Bromine measurements in ozone depleted air over the Arctic Ocean

J. A. Neuman1,2, J. B. Nowak1,2, L. G. Huey3, J. B. Burkholder2, J. E. Dibb4, J. S. Holloway1,2, J. Liao3, J. Peischl1,2, J. M. Roberts2, T. B. Ryerson2, E. Scheuer4, H. Stark1,2, R. E. Stickel3, D. J. Tanner3, and A. Weinheimer5

1Cooperative Institute for Research in Environmental Sciences, University of Colorado at Boulder, Boulder, CO, USA
2Earth System Research Laboratory, NOAA, Boulder, CO, USA
3School of Earth and Atmospheric Science, Georgia Institute of Technology, Atlanta, GA, USA
4Institute for the Study of Earth, Oceans, and Space, University of New Hampshire, Durham, NH, USA
5Atmospheric Chemistry Division, National Center for Atmospheric Research, Boulder, CO, USA

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Correspondence to: J. A. Neuman (andy.neuman@noaa.gov)

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Abstract

In situ measurements of ozone, photochemically active bromine compounds, and other trace gases over the Arctic Ocean in April 2008 are used to examine the chemistry and geographical extent of ozone depletion in the arctic marine boundary layer (MBL). Data were obtained from the NOAA WP-3D aircraft during the Aerosol, Radiation, and Cloud Processes affecting Arctic Climate (ARCPAC) study and the NASA DC-8 aircraft during the Arctic Research of the Composition of the Troposphere from Aircraft and Satellites (ARCTAS) study. Fast (1 s) and sensitive (detection limits at the low pptv level) measurements of BrCl and BrO were obtained from three different chemical ionization mass spectrometer (CIMS) instruments, and soluble bromide was measured with a mist chamber. The CIMS instruments also detected Br$_2$. Subsequent laboratory studies showed that HOBr rapidly converts to Br$_2$ on the Teflon instrument inlets. This detected Br$_2$ is identified as active bromine and represents a lower limit of the sum HOBr+Br$_2$. The measured active bromine is shown to likely be HOBr during daytime flights in the arctic. In the MBL over the Arctic Ocean, soluble bromide and active bromine were consistently elevated and ozone was depleted. Ozone depletion and active bromine enhancement were confined to the MBL that was capped by a temperature inversion at 200–500 m altitude. In ozone-depleted air, BrO rarely exceeded 10 pptv and was always substantially lower than soluble bromide that was as high as 40 pptv. BrCl was rarely enhanced above the 2-pptv detection limit, either in the MBL, over Alaska, or in the arctic free troposphere.

1 Introduction

Low ozone mixing ratios over the Arctic Ocean region in the spring have been observed for nearly three decades (Oltmans, 1981). Model studies and measurements of ozone, halogen compounds, and volatile organic compounds that react with halogens have shown that bromine from the Arctic Ocean causes ozone depletion in the springtime arctic boundary layer (Simpson et al., 2007, and references therein).
Atmospheric concentrations of photochemically active bromine species are elevated by heterogeneous chemistry that converts condensed phase bromine to volatile and photolabile species such as Br$_2$ (Fan and Jacob, 1992). Photolysis of either Br$_2$ or HOBr then forms Br atoms. In seawater, chloride is 700 times more abundant than bromide. Yet in the atmosphere over the arctic in spring, the selective liberation of bromine from sea ice causes Br atoms to exceed Cl atoms by factors of hundreds, as shown by measurements of VOC ratios (Jobson et al., 1994; Impey et al., 1999; Keil and Shepson, 2006). Br atoms react with ozone by

\[ \text{Br} + \text{O}_3 \rightarrow \text{BrO} + \text{O}_2, \]  

(R1)

and catalytic ozone destruction occurs if Br atoms are regenerated by BrO self-reaction or reaction with HO$_2$. BrO produced from the reaction of Br and ozone has been observed in arctic regions and provides further confirmation of bromine-induced ozone destruction (Hausmann and Platt, 1994). In the springtime arctic, a stable boundary layer isolates the air in the lowest several hundred meters of the atmosphere from the free troposphere above. This reduces dilution of ozone-depleted air in the marine boundary layer (MBL) with ozone-rich air in the free troposphere and further allows bromine species to build to high levels.

Most arctic studies of ozone depletion have been conducted over land from coastal ground sites near ocean source regions of halogens (Barrie et al., 1988). Back trajectories and ozone observations have shown that the predominant transport pathway for ozone depleted air arriving at arctic ground sites is over the Arctic Ocean, suggesting that ozone depletion chemistry occurs over the ocean (Bottenheim and Chan, 2006). The few measurements obtained directly over Arctic Ocean ice have shown that ozone depletion is persistent in the spring (Hopper et al., 1998). Sustained ozone depletion over arctic ice has been observed at high latitudes (81–88°N) from a ship that drifted while frozen in the Arctic Ocean (Bottenheim et al., 2009). Ozone depletion and bromine chemistry in the lower troposphere at northern high latitudes also have been studied using satellite measurements of BrO combined with a chemical transport model (Zeng et al., 2003).
Measurements from aircraft, which permit investigations over these ocean regions where ozone depletion originates, also have shown that ozone depletion is ubiquitous over the Arctic Ocean in spring (Leaitch et al., 1994; Hopper et al., 1998). Large regions with both major (O$_3$ < 4 ppbv) and partial (4 < O$_3$ < 20 ppbv) ozone depletions were detected by aircraft measurements over the Canadian Arctic, the Arctic Ocean north of Greenland, and occasionally at more southern latitudes (Ridley et al., 2003). In that study, ozone depletion was observed regularly and was associated with increased levels of soluble bromide. The largest levels of soluble bromide were in the northern-most ozone-depleted air, but interestingly not in the layers with the least ozone. The relationship between ozone and bromine is investigated further here.

Arctic ozone depletion chemistry needs to be better understood, since it may have important regional environmental consequences. The presence of bromine and the depletion of ozone alter the oxidizing characteristics of the atmosphere to make the arctic a sink for some compounds. Elemental mercury (Hg), which is long-lived and distributed globally, is oxidized to reactive forms by Br atoms (Donohoue et al., 2006). This reactive gaseous mercury deposits to surfaces more readily than elemental Hg and could collect in arctic regions (Steffen et al., 2008, and references therein).

Although severe ozone depletion caused by halogen chemistry is usually confined to a thin layer of the atmosphere over the polar oceans and occurs primarily in the spring, the same processes may occur throughout the year over all oceans. Understanding the processes that cause ozone depletion in the arctic can reveal whether these reactions substantially affect global ozone concentrations. Ozone reductions caused by reactive halogens have been modeled to occur throughout the troposphere (Yang et al., 2005) and in polluted marine air masses transported over the ocean (Sander and Crutzen, 1996). Experimental verification of these model results regarding halogen chemistry is challenging. Although several bromine-containing compounds are expected to be elevated in air masses where bromine-induced ozone destruction is important, measurement capabilities for low levels of inorganic bromine-containing molecules are limited primarily to BrO (Read et al., 2008; Platt and Honninger, 2003). In particular,
HBr, BrO, HOBr, Br₂, and Br are predicted to be elevated at some point during ozone depletion, and their abundance indicates the temporal evolution of the ozone depletion chemistry (Lehrer et al., 2004; Calvert and Lindberg, 2003; Martinez et al., 1999; Sander and Crutzen, 1996; Evans et al., 2003; Fan and Jacob, 1992). Extending the limited geographical range of bromine observations and number of measured bromine compounds will provide experimental evidence for the effect of reactive halogens upon global lower tropospheric ozone abundance.

This study examines ozone depletion and bromine abundance over the Arctic Ocean near Alaska. New techniques for rapidly detecting low concentrations of Br₂, HOBr, BrO, and BrCl are developed. The extent and causes of ozone depletion in the springtime arctic are investigated with expanded capabilities for speciating inorganic bromine.

2 Measurements

Aircraft measurements were performed over the Arctic Ocean in April 2008 as part of the Polar Study using Aircraft, Remote Sensing, Surface Measurements, and Models of Climate, Chemistry, Aerosols, and Transport (POLARCAT) during the International Polar Year. Data were obtained from the NOAA WP-3D aircraft during the Aerosol, Radiation, and Cloud Processes affecting Arctic Climate (ARCPAC) experiment and the NASA DC-8 aircraft during the Arctic Research of the Composition of the Troposphere from Aircraft and Satellites (ARCTAS) experiment. Flights targeted many different air mass types to address the multiple goals of the ARCPAC (Brock et al., 2010) and ARCTAS studies (Jacob et al., 2009). A small fraction of the WP-3D (20%) and DC-8 (10%) arctic flights were dedicated to sampling in or near the MBL over the Arctic Ocean to examine halogen chemistry. The WP-3D measurements are described in detail here and the data are available at http://www.esrl.noaa.gov/csd/arcpac/. The DC-8 measurements are briefly summarized, and the data are available at http://www-air.larc.nasa.gov/data.htm.
2.1 Aircraft sampling platforms

The NOAA WP-3D aircraft flew over the Arctic Ocean on five flights that departed from Fairbanks, Alaska. The flights were performed during the day on 12, 15, 18, 19, and 21 April 2008 (Fig. 1). Over sea ice, the WP-3D typically flew between 100–160 m altitude, with occasional dips to 60 m. Air was sampled above the MBL in frequent altitude profiles and higher altitude flight legs up to 7.4 km. Approximately 20–90 min of flight time was dedicated to sampling in the MBL on each eight hour flight. Most of the MBL studies were conducted over the Beaufort and Chukchi Seas adjacent to the northern coastline of Alaska.

The NASA DC-8 aircraft conducted seven flights in the arctic region between 4 and 17 April 2008. Flight altitudes in the MBL over the Arctic Ocean were typically 100 m. Although the DC-8 devoted less time to studies of halogen chemistry in the MBL, it covered a much larger area, with measurements at high latitudes all the way to the pole and further east over the Canadian Arctic.

In situ measurements were obtained from fast-response, high-sensitivity instruments. Complete instrument descriptions, as well as field intercomparison results, are detailed in ARCPAC and ARCTAS overview papers (Brock et al., 2010; Jacob et al., 2009). On the WP-3D, ozone was measured by chemiluminescence. The data were averaged to once per second with a 1-sigma uncertainty of ±(4%+0.05) ppbv. Additional fast response measurements used to characterize the air masses included SO₂ (uncertainty ±(13%+0.55) ppbv), CO (uncertainty ±3%), NO (uncertainty ±(8%+0.02) ppbv), NO₂ (uncertainty ±(10%+0.04) ppbv), and NOₓ (uncertainty ±(12%+0.05) ppbv). Measurements of actinic flux between 280 and 690 nm were performed both in the upward and downward looking directions at 1 nm resolution using two grating spectrometer/CCD systems (Stark et al., 2007). Photolysis rates for bromine compounds were calculated from the actinic flux measurements using literature values for the absorption cross-sections and quantum yields (Sander et al., 2006).
On the DC-8, soluble bromide was measured with a mist chamber. The species responsible for bromide in the mist chamber are uncertain but include inorganic bromine compounds HBr, HOBr, particulate bromide, and possibly BrO and Br₂ (Ridley et al., 2003; Dibb et al., 2010; Liao et al., 2010). Each measurement averaged over 1.5 min, with a 3-pptv detection limit and a 20–30% uncertainty. Ozone was measured by chemiluminescence with 5% uncertainty, <1 s time response, and <0.1 ppbv imprecision. The chemical ionization mass spectrometers (CIMS) (Huey, 2007) used to measured halogen species on both aircraft are described in detail below.

2.2 CIMS halogen detection

Fast response measurements of reactive halogen compounds were achieved using newly developed detection techniques. Gas phase reactive halogen compounds were measured by a CIMS instrument on the DC-8 and by two independent CIMS instruments on the WP-3D. The CIMS techniques used here evolve from instruments that measure atmospheric SO₂ (Kim et al., 2007) and peroxyacyl nitrates (Slusher et al., 2004), with modifications that exploit laboratory studies of ion-molecule reactions involving halogen compounds (Huey et al., 1995; Roberts et al., 2008). Reagent ions are made in-flight and mixed with ambient air in a low-pressure flow tube, where ion-molecule reactions form unique product ions used to determine ambient mixing ratios. Br₂, BrCl, and BrO are measured using either I⁻ or SF₆⁻ reagent ions. The reagent and product ions are mass-filtered with quadrupole mass spectrometers and then counted with electron multiplier detectors. The air is admitted into the instruments through separate inlets by pulling flows of several standard liters per minute (slm) through heated (≈30 °C) tubes. All wetted inlet parts are made from perfluoralkoxy (PFA) Teflon (abbreviated as Teflon below). The aircraft inlets are housed in winglets that extend perpendicular to the skin of the aircraft and beyond the aircraft boundary layer, with inlet tips that are 0.4 cm ID Teflon tubes. Total inlet lengths are less than 1 m for each instrument, resulting in inlet residence times of tens of ms.

Instrument operation and diagnostics, including sampling, calibration, and zeroing,
are given below for the WP-3D CIMS that uses I\(^-\) ion chemistry. I\(^-\) reagent ions are produced by flowing 2 slm N\(_2\) doped with 2.5 ppmv CH\(_3\)I through a \(^{210}\)Po radioactive source. The CH\(_3\)I is obtained from a 5 sccm flow of 0.1% CH\(_3\)I in N\(_2\). This ion flow is combined with 1.6 slm ambient air in a 46-cm long flow tube at 20 Torr. Adding a 20-sccm N\(_2\) flow bubbled through distilled H\(_2\)O humidifies the flow tube. Humidification increases the abundance of I\(^-\)·H\(_2\)O cluster ions, which provide better sensitivity to halogen detection than I\(^-\) alone.

I\(^-\)·H\(_2\)O reacts with Br\(_2\), BrCl, Cl\(_2\), and BrO to form clusters of I\(^-\) with the halogen molecules. Since atomic I has no isotopes and a mass given by 127 atomic mass units (amu), and atomic Br and Cl have two naturally occurring isotopes (\(^{79}\)Br (50.7%), \(^{81}\)Br (49.3%), and \(^{35}\)Cl (75.8%), \(^{37}\)Cl (24.2%)), the cluster ions have simple but distinct isotope ratios. For example, I\(^-\)·Br\(_2\) clusters have three isotopes: 285 amu (25%), 287 amu (50%), and 289 amu (25%) (Fig. 2). These isotope patterns facilitate unambiguous identification of the ions. However, instrument sensitivity is reduced, since only one mass can be measured at a time, and isotopes distribute the ion products over several masses simultaneously. Ions are monitored in flight in the following sequence: 127 amu (I\(^-\)) for 100 ms, 145 amu (I\(^-\)·H\(_2\)O) for 100 ms, 224 amu (I\(^-\)·BrO) for 500 ms, 243 amu (I\(^-\)·BrCl) for 500 ms, and 287 amu (I\(^-\)·Br\(_2\)) for 500 ms. Consequently, BrO, BrCl, and Br\(_2\) mixing ratios are determined once every 2 s. I\(^-\) and its water clusters are orders of magnitude more abundant than any other negative ions in the flow tube, with I\(^-\) and I\(^-\)·H\(_2\)O ion count rates typically 400 kHz (Figs. 2 and 4).

Entirely different ion chemistry and resulting product ions are used for halogen detection on the DC-8 and in the other WP-3D CIMS instrument. Nitrogen doped with SF\(_6\) is used to produce the SF\(_6\)^- reagent ions. SF\(_6\)^- is a reactive ion that was employed with the primary goal of measuring SO\(_2\), but it also reacts efficiently with halogen compounds to form negatively charged halogens by charge transfer. For example, SF\(_6\)^- reacts with BrO to form BrO\(^-\). The operation of these two instruments is not detailed here, but they perform similarly to the WP-3D I\(^-\) CIMS, with in-flight calibrations and background determinations as described below.
2.3 Sensitivity

Instrument sensitivity is determined in-flight by standard addition calibrations of Br$_2$. The Br$_2$ source is a commercial permeation device (Kin-tek USA). 20 sccm of N$_2$ continuously flows around a Br$_2$ permeation tube that is pressure controlled at 2 atm and temperature controlled at 25°C. The Br$_2$ output was measured to be 12 ng/min by optical absorption at 404 nm. At 40°C, the measured output was 56 ng/min, and 14% higher than the manufacturer’s gravimetric determination of 49 ng/min. The calibration gas is mixed with the 5-slm inlet airflow to give 340 pptv calibrations. Br$_2$ is admitted into the inlet for 1 min approximately once per hour. The response to the calibration is rapid, with a >90% rise and fall in <2 s (Fig. 3). The sensitivity to Br$_2$ detection averages 5 ion counts per second per pptv (Hz/pptv).

The sensitivity to other halogen compounds was determined in the laboratory both before and after the field study, and in-flight sensitivity is calculated from the laboratory ratios to Br$_2$ sensitivity. Cl$_2$ sensitivity was determined using permeation devices and compressed gas mixtures containing 3–5 ppmv Cl$_2$ in N$_2$. Placing Br$_2$ and Cl$_2$ permeation tubes in a single oven at 40°C made BrCl. The BrCl output was related to Br$_2$ by the reduction in Br$_2$ when Cl$_2$ was added and BrCl was formed, assuming quantitative conversion of Br$_2$ to BrCl. The sensitivities to Cl$_2$ and BrCl detection at the most abundant product isotopes (I$^-$·$^{70}$Cl$_2$ (58%), I$^-$·$^{116}$BrCl (50%), Fig. 2) are approximately equal to that for Br$_2$ (I$^-$·$^{160}$Br$_2$, 50%). BrO was made by flowing Br$_2$ in He through a microwave discharge to make Br atoms (Orlando et al., 1991), and then ozone was added after the discharge to make BrO by Reaction (R1). The sensitivity is determined by comparing BrO to Br$_2$ variations when the discharge and ozone were modulated. Ozone addition lowers the Br$_2$ output since Br recombination that regenerates Br$_2$ is reduced by the loss of Br to BrO (Reaction R1). Similarly, operation of the microwave discharge reduces Br$_2$ signals. These experiments place upper and lower bounds on the BrO sensitivity relative to Br$_2$, since Br atoms may be lost to surfaces without Br$_2$ reformation. Using I$^-$ ion chemistry, the sensitivity to BrO detection is approximately 3835.
half the Br$_2$ sensitivity. BrO and BrCl measurement uncertainties are larger since their calibration magnitudes are not independently measured but are derived from changes in Br$_2$ signals.

2.4 Background determination and measurement uncertainty

The instrument background is measured by sampling ambient air through a scrubber that removes halogen compounds without otherwise changing the ambient air. Every 15 min, the Teflon valve located at the inlet tip (see Fig. 1 in the work of Neuman et al., 2002) was actuated and ambient air was directed through a scrubber before entering the inlet. After 1 min, the valve was switched to bypass the scrubber and directly sample ambient air. The scrubbing material in the WP-3D I$^-$ CIMS is untreated fine Pyrex glass wool that readily removes all Br$_2$ and BrO. Other tested materials, including steel wool and nylon wool, do not remove Br$_2$ completely. Activated charcoal, which is used as a scrubbing material in the other two CIMS instruments, effectively removes Br$_2$ from a gas stream, but it also removes many other compounds.

Background signal levels at the ion masses used for halogen detection were low. The measurement sequence shown in Fig. 3 illustrates the response to in-flight background determinations at ion mass 287 used for Br$_2$ detection. The open red circles indicate when ambient air was directed through the glass wool filter. Importantly, the instrument background did not change with altitude and remained low ($\approx$2 pptv) and stable. BrO background levels were also consistently a few pptv. BrCl backgrounds were sometimes higher and more variable, causing greater uncertainty in the BrCl measurements. The measured background values for each ion mass are subtracted from the total signal at that mass to determine ambient mixing ratios. Although sensitivity to Cl$_2$ is high, occasional rapid and large changes in the background at 197 amu (I$^-$·Cl$_2$) prevented accurate quantification of Cl$_2$ at levels less than a few hundred pptv. Hence, Cl$_2$ mixing ratios are not determined.

Imprecision, uncertainty in calibration addition and response, and background uncertainty contribute to total measurement uncertainties. Overall measurement uncertainty
for 2 s data from the WP-3D I^− CIMS is ±(15%+2) pptv for Br₂ detection, and ±(40%+3) pptv for BrO and BrCl detection. Greater uncertainties for BrO and BrCl detection are caused by the additional uncertainty in determining BrO and BrCl calibration magnitudes (discussed above) and the higher and more variable background levels. The SF₆⁻ CIMS instruments on the WP-3D and DC-8 also detected BrO and Br₂. The sensitivity to Br₂ detection using SF₆⁻ was better than 4 Hz/pptv for both instruments. The DC-8 CIMS measured BrO on two flights and Br₂ on all seven arctic flights, with measurements once every 30 s.

### 2.5 Selectivity

Bromine speciation is achieved by exploiting ion chemistry to detect each bromine-containing compound at a unique mass, as described above. The distinct isotopic signatures from bromine-containing compounds reduce ambiguity in ion identification (Fig. 2). Additionally, the product ions are at high masses and far from interfering ions or fragments of ion clusters. However, measuring highly reactive species is subject to sampling artifacts, including reaction on instrument surfaces that cause loss or chemical conversion of compounds. Laboratory tests showed that sampling air through the Teflon inlet did not introduce temporary retention or loss of BrO, BrCl, or Br₂. But experiments conducted after the field campaign showed that HOBr readily converts to Br₂ on the aircraft inlets used here. This conversion is similar to the efficient uptake of HOBr onto salt solutions (Fickert et al., 1999) and other surfaces (Mochida et al., 1998) that have been shown to produce gas phase Br₂.

The conversion of HOBr to Br₂ is quantified by synthesizing and measuring HOBr and studying its behavior on inlet surfaces. Gas phase HOBr is produced by liberating HOBr formed in solution by Br₂ hydrolysis (Eigen and Kustin, 1962) in the reaction

\[ \text{Br}_2 + \text{H}_2\text{O} \rightleftharpoons \text{HOBr} + \text{Br}^- + \text{H}^+ . \]  

(R2)

Br₂ hydrolysis is performed by flowing 4 sccm humidified N₂ and 0.2 ppmv Br₂ in
30 sccm N₂ over glass beads coated with AgNO₃. The 3 mm diameter glass beads are contained in a 5-cm long, 1-cm ID, Teflon tube. Teflon fittings connect to 0.32 cm OD Teflon supply and delivery lines. Br₂ hydrolyzes on the wet glass beads, and the resulting Br⁻ is removed by the formation of AgBr. This drives the hydrolysis equilibrium toward HOBr formation, so that much of the Br₂ is converted to gas phase HOBr. At the exit of the cell containing the AgNO₃-coated beads, the 34-sccm N₂ flow typically has 20 ppbv Br₂ and 100 ppbv HOBr. Mixing this flow into a several slm inlet flow dilutes Br₂ and HOBr to hundreds of pptv in the instrument.

HOBr is detected by the CIMS instrument using the same I⁻ ion chemistry described above. Figure 4 shows laboratory mass spectra of I⁻ ion chemistry with HOBr and Br₂ in N₂. HOBr reacts with I⁻·H₂O to form I⁻·HOBr cluster ions (223 and 225 amu). Br⁻ and its water clusters are also formed (red peaks at masses 79 and 81 in Fig. 4). The efficient reaction channel that forms Br⁻ causes the sensitivity to HOBr (for detection at I⁻·HOBr) to be approximately one third that for Br₂ detection. Br⁻ ions are not used for HOBr detection since they do not uniquely represent a single bromine compound. For example, Br₂ also reacts with I⁻ to form Br⁻ (green peaks at masses 79 and 81 in Fig. 4), although with much less efficiency than the clustering of Br₂ with I⁻ (green peaks at masses 285, 287, 289 in Fig. 4).

HOBr conversion to Br₂ is quantified by comparing changes in HOBr and Br₂ signals caused by interactions with inlet surfaces. HOBr to Br₂ conversion occurs solely on the inlet walls. Br₂ signals are unaffected by HOBr addition directly into the flow tube, demonstrating that conversion of HOBr to Br₂ does not occur in the ion-molecule reaction region. In the flow tube, low Reynolds number and residence time (0.17 s) reduce surface interaction. Sampling HOBr through the 70 cm long aircraft inlet lowers HOBr and increases Br₂ signals (Fig. 5). Since Br₂ does not react on the Teflon inlet, HOBr conversion to Br₂ is quantitative in these laboratory studies. Figure 5 shows the measured signals from 0.4 ppbv Br₂ and 2.5 ppbv HOBr in 5 slm dry N₂ that are admitted directly into the instrument (labeled Instrument Response), into the inlet (Teflon Aircraft Inlet), or into a Pyrex tube coated with NaBr. The conversion of HOBr to Br₂ on Br⁻
coated surfaces is five times more efficient than conversion on the Teflon aircraft inlet (Fig. 5). On other materials, including glass, aluminum, stainless steel, polyvinylidene fluoride (PVDF), and several other types of Teflon, HOBr converts to Br₂ with equal or greater efficiency than with PFA Teflon. On nylon, however, HOBr is removed without substantial Br₂ production.

Since HOBr is converted to Br₂ on the aircraft inlet, Br₂ signals at the detector represent a sum of atmospheric Br₂ and HOBr. Conversion efficiency is enhanced by wall contact, which is not quantified for all flight conditions. Furthermore, conversion of HOBr to Br₂ increases dramatically on bromide-coated surfaces, yet the extent to which the aircraft inlets were bromide-coated is not known. Therefore, HOBr to Br₂ conversion efficiency is uncertain but no greater than unity. Consequently, Br₂ signals represent a lower limit of the sum of atmospheric HOBr+Br₂, which is called active bromine below.

In-flight diagnostics performed by the WP-3D I⁻ CIMS include mass scans from 215–295 amu once every 90 min. These were performed to assess mass spectrometer and ion chemistry stability, interferences from other compounds, and possible detection of other molecules. As an example, Fig. 6 shows mass spectra obtained over the Arctic Ocean north of Barrow on 12 April. Ozone mixing ratios were 12 ppbv at 150 m altitude and 50 ppbv at 1.7 km. In the ozone-depleted air, I⁻·Br₂ ions were elevated, but clusters of I⁻ with BrO, HOBr, and BrCl were not elevated above detection limits. The similarity of all spectra, absence of interferences near the product ion masses used for bromine detection, and unique isotopic signature of Br demonstrate that bromine was enhanced in ozone-depleted air and was detected as Br₂.

The two CIMS instruments that used SF₆⁻ chemistry recorded similarly large Br₂ signals. The Br₂ measurements from the two CIMS instruments on the WP-3D were in quantitative agreement. Both instruments use the same calibration sources, so the correlation slopes between the two measurements do not reveal absolute accuracy. But high correlation coefficients (e.g., r=0.86 on 19 April) indicate that elevated Br₂ signals are not artifacts of one instrument’s ion chemistry or scrubbing materials. Thus, either
atmospheric $\text{Br}_2$ or conversion of HOBr to $\text{Br}_2$ on the inlets of both CIMS instruments caused the elevated $\text{Br}_2$ signals (called active bromine here).

3 Results

3.1 Ozone depletion

Ozone depletion is identified and categorized by comparing ozone mixing ratios in the MBL to values in the lower free troposphere, where ozone was never substantially depleted and variability was low. For example, between 1 and 2 km altitude on the five arctic WP-3D flights, average ozone mixing ratios were $54 \pm 9$ ppbv, with a 33–73 ppbv range. In the MBL, ozone was often considerably lower than these free tropospheric levels. Major ozone depletion ($O_3 < 4$ ppbv, as defined in the work of Ridley et al., 2003) was encountered on three WP-3D flights: 12 April, 15 April, and 18 April, for 71, 25, and 13 min, respectively. Partial ozone depletion ($4 < O_3 < 20$ ppbv) was measured on all 5 arctic WP-3D flights, for a total of 10–67 min on each flight. In total, major ozone depletions were sampled for 1.8 h, and partial ozone depletions were sampled for 2.7 h.

The map showing the five WP-3D aircraft flights that sampled the arctic boundary layer illustrates the location of ozone-depleted air (Fig. 1). Ozone was usually depleted at low altitudes over the Arctic Ocean. The exception was on 18 April, when ozone was not depleted in the MBL (indicated by the gaps along the flight track in Fig. 1). Unique to this flight, winds were predominately from land to the south, and ozone-rich air was advected over the ocean. Similarly, ozone-depleted air was sometimes transported inland when winds were from the north, as shown by the colored markers over land on 15 April. Above 300 m altitude ozone was nearly always greater than 30 ppbv.

Observations from the DC-8 were similar. Ozone-depleted air was encountered in the MBL on six of the seven DC-8 flights in the arctic, with major ozone depletions sampled on 3 flights. In total, the DC-8 sampled major ozone depletions for 0.7 h and partial ozone depletions for 1.2 h. The DC-8 measurements also show that ozone
depletion ($O_3 < 20$ ppbv) was confined to the MBL. On all seven flights in the arctic, ozone was always greater than 33 ppbv above 1 km altitude.

Ozone depletion is identified here only when ozone was less than 20 ppbv. However, both aircraft regularly observed ozone levels both in and above the MBL that were between 20–40 ppbv and lower than the average lower tropospheric values. These slight reductions in ozone mixing ratios, which are not analyzed here, also may have been caused by bromine chemistry, followed by transport and dilution of ozone-depleted air. In the MBL, extremely low ozone values were occasionally observed. Both the DC-8 and WP-3D measured $O_3 < 1$ ppbv on two flights and $O_3 < 0.1$ ppbv on one flight.

### 3.2 Bromine distributions and speciation

The compounds and reactions involved in ozone depletion chemistry are investigated with speciated bromine measurements. Active bromine, which represents a lower limit of the sum $HOBr + Br_2$, was consistently enhanced in the MBL when ozone was depleted (Fig. 1). Active bromine measured along the WP-3D flight tracks (Fig. 1b) was co-located with ozone-depleted air (Fig. 1a). BrO measured from the WP-3D aircraft was usually less than active bromine, and BrCl levels were typically less than the 2-pptv detection limit (Fig. 7). Above 300 m altitude (gray lines in Fig. 1a), active bromine was below the 2-pptv detection limit (gray lines in Fig. 1b) indicating little transport of active bromine out of the MBL.

The relationship between active bromine and photolysis rates is examined to assess the relative contribution of $Br_2$ and $HOBr$ to active bromine. $Br_2$ and BrO photolysis rates calculated from actinic flux measurements varied from 0.02/s to 0.06/s, while $HOBr$ photolysis rates were approximately 30 times slower (e.g., Lehrer et al., 2004; Martinez et al., 1999). Models show that rapid $Br_2$ photolysis should restrict daytime $Br_2$ to the bottom few cm of the atmosphere (Lehrer et al., 2004). Since Br recombination or BrO self reaction that forms $Br_2$ is slow compared to photolysis, $Br_2$ is predicted to be elevated only in close proximity to its surface source. Since the time required for air to travel from the surface to aircraft altitudes ($\sim 100$ m) is many $Br_2$ photolysis
lifetimes, atmospheric Br$_2$ should be near zero in these daytime aircraft measurements. And if Br$_2$ were elevated, its abundance ought to depend upon the large Br$_2$ photolysis rate, $j_{Br_2}$, which varied with latitude, time, cloud cover, and surface albedo. For example, the aircraft flew over open ocean leads in the ice, which had much lower albedo than arctic ice. The reduction in reflected light over the leads caused rapid and large (>30%) reductions in $j_{Br_2}$ that should alter Br$_2$ mixing ratios. However, active bromine did not vary with $j_{Br_2}$, suggesting that there was little contribution from Br$_2$ to active bromine. Therefore, the measured active bromine is likely HOBr.

Bromine measurements from the DC-8 show similar distributions and enhancements. In ozone-depleted air sampled in the MBL over the Arctic Ocean, active bromine, BrO, and soluble bromide were elevated. BrO levels (typically <10 pptv) were low compared to soluble bromide, which ranged from 10–40 pptv. Although several compounds could be responsible for the measured soluble bromide, both the active bromine and soluble bromide observations are consistent with the presence of HOBr in ozone-depleted air.

### 3.3 Vertical distributions

Altitude profiles further illustrate the confinement of ozone depletion and bromine enhancement to the MBL. The WP-3D measurements shown in Fig. 7 illustrate that all bromine compounds were zero, within measurement uncertainty, just above the MBL. Very steep gradients in both ozone and active bromine occurred at the MBL top. For example, active bromine dropped from tens of pptv to zero and ozone increased from less than 10 ppbv to 40 ppbv between the MBL and above 600 m altitude (at 02:42 and before 02:35 in Fig. 7). The DC-8 also measured elevated active bromine in the MBL and values below detection limit above.

Boundary layer meteorology plays an important role in determining the vertical extent of ozone-depleted air. Figure 8 shows 1 s measurements of ozone, temperature, potential temperature, and CO, and 10 s averages of active bromine during a WP-3D descent from 4 km to 60 m altitude. These measurements were obtained on 12 April over the
Arctic Ocean at approximately 75° N, 145° W (Fig. 1) and approximately 560 km north of the Alaskan coast. The MBL was well-mixed, as indicated by near-constant potential temperature between 60–200 m. Above the well-mixed layer, a strong temperature inversion of about 5°C over 500 m (Fig. 8) formed a stable layer that inhibited mixing between the boundary layer and the free troposphere above. Below the temperature inversion over the ocean, ozone was depleted and active bromine was enhanced. Above the MBL ozone was 40–55 ppbv and active bromine was below detection limits. Similar altitude profiles were obtained on the five flights shown in Fig. 1, both at coastal locations and over the Arctic Ocean hundreds of km from the Alaskan coast. Other trace gases exhibit little difference in concentration between the MBL and the free troposphere. For example, CO mixing ratios averaged 158±2 ppbv over the entire vertical profile shown in Fig. 8.

Variations with altitude were not apparent for many species because their mixing ratios were usually near instrument detection limits below 1 km over the Arctic Ocean. BrO and BrCl were below detection limits throughout the vertical profile shown in Fig. 8, averaging −0.9±2.6 pptv and −0.5±2.3 pptv, respectively. Ozone depletion was observed in clean background air, and low mixing ratios of NO, NO₂, and SO₂ were typical for most of the sampling in the MBL. NO, NO₂, and SO₂ were equivalent to zero, within measurement uncertainties, over the time period shown in Fig. 7, with average values of 2±8 pptv, 16±17 pptv, and 13±270 pptv, respectively.

3.4 Bromine-ozone relationship

The relationship between ozone and each halogen compound was unique, and these relationships were consistent on all WP-3D flights into the arctic. Figure 9 shows 10 s averages of active bromine (A), BrCl (B), and BrO (C) vs. ozone. When ozone was greater than 40 ppbv, both active bromine and BrCl were at or below detection limits, while BrO increased at the highest ozone levels. Elevated ozone and BrO were measured at 7.4 km in air that was influenced by the stratosphere. In this stratospheric intrusion sampled by the WP-3D on 19 April, ozone reached 152 ppbv and
was anti-correlated with CO, which dropped to 91 ppbv. 30 s averages of BrO reached a maximum of 7 pptv.

In ozone-depleted air, which was measured only in the MBL, active bromine was consistently enhanced with values that reached 20 pptv. BrCl remained at or below detection limits, and BrO was usually less than 5 pptv. The largest enhancements in active bromine and BrO in the MBL were observed on 19 April, when the meteorology differed considerably from the other days. On 19 April, the temperature inversion was higher (500 m), the MBL air was warmer (−12 °C) and the winds were stronger (16 m/s) compared to the four other days, when the temperature inversion occurred between 200 and 250 m, the MBL temperatures were from −20 to −15°C, and wind speeds were 2–8 m/s. BrO mixing ratios measured here in ozone-depleted air were considerably lower than many previous reports. When O₃ < 4 ppbv, measurements of BrO from the WP-3D averaged −0.3±1 pptv, and those from the DC-8 averaged 2±3 pptv. When ozone depletion was less complete (4 < O₃ < 20 ppbv), BrO mixing ratios were higher, averaging 0.9±1.6 pptv (WP-3D) and 3±4 pptv (DC-8). In contrast, long-path measurements have observed BrO mixing ratios up to ten times greater in ozone-depleted air (e.g., Hausmann and Platt, 1994; Tuckermann et al., 1997).

4 Discussion and summary

4.1 Geographical extent of ozone depletion

Measurements here over the Alaskan Arctic are consistent with those from the Canadian Arctic. Ozone depletion in the springtime has been reported as a persistent feature of the MBL over the Canadian Arctic (Hopper et al., 1998; Bottenheim and Chan, 2006; Leaitch et al., 1994; Ridley et al., 2003). Similarly, measurements here show that ozone was regularly depleted in the MBL over the Arctic Ocean north of Alaska. The WP-3D aircraft measured ozone depletion above the Chukchi Sea (as far south and west as 67° N, 165° W), Beaufort Sea (as far east as 125° W), coastal areas (for
example, 71° N, 151° W near Barrow, Alaska), and hundreds of km from land (as far as 75° N, north of the Alaskan Coast). In all of these cases, ozone depletion was confined to the lower few hundred meters of the atmosphere in the MBL. Ozone depletion was observed in air masses with temperatures that ranged from −22 °C to −10 °C, suggesting that ozone depletion can occur over a wide range of temperatures, as has been noted previously (Simpson et al., 2007; Bottenheim et al., 2009).

4.2 HOBr and Br₂ measurements

This study might lead to new speciated measurements of inorganic bromine and assist with the interpretation of previous reports of photochemically active bromine compounds. HOBr is shown to convert rapidly to Br₂ on many types of Teflon, glass, and metal, even without bromide coatings applied to the materials’ surfaces. Sampling air without introducing wall contact eliminates this reaction, although this is not always feasible. Alternatively, rapid conversion can be exploited to quantitatively convert all HOBr to Br₂ by sampling air through a bromide-coated inlet to measure the sum HOBr+Br₂. And Br₂ can be measured independently by sampling air through a nylon wool filter, since HOBr is removed on nylon, but Br₂ is not. From these two measurements, HOBr can be determined from the difference between (HOBr+Br₂) and Br₂.

HOBr conversion on instrument surfaces may have contributed to previous observations of Br₂. For example, up to 27 pptv of Br₂ was observed after sunrise in ozone-depleted air by a CIMS technique that sampled a large gas flow through a 9 m long Teflon inlet (Foster et al., 2001). If HOBr were present in the atmosphere and the sampled air contacted the inlet walls, HOBr conversion likely contributed to the Br₂ signal. Similarly, measurements of HOBr and Br₂ using glass cells (Impey et al., 1999) also may have been affected by the conversion of HOBr to Br₂ on surfaces.
4.3 Bromine chemistry

Several inorganic bromine compounds participate in chemical reactions that lead to ozone depletion in the springtime over the Arctic Ocean, and several of those compounds were observed here. Previous observations of BrO and total bromine in ozone-depleted air show that BrO accounts for a widely varying fraction of total bromine (Martinez et al., 1999; Hausmann and Platt, 1994). But speciated measurements of inorganic bromine other than BrO are sparse (Barrie et al., 1994; Impey et al., 1999; Foster et al., 2001), and the relative contribution of each bromine compound to total bromine is not certain. HOBr is often comparable to or larger than BrO during the daytime measurements reported here, assuming that the measured active bromine is only HOBr. The predominance of HOBr over Br$_2$ is consistent with previous measurements (Impey et al., 1999), and HOBr has been predicted to be more abundant than BrO in ozone depleted air (Martinez et al., 1999). However, the BrO mixing ratios measured here (typically a few pptv in ozone-depleted air) from three different aircraft CIMS instruments were much lower than the tens of pptv that has been observed previously from ground-based instruments. The causes for low BrO values have not been determined, but several factors may explain these results. If BrO were confined to the lowest altitudes of the MBL, these aircraft observations at approximately 100 m altitude may have been above regions with enhanced BrO. Low BrO could be caused by losses on inlets exposed to ambient air, although BrO losses on Teflon inlets were measured to be insignificant in the laboratory. Lastly, the weak correlation between ozone and BrO (Hausmann and Platt, 1994) and BrO variations between 0–30 pptv in ozone-depleted air (Tuckermann et al., 1997) make it possible that these few aircraft flights may have sampled only those ozone depletions with low BrO. However, the observed increases in active bromine indicates the presence of bromine chemistry in the sampled ozone-depleted air masses, where both BrO and HOBr are predicted to be enhanced (e.g., Martinez et al., 1999; Evans et al., 2003; Calvert and Lindberg, 2003; Lehrer et al., 2004).
Bromine partitioning between HBr, BrO, and HOBr may reveal whether ozone depletion has terminated or is still occurring in a sampled air mass. Several models examined the temporal behavior of halogen-initiated ozone depletion and show HBr, BrO, and HOBr enhancements of many pptv during daytime in ozone-depleted air (Martinez et al., 1999; Calvert and Lindberg, 2003; Evans et al., 2003; Fan and Jacob, 1992; Shepson et al., 1996; Lehrer et al., 2004), with relative abundances that vary with the duration of ozone depletion chemistry. Bromine accumulates as HBr when ozone depletion has terminated (Lehrer et al., 2004; Martinez et al., 1999), and the presence of HOBr may indicate that ozone depletion chemistry is still in progress. Improved measurements of these halogen compounds as suggested above might allow for identification and better characterization of halogen chemistry that depletes ozone.

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**Fig. 1.** (A) 1 s measurements of ozone on five WP-3D flights originating from Fairbanks, Alaska. Red circles are major ozone depletions, blue are partial ozone depletions, and no symbol represents ozone >20 ppbv. Gray lines indicate flight tracks over 300 m altitude. Gaps in the flight tracks (on 18 April, for example) indicate flight altitudes less than 300 m and ozone greater than 20 ppbv. (B) 10 s averages of active bromine, colored according to mixing ratio. Red markers show active bromine over 6 pptv, blue indicate active bromine between 2 and 6 pptv, and gray lines represent active bromine less than 2 pptv. The gaps occur when in-flight diagnostics were performed and active bromine was not measured.
Fig. 2. Mass spectra for laboratory air admitted into a humidified flow tube with I⁻ ions (black line). The ordinate is linear up to 4000 Hz and logarithmic above. The red line includes addition of 0.3 ppbv Br₂, 0.1 ppbv BrCl, and 1.5 ppbv Cl₂.
Fig. 3. I⁻•Br₂ signals recorded once every 2 s during ambient measurements (gray), calibration (green), and background determinations (red) over the Arctic Ocean on 19 April. Ozone is shown in blue and aircraft altitude in black.
Fig. 4. Mass spectra for dry N\textsubscript{2} (black), dry N\textsubscript{2} with 1 ppbv Br\textsubscript{2} (green), and dry N\textsubscript{2} with 0.3 ppbv Br\textsubscript{2} and 1.5 ppbv HOBr (red) mixed with I\textsuperscript{−} in a humidified flow tube. The ordinate is linear up to 6000 Hz and logarithmic above.
**Fig. 5.** Br\(_2\) (black) and HOBr (diagonal lines) signals when 0.4 ppbv Br\(_2\) and 2.5 ppbv HOBr in dry N\(_2\) are admitted directly into the flow tube (labeled Instrument Response), flow through the Teflon aircraft inlet, and flow through a NaBr-coated tube.
Fig. 6. Mass spectra obtained in flight at 1.7 km (black) and 150 m (red) altitude over the Arctic Ocean north of Barrow on 19 April. The WP-3D aircraft was in the MBL and ozone was depleted ($O_3=12$ ppbv) at 150 m altitude.
Fig. 7. 1 h time series of measurements obtained over the Arctic Ocean 300 km north of Barrow, Alaska on 19 April. 1 s measurements of ozone (blue) and aircraft altitude (black), and 10 s averages of active bromine (red), BrO (green), and BrCl (pink) are shown.
Fig. 8. 1 s measurements of O₃ (blue), potential temperature (black), ambient temperature (pink), and CO (green) vs. altitude during a descent of the WP-3D aircraft from 4 km to 60 m altitude on 12 April 2008. Red symbols are 10 s averages of active bromine. The gap in bromine data occurred during a 60 s instrument zero.
Fig. 9. 10 s averages of active bromine (A), BrCl (B), and BrO (C) vs. ozone from five WP-3D flights that included sampling in the MBL over the Arctic Ocean.