

Coastal zone production of IO precursors: a 2-dimensional study

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Abstract. At Mace Head, Eire, in the coastal East Atlantic, diiodomethane has been identified as an important precursor of iodine oxide radicals. Peak concentrations of both CH₂I₂ and IO at low water indicate that the intertidal region is a strong source of organo-iodines. Atmospheric measurements of CH₂I₂ made in marine air are compared with the concentrations predicted by a 2-dimensional model incorporating horizontal and vertical dispersion of surface emissions. The model shows that micrometeorological variability, proximity of the site to emissions, and photolysis all play important roles in determining the CH₂I₂ concentrations at Mace Head. In addition to a tidal-height dependent intertidal flux, which was estimated from seaweed production data, a contribution from offshore (non-local) sources was required in order to reproduce the strong signature of photolysis in the CH₂I₂ observations. A combination of an offshore flux and an intertidal flux (of up to 1.4×10^9 molecules cm⁻² s⁻¹ at low water) results in good agreement between the measured and modelled CH₂I₂ concentrations. Although this study does not necessarily infer emission of CH₂I₂ from the open ocean, it suggests that air-sea exchange of CH₂I₂ in coastal waters does occur.

1 Introduction

During the last decade it was established that catalytic cycles involving halogen oxide radicals (BrO with smaller contributions from IO and ClO) were responsible for rapid ozone depletion events in the Arctic boundary layer during spring (Barrie et al., 1988; Bottenheim et al., 1990; Barrie and Platt, 1997). More recently, the IO radical has been identified in Antarctica (Friess et al., 2001) and in the mid-latitudes (Alicke et al., 1999; Allan et al., 2000) in conjunction with organoiodine precursors (Carpenter et al., 1999), confirming

the potential for the more widespread occurrence of boundary layer ozone depletion. Formation of IO is driven by ocean-atmosphere exchange of organo-iodines that are photolysed quickly within the marine boundary layer. Previous studies at Mace Head found that a range of photolabile iodine containing organics including CH₃I, C₂H₅I, CH₂ICl, CH₂IBr and CH₂I₂ were present in sufficiently high concentrations to sustain IO levels of a few parts per trillion (ppt), via the following reactions:



Of the organoiodines measured, CH₂I₂ was found to be the most important iodine precursor (Carpenter et al., 1999). The photodissociation lifetime of diiodomethane is only a few minutes at midday (Mössinger et al., 1998).

The impact of IO on ozone concentrations depends on the detailed gas phase and heterogeneous chemistry following reaction (2). Computational modelling investigations indicate that the rate of ozone destruction associated with iodine photochemistry in the marine boundary layer could equal that from HO_x photochemistry (Vogt et al., 1999; Stutz et al., 1999; McFiggans et al., 2000). Perhaps of even greater uncertainty are the emissions and distributions of iodine precursors. Shipboard experiments have identified CH₂ICl (Klick and Abrahamsson, 1992; Schall et al., 1997) and CH₂I₂ (Schall et al., 1997) in the open ocean, although it is not yet established whether these originate from microalgae (phytoplankton) or other (e.g. photochemical) sources. That coastal macroalgae are prolific emitters of a wide range of organic halogens is without doubt (e.g. Gschwend et al., 1985; Nightingale et al., 1995; Pedersén et al., 1996; Carpenter et al., 2000 and references therein), although the global budgets of seaweed emissions are very uncertain.

During the ACSOE (Atmospheric Chemistry Studies in the Oceanic Environment) experiment at Mace Head in 1997,

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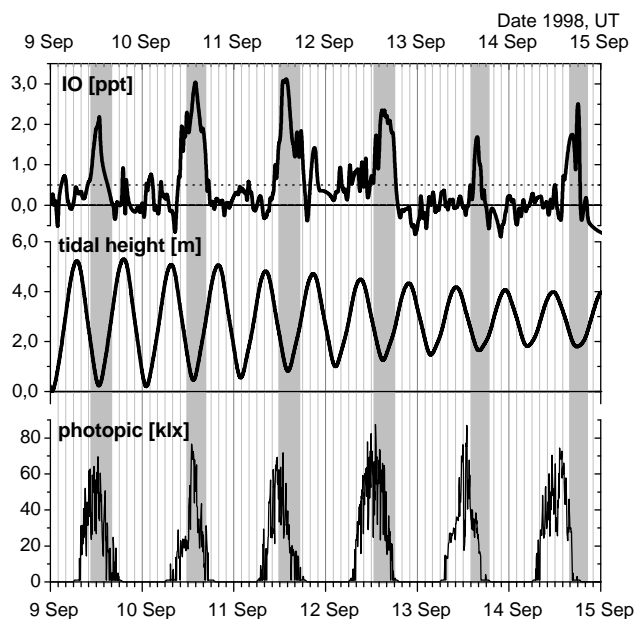


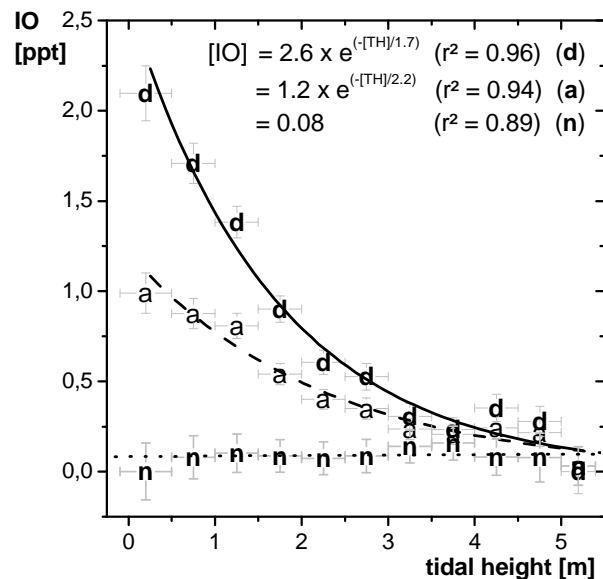
Fig. 1. IO, tidal height and solar radiation (as photopic flux) during 9–15 September 1998. The dotted line on the IO graph represents the average detection limit. The grey areas mark the low tide periods during the day.

macroalgal emissions were observed to have a direct impact on the local atmosphere, as reflected by the peak air concentrations of polyhalogenated halocarbons occurring at low water (Carpenter et al., 1999). These observations may be attributed to direct emission into the surrounding air from exposed macroalgae at low tide and/or increased emissions due to oxidative stress of algae upon exposure (Carpenter et al., 2000). At high tide, emissions from submerged beds are diluted in seawater and may undergo photolysis before exchange at the air-sea interface.

Measurements of organo-iodines and IO radicals at Mace Head were repeated during the 1998 EU project PARFORCE (New Particle Formation and Fate in the Coastal Environment). In this study we attempt to evaluate the relative importance of coastal and offshore emissions of CH_2I_2 , providing further information on sources of the IO radical. A 2-dimensional model is used to predict the horizontal and vertical distributions of CH_2I_2 resulting from coastal and offshore (non-local) emissions.

2 Experimental

Mace Head is located on the remote western coast of County Galway, Eire ($53^\circ 19' \text{ N}$, $9^\circ 54' \text{ W}$). The site is well known for background air measurements and receives relatively clean marine air from the prevailing westerly sector associated with the easterly tracking cyclonic systems of the North At-



n = night
d = day
a = all data

Fig. 2. Correlation of IO and tidal height (TH) during PARFORCE (whole data set). The data points are binned averages of 0.5 m segments of TH. The vertical error bars represent the relative frequency of the data, the horizontal error bars the length of the respective interval. The day-time data and all data fitted with an exponentially decreasing trend, the night-time data has a linear fit.

lantic. The rocky upper littoral zone and cold waters provide favourable conditions for seaweeds.

2.1 Halocarbons

During the period 5–24 September 1998, halocarbons in air were monitored in-situ every 40 minutes with a Hewlett Packard 6890/5973 gas chromatograph/mass spectrometer (GC/MS) system. The GC/MS was operated in a laboratory situated ca. 100 m from the high tide mark, with a sampling inlet located with an open fetch to the ocean at a height of approx. 12 m above mean sea level (MSL). The system is developed for automated air sampling and is described fully in Carpenter et al. (1999). Analysis was also performed on discrete samples of surface seawater and the water surrounding incubated seaweeds. For a full description of analytical procedures for air and seawater analyses during PARFORCE see Carpenter et al. (2000).

2.2 IO measurements by DOAS

Over a 1-month period from 8 September until 8 October 1998 LP DOAS (Long-Path Differential Optical Absorption Spectroscopy; Platt, 1994) measurements of the halogen ox-

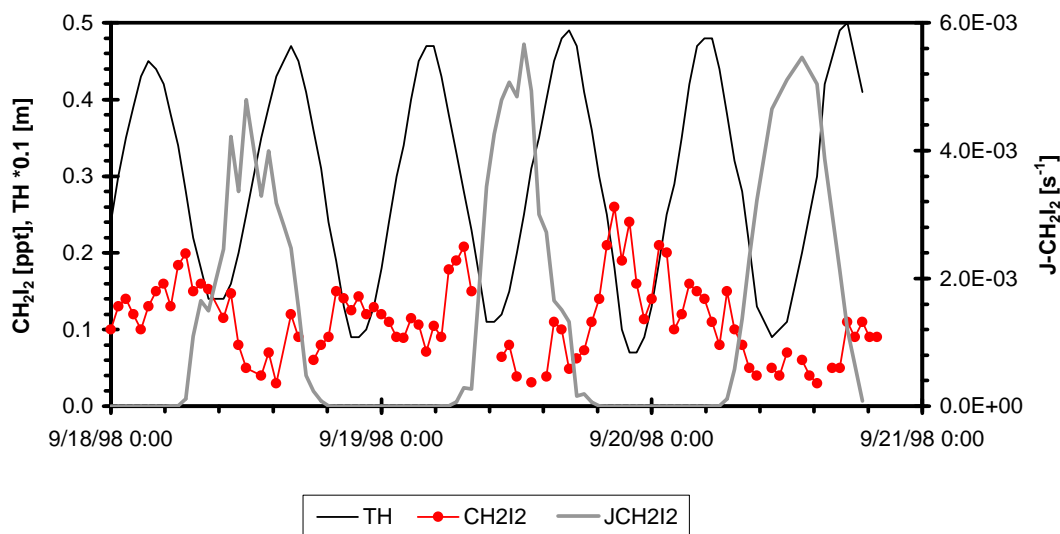


Fig. 3. CH_2I_2 , tidal height and calculated $\text{J-CH}_2\text{I}_2$ during clean marine conditions.

ides IO, OIO and BrO and of other atmospheric trace gases including NO_2 , O_3 , HCHO, HONO and NO_3 were carried out whenever the visibility allowed reasonable signal to noise ratios.

Briefly, the principle of DOAS is the identification and quantification of atmospheric trace gases by their specific narrow (< 5 nm) band optical absorption structure in the open atmosphere, separating trace gas absorption from broad band molecule and aerosol extinction processes, thus allowing very sensitive detection of many molecular species (see e.g. Platt and Perner, 1983). The identification of the gases is unambiguous since their specific absorption structure is, similar to a fingerprint, unique. Calibration of the instrument is not necessary as long as the absorption cross section is known.

A DOAS instrument based on the principle of Platt and Perner (Platt and Perner, 1983) was used at Mace Head. It incorporates a combination of two coaxially arranged Newtonian telescopes, one collimating the light of a Xe-short-arc lamp through the atmosphere, the second recollecting the beam reflected by an array of quartz prism retroreflectors. The telescopes were set up in a laboratory about 20 meters from the shore, while the retroreflector array was placed on the northern shore of the bay. The light beam was running at an average height of 10 meters above the ocean with a light path length of 2×7.27 km. A 0.5 m Czerny-Turner spectrograph ($f = 6.9$, 600 gr/mm grating, thermostated to $30 \pm 0.3^\circ\text{C}$), in combination with a 1024 pixel photodiode array detector (thermostated to $-15 \pm 0.3^\circ\text{C}$), was coupled to the telescope by a quartz fibre, which also performed the task of a mode mixer (Stutz and Platt, 1997). Iodine oxide was measured in the wavelength range from 414 to 437 nm with a spectral resolution of about 0.5 nm (a dispersion of

0.078 nm/pixel). Three further wavelength regions were chosen to detect BrO (335 ± 40 nm) (Hönninger, 1999), OIO (550 ± 40 nm) (Hebestreit, 2001), NO_3 (645×40 nm) and the other species absorbing in the respective wavelength regions (O_3 , SO_2 , NO_2 , HCHO, HONO). Besides NO_3 , all spectra were recorded using the multi channel scanning technique (Brauers et al., 1995). The concentrations of IO were derived using a least squares fit mixer (Stutz and Platt, 1997) of the reference spectra of H_2O , NO_2 and IO, together with a sixth order polynomial to the atmospheric absorption spectrum. The IO reference spectrum was measured in our laboratory, since no absorption cross section of sufficient resolution was available. NO_2 was measured in a reference cell in the field. Absolute values for both spectra were determined by comparison with literature cross sections (Cox et al., 1999; Harder et al., 1997; Hönninger, 1999). The H_2O reference was calculated by a convolution of the instrument function with the absorption cross section of the HITRAN database (HITRAN, 1987).

2.3 Model description

A 3-day period (18–20 September 1998, Julian days 261–263) of clean marine south-westerly air with stable temperatures ($17.2 \pm 1.9^\circ\text{C}$) and wind speed ($5.5 \pm 1.0 \text{ m s}^{-1}$) was selected as a case study for evaluation of marine emissions. Data were evaluated using a model written in the FACSIMILE language. The photolysis rate of diiodomethane ($\text{J-CH}_2\text{I}_2$) was computed at 40 minute intervals using a 2-stream radiative transfer model (Hough, 1988), with absorption cross-sections from Mössinger et al. (1998) and TOMS O_3 column data (<http://toms.gsfc.nasa.gov/ozone/ozone01.html>). Photolysis rates were corrected for cloud cover using the ratio of mea-

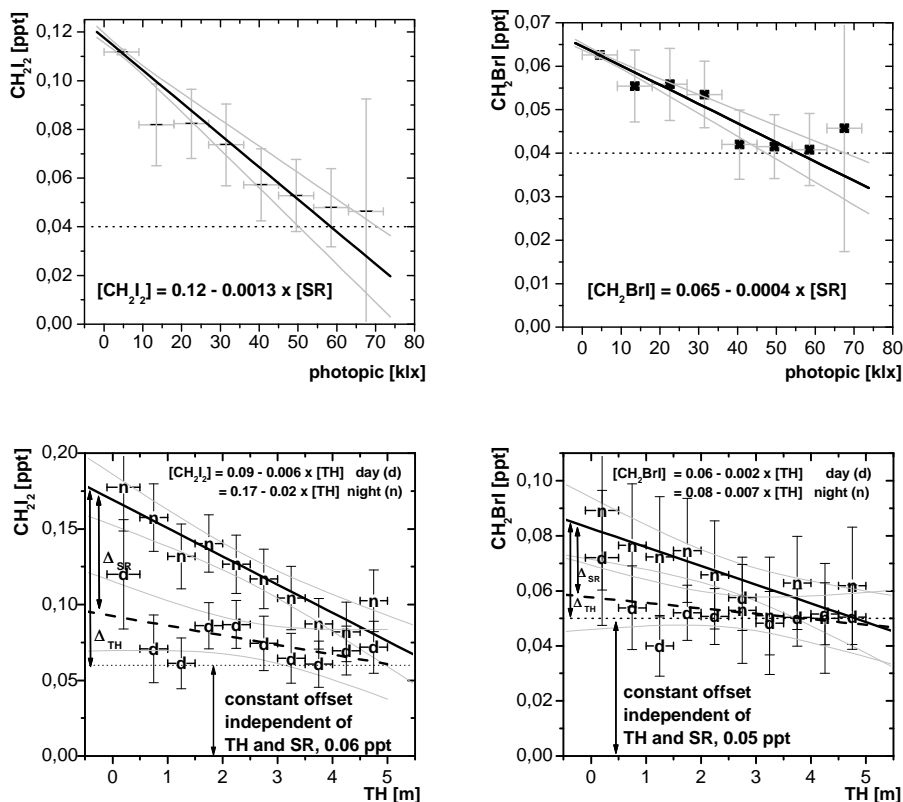


Fig. 4. Correlations of CH_2I_2 and CH_2IBr with solar radiation (SR) and tidal height (TH) for the whole measurement period. The data points are binned averages of 0.5 m segments of TH. The vertical error bars represent the relative frequency of the data, the horizontal error bars the length of the respective interval. The correlation with TH is divided into night-time (n) and day-time (d) values of CH_2I_2 and CH_2IBr , respectively.

sured/modelled UVA data. The 2-dimensional model incorporated a 5.2 km long by 50 m high slice through the atmosphere divided into cells of 100 m length and 2 m height. The first 5 km of the horizontal axis represented the “offshore” region, the next 100 m represented the intertidal region, and the Mace Head site was notionally located in the final 100 m cell, at 12 m height (the sampling height of CH_2I_2). CH_2I_2 was emitted into the bottom cell and transported across consecutive cell faces using the finite volume method to discretise the spatial partial derivatives, given by:

$$\frac{\partial C}{\partial t} = Q + D \frac{\partial^2 C}{\partial x^2} - u \frac{\partial C}{\partial x} \quad (3)$$

where C is the concentration of CH_2I_2 , Q is the net rate of production and destruction, D is the diffusion coefficient (set to zero for horizontal transport), the third term representing advection, with u the horizontal wind speed. Horizontal wind speed was parameterised with 40-minute averages of the measured mean values at 3 m height (de Leeuw et al., 2001). Measurements made at 18 m were typically $\sim 25\%$ higher (de Leeuw et al., 2001) as expected from the logarithmic dependence of wind speed with height and a roughness length, z_0 , of $\sim 10^{-4}$ m (Kunz et al., 2000). Locally,

the extent of the sloping terrain upwind of the site will vary with tidal height, adding to some variability in the roughness length.

The rate of transport across the vertical boundary faces of each cell was described by replacing ∂x with ∂z and the eddy diffusivity coefficient, K_z , in place of D . The diffusivity coefficient was assumed to vary with height in the surface layer according to (cf. Stull, 1988):

$$K_z = \frac{\kappa u_* z}{\phi(z/L)} \quad (4)$$

where κ is the von Karman constant ($\kappa = 0.4$), z is height, u_* is the friction velocity and $\phi(z/L)$ is a stability correction equal to unity during neutrally stratified boundary layer conditions, which prevailed at Mace Head (de Leeuw et al., 2001). During PARFORCE, de Leeuw et al. (2001) found that the friction velocity in marine sector air was approximately equal to 9% of the wind speed at 3 m, a relationship which was used to define u_* in the model. Formation of an internal boundary layer (IBL) would lead to abrupt changes in the diffusivity gradient at heights determined by the distance from the shoreline. It is difficult to assess the impact of

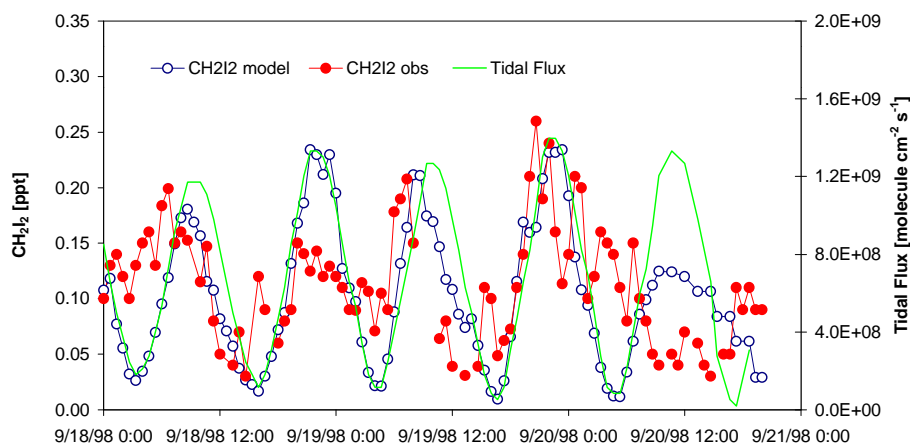


Fig. 5. Comparison of measured and modelled CH_2I_2 levels assuming only tidal emissions, shown with the tidal flux used in the model.

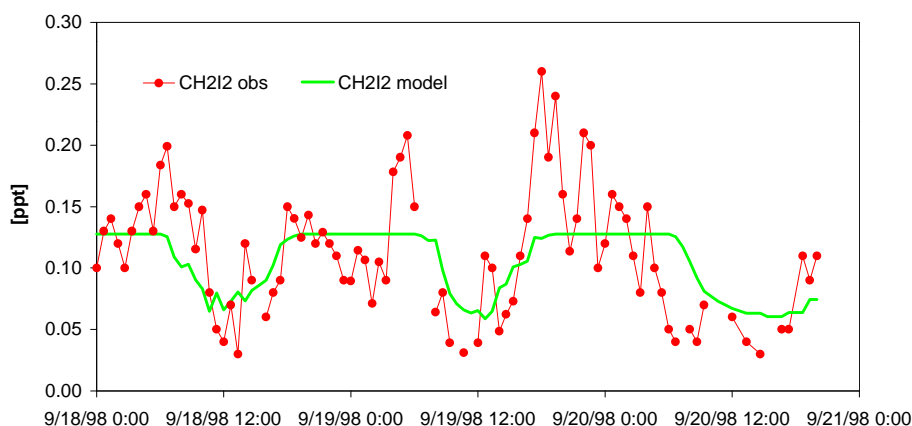


Fig. 6. Comparison of measured and modelled CH_2I_2 levels assuming only offshore emissions (constant wind speed and K).

IBL formation on the conclusions of this study, but we note the additional uncertainty.

3 Results and discussion

3.1 Relationships of iodine species with tidal height and solar radiation

Analysis of PARFORCE data showed that tidal height (TH) and solar radiation (SR) were correlated with CH_2I_2 , IO, and to a lesser extent, CH_2IBr air concentrations. Figure 1 shows IO, tidal height and solar radiation at 555 nm (photopic flux in $\text{kW/m}^2 = \text{klx}$) for the period 9–15 September. The figure shows that the maximum IO concentrations occurred during the day at low tide. The daily IO maximum shifted simultaneously with the minimum in TH (about one hour from day to day), indicating that both solar flux and tidal height controlled the time of the IO peak. There is an IO peak after sunset on the evening of the 14 September, however the IO signal during this period was below the detection limit,

which increased substantially at that time due to fog. Figure 2 shows correlations of IO versus TH for daytime values only (*d*), all data (*a*), and night-time values (*n*) of the 4 week measurement period. Under daylight conditions, high IO values were clearly associated with low tidal height, while at night no correlation between IO and TH was found. The correlation was described well by an exponential decrease with rising tide ($r^2 = 0.96$).

Figure 3 shows tidal height, CH_2I_2 concentrations and calculated $\text{J-CH}_2\text{I}_2$ during the clean marine south-westerly air period 18–20 September 1998. Scatter plots for CH_2I_2 and CH_2IBr concentrations with TH and SR for the whole campaign period are shown in Fig. 4. Strong evidence for these organic iodines as important photolytic precursors of IO is evident from their decreasing concentrations with solar flux and, during night-time, maxima at low tide. There is little correlation between TH and organo-iodine levels during the day, because of the strong influence of photolysis. Note that although peak concentrations of organo-iodines were observed at low water, both CH_2I_2 and CH_2IBr exhibited

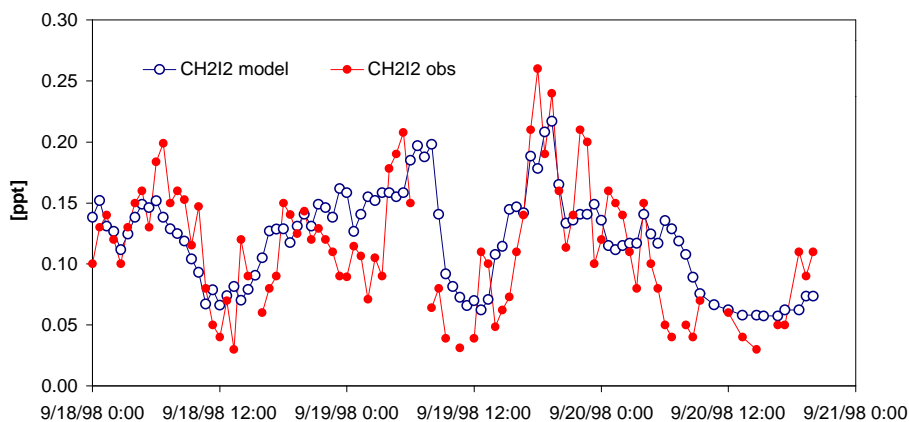


Fig. 7. Comparison of measured and modelled CH_2I_2 levels assuming only offshore emissions (40 minute averaged wind speed and K).

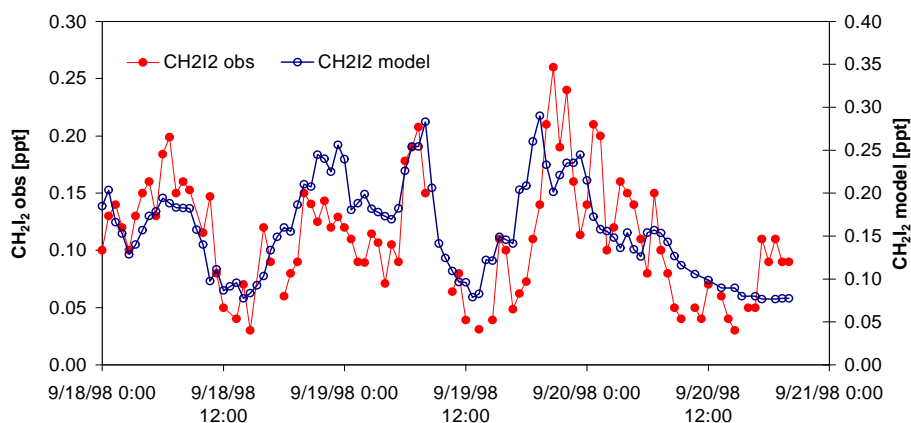


Fig. 8. Comparison of measured and modelled CH_2I_2 levels assuming both offshore and intertidal emissions (40 minute averaged wind speed and K).

non-zero concentrations above their detection limits of 0.03 pptv even at high tide, presumably due to local sources, as discussed later. The dependence of CH_2I_2 on tidal height and solar radiation was found to be $[\text{CH}_2\text{I}_2]_{\text{night}} = 0.17 - 0.02[\text{TH}]$ and $[\text{CH}_2\text{I}_2]_{\text{all}} = 0.12 - 0.0013[\text{SR}]$.

Given that the organoiodine precursors exhibited linear dependencies with TH at night, the cause of the exponential relationship of IO with TH (Fig. 2) does not appear to be explained by source variations and is possibly attributable to the photochemistry of IO. It should also be noted that the IO measurements represent averages over ~ 7 km, whereas the organoiodines were measured by point sampling.

3.2 2-dimensional model simulations

The initial model simulations incorporated only tidal emissions. The night-time CH_2I_2 vs TH negative trend, shown in Fig. 4, was presumed to be indicative of coastal production processes. Therefore the tidal dependence of the coastal flux was assumed to have the same qualitative form. A quan-

titative fix was available from independent data on seaweed emissions, described below.

In addition to atmospheric measurements made during PARFORCE, the release rates of organic bromines and iodines from seaweeds were determined from incubations in seawater of ten species of brown, red and green macroalgae collected in the intertidal or subtidal zones of the rocky shore (Carpenter et al., 2000). The most prevalent seaweeds present in the intertidal zone at Mace Head, in common with most Northern European rocky shores (Michanek, 1975), were the brown algae *Laminaria digitata*, *Laminaria saccharina* and *Ascophyllum nodosum*. These algae are also among the most productive in terms of CH_2I_2 emissions, with mean production rates of 8.3, 1.9 and 0.36 pmol g^{-1} fresh weight hr^{-1} , respectively (Carpenter et al., 2000). Estimates of total kelp density from data provided by the Irish Seaweed Industry Organisation (ISIO) are 11.6 kg m^{-2} (Carpenter et al., 2000). However this value includes *Laminaria hyboborea*, which is only present at depth, therefore the kelp density of the intertidal zone should be reduced by $\sim 25\%$ (Carpenter

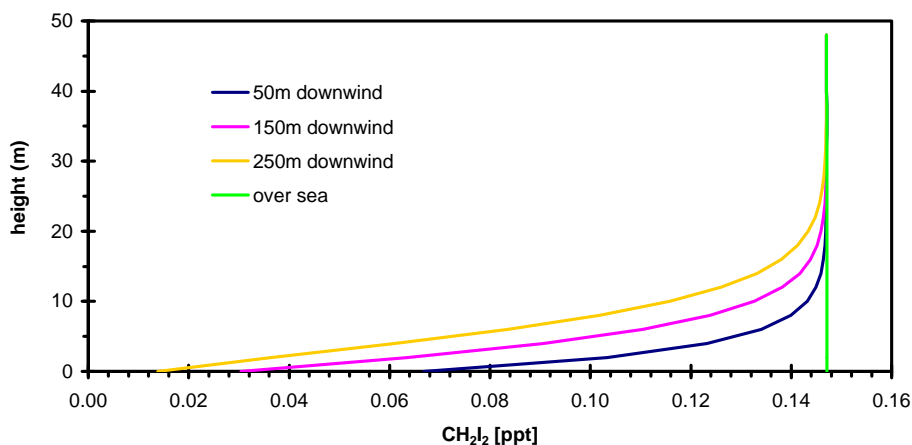


Fig. 9a. Gradient of CH_2I_2 mixing ratios predicted using the offshore flux scenario.

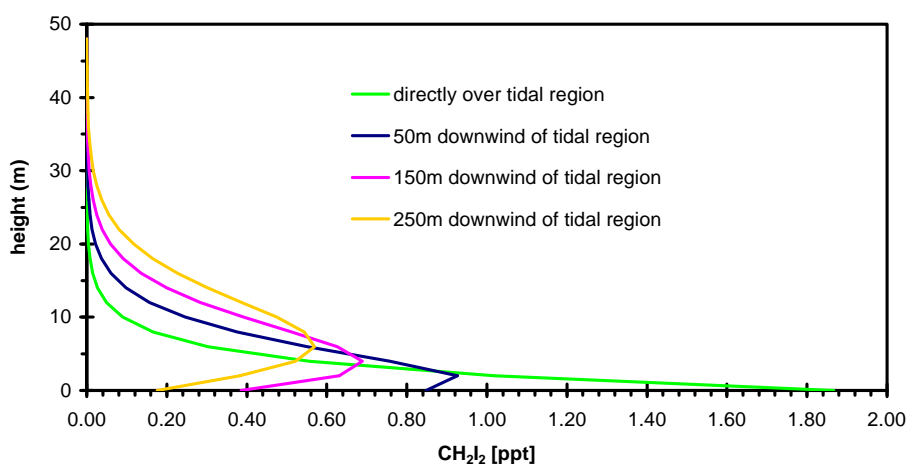


Fig. 9b. Gradient of CH_2I_2 mixing ratios predicted using the coastal flux scenario.

et al., 2000). Assuming the density is evenly spread between *L. digitata*, *L. saccharina* and *A. nodosum* and over the intertidal zone leads to a total CH_2I_2 emission rate of $\sim 1.5 \times 10^9$ molecules $\text{cm}^{-2} \text{s}^{-1}$. This rate was used as the maximum coastal flux at the lowest tide. The high tide flux was assumed to be equal to the offshore flux, described below.

Figure 5 shows that the overall magnitude of the CH_2I_2 concentrations predicted by the model was in good agreement with the observed levels, indicating that estimation of the CH_2I_2 flux from seaweed emissions was a valid approach. However, there were clear discrepancies in some of the features. Also shown in Fig. 5 is the modelled intertidal flux used in this scenario. At night, the flux is closely correlated with the CH_2I_2 concentration, with small differences due to the variability in wind speed and eddy diffusivity. During the day, the modelled CH_2I_2 concentration is reduced relative to the tidal flux because of photolysis. However, the reduction due to photolysis was not sufficient to simu-

late the observations, as is clearly seen during the last day of data. There are also other features not well represented by the model, such as the overestimation of $[\text{CH}_2\text{I}_2]$ during the night of the 18–19 September.

3.3 Simulation of offshore emissions

An alternative flux situation was investigated, wherein the offshore region was a source of CH_2I_2 . The atmospheric concentration of CH_2I_2 predicted by the 2-dimensional model is related to the product of the magnitude of the flux and the length of the grid. It is therefore not possible to put a meaningful value on the magnitude of the offshore flux without knowing precisely the length of the upwind fetch. Although this was known to a reasonable degree for the tidal scenario (i.e. the intertidal range), the degree of open ocean production of CH_2I_2 is uncertain.

Ocean production of CH_2I_2 up to 5 km offshore of Mace Head was assessed from discrete surface seawater measure-

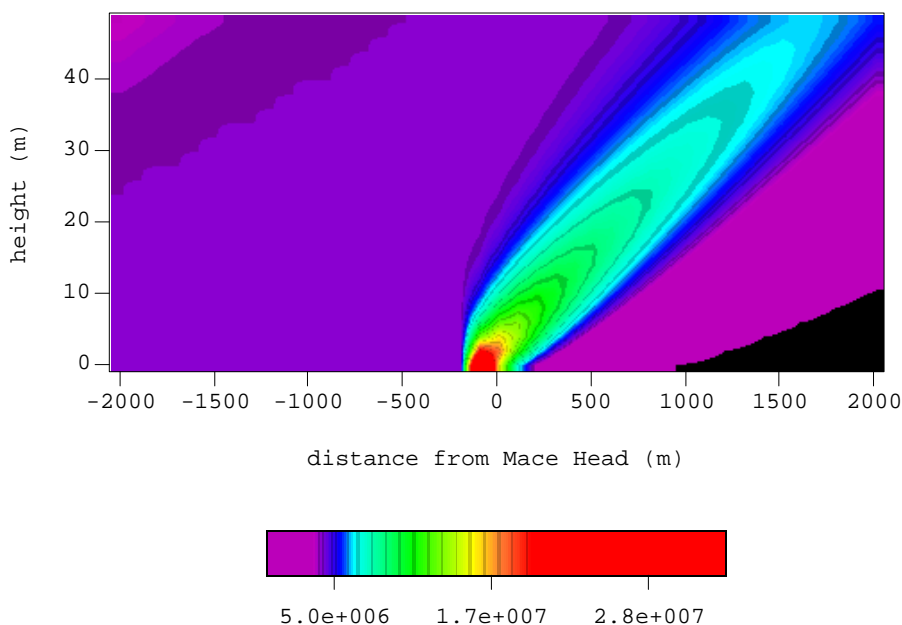


Fig. 10. Predicted variation in CH_2I_2 concentration gradients upwind (negative distances) and downwind (positive distances) of Mace Head at midnight. The colour scale represents the CH_2I_2 concentration in molecule cm^{-3} , with black representing concentrations less than 1×10^5 molecule cm^{-3} .

ments made during the 2 weeks immediately after the atmospheric measurements. Although there was a higher concentration of CH_2I_2 in seawater sampled directly over kelp beds than in areas with no seaweeds, there was no obvious relationship between distance from shore and dissolved CH_2I_2 concentration between 200 m and 5 km offshore. The average concentration over this region was 0.52 ± 0.26 pmol L^{-1} (mean of 6 measurements) and the mean seawater temperature was 12°C (Carpenter et al., 2000). The equilibrium air concentration at this temperature calculated using the Henry's law coefficient for CH_2I_2 reported by Moore et al. (1995) is 0.1 pptv. Thus, it is difficult to assess whether or not the offshore waters of Mace Head were a source of CH_2I_2 or simply in equilibrium. An upper limit to the flux can however be calculated using a CH_2I_2 concentration in air of zero. From the Liss-Merlivat expression (Liss and Merlivat, 1986) with a square root molecular weight correction for the transfer velocity (Liss and Slater, 1974) and the mean wind speed, the flux is 2×10^5 molecules $\text{cm}^{-2} \text{s}^{-1}$.

Because the offshore emissions and fetch are highly uncertain, we do not attempt here to provide an estimate of these emissions, but rather to provide an emission regime that results in good agreement with the measurements. Thus, the point of including an offshore scenario is to establish whether non-local sources of CH_2I_2 were present at Mace Head, and the possible contribution of these.

With a 5 km fetch of offshore emissions, a flux of 4×10^7 molecules $\text{cm}^{-2} \text{s}^{-1}$ was required for good agreement of the overall levels of measured and modelled $[\text{CH}_2\text{I}_2]$, as

shown in Fig. 6. Because a fetch of at least up to 5 km offshore was established by the seawater measurements, this flux can be taken as an upper limit. Note, however, that the same agreement would have been possible with the estimated flux of 2×10^5 molecules $\text{cm}^{-2} \text{s}^{-1}$ and a fetch of about 1000 km. The model predictions shown in Fig. 6 were made assuming constant wind speed and eddy diffusivity (averages for the period). The model reproduced the daytime reductions in $[\text{CH}_2\text{I}_2]$ due to photolysis, indicating that non-local sources were indeed an important contribution to the atmospheric levels at Mace Head. However, the peak levels of CH_2I_2 were not replicated well by the model. The simulation was repeated using 40-minute averages of wind speed and eddy diffusivity. Figure 7 shows that the inclusion of micrometeorological variability resulted in significantly improved agreement with the measurements. Clearly, meteorological parameters also played a large part in controlling the variability in CH_2I_2 concentrations at Mace Head.

The result of including both coastal and offshore fluxes in the model is shown in Fig. 8. Although the combination of fluxes resulted in an overestimation of modelled CH_2I_2 concentrations by $\sim 30\%$, the correlation between modelled and measured data was greater than that of either the separate flux scenarios. For ten-point averages of modelled vs measured data, the r^2 values were 0.89 (coastal + offshore), 0.78 (offshore only) and 0.75 (coastal only).

3.4 Prediction of spatial variation of CH₂I₂ concentrations

The horizontal domain of the model was extended by 2 km in order to examine the dispersion of CH₂I₂ downwind from the shore. Both the coastal and offshore flux scenarios gave rise to similar levels of CH₂I₂ at Mace Head at 12 m height. However, the vertical profiles predicted were very different, as shown in Figs. 9a and 9b. The offshore source resulted in a constant CH₂I₂ concentration vertical gradient over the open sea, but a rapidly increasing concentration with height over Mace Head, 150 m downwind of the shore. Conversely, over a few metres height, the tidal source gave rise to decreasing CH₂I₂ levels over Mace Head, 50 m downwind of the tidal region (i.e. the mid point of the 100 m cell adjacent to the 100 m tidal region). Internal boundary layer formation would lead to perturbations of these modelled gradients.

Figure 10 shows the predicted CH₂I₂ concentrations from 2 km upwind of the site to 2 km downwind at midnight in westerly (marine) air. Considerable heterogeneity is expected over the few hundred metres downwind. Measurements in both the horizontal and vertical domains at Mace Head are planned to test these predictions.

4 Summary and conclusions

Strong evidence for the organoiodines CH₂I₂ and CH₂IBr as photolytic precursors to the IO radical at Mace Head was shown from the dependence of the atmospheric concentrations of all three species on tidal height. Iodine oxide concentrations peaked at low water during midday hours and the reactive organoiodines, whose atmospheric lifetimes are less than 1 hour at midday, peaked when low water coincided with night.

The intertidal flux of CH₂I₂ upwind of Mace Head was estimated from seaweed emissions and used in a 2-dimensional model to predict the CH₂I₂ concentrations in marine air at Mace Head. This flux resulted in good agreement between the mean concentrations of measured and modelled CH₂I₂, but did not reproduce well the daytime depletion due to photolysis. A non-local, offshore, source of CH₂I₂ was invoked, which did reproduce the daytime depletion. Three separate scenarios of offshore sources alone, tidal sources alone, and offshore plus tidal sources were used as inputs to the model. The model showed that variability in wind speed and friction velocity, proximity to source, and photolysis all played important roles in determining the CH₂I₂ concentrations at the site. The best agreement with the CH₂I₂ observations was obtained using constrained micrometeorological fields, and a combination of a constant offshore flux and a tidal flux that peaked at 1.4×10^9 molecules cm⁻² s⁻¹ at low water.

Because the atmospheric concentration of CH₂I₂ predicted by the 2-dimensional model was related to the length of the upwind fetch, a meaningful value cannot be assigned to the offshore emission rate, although we estimate an upper

limit of 4×10^7 molecules cm⁻² s⁻¹. Rather, we conclude that a contribution from both intertidal and offshore sources is likely, with the tidal flux being several orders of magnitude higher. Although this study cannot predict whether CH₂I₂ is emitted from the open ocean, it suggests that it does undergo air-sea exchange in coastal waters despite its presumably rapid photolysis rate.

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