INVESTIGATION OF TROPOSPHERIC BRO USING SPACE-BASED TOTAL COLUMN BRO MEASUREMENTS

A Thesis
Presented to
The Academic Faculty

by

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In Partial Fulfillment
of the Requirements for the Degree
Doctor of Philosophy in the
School of Earth and Atmospheric Sciences

Georgia Institute of Technology
May 2012
INVESTIGATION OF TROPOSPHERIC BRO USING SPACE-BASED TOTAL COLUMN BRO MEASUREMENTS

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This thesis is dedicated to my father Byung-sik Choi,

my mother Boksoon Lee,

and my grandmother Euijin Lee.
ACKNOWLEDGEMENTS

First, I would like to thank Dr. Joanna Joiner, without whose supports including kind consideration, insightful advices and constant encouragement I would never ever have been able to graduate. I am also grateful to Dr. Ross J. Salawitch and Dr. Tim Canty for discussions and comments that improved the Arctic tropospheric BrO paper enormously.

I appreciate Dr. Yuhang Wang’s supports regarding ARCTAS and ARCPAC projects. I express my gratitude to my thesis committee members, Dr. Judith Curry, Dr. Greg Huey, Dr. Rodney Weber, and Dr. Athanasios Nenes for careful reading and inspirational questions to my thesis.

I appreciate Dr. Tao Zeng’s help for me to learn IDL and Fortran when I was totally new to computation. Talking to Burton Gray, Dasa Gu and Zhen Liu always made me feel refreshed and encouraged. I enjoyed discussions with Jin Liao about bromine measurements and chemistry. I also thank to Dr. Junsang Nam for his wise advices.

My parents, my sister and my aunts gave great support for my life of studying abroad. I could not realized how precious they are until I became apart from them.

Finally, I express special thanks to actors and makers of Infinite Challange, Tengen Toppa Gurren Lagann and Ookiku Furikabutte, which saved me from continuous frustration.
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SUMMARY

We derive tropospheric column BrO during the ARCTAS and ARCPAC field campaigns in spring 2008 using retrievals of total column BrO from the satellite UV nadir sensors OMI and GOME-2 using a radiative transfer model and stratospheric column BrO from a photochemical simulation. We conduct a comprehensive comparison of satellite-derived tropospheric BrO column to aircraft in-situ observations of BrO and related species. The aircraft profiles reveal that tropospheric BrO, when present during April 2008, was distributed over a broad range of altitudes rather than being confined to the planetary boundary layer (PBL). Perturbations to the total column resulting from tropospheric BrO are the same magnitude as perturbations due to longitudinal variations in the stratospheric component, so proper accounting of the stratospheric signal is essential for accurate determination of satellite-derived tropospheric BrO. We find reasonably good agreement between satellite-derived tropospheric BrO and columns found using aircraft in-situ BrO profiles, particularly when satellite radiances were obtained over bright surfaces (albedo >0.7), for solar zenith angle <80° and clear sky conditions. The rapid activation of BrO due to surface processes (the bromine explosion) is apparent in both the OMI and GOME-2 based tropospheric columns. The wide orbital swath of OMI allows examination of the evolution of tropospheric BrO on about hourly time intervals near the pole. Low surface pressure, strong wind, and high PBL height are associated with an observed BrO activation event, supporting the notion of bromine activation by high winds over snow. We also provide monthly climatological maps of free tropospheric BrO volume mixing ratio (VMR) derived using the so-called cloud slicing technique. In this
approach, the derived slope of the total column BrO versus cloud pressure is proportional to free tropospheric BrO VMR. Estimated BrO VMR shows a minimum in the tropics and greater values at higher latitudes in both hemispheres. High tropospheric BrO VMR at high latitudes in spring could be influenced by near-surface bromine activation.
Bromine radicals ($\text{BrO}_x = \text{Br} + \text{BrO}$) are important species owing to their ability to destroy ozone catalytically [Barrie et al., 1988, Hausmann and Platt, 1994]. They may also impact oxidative pathways of many trace chemicals including mercury [Schroeder et al., 1998], and dimethyl sulfide (DMS) [Toumi, 1994]. For example, bromine radicals are known to oxidize gaseous elemental mercury (GEM) to reactive gaseous mercury (RGM) [Schroeder et al., 1998, Lu et al., 2001, Douglas et al., 2005, Holmes et al., 2010]. Rapid depletion of GEM concurrent with high BrO concentrations has been reported in the polar boundary layer [Ariya et al., 2004] as well as in the tropical marine boundary layer [Saiz-Lopez et al., 2006]. In addition, bromine radicals oxidize DMS which may shift the DMS oxidation channel [Read et al., 2008, Mahajan et al., 2010, Saiz-Lopez et al., 2004, 2006]. DMS oxidation is normally dominated by OH, particularly in tropical marine boundary layer. However, high BrO concentrations can cause DMS oxidation to be dominated by BrO. This in turn may affect aerosol formation and cloud albedo [von Glasow and Crutzen, 2004]. Therefore bromine radicals could influence global climate through the production of cloud condensation nuclei [Read et al., 2008, Mahajan et al., 2010].

Bromine monoxide (BrO) is the most commonly observed active bromine species. BrO absorbs ultraviolet (UV) radiation, which enables measurement using remote sensing techniques. Reported in-situ and remotely-sensed ground-, balloon-, and aircraft-based BrO measurements include: (1) boundary layer mixing ratio with Long Path-Differential Optical Absorption Spectroscopy (LP-DOAS) [Hausmann and
Platt, 1994, Tuckermann et al., 1997, Martinez et al., 1999; (2) mixing ratio by aircraft in-situ Chemical-Ionization Mass Spectrometry (CIMS) [Neuman et al., 2010, Liao et al., 2011a,b]; (3) boundary layer and free tropospheric BrO column using DOAS on aircraft [McElroy et al., 1999, Prados-Roman et al., 2011]; (4) total, stratospheric, and tropospheric columns using ground-based zenith sky and direct sun DOAS [Frieß et al., 1999, Schofield et al., 2004, 2006, Hendrick et al., 2007, 2008, Theys et al., 2007]; (5) boundary-layer column with ground-based MAX-DOAS [Hönninger, 2004, Simpson et al., 2007a, Donohoue et al., 2010, Frieß et al., 2011]; (6) balloon profiles using a variety of spectroscopic methods [Fitzenberger et al., 2000, Pfeilsticker et al., 2000, Pundt et al., 2002, Dorf et al., 2008]. Despite the many measurements of BrO and related species obtained by various techniques, significant uncertainties remain regarding the importance of very short lived source compounds on the stratospheric bromine budget [e.g. Sect. 1.3.3.3 of WMO, 2011] as well as the magnitude of the global, ubiquitous, background level of tropospheric BrO [e.g. Sect. 5.3 of Theys et al., 2011].

Space-based observation of BrO offers an excellent tool for studying bromine chemistry. Satellite observations provide global coverage, far superior to the spatial coverage available from ground- and aircraft-based measurements. Limb-sounding observations from the Microwave Limb Sounder (MLS) [Livesey et al., 2006, Kovalenko et al., 2007] provide quantification of BrO profiles in the upper and middle stratosphere. The MLS observations of BrO imply a significant contribution to stratospheric bromine from sources other than long-lived CH$_3$Br and halons. Limb observations from the SCanning Image Absorption SpectroMeter for Atmospheric CHartographY (SCIAMACHY) [Sinnhuber et al., 2005, Sioris et al., 2006, Rozanov et al., 2011] extend into the lowermost stratosphere (LMS) and nadir measurements from SCIAMACHY constrain total column BrO, including contributions from the stratosphere and troposphere. The retrievals of SCIAMACHY BrO by Rozanov et al. [2011] and Sinnhuber
et al. [2005] imply a limited role for supply of bromine by very short lived (VSL) compounds to the LMS and a considerable burden of global, ubiquitous, background BrO whereas the retrieval of SCIAMACHY BrO described by Sioris et al. [2006] implies a larger role for supply of stratospheric inorganic bromine, Br$_y$, by VSL compounds and a much smaller level for background tropospheric BrO [e.g. Sect. 2.5.2.1 of WMO, 2007].

1.1 Analysis of satellite-derived Arctic tropospheric BrO columns in conjunction with aircraft measurements during ARCTAS and ARCPAC

Bromine plays an important role in tropospheric ozone chemistry and the resulting oxidation capacity of the polar boundary layer. Bromine radicals catalytically destroy ozone, leading to nearly complete removal near the surface that is termed an ozone (O$_3$) depletion event (ODE). Once O$_3$ is depleted, high levels of reactive halogen species including atomic bromine (e.g. Br) become the primary oxidants for many species, including methane (CH$_4$) and mercury (Hg) [Simpson et al., 2007b, Schroeder et al., 1998]. During mercury depletion events (MDEs), Br and BrO are thought to oxidize elemental mercury to more reactive gaseous mercury that deposit to the polar ecosystem [Schroeder et al., 1998, Lu et al., 2001, Ariya et al., 2004, Douglas et al., 2005, Holmes et al., 2010].

A primary source of active bromine in the polar boundary layer is thought to be bromide (Br$^-$) in the condensed phase, which is transformed to gaseous molecular bromine (Br$_2$) by heterogenous chemistry. Potential sources of bromide are sea salt aerosols [Fan and Jacob, 1992], surfaces of first year sea ice [Simpson et al., 2007a, Wagner et al., 2007] or newly formed sea ice [Jones et al., 2006], frost flowers [Kaleschke et al., 2004], or blowing snow triggered by strong winds [Yang et al., 2008, 2010, Jones et al., 2009, 2010, Begoin et al., 2010, Theys et al., 2011, Toyota et al., 2011]. Br$_2$ released to the atmosphere is rapidly photolyzed to yield bromine radicals
(Br), i.e.

\[
\text{HOBr} + \text{Br}^- + \text{H}^+ \rightleftharpoons \text{H}_2\text{O} + \text{Br}_2 \quad \text{(condensed phase)}
\]

(1)

\[
\text{Br}_2 + h\nu \rightarrow 2\text{Br} \quad \text{(gas phase)}.
\]

(2)

The bromine radicals then lead to catalytic ozone loss, i.e.

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

(3)

\[
\text{BrO} + \text{XO} \rightarrow \text{BrX} + \text{O}_2 \quad (\text{X} = \text{Br, Cl, I, and OH})
\]

(4)

\[
\text{BrX} + h\nu \rightarrow \text{Br} + \text{X}
\]

(5)

Barrie et al. [1988] found that filterable bromine is strongly anti-correlated with the abundance of O\textsubscript{3} in the polar boundary layer. Following this work, many additional measurements of bromine species along with O\textsubscript{3} have been made in polar environments [e.g. Bottenheim et al., 1990, Barrie et al., 1994, Hausmann and Platt, 1994, Tuckermann et al., 1997, Martinez et al., 1999, Ridley et al., 2003]. The association of elevated BrO and depleted surface O\textsubscript{3} during Arctic spring is well established [e.g. Platt and Höninger, 2003, and references therein].

Nadir-viewing instruments on polar-orbiting satellites provide multiple daily observations of total column BrO at high latitude. The high surface albedo of polar regions provides good sensitivity to tropospheric BrO, including that near the surface [Wagner and Platt, 1998, Theys et al., 2011]. Chance [1998] and Hegels et al. [1998] retrieved BrO total vertical column densities (VCD) from nadir radiiances in the UV obtained by Global Ozone Monitoring Experiment (GOME) [GOME Users Manual, 1995] on the European Space Agency (ESA) European Remote Sensing 2 (ERS-2) satellite. The early GOME observations showed large enhancements over Hudson Bay during spring 1997, which was attributed to bromine release from the surface [Chance, 1998]. Estimates of BrO total column amount have been subsequently derived from other nadir-viewing satellite sensors including the Ozone Monitoring
Instrument (OMI) [Levelt et al., 2006] on the US National Aeronautics and Space Administration (NASA) Aura satellite, SCIAMACHY [Bovensmann et al., 1999] on the ESA Environmental Satellite (EnviSat), and the second Global Ozone Monitoring Experiment (GOME-2) instruments [Munro et al., 2006] that are flying on the series of European Meteorological Operational Satellites (EuMetSat MetOp). Retrievals of VCD BrO from different satellite instruments and by various groups tend to be in fairly close agreement. All satellite instruments report levels of BrO considerably larger than predicted by models that account only for stratospheric supply from the decomposition of methyl bromide and halons. The focus of the scientific community since publication of early GOME observations has been quantifying the degree to which this “excess BrO” resides in the troposphere, the stratosphere, or perhaps both regions of the atmosphere.

Tropospheric BrO columns can be derived from satellite observations using the residual method, in which an estimate of the stratospheric BrO column is subtracted from the satellite-derived VCD of BrO. Wagner and Platt [1998] and Richter et al. [1998] estimated tropospheric VCD of BrO using this approach from GOME observations, based on simple assumptions for stratospheric BrO. These studies and many others assumed that stratospheric BrO is zonally symmetric and that bromine was supplied to the stratosphere solely by the decomposition of long-lived organic compounds. Theys et al. [2009] and Salawitch et al. [2010] showed that the stratospheric VCD of BrO can exhibit strong gradients with respect to longitude at high latitude during spring and Salawitch et al. [2010] questioned prior estimates of residual tropospheric BrO found assuming that the stratospheric burden was zonally symmetric. Theys et al. [2011] derived tropospheric BrO columns from GOME-2 spectra with a model-based stratospheric BrO climatology [Theys et al., 2009] similar to that used here and described below.
Many studies related to bromine and ozone chemistry have been conducted using satellite-derived tropospheric BrO columns. Wagner and Platt (1998) reported elevated regions of BrO vertical column density in the Arctic and Antarctic regions. They noted these enhancements were likely due to increased abundance of tropospheric BrO, rather than a stratospheric disturbance, based on a variety of factors including the correlation between enhanced columns of BrO and the O$_2$ – O$_2$ collision complex. Wagner et al. [2001] showed that elevated BrO column amounts observed by GOME were correlated with low ozone in the boundary layer observed in-situ at Ny-Ålesund (Spitsbergen), Norway. In other studies, spatial and temporal features of ODEs have been simulated using 3-dimensional regional chemical transport models and GOME-derived tropospheric BrO columns [Zeng et al., 2003, 2006]. Connections between BrO-rich air masses and first-year sea ice have been indicated with back-trajectory analyses using SCIAMACHY data [Wagner et al., 2007]. Transport of a large BrO plume near the North Pole is also reported by Begoin et al. [2010]. A back trajectory study using satellite-derived tropospheric BrO columns indicated that ODEs can be differentiated into locally activated and transport driven events [Koo et al., 2011].

Despite the numerous studies of tropospheric polar bromine chemistry using satellite BrO observations, estimation and interpretation of tropospheric BrO information from space presents ongoing challenges. To properly estimate tropospheric BrO column amounts, the stratospheric contribution to the satellite-derived total column must be accurately represented [e.g. Theys et al., 2009, Salawitch et al., 2010]. The global, ubiquitous background tropospheric level of BrO inferred from the satellite record is sensitive to the amount of Br$_y$ delivered to the stratosphere by VSL bromocarbons [Salawitch et al., 2005]. Furthermore, low solar elevation angles in the early polar spring lead to large uncertainties in satellite total BrO column retrievals (see below). The presence of clouds further complicates the retrieval of tropospheric
BrO from satellite observations. Theys et al. [2011] have recently addressed many of these issues. They showed maps of tropospheric BrO columns derived using a method similar to that described below, and evaluated these columns using ground-based measurements of the tropospheric and stratospheric contributions to the total column. Our study builds upon the work of Theys et al. [2011] by further exploring the issues that affect quantification of tropospheric BrO columns and by using aircraft measurements of BrO to evaluate the satellite-derived tropospheric columns.

Salawitch et al. [2010] provided an initial analysis of BrO observations obtained by instruments aboard the NASA and NOAA aircraft [Neuman et al., 2010, Liao et al., 2011b] and the OMI satellite instrument [Kurosu and Chance, 2011] during spring 2008. They simulated the stratospheric BrO column assuming VSL bromocarbons supply between 5 and 10 ppt of Br\textsubscript{y} to the stratosphere in the form of product gas injection (PGI), in addition to the \sim 2 ppt of Br\textsubscript{y} supplied by source gas injection (SGI) of the VSL species CH\textsubscript{2}Br\textsubscript{2}. On the other hand, Theys et al. [2011] simulated the stratospheric burden of BrO assuming 1 ppt of PGI and 5 ppt of SGI (all represented as CH\textsubscript{2}Br\textsubscript{2}). The stratospheric BrO burden found by Salawitch et al. [2010] for their "best case" simulation of 7 ppt from PGI and 2 ppt from SGI (see their Figure S7) is considerably larger (\sim 27\% overall difference, with quite a bit of geographic variability) than the stratospheric BrO burden used by Theys et al. [2011] as shown in Figure A.1. Although we do not compare to independent observations of BrO here, such comparisons to aircraft and balloon-borne observations of BrO in Salawitch et al. [2005] support the use of our values for the delivery of stratospheric bromine by PGI and SGI. Other observations of BrO, such as some of those discussed in Section 2.5.2 of WMO [2007], support smaller values for these parameters.

Salawitch et al. [2010] concluded that high column amounts of BrO derived from satellite observations could, in some cases, be attributed to compression of stratospheric air to low altitudes (high pressures) and that this condition, not previously
recognized in the analysis of the satellite BrO record, should be considered to properly relate the satellite record to surface events. The magnitude of the stratospheric enhancement is sensitive to the supply of stratospheric Br\textsubscript{y} from VSL bromocarbons as well as the pathways of this transport (i.e. PGI versus SGI; see Chapter 2 of WMO [2003] for a detailed discussion of these pathways). Salawitch et al. [2010] showed agreement, to within uncertainties, between total column BrO measured by OMI and the sum of modeled stratospheric and aircraft-measured tropospheric BrO partial columns. However, their treatment of the tropospheric column did not explicitly account for tropospheric air mass factors (AMFs) and thus must be viewed with caution. Our study builds on this prior work by using a radiative transfer model to calculate tropospheric AMFs and also by considering the effects of clouds, surface reflectivity, and viewing geometry on the evaluation of the bromine budget for Arctic spring 2008.

Validation of satellite-derived BrO columns and their relationship to ODEs was a goal of two Arctic field campaigns conducted in 2008 under the auspices of the International Polar Year (IPY). The Arctic Research of the Composition of the Troposphere from Aircraft and Satellite (ARCTAS) mission was conducted in April and June–July 2008 by NASA [Jacob et al., 2010] and the US National Oceanic and Atmospheric Administration (NOAA) Aerosol, Radiation, and Cloud Processes affecting Arctic Climate (ARCPAC) mission took place in April 2008 [Brock et al., 2011]. Various in-situ aircraft measurements of trace chemicals, including O\textsubscript{3} and reactive bromine species (BrO, Br\textsubscript{2}, and soluble bromide), were made during these campaigns. Neuman et al. [2010] and Liao et al. [2011a] provide an overview of the in-situ measurements of BrO and related species obtained during ARCTAS and ARCPAC. We derive Arctic tropospheric columns BrO and evaluate the columns in conjunction with aircraft measurements in Chapter 3.
1.2 Estimation of free tropospheric BrO using OMI

Model studies have reported potential consequences of tropospheric bromine to ozone, mercury, and sulfur chemistry. Bromine chemistry could reduce global ozone according to model studies assuming various source scenarios [von Glasow et al., 2004, Yang et al., 2010]. Also, tropospheric BrO could be a major global sink of mercury [Holmes et al., 2010]. 0.5 ppt of BrO concentration in the marine boundary layer can significantly influence DMS in the global scale [von Glasow et al., 2004]. However, accurate estimate of tropospheric bromine is necessary to assess impact of bromine to tropospheric trace gases.

The presence of tropospheric BrO has been reported in many studies under a variety of circumstances. High BrO levels (> 10 pptv) in the polar boundary layer during spring may result from rapid activation of bromine species (so-called bromine explosions) [e.g. Simpson et al., 2007b, and references therein]. Local activation of BrO is also reported near Salt lakes [Obrist et al., 2010]. Moreover, considerable amounts of BrO have been observed in tropical [Read et al., 2008, Mahajan et al., 2010] and mid-latitude [Saiz-Lopez et al., 2004, 2006] marine boundary layers as well as semi-polluted boundary layer [Mahajan et al., 2009]. The primary source of tropospheric BrO in the marine boundary layer is thought to be bromide in sea salts which is activated by multiphase reactions. [e.g. Fan and Jacob, 1992]

Ground-based BrO column observations imply free tropospheric BrO concentrations of between about 0.5 and 2 pptv, although there is significant uncertainty in absolute magnitudes [Van Roozendael et al., 2002, Harder et al., 1998, Schofield et al., 2004, 2006, Hendrick et al., 2007]. McElroy et al. [1999] suggested that a source of free tropospheric BrO is convection of bromine species that can be activated in polar regions, while von Glasow et al. [2004] postulates that downward transport of bromine species from stratosphere can contribute. However, the depiction of the free tropospheric BrO budget is still highly uncertain.
Satellite measurements offer an intriguing opportunity to quantify the tropospheric BrO budget. The use and interpretation of satellite measurements has many inherent challenges as discussed by Choi et al. [2012, etc.]. Here, we exploit a method known as cloud slicing to infer information about tropospheric BrO. The cloud-slicing technique, first introduced by [Ziemke et al., 2001], has been exploited to estimate free tropospheric ozone mixing ratios [Ziemke et al., 2009]. In Chapter 4, we apply this technique using satellite BrO and cloud pressure retrievals, building on the initial studies of Richter et al. [2002], Theys et al. [2011] that suggested the existence of global ubiquitous free-tropospheric BrO.
CHAPTER II

DATA DESCRIPTION

In this chapter, we introduce various data sets used in this study in order to calculate satellite-derived tropospheric column and free tropospheric BrO volume mixing ratio. We use a variety of aircraft and satellite measurements and model outputs to calculate and evaluate satellite-derived tropospheric BrO columns (Chapter 3). Tropospheric BrO columns are inferred from aircraft measurements of in-situ BrO for comparison with satellite-derived tropospheric column (henceforth referred to as “in-situ columns”, as described in Sect. 2.1). We derive tropospheric vertical column densities (VCDs) using BrO total slant column densities (SCD) retrieved from OMI and GOME-2 (see Sect. 2.2) and a model simulation of the stratospheric BrO column as detailed in Sect. 2.3. OMI rotational Raman scene pressures and MODIS cloud products (see Sect. 2.4) are used to assess cloud effects on the derived tropospheric BrO columns. Data from a global reanalysis, described in Sect. 2.5, are used to examine relationships between observed BrO enhancement events and meteorological conditions. OMI total column BrO and OMI cloud scene pressure are also used to estimate free tropospheric BrO volume mixing ratio (Chapter 4).

2.1 Aircraft in-situ measurements

Aircraft in-situ measurements of BrO and various other trace gas concentrations were made from the NASA DC-8 aircraft during ARCTAS and the NOAA WP-3D aircraft during ARCPAC. An overview of the instruments, flights, and mission goals and accomplishments are provided in the ARCTAS [Jacob et al., 2010] and ARCPAC [Brock et al., 2011] overview papers. BrO was measured using Chemical Ionization Mass Spectrometer (CIMS) instruments during both experiments [Neuman et al.,
DC-8 CIMS measurements of BrO were made once every 30 s, and WP-3D CIMS measurements were acquired once every 2 s. Measurement uncertainties for BrO are ±40 % with a detection limit of 3 pptv for WP-3D data and ±40 % and a detection limit of 2–5 pptv for DC-8 data [Neuman et al., 2010, Liao et al., 2011b]. In this study, we use 1 min averaged data for both DC-8 and WP-3D measurements. The spatial resolution of the 1 min averaged aircraft data is approximately 10 km.

We estimate tropospheric BrO columns from the aircraft BrO mixing ratio measurements for each ascent and descent. Figure 2.1 shows the flight tracks and locations of the 16 (29) profiles made from the DC-8 (WP-3D). BrO profiles are available for 16 and 17 April 2008 from the DC-8 and 12, 15, 18, 19 and 21 April 2008 from the WP-3D. We also examine measurements of other quantities for DC-8 flights of 5 and 8 April 2008, for which BrO was below the instrument detection limit. To calculate tropospheric columns from the aircraft data, the profiles were binned on an altitude grid with 500 m resolution from the surface to 7.5 km (BrO measurements are available only up to 7.5 km altitude). For each bin, the median value was selected as the BrO mixing ratio at that altitude. If the median value was less than 0, then we assumed the mixing ratio was 0.
Figure 2.1: Flight tracks (lines) and the locations (numbers) of BrO profiles measured by instruments on board the (a) NASA DC-8 aircraft during ARCTAS and (b) NOAA WP-3D aircraft during ARCPAC.
This sampling strategy was chosen to properly represent the signature of the ARCTAS and ARCPAC BrO profiles, which show that the tropospheric column is dominated by contributions from above the top of the convective boundary layer (i.e., the region of constant potential temperature; see Neuman et al. [2010] and Salawitch et al. [2010]) and, when BrO enhancements occur, they are present in relatively thick altitude segments (i.e., extent larger than 500 m). Our results are insensitive to use of mean BrO within the layers rather than median BrO.

Figure 2.2a shows the entire collection of DC-8 BrO measurements and the median DC-8 (henceforth referred to as the composite) profile. Figure 2.2b and 2.2c shows samples of different types of DC-8 BrO profiles collected during April 2008. Among the 16 DC-8 BrO profiles, 8 have shapes similar to the DC-8 composite profile, while the others show elevated BrO near the surface and/or in layers at altitudes from 2 to 4 km. WP-3D measurements of BrO exhibit larger variability than those from the DC-8 (Fig. 2.2d, e, and f).
Figure 2.2: (a) Collection of DC-8 measurements of BrO (blue dots) and DC-8 composite BrO profile (black line) and (b) DC-8 BrO profiles #7, #10, and #11, (c) DC-8 BrO profiles #12, #13, and #14, (d) similar to (a) but for WP-3D measurements of BrO, (e) WP-3D BrO profiles #5, #22, and #23, (f) WP-3D BrO profiles #25, #26, #27, and #28.
The DC-8 and WP-3D instruments reported much lower mixing ratios of BrO near the surface than have been measured in the past by LP-DOAS instruments in the springtime Arctic boundary layer [Hausmann and Platt, 1994, Tuckermann et al., 1997, Martinez et al., 1999]. The median value of surface BrO at Barrow, Alaska (71°N, 156°W) during spring 2009 was measured to be \(\sim 4\) ppt by a CIMS instrument and \(\sim 8\) ppt by a DOAS instrument (Figure A.2). We have no reason to suspect that conditions during spring 2008 were appreciably different than conditions during spring 2009. The low values of BrO observed during ARCTAS and ARCPAC could be due to preferential sampling of O\(_3\) depleted air by the two aircraft (which would titrate inorganic bromine from BrO to Br) or the fact the aircraft generally sampled to an altitude of 100 m above the surface [Neuman et al., 2010]. While it is tempting to argue that perturbations to the BrO profile due to surface release are confined to the lowest region of the marine boundary layer (a region inaccessible to the two aircraft), ground-based MAX-DOAS observations from Barrow, Alaska obtained during April 2008 suggest that when BrO was elevated, the perturbation extended to \(\sim 1\) km altitude [Salawitch et al., 2010]. In contrast, MAX-DOAS near Barrow obtained during April 2009 by another group suggest the enhancement to BrO usually occurs in the lowest 300 m, with one case where elevated BrO extended to 500 m altitude [Frieß et al., 2011]. A retrieval of the BrO profile using radiances measured in the Arctic during April 2007 by an airborne limb scanning mini-DOAS instrument also suggest elevated BrO is confined to the boundary layer [Prados-Roman et al., 2011]. The ARCTAS and ARCPAC in-situ profiles of BrO show, consistent with MAX-DOAS data from April 2008, that the highest mixing ratios of BrO tended to be observed above the boundary layer. Nonetheless, the profiles of BrO shown in Fig. 2.2 almost certainly reflect admixtures of air that has been influenced by recent surface release of active bromine with air that has achieved a background level of BrO, reflecting dilution of the surface signal. A plausible explanation for the
tendency for aircraft BrO to be lower than surface BrO is the influence of mixing. Satellite measurements of total column BrO will be influenced by this mixing process, as well. Below, we quantify the impact of elevated surface BrO on the total column.

To estimate the tropospheric BrO column from aircraft measurements, we must first make assumptions about mixing ratios between the surface and the lowest altitude sampled by the aircraft. We only take profiles with aircraft minimum altitudes less than 500 m. Then, we assume that the BrO mixing ratio in the lowest bin (surface to 500 m) is the median BrO mixing ratio between the lowest aircraft altitude and 500 m. When the aircraft did not sample up to the tropopause, we made assumptions about mixing ratios between the highest aircraft altitude and the tropopause. Here, we use the upper part of the DC-8 composite profile to fill empty upper bins. However, the DC-8 composite profile only goes up to 7.5 km, the highest altitude where BrO is sampled by the aircraft. We assume that the BrO mixing ratio between 7.5 km and the tropopause is zero. Aircraft data suggest that BrO mixing ratios at these altitudes are very small for the sampled air masses (see Fig. 2.2). It is possible that stratospheric to tropospheric transport of air could supply BrO to the upper troposphere, particularly along the western flank of Arctic low pressure systems (i.e. after low altitude tropopause systems pass over a region) [Salawitch et al., 2010]. This idea is speculative and is not considered below. Finally, layer column BrO amounts (BrO column amounts for vertical bins) are estimated using pressures and temperatures from aircraft measurements and integrated from the surface to the tropopause, to provide a tropospheric column BrO abundance. We use the tropopause height from the Modern Era Retrospective-Analysis for Research and Application (MERRA) data set (see Sect. 2.5).

Tables 2.1, 2.2, and 2.3 provide information related to the aircraft BrO profiles including the bottom and top altitudes of the aircraft in-situ profiles (Alt_B and
Alt_T, respectively), the MERRA tropopause height (H_TRP), and the in-situ tropospheric BrO column (Col.IN-SITU) obtained by integrating the aircraft profiles. Tables 2.4, 2.5, and 2.6 provide solar zenith angle (SZA), OMI reflectivity at 331 nm (R_331 nm), the difference between terrain and OMI cloud pressure (∆P_c, explained in Sect. 2.4), the satellite-derived tropospheric columns from OMI (Col.OMI) and GOME-2 (Col.GOME-2) corresponding to the in-situ aircraft BrO profiles, and the ratios of satellite-derived tropospheric BrO columns to the in-situ BrO columns (Ratio_{OMI} and Ratio_{GOME-2}) (see Section 2.2 and 3.2 for a description of the satellite data and related parameters in these tables).
Table 2.1: Information for NASA DC-8 BrO profiles shown in Fig. 2.1. “Day” means day of April 2008. \( \text{Alt}_B \) (\( \text{Alt}_T \)) means the bottom (top) altitude of the profile. “Col.” stands for BrO vertical column density (VCD). Units of BrO VCD are \( 10^{13} \text{ molecules cm}^{-2} \). “Col.\_ADDED” is the estimated amount of BrO that was added to the measured in-situ column to provide an estimate of the complete tropospheric column.

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Table 2.2: Similar to Table 2.1 but for NOAA WP-3D profiles of BrO, measured on 12 to 15 April 2008.

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Table 2.3: Similar to Table 2.1 but for NOAA WP-3D profiles of BrO, measured on 18 to 21 April 2008.

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<td>-144.2</td>
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<tr>
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Table 2.4: Satellite-derived information corresponding to DC-8 BrO profile locations shown in Fig. 2.1. Units and abbreviations are as used in Tables 2.1, 2.2 and 2.3. “Ratio\textsubscript{OMI}” means ratio of the OMI-derived tropospheric column to the in-situ column. “Ratio\textsubscript{GOME-2}” is the same but for GOME-2.

<table>
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<th>$\Delta P_c$ [hPa]</th>
<th>Col. \textsubscript{OMI}</th>
<th>Col. \textsubscript{GOME-2}</th>
<th>Ratio \textsubscript{OMI}</th>
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**Table 2.5:** Similar to Table 2.4 but for NOAA WP-3D profiles of BrO, measured on 12 to 15 April 2008.

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<th>Col. GOME-2</th>
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<th>Ratio GOME-2</th>
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<td>1.22</td>
<td>−</td>
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Table 2.6: Similar to Table 2.4 but for NOAA WP-3D profiles of BrO, measured on 18 to 21 April 2008.

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The lowest altitudes sampled by the DC-8 and WP-3D during their descents over BrO-enhanced regions were 75 and 77 m, respectively. In our analysis we use the composite DC-8 or WP-3D profile to extrapolate between the lowest sampled altitude and the surface, for each formulation of in-situ column BrO shown in Tables 2.1, 2.2, and 2.3. At times, surface BrO can reach mixing ratios as high as 40 pptv [Liao et al., 2011a]. We have assessed the impact of elevated surface BrO on our analysis of aircraft, satellite, and modeled stratospheric columns by conducting a probability distribution function for daytime surface BrO, observed at Barrow, Alaska. Two thirds of the time, surface BrO is below 8 pptv (Figure A.2). A uniform distribution of 8 pptv of BrO between the surface and 75 m altitude would contribute $0.18 \times 10^{13}$ cm$^{-2}$ to the column, an amount much smaller than the in situ and satellite-based columns discussed throughout the paper (see caption, Figure A.2). Levels of BrO reaching 40 pptv below the aircraft would contribute $1 \times 10^{13}$ cm$^{-2}$ to the column if the BrO were present, at this amount, uniformly between the surface and 75 m altitude. The PDF analysis shows that while surface BrO did reach 40 ppt in spring 2009, such occurrences were rare. While layers of highly elevated BrO below the aircraft could on occasion compromise our comparisons, our overall conclusions are robust because surface measurements indicate only on rare occasion are BrO enhancements large enough to significantly perturb the column.

In addition to BrO, other aircraft trace gas measurements are useful for inferring halogen chemistry and air mass characteristics. For example, $r_2$ was measured by CIMS instruments from both aircraft. Laboratory studies revealed that the Br$_2$ signals include contributions from HOBr that reacted on inlet surfaces. Therefore, Br$_2$ represents the lower limit of HOBr + Br$_2$ [Neuman et al., 2010]. Henceforth, we refer to this measurement as “active bromine”. Soluble bromide was also measured from the DC-8 using mist chamber/ion chromatography. Soluble bromide may include inorganic bromine compounds such as HBr, HOBr, particulate bromine, and possibly
BrO and Br$_2$ [Ridley et al., 2003, Dibb et al., 2010, Neuman et al., 2010, Liao et al., 2011b]. Measurements of O$_3$ mixing ratios by chemiluminescence were made from both aircraft. CO was measured using tunable diode laser absorption spectroscopy on the DC-8 and UV fluorescence on the WP-3D [Jacob et al., 2010, Brock et al., 2011].

### 2.2 Total BrO slant column density from OMI and GOME-2

OMI is a nadir-viewing ultraviolet and visible (UV/Vis) sensor [Levelt et al., 2006] aboard the NASA Aura satellite that is in a sun-synchronous orbit with an overpass of 01:38 p.m. local time. The spectral resolution in the OMI UV-2 channel used to retrieve BrO columns is approximately 0.5 nm. The OMI swath width is about 2600 km. The pixel size of OMI UV-2 channel is approximately $13 \times 24$ km$^2$ at the swath center and significantly larger at the swath edges. With its wide swath, OMI provides multiple daily observations at high latitudes in spring and daily global coverage at low and middle latitudes. An obstruction outside the instrument that produces radiance errors (known as the “row anomaly”) reduced the swath coverage mainly after May 2008; it does not significantly affect the observations shown here [Claas et al., 2010].

OMI BrO SCDs are retrieved by directly fitting backscattered UV radiances to absorption cross-sections of BrO (the target gas), NO$_2$, HCHO, and SO$_2$ as well as inelastic rotational-Raman scattering (also known as the Ring effect) using a non-linear least-squares approach [Chance, 1998]. The spectral fitting window for the OMI algorithm is 319 to 347.5 nm. BrO cross sections from Wilmouth et al. [1999] are used. The total fitting uncertainty of OMI BrO total column typically ranges from 15 to 51% [Salawitch et al., 2010, Kurosu and Chance, 2011].

GOME-2 is a 4 channel UV/Vis nadir viewing instrument operating on Metop-A since January 2007 [Callies et al., 2000]. It has a local equator crossing time of 9:30 am in the descending node, a swath width of 1920 km, and a spatial resolution of
40 \times 80 \text{km}^2. For the BrO SCD retrieval used here [Begoin et al., 2010], measurements in the window 336 to 347 nm are used, where GOME-2 has a spectral resolution of about 0.3 nm. SCDs of BrO are retrieved using the standard DOAS approach. Absorption due to BrO [Wahner et al., 1988], O$_3$, NO$_2$, and the effects of rotational-Raman scattering are included. The uncertainty of the GOME-2 BrO total SCDs is 10 to 30\% depending on solar zenith angle (SZA) and surface albedo. The uncertainty has both random and systematic contributions from spectral interferences and the cross-sections. As a result of throughput loss of the UV channels, random errors have increased since launch. However, this effect is not significant during the time period studied here [Dikty et al., 2011].

All of the analyses conducted in this study are based on the assumption that the magnitude of satellite-derived slant column is correct. A detailed analysis of the errors involved in deriving satellite-derived slant columns of BrO is beyond the scope of this paper. There are potentially large uncertainties associated with the derivation of both slant and vertical absolute column BrO related to the choice of spectral fitting windows and various DOAS parameters (e.g., polynomial order, orthogonalizations) used in retrievals as well as assumed BrO vertical profiles (G. Mount, private communication, 2011; DLR, 2009).

For all analyses conducted in this study, we use level 2 (time-ordered) satellite data. Level 2 data provide the best temporal match between satellite observations and aircraft in-situ and ground-based observations.

### 2.3 Stratospheric BrO column

A model simulation of stratospheric bromine species is used to estimate the spatial structure of the stratospheric BrO column. Our approach is similar to that described by Salawitch et al. [2010], but differs in that they showed results for a fixed local solar time of 01:30 p.m. whereas here we calculate BrO along each OMI or GOME-2
orbit, which spans a range of local solar times near the pole. A photochemical steady state (PSS) model is constrained to profiles of temperature, O$_3$, NO$_y$, Cl$_y$, H$_2$O, CH$_4$, sulfate aerosol surface area, etc. output from a run of Whole Atmosphere Community Climate Model (WACCM) [Garcia et al., 2007] conducted using meteorological fields for spring 2008. Salawitch et al. [2010] provide a description of this WACCM run, which was conducted to support the Stratosphere-Troposphere Analyses of Regional Transport 2008 campaign.

We specify the vertical distribution of Br$_y$ (the total bromine content of all inorganic bromine species) input to the PSS model, based on a relation between Br$_y$ and CFC-12. Profiles of CFC-12 are from the NASA Global Modeling and Assimilation Office (GMAO) Goddard Earth Observing System Data Assimilation System Version 5 (GEOS-5) [Rienecker et al., 2007] assimilation for spring 2008. Salawitch et al. [2010] evaluated the sensitivity of stratospheric partial column BrO to the amount of stratospheric bromine supplied by VSL bromocarbons, termed VSL Br$_y$. The value of VSL Br$_y$ is quite uncertain: Table 1-9 of [WMO, 2011] gives a range of 1 to 8 ppt for this quantity.

The PSS model is used to find BrO, at the local solar time of each OMI or GOME-2 orbit, assuming production and loss of all species are in balance over a 24 h period of time for a stationary air parcel. Then, BrO abundance is integrated from the pressure of the tropopause (WMO definition of the thermal tropopause, based on GEOS-5 temperature) to 0.01 hPa, yielding stratospheric column BrO. The diurnal variation of the stratospheric column BrO must be considered in estimating tropospheric BrO column, especially for the high SZAs of the level 2 data examined below. Further details of the PSS model are provided in Sect. 6.3.2 of SPARC CCMVal [2010], and references therein.

As noted above, the Br$_y$ versus CFC-12 relation used here consists of a baseline value of Br$_y$ plus 7 ppt (representing stratospheric injection of bromine in the form of
product gases of VSL bromocarbons). Baseline Br\textsubscript{y} is set to zero at the tropopause and represents, above the tropopause, supply of bromine from the decomposition of methyl bromide (CH\textsubscript{3}Br), halons, as well as dibromomethane (CH\textsubscript{2}Br\textsubscript{2}). Other groups place CH\textsubscript{2}Br\textsubscript{2} into the definition of VSL Br\textsubscript{y}. We consider CH\textsubscript{2}Br\textsubscript{2} to be part of baseline Br\textsubscript{y}, however, because this compound is observed above the tropical tropopause (e.g. Wamsley et al., 1998). The total stratospheric Br\textsubscript{y} burden of 26 ppt used here is at the upper end of the present range of uncertainty (e.g. WMO, 2011). This formulation (7 ppt for VSL Br\textsubscript{y} plus baseline that includes CH\textsubscript{2}Br\textsubscript{2}) was chosen because it results in best agreement between OMI total column BrO and the sum of stratospheric (modeled) and tropospheric (measured) partial column BrO. A wide range of values for VSL Br\textsubscript{y} yields “reasonable agreement” between these two terms when all uncertainties are considered (Fig. 6, Auxiliary Material, Salawitch et al., 2010). Despite the considerable uncertainty in stratospheric Br\textsubscript{y} and the impact of this parameter on tropospheric BrO column inferred from satellite observations of total column BrO, our approach of using a value for stratospheric Br\textsubscript{y} near the upper end is supported by the generally close quantitative agreement between inferences of tropospheric BrO column from the satellites and the in-situ data, described below. As noted previously, the fields of stratospheric BrO used here are approximately 27\% larger than those reported by Theys et al. [2011] (with percentage differences that vary considerably with respect to location as shown in Figure A.1).

2.4 Cloud parameters

Optically thick clouds shield the underlying atmosphere from satellite sensors. The OMI rotational Raman (RR) cloud product [Vasilkov et al., 2008] is used to infer information about the shielding effects of clouds over snow and ice [Vasilkov et al., 2010]. This product provides an estimate of the scene (combined cloud and surface) pressure over snow and ice surfaces. The Near-real-time SSM/I EASE-grid daily
global Ice and snow concentration and Snow Extent (NISE) data set [Nolin et al., 1998] is used to identify snow and ice-covered pixels. When the difference between scene and terrain pressure ($\Delta P_c$) exceeds 250 hPa we infer that clouds have led to significant shielding of tropospheric BrO from the satellite sensor [Vasilkov et al., 2010], as discussed below.

We use level 3 (gridded) cloud optical thickness and cloud-top pressure retrievals from the MODerate-resolution Imaging Spectroradiometer (MODIS) [Platnick et al., 2003] on the NASA Aqua satellite as a secondary check on our detection of shielding clouds. Analysis of cloud effects on the satellite-derived tropospheric BrO columns is presented in Sect. 3.1.2.

### 2.5 Meteorological data sets

Tropopause heights are inferred along the flight tracks using tropopause pressure and geopotential height profiles from MERRA, a reanalysis based on GEOS-5 system [Rienecker et al., 2007] provided by NASA Global Modeling and Assimilation Office (GMAO). These tropopause heights are used to integrate the aircraft BrO profiles and are given in Tables 2.1, 2.2 and 2.3. MERRA also provides meteorological parameters including sea level pressure, wind speed, and planetary boundary layer height. These meteorological data are used to infer relationships between BrO enhancements and the meteorological conditions. MERRA parameters are provided at 0.5° latitude × 0.667° longitude resolution. All parameters except geopotential height profiles are provided hourly; geopotential height is given every 6 h.
CHAPTER III

ANALYSIS OF SATELLITE-DERIVED ARCTIC TROPOSPHERIC BRO COLUMNS IN CONJUNCTION WITH AIRCRAFT MEASUREMENTS DURING ARCTAS AND ARCPAC

3.1 Derivation of tropospheric BrO vertical column density

3.1.1 Residual method to obtain tropospheric BrO vertical column density

Tropospheric column BrO can be obtained from satellite total column BrO retrievals using the residual method [e.g. Theys et al., 2011, and references therein]. Here, as in Theys et al. [2011], we calculate tropospheric BrO vertical column densities (VCD\textsubscript{Trop}) for each OMI or GOME-2 pixel using the derived BrO total slant column (SCD\textsubscript{Total}), an estimate of stratospheric BrO vertical column (VCD\textsubscript{Strat}), stratospheric air mass factors (AMF\textsubscript{Strat}), and an estimate of the tropospheric air mass factor (AMF\textsubscript{Trop}) as follows:

\[
\text{VCD}_{\text{Trop}} = \frac{\text{SCD}_{\text{Trop}}}{\text{AMF}_{\text{Trop}}} = \frac{(\text{SCD}_{\text{Total}} - \text{SCD}_{\text{Strat}})}{\text{AMF}_{\text{Trop}}} = \frac{(\text{SCD}_{\text{Total}} - \text{VCD}_{\text{Strat}} \cdot \text{AMF}_{\text{Strat}})}{\text{AMF}_{\text{Trop}}}.
\]

(6)

The SCD of a given absorber seen by a satellite sensor is defined as the amount of the absorber along an average light path taken by photons as they travel from the sun, through the atmosphere, and back to the sensor. The SCD is affected by scattering and absorption within the atmosphere as well as reflection off the surface and clouds.

For a given altitude range (denoted by a subscript \(z\)), The air mass factor (AMF\textsubscript{z})
is used to convert SCD$_z$ to VCD$_z$ as follows:

$$\text{AMF}_z = \frac{\text{SCD}_z}{\text{VCD}_z}. \quad (7)$$

The sensitivity of UV radiance measurements to the BrO layer amounts varies with altitude. This variation depends on viewing geometry (mainly solar zenith angle, SZA, and view zenith angle, VZA), surface albedo, cloud effects, and the vertical BrO profile. We must account for this varying sensitivity in the AMF. Using the optically thin absorber assumption, the AMF$_z$ can be formulated as

$$\text{AMF}_z = \frac{\text{SCD}_z}{\text{VCD}_z} = \frac{\int W(z)N(z)dz}{\int N(z)dz}, \quad (8)$$

[Palmer et al., 2001, Theys et al., 2011], where $z$ is altitude, $N(z)$ is the number density profile of the absorber, and $W(z)$ is the weighting function profile that represents all the parameters influencing the AMF except the vertical profile of the absorber.

Here, we use estimates of AMF$_{\text{Strat}}$ provided in the OMI and GOME-2 total BrO column products. AMF$_{\text{Strat}}$ for GOME-2 is computed using the SCIATRAN radiative transfer model [Rozanov et al., 2005]. The OMI algorithm uses a different approach to calculate AMF$_{\text{Strat}}$; BrO VCDs are obtained by applying wavelength- and albedo-dependent AMFs to the BrO absorption cross sections prior to fitting. It is assumed that all BrO resides in the stratosphere in this procedure. An effective AMF is then defined as the ratio of the SCD (derived independently as described in Sect. 2.2) to this VCD. We assume that AMF$_{\text{Strat}}$ is represented by this effective air mass factor.

We compute the tropospheric AMF$_{\text{Trop}}$ at 344.6 nm using the LInearized Discrete Ordinate Radiative Transfer (LIDORT) model [Spurr et al., 2001]. We use a clear scene assumption in the AMF calculation. The DC-8 composite profile shown in Fig. 2.2a is taken as the default tropospheric BrO profile in the AMF$_{\text{Trop}}$ calculation. However, we use individual aircraft profiles of BrO, from both the DC-8 and WP-3D, to calculate AMF$_{\text{Trop}}$ for point-to-point comparisons between aircraft in-situ and satellite-derived tropospheric columns.
We have generated a look-up table of AMF$_{\text{Trop}}$ for various representative surface albedos and viewing geometries. A 5° interval for SZA and a 2.5° interval for VZA are used. Because the weighting functions are highly dependent on surface albedo, we use step-widths of 0.1 in the range [0,1] for surface albedo.

The calculation of AMF$_{\text{Trop}}$ uses the derived OMI reflectivity at 331 nm for each pixel, from the OMI Total Ozone Mapping Spectrometer (TOMS) total O$_3$ product, as a proxy for surface albedo. We thus avoid using a surface albedo climatology, which can lead to AMF errors when the climatology differs from the actual surface albedo. Such errors can occur when the snow or sea ice distribution differs from the climatological mean, which is of particular concern at high latitudes during spring.

3.1.2 Sensitivity of the derived tropospheric column to SZA, surface albedo, and clouds

Here, we investigate the sensitivity of the satellite-derived tropospheric BrO retrieval to SZA, surface albedo, and clouds. Theys et al. [2011] analyzed the dependence of the weighting function on surface albedo and clouds for SZA = 45°. Here, we extend this analysis to a wider SZA range (40°–90°) relevant to polar observing conditions; at high latitudes (>60° N) during the day in the early spring SZA is >50°. Figure 3.1a shows that when the sun is relatively high in the sky (SZA ≤~60°), OMI and GOME-2 should have good sensitivity to tropospheric BrO for surface albedos >0.5. Sensitivity is significantly reduced for darker surfaces (e.g. albedo = 0.1). Sensitivity to tropospheric BrO decreases with increasing SZA; there is significantly lower sensitivity at 80° even for a surface albedo of 0.5. Figure 3.1b shows that at high surface albedo (0.9), there is increased sensitivity to BrO near the surface in addition to good overall tropospheric sensitivity for SZA up to 80°.
Figure 3.1: BrO weighting functions at 344.6 nm for various solar zenith angles (SZA) and surface albedos (A), for (a) clear sky conditions and (b) clear and cloudy sky conditions, where the cloudy case is for an optically thick cloud (cloud extinction coefficient = 300) at 2.5 km altitude.
The effect of optically thin clouds and aerosols on the sensitivity of OMI and GOME-2 to tropospheric BrO is dependent on surface reflectivity and viewing geometry. Vasilkov et al. [2010] showed that the sensitivity of UV satellite measurements to trace gas absorption near the surface in clear skies with moderately high surface albedo (70%) is approximately the same as for substantially cloudy conditions (optical thicknesses up to about 30) over a higher albedo surface (90%); the cloud shielding effect is much reduced for high albedo surfaces. In addition, enhanced absorption takes place above a cloud of moderate to high optical thickness. According to Vasilkov et al. [2010], the UV cloud shielding effect is generally reduced over bright surfaces as compared to dark surfaces for nadir-viewing satellite sensors. For example, there is good sensitivity to tropospheric absorbers below thin clouds over bright surfaces near the nadir swath positions for SZAs ≤∼67°. On the other hand, their simulation shows that there can be UV cloud shielding at the swath edge at high solar zenith angles, even for a thin cloud (τ ∼ 0.5). The clear scene assumption in the AMF calculation is appropriate over bright surfaces (i.e. snow/ice) when the derived scene pressure is close to the terrain pressure, which indicates that any particles along the line of sight must either be a thin, non-shielding cloud or an aerosol layer (the so-called Arctic haze) common in polar regions. The optical thickness of Arctic haze during ARCTAS was reported to be up to ∼0.2 at 354 nm [Shinozuka et al., 2011]. Based on the calculations of Vasilkov et al. [2010], OMI and GOME-2 should have good sensitivity to surface BrO in Arctic haze conditions at the near-nadir swath positions, but may have reduced sensitivity at the swath edge. However, as shown by Theys et al. [2011] and in Fig. 3.1b, optically thick clouds can shield satellite measurements from absorbers including BrO.
Figure 3.2: (a) Tropospheric air mass factors (AMFs) at 344.6 nm computed using the DC-8 composite BrO profile for various surface albedos as a function of SZA; (b) similar to (a) but also showing AMFs found assuming all of the BrO is below 500 m (dotted lines). A refers again to surface albedo.
Figure 3.2a shows computed tropospheric air mass factors. As shown for the weighting functions in Fig. 3.1, tropospheric air mass factors are higher over brighter surfaces owing to increased near-surface sensitivity. However, tropospheric sensitivity begins to be lost as SZA increases; tropospheric sensitivity almost disappears when SZA > 80° even for relatively bright surfaces. In this study, we present satellite-derived BrO tropospheric columns only for SZA < 80°.

The sensitivity of AMF$_{\text{Trop}}$ to the tropospheric BrO profile is assessed in Fig. 3.2b. For a bright surface (albedo = 0.9), no significant difference exists between tropospheric AMFs computed using the DC-8 composite profile and a profile where all tropospheric BrO is contained below 500 m for SZA ≤ ∼60°. This is a consequence of good sensitivity at all tropospheric altitudes for bright surfaces and low SZA as shown in Fig. 3.1. For 60° < SZA < 75°, the sensitivity to middle- and upper-tropospheric BrO is maintained while the near-surface sensitivity drops; this leads to a small sensitivity to the profile shape under these conditions. Profile sensitivity decreases for higher SZAs as the sensitivity to the entire troposphere drops. For a darker surface (albedo = 0.4), the retrieval has lower sensitivity to BrO near the surface even when the sun is relatively high (SZA ∼ 40°). In this case, the retrieval of total column BrO and our inference of the tropospheric column will be sensitive to the shape of the profile of BrO in the troposphere. If most of the tropospheric column happened to originate from BrO in the lowest 500 m of the atmosphere, then our inference of tropospheric column would be biased low over dark surfaces.

We next examine tropospheric column BrO estimated from OMI in the context of the different sensitivities discussed above. Figure 3.3 shows OMI total, the model stratospheric, and the derived tropospheric BrO columns for OMI orbit 20050 on 22 April 2008 along with several parameters that impact the sensitivity of the radiance observation to the tropospheric BrO column. These parameters include 331 nm reflectivity (a proxy for surface albedo), Δ$P_c$, and SZA.
Figure 3.3: Measured total vertical column BrO, reflectivity, and cloud pressure for OMI orbit 20050 at 03:23 UTC on 22 April 2008 as well as calculated stratospheric column BrO for the time of OMI overpass and the inferred tropospheric residual column BrO. The panels highlight reduced tropospheric BrO over low albedo surfaces (black box) and in the presence of shielding clouds (orange box). See text for more details and discussion.
OMI total column BrO in Fig. 3.3a, within the region of the Barents Sea (black box), shows significant gradient. The region of high total BrO towards the upper left hand corner of the black box is associated with enhanced stratospheric burden (Fig. 3.3b, orange). Another region of large total BrO (lower portion of black box) is associated, by our analysis, with an enhanced tropospheric burden (Fig. 3.3c, red). This region of enhanced tropospheric BrO occurs over a bright portion of the Barents Sea (Fig. 3.3d, crimson, indicating snow or ice). Nearly zero tropospheric BrO column amounts are obtained over low surface albedo areas (OMI reflectivity < 0.5) of the Barents Sea (Fig. 3.3d, blue). Here, the retrieved total columns are generally less than retrieved columns over adjacent areas with higher reflectivity, leading to low tropospheric column BrO over parts of the Barents Sea. However, satellite-derived tropospheric BrO may not be reliable when the surface albedo is low. While it is possible tropospheric column BrO was truly low on 22 April over this region of the Barents Sea due to the lack of snow or ice leads that may be needed for bromine activation, it is also possible that our inference of low tropospheric BrO could either be a result of limited sensitivity over dark surfaces or an over-estimation of the stratospheric burden. The complicated sensitivity of satellite-derived tropospheric BrO to surface reflectivity and stratospheric burden requires concerted future study.

In Fig. 3.3, the cloud shielding effect is shown over Siberia (marked with orange rectangles), where the OMI rotational Raman cloud product reports $\Delta P_c > 250$ hPa. Aqua MODIS also indicates cloud cover, with cloud top pressures between 450 and 500 hPa (not shown). Our derived tropospheric columns over this region are significantly less than those from adjacent areas without cloud cover. This supports the presence of tropospheric BrO mainly at altitudes below 450–500 hPa and the ability of OMI and other nadir viewing sensors to capture information about tropospheric BrO over bright polar regions, with SZA up to $\sim 80^\circ$ when the sky is clear. Theys et al. [2011] also used the cloud shielding effect to infer an amount of BrO in the
troposphere, but applied this approach only in the tropics.

For completeness, we include all data in our analysis of aircraft measurements. However, we note aircraft soundings for which observing conditions likely produce large errors in the satellite-derived tropospheric BrO columns. In mapping surface BrO enhancement events with satellite retrievals, we use only data that has good sensitivity to the troposphere: SZA < 80°, ∆Pc < 250 hPa, and surface albedo > 0.7.

3.1.3 Uncertainty analysis

For a single pixel, we express the uncertainty of the tropospheric VCD, σ_{VCD_{Trop}}, in a simplified form:

$$\sigma_{VCD_{Trop}}^2 = \left( \frac{\sigma_{SCD_{Total}}}{AMF_{Trop}} \right)^2 + \left( \frac{\sigma_{VCD_{Strat}}}{AMF_{Trop}} \right)^2$$

[Boersma et al., 2004, De Smedt et al., 2008, Theys et al., 2011], where σ_{SCD_{Total}} is the uncertainty of total slant column, σ_{SCD_{Strat}} is the uncertainty of stratospheric slant column, and σ_{AMF_{Trop}} is the uncertainty of tropospheric air mass factor. This representation assumes that the different types of uncertainties are uncorrelated with each other.

We take the fitting uncertainty derived from observed minus fitted radiances for each pixel as σ_{SCD_{Total}}. This assumes that the SCD_{Total} error has a zero mean. The average fitting uncertainty at latitudes greater than 60°N is about 18%. Here, we do not consider systematic errors in SCD_{Total} or VCD_{Total}. Systematic error will generally result in either a geographically uniform over-estimate or under-estimate of total column BrO (VCD_{Total}). There is synergy between systematic error in VCD_{Total} and our prescription of the contribution of VSL species to stratospheric Br_y. If subsequent analysis shows the estimates of VCD_{Total} BrO used here are biased high by a considerable margin, then clearly we must use a smaller contribution to stratospheric Br_y to derive similar overall magnitude of tropospheric BrO. However, the geographic distribution of tropospheric BrO will not be strongly altered due to this synergy. An
exploration of the systematic error in VCD_{Total} and the implication for tropospheric BrO will occur following analysis of a ground-based, OMI BrO validation campaign conducted in Fairbanks, Alaska during April 2011.

We estimate the uncertainty of the stratospheric slant column (\(\sigma_{\text{SCD}_{\text{Strat}}}\)) by multiplying the uncertainty of the stratospheric column BrO by the stratospheric AMF. The uncertainty of the stratospheric column BrO results from a root-sum-squares combination of three terms: the uncertainty in chemical kinetics that govern the BrO to Br\(_y\) ratio, the uncertainty in the dynamics that govern CFC-12 (and hence Br\(_y\) due to Source Gas Injection), and the uncertainty in VSL Br\(_y\). The uncertainty in chemical kinetics is evaluated by varying the rate constant of individual chemical reactions (including \(J\) values) by the 1-sigma estimate of uncertainty given by S. P. Sander et al. [2006]. The most important chemical term is BrO + NO\(_2\) forming BrNO\(_3\); this rate constant is uncertain by about a factor of 2 at 220 K. The uncertainty due to dynamics is found by repeating calculations for plus and minus 4\% variations in the abundance of CFC-12, because comparison to aircraft observations showed CFC-12 from GEOS-5 was accurate to within \(\pm 4\%\) in the lower stratosphere [Salawitch et al., 2010]. Finally, the uncertainty due to VSL Br\(_y\) is set at \(\pm 27\%\), which represents the mean difference in stratospheric column BrO resulting from our approach to handling this term compared to the approach of Theys et al. [2011].

Here, we neglect the uncertainty in the tropospheric air mass factor as it is relatively small compared with the other error sources. For example, Theys et al. [2011] describe sensitivity tests that show use of a single wavelength for the weighting function profiles leads to an error of less than 5\%. We find that the profile dependence produces error between 7 and 13\% for bright surfaces.
3.2 **Comparisons of aircraft in-situ measurements with satellite retrievals**

In this section, comparisons between aircraft and satellite inferences of tropospheric column BrO are shown for cases where agreement is good, agreement is poor but the cause for disagreement is understood, and agreement is poor and not well understood. These comparisons are performed to evaluate the veracity of the satellite-derived tropospheric column BrO product. We present a series of figures for various ARCTAS and ARCPAC flights. For each, we show maps of the satellite-derived total and tropospheric columns and time series of various aircraft measurements, collocated satellite-derived BrO data, and estimated stratospheric BrO amounts with vertical error bars derived using the formulation given in Sect. 3.1.3. Satellite-derived data are shown for all conditions (e.g. even in the presence of thick clouds that likely shield the surface). We also show plots of various parameters that affect the sensitivity of the satellite measurements to tropospheric BrO and note times and places where satellite observing conditions are not ideal for inferring reliable tropospheric BrO amounts.

### 3.2.1 Cases of good agreement between satellite and in-situ data

Figure 3.4 shows maps of the OMI and GOME-2 measurements of total column BrO as well as maps of tropospheric column BrO inferred from each satellite sensor on 17 April 2008. The track of the DC-8 and locations of BrO profiles, for the ARCTAS flight on this date, are marked. The bottom panel shows geographic regions of interest. OMI and GOME-2 orbits closest in time to the ARCTAS flight are used.
Figure 3.4: Maps on 17 April 2008 of (top left) OMI total vertical column density (VCD) of BrO, (top right) inferred OMI tropospheric VCD of BrO, (middle left) GOME-2 total VCD of BrO, and (middle right) inferred GOME-2 tropospheric VCD of BrO column. Negative values are included as dark blue. The flight track of the DC-8 on 17 April 2008 is indicated by black lines; aircraft profile locations where BrO was measured are marked with black diamonds. Geographic regions used in the paper are denoted in the bottom panel.
The tropospheric column BrO maps have different spatial structure than maps of total column BrO. For example, total column BrO is highest over the north coast of Canada. These local maxima are significantly reduced in the tropospheric columns. A long tail of enhanced values near the North Pole is pronounced in the maps of tropospheric column BrO and is not as prominent in total column BrO. As discussed by Salawitch et al. [2010] and Theys et al. [2011], inferring tropospheric BrO from the total BrO column requires an accurate estimate of the stratospheric contribution to the total column, because variability induced by stratospheric dynamics is comparable in magnitude to variability induced by the surface release of bromine. Figure 3.4 is a perfect illustration of this point, which as noted above has been overlooked in many prior studies. Essentially all derivations of tropospheric column BrO prior to Salawitch et al. [2010] and Theys et al. [2011] relied on the use of zonally fixed stratospheric BrO, which, as first pointed out by Theys et al. [2009], will lead to large errors.
Figure 3.5: Time series plots for NASA ARCTAS DC-8 flight of 17 April 2008. For bottom three panels, line colors corresponding to the same color y axis. The top panel shows various estimates of VCD of BrO along the DC-8 flight track, including OMI total column, the modeled stratospheric column, and tropospheric columns based on aircraft in-situ measurements (black dots) as well as our analysis of OMI retrievals (red squares). The error bars of the tropospheric columns are described in the text. The two middle panels show various aircraft in-situ measurements along with the flight altitude. The bottom panel shows OMI reflectivity at 331 nm and cloud retrieved parameters from OMI and MODIS, as described in the text. Soluble bromide is not plotted when the reported mixing ratio was below the detection limit (1–6 pptv).
Figure 3.5 shows time series plots of DC-8 flight data and collocated satellite measurements on 17 April 2008. We indicate time in hours relative to the starting date of the flight; therefore times greater 24:00 refer to the following day. DC-8 data include aircraft altitude, in-situ O$_3$, CO, BrO, active bromine and soluble bromide. Satellite data include OMI-derived BrO columns (total and tropospheric), $\Delta P_c$, OMI 331 nm reflectivity, and Aqua MODIS cloud optical thickness. Estimated errors of satellite-derived BrO columns are presented as vertical error bars. The tropopause height, as given in the MERRA data set, is at approximately $\sim$7 km during the flight (not shown). CO and O$_3$ data suggest that there were no stratospheric intrusions for any of the collected tropospheric BrO profiles. Obvious signatures of stratospheric air (low CO and high O$_3$) were occasionally seen, but only when the aircraft was flying near or above the tropopause.

Satellite-derived BrO tropospheric columns (orange line) are $\sim$zero near Fairbanks (see top panel of Fig. 3.5). As discussed in Sects. 3.1.1 and 3.1.2, tropospheric BrO information is not reliably retrieved when surface reflectivities are less than $\sim$0.4, which is the case near Fairbanks (see the fourth panel of Fig. 3.5).

Both total and tropospheric column BrO show a slight dip near the pole, where the OMI and MODIS cloud products indicate shielding clouds (middle of Fig. 3.5). Here, the low values of derived tropospheric BrO over optically thick clouds likely result from the cloud shielding effect [Theys et al., 2011]. Such clouds may also reduce tropospheric BrO columns by slowing photochemical production of active bromine. On the other hand, high values of satellite-derived tropospheric column are found along the DC-8 flight path both before the descent in altitude at $\sim$24:15 UTC and after the plane has ascended. Data from both OMI and MODIS indicate optically thick clouds were present during and after the ascent. The high value of satellite-derived tropospheric column during and after ascent could be due to the presence of BrO above or within these clouds. Profile #12 (see Fig. 2.2) shows a plume of
enhanced BrO at about 3 km altitude, which could have been above the cloud deck.

In-situ tropospheric column BrO, marked as black dots in the time series plot (the first panel of Fig. 3.5), are obtained by integrating in-situ BrO mixing ratio profiles as described in Sect. 2.1. Uncertainties in the in-situ columns are shown as black vertical error bars. The horizontal bars indicate the distance covered by corresponding flight ascents/descents. OMI tropospheric BrO columns for point-to-point comparisons with aircraft data, shown as red squares, are obtained by averaging pixels over the corresponding flight segment.

The OMI tropospheric BrO columns exhibit a magnitude and variability similar to that of the DC-8 in-situ columns. For example, profile #12 (see Figs. 2.1, 2.2 and 3.5) reports high BrO mixing ratios from the surface to ~4 km; the OMI tropospheric BrO column (as well as GOME-2, shown in Fig. 3.4 but not in Fig. 3.5) is also relatively high at the same location.

However, some discrepancies appear, perhaps because the aircraft captures small scale features that the satellite observations cannot resolve. For example, in-situ profile #13 (see Figs. 2.2 and 3.5) shows a relatively large value for tropospheric column BrO, caused by enhanced BrO mixing ratios in a layer near 2 km. This could be a local event as the nearby aircraft column #14 does not show such an enhancement. The satellite retrieval agrees well with profile #14 but underestimates the abundance of BrO measured in profile #13. Overall, OMI tropospheric column BrO quantitatively agrees with the DC-8 in-situ column BrO to within the respective estimated uncertainties. The comparison between satellite and in-situ tropospheric column BrO for this flight is further quantified by Liao et al. [2011b]. These comparisons demonstrate the ability of OMI and GOME-2 to capture the magnitude and spatial distribution of tropospheric column BrO over bright surfaces for clear conditions.
3.2.2 Examination of differences between satellite-derived and in-situ tropospheric columns of BrO

We next describe two cases of apparent inconsistency between satellite and in-situ observations. Figures 3.6 and 3.7 show data corresponding to the DC-8 flight on 5 April 2008. In-situ measurements did not detect ozone depletion or high bromine over Hudson Bay, whereas OMI and GOME-2 both report high total column BrO over vast regions of eastern Canada including Hudson Bay (Salawitch et al., 2010). BrO measurements were not available to produce profiles for this flight, but active bromine and soluble bromide were reported. Figure 3.7 shows that the derived tropospheric BrO column from OMI is low over Eastern Canada, where the total column is high. Similar results are found for GOME-2 (Fig. 3.6). The OMI measurement of total column BrO shows very similar magnitude and structure as the calculated stratospheric column BrO. This suggests that the elevated total column BrO over Eastern Canada could be a consequence of high stratospheric columns. Furthermore, the in-situ CO and O₃ measurements indicate no evidence of stratospheric air below 6 km altitude in ascending or descending flight tracks near Hudson Bay.
Figure 3.6: Similar to Fig. 3.4 but for the DC-8 flight on 5 April 2008. The GOME-2 measurement of total vertical column BrO is quite similar to OMI measurement and is therefore not shown.
Figure 3.7: Similar to Fig. 3.5 but for the DC-8 flight on 5 April 2008. Aircraft in-situ BrO measurements are not available for this flight because BrO was below the detection limit.
Figures 3.8 and 3.9 show similar maps and time series, respectively, for the DC-8 flight on 8 April 2008. GOME-2 observations during this flight are not available. Aircraft in-situ BrO measurements are also not available for this flight. This makes it difficult to evaluate the validity of the OMI-derived BrO tropospheric columns. The aircraft observed a severe ozone depletion event ([O$_3$] < 1 ppbv) and high levels of active bromine and soluble bromide near Alert (marked as black square), whereas the OMI total column BrO is relatively low and the inferred OMI tropospheric column BrO is at or near background levels. The ODE observed by the DC-8 on 8 April 2008 is thus not apparent in the satellite measurement of BrO on this date.

There are segments where inferred OMI tropospheric BrO is negative, such as the early time period of 8 April 2008 (see Fig. 3.9). These values of negative BrO could be indicative of an over-correction for the stratospheric column, measurement noise for total column BrO, or some other error in calculation. Overall, we do not find extended regions of negative tropospheric BrO (either spatially or temporally).
Figure 3.8: Similar to Fig. 3.4 but for the DC-8 flight on 8 April 2008. GOME-2 data are not available for this day. The location of severe ozone depletion and high active bromine sampled by the DC-8 is marked by the black squares.
Figure 3.9: Similar to Fig. 3.7 but for the DC-8 flight on 8 April 2008.
As described in Eqs. (3), (4), and (5), BrO catalyzes the loss of ozone and, at the same time, BrO is produced by reaction of Br with O₃. This suggests that high BrO concentrations cannot be maintained if O₃ is completely depleted, even though plenty of active bromine may be available. This explanation is consistent with other observations [e.g. Hausmann and Platt, 1994, Tuckermann et al., 1997, Neuman et al., 2010, Frieß et al., 2011] that show high BrO concentrations only when ozone is partially depleted. The presence of a severe ozone depletion event is therefore indicative of an expectation of little or no enhancement of tropospheric BrO, despite the presence of large amounts of active bromine and soluble bromide. Liao et al. [2011a] also discuss the relationship between O₃ and BrO; little BrO exists when O₃ is low due to the partitioning between Br and BrO.

OMI tropospheric column BrO shows enhancements in the vicinity of the 8 April 2008 ozone depletion event, during the 36h time period prior to this flight (not shown). This enhancement is likely connected with the nearly complete removal of ozone observed by the DC-8 aircraft.

3.2.3 Cases of disagreement between satellite and in-situ data

Figure 3.10 shows maps of satellite-derived BrO columns and the flight track of the NOAA ARCPAC WP-3D on 19 April 2008. The OMI and GOME-2 data both indicate enhanced total and tropospheric columns of BrO over the Chukchi Sea. The aircraft flew into the area of elevated satellite-derived tropospheric column BrO at the northernmost extent of the track.

Figure 3.11 shows time series plots for the flight of 19 April 2008. Two types of disagreement between in-situ and satellite tropospheric column BrO are observed. First, satellite-derived tropospheric column BrO is considerably less than the in-situ column over surfaces with moderate reflectivity in the presence of clouds. For example, the OMI estimate is less than the WP-3D columns (#19 and #23) near
Fairbanks. These observations were obtained in a region where the surface albedo is between 0.4 and 0.6, in the presence of low level clouds. This type of disagreement may be expected from the sensitivity studies described in Sect. 3.1.2. It is a challenge for satellite sensors to accurately retrieve tropospheric BrO over a moderately reflective surface in the presence of clouds.
Figure 3.10: Similar to Fig. 3.6 but for the WP-3D flight of 19 April 2008.
Figure 3.11: Similar to Fig. 3.5 but for the WP-3D flight on 19 April 2008. Soluble bromide was not measured from the WP-3D.
The second type of disagreement during this flight is illustrated by profile #21 (see Figs. 2.1, 2.2 and 3.11), where both OMI and GOME-2 tropospheric column BrO are significantly enhanced (only OMI is shown in the first panel of Fig. 3.11), whereas the in-situ BrO column does not show evidence of enhancement. Artifacts from clouds or surface albedo are not the likely explanation for this disagreement, as the surface was bright and no clouds are evident in ∆Pc (fourth panel, Fig. 3.11). The highest level of active bromine is observed from the aircraft sensor at the same location where tropospheric column BrO from both satellite sensors show enhancements. O3 mixing ratios of ∼10 ppbv suggest the aircraft sampled a partial ozone depletion event [e.g. Ridley et al., 2003]. Prior DOAS observations in the Arctic suggest that BrO should still be present for O3 levels of ∼10 ppbv [Hausmann and Platt, 1994, Tuckermann et al., 1997]. The disagreement between satellite and aircraft BrO could be related to the different spatial scales, and/or vertical coverage spanned by the respective instruments: the aircraft does not observe the complete profile and the field of view encompassed by the satellite covers a larger area than that sampled by the airplane. The aircraft may have missed an important part of the BrO profile as the minimum sampled height was 151 m above the surface. Timing may also be a possible explanation for the disagreement, as the aircraft flew near the location of the enhancement about five hours after the OMI overpass and about three hours after that of GOME-2.

Our analysis indicates that the large enhancement in total column BrO seen by OMI and GOME-2 over the Chukchi Sea on 19 April 2008 was tropospheric in origin, rather than stratospheric. The WP-3D aircraft recorded highly elevated active bromine and partially depleted O3 near the surface at this precise location, indicating recent association with elevated BrO. It is possible the satellite perturbation was caused by the presence of BrO at higher altitudes than those sampled by the aircraft at this location. By the time the aircraft reached 3 km altitude, it had left the
Figure 3.12: Similar to Fig. 3.6 but for the WP-3D flight of 21 April 2008.

region of highly elevated satellite BrO (terminal point of profile #23 (see Figs. 2.2 and 3.11)). Salawitch et al. [2010] presented an analysis of ground-based Max DOAS observations of BrO for April 2008 that, together with ARCTAS and ARCPAC BrO profiles, showed important contributions to column BrO often originate within the troposphere from altitudes above the top of the planetary boundary layer.

Figures 3.12 and 3.13 show maps of the BrO columns and time series plots for the NOAA WP-3D flight of 21 April 2008, respectively. The maps show enhanced OMI total column BrO over the north shore of Canada and the Canadian Archipelago. The stratospheric model shows only a slight enhancement for these regions; the subtraction of the stratospheric model from the OMI column shows significant levels of tropospheric column BrO for much of the flight segment. Time series plots indicate that WP-3D in-situ columns #25, #26, and #27 (see Figs. 2.1, 2.2 and 3.13) are lower than the OMI tropospheric column, but the differences are not significant given the uncertainties in OMI and in-situ tropospheric columns (i.e., error bars overlap). However, the absolute magnitude of the OMI tropospheric column is about a factor of 3 larger than the in-situ BrO column. There is, however, good agreement between in-situ column #28 and the satellite-based estimate.
Figure 3.13: Similar to Fig. 3.11 but for the WP-3D flight on 21 April 2008, with tropopause pressure added (bottom panel).
WP-3D profiles #26, #27, and #28 (see Figs. 2.1, 2.2 and 3.13) were collected in close proximity (see Fig. 2.1), but show a large variation in the column amounts. As noted above, the profile #28 column agrees well with the satellite-based column whereas the other two in-situ based BrO columns are lower than the satellite-based estimates. As discussed above for other flights, this result may be explained by the fact that the aircraft captures small-scale spatial features while the satellite observes larger scales.

The overestimation of the tropospheric column BrO by OMI, compared to in-situ columns for profile #25, #26, and #27 (see Figs. 2.1, 2.2 and 3.13), could potentially be explained by a stratospheric column that is not fully removed from the total column. One factor not considered in our analysis is irreversible, cross-tropopause exchange of air parcels with elevated levels of Br_y from the stratosphere to the troposphere (STE). Such transport events occur on the western flank of Arctic low pressure systems [Salawitch et al., 2010, and references therein]. If the satellite signal were to originate from STE of Br_y, the BrO signal associated with such air parcels may not have been sampled by the WP-3D because the maximum altitudes of the WP-3D profiles #25, #26 and #27 (6 km) is well below the height of the tropopause (10.5 km). Future measurement of profiles of BrO from near the surface to the lowermost stratosphere, in the footprint of a satellite sensor after passage of an airmass with a low altitude tropopause, are needed to assess the importance of STE of bromine on the interpretation of the satellite record.

3.3 BrO explosions observed with satellites

Events of rapid enhancement of tropospheric column BrO are apparent from OMI and GOME-2 observations after adjustment for the stratospheric burden of BrO, for the time period mid-March to late April 2008. The NASA DC-8 aircraft flew into an area of enhanced tropospheric column BrO near the North Pole on 17 April 2008 (Fig. 3.4)
and the NOAA WP-3D aircraft flew near another tropospheric BrO enhancement on 19 April 2008 (Fig. 3.10). Here we examine the event near the North Pole on 17 April 2008 and similar events in more detail using only OMI retrievals. The wide orbital swath and high spatial resolution of OMI, in addition to its frequent observations at high latitudes, provide a unique view of the temporal evolution of these events.

Figure 3.14 shows the evolution of a tropospheric BrO enhancement event (“BrO explosion”) observed from 16 to 18 April 2008. Here, we only show observations when the following conditions are met, to provide reliable tropospheric BrO information as discussed in Sect. 3.1.2: SZA < 80°, reflectivity > 0.7, and ∆Pc < 250 hPa. The stratospheric column has been removed, as discussed in Sect. 3.1.1, using photochemical model output for the VSL Bry = 7 ppt simulation, for the local solar time of each OMI pixel. The major activation of BrO starts at ~22:00 UTC on 16 April and lasts for ~30 h. We see activations near the North Pole and Canadian Archipelago. The spatial features of the elevated tropospheric column BrO change rapidly, with significant variations over the course of a day.
Figure 3.14: OMI tropospheric BrO VCD for different orbits (UTC time as indicated) from 16 to 18 April 2008.
We have ruled out a stratospheric origin for the enhanced total column BrO observations near the North Pole and Canadian Archipelago on these days. Since stratospheric BrO and O$_3$ columns exhibit a significant correlation [Theys et al., 2009, Salawitch et al., 2010], the OMI measurement of total O$_3$ column can be used as a proxy for the spatial pattern of stratospheric column BrO. The comparison of the OMI tropospheric BrO column with the OMI total O$_3$ column shown in Fig. 3.15 indicates lack of correlation. Thus, the elevated region of total column BrO is not of stratospheric origin. Figure 3.15 also shows few shielding clouds near the areas with high BrO columns.

Figure 3.16 shows three tropospheric BrO enhancement events during April 2008 along with maps of sea level pressure, wind speed at 2m altitude, and planetary boundary layer height from MERRA for the closest synoptic hour. For these events, the locations of enhanced BrO columns are coincident with high near-surface wind speeds (3rd column) that are geostrophically consistent with localized low pressure systems. Neuman et al. [2010] also note high wind speed along the flight paths in conjunction with high concentrations of active bromine. Our analysis is consistent with the suggestion that strong surface winds associated with low pressure systems can trigger bromine activation via blowing snow [Yang et al., 2008, 2010, Jones et al., 2009, 2010].
Figure 3.15: (a) OMI tropospheric BrO VCD; (b) OMI total ozone column; (c) terrain minus cloud pressure for OMI orbit 19976 on 17 April 2008.
Figure 3.16: First column: OMI tropospheric BrO VCD; Second column: MERRA sea level pressure; third column: MERRA wind speed at 2 m altitude; fourth column: MERRA planetary boundary layer height; fifth column: aircraft in-situ profiles of BrO (red), active bromine (blue) and temperature (black). The rows correspond to OMI orbits. We show results for orbit 19979 on 17 April 2008 (top row), orbit 19990 on 18 April 2008 (middle row), and orbit 20018 on 19 April 2008 (bottom row). Aircraft profiles were not obtained during the time that the satellite data shown in the top row were acquired. An animated GIF of this figure is available in Supplemental Material.
Figure 3.16 also shows that the spatial structure of high tropospheric column BrO is similar to that of the planetary boundary layer (PBL) height, although there is not always a precise alignment of these features. At high latitudes where the meteorological analysis is driven primarily by satellite data, the MERRA fields may contain displacement or other errors, particularly in near-surface fields. The following discussion should therefore be considered somewhat speculative in light of these uncertainties. We provide an animated visualization as a Supplement to depict evolution of enhanced tropospheric column BrO, sea level pressure, wind speed at 2 m and planetary boundary layer height from 16 to 18 April 2008.

Neuman et al. [2010] reported a temperature inversion at 500 m during the BrO activation event of 19 April 2008, more or less consistent with the MERRA PBL height. Bromine chemistry may be related to PBL height as follows: BrO concentrations are controlled in large part by O\textsubscript{3}; a sufficient amount of O\textsubscript{3} is necessary to maintain high BrO concentrations as explained in Sec. 3.2.2. Ozone is quickly consumed in a shallow boundary layer during bromine activation [Lehrer et al., 2004, Anderson and Neff, 2008]. When O\textsubscript{3} is substantially depleted, production of BrO will cease and BrO will be destroyed by various reactions, including those in Eq. (4). On the other hand, the O\textsubscript{3} loss rate is quadratic in BrO, so a deeper boundary layer could slow down the chemical removal of O\textsubscript{3} as BrO is diluted.

The second row of Fig. 3.16 illustrates an event at 00:30 UTC on 18 April 2008 and the third row is for an event at 22:40 UTC on 19 April 2008. The two OMI orbits presented are closest in time to airplane flights into the elevated BrO and active bromine layers (profile #12 from DC-8 and #21 from WP-3D, respectively (see Figs. 2.1, 2.2, 3.5 and 3.11)). The locations of the profiles are shown as black diamonds on the maps of tropospheric BrO. The aircraft profiles of temperature, BrO mixing ratio, and active bromine mixing ratio are presented in the last column. Aircraft measurements show enhanced active bromine in the near-surface layer for both flights.
and enhanced amounts of BrO for the 18 April flight. Satellite-derived tropospheric BrO enhancements are closely related to near-surface parameters including sea level pressure, wind speed at 2 m, and PBL height. This observation suggests that the BrO activations originate at the surface, which of course is consistent with prior expectation. For the event on 18 April, a tail of enhanced BrO column exists parallel to the wind direction inferred from sea level pressure. Begoin et al. [2010] suggested long range transport of tropospheric Br\textsubscript{y} for plumes in which recycling of bromine from condensed to gas phase sustains elevated BrO. The profile for BrO measured on 18 April by the DC-8 instrument (Fig. 3.16) also shows enhancements in the free troposphere, above the top of the PBL height (∼500 m). Simultaneous profiles of CO and O\textsubscript{3} (not shown here, see Fig. 3.5b) imply no stratospheric influence. The existence of elevated BrO above the top of the PBL could be due to vigorous convection over ice leads driven by warm exposed water, with BrO then dispersed horizontally by prevailing winds [Simpson et al., 2007b, Salawitch et al., 2010]. Our results are consistent with a surface origin of elevated BrO as well as transport of Br\textsubscript{y} enriched air parcels away from the source of origin. Indeed, an animation of the panels in Fig. 3.16 (see Supplement) provides compelling evidence for this behavior.
CHAPTER IV

ESTIMATION OF FREE TROPOSPHERIC BRO USING CLOUD-SLICING TECHNIQUE

4.1 Derivation of free-tropospheric BrO volume mixing ratios (VMR) using the ensemble cloud slicing technique

Optical centroid cloud pressures (OCCP, denoted as \( P_{cld} \)) can be derived from photon-path-sensitive measurements including those sensitive to rotational Raman scattering [e.g. Joiner et al., 2004], oxygen dimer (\( O_2-O_2 \)) absorption [e.g. Sneep et al., 2008] and oxygen A-band absorption [e.g. Rozanov et al., 2004]. OCCPs are significantly different from cloud-top pressures retrieved from thermal infrared observations [Vasilkov et al., 2008, Joiner et al., 2009, 2010]; usually OCCPs are located significantly inside of clouds, while thermal IR cloud-top pressures are located near the physical tops of clouds. OCCP is more appropriate than cloud-top pressure for radiative calculations in the UV; Solar radiation at these wavelengths penetrates inside clouds to reach pressures near the OCCP value [Ziemke et al., 2009, Vasilkov et al., 2010].

BrO column measurements from nadir-viewing spectrometers such as OMI extend down to the OCCP. For example, an OMI-retrieved vertical column of BrO observed in a fully cloudy scene can be interpreted approximately as the BrO column from the OCCP to the top of atmosphere. For a partly cloudy scene, the retrieved BrO VCD can be approximated as the BrO VCD from a fully cloudy pixel with the cloud placed at an effective cloud scene pressure \( P_{scene} \). \( P_{scene} \) is defined for a given pixel as the pressure at which a Lambertian surface must be placed in order to produce the observed amount of rotational-Raman scattering [Vasilkov et al., 2010].

Over dark and moderately bright surfaces (i.e. not covered by snow and ice), we
calculate the effective cloud scene pressure, $P_{\text{scene}}$, using

$$P_{\text{scene}} = \text{CldF} \times P_{\text{cld}} + (1 - \text{CldF}) \times P_{\text{terrain}},$$  \hspace{1cm} (10)$$

where CldF is effective cloud fraction and $P_{\text{terrain}}$ is the surface pressure [Joiner et al., 2009]. This formulation assumes that tropospheric BrO is vertically well-mixed. Over snow/ice covered surfaces, the OMI rotational Raman cloud algorithm retrieves effective cloud/surface scene pressure (equivalent to $P_{\text{scene}}$) instead of OCCP, assuming an effective cloud fraction of unity.

Here, we use the cloud slicing technique to obtain monthly climatological maps of free tropospheric BrO VMR using OMI data from Oct 2004 to Apr 2008. These maps are obtained for a 5° latitude x 5° longitude grid over the globe. For each grid box, we fit the observed BrO column over clouds versus effective cloud scene pressure ($P_{\text{scene}}$) using simple linear regression, i.e.,

$$\text{VCD from } P_{\text{scene}} \text{ to TOA} = A \times P_{\text{scene}} + B.$$  \hspace{1cm} (11)$$

BrO VMR can be obtained from the slope, $A$, of this fit using

$$\text{BrO VMR (pptv)} = 417.7 \times A,$$  \hspace{1cm} (12)$$

where the unit of $A$ is $10^{13}$ molec cm$^{-2}$ per hPa. OMI pixels with an effective cloud fraction $> 0.7$ and reflectivity $> 0.6$ are used for this calculation where cloud retrievals are most accurate [Joiner et al., 2012] and sensitivity to tropospheric BrO is highest [Theys et al., 2011, Choi et al., 2012]. The BrO VMR obtained is valid for the range of altitudes spanned by $P_{\text{scene}}$ and assumes vertical homogeneity within that range. We also assume that both stratospheric column BrO and tropospheric mean BrO VMRs are not highly variable over a given ensemble grid box and time period (1 month).
4.2 Results

Figure 4.1 shows examples of ensembles collected at various $5^\circ \times 5^\circ$ grid boxes for April of 2005–2008. Sample locations are shown in Fig. 4.2. As the BrO VCD has large dispersion, we need a relatively large number of collocated pixels as well as a significant amount of $P_{\text{scene}}$ variability to reliably determine the relationship between BrO VCD and effective cloud scene pressure.

Fig. 4.1a shows a 2-dimensional (2D) histogram of BrO VCD versus for a grid box in the tropics. The BrO VCD is positively correlated with $P_{\text{scene}}$, although it shows a fairly large dispersion. $P_{\text{scene}}$ also shows relatively large variability due to tropical deep convection. The linear regression fit is well-centered about the distribution. While the correlation coefficient is small due to large variations in the BrO measured column, the data clearly show a relationship between the BrO VCD and $P_{\text{scene}}$ that is consistent with BrO located in the free troposphere.
Figure 4.1: Examples of 2-dimensional histograms of total column BrO versus effective cloud scene pressure ($P_{\text{scene}}$) for April 2005-2008 for various representative regions indicated in Fig. 4.2. Number of observations (N), correlation coefficient (R), p-value of the regression, estimated BrO volume mixing ratio (VMR), and 99% confidence interval are indicated in each panel. The BrO volume mixing ratio is derived from the slope of the fitting line of total column BrO versus $P_{\text{scene}}$. 
Fig. 4.1b shows similar data for a grid box located in a desert region (± 20–30 N/S), where deep-convective clouds rarely form. The data in this grid box gives a negative value of BrO VMR, and its confidence interval includes 0. The number of pixels in this ensemble is very small owing to the fact that effective cloud fractions are generally small in this region. Variability of P_{scene} is small as well. The correlation between BrO VCD and P_{scene} is not significant in this ensemble. The width of the confidence interval (proportional to the standard error) of BrO VMR is very high (see Fig. 4.2). The large p-value (0.167) also implies that the derived BrO VMR is not reliable.

Fig 4.1c shows a 2D histogram for a grid box in a mid-latitude oceanic region. Similar to 4.1a, this region also shows large cloud variability although the clouds form at somewhat lower altitudes/high pressure levels. Variability in cloud pressures in this region is produced by frontal storms. Here, the distribution is skewed about the fit. The distribution could represent a sum of two components: A low slope part obtained from P_{scene} = \sim 400 - \sim 750 hPa, and a higher slope for P_{scene} > 750 hPa. This can be explained by free tropospheric BrO that is not well-mixed. Vertical inhomogeneity of BrO VMR may occur in this region owing to less frequent convection in mid-latitudes as compared with the tropics.

Fig 4.1d shows a 2D histogram over high-latitude land. Here again we have a relatively low number of pixels as well as low cloud variability as compared with some of the oceanic regions, because high, optically thick cloud do not form very often over land. BrO VCD is well centered about the linear regression fit at most P_{scene} levels. The low p-value and relatively high correlation coefficient implies that the regression analysis here is reliable. This supports presence of free tropospheric BrO at high latitudes in spring, agreeing with previous studies by Salawitch et al. [2010], Neuman et al. [2010], Choi et al. [2012].

Figure 4.1e shows an example in a high-latitude ocean region. Very high numbers
of pixels (>30000) and low $P_{\text{scene}}$ indicates that this regions contains sea-ice. Even though Figs. 4.1 d and e are at similar latitude regions, they show significant difference owing to variations in characteristics of surface. Over snow/ice, the OMI rotational Raman algorithm no longer retrieves cloud fraction, so we cannot screen data for cloud fraction > 0.7. As a result, the ensemble here includes all measurements regardless of cloud fraction. Observed BrO VCDs here are mixture of boundary layer column and free tropospheric & stratospheric column instead of VCD solely from $P_{\text{scene}}$ to TOA, (and it is impossible to separate them) which is indicated by high median $P_{\text{scene}}$. This ensemble also consist of two parts; one is low slope part between 600 and 800 hPa in free troposphere, and the other is high slope part over 800 hPa due to surface activation of BrO. Even though the collection is affected by near-surface bromine activation, we can still derive BrO VMR between ground and $\sim$600 hPa as the number of points effected by bromine activation is relatively low. The correlation coefficient of the case of Fig. 4.1e is $\sim$0.3; it is not very high, but still reasonable range regarding this analysis. However, in case of Fig. 4.1f, we cannot derive free tropospheric BrO VMR as most clouds are below altitude level of 800 hPa. And the correlation coefficient in Fig. 4.1f is $\sim$0.08. Therefore, we filter collections with correlation coefficient less than 0.15.
Figure 4.2: Top panel: Number of pixels used in $5 \times 5$ degree$^2$ grid boxes (contoured); Bottom panel: (bottom panel) median effective cloud scene pressure ($P_{\text{scene}}$) in each grid box for April 2005-2008.
Figure 4.2 shows variables used to diagnose reliability of the estimated free tropospheric BrO VMR obtained during April 2005-2008. An important variable is the number of the collocated pixels used in a grid box. As the retrieved BrO VCD has a relatively large dispersion, we need a large number of cloudy pixels (cloud fraction > 0.7) to reliably estimate the free tropospheric BrO VMR. For grid boxes with < 1000 pixels, the uncertainties in the estimated BrO VMR are large as shown in Fig. 4.1b. The first panel of Fig. 4.2 shows the number of pixels used for each grid box. Near the equator, there is a narrow band of grid boxes with large number (>∼10000) of cloudy pixels corresponding to inter-tropical convergence zone (ITCZ). Also, regions with low numbers of cloudy pixels appear at latitudes of ∼20° both in both the northern and southern hemispheres, predominantly over northern Africa, off the west coasts of Africa, and South America. In the Arctic, we see an expansive region with large number of pixels; this is indicative of a snow/ice-covered surface as we explained regarding Figs. 4.1e and f. Due to low solar elevation angles, we do not have good observations in the Antarctica during April.

We show the median effective cloud scene pressure (P$_{\text{scene}}$) in the bottom panel of Fig. 4.2. The median P$_{\text{scene}}$ indicates the predominant pressure level of the estimated BrO VMR. The median P$_{\text{cld}}$ is greater than 600 hPa in most low latitude regions (latitude < 30 deg) both in the northern and southern hemisphere (NH and SH). In the tropics, the estimated BrO VMR is mostly sensitive to the altitudes of deep convection. In mid-latitude regions, the median P$_{\text{scene}}$ is ∼750 hPa in northern hemisphere, while it is about 700 hPa in southern hemisphere. In mid-latitudes, the BrO VMR is mostly sensitive to the altitudes of frontal storm tracks. The median P$_{\text{scene}}$ is over 900 hPa in most regions of Arctic, which indicates such regions are covered by snow and/or ice similar to large numbers of pixels with effective cloud fraction > 0.7.
Figure 4.3: Top panel: Width of 99% confidence interval of estimated BrO VMR (see Fig. 4.4) for April 2005-2008; Bottom panel: similar to the top panel but for October 2004-2007.
Figure 4.3 shows the width of the 99% confidence interval (CI) of the BrO VMR (as a fractional value) for April 2005-2008 and October 2004-2007. **The width of the CI is proportional to the standard error. Thus, large CI means large error in the estimated BrO VMR.** Regions of large CI overlap where the number of cloudy pixels is low both in April (see the top panel of Fig. 4.2) and October (not shown). This shows this analysis requires large number of cloudy pixels due to large dispersion of BrO VCD. We also see high CI in polar regions because of low cloud variability and near-surface bromine activation. Overall, the CI of BrO VMR does not exceed 20% except in those regions.

Figure 4.4 shows maps of the estimated BrO VMR for April and October. We do not show BrO VMR where CI is greater than 25%, or where the correlation coefficient between BrO VCD and P\textsubscript{scene} is less than 0.15 (discussed regarding Fig. 4.1). In both months, high BrO VMR (>2.5 pptv) at the polar regions in spring (Arctic in April and Antarctic in October) is apparent. This may result from bromine explosions in polar boundary layer during springtime. In the NH, the high BrO VMR region in the Arctic extends to $\sim 40^\circ$ N, covering northern parts of Eurasia, North Pacific and North America. On the other hand, in the SH, enhanced BrO VMR due to polar bromine explosions is mostly limited to near the coastline of the Antarctica. However, we also observe enhanced BrO in the southern part of South Atlantic and south of Africa. High BrO VMR in North America and Antarctic coastline maintains in fall (NH in October and SH in April).
Figure 4.4: Top panel: Maps of free tropospheric BrO VMR for April 2005-2008; Bottom panel: similar to the top panel but for October 2004-2007. Values are only shown where the width of 99% confidence interval is less than 25% of the derived BrO VMR.
Also apparent are bands of high BrO (\(~2\) pptv) centered at 40° latitude in both hemispheres. This pattern is year-round although the magnitude varies. High BrO near 30°N, 120°W is also appears regardless of the season. The sources of the inferred free tropospheric BrO in these regions are uncertain. Stratospheric intrusion is a possible source, as is the case for tropospheric ozone [Schoeberl et al., 2007].

Figure 4.5 shows the zonal mean and standard deviation of BrO VMR calculated for 5 degree binned zonal bands. The standard deviation is shown for every 10 degrees. The standard error of the mean (not shown) is less than 0.1 pptv in most regions except in polar regions, and is less than 0.2 pptv in polar regions. In the tropics, BrO VMR is almost constant at \(~1\) pptv, but BrO VMR is slightly higher in spring in both hemispheres. This trend is appears in mid-latitudes as well. During springtime in the NH, free tropospheric BrO is enhanced in wide regions at latitudes between 45° and 80°. On the other hand, in the SH, a sharp peak of enhanced BrO in springtime (October) is centered at \(~65°\) S, possibly connected with BrO activation in the coastline of the Antarctica. As described in previous paragraph, small peaks of BrO VMR in fall are apparent near \(~40°\) latitude in fall of both hemispheres. Relatively high BrO VMR are maintained in autumn at the regions of polar BrO activation.
Figure 4.5: Zonal mean and standard deviation of BrO VMR as a function of latitude for April 2005-2008 (blue) and for October 2004-2007 (red). The standard error of the mean (not shown) is less than 0.1 pptv in most regions except polar regions, and is less than 0.2 pptv in polar regions.
CHAPTER V

CONCLUSIONS

5.1 Summary of research

We estimated Arctic tropospheric column BrO for April 2008 from OMI and GOME-2 retrieved total column BrO retrievals and a model simulation of the stratospheric column BrO. The sensitivity of satellite radiances to tropospheric BrO depends on various parameters, such as surface albedo, solar zenith angle (SZA), viewing zenith angle (VZA), the shape of the BrO profile, and the presence of clouds. We have quantified these dependences with a radiative transfer model. A detailed comparison is provided between tropospheric column BrO found from OMI and GOME-2 measurements of total column BrO and tropospheric column BrO computed from in-situ observations of BrO and related species acquired by instruments aboard the DC-8 and WP-3D aircraft during the ARCTAS and ARCPAC field campaigns.

Our analysis shows that retrievals of total column BrO from OMI and GOME-2, combined with a model estimate of stratospheric column BrO, can be used to retrieve realistic estimates of the magnitude and spatial variations in tropospheric column BrO provided that proper observing conditions are met. These conditions include bright surfaces (albedos $\geq 0.7$), low SZAs ($\leq 80^\circ$), and the absence of optically thick clouds. For the flight of 17 April 2008, satellite-derived tropospheric column BrO shows enhancements where the DC-8 reports intense bromine activation and moderate ozone depletion. The satellite and aircraft based measurements of tropospheric column BrO are in close quantitative agreement on 17 April 2008, particularly for DC-8 flight segments over bright surfaces in clear conditions. In contrast to prior expectation based on DOAS measurements, the tropospheric burden of BrO based
on the aircraft profiles was distributed over a broad range of altitudes and was not restricted to the planetary boundary layer. The presence of elevated BrO above the PBL might be driven by convection over warm exposed water in regions of ice leads, with BrO then dispersed by prevailing winds [Simpson et al., 2007b, Salawitch et al., 2010].

Many prior studies have used a zonally symmetric representation of stratospheric BrO to derive tropospheric BrO from satellite observations. As put forth by Theys et al. [2009], Salawitch et al. [2010], and Theys et al. [2011] and reinforced here, this is a deeply flawed assumption. For the DC-8 flight of 5 April 2008, regions of enhanced total column BrO observed by OMI and GOME-2 near Hudson Bay are attributed to compression of stratospheric air associated with a low altitude tropopause, as neither the aircraft nor the satellite based measurements of tropospheric column BrO show enhancements. The regions of enhanced tropospheric column BrO inferred from OMI and GOME-2 on 5 and 17 April 2008 are not readily apparent in maps of total column BrO because stratospheric variability imparts variations in column BrO comparable in magnitude to the tropospheric signal. Use of a zonally symmetric representation of stratospheric column BrO prevents the proper identification of the tropospheric BrO burden [e.g. Theys et al., 2009, Salawitch et al., 2010, Theys et al., 2011]. Caution should be applied when interpreting satellite-derived tropospheric BrO for SZA greater than ∼80 degrees because, under these conditions satellite radiances have decreased sensitivity to absorption by tropospheric BrO (Fig. 3.2).

Satellite and aircraft measurements of tropospheric column BrO do not always exhibit good agreement, at times for reasons that seem well understood and at other times for reasons that are unclear but may be related to differences in the timing or spatial coverage of the respective observations. For the severe ozone depletion event observed by the DC-8 near Alert on 8 April 2008, neither OMI total column BrO column nor OMI tropospheric column BrO were elevated. Atmospheric conditions
were favorable for the remote sensing from space of tropospheric BrO near Alert on this date (e.g. clear skies, high surface reflectivity). We believe the lack of a signal for OMI tropospheric column BrO near Alert on 8 April is consistent with our understanding of bromine chemistry: the production of BrO diminishes when ozone is severely depleted [e.g. Hausmann and Platt, 1994, Tuckermann et al., 1997]. Hence, the association of elevated tropospheric column BrO and depleted O$_3$ is expected to be much stronger for a partial ozone depletion event (ODE) than a major ODE. Tropospheric column BrO from OMI does show an enhancement, near Alert, 36 h prior to the major ODE. The aircraft may have been capturing very small scale variability (several nearby profiles showed large differences) compared to space-based observations. Finally, aircraft observations generally did not sample the altitude region where stratosphere to troposphere transport of active bromine associated with air parcels with elevated levels of inorganic bromine could potentially be affecting the satellite measurement of column BrO.

We examined several events of rapid enhancement of tropospheric column BrO observed by OMI. Observations at high latitudes from this polar-orbiting, sun-synchronous satellite provide a unique illustration of the rapid temporal evolution of BrO activation because orbital periods are about 100 min apart. Our analysis indicates that BrO activation events occurring on 16, 17, and 18 April 2008 are related to near-surface parameters such as low pressure systems, strong surface winds, and/or high planetary boundary layer. The satellite measurements reveal horizontal transport of activated bromine away from the source of origin, and the aircraft measurements show disbursement of BrO within the free troposphere. The strong quantitative agreement between OMI and aircraft tropospheric column BrO on 17 April 2008 supports the validity of the rapid time evolution, on synoptic scales, revealed by the OMI tropospheric BrO product. The events on 17, 18, and 19 April suggest bromine activation via high winds over snow [Yang et al., 2008, Jones et al., 2009, Yang et al., 2010, Jones et al.,
2010] as well as long range transport of Br by surface winds [Begoin et al., 2010].

We estimated free tropospheric BrO VMR from total column BrO and effective cloud scene pressure retrieved from OMI. Existence of free tropospheric BrO was already suggested by previous studies that pointed out the relationship between total column BrO and cloud pressure in tropical regions [Richter et al., 2002, Theys et al., 2011]. Our analysis extends this approach to capture spatial and seasonal variations of free tropospheric BrO VMR outside the tropics. We confirmed that the total column BrO and cloud pressure have a positive correlation throughout the globe. The estimated free tropospheric BrO VMR has spatial structure; BrO VMR has a minimum in low latitude regions, and increases at higher latitudes. Seasonally, free tropospheric BrO VMR is high in spring due to bromine activation in polar regions. In addition, we observe a peak of high BrO VMR at 40° latitude in both hemispheres throughout a year. This feature is currently not understood and is being further investigated.

This portion of the study is currently ongoing and awaits results from various modeling groups to aid in interpretation. There are only a few modeling efforts worldwide that include tropospheric bromine chemistry/transport. Tropospheric bromine sources and detailed bromine chemistry in these models is still highly uncertain. The only process captured in the models is polar bromine activation.

The current study presented here represents a first and significant step forward in deriving spatial and temporal variations in free-tropospheric BrO VMRs, though work in this area is ongoing. For example, systematic errors in the retrieved total column BrO are currently under investigation. In particular, the use of different fitting windows can affect the absolute magnitude of the retrieved total column BrO and thus the estimated value of BrO VMR. Lastly, there are possible interferences including that might bias the observed relationship between the total column BrO and the cloud pressure. For example, the oxygen dimer (O₂-O₂) is not currently
included in the OMI fitting algorithm, because its inclusion produced noisier results. We plan to work with the OMI BrO data producers to further investigate the impact of different fitting windows and parameters on derived tropospheric BrO.

5.2 Recommendations for future studies

Although sources of active bromine remain unresolved, several possible sources have been proposed. Bromide (Br\textsuperscript{−}) in sea salt is thought to be transformed to active bromine by a series of acid-catalyzed multiphase reactions [e.g. Fan and Jacob, 1992]. As stated in the introduction, aerosols, first year sea ice, blowing snow triggered by strong winds are thought to provide the condensed phase for the reactions [e.g. Simpson et al., 2007b]. It is believed that saline surfaces are required for the bromine release [Nghiem et al., 2012]. First year (seasonal) sea ice has more saline surface than perennial ice, together with more refrozen leads, polynyas, and frost flowers. Sea spray can be a mechanism that deposits sea salt on snow-covered sea ice. Sea ice deformation in winter and spring can create open water areas (leads, polynyas) where new thin ice and frost flowers form highly saline surfaces [Nghiem et al., 2012]. Aerosol production from blowing snow events has been suggested as a possible source of active bromine [Yang et al., 2008]. Remotely sensed sea ice maps were used to investigate bromine source regions [Simpson et al., 2007a, Nghiem et al., 2012]. Identification of regions with enhanced tropospheric BrO columns, derived from satellite measurements, with regions of first year sea ice in sea ice maps could be a promising avenue to develop a deeper understanding of the sources of polar bromine activation.

The series of reactions to activate sea salt bromide to molecular bromine require acidic conditions. Since seawater is basic, the source of acidity is another issue that needs further investigation. When sea ice forms, rejection of salts from the freezing water creates highly saline brine. As the brine cools with the ice, some forms of salts including calcium carbonate (CaCO\textsubscript{3}·6H\textsubscript{2}O) reach solubility limits and precipitate,
altering the chemical composition of the residual unfrozen brine [Simpson et al., 2007b]. Precipitation of carbonates and other salts could provide the acidic condition required for the reaction in condensed phase [R. J. Sander et al., 2006, Sander and Morin,, 2009, Dieckmann et al., 2010]. Anthropogenic acidic aerosols also have been suggested as source of the acidity [e.g. Fan and Jacob, 1992].

HO\textsubscript{x} chemistry has significant impacts on bromine chemistry, and vice versa. The reaction of two BrO molecules (Reaction 4, X=Br) has been considered the most important reaction to decide the rate of ODEs [e.g. Zeng et al., 2003, 2006]. However, the reaction of HO\textsubscript{2} and BrO (Reaction 4, X=OH) also destroys ozone as the reaction transforms BrO to Br without producing odd oxygen. According to previous model studies [Sander et al., 1997, Evans et al., 2003], the self-reaction of BrO is the dominant mechanism for ozone loss (∼70%) and BrO + HO\textsubscript{2} plays a minor role (∼25%) during ODEs. The reaction of BrO and HO\textsubscript{2}, however, is more important in ODEs when ozone is depleted, because BrO is not produced in low-ozone conditions (described in Sect. 3.2.2 of Chap. 3.). More importantly, the reaction of BrO and HO2 produces HOBr that deposits on saline surfaces and generates Br\textsubscript{2}, providing twice as many Br atoms back to the gas phase. Active bromine also affects the HO\textsubscript{x} budget, as oxidation of hydrocarbons and carbonyls by Br is a significant source of HO\textsubscript{2}. Mutual impacts between bromine and HO\textsubscript{x} budget need to be further investigated.
Figure 5.1: CALIPSO browse image for a BrO explosion event on April 16 2008 over northern coast of Canada (red box) shown in the fourth column of the second row of Fig. 3.14.
Polar dynamics related to bromine activation is important in understanding the observed features of the tropospheric BrO. Aircraft measurements during ARCTAS and ARCPAC reported enhanced BrO plumes at high altitudes (Fig. 2.2). Convection near open leads have been proposed as a possible mechanism to transport active bromine near surface to high altitudes [McElroy et al., 1999]. Recycling of BrO through aerosols would be necessary in vertical transport of active bromine [Begoin et al., 2010]. In addition to aerosols, active bromine might interact with cloud particles generated by the convection from leads. Figure 5.1 shows CALIPSO browse image for a BrO explosion event over northern coast of Canada (see the fourth column in the second row of Fig. 3.14). CALIPSO image indicates the existence of clouds generated by convection up to $\sim 7$ km altitude as well as polluted continental aerosols where the BrO explosion event occurred (red box). Using space-based lidar (e.g. CALIPSO) data in combination with tropospheric column BrO will be a promising tool to study dynamics related to the high BrO events.

Active bromine species can impact atmospheric oxidation potential and thus change the fates of many atmospheric trace gases. Polar bromine explosions might not only affect polar regions, but global environments as well. We hope this study will initiate further studies on the precise sources of bromine, mechanism of bromine activation, and the environmental impacts of active bromine utilizing data from space-based observations.
APPENDIX A

SUPPLEMENTAL FIGURES

For the animation of Fig. 3.16, see the link below:

http://www.prism.gatech.edu/~schoi41/broexplosion.gif
Figure A.1: Top panel: Maps of total vertical column density of BrO measured on 17 April 2008 by OMI (left) and GOME-2 (right). Middle panel: Estimates of stratospheric vertical column density of BrO (Strat BrO) found using the formulation adopted in the main body of our paper (UMCP; see main text) (left) and found using the formulation of [Theys et al., 2011] (right), all for OMI overpass time. Bottom panel: Map of tropopause pressure at time of OMI overpass from MERRA (see main text) (left) and the ratio of Strat BrO from UMCP divided by Strat BrO from Theys formulation (right). The ratio of these two formulations of Strat BrO bears a relation to tropopause pressure, because the UMCP-based estimate has a higher contribution to total column from air in the lowermost stratosphere (LMS) than the Theys-based estimate, due to the greater emphasis of Product Gas Injection of bromine in the UMCP formulation. When air in the LMS is compressed to high density (high tropopause pressure), the difference in the two formulations grows. Variations in solar zenith angle with respect to latitude and height profiles of total inorganic bromine as a function of altitude also factor into the ratio. For the day shown, the mean value of the ratio of these two estimates of Strat BrO is 1.37. For the 8 days for which data appear in the main paper (5, 6, 7, 8, 17, 19, 21, and 22 April), the mean and standard deviation of the value of the ratio is 0.27 ± 0.09.
Figure A.2: Probability Distribution Function (PDF) of surface BrO measured by two ground-based instruments at Barrow, Alaska in spring 2009: a chemical ionization mass spectrometer (CIMS) and a long path-differential optical absorption spectrometer (LP-DOAS). Only data acquired during daytime are represented. This PDF corresponds to the data shown in Figure 5 of Liao et al. [2011a], JGR, doi:10.1029/2010JD014788, 2011. While surface BrO measured by CIMS and LP-DOAS peaked at 41 and 42 pptv, respectively, the PDF shows that 2/3 of the time, BrO at the surface as measured by CIMS was below 8 pptv. This determination is important for the main body of the paper due to the following: for a surface temperature of 245 K, density at the surface would be $\sim 3 \times 10^{19}$ molecules/cm$^3$. The lowest altitude sampled by the CIMS instrument on board the DC-8, for descents in regions BrO-enhanced regions, was 75 meter above the surface. In the main body of the paper, we use a composite DC-8 profile to estimate the in-situ tropospheric column. Suppose the surface BrO measurements of Liao et al. [2011a] were present, uniformly, between the surface and 75 meters. Then the resulting column, for 8 pptv of BrO, would be $0.18 \times 10^{13}$ molecules/cm$^2$ ($8 \times 10^{-12} \times 3 \times 10^{19}$ molecules/cm$^3 \times 7500$ cm), which is a small contribution to the total in-situ column as well as the total vertical column of BrO. The PDF shown in Figure A.2 supports the view that most of the time, the perturbation to our analysis due to an enhanced layer of BrO present below the aircraft would be small and inconsequential. If 40 pptv of BrO had been uniformly distributed between the surface and 75 meter altitude would contribute $\sim 1 \times 10^{13}$ molecules/cm$^2$ to these columns, which would be consequential. The PDF shows 40 pptv of BrO, while seen, was rarely present. As a result, we state in the main paper “while layers of highly elevated BrO below the aircraft could on occasion compromise our comparisons, our overall conclusions are robust because surface measurements indicate only on rare occasion are BrO enhancements large enough to significantly perturb the column.”


Pfeilsticker, K., Sturges, W. T., Bosch, H., Camy-Peyret, C., Chipperfield, M. P., Engel, A., Fitzenberger, R., Muller, M., Payan, S., and Sinnhuber, B. M.: Lower


Investigation of Tropospheric BrO Using Space-based Total Column BrO Measurements

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107 Pages

Directed by Professor Judith Curry

We derive tropospheric column BrO during the ARCTAS and ARCPAC field campaigns in spring 2008 using retrievals of total column BrO from the satellite UV nadir sensors OMI and GOME-2 using a radiative transfer model and stratospheric column BrO from a photochemical simulation. We conduct a comprehensive comparison of satellite-derived tropospheric BrO column to aircraft in-situ observations of BrO and related species. The aircraft profiles reveal that tropospheric BrO, when present during April 2008, was distributed over a broad range of altitudes rather than being confined to the planetary boundary layer (PBL). Perturbations to the total column resulting from tropospheric BrO are the same magnitude as perturbations due to longitudinal variations in the stratospheric component, so proper accounting of the stratospheric signal is essential for accurate determination of satellite-derived tropospheric BrO. We find reasonably good agreement between satellite-derived tropospheric BrO and columns found using aircraft in-situ BrO profiles, particularly when satellite radiances were obtained over bright surfaces (albedo >0.7), for solar zenith angle <80° and clear sky conditions. The rapid activation of BrO due to surface processes (the bromine explosion) is apparent in both the OMI and GOME-2 based tropospheric columns. The wide orbital swath of OMI allows examination of the evolution of tropospheric BrO on about hourly time intervals near the pole. Low surface pressure, strong wind, and high PBL height are associated with an observed BrO activation event, supporting the notion of bromine activation by high winds over snow. We also provide monthly climatological maps of free tropospheric BrO volume mixing ratio (VMR) derived using the so-called cloud slicing technique. In this
approach, the derived slope of the total column BrO versus cloud pressure is pro-
tional to free tropospheric BrO VMR. Estimated BrO VMR shows a minimum in the
tropics and greater values at higher latitudes in both hemispheres. High tropospheric
BrO VMR at high latitudes in spring could be influenced by near-surface bromine
activation.