sherwen et al.,
39 2016b; Wang et al., 2021b; Pound et al., 2023; Caram et al., 2023). The dominant source of atmospheric iodine is from the
40 oceans, primarily from the reaction of O₃ and iodide on the sea surface (Garland et al., 1980; Carpenter et al., 2013; Macdonald
41 et al., 2014), resulting in the emission of molecular iodine (I₂) and hypoiodous acid (HOI). HOI is thought to be the largest
42 source, representing ≈75% of iodine emissions into the troposphere (Sherwen et al., 2016a). In the atmosphere, I₂ and HOI
43 rapidly photolyse, producing reactive iodine radicals that catalytically destroy O₃ (Saiz-Lopez and Von Glasow, 2012; Simpson
44 et al., 2015). This cycle represents a negative feedback mechanism, whereby increased O₃ concentrations are offset by
45 increased iodine emissions (Carpenter et al., 2013; Prados-Roman et al., 2015). In fact, model calculations project that future
46 increases in anthropogenic O₃ will be buffered by iodine (Iglesias-Suarez et al., 2020), which has implications for future air
47 quality. Ice core records have shown that there has been a tripling of atmospheric iodine since 1950 (Cuevas et al., 2018;
48 Legrand et al., 2018), likely due to increased anthropogenic O₃ production. Recent measurements of iodine radicals in the
49 lower stratosphere (Koenig et al., 2020) have also demonstrated the potential for iodine to contribute to stratospheric ozone
50 depletion. Iodine has a far higher O₃ destruction potential than chlorine (Klobas et al., 2021) and has been implicated as a
51 potential reason for the unexplained decrease in extra-polar lower stratospheric O₃ (Koenig et al., 2020), which is a region
52 where O₃ changes exert strong radiative effects on climate (Neale et al., 2025).

53
54 Despite the significance of HOI and I₂ as the main emission source of atmospheric iodine, and of HOI as a sink for the iodine
55 oxide (IO) radical, there have been few measurements of these compounds. Measurements of I₂, which has been detected by
56 both Differential Optical Absorption Spectroscopy (DOAS) and by Chemical Ionisation Mass Spectrometry (CIMS), have
57 typically been restricted to coastal areas (Saiz-Lopez and Plane, 2004; Bitter et al., 2005; Peters et al., 2005; Mahajan et al.,
58 2009; Huang et al., 2010) with only one measurement in the remote ocean region (Lawler et al., 2014). There has so far been
59 only one directly quantified measurement of HOI, by CIMS, also in a coastal region rather than the open ocean (Tham et al.,
60 2021). There are several challenges that contribute to the scarcity of measurements of these molecules. The mixing ratios of
61 iodine compounds are low, in the sub-ppt to 10 ppt range over the ocean, due to their short atmospheric lifetimes (Saiz-Lopez
62 et al., 2012; Lawler et al., 2014). In addition, halogen compounds can undergo heterogeneous chemistry on the surfaces of
63 instrument inlets, resulting in loss or potentially addition of signal (Neuman et al., 2010; Liao et al., 2012; Liao et al., 2014;

64 Le Breton et al., 2017; Peng et al., 2022). Further, there is a lack of highly sensitive and selective measurement techniques and
65 instrumentation to measure and quantify these compounds. Given that reactive halogen chemistry is increasingly being
66 incorporated into global chemistry transport models, there is an urgent requirement for observations to help provide constraints
67 and improve model capabilities.

68
69 Chemical ionisation mass spectrometry is increasingly being used in the measurement of trace atmospheric gases due to its
70 high sensitivity and selectivity, high time-resolution, soft ionisation, and ability to directly measure compounds in the field
71 with minimal sample preparation (Huey, 2007; Lee et al., 2014; Zhang et al., 2023; Riva et al., 2024). Reagent ions are
72 generated and interact with sample molecules in an ion molecule reactor (IMR) to produce analyte ions, such as through charge
73 transfer or adduct formation reactions. Various reagent ions have been used to measure atmospheric halogen species such as
74 SF_5^- to measure HCl and ClONO_2 (Marcy et al., 2005), NO_3^- to measure HIO_3 (Sipilä et al., 2016; Finkenzeller et al., 2023),
75 and SF_6^- to measure bromine and chlorine compounds such as Br_2 , Cl_2 , BrCl , IBr , and BrO (Sjostedt and Abbatt, 2008; Buys
76 et al., 2013; Roscoe et al., 2014). More recently, I^- has been used for a wide range of chlorine and bromine atmospheric species
77 including Cl_2 , Br_2 , ClNO_2 , BrNO_2 , ClONO_2 , ClO , BrO , HCl , HOCl , HOBr , BrCl , and various chlorine-containing OVOCs
78 (Kercher et al., 2009; Lee et al., 2014; Custard et al., 2016; Le Breton et al., 2017; Lee et al., 2018; Priestley et al., 2018).
79 However, the I^- reagent ion is not effective in measuring trace levels of iodine species as they can be formed during the
80 generation of the reagent gas (Wang et al., 2021a). An alternative is to use Br^- as a reagent ion. This has previously been used
81 to measure Cl_2 , HO_2 , H_2SO_4 and HNO_3 (Lawler et al., 2011; Sanchez et al., 2016; Rissanen et al., 2019; Wang et al., 2020)
82 and more recently to detect various iodine species such as I_2 , HOI , ICl and IBr using a chemical ionisation atmospheric pressure
83 interface time-of-flight (CI-API-TOF) mass spectrometer (Tham et al., 2021; Wang et al., 2021a; He et al., 2023). A reduced
84 pressure Br-CIMS has also been developed which can produce analyte ions through an AIM (adduct ionisation mechanism)
85 IMR (Riva et al., 2024). The instrument operates at medium pressures (50–500 mbar) producing low electric fields ($E/N < 10$
86 Td or field-free conditions), which reduces the chances of fragmentation and promotes adduct formation as the primary method
87 of analyte formation. This loses some of the sensitivity compared to higher pressure CIMS instruments due to the reduced
88 numbers of collisions occurring in the IMR. However, this is compensated for with an increased linear range of detected
89 compounds through promotion of adduct formation.

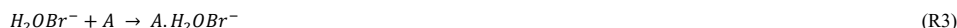
90
91 It has previously been shown that water can form an adduct with halide (I^- , Br^-) reagent ions (R1), which can subsequently act
92 as a reagent ion itself (R3) (Lee et al., 2014; Wang et al., 2021a). The formation enthalpy between a sample molecule and the
93 hydrated reagent ion is lower than with the dry reagent ion (He et al., 2023), and so increasing humidity should result in a
94 decrease in sensitivity, as the proportion of dry reagent ions decreases. However, for small molecules (<8 atoms), increasing
95 humidity has generally been shown to increase sensitivity (Lee et al., 2014; Iyer et al., 2016). This was explained by Quantum
96 Rice-Ramsperger-Kassel (QRRK) theory (Rice and Ramsperger, 1927; Kassel, 1928) which treats the halide adduct cluster as
97 a collection of identical harmonic oscillators (vibrational modes) at an identical the same frequency. On formation of the cluster,

98 the energy produced is distributed amongst the cluster vibrational modes. Not all vibrational modes are considered accessible
99 with experimental evidence suggesting the number of accessible vibrational modes is about half of the total number of
100 vibrational modes (Laidler, 1987). The cluster will fragment if energy above a critical threshold is localised in a single
101 oscillator vibrational mode. The rate at which this occurs for a cluster (the decay rate) is calculated by Eq. (1) (Kurtén et al.,
102 2010; Lee et al., 2014).

$$104 \quad k = \nu \frac{j!(j-m+s-1)!}{(j-m)!(j+s-1)!} \quad (1)$$

106 where k is the decay rate (s^{-1}), ν is the average vibrational frequency of the cluster (cm^{-1}), s is the number of accessible
107 vibrational modes, m is the quantised binding energy and j is the quantised total thermal energy of the cluster (binding energy
108 plus thermal energy of the individual molecules). m and j are quantised by dividing their calculated energies (cm^{-1}) by ν .

110 Adding a water molecule to the cluster increases the number of harmonic oscillators vibrational modes, which decreases the
111 probability that the critical energy threshold is reached in a single oscillator (Kurtén et al., 2010)(Kurtén et al., 2010), resulting
112 in a decrease in the rate of fragmentation decay rate of the newly formed clusters (R4 and R5). Additionally, a lower energy
113 fragmentation pathway is available for the hydrated adduct (R6), which kicks out the water molecule and keeps the sample
114 molecule and reagent ion together.



123 where A is the sample molecule.

124
125 This kinetic enhancement via the water molecule is in competition with the thermodynamic effect of lower formation enthalpy
126 and will affect different sample molecules to different degrees, requiring the humidity dependence of every molecule of interest
127 to be quantified. Additionally, the humidity dependence will vary between instruments due to differences in IMR conditions
128 and instrument tuning settings.

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130 In this work, the ability for the Br-CIMS to detect and quantify the iodine species I₂ and HOI is demonstrated, and the impact
131 of changing humidity on the instrument sensitivity to these molecules is quantified.

132 2 Methods

133 2.1 Bromide Time-of-Flight Chemical Ionisation Mass Spectrometry (CIMS)

134 The measurement of I₂ and HOI were performed using a Vocus S Chemical Ionisation Time-of-Flight (CI-TOF) Mass
135 Spectrometer (Toferwerk, Switzerland), a high resolution (R ≈ 5000) instrument that can produce Br⁻ reagent ions and Br-adduct
136 analyte ions through a Vocus Aim reactor. The operational principles of the instrument and the AIM reactor are described in
137 detail by Riva et al. (2024).

138
139 The Br⁻ reagent ions are generated by passing 0.25 SLPM ultra-high purity N₂ gas over a permeation tube held at 80 °C
140 containing >99% benzene, C₆H₆, and trace amounts of bromoethane, C₂H₅Br. This gas mixture is passed into a vacuum
141 ultraviolet (VUV) ion source where UV light is emitted from a Kr lamp at 116.487 nm and 123.584 nm which is absorbed by
142 the benzene, generating photoelectrons (R7) (Ji et al., 2020; Breitenlechner et al., 2022). These photoelectrons react with the
143 C₂H₅Br, generating Br⁻ ions (R8) (Riva et al., 2024).



147
148 The Br⁻ reagent ions are passed into an ion-molecule reactor (IMR) where ~~they are~~ joined by 1.8 SLPM of sample gas
149 which has been passed into the sampling inlet and through a 0.475 mm critical orifice. The IMR is temperature controlled at
150 50 °C and pressure controlled at 50 mbar using a vacuum pump (IDP3, Agilent Technologies). The Br⁻ reacts with sample
151 molecules to form adduct ions. These are drawn through another critical orifice and travel through four differentially pumped
152 chambers which remove neutral molecules and focus the ions into a narrow beam before entering the drift region of the time-
153 of-flight (ToF) chamber. Ions in the TOF chamber are extracted in discrete packets at a frequency of 18.02 kHz and converted
154 into mass spectra using an MCP detector with a preamplifier over a range of 7–510 Th. The extracted packets are averaged
155 over a period of 1 second which is also the speed of data collection. Data averaging, mass calibration, peak assignment, peak
156 fitting and peak integration were all performed using the software package Tofware (version 4.0.1, Aerodyne Research Inc.)
157 used in Igor Pro 9 software (Wavemetrics). This data was then exported and analysed further using the R language for statistical
158 computing (R Core Team).

2.2 Humidity generation

The calibrations performed in this work were made over a humidity range from near complete dryness to almost 100% RH in order to be able to correct for effects caused by variations in ambient humidity in field measurements. This was achieved by humidifying the N₂ gas entering the instrument to various degrees using an in-house dynamic liquid calibration unit (LCU). The operation of this liquid calibration system has been described elsewhere (Yeoman et al., 2022) but is briefly covered here. The LCU is comprised of a proportional liquid-gas mixing valve (Bronkhorst) which controls the mass flow of liquid measured by a mini-Coriolis flow meter (Bronkhorst). It introduces a mass flow controlled zero-air dilution gas to aerosolize and fully evaporate the liquid into a temperature-controlled mixing region. The liquid is pressurized without gas contact, using a custom-built pneumatic cylinder with wetted materials of glass and PTFE.

The gas output of the LCU was maintained at 3.5 SLPM and the water concentration ranged from 0–5 g h⁻¹. During the I₂ calibrations, an additional flow of N₂ gas was used to dilute the humidified flow further increase the gas output. The tubing between the LCU outlet and the additional flow was heated to prevent saturation and condensation of the water vapour.

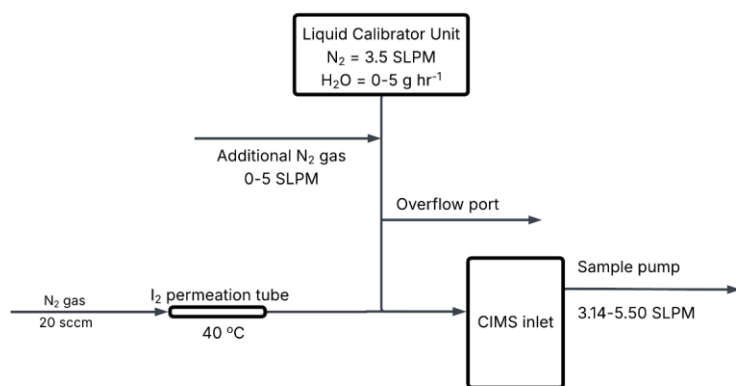
2.3 Calibration setup

2.3.1 I₂ calibration

An I₂ permeation tube was constructed by adding solid I₂ crystals (99.99%, Sigma-Aldrich) into a ¼" o.d. thin-walled PFA tube with the ends heat sealed closed. The permeation tube was held in a temperature-controlled permeation holder at 40 ± 0.1 °C. The temperature variation in the holder was minimal, at ± 0.1 °C, and is included in the uncertainty calculation. The permeation tube and was continuously swept by 20 sccm high-purity oxygen-free N₂ gas. The emission rate was calculated gravimetrically by measuring the mass loss of the permeation tube over a period of 6 months. Measurements were taken on average every 3 weeks with each data point representing the average of 6 sequential mass measurements, as shown in Fig. A1. several months and The emission rate was calculated from the gradient of Fig. A1 and corresponded to an emission rate of 34.7 ± 0.21 ng min⁻¹, with little variation seen during the measurement period.

A schematic of the I₂ calibration setup is shown in Fig. 1. Humidified N₂ gas from the LCU was used to dilute the permeation gas flow, and the CIMS internal sample pump was used to sample this diluted I₂ permeation gas. Before mixing with the permeation gas, the dilution gas was first split, with some diluting the I₂ permeation gas and some passing through an overflow port. The amount of dilution gas mixing with the permeation gas depends on how much flow the CIMS sample pump pulls towards the instrument. By changing this sample flow rate and hence the dilution of the permeation gas, the concentration of I₂ was altered, allowing for the construction of calibration curves. The quantity of dilution flow used for the calibrations was varied by altering the speed of the sample pump, with excess dilution gas from the LCU flowed through an overflow port. Using different pump speeds altered the concentration of I₂, allowing for calibration curves to be constructed at different

191 humidity levels. Concentrations of I₂ used-generated during the calibrations ranged from 1–3 x10¹⁰ molecules cm⁻³ (600–1000
 192 ppt at SATP).



193
 194 **Figure 1: Schematic of the I₂ calibration setup.**

196 **2.3.2 HOI calibration**

197 HOI was generated from the hydrolysis of I₂ (R9) in an experimental setup similar to that previously developed for the
 198 generation of HOBr in a study by Liao et al. (2012). 500 sccm of N₂ gas was passed over the same I₂ permeation tube from the
 199 I₂ calibration and mixed with 500 sccm N₂ gas which had been humidified by passing through a bubbler containing Milli-Q
 200 water. This was passed over wetted AgNO₃ crystals (99% Fluorochem) in a 9.5 cm long ½" o.d. PFA tube. A small aqueous
 201 layer forms on the AgNO₃ crystal surface, in which I₂ hydrolysis occurs (R9). The AgNO₃ reacts with the generated I⁻ and H⁺
 202 (R-10–12), pushing the (R9) equilibrium in favour of HOI production.

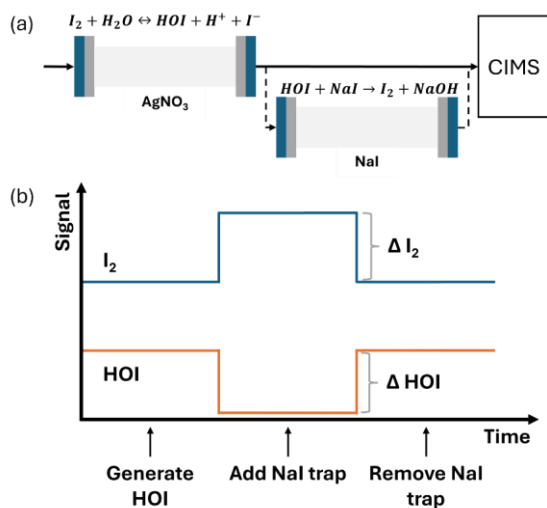


208
 209 The resulting I₂ and HOI gas mixture was diluted by humidified N₂ gas from the LCU before being drawn into the CIMS at a
 210 flow rate of ≈ 4.1 SLPM. The AgNO₃ reactor tube and instrument inlet were wrapped in heating tape and heated to 50 °C to

211 avoid HOI loss to the tube and inlet walls. The quantification of HOI is achieved by the inclusion of a trap containing sodium
212 iodide (NaI) crystals (99%, Sigma-Aldrich). When the trap is in place, HOI is converted back into I₂ (R13).



215
216 The sensitivity of HOI relative to I₂ was found by comparing the HOI signal loss and I₂ signal gain from R13. A 1:1 conversion
217 of HOI to I₂ was assumed, consistent with the approach of Liao et al. (2012) for the equivalent HOBBr–Br₂ system. Figure 42
218 shows the simplified schematic of the calibration setup and an idealised example of the relative change between I₂ and HOI.
219 As water is required in the HOI generation step, the calibrations are performed at high humidity. Lowering the humidity of the
220 N₂ gas should allow for lower humidity calibrations, but this will reduce the yield of HOI produced by (R9), making it more
221 difficult to observe the change in I₂ signal.



223
224 **Figure 24:** (a) A simplified schematic of the HOI calibration setup with reactions shown for HOI generation and destruction. (b) An
225 idealised example of the change in HOI and I₂ signal in the CIMS on addition and removal of the NaI trap.

226 2.4 Field measurements at Tudor Hill Marine Atmospheric Observatory, Bermuda

227 The CIMS instrument was deployed during the Bermuda boundary Layer Experiment on the Atmospheric Chemistry of
228 Halogens (BLEACH) field campaign in June 2022. The instrument was stationed at the Tudor Hill Marine Atmospheric
229 Observatory (THMAO) on the west coast of the island of Bermuda (32.26° N, 64.88° W). At this site, the instrument was

placed on top of a temporary sampling tower at 10 metres above ground level and 40 metres above sea level. The ion source gas was supplied by an air compressor connected to a nitrogen membrane and connected to the instrument by ¼" stainless steel tubing. The instrument was contained in an air-conditioned water-resistant enclosure and located at the top of the tower, minimising the potential for inlet effects to occur. The sample inlet was a short (15 cm, ½ in. outer diameter) length of PFA tube. A PTFE guard was placed at the entrance of the inlet tube, with an opening at the bottom, orientated 90° to the inlet. This allowed air to enter at a 90° bend and continue to the instrument while reducing the number of particles entering the CIMS as they impact onto the inlet walls. The sampling rate was 4.1 L min⁻¹, resulting in an inlet residence time of 0.59 s. The Reynolds number was 155, indicating laminar flow once the air was past an 8.5 cm entrance length.

2.5 Quantum chemical calculations

Quantum chemical calculations were used to find the vibrational frequency, binding energy and number of vibrational modes for the adducts I₂.Br⁻ and HOI.Br⁻. These were performed using Gaussian 16, Revision C.02 (Frisch et al., 2019), including all electrons in the correlation calculations through "MP2(Full)" and "CCSD(T,Full)". The geometries of the adducts were optimised at the MP2 level and then single-point energies were calculated, at these geometries, at the CCSD(T) level. All geometry optimisations were carried out under the "VeryTight" convergence criteria, and the optimised geometries were confirmed as local minima through subsequent frequency calculations. All calculations used the standard aug-cc-pVTZ basis set for all atoms except iodine, and the aug-cc-pVTZ-PP ECP basis from the Basis Set Exchange (Pritchard et al., 2019) for iodine. We refer to this combination as "aug-cc-pVTZ-PP". Binding enthalpies were obtained by adding together the energy difference E (complex) – E (reactants) at the CCSD(T) level to the corresponding difference between the thermal corrections to the enthalpy at the MP2 level (the thermal corrections to the enthalpy involve vibrational contributions, hence the need to have these at the level at which the geometries were optimised). The values are shown in Table 1.

Table 1: The calculated vibrational frequencies, binding energy, total thermal energy and vibrational modes of the I₂.Br⁻ and HOI.Br⁻ clusters. The decay rate is found using Eq. (1).

Cluster	Vibrational frequency / Hz	Binding energy / kcal mol ⁻¹	Total thermal energy / kcal mol ⁻¹	Vibrational modes (accessible)	Decay rate / s ⁻¹
I ₂ .Br ⁻	3.06 × 10 ¹²	30.8	33.5	4 (2)	2.66 × 10 ¹¹
HOI.Br ⁻	1.19 × 10 ¹³	24.3	27.1	6 (3)	2.38 × 10 ¹¹

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254 3. Results and Discussion

255 3.1 Detection of iodine compounds

256 I₂ and HOI were detected as adducts with Br⁻, with the two I₂.Br⁻ isotopes observed at m/Q 332.73 and 334.73 Th and the two
257 HOI.Br⁻ isotopes at 222.83 and 224.83 Th. The single peak mass spectra fits [for I₂](#) are shown in Fig. 23, [and in Fig. 4 for HOI](#),
258 taken from an HOI calibration experiment and of ambient air at Tudor Hill, Bermuda. At the instrument resolution of 5000,
259 the high mass defect of the I₂.Br⁻ adduct allows it to be clearly differentiated from other peaks ~~at 333 m/Q as seen in Fig. 2 d-~~
260 ~~f~~, even during ambient conditions at low mixing ratios. The HOI.Br⁻ adduct has a smaller mass defect, and subsequently has
261 more overlap with interferent peaks. During the HOI generation stage of the calibration (Fig. 42a), the HOI signal is
262 significantly greater than any interferent peaks. However, on addition of the NaI trap (Fig. 42b), or during ambient
263 measurements (Fig. 42ce), the HOI signal overlaps with other interferent peaks. Despite this, the resolution is sufficient for
264 Tofware's multi-peak fitting algorithm (Stark et al., 2015) to identify HOI even at very low mixing ratios. [The signal can be](#)
265 [further verified by examining the fit of the HOI isotope peak \(Fig 4. d-f\), calculated by the Tofware software.](#) Detection limits
266 during field conditions are discussed in [Sect.section](#) 3.7.
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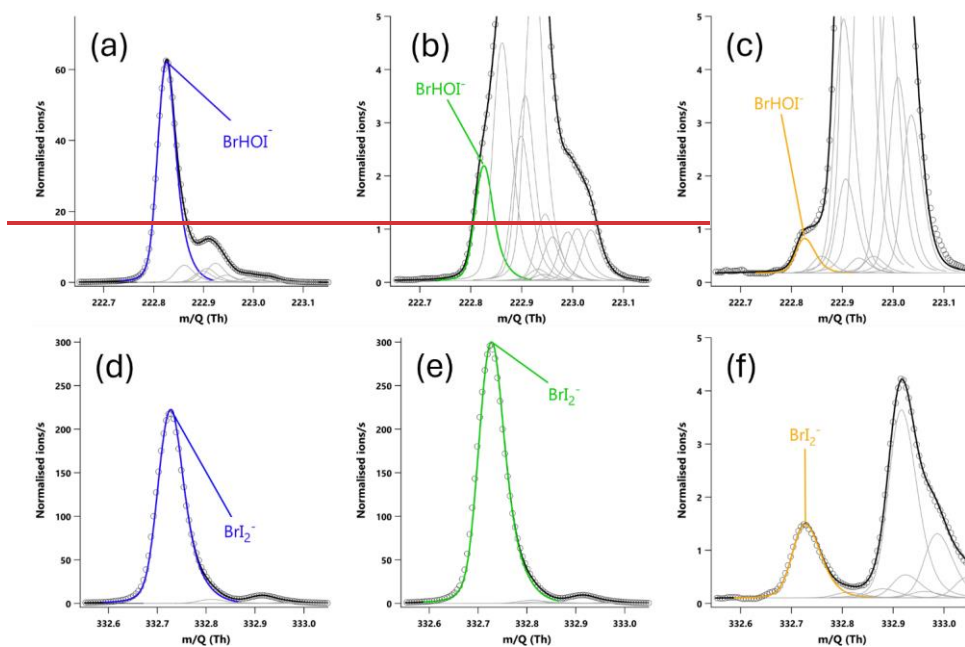
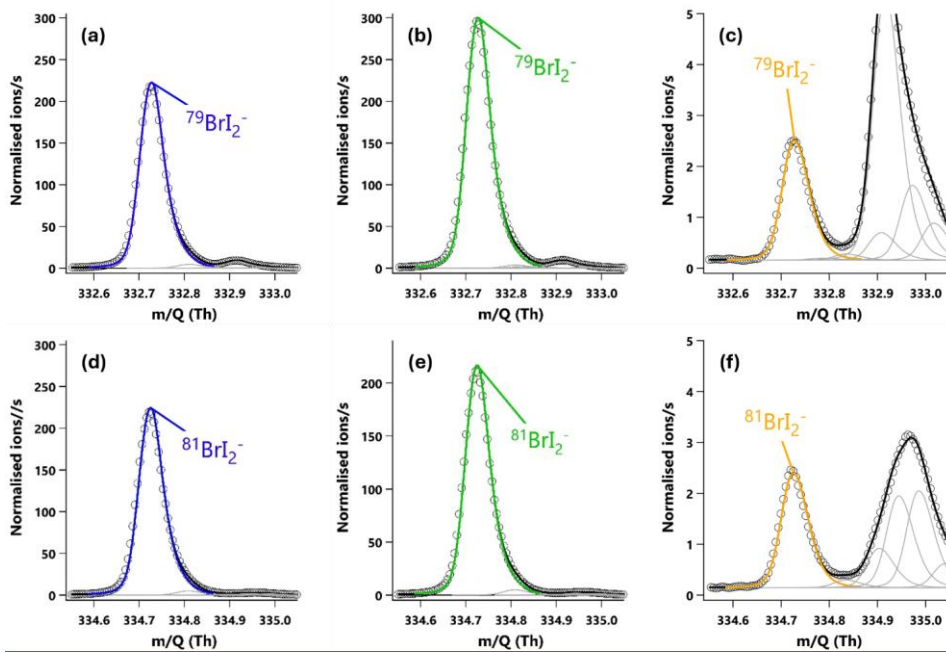


Figure 2: 20-minute averaged high-resolution single-peak mass spectra fits for HOI (a,b,c) and I_2 (d,e,f). The signal intensities correspond to mixing ratios of (a) 60.8 ppt, (b) 2.2 ppt, (c) 0.87 ppt, (d) 190.2 ppt, (e) 254.0 ppt and (f) 1.37 ppt. (a) The signal of HOI and (d) I_2 during an HOI calibration run without the NaI trap. (b) The signal of HOI and (e) I_2 during an HOI calibration run on addition of the NaI trap. (c) Ambient air measurements of HOI and (f) I_2 in a high humidity environment.

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275 **Figure 3:** 20-minute averaged high resolution single peak mass spectra fits for the molecular ion peak (a,b,c) and primary isotope
276 peak (d,e,f) for I_2 . The signal intensities correspond to mixing ratios of (a) 190.2 ppt, (b) 254.0 ppt and (c) 1.37 ppt. (a, d) The I_2
277 signal during an HOI calibration run without the NaI trap. (b, e) The I_2 signal during an HOI calibration run on addition of the NaI
trap. (c, f) Ambient air measurements of I_2 in a high humidity environment.

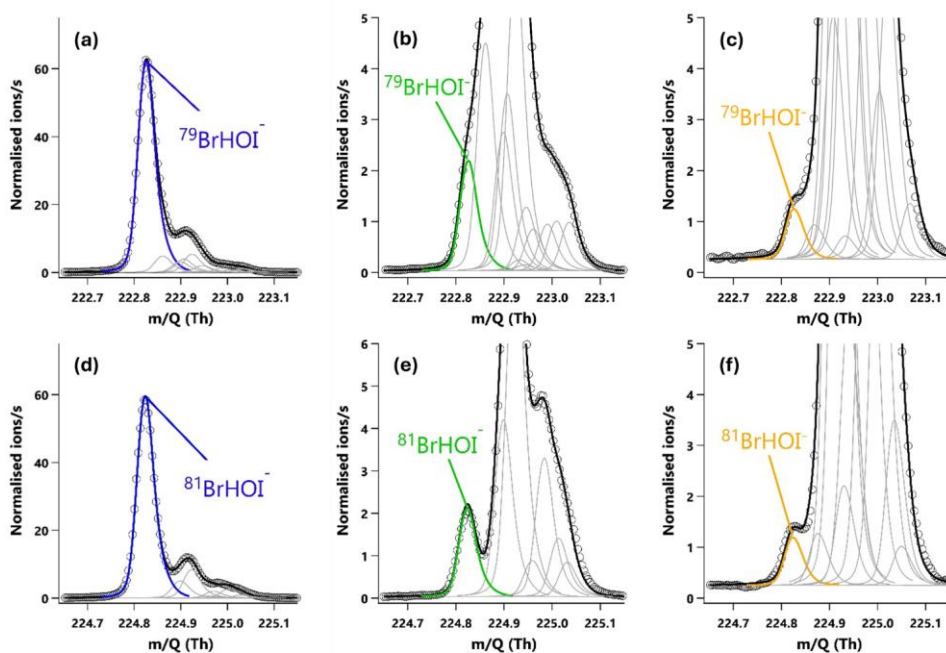


Figure 4: 20-minute averaged high resolution single peak mass spectra fits for the molecular ion peak (a,b,c) and primary isotope peak (d,e,f) for HOI. The signal intensities correspond to mixing ratios of (a) 60.8 ppt, (b) 2.2 ppt and (c) 0.87 ppt. (a, d) The HOI signal during an HOI calibration run without the NaI trap. (b, c) The HOI signal during an HOI calibration run on addition of the NaI trap. (c, f) Ambient air measurements of HOI in a high humidity environment.

3.2 Signal normalisation

The I_2 and HOI adducts are formed from a pseudo first order reaction between the sample molecules and the Br^-/H_2OBr^- reagent ions, and the adduct signal intensity is proportional to the reagent ions available. During experiments, the quantity of available reagent ions fluctuates over time. This can be due to variations in the efficiency of the ion source in generating the reagent ions, or with slight changes within the instrument. This variation can be corrected for by normalising the analyte signal against the sum of the reagent ions measured by the instrument, shown in Eq. (42). For CIMS instruments, the normalised signal is typically reported per million reagent ions, with units of normalised counts per second per million reagent ion counts per second (ncps). It has been reported that normalisation can compensate up to 50% reagent ion depletion from sample molecules (Riva et al., 2024).

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$$\text{Normalised signal (ncps)} = \frac{A \cdot \text{Br}^- \text{ (cps)}}{\text{Br}^- \text{ (cps)} + \text{H}_2\text{OBr}^- \text{ (cps)}} * 10^6 \quad (42)$$

where A is the sample molecule.

3.3 I₂ humidity dependence

The humidity of the instrument IMR, where the reagent ion collisions occur, is not directly measured but has been related to the sample relative humidity (Wang et al., 2021a; He et al., 2023) and/or vapour pressure of water (Lee et al., 2014; Lee et al., 2018). An alternative method is to use the ratio between the first reagent ion water cluster, H₂OBr⁻, and the dry reagent ion, Br⁻ (Dörich et al., 2021) (Dörich et al., 2021), Eq. (23). This has the advantage of not requiring further equipment to measure humidity levels during calibrations and accounts for any changes that may occur when the gas flows through the inlet. The so-called “water ratio” is used hereon to represent humidity. The water ratio is compared against absolute humidity measured during the BLEACH campaign in Fig. A4. Previous studies have shown that IMR temperature can impact the proportions of the hydrated and dry reagent ions, though this has been performed over a wide range of temperatures (Robinson et al., 2022). Fig. A4 shows that there was little variation in the IMR temperature (±0.6 °C) and pressure (±0.35 mbar) during the BLEACH campaign and that these changes did not noticeably affect the water ratio at different humidities.

$$\text{water ratio} = \frac{\text{H}_2\text{OBr}^- \text{ (cps)}}{\text{Br}^- \text{ (cps)}} \quad (23)$$

I₂ calibrations were performed at a range of different humidities generated by changing the water mass supplied by the LCU. The range of I₂ concentrations for each humidity was between 1–3 × 10¹⁰ molecules cm⁻³; about two orders of magnitude higher than what would typically be observed in the atmosphere. Ideally, the concentration range would be comparable to atmospheric levels, but limitations in the amount of gas dilution and water mass flow restricted the minimum concentrations that could be used. Low pressure CIMS instruments reportedly have large linear ranges, particularly when normalising to the reagent ion (Riva et al., 2024), and so it is assumed in this work that the calibration range is within the linear range of the instrument. The I₂ sensitivity was determined by calculating the gradient of the calibration curves with units of ncps cm³ molecules⁻¹. The instrument response over the calibration range demonstrated a high degree of linearity, an example of which is shown in Fig. 35.

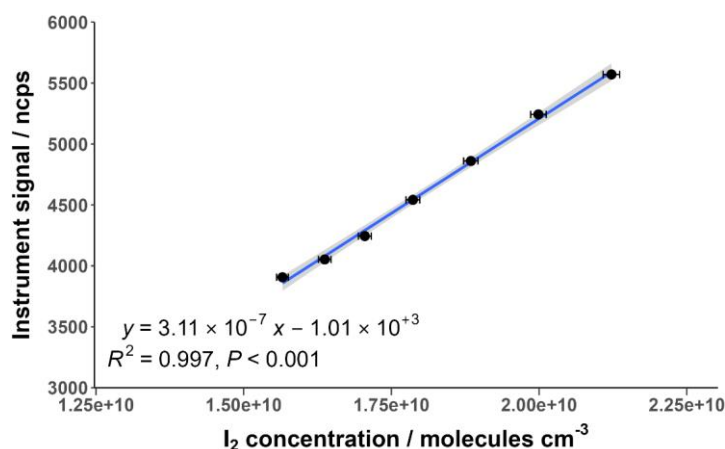
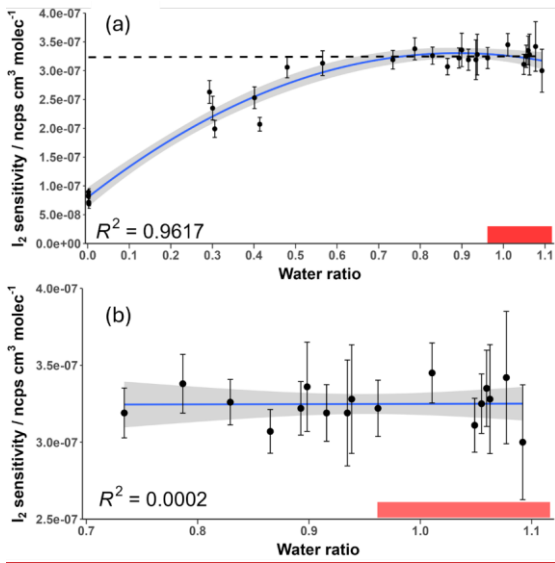


Figure 5: Normalised instrument signal against I₂ concentration for an example calibration at a water ratio of 1.05. The shaded area represents the 95% confidence interval of the fitted curve. Error bars indicate the calculated propagated uncertainty of the I₂ concentration, shown in [Sect. section 3.5](#).

The change in I₂ sensitivity at different humidities is demonstrated in Fig 4.6. Above a water ratio of ~0.7, the sensitivity of I₂ is effectively humidity-independent with a sensitivity of $3.25 \times 10^{-7} (\pm 6.30 \times 10^{-9})$ ncps cm³ molecules⁻¹. This is four times more sensitive than the average sensitivity of $7.92 \times 10^{-8} (\pm 3.18 \times 10^{-9})$ ncps cm³ molecules⁻¹ at near dryness (average water ratio = 0.0022), which can be attributed to the stabilising effect of the H₂O molecule on the formed adduct. As humidity is increased, at high humidities, this stabilising effect is balanced out by more adduct formation occurring via the the lower formation enthalpy of adducts formed with the BrH₂O⁻ reagent ion, which is a less exothermic reaction (Wang et al., 2021a). There is also an increase in the formation of the second water cluster, Br(H₂O)₂⁻, at very high humidities, shown in Fig 7. There appears to be no literature on whether this cluster also acts as a reagent ion for I₂. However, it is unlikely to act as one due to an additional collisional reaction needed for formation of Br(H₂O)₂⁻ and the likely even lower formation enthalpy between the second water cluster and a sample molecule. The second water cluster may have an indirect effect on the reduction of the increase in sensitivity. At very high humidities, the proportion of dry reagent ion continues to decrease with the first water cluster proportion remaining steady while the second water cluster increases. This suggests the formation rate of the first water cluster is being matched by its conversion to the second water cluster. This will still reduce the amount of adduct formation occurring through the more exothermic dry reagent ion pathway but will not produce an increase in sensitivity via water stabilisation. These factors potentially explain the emergence of the humidity-independent region above the 0.7 water ratio.

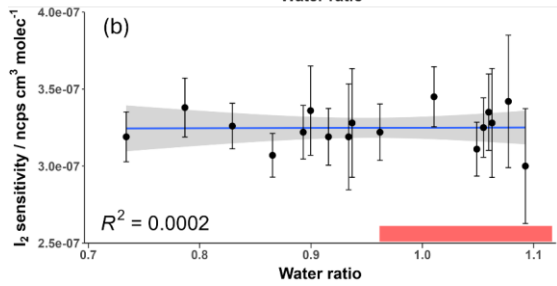
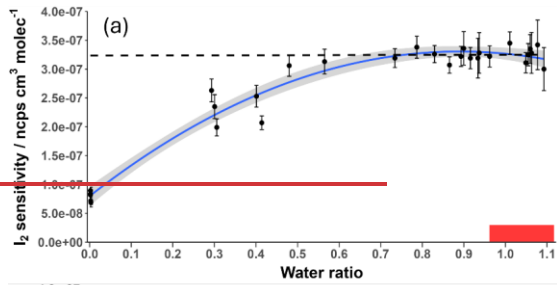
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341
 342 **Figure 6: Measured I₂ sensitivity at different humidity levels over (a) the full range and (b) the humidity-independent range. Error**
 343 **bars represent the calculated error for each sensitivity value described in Sect. 3.5. The shaded grey area represents the 95%**
 344 **confidence interval of the fitted curve. The shaded red region represents the ambient water ratio observed in the marine boundary**
 345 **layer at Tudor Hill, Bermuda. The dotted line in (a) is the average of the data points in (b).**

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Figure 4: Measured I_2 sensitivity at different humidity-levels over (a) the full range and (b) the humidity-independent range. Error bars represent the calculated error for each sensitivity value described in section 3.5. The shaded grey area represents the 95% confidence interval of the fitted curve. The shaded red region represents the ambient water ratio observed in the marine boundary layer at Tudor Hill, Bermuda. The dotted line in (a) is the average of the data points in (b).

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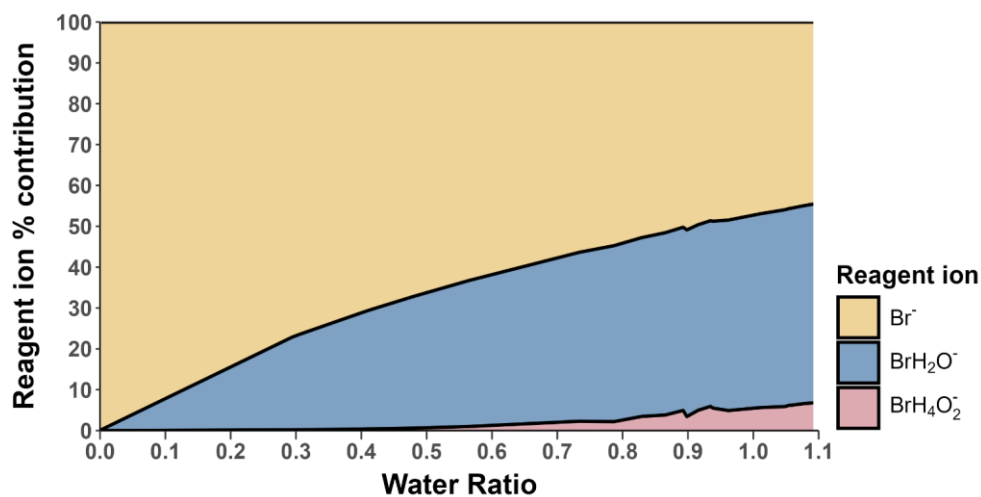


Figure 7: The percentage contribution of the Br⁻ ion (yellow), BrH₂O⁻ ion (blue) and BrH₄O₂⁻ ion (pink) at different water ratios. The values shown are the average contributions of these ions detected during the I₂ calibration runs.

3.4 HOI calibration and humidity dependence

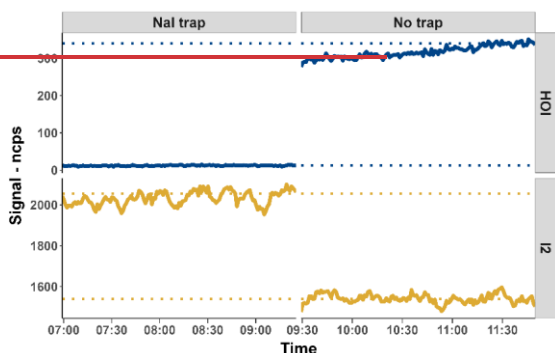
The relative sensitivity of HOI compared to I₂ (termed the HOI/I₂ ratio) was quantified by comparing the change in signal of HOI and I₂ with and without the NaI trap in place, with an example calibration shown in Fig. 58. With no trap in place, the ratio of HOI to I₂ signal was found to be around 1:5. On addition of the NaI trap, almost all of the HOI signal was removed with a corresponding increase in I₂ signal observed. For the equivalent HOBr–Br₂ system in the Liao et al. (2012) system, the conversion of HOBr to Br₂ was assumed to be 1:1. Here, the 1:1 HOI to I₂ conversion assumption was tested by measuring the signal of various other iodine-containing compounds, to determine whether any other significant reaction pathways were occurring during the production and destruction of HOI. The mass spectra of these other iodine compounds are shown without including the NaI trap in Fig. A24 and including without the trap in Fig. A32. Some of these compounds have substantial overlapping interfering peaks, and it is not possible at the instrument resolution to determine how real their signals are. Of these compounds, only HI and IBr had both distinct enough peaks and showed a small increase on an addition of the NaI trap, a small increase in signal was observed for HI and IBr. These quantities increase in HI and IBr signals were much smaller than for I₂, which represented 96% of the change in signal, as shown in Table A1. The production of HI is likely from the reaction of NaI and HNO₃, which is produced via (R12) in the HOI generation step, rather than from HOI. It is possible that the HI and IBr signal may be produced from HOI, for example if there was some contaminant NaBr also present in the trap.

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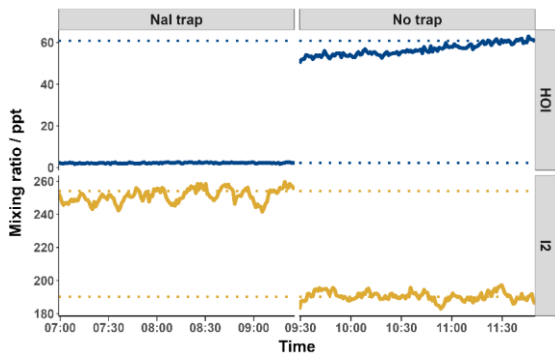
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371 However, conversely, any contaminant bromine compounds in the gas stream may react with the NaI trap, producing the IBr.
 372 However, the average standard error of the I₂ signal change is already higher than the increase in IBr signal, and so the
 373 production of IBr can be considered negligible against the other uncertainties of the calibration and the assumed 1:1 conversion
 374 of HOI to I₂ can be maintained. It is also possible that the presence of NaI is the cause of these compounds, such as any
 375 contaminant bromine compounds reacting with NaI to produce IBr. Either way, the intensity of the signals for HI and IBr can
 376 be considered negligible and therefore substantiate the assumption of 1:1 conversion of HOI to I₂.



378 Figure 5: The 1-minute averaged HOI and I₂ normalised signal with and without the inclusion of the NaI trap. Dotted lines indicate
 379 the average signal of the final 20 minutes of each run.



381 Figure 8: 1-minute averaged HOI and I₂ mixing ratios with and without the inclusion of the NaI trap. Dotted lines indicate the
 382 average mixing ratio of the final 20 minutes of each run. Values were calculated according to the calibration curve from Fig 9.

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385 The HOI calibration experiments were performed at different water ratios to determine how the HOI/I₂ ratio changes with
 386 humidity. As water was required to generate HOI in the first stage of the calibration, the experiments were only able to be
 387 performed at relatively high humidity, between water ratios of 0.7 and 1.1. This range corresponds to the humidity-independent
 388 region found for I₂. The humidity dependence of the HOI/I₂ ratio is shown in Fig. 69.

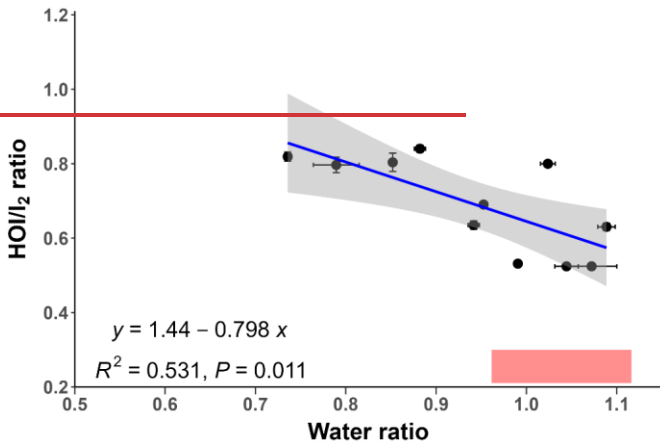


Figure 6: The relative sensitivity of HOI compared to I₂ (HOI/I₂ ratio) at different water ratio values. The shaded grey area represents the 95% confidence interval of the fitted regression line. The shaded red region represents the ambient water ratio observed in the marine boundary layer at Tudor Hill, Bermuda.

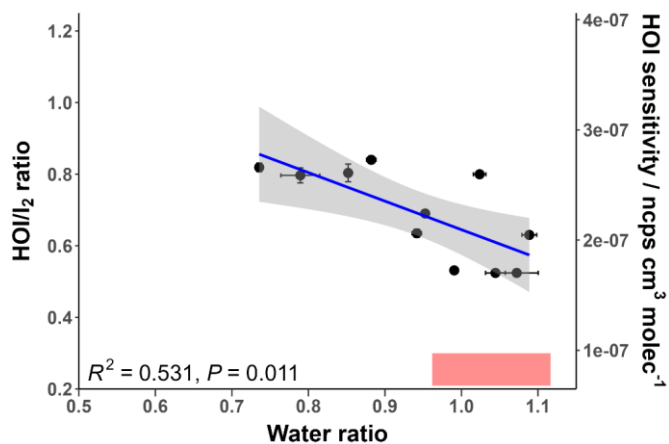


Figure 9: The relative sensitivity of HOI compared to I₂ (HOI/I₂ ratio) at different water ratio values. The regression line follows the equation $y = 1.44 - 0.798x$. The secondary y-axis shows the corresponding HOI sensitivity compared to the HOI/I₂ ratio. The shaded grey area represents the 95% confidence interval of the fitted regression line. The shaded red region represents the ambient water ratio observed in the marine boundary layer at Tudor Hill, Bermuda.

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Figure 6-9 shows that the HOI/I₂ ratio moderately correlates with increasing humidity; the correlation is statistically significant with a P-value below 0.05. As the humidity dependence of I₂ is independent over this measurement range, this decrease is attributed to a decreasing sensitivity for HOI. This can be explained from a QRRK theory standpoint. As seen in Table 1, the HOI.Br⁻ cluster has more accessible vibrational modes harmonic oscillators than I₂.Br⁻, so the increase of harmonic oscillators in vibrational modes from the presence of water has will have a smaller stabilising effect for HOI.Br⁻ than for I₂.Br⁻. Additionally, as the formation enthalpy for the HOI.Br⁻ cluster is lower than for I₂.Br⁻ (He et al., 2023), and so the effect of reducing the amount of dry cluster formation via the dry reagent ion has a more pronounced effect on the HOI.Br⁻ cluster.

3.5 Sensitivity measurement uncertainty

Propagation of error was used to calculate the uncertainty of the I₂ and HOI sensitivities. This can be calculated using the exact formula of propagation, assuming that variables are independent of each other, as shown in Eq. (34).

$$\sigma_x = \sqrt{\left(\frac{\partial x}{\partial a}\right)^2 \cdot \sigma_a^2 + \left(\frac{\partial x}{\partial b}\right)^2 \cdot \sigma_b^2 + \left(\frac{\partial x}{\partial c}\right)^2 \cdot \sigma_c^2 + \dots + \left(\frac{\partial x}{\partial n}\right)^2 \cdot \sigma_n^2} \quad (34)$$

412 where a, b, c, \dots, n are the variables of the function x , $\partial x/\partial a$ is the partial derivative of the variable with respect to x , and σ_a is
413 the error of the individual variable.

414 3.5.1 I₂

415 The I₂ concentration error was calculated from the propagation of the uncertainties of the emission of the I₂ permeation tube,
416 the temperature of the permeation tube holder, and the total gas flow from the sample pump to give a relative error of 0.7%.
417 The uncertainty of the sensitivity from a specific humidity calibration was calculated from the propagated uncertainty of the
418 I₂ concentration and the instrument signal for each point in the calibration curve, as shown in Fig. 53. The uncertainties of the
419 sensitivities in the humidity independent region can be averaged using Eq. (45) as these can be considered as repeats.

420

$$421 \sigma_{\text{humidity independent sensitivity}} = \frac{\sqrt{\sum_{i=1}^n (\sigma_i)^2}}{n} \quad (45)$$

422

423 where σ_i is the uncertainty of a particular sensitivity measurement and n is the number of measurements made in the humidity
424 independent region.

425

426 This resulted in a sensitivity uncertainty of 6.30×10^{-9} ncps cm³ molecules⁻¹ for the humidity-independent region, corresponding
427 to a relative error of 1.9%. This was repeated for the dry calibration sensitivities to produce an uncertainty of 3.18×10^{-9} ncps
428 cm³ molecules⁻¹ which represents a relative error of 4.0%. This uncertainty encompasses both calibration and humidity-
429 dependence uncertainties, but not background (zero) or inlet effects (see Sect. 3.6).

430 3.5.2 HOI

431 The sensitivity of HOI was determined relative to I₂ based on the linear model in Fig. 59. The uncertainty of the HOI sensitivity
432 at a specific water ratio was found from the uncertainty of the I₂ sensitivity at that water ratio propagated with the uncertainty
433 of the gradient and intercept of the linear model. This results in an uncertainty of 4.29×10^{-8} ncps cm³ molecules⁻¹ for the HOI
434 sensitivity, corresponding to an average relative error of 19.8% with a range of 16-25%, neglecting any background and inlet
435 effects.

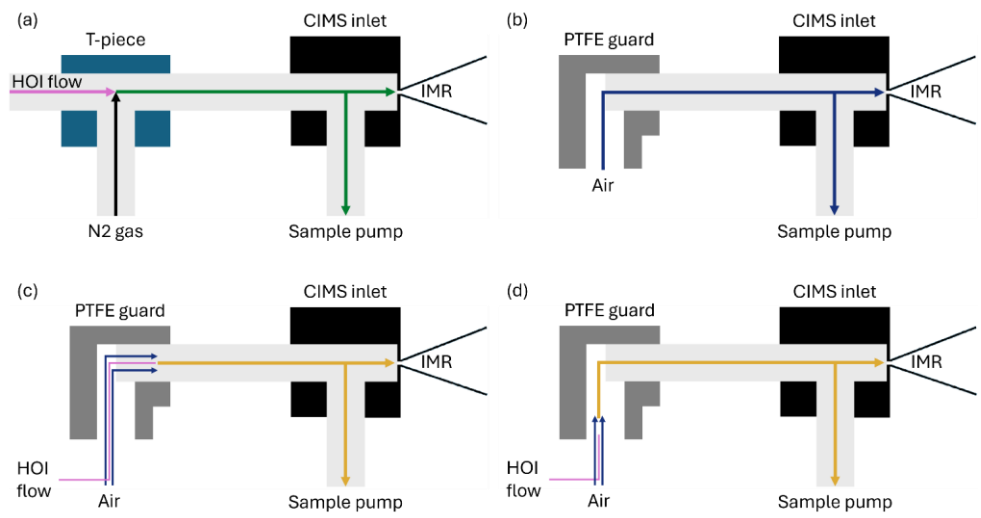
436 3.6 Inlet loss considerations

437 Measuring gas-phase compounds through an instrument inlet will introduce gas-wall interactions that can lead to loss of signal
438 or require conditioning to reach a steady state (Krechmer et al., 2016; Huang et al., 2018; Deming et al., 2019). Additionally,
439 halogen compounds can undergo heterogeneous chemistry on inlet walls, resulting in conversion to other halogen compounds.
440 This is particularly the case for the hypohalous acids and has been observed for HOBr and HOCl when measured by CIMS
441 instruments (Neuman et al., 2010; Liao et al., 2014; Le Breton et al., 2017; Peng et al., 2022). While all halogen compounds

442 experience these wall interactions, it is particularly pronounced for iodine compounds, leading to large losses and uncertainties
443 in measurements. Previous measurements of I₂ have found sample line losses between 18–40% (Shaw and Carpenter, 2013;
444 Carpenter et al., 2013). For HOI, the calibration performed by Tham et al. (2021) modelled the loss of HOI through their inlet
445 system via diffusion to the inlet walls which contributed to their total HOI uncertainty of $\pm 55\%$.

446
447 The effect of inlet losses is heavily influenced by the instrument setup. This means that any differences in inlet configuration
448 between calibrations and field measurements need to be accounted for. When deployed in the field, the inlet for this instrument
449 had a PTFE guard, allowing sample air to enter the inlet at a 90° angle, shown in Fig. 7b10b. The guard was implemented with
450 the intention that light gas-phase molecules could navigate around the bend without loss to the walls, but heavier aerosol
451 particles could not, reducing the potential for aerosol to block the entrance to the CIMS instrument. However, when calibrating,
452 the calibration gas was directed linearly into the sampling inlet, shown in Fig. 7a10a. The effect of this difference was
453 investigated.

454
455 Two inlet loss experiments were performed, one using a PFA T-piece to investigate the effect of physically having a bend in
456 place, and another with the PTFE guard that had been salted with sea water to mimic field conditions. A line of 1/8" PFA
457 tubing was attached to the output of the HOI calibration system and held in two configurations: one where it is pushed past the
458 bend as shown in Fig. 7e-10c and one where it is held before it, Fig. 7d10d. This required altering the dilution flow to laboratory
459 air to accommodate this configuration. The effect of the T-piece and PTFE guard on the normalised signal of HOI and I₂, along
460 with other iodine compounds that are potentially present, are shown in Table 42.



462
463
464
465

Figure 10: Simplified diagrams of the CIMS inlet setup during (a) the HOI calibration experiments, (b) sampling of air at Tudor Hill in Bermuda, (c) the line loss experiment bypassing the PTFE guard bend, (d) the line loss experiment including the bend from the PTFE guard.

Table 2: The normalised signal ± 1 standard deviation of various iodine species measured during the direct and bended inlet configurations for the T-piece and PTFE guard inlet loss experiments.

Species	T-piece			PTFE guard		
	Direct configuration $\pm 1\sigma$ / ncps	Bend configuration $\pm 1\sigma$ / ncps	Signal difference / ncps (% change)	Direct configuration $\pm 1\sigma$	Bend configuration $\pm 1\sigma$	Signal difference / ncps (% change)
HOI	381.45 \pm 5.85	132.79 \pm 6.38	-248.66 (-65%)	335.05 \pm 5.79	85.13 \pm 2.59	-249.92 (-75%)
I ₂	2576.62 \pm 25.92	2985.40 \pm 49.75	+408.78 (+16%)	2149.51 \pm 23.18	2472.60 \pm 25.23	+323.09 (+15%)
ICl	371.44 \pm 8.45	375.98 \pm 10.53	+4.54 (+1%)	400.73 \pm 8.62	531.58 \pm 15.02	+130.85 (+33%)
IBr	20.82 \pm 1.47	17.34 \pm 1.81	-3.48 (-17%)	20.32 \pm 1.51	185.06 \pm 4.57	+164.74 (+811%)
HI	0.16 \pm 0.42	0.20 \pm 0.44	+0.04 (+25%)	0.17 \pm 0.43	0.27 \pm 0.47	+0.10 (+59%)
IO	1.04 \pm 0.37	1.42 \pm 0.71	+0.38 (+37%)	1.23 \pm 0.35	1.02 \pm 0.50	-0.21 (-17%)
HIO ₂	27.67 \pm 3.77	30.36 \pm 9.74	+2.69 (+10%)	26.73 \pm 5.01	28.12 \pm 4.75	+1.39 (+5%)
INO ₂	2.30 \pm 0.73	1.00 \pm 0.58	-1.30 (-57%)	2.36 \pm 0.74	1.17 \pm 0.50	-1.19 (-50%)
HIO ₃	0.79 \pm 0.50	0.58 \pm 0.64	-0.21 (-27%)	1.02 \pm 0.56	0.93 \pm 0.63	-0.09 (-9%)
IONO ₂	6.75 \pm 1.39	2.51 \pm 1.28	-4.24 (-63%)	6.48 \pm 1.57	3.30 \pm 1.33	-3.18 (-49%)

469 For both the T-piece and the guard, the direct signal is comparable in intensity to the signal seen during the HOI calibrations,
470 with the exception of ICl, which is much higher than previously seen. When including the bend into the T-piece experiment,
471 little variation in signal is observed for ICl, HI, IO, HIO₂, and HIO₃, with changes smaller than the standard deviation of the
472 signal. There is a slight decrease in the IBr, INO₂, and IONO₂ signal, but the total quantity of loss from these compounds is
473 small compared to the total loss (<4%) The most noticeable loss is observed with HOI, along with an accompanying increase
474 in I₂. When converted into mixing ratios this corresponded to a 42.7 ppt (65%) HOI decrease and a 50.5 ppt (16%) increase in
475 I₂. This conversion of HOI to I₂ likely proceeds via the reverse iodine hydrolysis reaction (R14). The I⁻ required is most likely
476 from the HOI generation reaction in (R8), which becomes coated on the inlet walls. It is unknown how much I⁻ is present
477 during field measurements, and so the observed I₂ increase found during the inlet loss tests may be an overestimate compared
478 to atmospheric conditions.



481
482 With the salted PTFE guard, a higher proportion of HOI was lost (75% decrease) compared to the T-piece although the amount
483 lost was similar at 43.4 ppt. Meanwhile the proportion of I₂ gain remained about the same (15% increase) but the amount has
484 decreased to 40.0 ppt. There is a marked increase in IBr and ICl signal and the other iodine species remain at similar levels to
485 the T-piece run. Cl⁻ and Br⁻ are abundant in sea salt aerosol and are known to react with HOI (Vogt et al., 1999; Braban et al.,
486 2007; Tham et al., 2021), which is the likely reason for the increase in ICl and IBr and further loss of HOI compared to the T-
487 piece experiment. The sensitivities for ICl and IBr have not been measured for Br-CIMS instruments. However, it has been
488 suggested that sensitivities should be similar to that of I₂ (Wang et al., 2021a). If it is assumed that ICl and IBr have the same
489 response to humidity as I₂, the signal change would correspond to a 16.9 and 20.2 ppt increase, respectively. This is far higher
490 than would be expected from the additional HOI loss and may suggest there are additional pathways present to produce ICl
491 and IBr. ~~This again could be from the I⁻ present from the HOI production step though this likely requires oxidation before
492 reacting with Cl⁻ and Br⁻.~~

493 3.7 Application to atmospheric HOI and I₂ data.

494 I₂ and HOI were measured by the Br-CIMS during the BLEACH campaign in June 2022 with the inlet configuration shown
495 in Fig. 6b10b. Background signals were measured by flowing dry nitrogen (N₂) through a zero port in the CIMS inlet and
496 through the instrument critical orifice to the IMR, with the rest of the sampled air directed through the sample pump. These
497 zero measurements were repeated hourly, and the signal was linearly interpolated between measurements, providing a value
498 for the background signal during sampling periods. The limit of detection (LoD) for I₂ and HOI were calculated from an
499 extended zeroing period during the campaign as 3 standard deviations of the Allen variance of the zeroing period, similarly to
500 that described in Riva et al. (2024). These corresponded to detection limits of 0.14 ppt for I₂ and 0.27 ppt for HOI.

501

502 The HOI calibration could not be performed at dryness, making zero subtraction difficult to quantify. The HOI/I₂ ratio from
 503 Fig. 9 could be extrapolated to dryness at the y-intercept which would result in a dry HOI sensitivity that is 1.44 times higher
 504 than I₂. However, the 95% confidence interval of the intercept is ±0.43, suggesting a low precision when extrapolating from
 505 this calibration. An alternative method is to use the quantum chemical calculations to compare the decay rate between the
 506 HOI.Br and I₂.Br clusters. Using the values from Table 1, the decay rate of HOI.Br is slightly slower than I₂.Br, which would
 507 suggest a slightly higher sensitivity at dryness. However, this value is still similar to the decay rate of I₂.Br, and so the dry
 508 sensitivity of I₂ could be used as a conservative estimate of the dry HOI sensitivity.

509

510 ~~The HOI calibration could not be performed at dryness, making zero subtraction difficult to quantify. It was expected also~~
 511 ~~possible that any signal observed in the zero for HOI and I₂ was due to instrument noise rather than any interferent~~
 512 ~~HOI compounds. This was tested by comparing the relative signal intensities of two major isotope peaks of the I₂.Br and~~
 513 ~~HOI.Br adducts. The theoretical ratios between I₂.⁷⁹Br and I₂.⁸¹Br and HOI.⁷⁹Br and HOI.⁸¹Br are 1:0.975 and 1:0.977,~~
 514 ~~respectively, and so the zero data was tested to see whether the ratio between the two isotope peaks were within 10, 20 or~~
 515 ~~30% of this ratio or whether it fell outside these limits. These values are shown in Table 2. In total, for I₂, 72% of the zero~~
 516 ~~data fell within the 30% ratio limit, suggesting that the signal is generally real and needs to be accounted for by zero subtraction.~~
 517 ~~For HOI, 76.74% of the zero data had isotope ratios that were greater than the 30% tolerance, indicating that the signal generally~~
 518 ~~may not be real. Both the sample and zero timeseries for HOI and I₂ are shown in Fig. A5. The sample timeseries, without zero~~
 519 ~~subtraction, can be considered as an upper limit of the HOI and I₂ mixing ratios, and so it was decided to not background~~
 520 ~~subtract the HOI data. Therefore, the HOI data shown should be considered as an upper limit.~~

521

522 **Table 2: The instrument zero data points during the BLEACH campaign that fell within various limits of the theoretical isotope**
 523 **ratio between the HOI.⁷⁹Br and HOI.⁸¹Br adduct signals. A ratio value that was greater than 30% of the theoretical value was**
 524 **considered to be outside the ratio tolerance.**

	Total data points	Ratio limits			
		<10%	<20%	<30%	>30%
Zeroing data	242	46	33	58	184
	(100%)	(6.6%)	(13.6%)	(24.0%)	(76.0%)

525 **Table 3: The instrument zero data points during the BLEACH campaign that fell within various limits of the theoretical isotope**
 526 **ratio between the I₂.Br and HOI.Br adduct signals. A ratio value that was greater than 30% of the theoretical value was considered**
 527 **to be outside the ratio tolerance.**

	Total data points	Ratio limits			
		<10%	<20%	<30%	>30%
I ₂ zeroing data	412	129	232	297	115
	(100%)	(31.3%)	(56.3%)	(72.1%)	(27.9%)

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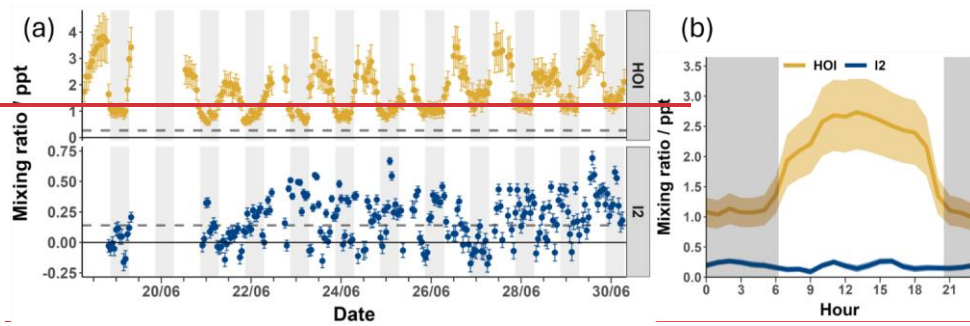
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<u>HOI zeroing data</u>	<u>412</u>	<u>34</u>	<u>62</u>	<u>108</u>	<u>304</u>
	(100%)	(8.3%)	(15.0%)	(26.2%)	(73.8%)

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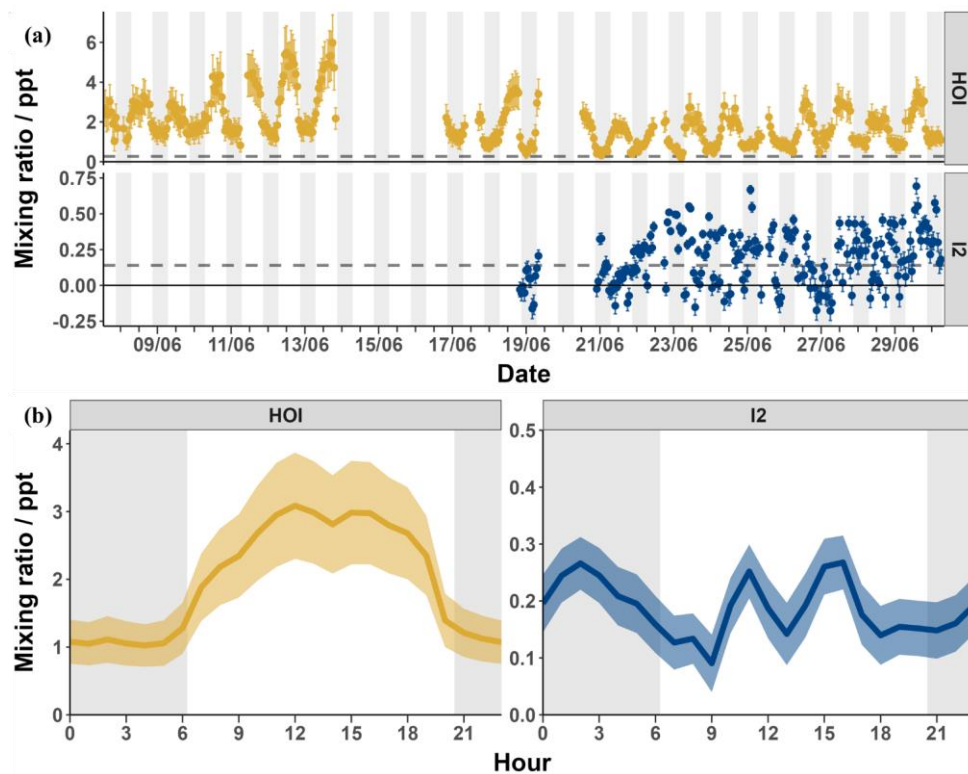
528
529 Measured signals were converted into mixing ratios using the calibrations from Sect. 3.3 and 3.4 and applying the loss
530 corrections described in Sect. 3.5. The I_2 data was then zero subtracted using the linearly interpolated instrument zeros. Figure
531 [8-11](#) shows the loss-corrected timeseries and diurnal cycles for I_2 and HOI. After zero subtractions and loss corrections, I_2 was
532 detected between 0–0.7 ppt with some mixing ratios dropping to negative values. HOI ranged from 0.52–3.96.0 ppt after its
533 [zero subtractions and](#) loss corrections. There was little pattern in the diurnal cycle for I_2 with the average signal being between
534 0.1–0.3 ppt during both the day and night. The observed mixing ratios of I_2 are similar to previous open ocean measurements
535 at the Cape Verde Atmospheric Observatory of <0.02–0.6 and <0.03–1.67 ppt in May 2007 and 2009 respectively (Lawler et
536 al., 2014). These previous measurements observed a diurnal cycle for I_2 peaking at night, as expected due to its rapid photolysis.
537 A diurnal cycle was not observed in our data, indicating either inlet or background effects which were not adequately quantified
538 by our experiments or an unknown [or poorly quantified](#) daytime source of I_2 . [Potential daytime sources include heterogeneous](#)
539 [reactions of photochemically produced oxidants such as HOI with iodide on the surface of aerosols \(Vogt et al., 1999; Moon](#)
540 [et al., 2026\) or on the sea surface \(Pound et al., 2024\), photochemical oxidation of iodide \(Raso et al., 2017\) or photochemical](#)
541 [reduction of iodate \(Reza et al., 2024\). However, these would have to be occurring close to the measurement site due to the](#)
542 [fast photolytic lifetime of \$I_2\$. Additionally, the release of \$I_2\$ from aerosol is also a likely contributor to the inlet effects, due to](#)
543 [buildup of aerosol on the inlet. With the inlet setup used in this work, these processes are indistinguishable.](#) The overall
544 uncertainty of the I_2 measurements made during the BLEACH campaign, as seen in Sect. 3.7.1, and the fact that they were
545 near to the detection limit of 0.14 ppt means that caution should be applied in interpreting the data.

546
547 [In contrast, aA](#) regular diurnal pattern was observed for HOI, with the signal consistently around 1 ppt during the night and
548 increasing during the day to an average peak of around 2.753 ppt. There are no reported mixing ratios of HOI in the open
549 ocean to compare to these values. The closest comparison is to the measurement of HOI at the coastal site of Mace Head (Tham
550 et al., 2021). There, a diurnal cycle was also observed, with low nighttime mixing ratios and an increase during the day.
551 However, the amounts seen at Mace Head during the day were much higher (up to 66.6 ppt), likely due to photochemical
552 reactions caused by I_2 emission by macroalgae (Tham et al., 2021).



553 -
 554 Figure 8: The timeseries (a) and diurnal cycle (b) of the mixing ratios of I₂ (blue) and HOI (yellow) from the BLEACH
 555 campaign in June 2022 after zero subtraction (for I₂) and loss correction (for both). The dashed line indicates the limit of

556 detection for each compound. The average relative uncertainty of I₂ was ± 29.6% and for HOI was ± 26.2%



557
558 **Figure 11: The timeseries (a) and diurnal cycle (b) of the mixing ratios of I₂ (blue) and HOI (yellow) from the BLEACH campaign**
559 **in June 2022 after zero subtraction and loss correction. The dashed line indicates the limit of detection for each compound. The**
560 **average relative uncertainty of I₂ was ± 28.7% and for HOI was ± 27.7%.**

561 3.7.1 Measurement uncertainties of atmospheric data

562 The uncertainty of the I₂ and HOI mixing ratios were again calculated by propagation of uncertainty. For I₂, this was
563 calculated from the uncertainty of the humidity independent sensitivity, twice the uncertainty of the dry sensitivity from the
564 linear interpolation calculation, and the uncertainty of the instrument signal during the inlet loss experiment. This resulted in
565 an average relative error throughout the BLEACH campaign of 29.628.7% for I₂ and 27.7% for HOI. There are a number of
566 other uncertainties from the inlet loss experiment that cannot be quantified and so the uncertainty for I₂ and HOI should be

567 considered a lower limit. In future work, repeated experiments may be able to account for these uncertainties by finding a
568 standard deviation of the loss correction value.

~~569 The uncertainty of HOI was calculated with just the uncertainty from the humidity dependent sensitivity and the uncertainty
570 of the instrument signal during the inlet loss experiment, resulting in an average relative error of 26.2% throughout the
571 BLEACH campaign. The HOI uncertainty value should also be considered a lower limit as there are the other unquantified
572 uncertainties of the inlet loss experiment and a lack of a zero subtraction.~~

573 4. Summary and conclusions

574 There have been few atmospheric measurements of HOI and I₂ due to the requirement of highly sensitive and selective
575 instrumentation and the lack of readily available calibration methods for HOI. In this study, we demonstrate a novel method
576 for the generation and calibration of HOI at ppt levels, utilising its interconversion to I₂ via chemical traps. The presence of
577 potential interferent iodine compounds was found to be negligible and the calibration was shown to be repeatable over a range
578 of humidities. The developed calibration method was utilised to ascertain the humidity dependence of the CIMS for HOI and
579 I₂. It was found that at humidities typical of the marine boundary layer, I₂ exhibited a humidity-independent sensitivity whereas
580 HOI showed a slight negative dependency. A possible explanation, supported by QRRK theory, is that higher humidities
581 increase the proportion of the Br(H₂O)⁺ reagent ion, which has a lower adduct formation enthalpy with HOI and I₂ compared
582 to the Br⁺ reagent ion, resulting in decreased sensitivities. However, the presence of H₂O increases the available ~~harmonic~~
583 ~~oscillators~~ vibrational modes of the adduct, improving the energy distribution and stabilisation of the adduct and increasing its
584 sensitivity. This effect is greater for I₂.Br⁺ as it has fewer harmonic oscillators than HOI.Br⁺ and these two opposing factors
585 mean that the sensitivity of I₂.Br⁺ is humidity independent. However, this stabilisation is not sufficient in the case of the HOI.Br⁺
586 adduct to mitigate the increasing proportion of Br(H₂O)⁺ vs Br⁺ reagent ions, resulting in a decreased sensitivity at high
587 humidities.

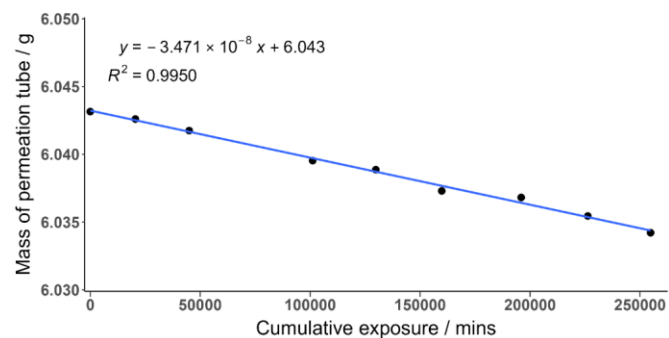
588
589 Heterogeneous chemistry and inlet wall losses are a perennial problem in the measurement of acidic and reactive species,
590 which can be exacerbated depending on the inlet configuration. When using the inlet configuration made for field
591 measurements, up to 75% of the HOI signal was lost compared to the calibration setup. This was accompanied by a
592 corresponding increase in I₂ signal, suggesting that this loss occurred via the reverse hydrolysis of iodine.

593
594 After correcting for humidity sensitivities and inlet losses, we detected I₂ and HOI in the marine boundary layer during the
595 BLEACH campaign at Tudor Hill, Bermuda in June 2022 at mixing ratios of between 0–0.7 ppt for I₂ and ~~an upper limit of~~
596 ~~0.52–6.03–8~~ ppt for HOI. The overall uncertainty of these measurements was calculated as $\pm 29.628.7\%$ for I₂ and $\pm 26.327.7\%$
597 for HOI. These uncertainties can be considered as lower limits as there were additional uncertainties from the inlet loss

598 experiment that were not quantifiable. ~~Additionally, zero subtraction could not be performed for the HOI data which could~~
599 ~~further increase the relative error.~~

600
601 In future work, the accuracy for I₂ and HOI could be improved by performing background measurements at similar humidity
602 levels to the ambient measurements, reducing the need to account for large differences in sensitivities from different humidities.
603 Additionally, further work on the HOI calibration can further constrain the uncertainty of the humidity dependent HOI/I₂ ratio
604 over a wider range of humidities. Finally, development of atmospheric detection of HOI and I₂ should focus on using
605 instrument inlets that can minimise inlet effects for these compounds, reducing the need to perform correction tests. Any
606 correction tests that are performed should be done before, during, and after field measurement collection to best characterise
607 any inlet artifacts and changes during the measurements.

608 Appendix



610
611 Figure A1: The mass loss of an I₂ permeation tube over time. The gradient represents the emission rate of the permeation tube. Each
612 data point is the average of 6 measurements.

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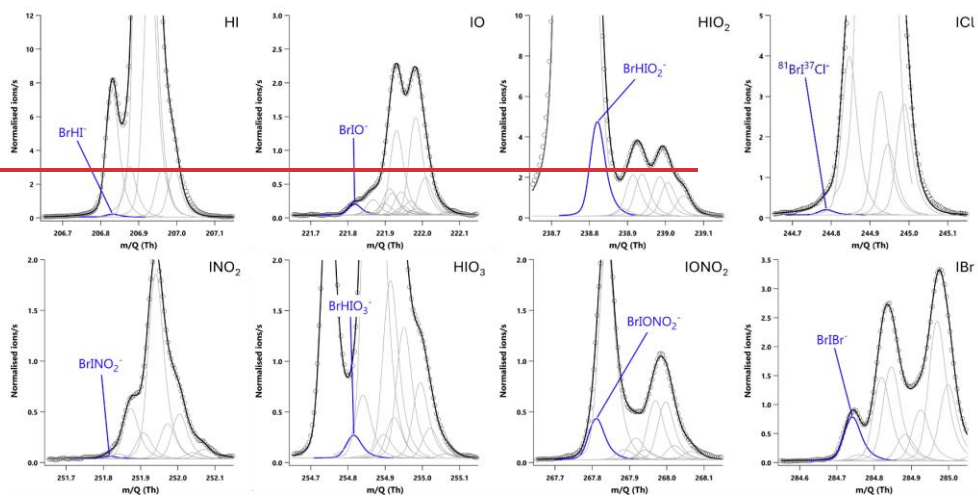
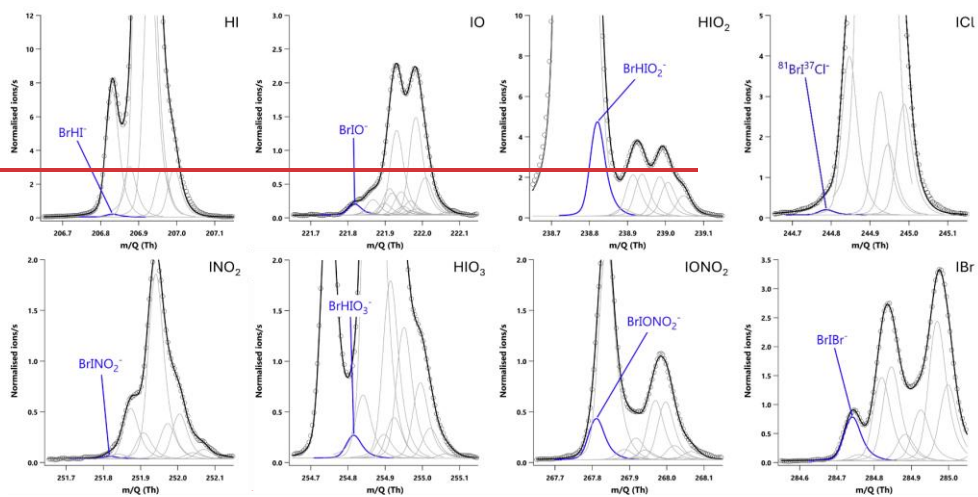


Figure A1: 20-minute averaged high-resolution single-peak mass spectra fits for potential iodine-containing compounds during the HOI-calibrations with the NaI-trap in place.



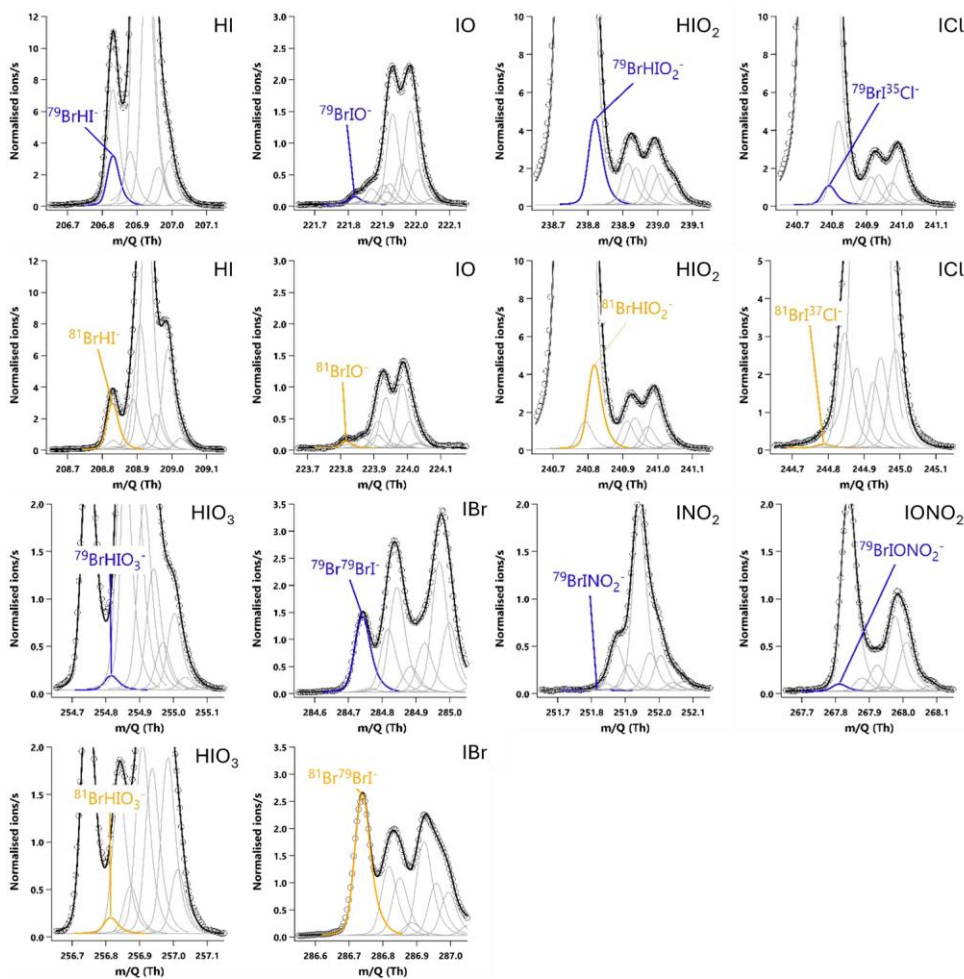
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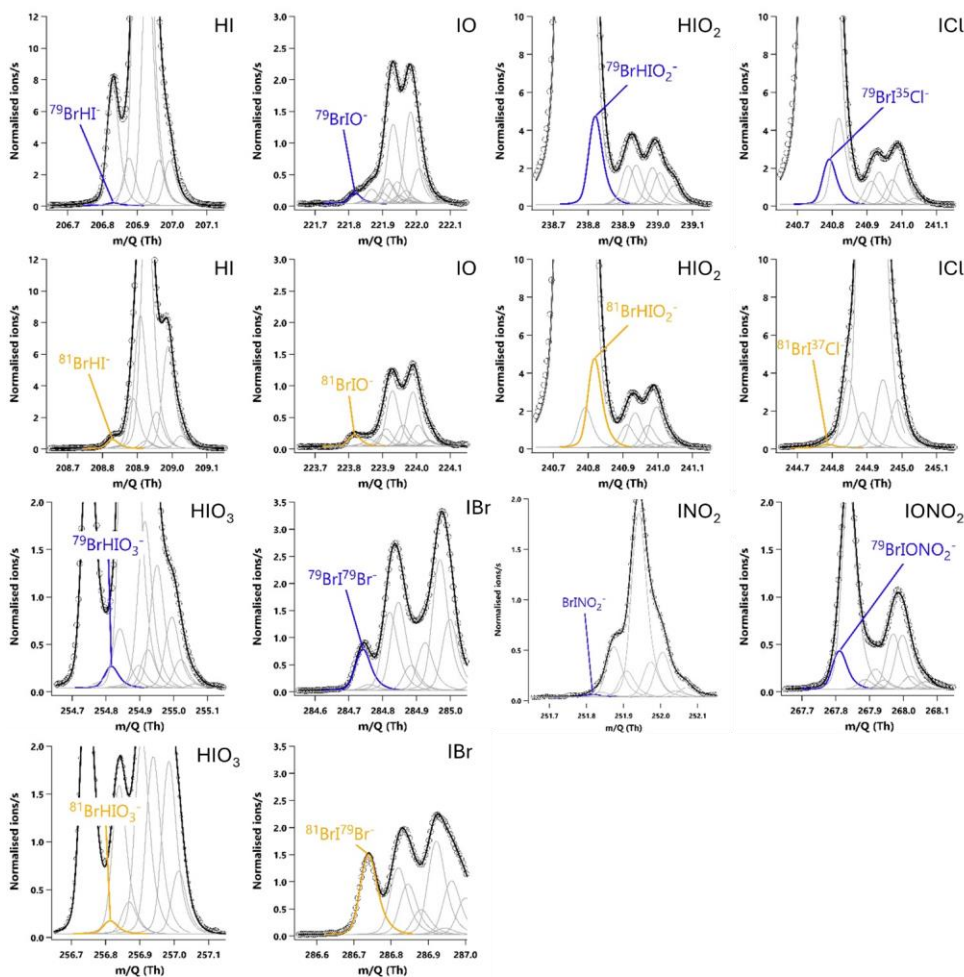
Figure A2: 20-minute averaged high-resolution single-peak mass spectra fits for potential iodine-containing compounds during the HOI calibrations after the removal of the NaI trap.

622



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623
624 **Figure A2: 20-minute averaged high resolution single peak mass spectra fits for potential iodine containing compounds during the**
625 **HOI calibrations with the NaI trap in place. The molecular ion peak is shown in blue with the primary isotope peak in gold with an**
626 **exception for $^{81}\text{Br}^{37}\text{Cl}^-$ as this was the isotope used to calculate the BrICl signal. Isotope peak shapes were determined using the**
627 **Tofware software. The isotope peaks of BrINO_2^- and BrIONO_2^- were not able to be visualised by the software.**



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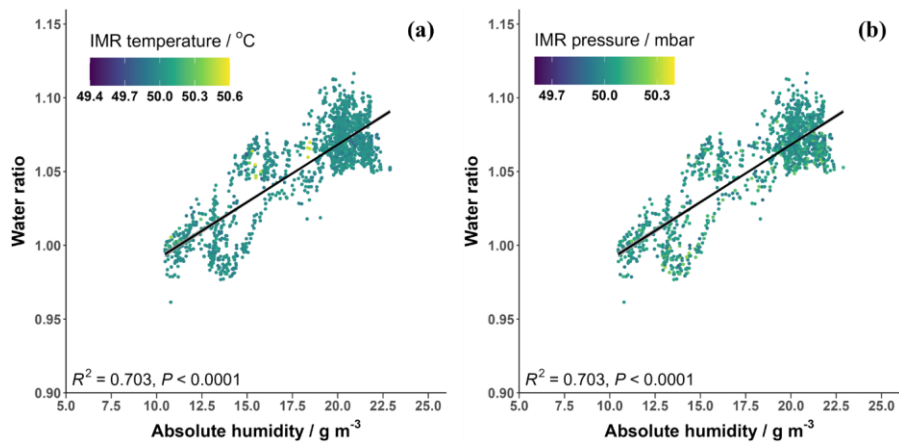
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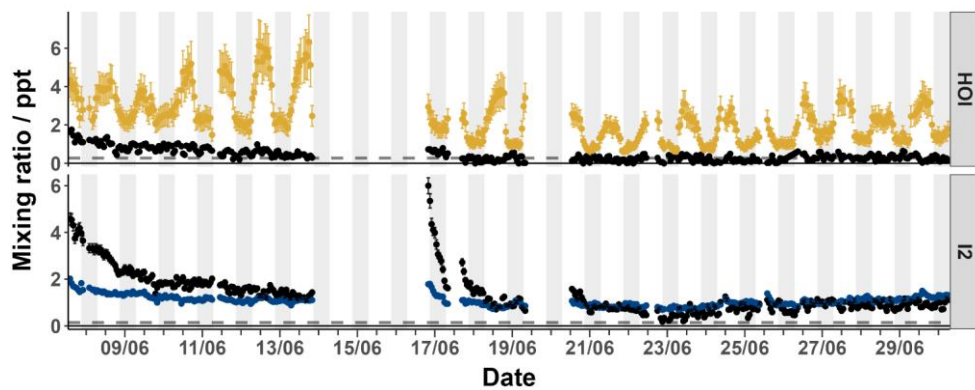
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Figure A3: 20-minute averaged high resolution single peak mass spectra fits for potential iodine containing compounds during the HOI calibrations after the removal of the NaI trap. The molecular ion peak is shown in blue with the primary isotope peak in gold with an exception for $^{81}\text{Br}^{37}\text{Cl}^-$ as this was the isotope used to calculate the BrICl signal. Isotope peak shapes were determined using the Tofware software. The isotope peaks of BrINO_2^- and BrIONO_2^- were not able to be visualised by the software.



634
635 **Figure A4: The water ratio in the CIMS IMR plotted against the absolute humidity measured during the BLEACH campaign**
636 **coloured by (a) the variation in IMR temperature and (b) the variation in IMR pressure.**



638
639 **Figure A5: The timeseries of the sample mixing ratios of HOI (yellow) and I₂ (blue) along with their corresponding zero mixing**
640 **ratios (black) from the BLEACH campaign in June 2022. The dashed line indicates the limit of detection for each compound.**

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Table A1: The signal change (in normalised ion counts per second) of HNO₃, HI, IBr and I₂ along with their calculated standard errors during the HOI calibration runs upon addition of the NaI trap.

Run number	Signal change (\pm standard error) on addition of NaI trap / neps			
	HNO ₃	HI	IBr	I ₂
<u>1</u>	<u>-4902.1 (\pm 71.5)</u>	<u>36.49 (\pm 0.93)</u>	<u>6.84 (\pm 0.51)</u>	<u>430.04 (\pm 10.89)</u>
<u>2</u>	<u>-5274.0 (\pm 164.4)</u>	<u>20.25 (\pm 0.80)</u>	<u>5.07 (\pm 0.46)</u>	<u>451.62 (\pm 13.60)</u>
<u>3</u>	<u>-4543.7 (\pm 43.7)</u>	<u>11.96 (\pm 0.69)</u>	<u>4.73 (\pm 0.40)</u>	<u>519.53 (\pm 5.76)</u>
<u>4</u>	<u>-4129.3 (\pm 58.4)</u>	<u>10.72 (\pm 1.02)</u>	<u>5.82 (\pm 0.49)</u>	<u>573.95 (\pm 6.90)</u>
<u>5</u>	<u>-4051.3 (\pm 116.0)</u>	<u>13.98 (\pm 1.41)</u>	<u>7.24 (\pm 0.51)</u>	<u>517.84 (\pm 8.79)</u>
<u>6</u>	<u>-4865.5 (\pm 48.7)</u>	<u>6.77 (\pm 0.95)</u>	<u>6.98 (\pm 0.53)</u>	<u>662.69 (\pm 9.09)</u>
<u>7</u>	<u>-2812.2 (\pm 43.3)</u>	<u>6.57 (\pm 0.82)</u>	<u>10.66 (\pm 0.52)</u>	<u>424.65 (\pm 6.39)</u>
<u>8</u>	<u>-3742.2 (\pm 64.4)</u>	<u>24.76 (\pm 1.09)</u>	<u>6.89 (\pm 0.49)</u>	<u>391.72 (\pm 5.24)</u>
<u>9</u>	<u>-2413.3 (\pm 63.5)</u>	<u>6.67 (\pm 0.55)</u>	<u>11.63 (\pm 0.55)</u>	<u>553.02 (\pm 7.72)</u>
<u>10</u>	<u>-4725.2 (\pm 50.5)</u>	<u>4.26 (\pm 0.82)</u>	<u>7.35 (\pm 0.40)</u>	<u>516.05 (\pm 4.749)</u>
<u>11</u>	<u>-4555.5 (\pm 52.7)</u>	<u>2.27 (\pm 0.48)</u>	<u>8.04 (\pm 0.40)</u>	<u>884.80 (\pm 7.62)</u>

645

646 **Data availability**

647 Datasets for the figures shown are available at: <https://doi.org/10.15124/48ade8f3-d750-4fe6-a2d9-b8f69bf86262>

648 **Author contribution**

649 LM, MS, LJC and SJA designed and developed the humidity dependence calibrations for I₂ which were conducted by LM.
650 MS, PR and JT developed the HOI calibration method and was adapted and conducted in lab experiments by LM and MZ.
651 BA was responsible for the management and coordination of the BLEACH field campaign and AP was responsible for the
652 management and usage of the Tudor Hill Marine Atmospheric Observatory.
653 [PBK performed the quantum chemical calculations.](#)
654 LM and LJC prepared the paper, and all authors reviewed the paper.

655 **Competing interests**

656 The authors declare that they have no conflict of interest.

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