

# On the vertical distribution of boundary layer halogens over coastal Antarctica: implications for O<sub>3</sub>, HO<sub>x</sub>, NO<sub>x</sub> and the Hg lifetime

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Abstract. A one-dimensional chemical transport model has been developed to investigate the vertical gradients of bromine and iodine compounds in the Antarctic coastal boundary layer (BL). The model has been applied to interpret recent year-round observations of iodine and bromine monoxides (IO and BrO) at Halley Station, Antarctica. The model requires an equivalent I atom flux of  $\sim 10^{10}$  molecule cm<sup>-2</sup> s<sup>-1</sup> from the snowpack in order to account for the measured IO levels, which are up to 20 ppt during spring. Using the current knowledge of gas-phase iodine chemistry, the model predicts significant gradients in the vertical distribution of iodine species. However, recent ground-based and satellite observations of IO imply that the radical is well-mixed in the Antarctic boundary layer, indicating a longer than expected atmospheric lifetime for the radical. This can be modelled by including photolysis of the higher iodine oxides  $(I_2O_2, I_2O_3, I_2O_4 \text{ and } I_2O_5)$ , and rapid recycling of HOI and INO3 through sea-salt aerosol. The model also predicts significant concentrations (up to 25 ppt) of I<sub>2</sub>O<sub>5</sub> in the lowest 10 m of the boundary layer. Heterogeneous chemistry involving sea-salt aerosol is also necessary to account for the vertical profile of BrO. Iodine chemistry causes a large increase (typically more than 3-fold) in the rate of O<sub>3</sub> depletion in the BL, compared with bromine chemistry alone. Rapid entrainment of O<sub>3</sub> from the free troposphere appears to be required to account for the observation that on occasion there is little O<sub>3</sub> depletion at the surface in the presence of high concentrations of IO and BrO. The halogens also cause significant changes to the vertical profiles of OH and HO<sub>2</sub> and the NO<sub>2</sub>/NO ratio. The average  $Hg^0$  lifetime



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against oxidation is also predicted to be about 10h during springtime. An important result from the model is that very large fluxes of iodine precursors into the boundary layer are required to account for the observed levels of IO. The mechanisms which cause these emissions are unknown. Overall, our results show that halogens profoundly influence the oxidizing capacity of the Antarctic troposphere.

## 1 Introduction

Reactive halogen species (RHS=X, X2, XY, XO, OXO, HOX, XNO<sub>2</sub>, XONO<sub>2</sub> where X, Y is a halogen atom: Br, Cl, or I) play important roles in a number of atmospheric processes. One major impact is the depletion of ozone through catalytic cycles involving halogen radicals (e.g. Br, IO, BrO, Cl). In the 1970s, interest was focused mainly on stratospheric ozone depletion (e.g. Molina and Rowland, 1974; Stolarsky and Cicerone, 1974), whereas since the 1980s there have been reports of complete O<sub>3</sub> depletion events (ODEs) in the polar tropospheric boundary layer (BL) of both the Arctic and Antarctic (e.g., Bottenheim et al., 1986; Oltmans and Komhyr, 1986; Murayama et al., 1992; Kreher et al., 1997; Tuckermann et al., 1997; Wessel et al., 1998; Spicer et al., 2002; Brooks et al., 2006; Jones et al., 2006). These events occurred at polar sunrise in the spring and were explained by the influence of bromine-catalyzed chemical cycles (e.g. Barrie et al., 1988; McConnell et al., 1992; Tuckermann et al., 1997; Kreher et al., 1997; Frieß et al., 2004; Kaleschke et al., 2004). Bromine chemistry also plays a central role in the oxidation of atomic mercury (Hg<sup>0</sup>) in the polar atmosphere (Schroeder et al., 1998; Brooks et al., 2006), which seems to provide an important pathway for this element to enter



Fig. 1. Simplified scheme of bromine and iodine chemistry in the Antarctic boundary layer.

the Arctic food chain (Scott, 2001). It has been suggested that the presence of iodine can trigger this bromine chemistry and enhance the removal of ozone and mercury (Sander et al., 1997; Calvert and Lindberg, 2004a, b; O'Driscoll et al., 2006).

Major components of the atmospheric chemistry of bromine and iodine in the polar BL are illustrated in Fig. 1. The release of bromine-containing compounds proceeds through the so-called "bromine explosion", which involves bromide ions in brine-coated sea ice or snow being converted to gas-phase inorganic bromine (Br, BrO, HOBr etc.) in the BL (Hönninger and Platt, 2002). The presence of BrO has been reported by several groups such as Tuckermann et al. (1997) at Ny Ålesund, Hönninger et al. (2004a) at Hudson Bay, Canada and Brooks et al. (2006) at Barrow, Alaska. Ground-based measurements of BrO have also been made in coastal Antarctica using a passive differential optical absorption spectrometer (DOAS) during the spring of 1999 and 2000 (e.g. Frieß et al., 2004). Satellite observations have shown the presence of BrO in both the northern and southern polar regions (Richter et al., 1998; Wagner and Platt, 1998; Wagner et al., 2001; Richter et al., 2002; Hollwedel et al., 2004).

The vertical extent of bromine chemistry has been studied using a combination of the long path DOAS and multi-axis DOAS measurement techniques (Tuckermann et al., 1997; Martinez et al., 1999; Hönninger et al., 2004b). These studies showed that the enhanced BrO during ODEs was confined to the BL. It was also concluded that there was a very small vertical gradient of BrO within the BL (Hönninger and Platt, 2002; Hönninger et al., 2004b). These observations of BrO being well-mixed within the BL are consistent with a one-dimensional model study by Lehrer et al. (2004), which concluded that ODEs in the Arctic are caused by halogen chemistry confined to the BL by an inversion layer. This study also showed that emission from brine-covered sea ice was not sufficient to explain the observed ODEs, and suggested that recycling of halogen radicals through heterogeneous chemistry on aerosols was also required. A multiphase model study of bromine chemistry at polar sunrise concluded that  $O_3$  depletion cannot be simulated without the presence of heterogeneous halogen chemistry involving the snowpack (Michalowski et al., 2000).

Besides removing O<sub>3</sub>, RHS can also affect the oxidizing capacity of the troposphere in other ways. X and XO radicals reach a photochemical steady state during the day, essentially governed by reaction with  $O_3$  and photolysis (Fig. 1). The X atoms can react with non-methane hydrocarbons (NMHC), leading to hydrogen atom abstraction (analogous to OH). In fact, indirect measurements of the vertical extent of RHS have been carried out by measuring hydrocarbon destruction patterns (Solberg et al., 1996; Ramacher et al., 1999). Br atoms also recombine with Hg<sup>0</sup> to produce HgBr, which is relatively stable at the low temperatures of the polar spring BL; HgBr can then add a further radical (Br, I etc.) to yield mercury in its stable +2 oxidation state (Goodsite et al., 2004). Halogen oxides such as BrO and IO can also act directly as oxidizing radicals. For example, the rate of oxidation of dimethyl sulphide (DMS) by BrO can be up to an order of magnitude higher than the oxidation by OH (Saiz-Lopez et al., 2004; von Glasow et al., 2004).

RHS also affect the  $HO_x$  (i.e.,  $[HO_2]/[OH]$ ) and  $NO_x$  ratios (i.e.,  $[NO_2]/[NO]$ ) (e.g. von Glasow and Crutzen, 2003; Saiz-Lopez and Plane, 2004a; Bloss et al., 2005). XO

radicals oxidize NO to NO<sub>2</sub>, thus increasing the NO<sub>x</sub> ratio. In contrast, they react with HO<sub>2</sub> to yield HOX, which then photolyses efficiently to OH (particularly in the case of HOI), thus decreasing the HO<sub>x</sub> ratio (Bloss et al., 2005). A detailed study of  $BrO_x$ - $ClO_x$ -HO<sub>x</sub>- $NO_x$  chemistry during brominecatalysed ozone depletion events in the Arctic boundary layer has been performed by Evans et al. (2003).

Lastly, the role of iodine oxides in forming ultra-fine aerosol has been investigated by laboratory, field and modeling experiments (e.g. O'Dowd et al., 1998; Hoffmann et al., 2001; O'Dowd et al., 2002; Jimenez et al., 2003; Saiz-Lopez and Plane, 2004b; McFiggans et al., 2004; Burkholder et al., 2004; Sellegri et al., 2005; Saunders and Plane, 2005, 2006; Saiz-Lopez et al., 2006). The process of iodine oxide particle (IOP) production is thought to involve the recombination reactions IO+IO, IO+OIO, and OIO+OIO to yield  $I_2O_y$ , where y=2, 3 or 4, respectively. Further oxidation with  $O_3$ may then produce the most stable higher oxide, I<sub>2</sub>O<sub>5</sub>, which can then polymerize to form solid particles of  $I_2O_5$  (Saunders and Plane, 2005, 2006). Thus the presence of IO in the atmosphere points to the possibility of IOP formation; these particles could then provide condensation nuclei for other condensable vapours and grow to the point of becoming cloud condensation nuclei.

In Antarctica, the vertical column of IO was measured over a year using a ground-based differential optical absorption spectrometer (DOAS) (Frieß et al., 2001), and very recently from the SCIAMACHY instrument on the ENVISAT satellite (Saiz-Lopez et al., 2007b; Schöenhardt et al., 2007). However, the most comprehensive dataset of direct measurements of Antarctic IO (and BrO) in the BL was obtained during the Chemistry of the Antarctic Boundary Layer and Interface with Snow (CHABLIS) field measurement campaign at Halley Station, shown on the map in Fig. 2 (Saiz-Lopez et al., 2007a). In the present paper we investigate, using a 1-D model, the source strengths and likely vertical distributions of halogens within the Antarctic BL. The effects of the combined iodine and bromine chemistry on O<sub>3</sub> depletion, the HO<sub>x</sub> and NO<sub>x</sub> ratios and the lifetime of Hg within the BL are then explored.

#### 2 Model description

The Tropospheric Halogen Chemistry model (THAMO) is a one-dimensional chemical and transport model that uses time-implicit integration (Shimazaki, 1985). It has three main components: i), a chemistry module that includes photochemical, gas-phase and heterogeneous reactions; ii), a transport module that includes vertical eddy diffusion; and iii), a radiation scheme which calculates the solar irradiance as a function of altitude, wavelength and solar zenith angle. The continuity equations to account for the change in con-



**Fig. 2.** Map showing the location of Halley Station on the Brunt Ice Shelf in Antarctica during 2005. Antarctic coastline generated using the Coastline Extractor page hosted by the National Oceanic & Atmospheric Administration (NOAA)/National Geophysical Data Center, Marine Geology and Geophysics Division.

centration  $n_i$  of a species *i* at an altitude *z* and a time *t* are given by:

$$\frac{\partial n_i}{\partial t} = P_i - L_i n_i - \frac{\partial \phi_i}{\partial z} \tag{1}$$

where  $P_i$  and  $L_i$  are the production and loss rates of species *i*, respectively.  $\phi_i$  is the vertical flux due to eddy diffusion:

$$\phi_i = K_z \left[ \frac{\partial n_i}{\partial z} + \left( \frac{1}{T} \frac{\partial T}{\partial z} + \frac{1}{\overline{H}} \right) n_i \right]$$
(2)

where  $\overline{H}$  is the scale height of the atmosphere and  $K_z$  is the eddy diffusion coefficient (the second and third terms in Eq. (2) cast the flux in terms of the vertical gradient in mixing ratio, rather than just concentration (Shimazaki, 1985)). These equations are solved in the model with an integration time step of 2 min, which was chosen as a compromise between computational efficiency and compliance with the Courant-Levy criterion (Courant et al., 1928): reducing the time step to 15 s changed the calculated IO and BrO after 48 h by less than 1%. In this study, the model extends from the ground up to the upper boundary at 200 m with a spatial resolution of 1 m. To account for the entrainment of free tropospheric  $O_3$ , a flux into the top-most level of the model at the upper boundary can be turned on (or set to zero for the case of a stable BL). The lower boundary is the snowpack surface, where deposition from the gas phase and an upward trace gas flux from the snowpack can occur.

#### 2.1 Vertical transport parameterization

This is described by the turbulent diffusion coefficient  $K_z(z, t)$ , which will usually be a function of time t and height z. Near the surface, the layer where there is minimal change in momentum flux, turbulence is generated predominantly by wind shear. This "surface layer" is well described by Monin-Obhukov similarity theory, from which a function for the diffusion coefficient can be derived (Stull, 1988):

$$K_z(z,t) = \kappa . z.u * (t) \tag{3}$$

where  $\kappa$  is the von Kármán constant=0.4, z is the height above the surface and  $u^*(t)$  is the surface friction velocity. In the stable BL, where buoyancy can be neglected,  $u^*$  can be derived to a good approximation from:

$$\frac{\kappa U}{u*} = \ln\left(\frac{z}{z_0}\right) \tag{4}$$

where U is wind speed, z is the measurement height of the wind, and  $z_0$  is the surface roughness length. For Halley, long-term measurements indicate that  $z_o \sim 5 \times 10^{-5}$  m (King and Anderson, 1994). Note that this form of  $K_z(z, t)$  is relevant to the surface layer, where  $u^*(t)$  is virtually constant. Equation (4) implies that  $K_z(z, t)$  is linearly dependent on z, and tends to zero at the surface. This in turn would imply that a trace gas released at the surface, z=0, would never diffuse upwards, a situation which is clearly unrealistic. Hence, a surface condition is used which assumes that this form of  $K_{z}(z,t)$  is only valid for  $z \ge z_{0}$  (Stull, 1988). The model was run for two scenarios: stable neutral conditions, where  $K_z$  decreased from  $4.7 \times 10^3$  cm<sup>2</sup> s<sup>-1</sup> at 1 m to 2.0 cm<sup>2</sup> s<sup>-1</sup> at the top of the BL (200 m); and convective conditions, where  $K_z$  varied from 700 cm<sup>2</sup> s<sup>-1</sup> at 1 m to  $1 \times 10^4$  cm<sup>2</sup> s<sup>-1</sup> above 30 m.

## 2.2 Chemical scheme

The major objective of this modelling study is to simulate the IO observations, and thus explore the impact of iodine chemistry in the polar BL. The THAMO standard model contains the gas-phase iodine chemistry scheme we have used in previous modelling studies (McFiggans et al., 2000; Saiz-Lopez et al., 2006). The sources of the high concentrations of halogens at Halley are an open question. In fact, an explicit investigation of the microphysical processes that occur at the ice-atmosphere interface and lead to halogen release from ice surfaces is beyond the scope of this paper. The DOAS observations of IO and BrO indicate that oceanic air from the sea ice zone provides the primary halogen source (Saiz-Lopez et al., 2007a), although there is evidence that efficient snowpack recycling provides a secondary source. The rationale for this is that during CHABLIS significant concentrations of both IO and BrO were observed in air masses that had been in the continental boundary layer for several days (Saiz-Lopez et al., 2007a). This indicates (though does not prove) an efficient recycling of halogens following deposition onto the snowpack of either gas-phase species or wind-blown sea salt/frost flowers (since Halley is a coastal station – Fig. 2).

However, our knowledge about the details of these halogen sources is poor. Since horizontal transport is also not considered in this 1-D vertical model, we use the model to estimate the source strengths required to account for the observed IO and BrO concentrations (the model uses state-ofthe-art halogen gas-phase chemistry). In order to simulate the largest concentrations of IO and BrO observed (during spring), the emission fluxes of I<sub>2</sub> and Br<sub>2</sub> from the snowpack were required to be  $1 \times 10^{10}$  molecule cm<sup>-2</sup> s<sup>-1</sup> and  $1 \times 10^9$  molecule cm<sup>-2</sup> s<sup>-1</sup> at midday, respectively. The recycling in the snowpack is also assumed to be photochemical (for reducing iodine from the +3, +4 or +5 oxidation states). These fluxes are therefore assumed to depend on actinic flux, and hence their diurnal variation is described by a Gaussian profile peaking at midday.

Heterogeneous chemistry is treated in the following way. The uptake and subsequent hydrolysis of IONO<sub>2</sub>, HOI and INO<sub>2</sub> on aerosols produces HOI, which equilibrates between gas and aqueous phase according to its Henry's law solubility. The processing of aqueous HOI to IBr, ICl and I<sub>2</sub>, via reaction with Br<sup>-</sup>, Cl<sup>-</sup> and I<sup>-</sup>, respectively, takes only between 10 and 15 min in fresh sea-salt aerosol (McFiggans et al., 2000). The di-halogen molecules are insoluble and so are released rapidly to the gas phase. Hence, uptake of the inorganic iodine species onto aerosols is the rate-limiting step of the process (McFiggans et al., 2000). Note, however, that aged sea-salt aerosols become depleted in Br<sup>-</sup> and Cl<sup>-</sup>, which will slow down the aerosol processing time. Aged aerosols also become progressively acidified by uptake of HNO<sub>3</sub>, H<sub>2</sub>SO<sub>4</sub> and SO<sub>2</sub> (e.g. Fickert et al., 1999; von Glasow et al., 2002) which increases the rate of processing. However, for this study we do not explicitly treat the aqueous phase chemistry in the bulk of the aerosol. Instead, we assume that the limiting step for halogen heterogeneous recycling on aerosols is the first-order rate of uptake which we compute, using the free-regime approximation, for a number of gas phase species (Table 1 of the Supplementary Material: http://www.atmos-chem-phys.net/8/887/ 2008/acp-8-887-2008-supplement.pdf).

The gas-phase chemistry of IO in very clean air (NO<sub>x</sub> <30 ppt) is dominated by the reactions of IO with itself to form I<sub>2</sub>O<sub>2</sub> and OIO+I (Sander et al., 2006), IO+OIO to form I<sub>2</sub>O<sub>3</sub> (Gomez Martin et al., 2007), and OIO+OIO to form I<sub>2</sub>O<sub>4</sub> (Gomez Martin et al., 2007). These reactions proceed rapidly in the gas phase with rate coefficients of  $\sim 1 \times 10^{-10}$ ,  $5 \times 10^{-11}$  and  $1 \times 10^{-10}$  cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup>, respectively. The model also includes the formation of gasphase I<sub>2</sub>O<sub>5</sub> through a series of oxidation reactions of I<sub>2</sub>O<sub>2</sub>, I<sub>2</sub>O<sub>3</sub> and I<sub>2</sub>O<sub>4</sub> by O<sub>3</sub> up to the +5 oxidation state; we adopt a possible lower limit to the rate coefficients of these reactions of  $6 \times 10^{-13}$  cm<sup>3</sup> molecule<sup>-1</sup> s<sup>-1</sup> (Saunders and Plane, 2005). We also assume in the standard model that I<sub>2</sub>O<sub>2</sub>,  $I_2O_3$ ,  $I_2O_4$  and  $I_2O_5$  do not undergo photolysis or other reactions which would reduce them to IO or OIO, and that these molecules are lost by dry deposition to the snowpack and uptake onto pre-existing aerosol surfaces. Hence, any newly formed IO and OIO will be rapidly converted into higher-order iodine oxides, effectively limiting the atmospheric lifetime of both radicals. This means that when air masses arrived at Halley directly from the sea ice – when the measured IO and BrO concentrations were significantly higher (Saiz-Lopez et al., 2007a) – any IO produced from emissions over the sea ice would not have remained in the gas phase while being transported (transit time typically 2h). However, using the revised model chemistry where photolysis of  $I_xO_y$  is included (see below), IO could have been recycled in the gas phase during transit.

In addition to halogen chemistry, the THAMO model contains a set of odd-hydrogen, odd-nitrogen and methane chemical reactions, and a limited treatment of non-methane hydrocarbon (NMHC) chemistry. It also includes a detailed chemical scheme of reduced sulphur oxidation. Specific details of the iodine and bromine chemistry schemes are given in Sects. 4.1 and 4.2, respectively. A simplification of the chemistry described in the model is illustrated in Fig. 1, whereas the full reaction scheme is listed in Table 1 of the Supplementary Material (http://www.atmos-chem-phys. net/8/887/2008/acp-8-887-2008-supplement.pdf). The model is constrained with typical measured values of other chemical species via a time-step method. This requires that the field data were first averaged or interpolated to the two-minute frequency. Then, the concentrations of the constrained species were read in at the appropriate integration time step. Hence, these species were not assigned continuity equations and integrated, and their fluctuations do not determinate the size of the integration time-step. The species used to constrain the model were measured during the CHABLIS campaign, and had the following peak mixing ratios: [CO]=35 ppb; [DMS]=80 ppt; [SO<sub>2</sub>]=15 ppt; [CH<sub>4</sub>]=1750 ppb; [CH<sub>3</sub>CHO]=150 ppt; [HCHO]=150 ppt; [isoprene]=60 ppt; [propane]=25 ppt; [propene]=15 ppt (Jones et al., 2007a; Read et al., 2007). The model is also updated at every simulation time-step with measurements of temperature and relative humidity made during CHABLIS.

## 2.3 Photochemistry

The rate of photolysis of species is calculated on-line using an explicit two-stream radiation scheme from Thompson (1984). The irradiance reaching the surface is computed after photon attenuation through 50 1-km layers in the atmosphere as a function of solar zenith angle (SZA), location and timeof-year. The absorption cross-section and quantum yield data used in this model are summarized in Table 1 (Supplementary Material – http://www.atmos-chem-phys.net/8/887/ 2008/acp-8-887-2008-supplement.pdf). The rate of photolysis of species is computed by including snowpack albedo measurements (typical measured albedo=0.85) made with an actinic flux spectrometer during the CHABLIS campaign (Jones et al., 2007a).

#### 2.4 Sea ice/snowpack surface and aerosol uptake

Dry deposition to the surface occurs from the lowermost level of the model. Deposition was included for the following species: HOBr, HOI, HBr, HI, IONO<sub>2</sub>, BrONO<sub>2</sub>, OH, HO<sub>2</sub>, NO<sub>3</sub>, N<sub>2</sub>O<sub>5</sub>, CH<sub>3</sub>O<sub>2</sub>, HNO<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub>. The deposition flux of each gaseous species i was calculated as  $V_d$  $C_i$ , where  $V_d$  is the deposition velocity and  $C_i$  is the concentration of *i*. Deposition velocities were set to  $0.5 \text{ cm s}^{-1}$ . Here we assume that the ground is a flat surface. Similarly, the uptake of these species onto sea-salt aerosol surfaces was included in the model. The uptake of a gas species onto an aerosol surface was computed using the volumetric aerosol surface area (ASA) and the free molecular transfer approximation (Fuchs, 1964). The ASA used in this work is  $10^{-7}$  cm<sup>2</sup> cm<sup>-3</sup> (von Glasow et al., 2002), chosen to be typical of remote oceanic conditions, and is assumed constant in each vertical level of the model (i.e. the aerosol is well-mixed in the BL). Note that Halley is a remote coastal station and in the absence of aerosol measurements during CHABLIS we consider this ASA is the best estimate available. The uptake coefficients (see Table 1 of the Supplementary Material - http://www.atmos-chem-phys.net/8/887/ 2008/acp-8-887-2008-supplement.pdf) are taken from the recommendations of Sander et al. (2006) and Atkinson et al. (2000), unless otherwise stated.

## **3** DOAS observations at Halley Station

BL observations of IO and BrO were carried out from January 2004 to February 2005 using the technique of long-path DOAS (Platt, 1994; Plane and Saiz-Lopez, 2006). The measurements were performed during the CHABLIS campaign at Halley Station ( $75^{\circ}35'$  S,  $26^{\circ}30'$  W) situated on the Brunt Ice Shelf, about 35 m above sea level (Fig. 2). The ice edge is some 12 km north, 30 km west and 20 km south-west of the station. A detailed description of the CHABLIS campaign can be found elsewhere in this issue (Jones et al., 2007a).

The instrument was located in the Clean Air Sector Laboratory (CASLAB). An effective light path of 8 km at a height of 4 to 5 m above the snowpack (varying through the year as a result of snow accumulation) was set up between the CASLAB and a retro-reflector array positioned 4 km to the east. Further information on the instrumental design and spectral de-convolution procedures can be found in Plane and Saiz-Lopez (2006).

The measurements provide comprehensive observations of the diurnal and seasonal trends of both radicals at the base of the coastal Antarctic BL (Saiz-Lopez et al., 2007a). The IO and BrO concentrations exhibit a diurnal cycle with a clear



**Fig. 3.** Modelled diurnal variations of a), the IO mixing ratio profile, and b), the OIO mixing ratio profile during austral spring at Halley, using the standard model (see text for details). The mixing ratios at a height of 5 m agree well with the measurements made during the CHABLIS campaign using a DOAS with a beam  $\sim$ 5 m above the snowpack (Saiz-Lopez, 2007a).

dependence on solar irradiance. Higher concentrations were also measured in air that had passed over sea ice within the previous 24 hours. However, even in continental air that had spent at least four days over the interior of Antarctica, both radicals were still measured at mixing ratios up to  $\sim$ 6 ppt during sunlit periods, significantly above the detection limit of the instrument (1–2 ppt).

The seasonal trends of both radicals are remarkably similar, both in timing and absolute concentration. The radicals first appeared above the DOAS detection limit during twilight (August), and were then present throughout the sunlit part of the year. The peak mixing ratios of IO and BrO (20 ppt) were measured in springtime (October), followed by a possible smaller peak in autumn (March–April). Regarding the DOAS detection of OIO, the molecule was not conclusively measured above its DOAS detection limit (6–7 ppt) during 21 days of observations between January 2004 and February 2005. For more details about the DOAS measurements and correlation with meteorology during CHABLIS, see Saiz-Lopez et al. (2007a).

## 4 Results and discussion

## 4.1 Iodine chemistry

Here we use the model to investigate two important questions. First, what is the iodine source strength required to simulate the IO mixing ratios observed at the height of the DOAS measurements (i.e., 4–5 m above the snowpack)? Second, what is the vertical extent of the iodine chemistry in this environment: is its impact limited to the near surface or throughout the entire BL? During the CHABLIS campaign, the highest mixing ratios of IO, up to 20 ppt, were observed in springtime, following halogen activation mechanisms that, for the case of iodine, are not yet well understood. Figure 3 shows the modelled mixing ratios of IO and OIO in spring using the standard model of iodine chemistry, with an I atom flux out of the snowpack of  $1 \times 10^{10}$  molecule cm<sup>-2</sup> s<sup>-1</sup>. The calculated IO mixing ratio maximum of 16–18 ppt is in good accord with the DOAS observations. Sunrise is at 04:30 and sunset at 23:30 GMT. The days which match these conditions during CHABLIS are 20 and 21 October 2004, which are presented in Saiz-Lopez et al. (2007a). For OIO the mixing ratios are below 6 ppt, which was the instrumental detection limit during the CHABLIS campaign. The diurnally-averaged lifetime for  $I_xO_y$  is ~2 h. The modelled summer IO mixing ratio at midday peaking at 6 ppt, in accord with the DOAS observations, requires an I atom flux out of the snowpack of  $1 \times 10^9$  molecule cm<sup>-2</sup> s<sup>1</sup>.

Figure 3a and b show that both IO and OIO exhibit a strong vertical gradient in the BL: the concentration of IO at a height of 30 m is only 10% of that at 5 m. In the standard model, the transport of reactive iodine to the top of the BL occurs only via iodine recycling through sea-salt aerosol. However, the result of such a steep IO gradient is that the column abundance of the radical is predicted to be only  $7 \times 10^{11}$  molecule cm<sup>-2</sup> for the springtime simulation in Fig. 3. This is very much smaller than satellite observations of IO by SCIAMACHY in October 2005, where vertical columns  $>3 \times 10^{13}$  molecule cm<sup>-2</sup> were observed over Antarctic sea ice (Saiz-Lopez et al., 2007b; Schönhardt et al., 2008), or zenith-pointing DOAS measurements of the IO slant columns up to  $1 \times 10^{14}$  molecule cm<sup>-2</sup> made at the coastal Antarctic station of Neumayer (Frieß et al., 2001). Hence, vertical column measurements strongly suggest that IO does not have a steep gradient within the BL. The most likely explanation is that the standard model is missing a recycling mechanism from the higher iodine oxides to IO, which would increase the IO lifetime and allow it to be wellmixed in the BL.

The most likely recycling mechanism is photodissociation of  $I_xO_v$ . This would have two effects: i) increase the lifetime of IO<sub>x</sub> and therefore impact on the vertical distribution of inorganic iodine, ii) reduce the required atomic I flux from the snowpack into the gas phase to sustain the observed levels of IO. The model was therefore run with the photolysis of I2O2, I2O3, I2O4 and I2O5, set to a frequency at noon of  $1 \times 10^{-2}$  s<sup>-1</sup>. This photodissociation frequency is 2-3 times faster than  $J(IONO_2)$  at this location in spring, calculated using a new measurement of the IONO<sub>2</sub> photolysis cross section (Joseph et al., 2007). Preliminary measurements of the absorption cross sections of these higher iodine oxides (Gomez Martin et al., 2005) indicate that their magnitudes and long wavelength thresholds are similar to those of IONO2. Nevertheless, further laboratory work on the photochemistry of  $I_x O_y$  species is urgently needed to advance our understanding of this aspect of atmospheric iodine chemistry.

The inclusion of  $I_xO_y$  photolysis is now referred to as the *revised model*. Figure 4a shows the computed IO mixing ratio for the springtime scenario using the revised model



**Fig. 4.** (a) Modelled diurnal IO mixing ratio profile during austral spring at Halley, and (b) vertical profiles at noon of the main gasphase iodine species, using the revised model (photolysis of  $I_xO_y$ ). The mixing ratios at a height of 5 m agree well with the measurements made during the CHABLIS campaign using a DOAS with a beam ~5 m above the snowpack (Saiz-Lopez, 2007a).

(note the different height scale with respect to Fig. 3). The calculated IO at the top of the BL (200 m) is now 20% of that close to the surface, and the vertical column of  $5 \times 10^{12}$  molecule cm<sup>-2</sup> is now approaching reasonable accord with the satellite column measurement. Thus,  $J(I_xO_y)=1\times 10^{-2} \text{ s}^{-1}$  is the lower limit required to sustain high IO levels throughout the boundary layer. Figure 4b shows vertical profiles of the major gas-phase iodine species throughout the BL. Note that the main reservoir of iodine is  $I_2O_5$  with mixing ratios ranging from 25 ppt to 8 ppt from the surface to the top of the BL, respectively. Figure 5a illustrates a comparison between the measurements of IO and the model. The lack of agreement in the decrease of IO after midday may indicate that the chemistry in the model requires further development, or simply that this is a 1-D model applied to a spatially heterogeneous environment.

Figure 6 shows the modelled diurnal variation of the  $I_2O_5$  mixing ratio. Since the formation rate of IOPs is highly non-linear in  $I_xO_y$  mixing ratio (Saunders and Plane, 2005), most ultra-fine particles are predicted to form in the first 10 m above the snowpack, in the early afternoon (12:00–16:00 LT). This may be the source of ultra-fine particles that have been measured over sea-ice around Antarctica (Davison et al., 1996). Once IOPs form, there will be a competition between uptake onto pre-existing aerosol and further growth by the uptake of condensable vapours and coagulation.

The precise mechanism by which IO and OIO form IOPs is a subject of ongoing research, and IOP formation cannot be modelled with a great degree of confidence. There are two problems. The first is that the solid particles produced in laboratory studies have the stoichiometry of  $I_2O_5$ , but it is not clear whether this is produced in the gas phase by  $O_3$  oxidising  $I_2O_y$  (y<5), or by the polymerization of these lower oxides into particles which then rearrange and eject  $I_2$  (Saunders and Plane, 2005). The second is that although IOP production in the laboratory is a rapid process (Burkholder et



**Fig. 5.** Comparison of diurnal halogen oxides profiles at a height of 5 m measured by the DOAS instrument during the CHABLIS campaign and calculated using the revised model: (a) IO on 21 October 2004; (b) BrO on 20 October 2004.



Fig. 6. Modelled diurnal  $I_2O_5$  mixing ratio profile during the Antarctic springtime, using the revised model.

al., 2004; Saunders and Plane, 2006), there may be free energy barriers to the formation of small  $I_xO_y$  clusters which are not apparent at the relatively high concentrations of IO employed (compared with the atmosphere). The influence of humidity and other condensable vapours in the marine environment (e.g.  $H_2SO_4$ ) also remains to be explored. Since there were no aerosol measurements during CHABLIS, we



**Fig. 7. (a)** Two-day simulation of the boundary layer distribution of BrO during Antarctic springtime; (b) vertical profiles of the major gas-phase bromine species at noon on day 2. The mixing ratios at a height of 5 m agree well with the measurements made during the CHABLIS campaign using a DOAS with a beam  $\sim$ 5 m above the snowpack (Saiz-Lopez, 2007a).

have not explicitly examined IOP production in this study, and IOP nucleation was not included in the model (i.e.  $I_2O_5$ formation acts here as termination step for iodine chemistry in the standard model). It is worth noting that if polymerization is very rapid and without free energy barriers, then nucleation would compete with photolysis of  $I_xO_y$ , and an even larger iodine flux would be required to explain the IO observations.

IO was also observed at mixing ratios up to 6 ppt in air masses that had been over the continent for several days (Saiz-Lopez et al., 2007a). As mentioned earlier, this indicates an efficient iodine recycling mechanism capable of sustaining iodine radical chemistry over the snowpack. One possibility is the transport of sea-salt aerosol, and frost-flower fragments coated with sea-salt, from the ice front into the interior of the continent followed by deposition onto the snowpack; subsequent heterogeneous reactions would then recycle photolabile iodine to the gas phase, as we assume in THAMO. In fact, recent satellite measurements show that IO is widespread around coastal Antarctica, including areas over the snowpack (Saiz-Lopez et al., 2007b; Schönhardt et al., 2008). It is worth reiterating at this point that the sources of iodine are not well understood and hence in this study we have focused on finding the source strength of halogens needed, rather than on the nature of the source itself.

#### 4.2 Bromine chemistry

Figure 7a shows a two-day model simulation of the BrO vertical profile in springtime. The initial Br<sub>2</sub> flux from the snowpack is set to be only  $1 \times 10^9$  molecule cm<sup>-2</sup> s<sup>-1</sup> (i.e. a factor of 5 smaller than the I atom flux (see above)), because of the effect of the bromine autocatalytic mechanism. It can be seen that on the second day of the simulation the BrO mixing ratios at the height of the DOAS measurements are similar to the observations, peaking at ~20 ppt.

Figure 5b shows a comparison between the measured and modelled BrO at 5 m height. The observed diurnal asymmetry in the BrO is quite well reproduced by the model. During this season it is predicted that BrO will be better mixed within the boundary layer than IO (cf. Figs. 3 and 4). Figure 7b shows the noon vertical profiles of gas-phase bromine species during spring. Note that significant concentrations of inorganic bromine species reach the top of the BL, suggesting that BL ventilation under convective conditions may provide a source of inorganic bromine to the free troposphere. The predicted BrO column abundance for a 200 m BL height is  $1 \times 10^{13}$  molecule cm<sup>-2</sup>, while satellite observations have reported averaged tropospheric vertical columns of  $4 \times 10^{13}$  molecule cm<sup>-2</sup> in the austral spring over coastal Antarctica (Hollwedel et al., 2004). The difference could arise from the presence of a free tropospheric component in the satellite column measurement (Salawitch, 2006).

For the summertime case, a modelled BrO peak mixing ratio of 6 ppt at 4–5 m, in accord with the DOAS measurements, requires a Br<sub>2</sub> flux of  $2 \times 10^8$  molecule cm<sup>-2</sup> s<sup>-1</sup>. For both summer and spring model runs the predicted concentrations essentially track the solar irradiance profile and therefore peak at local noon when photodissociation of photolabile bromine (e.g. Br<sub>2</sub>, BrCl) is most efficient. In summer the BrO mixing ratio for SZA>90° does not decrease to zero since there is enough solar radiation reaching the surface during twilight for halogen activation to occur.

If heterogeneous reprocessing of bromine on sea-salt aerosol is switched off in the model, only a small fraction  $(\sim 10\%)$  of BrO at the surface will be transported by convection to the top of the BL, producing a pronounced vertical gradient. In contrast, including heterogeneous processing of bromine yields a vertical distribution of BrO that is better mixed through the BL, in good accord with observations in the Arctic which have shown elevated levels of BrO at heights tens of meters above the snow surface (e.g. Tuckermann et al., 1997; Martinez et al., 1999; Hönninger et al., 2004a). This shows that aerosols have a significant effect on the vertical distribution of inorganic bromine, although the actual efficiency of the halogen aerosol processing will depend on the vertical profile of the aerosol size distribution and chemical composition (in particular, the degree of halide ion depletion and the pH of the aerosol). A final point is that although the model with heterogeneous processing predicts better vertical mixing for BrO, there is still a gradient in BrO measurements where the BrO mixing ratio at the top of the boundary layer is  $\sim$ 50% lower than at the surface. This is in accord with observations of a negative vertical gradient in BrO reported by Avallone et al. (2003), where measurements of BrO in the Arctic were made using an in situ instrument near the surface (0.25-1 m) and a DOAS instrument higher up (20-200 m).

#### 4.3 Impact of bromine and iodine chemistry on ozone

For a case where the noon peak of BrO is 10 ppt, a simple photochemical box model shows the diurnally-averaged O<sub>3</sub> loss rate is  $0.14 \text{ ppb} \text{ h}^{-1}$  arising from bromine chemistry alone (the averaged background O<sub>3</sub> level during spring was  $\sim 20 \text{ ppb}$ ). An O<sub>3</sub> depletion rate of 0.25 ppb h<sup>-1</sup> is calculated for a noon peak of IO=10 ppt due to iodine chemistry only. When the halogens couple through the cross reaction

$$BrO + IO \rightarrow OIO + Br(R1a)$$
 (R1a)

$$\rightarrow$$
 I + Br + O<sub>2</sub> (R1b)

the total  $O_3$  loss rate is 0.55 ppb h<sup>-1</sup>, more than the sum of each component in isolation (Saiz-Lopez et al., 2007a). Thus, the  $O_3$  depletion rate is almost four times as fast as that predicted from bromine chemistry alone, demonstrating the central role that iodine plays in  $O_3$  depletion during Antarctic polar sunrise.

Figure 8a illustrates a THAMO run during springtime, for the case of a strong temperature inversion and stably buoyant BL, with no  $O_3$  entrainment from the free troposphere. It can be seen that after 36 hours the computed  $O_3$  levels in the BL drop to below the instrumental detection limit (~1 ppb) due to the combined impact of iodine and bromine chemistry. This is similar to the complete removal of  $O_3$ , which has been reported in the Arctic when a strong temperature inversion and high levels of BrO occur (e.g. Barrie et al., 1988; McConnell et al., 1992; Hönninger and Platt, 2002).

However, during the CHABLIS campaign it was observed that elevated levels of IO and BrO occurred without complete removal of O<sub>3</sub>. We now use the THAMO model to examine whether the occurrence of high halogen oxide concentrations for a prolonged period of time without complete  $O_3$  destruction can be explained by entrainment of  $O_3$ -rich air from aloft. Figure 8b shows the calculated O<sub>3</sub> profile with a downward flux from the free troposphere of O3 of  $3 \times 10^{11}$  molecule cm<sup>-2</sup> s<sup>-1</sup>, and a convective BL. The predicted O<sub>3</sub> levels after 36 h have now only decreased by 20%, consistent with the observations (although after 48 hours the  $O_3$  levels have decreased by ~50%, so that significant  $O_3$ does eventually occur).  $O_3$  is well-mixed throughout the BL, without a pronounced vertical gradient, as observed by Arctic and Antarctic ozonesonde observations during ODEs (e.g. Wessel et al., 1998; Tarasick and Bottenheim, 2002).

Note that the model does not include a parameterization of BL ventilation. Hence, the halogen flux used in the model can be considered as a *lower limit* under convective boundary layer conditions. In addition to the difficulty of describing turbulent transport into the free troposphere, knowledge of halogen concentrations (and speciation) in the lower free troposphere is needed to properly quantify exchange with the BL, and the distribution of bromine and iodine in the free troposphere still remains an open question. Therefore, for a scenario with a prescribed halogen flux constant during each



**Fig. 8.** Diurnal variation of the boundary layer  $O_3$  profile in the presence of springtime BrO and IO levels for two scenarios: (a) no entrainment of  $O_3$  and (b) entrainment of  $O_3$  from the free troposphere.

sunlit period the model does not reach steady state, in part because BL ventilation is not included. However, an approach to steady-state conditions in the same air mass over several days is unlikely to occur in coastal Antarctica. A final point to consider here is an alternative explanation for the absence of complete O<sub>3</sub> destruction is the presence of high halogen concentrations. This is that air masses were being sampled where IO and BrO had been released so recently that significant O<sub>3</sub> depletion had not yet occurred. However, while this might explain the observations of O<sub>3</sub> and halogen oxides in air from the coastal sector, simultaneously high O<sub>3</sub> and halogen oxides were also seen in air masses that had been over the continent for several days (Saiz-Lopez et al., 2007a).

#### 4.4 Impact of halogens on HO<sub>x</sub>, NO<sub>x</sub> and the Hg lifetime

During January and February 2005 (austral summer), in situ measurements of OH and HO<sub>2</sub> were performed using the FAGE technique (Bloss et al., 2007). The measurements were made at the same height above the snowpack ( $\sim 5 \text{ m}$ ) as the DOAS beam. Typical peak noon values of  $4 \times 10^{-2}$  ppt and 1.50 ppt were measured for OH and HO<sub>2</sub> respectively. IO and BrO react with HO2 to form HOI and HOBr, whose subsequent photolysis produces OH, thus reducing the HO<sub>2</sub>/OH ratio. In order to model the OH and HO<sub>2</sub> mixing ratios, we use the  $J(O^{1}D)$  values measured during CHABLIS (typical noon photolysis frequency of  $4 \times 10^{-5}$  s<sup>-1</sup>) and the measured mixing ratios of relevant species (see Sect. 2.2). Figure 9a and b show the effect of halogen chemistry on the vertical distributions of OH and HO<sub>2</sub>, respectively, for runs with the standard model, the revised model, and no halogens. At the height of the measurements, the model runs without halogen chemistry over-predict HO<sub>2</sub> by  $\sim$ 3 times the measured concentration, and underpredict OH by  $\sim 40\%$ . Thus, the modeled HO<sub>x</sub> (HO<sub>2</sub>/OH) ratio is 115, compared with the measured ratio of 37. When halogen chemistry is included (i.e. typical summer noon BrO and IO mixing ratios of 5 ppt), the calculated OH and HO<sub>2</sub> levels and diurnal profile are in very



Fig. 9. Vertical profiles of (a) modelled OH and (b) modelled  $HO_2$  for three scenarios: without halogen chemistry, with halogens (standard model), and including photolysis of  $I_xO_y$  species (revised model).

good agreement with the observations: the modeled ratio is now 33. Figure 9 also shows that above about 10 m height there is a significant difference in the modeled OH and  $HO_2$ profiles depending on whether  $I_xO_y$  photochemistry is included.

NO and NO<sub>2</sub> were also measured during the Antarctic summer period with noon average mixing ratios of 14 ppt and 7.5 ppt, respectively, so that the NO<sub>x</sub> (NO<sub>2</sub>/NO) ratio was 0.5. IO and BrO increase the NO<sub>2</sub>/NO ratio by converting NO to NO<sub>2</sub>. NO<sub>x</sub> is produced in the model by photochemistry of snowpack nitrate ions. The resulting summer and spring NO<sub>x</sub> fluxes from the snowpack were set to  $2 \times 10^8$  and  $1.2 \times 10^7$  molecule cm<sup>-2</sup> s<sup>-1</sup>, respectively, which were measured during CHABLIS (Jones et al., 2007b). Ventilation of NO<sub>x</sub> to the free troposphere, or entrainment from

the free troposphere into the BL, were not included. The model was constrained with summer  $J(NO_2)$  (typical noon maximum values of  $0.015 \text{ s}^{-1}$ ) values measured with an actinic flux spectrometer during CHABLIS. Without halogens in the model, the NO<sub>x</sub> ratio at noon is only 0.3, whereas when halogens are included the ratio is 0.54.

Lastly, we use the measured XO concentrations and the XO:X ratio from THAMO to assess the impact of bromine and iodine chemistry on the lifetime of elemental mercury  $(Hg^0)$  over coastal Antarctica. To calculate the lifetime of  $Hg^0$  against oxidation to  $Hg^{II}$ , we use the formalism given in Goodsite et al. (2004):

$$\tau = \frac{(k_{-2} + k_3[Br] + k_4[I])}{k_2[Br](k_3[Br] + k_4[I])}$$
(5)

where, at a pressure of 1 bar in air,

$$k(\text{Hg} + \text{Br} \rightarrow \text{HgBr}, 180-400 \text{ K}) =$$
  
 $1.1 \times 10^{-12} (T/298 \text{ K})^{-2.37} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$  (R2)

$$k(\text{HgBr} \rightarrow \text{Hg} + \text{Br}, 180-400 \text{ K}) =$$
  
 $1.2 \times 10^{10} \exp(-8360/T) \text{ s}^{-1}$  (R-2)

$$k(\text{HgBr} + \text{Br} \rightarrow \text{HgBr}_2, 180-400 \text{ K}) =$$
  
 $1.4 \times 10^{-10} + 2.6 \times 10^{-13} \cdot T - 8.6 \times 10^{-16} \cdot T^2$  (R3)

$$k(\text{HgBr} + \text{I} \rightarrow \text{HgBrI}, 180-400 \text{ K}) =$$
  
 $1.2 \times 10^{-10} + 4.2 \times 10^{-13} \cdot T - 1.0 \times 10^{-15} \cdot T^2$  (R4)

During springtime, the model shows that the diurnallyaveraged mixing ratio of BrO of ~8 ppt measured by DOAS would have coexisted in steady state with a Br mixing ratio of  $\sim 0.7$  ppt for a springtime diurnally-averaged O<sub>3</sub> mixing ratio of 12 ppb. The computed lifetime of Hg<sup>0</sup>, against oxidation by bromine chemistry alone (Reactions 2-3) is then about  $\sim$ 13 h at an average temperature of 260 K. The average IO measured by DOAS during springtime ( $\sim 8 \text{ ppt}$ ) would have coexisted with a calculated I mixing ratio of  $\sim$ 5 ppt . The smaller IO/I ratio, compared to that of BrO/Br, arises from the self-reaction of IO to form OIO+I and I<sub>2</sub>O<sub>2</sub> (Sander et al., 2006), and from the thermal decomposition of  $I_2O_2$  to OIO+I and IO+IO. Including the role of atomic I through Reaction (R4) leads to a 40% reduction of the Hg lifetime. In addition, iodine chemistry decreases the calculated BrO/Br ratio from 11 to 5 via conversion of BrO back to Br through Reaction (R1).

The Hg<sup>0</sup> lifetime is predicted to be reduced to only 2 h under conditions of high XO mixing ratios (i.e. 20 ppt), as observed as Halley during springtime, when the Br and I mixing ratios would have been 4 ppt and 13 ppt, respectively. Note that these calculations are made for high levels of halogens *without* severe  $O_3$  depletion, as observed on occasions during CHABLIS (Saiz-Lopez et al., 2007a). If  $O_3$  were completely removed, then the BrO/Br and IO/I ratios would decrease to 0.15 and 0.01, respectively, and the Hg<sup>0</sup> lifetime would be reduced to ~10 min. Our measurements of BrO and IO therefore indicate that there should be sustained removal of Hg into the snowpack throughout the sunlit period in coastal Antarctica. Unfortunately, mercury measurements were not made during CHABLIS, but should certainly form a central component of any future campaigns in coastal Antarctica.

## 5 Summary and conclusions

The THAMO chemical transport model has been used to investigate the vertical gradients of halogens in the Antarctic coastal BL with a parameterization of the vertical transport under stable and convective BL conditions. The standard iodine chemistry model predicts a very steep gradient of iodine gas-phase species in the first 20 m of the BL. The result is that, although the predicted IO concentration at 5 m height is in good accord with the boundary layer DOAS observations during the CHABLIS campaign, the modelled column abundance is much less than the IO column abundance measured from ground-based and satellite-borne instruments. This indicates that IO is much better mixed in the BL than predicted by the standard model. We have therefore revised the standard model to including the photolysis of the higher iodine oxides  $I_2O_v$ , where y=2–5. These reactions recycle IO efficiently throughout the BL, producing much better agreement between the model and observations. The photochemistry of these species needs to be studied in the laboratory.

The revised THAMO model was then used to try and explain the surprising occurrence of close-to-average  $O_3$  concentrations in the presence of high levels of IO and BrO. Within the constraints of a 1-D model, this can be achieved by replenishment of  $O_3$  through entrainment from the free troposphere, to ensure that complete depletion of  $O_3$  does not occur. The model then predicts a well-mixed vertical profile for  $O_3$  within the BL, which is in agreement with ozonesonde observations. In some cases the simultaneous observation of high levels of halogen oxides and  $O_3$  may also be explained by the very recent injection of the halogens into an air mass moving from the sea ice zone to Halley Bay.

The model is able to account for the measured perturbations in the  $HO_x$  and  $NO_x$  ratios by the observed concentrations of IO and BrO, and demonstrates that the presence of high atomic I concentrations leads to a significant enhancement in the oxidation rate of elemental  $Hg^0$  by atomic Br.

Finally, the model requires very large fluxes of iodine precursors from the snowpack in order to account for the boundary layer observations of IO. This indicates an efficient recycling mechanism in the snowpack which is still to be understood. Acknowledgements. The authors would like to thank R. L. Jones and R. A. Cox (University of Cambridge), and R. W. Saunders (University of Leeds) for helpful discussions. This work was supported by the NERC's Antarctic Funding Initiative. We thank the School of Chemistry, University of Leeds for a research fellowship (A. Saiz-Lopez) and a research studentship (A. S. Mahajan).

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