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Recommended Citation

Salawitch, R. J., et al. (2010), A new interpretation of total column BrO during Arctic spring, Geophys. Res. Lett., 37, L21805, doi:10.1029/2010GL043798.

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A new interpretation of total column BrO during Arctic spring

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Received 30 April 2010; revised 24 August 2010; accepted 27 August 2010; published 3 November 2010.

[1] Emission of bromine from sea-salt aerosol, frost flowers, ice leads, and snow results in the nearly complete removal of surface ozone during Arctic spring. Regions of enhanced total column BrO observed by satellites have traditionally been associated with these emissions. However, airborne measurements of BrO and O₃ within the convective boundary layer (CBL) during the ARCTAS and ARCPAC field campaigns at times bear little relation to enhanced column BrO. We show that the locations of numerous satellite BrO "hotspots" during Arctic spring are consistent with observations of total column ozone and tropopause height, suggesting a stratospheric origin to these regions of elevated BrO. Tropospheric enhancements of BrO large enough to affect the column abundance are also observed, with important contributions originating from above the CBL. Closure of the budget for total column BrO, albeit with significant uncertainty, is achieved by summing observed tropospheric partial columns with calculated stratospheric partial columns provided that natural, short-lived biogenic bromocarbons supply between 5 and 10 ppt of bromine to the Arctic lowermost stratosphere. Proper understanding of bromine and its effects on atmospheric composition requires accurate treatment of geographic variations in column BrO originating from both the stratosphere and troposphere. Citation: Salawitch, R. J., et al. (2010), A new interpretation of total column BrO during Arctic spring, Geophys. Res. Lett., 37, L21805, doi:10.1029/2010GL043798.

Technology, Atlanta, Georgia, USA.

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1. Introduction

- [2] Regions of elevated BrO at high northerly latitudes during spring associated with the autocatalytic release of bromine from sea-salt aerosol, frost flowers, ice leads, and snow, commonly called the "bromine explosion", cause complete removal of surface ozone [e.g., Barrie et al., 1988; Platt and Hönninger, 2003]. Satellite observations of enhanced column BrO during spring, which we term "BrO hotspots" (regions where total column BrO is elevated by 2 to 3×10^{13} cm⁻² relative to the zonal mean), have long been associated with the surface release of bromine and ozone depletion events (ODEs) [e.g., Chance, 1998; Richter et al., 1998; Wagner et al., 2001].
- [3] There is also widespread interest in atmospheric BrO due to its role as a catalyst for loss of ozone in the stratosphere [e.g., Salawitch et al., 2005] and upper troposphere [e.g., von Glasow et al., 2004]. Oxidation by reaction with atomic Br could be the dominant sink for elemental mercury, with important consequences for mercury deposition [e.g., Holmes et al., 2006]. Reaction with BrO could be a significant sink for dimethlysulfide in the marine boundary layer, reducing subsequent production of SO₂ and new cloud condensation nuclei [e.g., von Glasow et al., 2004].
- [4] Satellite observations provide the best constraint on the global distribution of BrO. Measurements of the vertical column abundance of BrO (BrOVC) reveal much higher amounts (i.e., factor of 2 or 3 more) than found in standard

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models. Considerable debate has centered on the relative role of contributions from the stratosphere and troposphere to this difference [e.g., *Richter et al.*, 1998; *Salawitch et al.*, 2005].

[5] Traditionally, the tropospheric BrO burden has been obtained by subtracting a background from the measured satellite signal, accounting for different air mass factors (ratio of light path through the atmosphere to a vertical path) of the stratospheric and tropospheric components [e.g., Wagner et al., 2001]. The background is commonly based on a longitudinally invariant stratosphere, although potential errors of this approach have been noted [e.g., Richter et al., 1998]. Significant effort has been devoted to measuring the near surface mixing ratio of BrO during Arctic spring [e.g., Platt and Hönninger, 2003, and references therein]. Prior measurements in the Arctic from aircraft [McElroy et al., 1999] and ground-based [Hönninger et al., 2004] instruments suggest the presence of significant levels of tropospheric BrO above the top of the convective boundary layer (CBL, characterized by constant potential temperature with respect to altitude), further complicating our ability to relate satellite observations of BrO^{VC} to surface ODEs. Quantitative closure of the budget for BrO^{VC} has heretofore not been achieved [e.g., Ridley et al., 2007].

[6] This paper is focused on quantification of contributions from the troposphere and the stratosphere to BrOVC during Arctic spring. Enhancements of BrO large enough to be recorded as a satellite "hotspot" are associated with the compression of stratospheric air to high pressure in regions of a low altitude tropopause. Consequently, the notion that the stratospheric contribution to total column BrO can be approximated by a constant background is flawed. Our observations demonstrate that the tropospheric burden of BrO can also contribute to BrOVC at a magnitude consistent with satellite "hotspots" and that a significant portion of the tropospheric signal originates from above the top of the CBL, a region not typically sampled by ground based instruments. We show that closure of the budget for BrOVC can be achieved by summing observed tropospheric partial column BrO with calculated stratospheric partial column BrO, albeit with significant uncertainty in each term. This represents a significant step forward in our understanding of atmospheric BrO.

2. Observations and Model Description

2.1. OMI BrO

[7] OMI is on the NASA Aura platform in a sunsynchronous orbit with a 1:38 pm equator crossing time (ascending node). BrOVC is retrieved from reflected sunlight observed in nadir. The algorithm is based on non-linear, least-squares fitting of radiances in the 319 to 347.5 nm window [Chance, 1998]. BrOVC is found using wavelength dependent air mass factors, computed with a multiple scattering radiative transfer model. Contributions from the O_2 dimer have been neglected, resulting in lower noise and smaller fitting uncertainties than the operational OMI BrO product. Surface albedo is based on a geographically varying, monthly mean climatology derived from OMI observations [Kleipool et al., 2008]. Typically, the fitting residual leads to a $\pm 22\%$ uncertainty (1σ) for BrOVC.

[8] Retrievals of BrO^{VC} from OMI compare extremely well with estimates from ground-based instruments located in Harestua, Norway (60.2°N, 11°E) and Lauder, New Zealand (45.0°S, 169.7°E). The ground-based and satellite measurements agree within 15%, with no discernable bias. The auto-correlation of errors in the state vector elements of BrO^{VC} and O_3 column, from a simultaneous retrieval, is negligible. We therefore conclude regions of enhanced BrO^{VC} are not an artifact caused by the treatment of O_3 in the retrieval algorithm. Further details of the retrieval and these comparisons are given in the auxiliary material. ¹

2.2. Aircraft BrO and Related Species

[9] A variety of aircraft observations are used. The DC-8 and WP-3D aircraft flown during the NASA ARCTAS (Arctic Research of the Composition of the Troposphere from Aircraft and Satellites) and NOAA ARCPAC (Aerosol, Radiation, and Cloud Processes affecting Arctic Climate) campaigns carried in situ instruments that measured BrO, BrCl, and Br₂ using chemical ionization mass spectrometry (CIMS) [Neuman et al., 2010]. The measurements of BrO are accurate to $\pm 40\% + 1$ ppt with a precision of 3 ppt for a 2 sec integration time. The DC-8 measurements are reported with 30 sec time resolution and the WP-3D measurements are reported with 2 sec resolution. The detection limit was typically ~2 ppt for the WP-3D instrument and between 2 to 5 ppt for the DC-8 instrument. Laboratory [Neuman et al., 2010] and field comparisons [Liao et al., 2010] indicate that BrO is not produced or lost on the inlets of the CIMS instruments. The presence of Br₂ during daylight is thought to result from the conversion of HOBr to Br2 on the instrument inlet [Neuman et al., 2010]. We show time series of BrO + BrCl + 2 \times Br₂, which we term BrO_x. Soluble bromide was measured in a mist chamber and includes numerous condensable species. Detailed descriptions of the various instruments are given by Neuman et al. [2010].

[10] Measurements of organic bromocarbons and CFC-12 were acquired by Whole Air Sampler (WAS) instruments onboard the NASA DC-8 aircraft during ARCTAS and TC⁴ (Tropical Composition, Cloud and Climate Coupling) and the NASA WB-57 aircraft during TC⁴. Details of the WAS instruments are given by *Schauffler et al.* [1999]. The WAS data are discussed primarily in the auxiliary material. In situ O₃ was measured using chemiluminescence on the WP-3D and DC-8 and using a dual beam UV photometer on the WB-57.

2.3. Ground Based BrO

[11] Observations of tropospheric BrO over Barrow, Alaska (71.3°N, 156.8°W) are provided by a Max-DOAS (Multi Axis Differential Optical Absorption Spectroscopy) instrument. The differential slant column density of BrO (BrO^{dSCD}) is found as a function of elevation angle (EA) of the acquired spectra [*Hönninger et al.*, 2004]. A radiative transfer program is used to model the variation of BrO^{dSCD} with EA, for various assumptions regarding the height distribution of BrO. This technique provides a strong constraint on the distribution of BrO within the lowest several km of the troposphere.

¹Auxiliary materials are available in the HTML. doi:10.1029/2010GL043798.

2.4. Stratospheric Model BrO

[12] Calculation of the vertical column abundance of stratospheric BrO (BrO $^{\rm STRAT}$) is central to this paper. The calculation begins with global profiles of CFC-12 found by a GEOS-5 assimilation conducted for ARCTAS, with surface emission and stratospheric destruction of CFC-12 [Liang et al., 2008]. CFC-12 is output on a $0.5^{\circ} \times 0.67^{\circ}$ (lat/lon) grid for 72 pressure levels, from the surface to 0.01 hPa, every 6 hours. Comparison to aircraft observations (see auxiliary material) demonstrates that modeled CFC-12 is accurate to within $\pm 4\%$ in the lower stratosphere. Values of Br_y (total inorganic bromine) are found from CFC-12, using the method of *Wamsley et al.* [1998]. The baseline value for Br_y assumes supply of stratospheric inorganic bromine (Br_y) from CH₃Br, halons, and CH₂Br₂.

[13] We also conduct simulations of BrO assuming supply of an additional 5 and 10 ppt of Br_v from very short lived (VSL) bromocarbons, termed Br_vVSL. Since our baseline value of Br_v includes a contribution from CH₂Br₂ by source gas injection (SGI) into the stratosphere, our definition of Br_v^{VSL} differs from that used by World Meteorological Organization (WMO) [2007]. The relation of our definition of Br_v^{VSL} to WMO [2007] and justification for use of 5 and 10 ppt levels of Br_v^{VŠL} based on the WAS measurements are discussed in the auxiliary material. Briefly, CBr_v (total organic bromine) was observed to reach upwards of 30 ppt in the tropical marine boundary layer and to exceed 25 ppt in the region of convective outflow in the tropical upper troposphere during TC⁴. These observations, together with WAS measurements that show direct injection of VSL species into the Arctic lowermost stratosphere (LMS), support the plausibility that Br_v in the Arctic LMS was 5 to 10 ppt higher than our baseline value of Br_v.

[14] Once Br_v is specified, BrO is found using the BrO/ Br_v ratio from a run of WACCM (Whole Atmosphere Community Climate Model) [Garcia et al., 2007] conducted for the START08 (Stratosphere-Troposphere Analyses of Regional Transport 2008) campaign (April to June 2008). WACCM output, provided every 3 hrs on a 1.9° × 2.5° (lat/lon) grid for 89 pressure levels ranging from 1000 to 4.5 \times 10⁻⁶ hPa, is interpolated to the finer GEOS-5 grid for the time of OMI overpass. A single run of WACCM that considered supply of Br_v from only CH₃Br and halons is used; we adjust Br_v outside of WACCM and rely on WACCM for the BrO/Br_v ratio (which is insensitive to Br_v for the range of variations used in this study). We use this procedure because BrO/Br_v is sensitive to O₃ and NO₂ [e.g., Theys et al., 2009] and the START08 WACCM run provides an estimate of O₃ and NO₂ for conditions specific to spring 2008. WACCM calculations of O₃, NO_x and NO compare extremely well to observations of these species obtained during ARCTAS and START08. A description of WACCM and demonstration of its performance, including excellent evaluation of the BrO/ Br_v ratio, is given by Chipperfield and Kinnison [2010]. BrOSTRAT is found by integrating model profiles of BrO from the tropopause (WMO definition of thermal tropopause) to the top of the model atmosphere, for the time of OMI overpass.

[15] A photochemical steady state model is used to assess the uncertainty in BrO^{STRAT}. This model has been constrained to profiles of O₃, NO_y, Br_y, etc. from WACCM as described by *Chipperfield and Kinnison* [2010]. The rate

constant of each reaction that affects BrO/Br_y is varied using uncertainties of *Sander et al.* [2006]. The total uncertainty in BrO^{STRAT} is found from a root-sum-squares combination of the individual chemical kinetics terms (including J values) and propagation, through the model, of the impact of a 4% error in the specification of CFC-12 in the lowermost stratosphere. The reaction of BrO + NO₂ forming BrNO₃, which is uncertain by a factor of 2 at 220 K, contributes most to the overall uncertainty [e.g., *Hendrick et al.*, 2008]. The 1σ uncertainty in BrO^{STRAT} for the Br^{VSL}_y = 10 ppt simulation is $\pm 30\%$ (globally) and $\pm 45\%$ (region of low altitude tropopause), with larger uncertainties near the low altitude tropopause due to increased importance of BrO + NO₂.

3. Results

3.1. Comparison of Airborne and Satellite BrO

[16] Observations of enhanced BrO_x and soluble bromide as well as depleted ozone obtained by the DC-8 aircraft during ARCTAS often bear little relation to the location of OMI BrO^{VC} "hotspots". Measurements obtained on 5 and 8 April 2008 are shown in Figure 1. Several extended low altitude legs (aircraft below 0.3 km) were targeted for regions of elevated BrO, based on analysis of OMI observations from the prior day. These flight portions are denoted by purple line segments for the flight track and altitude traces. The instruments recorded little perturbation to ambient O₃, BrO_y, and soluble bromide in the regions where OMI BrOVC was highly enhanced. Furthermore, a major ODE co-located with highly enhanced BrO_x and soluble bromide was observed on 8 April 2008 near 83°N, 65°W, far from the region of elevated BrOVC (pink circle). Interpolation of OMI BrOVC along the DC-8 flight track (bottom panel) exhibits no meaningful correlation with any of the DC-8 measurements. This set of observations, representative of the majority of data obtained during ARCTAS and ARCPAC, challenges pre-conceived notions of the relation between ODEs and satellite measurements of enhanced column BrO.

3.2. Importance of the Stratosphere

[17] Figure 2 shows total column O₃, tropopause pressure, BrO^{VC}, and modeled BrO^{STRAT} for 5 to 9 April 2008. Column O₃ is from the standard OMI product, available at http://disc.sci.gsfc.nasa.gov/Aura/data-holdings/OMI/omto3d_v003. shtml. The region of elevated BrO over Hudson Bay is coincident with a low tropopause (~5 km altitude or ~450 hPa pressure) and high total column O₃ (~450 DU). These features progress in a similar counter-clockwise fashion, suggesting the enhancements in BrO^{VC} originate from above the tropopause. Enhanced BrO^{STRAT} is associated with a synoptic weather pattern characterized by a low altitude tropopause and high column O₃. The preponderance of prior observations of elevated column BrO over Hudson Bay during spring may be related to a weather pattern known as the Hudson Bay low that is responsible for the depressed tropopause [Liu and Moore, 2004].

[18] An important feature of the satellite data is the compact, monotonic relation between BrO^{VC} and column O_3 for data acquired with solar zenith angle (SZA) $\leq 80^\circ$ (Figure 3). The monotonic nature of this relation (i.e., BrO^{VC} rises as column O_3 increases) provides strong

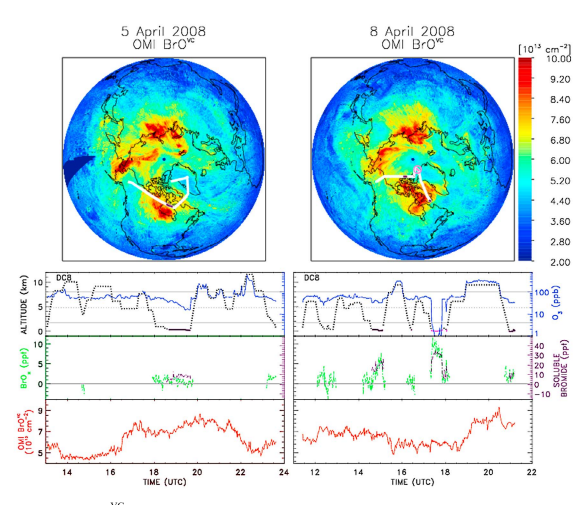


Figure 1. (top) OMI BrO^{VC} on 5 and 8 April 2008; (middle) time series of O_3 (blue), BrO_x (green), and soluble bromide (purple) measured by instruments onboard the NASA DC-8 on these dates; and, (bottom) an interpolation of OMI BrO^{VC} along the DC-8 flight track. The 5 April flight landed in Fairbanks and the 8 April fight originated from Fairbanks. BrO was below the detection limit of ~5 ppt (later flights had lower detections limits) and BrO_x for these dates reflects the presence of other gas phase bromine-bearing species. The DC-8 flight track is shown by white lines on the OMI images and DC-8 altitude is shown by the black dotted line; purple portions denote altitude < 0.3 km and pink indicates altitude < 0.3 km in the presence of a major ODE (O_3 < 4 ppb). Location of a major ODE on 8 April is denoted by the pink circle on the OMI image.

empirical support for the notion that a significant portion of the enhancement in BrO VC originates from above the tropopause. Figure 3 also shows the calculated relation between BrO STRAT and column O_3 ; thin error bars represent 1σ overall uncertainty. Between 5 and 10 ppt of Br $_y$, in excess to that of the baseline, must be present in the Arctic lowermost stratosphere to obtain a theoretical relation with a slope [dBrO VC /d(O_3 column)] similar to that observed. The relation between BrO VC and column O_3 breaks down for SZA $\geq \sim 85^\circ$ (not shown) as BrO goes into its nighttime reservoirs, further supporting our confidence that the relation between enhanced BrO and elevated O_3 is not a retrieval artifact. Departures from linearity of the observed relation, compared to the near linear theoretical expectation, could be due to the influence of the troposphere (i.e., high column O_3 tends to occur at Arctic latitudes, where the tropospheric influence is likely largest) or could represent a shortcoming of the theoretical relations.

shortcoming of the theoretical relations. [19] Modeled BrO^{STRAT} for $Br_y^{VSL} = 10$ ppt resembles OMI BrO^{VC} (Figures 2 and 3). The $Br_y^{VSL} = 5$ ppt simulation of BrO^{STRAT} shows similar features, whereas BrO^{STRAT}

found for $Br_y^{VSL}=0$ (standard for many ozone loss simulations) is quite different than the data. The last row of Figure 2 shows measured BrO^{VC} on 6 April 2008 and modeled BrO^{STRAT} for $Br_y^{VSL}=5$ and 10 ppt, with each quantity perturbed by its 1σ uncertainty. This row reinforces the notion that OMI BrO^{VC} is consistent with Br_y^{VSL} between 5 and 10 ppt. [20] Modeled BrO^{STRAT} assumes no contribution from

below the tropopause. The high BrO^{VC} observed to persist over Hudson Bay on 8 and 9 April 2008 could be due to irreversible mixing of stratospheric air into the UT after passage of the frontal system. Irreversible, cross tropopause exchange of air from the stratosphere to the troposphere (STE) often occurs on the western flank of Arctic low pressure systems [Gettelman and Sobel, 2000]. There is strong similarity between the crescent shaped region of elevated BrO^{VC} seen by many satellite instruments and the areas where STE is thought to occur [Wernli and Sprenger, 2007]. The presence of significant levels of BrO near the surface will also affect the comparison of modeled and measured BrO^{VC}. Quantification of the budget for BrO^{VC}

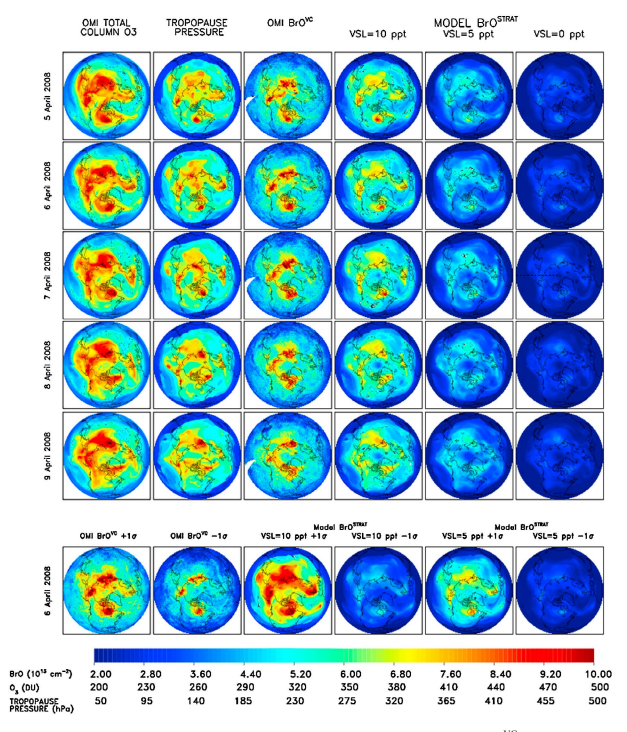


Figure 2. Measured column O_3 (OMI), calculated tropopause pressure (GEOS-5), measured BrO^{VC} (OMI), and calculated BrO^{STRAT} for the $Br_y^{VSL} = 10$, 5, and 0 ppt cases, for 5 to 9 April 2008. The bottom row shows BrO^{VC} and BrO^{STRAT} for the $Br_y^{VSL} = 10$ and 5 ppt cases, for 6 April 2008, with each quantity perturbed by its 1σ uncertainty. Data and model have been filtered for $SZA \le 80^{\circ}$.

requires consideration of contributions from the troposphere as well as the stratosphere.

3.3. Importance of the Troposphere

[21] Airborne observations of BrO show that the tropospheric burden also makes significant contributions to total

column BrO. We have computed vertical column tropospheric BrO (BrO^{TROP}) for 29 profiles of BrO measured during ARCTAS and ARCPAC. Further discussion of measurement uncertainty, which approaches the value of BrO^{TROP}, is given in the auxiliary material. Many profiles occurred near Barrow, Alaska, where elevated surface BrO

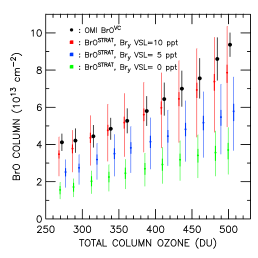


Figure 3. BrO^{VC} versus total column O_3 from OMI for data acquired for 5 to 9 April 2008, the five days shown in Figure 2 (black points). The error bars on BrO^{VC} represent the standard deviation, about the mean, of the values in the respective total column O_3 bins (250 to 525 DU, every 25 DU). The green, blue, and red points represent calculated BrO^{STRAT} for Bry^{SL} of 0, 5, and 10 ppt, respectively. The thick error bar for BrO^{STRAT} represents the uncertainty in model BrO due to errors in GEOS-5 CFC-12; the thin error bar represents the total 1σ uncertainty in BrO^{STRAT}, which is dominated by the factor of 2 uncertainty in $k_{\rm BrO+NO2+M}$. The BrO^{STRAT} points have been displaced slightly, with respect to the mean O_3 of each bin, for clarity of error bars.

and ODEs are often observed. Most profiles were obtained in clear sky. Profiles acquired on 16 April 2008 by the DC-8 are shown in Figure 4. The flight track is superimposed on OMI BrO $^{\rm VC}$ (Figure 4a) and modeled BrO $^{\rm STRAT}$ (Figure 4b). Elevated OMI BrO $^{\rm VC}$ is remarkably well aligned with the underlying sea, suggesting an association with surface release of bromine. The largest value of BrO $^{\rm TROP}$ observed this day, $3.9\times10^{13}~{\rm cm}^{-2}$, is co-located with a local maximum in OMI BrO $^{\rm VC}$.

[22] The most significant contribution to BrO^{TROP} on 16 April 2008 was from altitudes well above the CBL. For the highlighted profile, the CBL had a ceiling of ~0.2 km (Figure 4e). The WP-3D aircraft sampled extensively above and below this altitude on 16 April and other days, at times descending to 60 m above the surface [Neuman et al., 2010]. The importance of BrO above the CBL to BrO^{TROP} is common to all profiles acquired during ARCTAS and ARCPAC (auxiliary material). ARCTAS and ARCPAC aircraft measurements reveal lower abundances of BrO in the CBL than reported in past ground-based studies [e.g., Platt and Hönninger, 2003]. The aircraft flights, by design, often sampled O₃ depleted air within the CBL. Inorganic bromine shifts from BrO into other inorganic species as ambient O₃ falls below ~4 ppb, which was often the case for CBL air sampled during ARCTAS and ARCPAC [Neuman et al., 2010], perhaps accounting for the tendency for our measurements of BrO to be lower than prior observations in the CBL.

[23] Figure 5 shows BrO^{dSCD} vs EA for Max-DOAS data acquired on 20 April 2008. Best agreement between mod-

eled and measured BrOdSCD is found assuming a constant concentration of BrO within the lowest 1 km of the atmosphere, with a column of 2.3×10^{13} cm⁻². An ozonesonde showed a classic CBL extending to 0.3 km, with O₃ present uniformly at ~10 ppb. A model placing all of the BrO within the CBL overestimates observed BrOdSCD at low EA and underestimates BrO^{dSCD} at high EA, suggesting BrO had vented above the CBL. A significant amount of BrO was also present within the CBL, as demonstrated by the inability to properly simulate BrO^{dSCD} by placing all of the BrO between 0.3 and 1 km. Max-DOAS observations conducted during clear sky conditions throughout March and April 2008 (not shown) reveal contributions to BrO^{VC} of 0 to 3 \times 10¹³ cm⁻², with much of the day-to-day variability related to the direction of prevailing surface winds (higher BrOVC observed when air parcels originate from the nearby sea) (D. Donohoue et al., manuscript in preparation, 2010).

[24] The Max-DOAS and aircraft observations of tropospheric BrO are consistent in that they both show significant contributions to BrO^{TROP} from above the CBL. The prevalence of elevated BrO above the CBL may be due to vigorous convection over ice leads driven by warm exposed water, with BrO then dispersed horizontally by prevailing winds. This would be consistent with distribution of BrO throughout the polar boundary layer, a region in which surface emissions can be vertically mixed even if the atmosphere appears to be stable with respect to local convection [e.g., Simpson et al., 2007]. Attempts to relate satellite BrO^{VC} to ODEs and surface BrO, common in the literature, are complicated by the finding that BrO^{TROP} appears to be dominated by contributions from above the CBL.

3.4. Budget of Column BrO

[25] We examine here the budget for BrO^{VC} using the stratospheric model and tropospheric aircraft profiles. Figures 4f and 4g show regression plots of BrO^{MODEL} versus OMI BrO^{VC}, where BrO^{MODEL} is set equal to BrO^{STRAT} (Figure 4f) or BrO^{STRAT}+BrO^{TROP} (Figure 4g). Data are shown for all locations where BrO^{TROP} can be estimated from the two aircraft: 29 profiles encompassing observations acquired on 8 days. Neglecting the tropospheric contribution to column BrO, all modeled BrO columns show slopes below the 1:1 line, although the model for the largest VSL bromine contribution lies close to the line (Figure 4f).

[26] Accounting for contributions from both the troposphere and stratosphere, budget closure is achieved if BrySL lies between 5 and 10 ppt (Figure 4g). Closure of the budget is supported quantitatively by the ratio BrOMODEL/OMI BrOVC encompassing unity, within the standard deviation of the mean, for these two simulations. Strictly speaking, budget closure is achieved for values of BrySL ranging from ~1.4 to 13.2 ppt (auxiliary material). Considerable uncertainty exists, leading to a wide range of BrySL that could be consistent with OMI BrOVC, because BrOVC is uncertain at $\pm 22\%$, BrOSTRAT is uncertain at $\pm 45\%$ due to chemical kinetics, and BrOTROP is uncertain at an amount approaching the measured abundance (all 1σ). The effects of clouds, which potentially shield a portion of the tropospheric column from view of OMI, have not been considered. Nonetheless, Figure 4g demonstrates a plausible means to achieve budget

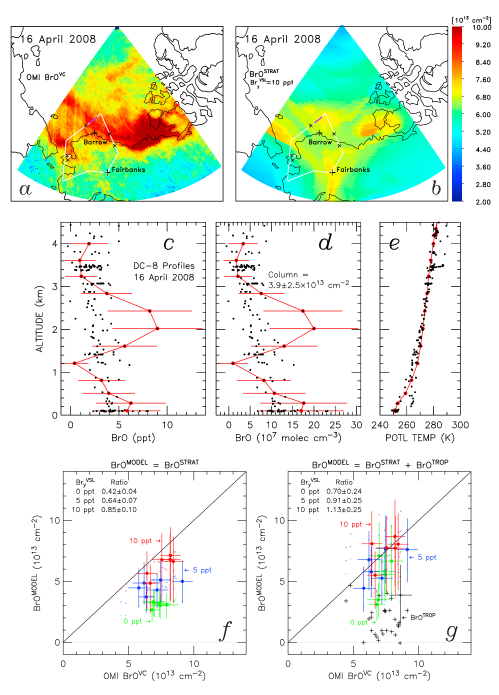


Figure 4. (a and b) Flight track of the DC-8 on 16 April 2008 (white line) superimposed on OMI measurements of BrO^{VC} and calculated BrO^{STRAT} (Br_y^{VSL} = 10 ppt), respectively. (c–e) Profiles of BrO volume mixing ratio, BrO concentration, and potential temperature (θ) on 16 April 2008 (black dots); profile shown in red, with BrO^{TROP} = $3.9 \pm 2.5 \times 10^{13}$ cm⁻², was measured at the segment marked in purple in Figures 4a and 4b; × denotes location of other profiles. All data were acquired for O₃ < 120 ppb, assuring the sampling of tropospheric air. (f and g) Scatter plot of BrO^{MODEL} versus OMI BrO^{VC} for locations of the DC-8 and WP-3D measurements of BrO^{TROP}, where BrO^{MODEL} = BrO^{STRAT} (Figure 4f) and BrO^{MODEL} = BrO^{STRAT} (Figure 4g), for values of BrO^{STRAT} from the Br_y^{VSL} = 0, 5, and 10 ppt simulations. Determinations of BrO^{TROP} (+) are shown in Figure 4g. Error bars for OMI BrO^{VC} represent 1σ uncertainty due to residuals in the spectral fit. Error bars for BrO^{MODEL} represent 1σ uncertainty in BrO^{STRAT} (Figure 4f) and RSS combination of 1σ uncertainty in BrO^{STRAT} and BrO^{TROP} (Figure 4g). For clarity, error bars are shown for every 5th data point and just one measurement of BrO^{TROP}. Values of the ratio BrO^{MODEL}/OMI BrO^{VC} are indicated on Figures 4f and 4g.

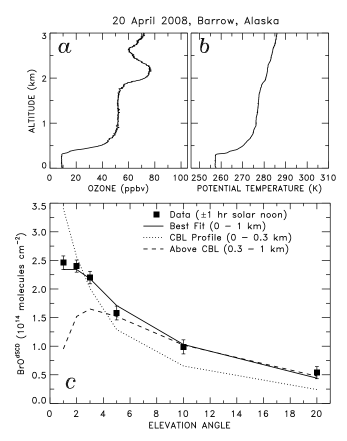


Figure 5. Measurements at Barrow, Alaska on 20 April 2008. (a and b) Profiles of θ and O_3 measured by an ozone-sonde. (c) Differential slant column density of BrO (BrO^{dSCD}) versus Elevation Angle. Black squares are average BrO^{dSCD} within 1 hour of local solar noon. Lines show modeled BrO^{dSCD} for three BrO profiles, all with uniform concentration with respect to altitude between: the surface and 1 km (solid); the surface and 0.3 km (dotted); 0.3 to 1 km (dashed); BrO concentration is otherwise zero. The solid and dashed profiles have a BrO column of 2.3×10^{13} cm⁻²; the dotted profile has a column of 1.3×10^{13} cm⁻².

closure for ${\rm BrO}^{\rm VC}$ that emphasizes the importance of both the troposphere and supply of stratospheric bromine from VSL sources.

4. Concluding Remarks

[27] Our analysis indicates significant contributions to the geographic variation in column BrO originate from both the stratosphere and troposphere, including regions of the troposphere above the convective boundary layer. Numerous OMI BrO "hotspots" exhibit spatial patterns similar to column O₃, tropopause pressure, and modeled stratospheric BrO, provided that VSL bromocarbons make a considerable contribution, 5 to 10 ppt, to stratospheric Br_y. Future attempts to relate satellite measurement of column BrO to surface events should account for the substantial contributions to geographic variability in BrO^{VC} that can originate from the stratosphere. *Theys et al.* [2009] have developed a stratospheric BrO climatology, using model estimates of Br_y and satellite observations of O₃ and NO₂, that could be applicable.

[28] Total column O₃ tends to exhibit highest values during Arctic spring, often in the vicinity of Hudson Bay. Based on our examination of OMI measurements of total column BrO and O₃ over Hudson Bay for the entirety of March and April 2008, it is likely that column BrO in this region is responding, at times, to tropospheric perturbations originating from surface release. Nonetheless, the community has likely been overestimating the extent of elevated tropospheric BrO at Hudson Bay, and perhaps throughout the Arctic, by associating all satellite BrO hotspots with high-latitude surface emission of bromine.

[29] There has been a prior attempt to relate changes in the tropospheric burden of Arctic BrO to global warming [Hollwedel et al., 2004]. Their findings may have to be reconsidered because a longitudinally invariant stratosphere was used for their estimate of tropospheric BrO. Biogenic processes in the tropical oceans [e.g., von Glasow et al., 2004] are the source of the BrO that ultimately drives some of the satellite hotspots seen at high-latitude during spring. The production of bromocarbons in the tropical oceans and transport of these species to the stratosphere responds to a number of forcings, such as oceanic upwelling and atmospheric convection [Salawitch, 2006, and references therein]. It is important to properly quantify the tropospheric and stratospheric contributions to column BrO to accurately assess the impact of bromine on ozone and to progress towards the eventual quantification of any link between climate change and atmospheric bromine.

[30] Acknowledgments. Research of many of the investigators has been supported by the ARCTAS, ACMAP, Aura, MAP, and Tropospheric Chemistry programs of the National Aeronautics and Space Administration, the ARCPAC program of the National Oceanic and Atmospheric Administration, and the START08 program of the National Science Foundation. The ground-based BrO activities at the Belgian Institute for Space Aeronomy (BIRA-IASB) are funded by the PRODEX contract SECPEA and the EC projects GEOmon (FP6-2005-Global-4-036677) and SHIVA (226224-FP7-ENV-2008-1); BIRA-IASB thanks M. P. Chipperfield for providing SLIMCAT output used in the retrieval. We thank the pilots, flight crews, and OMI scientific leadership and data processing teams for their wonderful efforts. We appreciate the three extensive and careful reviews that led to a much improved manuscript.

References

Barrie, L. A., et al. (1988), Ozone destruction and photochemical reactions at polar sunrise in the lower Arctic atmosphere, *Nature*, 334, 138–141, doi:10.1038/334138a0.

Chance, K. (1998), Analysis of BrO measurements from the Global Ozone Monitoring Experiment, *Geophys. Res. Lett.*, 25, 3335–3338, doi:10.1029/98GL52359.

Chipperfield, M., and D. Kinnison (2010), SPARC CCMVal report on the evaluation of chemistry-climate models, *SPARC Rep.* 5, edited by V. Eyring, T. G. Shepherd, and D. W. Waugh, chap. 6, World Meteorol. Organ., Geneva, Switzerland.

Garcia, R. R., D. R. Marsh, D. E. Kinnison, B. A. Boville, and F. Sassi (2007), Simulations of secular trends in the middle atmosphere, 1950–2003, *J. Geophys. Res.*, 112, D09301, doi:10.1029/2006JD007485.

Gettelman, A., and A. H. Sobel (2000), Direct diagnoses of stratosphere-troposphere exchange, *J. Atmos. Sci.*, *57*, 3–16, doi:10.1175/1520-0469(2000)057<0003:DDOSTE>2.0.CO;2.

Hendrick, F., P. V. Johnston, M. De Mazière, C. Fayt, C. Hermans, K. Kreher, N. Theys, A. Thomas, and M. Van Roozendael (2008), One-decade trend analysis of stratospheric BrO over Harestua (60°N) and Lauder (45°S) reveals a decline, *Geophys. Res. Lett.*, 35, L14801, doi:10.1029/2008GL034154.

Hollwedel, J., et al. (2004), Year-to-year variations of spring time polar tropospheric BrO as seen by GOME, Adv. Space Res., 34, 804–808, doi:10.1016/j.asr.2003.08.060.

- Holmes, C. D., D. J. Jacob, and X. Yang (2006), Global lifetime of elemental mercury against oxidation by atomic bromine in the free troposphere, Geophys. Res. Lett., 33, L20808, doi:10.1029/2006GL027176.
- Hönninger, G., et al. (2004), Ground-based measurements of halogen oxides at Hudson Bay by active longpath DOAS and passive MAX-DOAS, *Geophys. Res. Lett.*, 31, L04111, doi:10.1029/2003GL018982.
- Kleipool, Q. L., M. R. Dobber, J. F. de Haan, and P. F. Levelt (2008), Earth surface reflectance climatology from 3 years of OMI data, *J. Geophys. Res.*, 113, D18308, doi:10.1029/2008JD010290.
- Liang, Q., R. S. Stolarski, A. R. Douglass, P. A. Newman, and J. E. Nielsen (2008), Evaluation of emissions and transport of CFCs using surface observations and their seasonal cycles and the GEOS CCM simulation with emissions-based forcing, *J. Geophys. Res.*, 113, D14302, doi:10.1029/ 2007JD009617.
- Liao, J., et al. (2010), A comparison of Arctic BrO measurements by chemical ionization mass spectrometery (CIMS) and long path-differential optical absorption spectroscopy (LP-DOAS), *J. Geophys. Res.*, doi:10.1029/2010JD014788, in press.
- Liu, A. Q., and G. W. K. Moore (2004), Lake-effect snowstorms over southern Ontario and their synoptic-scale environment, *Mon. Weather Rev.*, 132, 2595–2609, doi:10.1175/MWR2796.1.
- McElroy, C. T., C. A. McLinden, and J. C. McConnell (1999), Evidence for bromine monoxide in the free troposphere during the Arctic polar sunrise, *Nature*, 397, 338–341, doi:10.1038/16904.
- Neuman, J. A., et al. (2010), Bromine measurements in O₃ depleted air over the Arctic Ocean, Atmos. Chem. Phys., 10, 6503–6514, doi:10.5194/acp-10-6503-2010.
- Platt, U., and G. Hönninger (2003), The role of halogen species in the troposphere, *Chemosphere*, *52*, 325–338, doi:10.1016/S0045-6535(03) 00216-9
- Richter, A., F. Wittrock, M. Eisinger, and J. P. Burrows (1998), GOME observations of tropospheric BrO in Northern Hemispheric spring and summer 1997, *Geophys. Res. Lett.*, 25, 2683–2686, doi:10.1029/98GL52016.
- Ridley, B. A., et al. (2007), An ozone depletion event in the sub-Arctic surface layer over Hudson Bay, Canada, J. Atmos. Chem., 57, 255–280, doi:10.1007/s10874-007-9072-z.
- Salawitch, R. J. (2006), Biogenic bromine, *Nature*, 439, 275–277, doi:10.1038/439275a.
- Salawitch, R. J., D. K. Weisenstein, L. J. Kovalenko, C. E. Sioris, P. O. Wennberg, K. Chance, M. K. W. Ko, and C. A. McLinden (2005), Sensitivity of ozone to bromine in the lower stratosphere, *Geophys. Res. Lett.*, 32, L05811, doi:10.1029/2004GL021504.
- Sander, S. P., et al. (2006), Chemical kinetics and photochemical data for use in atmospheric studies, *JPL Publ.*, 06-02.
- Schauffler, S. M., E. L. Atlas, D. R. Blake, F. Flocke, R. A. Lueb, J. M. Lee-Taylor, V. Stroud, and W. Travnicek (1999), Distributions of brominated organic compounds in the troposphere and lower stratosphere, *J. Geophys. Res.*, 104, 21,513–21,535, doi:10.1029/1999JD900197.
- Simpson, W. R., et al. (2007), Halogens and their role in polar boundary-layer ozone depletion, Atmos. Chem. Phys., 7, 4375–4418, doi:10.5194/acp-7-4375-2007.
- Theys, N., et al. (2009), A global stratospheric BrO climatology based on the BASCOE chemical transport model, *Atmos. Chem. Phys.*, 9, 831–848, doi:10.5194/acp-9-831-2009.
- von Glasow, R., et al. (2004), Impact of reactive bromine chemistry in the troposphere, *Atmos. Chem. Phys.*, 4, 2481–2497, doi:10.5194/acp-4-2481-2004.

- Wagner, T., C. Leue, M. Wenig, K. Pfeilsticker, and U. Platt (2001), Spatial and temporal distribution of enhanced boundary layer BrO concentrations measured by GOME aboard ERS-2, *J. Geophys. Res.*, 106, 24.225–24.235, doi:10.1029/2000JD000201.
- Wamsley, P. R., et al. (1998), Distribution of halon-1211 in the upper troposphere and lower stratosphere and the 1994 bromine budget, *J. Geophys. Res.*, 103, 1513–1526, doi:10.1029/97JD02466.
- Wernli, H., and M. Sprenger (2007), Identification and ERA-15 climatology of potential vorticity streamers and cutoffs near the extratropical tropopause, J. Atmos. Sci., 64, 1569–1586, doi:10.1175/JAS3912.1.
- World Meteorological Organization (WMO) (2007), Global ozone research and monitoring project, in *Scientific Assessment of Ozone Depletion*: 2006, Rep. 50, World Meteorol. Organ., Geneva, Switzerland.
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