A comparison of spectrophotometric and denuder based approaches for the determination of gaseous molecular iodine

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Abstract. The presence of molecular iodine in the atmosphere is thought to have implications for both climate and human nutritional health, but measurement of the gas at low concentrations requires technically demanding techniques that are not widely accessible. Here, amylose coated denuder tubes and solvent traps coupled with spectrophotometric detection are evaluated and compared as relatively cheap and straightforward methods to measure gaseous molecular iodine at environmentally relevant concentrations. Denuder tubes were found to give unacceptably low and highly variable recoveries of molecular iodine from a test gas source, with values ranging from 1 to 62%. Blank concentrations were also high, being equivalent to a gas phase concentration of 5 pptv under typical operating conditions. Ethanol and hexane solvent traps gave much better performance. Optimisation of the hexane solvent trap method gave 100% recovery and an atmospheric limit of detection of 70 pptv, which is within the range of concentrations observed in the coastal marine atmosphere.

1 Introduction

Understanding the biogeochemical cycle of iodine is important for three reasons: volatile iodine compounds are photoylsed in the atmosphere to give iodine atoms which are implicated in ozone destruction and particle formation reactions and may thus impact climate (O’Dowd and Hoffmann, 2005); iodine is an essential human nutrient, deficiency of which remains the leading cause of preventable brain damage and mental retardation worldwide (WHO, 2004); the long-lived radionuclide iodine-129 is released to the environment by the nuclear industry (Raisbeck and Yiou, 1999). Marine formation (biotic and abiotic) followed by sea-air exchange is the main route by which iodine enters the atmosphere. AtmospHERically processed iodine is subsequently deposited on land and incorporated into terrestrial food chains.

Molecular iodine has been observed in the coastal marine atmosphere at concentrations ranging from <0.2 to 93 pptv (Saiz-Lopez et al., 2006b; Finley and Saltzman, 2008). It is thought to be either released directly by exposed macroalgae at low tide or formed by the reaction of ozone and iodide on the surface of the algal fronds (Palmer et al., 2005; Kupper et al., 2008; Dixneuf et al., 2009). It may also be formed by the reaction of ozone and iodide at the ocean surface (Garland and Curtis, 1981), but as yet no open ocean measurements of atmospheric molecular iodine have been reported. Under-ice microalgal production of molecular iodine has also been suggested as a source of atmospheric reactive iodine in polar oceanic regions (Saiz-Lopez and Boxe, 2008). For all of these environments, the processes and fluxes by which molecular iodine may be formed and volatilised are not yet fully understood.

One of the hindrances in this field is the lack of a readily accessible, straightforward yet robust means of measuring molecular iodine in the gas phase at low concentrations approaching those encountered in the atmosphere. Typically, atmospheric measurements of molecular iodine are made using long path differential optical absorption spectroscopy (LP-DOAS; Saiz-Lopez et al., 2006b). However, this yields concentrations averaged over a path length of several kilometres so is unsuitable for point measurements and cannot be used on board a ship for open ocean measurements. Advances have been made in the use of broadband cavity enhanced absorption spectroscopy (BBCEAS) for the measurement of gaseous molecular iodine in situ (Saiz-Lopez et al., 2006a; Dixneuf et al., 2009), but this approach is also technically demanding and is unavailable to the majority of
researchers. Recently, atmospheric pressure chemical ionisation tandem mass spectrometry (APCI-MS-MS) has also been applied to the measurement of molecular iodine in air (Finley and Saltzman, 2008).

The use of diffusion denuders with either a starch (Chen et al., 2006; Saiz-Lopez et al., 2006a) or an alpha-cyclodextrin coating (Huang and Hoffmann, 2009) to measure molecular iodine in the coastal atmosphere has been reported. At higher concentrations, gas phase molecular iodine can be measured using a solvent trap followed by spectrophotometric detection (Palmer et al., 2005). These methods have the advantage of being straightforward, specific, relatively cheap and able to produce time-averaged rather than spatially-averaged measurements. Additionally, if using denuders the possibility exists for samples to be collected and analysed at a later date. Here we evaluate and compare the applicability of starch coated denuders and ethanol and hexane solvent traps for the measurement of molecular iodine at atmospherically relevant concentrations.

2 Experimental

2.1 Generation of a molecular iodine test source

A permeation oven (Kin-Tek™, USA) fitted with a commercial molecular iodine permeation tube (Kin-Tek™, USA) was used to provide a constant source of gaseous iodine for evaluation of the detection methods. The oven temperature was either 30 or 60 °C according to the required concentration. Iodine emission rates at these operating temperatures were determined directly by repeated weighing of the tube at approximately two week intervals. The permeation chamber was flushed with zero grade nitrogen (BOC) at a flow rate of 0.05 L min$^{-1}$. The outlet flow was further diluted with nitrogen, the flow rate of which was varied to achieve the desired final concentration of molecular iodine. Where required, a needle valve and T-piece was used to ensure that the outlet flow rate to the solvent trap or denuder remained constant. The range of concentrations obtainable was 190 pptv to approximately six months earlier. Recovery and trapping efficiency (T. E.) were calculated according to Eqs. (1) and (2), respectively.

\[
\text{Recovery}(\%) = \frac{\text{iodine measured}}{\text{iodine expected}} \times 100
\]

\[
\text{T. E.}(\%) = 100 \times \frac{[\text{iodine}]_{\text{trap1}}}{[\text{iodine}]_{\text{trap1}} + [\text{iodine}]_{\text{trap2}}}
\]

Three freshly coated and three older tubes which had not been attached to the iodine source were analysed as blanks. A preliminary blank experiment in which six starch coated tubes were flushed with compressed air at a flow rate of 0.5 L min$^{-1}$ was also conducted. A number of other preliminary experiments using the same conditions as described here, but differing test gas concentrations and starch types were also conducted.
In order to determine whether the concentration of iodine in the gas stream affects recovery, the dilution flow rate on the permeation oven was varied to give four different concentrations (ranging from 14 to 21 ppbv; see Table 1). The outlet flow was split and a needle valve used to maintain a constant flow rate of 0.5 L min\(^{-1}\) through the denuders. The duration of exposure was adjusted so that the total amount of iodine passing through the tubes was the same for each concentration. As before two tubes were used in series. Replicate experiments were conducted at each gas concentration.

The stability of iodine while on the denuder tubes was evaluated by exposing six pairs of tubes to the iodine test gas as described above, and analysing three pairs immediately and three pairs after storage for one week at room temperature.

### 2.2.3 Elution and analysis

Following the method of Chen et al. (2006), denuder tubes were eluted with 4 mL of 5% (v/v) tetramethyl ammonium hydroxide (TMAH; Riedel-de Haën). Each tube containing the TMAH was capped and gently inverted and rotated for 20 min before transfer of the eluent to a plastic sample tube. The tube was rinsed with a further 4 mL of ultra-pure water (MilliQ) to give a final TMAH concentration of 2.5% in the sample. Serial elutions gave iodine contents indistinguishable from solvent blanks in the second elution, suggesting this protocol was sufficient to remove the iodine from the inside of the tube. Chen et al. (2006) heated the eluate at 90 °C for three hours, but we found this to make no difference to the iodine content of extracts, blanks or iodine standards.

The iodine content of the denuder tube eluate was determined using an Elan 6000 inductively coupled plasma-mass spectrometer (ICP-MS) at the Food and Environmental Research Agency (formerly Central Science Laboratory), UK. Samples were analysed by flow injection with aspiration into the plasma via a PFA concentric nebuliser/water cooled cyclonic spray chamber combination. Sampler and skimmer cones were both platinum tipped. Instrument conditions were as follows: nebuliser gas flow ∼1.35 L min\(^{-1}\); dwell time 100 ms; 4 sweeps per reading; 55 readings per replicate. Calibration was performed using eight external standards (0 to 200 µg L\(^{-1}\) iodine) prepared from potassium iodate in 2.5% TMAH. A bulk diluent (0.5% TMAH) containing tellurium and antimony as internal standards (to correct for any instrument drift over the course of a run) was used to make 1:1 dilutions of all stock standards and samples prior to analysis. This ICP-MS method for iodine has been thoroughly validated using certified reference materials extracted into TMAH and is UKAS accredited to ISO 17025.

### Table 1. Recoveries of iodine by denuder tubes at different iodine concentrations. Values are for the sum of two tubes in series, blank corrected and averaged for two repeat experiments (range given in parentheses).

<table>
<thead>
<tr>
<th>Iodine concentration, ppbv</th>
<th>Exposure time, min</th>
<th>Recovery, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>21</td>
<td>8.4</td>
<td>1.6 (1.4 to 1.7)</td>
</tr>
<tr>
<td>18</td>
<td>9.8</td>
<td>6.1 (0.8 to 11.4)</td>
</tr>
<tr>
<td>15</td>
<td>11.2</td>
<td>0.7 (0.6 to 0.8)</td>
</tr>
<tr>
<td>14</td>
<td>12.6</td>
<td>24 (21.0 to 27.5)</td>
</tr>
</tbody>
</table>

### 2.3 Solvent traps

#### 2.3.1 Comparison of hexane and ethanol

Molecular iodine is very soluble in both hexane and ethanol, so these solvents are both effective in stripping it from the gas phase. In \(n\)-hexane, molecular iodine is stable and absorbs strongly at ∼520 nm (Fig. 1a) to give a purple coloured solution. In ethanol, molecular iodine disproportionate to give a number of species including iodide (\(I^–\)), which can be quantified by its absorbance at ∼225 nm (Fig. 1b). Both hexane and ethanol solvent traps have been used to measure the evolution of iodine gas in experiments (Palmer et al., 2005; Rajendran, 2008).

Absorption spectra from 190 to 650 nm were measured using a PerkinElmer Lambda 25 UV/VIS spectrophotometer fitted with a 1 cm cell. The instrument was autozeroed using the appropriate solvent. Calibration curves were constructed using standard solutions prepared from ground molecular iodine (puriss p.a., Riedel-de Haën) dissolved in either \(n\)-hexane (HPLC grade, Fisher UK) or absolute ethanol (Analytical Grade, Fisher, UK); the concentration range used was typically 500 to 5000 nM. To avoid any memory effects, the spectrophotometer cuvettes were rinsed thoroughly with solvent between each scan and solvent blanks were run between each sample or standard. The spectrophotometric limit of detection (LoD\(_{\text{spec}}\)) was calculated using Eq. (3) (Harris, 2002), where \(\sigma_{\text{STD}}\) is the standard deviation of the absorbance for repeat measurements of a mid-range standard (\(n=3\)) and the sensitivity is defined as the slope of the calibration curve.

\[
\text{LoD}_{\text{spec}} = 3 \cdot \sigma_{\text{STD}} / \text{sensitivity}
\] (3)

Initial trials used a prototype solvent trap consisting of a 100 mL round bottomed flask containing 50 mL of solvent. The prototype flasks were fitted with ground glass stoppers with an inlet tube reaching below the solvent surface and an outlet tube remaining above the solvent surface. The iodine test gas was bubbled through the solvent at a flow rate of 0.05 L min\(^{-1}\). The prototype traps were wrapped in foil to
prevents photolysis of molecular iodine and immersed in an ice/sodium chloride (−9°C) cold bath to minimise solvent evaporation.

The traps were deployed for periods of up to three hours and the iodine content determined spectrometrically. The recovery and trapping efficiency were determined using two traps in series as for the denuder tubes. The evaporation rate of the solvents under the trapping conditions was determined by bubbling with compressed air at 0.05 L min−1 and measuring the volume lost over time.

2.3.2 Optimisation of the hexane trap method

Upon establishing that hexane was the more suitable solvent (see Sect. 3.2.1), various improvements were made to the method to reduce the LoD as far as possible. The optimised solvent traps were set up as shown in Fig. 2. To increase the sensitivity of the spectrophotometric detection a 10 cm pathlength cell was used instead of a 1 cm cell. To further reduce solvent evaporation, and thus increase the length of time for which trapping can occur, an acetone/dry ice bath (−50°C) was used in the place of the ice/sodium chloride bath. In order to enhance contact between the gaseous analyte and the solvent, the round bottomed flasks were replaced with 25 mL capacity gas Midget bubblers (Supelco) with glass frits fitted to the end of the inlet tube. The traps were wrapped in black tape to prevent photolysis of molecular iodine. The volume of hexane used as trapping solvent was decreased to 20 mL while the test gas flow rate was increased. Volumetric losses of hexane by evaporation within this new experimental set-up were investigated as a function of nitrogen flow rate.

2.3.3 Derivatisation of molecular iodine using leuco-crystal violet

Leuco-crystal violet (4,4′,4″-methylidynetris (N,N-dimethylaniline)) reacts with molecular iodine to give a violet coloured product, and consequently has been used as a reagent in iodine assays (Black and Whittle, 1967). The method was adapted for use in a hexane matrix to determine whether it could further reduce the limit of detection of the solvent traps. Briefly, 0.2 mL of leuco-crystal violet indicator (∼1.5 g L−1) was added to 20 mL of hexane in an amber glass bottle, mixed for exactly one minute and the absorbance at 592 nm recorded.

3 Results and discussion

3.1 Denuder tubes

3.1.1 Blanks

The preliminary blank experiment in which coated denuder tubes were flushed with compressed air found high concentrations of iodine in the TMAH eluate (5.2 ng mL−1). This was traced to contamination of the TMAH during storage and handling at the University of York, despite taking all appropriate precautions (e.g. storing in an air-tight box away from known sources of iodine compounds, using dedicated, acid-washed glassware). Subsequently, elution of the denuder tubes was conducted in laboratories at the Food and Environment Research Agency, which are thought to have lower background levels of iodine compounds due to the differing nature of work conducted there. Blank values of around 2 to 3 ng mL−1 or less were then achieved. For a denuder tube deployed for 12 h and sampling at a flow rate of 0.5 L min−1, this is equivalent to an atmospheric iodine concentration of 5 pptv. Occasionally, extremely high iodine contents were observed in blanks although the source of the contamination could not be identified. As the blank values quoted here fall within the lower range of atmospheric molecular iodine concentrations (Saiz-Lopez et al., 2006b), the method may only be suitable for measurements where iodine levels are expected to be high (over kelp beds for example) and even then great care must be taken to minimise contamination. All results presented here are blank corrected unless otherwise noted.

3.1.2 Recovery and reproducibility

The total amount of iodine recovered from the permeation oven outlet by two denuder tubes in series was very low, with average values of 5 and 17% for freshly coated and older tubes, respectively (Fig. 3). The recovery was also poorly reproducible – not only was there a large difference between batches (a) and (b), but within each batch the relative standard deviations were 70 and 22%, respectively. Furthermore,
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Fig. 2. Optimised solvent trap experimental set up.

Fig. 3. Percentage recovery of molecular iodine from denuder tubes 1 and 2 in series for (a) freshly coated tubes and (b) tubes coated six months previously. Note that in (a), replicate 4, there is no data for tube 1.

the trapping efficiency was highly variable, with no consistent trends in the proportion of iodine retained on the first and second tube in series. Preliminary trials of the denuder tubes (results not shown) also indicated highly variable and generally low recovery of iodine from the gas stream, with values ranging from 0 to 27% for single tubes exposed for varying lengths of time and 46 to 62% for a separate batch in which two tubes in series were combined.

These results suggest that amylose-coated denuders used as described are not suitable for the quantification of molecular iodine in air. They contrast strongly with the findings of Chen et al. (2006), who report 92% recovery and 85% trapping efficiency using the same method. Given that the elution method appeared satisfactory and the ICP-MS determination step was well validated (see Sect. 2.2.3), the most likely cause of this major discrepancy, as well as the high levels of variation within the results presented here, is thought to be related to the starch coating.

The poor recoveries are not thought to be the result of saturation of the starch coating. Dry amylose can absorb 26% of its mass in iodine (Rundle and French, 1943), which equates to 1 mg of iodine for a denuder tube coated as described here, while the total amount of iodine passed through the tubes was only 800 ng. The iodine loading was double that used in the validation of Chen et al. 2006 (480 ng), but even if the denuders were completely saturated at 480 ng iodine, a recovery of ~55% should still be observed at the higher loading. Similarly, there was no apparent relationship between recovery and concentration of the iodine test gas over the range of concentrations investigated here (Table 1). The test gas concentration used here (11 ppbv) was approximately half that used by Chen et al. 2006 (46 ppbv) but the recoveries were much lower.

Amylose may take a number of structures, of which only the single helix “V” formation is able to complex with molecular iodine under anhydrous conditions, while the more ordered “A” and “B” configurations do not (Rundle and French, 1943; Rendleman, 2003). Consequently, the protocol used to isolate amylose from starch influences the ability of the amylose to interact with iodine (Rendleman, 2003), and thus the exact type and brand of starch used is likely to affect iodine recovery by the denuder. Amylose was used here, as it is the component of starch with the highest iodine binding capacity (Bates et al., 1943) and preliminary trials using ground starch (Fluka) as per the method of Chen et al. (2006) gave low recoveries and poor reproducibility. However, it is possible the amylose had undergone retrogradation from the “V” form at some stage, rendering it less reactive to iodine. For some starches and amyloses, the presence of moisture is required for iodine binding to occur (Rendleman, 2003). This is thought to be because water loosens the starch macrostructure in a manner that aids iodine absorption. As the denuder tubes were dried and capped after coating until use and the test gas was dry, little moisture was available to facilitate iodine absorption by the starch. It is possible that the slightly better recoveries obtained using denuder tubes that had been stored for six months between coating and use (Fig. 3b) occurred because these tubes had been exposed to some moisture during storage. In summary, as absorption of iodine by starch is a function of both macrostructure (Rundle and French, 1943) and relative humidity (Rendleman,
2003) both these factors must be strictly controlled if starch coated denuder tubes are to be used to make reproducible measurements of atmospheric iodine.

### 3.1.3 Stability

There was not a statistically significant difference in iodine content between denuders stored for one week and those analysed immediately following exposure (using a student’s t-test $p=20\%$). However, the lack of any significant difference was likely the result of the very high variance in both sample sets (113 and 52% RSD, respectively) so it is not possible to draw conclusions about the stability of exposed denuder tubes from this experiment.

### 3.2 Solvent traps

#### 3.2.1 Comparison of hexane and ethanol

The properties of the prototype hexane and ethanol solvent traps are summarised in Table 2. The performance of the two solvents was similar, with hexane exhibiting a lower spectrophotometric limit of detection and higher trapping efficiency. Note that while $\sim95\%$ ethanol is more typically used for UV spectrophotometry because it has lower levels of benzene contaminants (Williams and Fleming, 1995), absolute ethanol was used here because molecular iodine is more soluble in it (Nakanishi and Asakura, 1977). Using $\sim95\%$ ethanol (Fluka), an improved spectrophotometric limit of detection of $170 \text{ nM}$ was achieved but this is still greater than the value obtained for hexane.

The prototype hexane trap gave 84% recovery (Fig. 4) and minimum trapping efficiency of 86% (Table 2). Calculating trapping efficiency from recovery also gives a value of 86%; the good agreement between measured and calculated suggests that losses of iodine within the permeation system were minimal. The prototype ethanol trap appeared to give recoveries greater than 100%. This is thought to be because the action of passing the test gas through the solvent trap purges it of other dissolved gases, and consequently disrupts the complex equilibria that govern the disproportionation of molecular iodine (Lengyel et al., 1993). Comparison of calibration curves prepared using pre-purged and unpurged ethanol suggested that the iodine signal at $\sim225 \text{ nm}$ (iodide) in purged ethanol may be 1.2 to 1.8 fold greater than in unpurged ethanol, resulting in overestimation of the amount of iodine trapped. Therefore, it is necessary to ensure that the calibration solution has been purged to exactly the same extent as the trap at the point of sampling, and that identical conditions of pH are maintained. It was not possible to correct the recovery results presented here for the effect of purging and so instead the trapping efficiency of 67% is taken as an indicator of minimum recovery.

The utility of a solvent trap also depends on the rate of solvent evaporation – the lower the evaporation rate, the longer the trap may sample for and thus the lower the limit of detection achievable. Under the prototype trapping conditions, the evaporation rates of hexane and ethanol were similar, at 1–2 mL per hour. Assuming a 50 mL starting volume, this means the traps can be used for up to four hours with less than 10% solvent evaporation. A fraction of the trapped iodine evaporates with the solvent. By evaporating iodine in hexane solutions under a stream of gently warmed N$_2$ to simulate the warm gas stream produced by the permeation oven, we found that the ratio of % I$_2$ loss to % solvent loss was $0.47 \pm 0.03$ ($n=3$). Therefore, 10% hexane loss equates to approximately 5% iodine loss, a level considered sufficiently low as to not be corrected for in this work.

On the basis of the performance of the two methods, it was concluded that hexane was the better solvent for the trapping of gaseous molecular iodine. Additional problems were also encountered with ethanol condensing in the fittings and tubing around the top of the solvent trap that militated against it. Using the prototype hexane trap with an ice/salt cold bath, a 1 cm spectrophotometer cell and a six hour trapping time (~10% solvent evaporation) the limit of detection is estimated to be around 20 to 40 ppbv.

The trap is thought to be selective for molecular iodine over volatile organohalogens, HOI and I$_3^-$, which all have absorption maxima at shorter wavelengths (Wall et al., 2003; Palmer and Lietzke, 1982; Paquette, 1985). However, there may be some overlap in absorption spectra between I$_2$ and the dihalogens IBr and ICl, which absorb 492 and 465 nm, respectively in CCl$_4$ (Augdahl and Klaeboe, 1965). However, the absorption cross sections of ICl and IBr are red-shifted and weaker than that of I$_2$, so that their concentrations would have to be similar to or in excess of I$_2$ to comprise a significant interference. Therefore care would need to be
Table 2. Comparison of prototype solvent trap properties.

<table>
<thead>
<tr>
<th></th>
<th>Hexane</th>
<th>Ethanol</th>
</tr>
</thead>
<tbody>
<tr>
<td>LoD&lt;sub&gt;spec&lt;/sub&gt;, nM</td>
<td>140</td>
<td>580</td>
</tr>
<tr>
<td>Evaporation rate&lt;sup&gt;b&lt;/sup&gt;, mL hr&lt;sup&gt;−1&lt;/sup&gt;</td>
<td>1.2</td>
<td>1.7</td>
</tr>
<tr>
<td>T. E.&lt;sup&gt;c&lt;/sup&gt;, %</td>
<td>86</td>
<td>67</td>
</tr>
<tr>
<td>Recovery&lt;sup&gt;d&lt;/sup&gt;, %</td>
<td>84±8</td>
<td>&gt;100</td>
</tr>
</tbody>
</table>

<sup>a</sup> 1 cm cell  
<sup>b</sup> Trap held at ∼ −9°C, flow rate of 0.05 L min<sup>−1</sup>  
<sup>c</sup> Calculated taking LoD (spec) as the minimum concentration  
<sup>d</sup> Average from 1 to 3 h

exercised in experiments or environments where relatively high concentrations of the interhalogens are expected to be present.

### 3.2.2 Optimised method

Using a 10 cm cell rather than a 1 cm cell, the LoD was reduced approximately ten-fold to 14 nM as expected. Derivatisation using leucocrystal violet gave a further increase in sensitivity of approximately 20-fold. However, the reproducibility of the absorbance for a given standard concentration using leucocrystal violet was worse, with relative standard deviations in the range 4 to 16%. Because of the poor reproducibility, the LoD achieved with leucocrystal violet derivatisation was higher than with hexane alone. With further work, it may be possible to develop a derivatisation method that yields reproducible results in organic solvent and a lower LoD.

Hexane evaporation rate as a function of gas flow was investigated for the improved solvent trap apparatus. At flow rates of up to 0.2 L min<sup>−1</sup> hexane showed no appreciable loss other than that caused by uptake to the glass frit of the gas inlet (Fig. 5). Above this flow rate, evaporative losses of 0.17 mL per 0.1 L min<sup>−1</sup> increase in flow rate were observed (Fig. 5). Therefore, an optimum gas flow rate of 0.2 L min<sup>−1</sup> was selected. At this gas flow, evaporative losses were linear with time with a hexane loss rate of 0.19 mL hr<sup>−1</sup>. For an initial hexane volume of 20 mL, under the optimised conditions it would take nearly 11 h for 10% of the trapping solvent to be lost. While the colder trapping temperature offers improved performance in terms of solvent evaporation, it also causes any moisture in the sample gas stream to condense and subsequently freeze on the scinted frit of the inlet, blocking the system. Therefore, great care must be taken to ensure the gas stream entering the trap is completely dry. While this is practicable under many experimental situations, it may limit the applicability of the traps for sampling ambient air. Use of a Nafion box drier upstream of the solvent traps led to

Fig. 5. Volume of hexane lost over 60 min at a trapping temperature of −50°C, as a function of nitrogen flow rate.

~60% I<sub>2</sub> loss, while chemical drying agents such as K<sub>2</sub>CO<sub>3</sub>, Drierite and molecular sieve gave 50 to 80% I<sub>2</sub> loss. The most suitable means of removing moisture but not molecular iodine from the gas stream identified to date has been two spiral condensers in series held at 0 and −10°C, but some iodine losses were still incurred.

Under the optimised trapping conditions, recovery of iodine from the permeation oven source was found to be 105±6% (Fig. 6), with no iodine detectable in the second trap (T. E.=100%). This demonstrates that the trapping performance of the optimised trap is improved from that of the basic trap set up given in Table 2.

Combining the improved spectrophotometric limit of detection with the enhanced trapping efficiency, longer trapping times and higher gas flow rates obtainable with the optimised trap yields a considerably improved atmospheric limit of detection for molecular iodine. For a trapping time of 10 h 45 min (10% solvent loss), the atmospheric LoD is of 69 pptv and the limit of quantification (LoQ; calculated using 10 times the standard deviation of the blank) is 230 pptv. The
LoD is comparable to peak concentrations of molecular iodine in coastal air observed at low tide (93 pptv; Saiz-Lopez et al., 2006b), suggesting the traps may be used in laboratory investigations (for example, seaweed chamber studies) at concentrations close to those observed in the atmosphere.

4 Conclusions

On the basis of the results presented here, it is concluded that amylose coated diffusion denuders are not suitable for the determination of molecular iodine in air. The recovery is very low and highly variable, and these problems are compounded by the difficulty of obtaining a sufficiently clean blank. If a denuder approach is required, the cyclodextrin/iodide coating (Huang and Hoffmann, 2009) is expected to give more reliable results.

Hexane or ethanol solvent traps with spectrophotometric detection of iodine species were found to give high recoveries and good reproducibility. Of the two solvents, hexane gave the better performance. Using the optimised hexane trap method, a gas phase limit of detection of 70 pptv is achievable, which is within the upper limit of ambient levels of molecular iodine observed in the coastal marine atmosphere (Saiz-Lopez et al., 2006b). The optimised hexane trap is thus suitable for measuring molecular iodine at atmospherically relevant concentrations in experimental investigations. If a means of drying the sample gas stream without removing molecular iodine can be found, it may also be possible to use hexane traps to measure molecular iodine in air where naturally occurring concentrations are very high, such as in the vicinity of kelp beds.

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