

OMI total bromine monoxide (OMBRO) data product: algorithm, retrieval and measurement comparisons

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Abstract. This paper presents the retrieval algorithm for the operational Ozone Monitoring Instrument (OMI) total bromine monoxide (BrO) data product (OMBRO) developed at the Smithsonian Astrophysical Observatory (SAO) and shows comparisons with correlative measurements and retrieval results. The algorithm is based on direct nonlinear least squares fitting of radiances from the spectral range 319.0-347.5 nm. Radiances are modeled from the solar irradiance, attenuated by contributions from BrO and interfering gases, and including rotational Raman scattering, additive and multiplicative closure polynomials, correction for Nyquist undersampling and the average fitting residual spectrum. The retrieval uses albedo- and wavelength-dependent air mass factors (AMFs), which have been pre-computed using a single mostly stratospheric BrO profile. The BrO cross sections are multiplied by the wavelength-dependent AMFs before fitting so that the vertical column densities (VCDs) are retrieved directly. The fitting uncertainties of BrO VCDs typically vary between 4 and 7×10^{12} molecules cm⁻² (~ 10 %-20 % of the measured BrO VCDs). Additional fitting uncertainties can be caused by the interferences from O₂-O₂ and H₂CO and their correlation with BrO. AMF uncertainties are estimated to be around 10 % when the single stratosphericonly BrO profile is used. However, under conditions of high tropospheric concentrations, AMF errors due to this assumption of profile can be as high as 50 %.

The retrievals agree well with GOME-2 observations at simultaneous nadir overpasses and with ground-based zenith-sky measurements at Harestua, Norway, with mean biases less than $-0.22 \pm 1.13 \times 10^{13}$ and $0.12 \pm 0.76 \times$

 10^{13} molecules cm⁻², respectively. Global distribution and seasonal variation of OMI BrO are generally consistent with previous satellite observations. Finally, we confirm the capacity of OMBRO retrievals to observe enhancements of BrO over the US Great Salt Lake despite the current retrieval setup considering a stratospheric profile in the AMF calculations. OMBRO retrievals also show significant BrO enhancements from the eruption of the Eyjafjallajökull volcano, although the BrO retrievals are affected under high SO₂ loading conditions by the sub-optimum choice of SO₂ cross sections.

1 Introduction

Bromine monoxide (BrO) is a halogen oxide, predominantly located in the stratosphere and upper troposphere, where, like chlorine monoxide (ClO), it is a catalytic element in the destruction of stratospheric ozone (von Glasow et al., 2004; Salawitch et al., 2005) but with higher efficiency per molecule. Sources of tropospheric BrO include bromine release ("explosions") during the polar spring (Hausmann and Platt, 1994; Hollwedel et al., 2004; Simpson et al., 2007; Begoin et al., 2010; Salawitch et al., 2010; Abbatt et al., 2012; Blechschmidt et al., 2016), volcanic eruptions (Bobrowski et al., 2003; Chance, 2006; Theys et al., 2009), salt lakes (Hebestreit et al., 1999; Hörmann et al., 2016) and stratospheric transport (Salawitch et al., 2010). Global BrO measurements from space were first proposed for the Scanning Imaging Absorption Spectrometer for Atmospheric Cartography (SCIA- MACHY) instrument (Chance et al., 1991) and were first demonstrated with Global Ozone Monitoring Experiment (GOME-1) measurements (Chance, 1998; Platt and Wagner, 1998; Richter et al., 1998; Hegels et al., 1998), and since with SCIAMACHY nadir (Kühl et al., 2008), Global Ozone Monitoring Experiment 2 (GOME-2) measurements (Theys et al., 2011; Toyota et al., 2011) and TROPOMI (Seo et al., 2018). Initial observations of BrO by the Ozone Monitoring Instrument (OMI) were first reported by Kurosu et al. (2004). Polar spring BrO enhancements are known to be associated with boundary layer O₃ depletion (Hausmann and Platt, 1994; von Glasow et al., 2004; Salawitch et al., 2005, 2010; Simpson et al., 2007; Abbatt et al., 2012). OMI measurements of BrO have been used together with chemical and dynamical modeling to investigate stratospheric versus tropospheric enhancements of atmospheric BrO at high northern latitudes (Salawitch et al., 2010). OMI BrO retrieval using the Differential Optical Absorption Spectroscopy (DOAS) method has been used to study the seasonal variations of tropospheric bromine monoxide over the Rann of Kutch salt marsh (Hörmann et al., 2016). The Arctic Research of the Composition of the Troposphere from Aircraft and Satellites (ARCTAS) campaign (Choi et al., 2012) found consistency between BrO column densities calculated from Chemical Ionization Mass Spectrometer (CIMS) measurements with the tropospheric BrO columns derived from OMI using our operational retrieval algorithm. BrO has been observed from the ground in Harestua, Norway (Hendrick et al., 2007); Lauder, New Zealand (Schofield et al., 2004a, b); Antarctica (Schofield et al., 2006); Utqiagvik (formerly Barrow), Alaska (Liao et al., 2011, 2012; Frieß et al., 2011; Sihler et al., 2012; Peterson et al., 2016); Eureka, Canada (Zhao et al., 2015); Summit, Greenland (Stutz et al., 2011); and the Arctic Ocean (Burd et al., 2017).

Enhancement of BrO in the vicinity of salt lakes like the Dead Sea and the Great Salt Lake has been observed from ground-based measurements (Hebestreit et al., 1999; Matveev et al., 2001; Stutz et al., 2002; Tas et al., 2005; Holla et al., 2015). The active bromine compound release is due to the reaction between atmospheric oxidants with salt reservoirs. Satellite observation of salt lake BrO was first reported over the Great Salt Lake and the Dead Sea by OMI (Chance, 2006; Hörmann et al., 2016). Seasonal variations of tropospheric BrO over the Rann of Kutch salt marsh have been observed using OMI from an independent research BrO product (Hörmann et al., 2016). Bobrowski et al. (2003) made the first ground-based observations of BrO and SO₂ abundances in the plume of the Soufrière Hills volcano (Montserrat) using multi-axis DOAS (MAX-DOAS). BrO and SO₂ abundances as functions of the distance from the source were measured by MAX-DOAS in the volcanic plumes of Mt. Etna in Sicily, Italy, and Villarica in Chile (Bobrowski et al., 2007). The BrO/SO₂ ratio in the plume of Nyiragongo and Etna was also studied (Bobrowski et al., 2015). The first volcanic BrO measured from space was from the Ambrym volcano, measured by OMI (Chance, 2006). Theys et al. (2009) reported on GOME-2 detection of volcanic BrO emission after the Kasatochi eruption. Hörmann et al. (2013) examined GOME-2 observations of BrO slant column densities (SCDs) in the vicinity of volcanic plumes; the study showed clear enhancements of BrO in \sim one-quarter of the volcanos and revealed large spatial differences in BrO/SO₂ ratios.

The purpose of this paper is to describe the OMI BrO operational algorithm and the data product, compare it with ground-based and other satellite measurements and analyze its spatiotemporal characteristics. This paper is organized as follows: Sect. 2 describes the OMI instrument and the data product. Section 3 gives a detailed description of the operational algorithm including algorithm and product history, spectral fitting, air mass factor (AMF) calculations, destriping and fitting uncertainties. Section 4 presents results and discussion including comparison with GOME-2 and groundbased zenith-sky measurements at Harestua, Norway, and global distribution, seasonality and enhanced BrO from the US Great Salt Lake and Iceland's Eyjafjallajökull volcano. Section 5 concludes this study.

2 OMI instrument and OMBRO data product

2.1 OMI instrument

OMI (Levelt et al., 2006) was launched on the NASA Earth Observing System (EOS) Aura satellite into a sunsynchronous orbit on 15 July 2004. It is a push-broom imaging spectrometer that observes solar backscattered radiation in the visible and ultraviolet from 270 to 500 nm in three channels (UV1: 270–310 nm, UV2: 310–365 nm, visible: 350–500 nm) at a spectral resolution of 0.42–0.63 nm and spatial resolution in the normal (global sampling) mode ranging from $13 \text{ km} \times 24 \text{ km}$ at direct nadir to about $28 \text{ km} \times 150 \text{ km}$ at the swath edges. The global mode (GM) has 60 ground pixels with a total cross-track swath of 2600 km.

Since June 2007, certain cross-track positions of OMI data have been affected by the row anomaly (http://projects.knmi. nl/omi/research/product/rowanomaly-background.php, last access: 20 November 2017): some loose thermal insulating material likely appeared in front of the instrument's entrance slit, which can block and scatter the light, thus causing errors in level 1b data and subsequently the level 2 retrievals (Kroon et al., 2011). Initially, the row anomaly only affected a few positions, and the effect was small. But since January 2009, the anomaly has become more serious, spreading to \sim onethird of the positions, and retrievals at those positions are not recommended for scientific use. A flagging field has been introduced in the OMI level 1b data to indicate whether an OMI pixel is affected by this instrument anomaly.

OMI measures O_3 and other trace gases, aerosols, clouds and surface properties. Products developed at the SAO include operational BrO, chlorine dioxide (OClO), and formaldehyde (H₂CO; González Abad et al., 2015) that are archived at NASA Goddard Earth Sciences (GES) Data and Information Services Center (DISC) and offline ("preoperational") O₃ profile and tropospheric O₃ (Liu et al., 2010; Huang et al., 2017, 2018), glyoxal (C₂H₂O₂) (Chan Miller et al., 2014, 2016) and water vapor (H₂O) (Wang et al., 2014, 2016) that are available at the Aura Validation Data Center (AVDC). All the products except for the O₃ profile product are produced using nonlinear least squares (NLLS) fitting methods based on those previously developed at the SAO for the analysis of measurements from the GOME (now GOME-1) (Chance, 1998; Chance et al., 2000) and SCIA-MACHY instruments (Burrows and Chance, 1991; Chance et al., 1991; Martin et al., 2006).

2.2 OMBRO data product

The current operational BrO product, OMBRO version 3, contains BrO vertical column densities (VCDs), slant column densities (SCDs), effective air mass factors (AMFs) and ancillary information retrieved from calibrated OMI radiance and irradiance spectra. Each BrO product file contains a single orbit of data, from pole to pole, for the sunlit portion of the orbit. The data product from 26 August 2004 through the present is available at GES DISC. Data used in this study cover the period from 1 January 2005 to 31 December 2014.

3 Retrieval algorithm

3.1 Algorithm and product history

OMBRO Version 1.0 was released on 1 February 2007, based on a spectral fitting window of 338-357 nm. Version 2.0 was released on 13 April 2008. It included major adjustments for Collection 3 level 1b data, improved destriping measures, a change of the fitting window to 340-357.5 nm, improvements to radiance wavelength calibration, and several improvements for processing near-real-time data. In both Version 1 and 2, total BrO VCDs were retrieved in two steps: first performing spectral fitting using the basic optical absorption spectroscopy (BOAS) method to derive SCDs from OMI radiance spectra and then converting from SCDs to VCDs by dividing AMFs. This is similar to current SAO H₂CO, H₂O and C₂H₂O₂, as mentioned previously. The latest Version 3.0.5, released on 28 April 2011, includes major algorithm changes: the fitting window was moved to 319.0-347.5 nm, and BrO cross sections are multiplied by wavelength-dependent AMFs, which are a function of albedo, before fitting, for a direct retrieval of BrO VCDs. SCDs are similarly retrieved in a separate step by fitting BrO cross sections that have not been multiplied with wavelength-dependent AMFs, and an effective AMF =

SCD/VCD is computed. Diagnostic cloud information from the OMCLDO2 product (Acarreta et al., 2004) was added, and the row-anomaly-indicating flags were carried over from the level 1b product. We recommend not using pixels affected by the row anomaly despite being processed by the retrieval algorithm.

The current algorithm is described in detail in the rest of this section, with spectral fitting in Sect. 3.2, AMF calculation prior to spectral fitting in Sect. 3.3, post-processing destripping to remove cross-track-dependent biases in Sect. 3.4 and fitting uncertainties and error estimates in Sect. 3.5.

3.2 Spectral fitting

Most aspects of the algorithm physics for the direct fitting of radiances by the BOAS method were developed previously at SAO for analysis of GOME and SCIAMACHY satellite spectra (Chance, 1998; Chance et al., 2000; OMI, 2002; Martin et al., 2006) and in the various algorithm descriptions of other SAO OMI products (Wang et al., 2014; Chan Miller et al., 2014; González Abad et al., 2015).

The spectral fitting in the SAO OMI BrO retrieval is based on a Gauss–Newton NLLS fitting procedure, the CERN EL-SUNC procedure (Lindström and Wedin, 1987), which provides for bounded NLLS fitting. Processing begins with wavelength calibration for both irradiance and radiance. In each case the wavelength registration for the selected fitting window is determined independently for each cross-track position by cross-correlation of OMI spectra with a high spectral resolution solar irradiance (Caspar and Chance, 1997; Chance, 1998; Chance and Kurucz, 2010) using the preflight instrument slit functions (Dirksen et al., 2006). Radiance wavelength calibration is performed for a representative swath line of radiance measurements (usually in the middle of the orbit) to determine a common wavelength grid for reference spectra.

Following wavelength correction, an undersampling correction spectrum is computed to partially correct for spectral undersampling (lack of Nyquist sampling: Chance, 1998; Slijkhuis et al., 1999; Chance et al., 2005). The calculation of the corrections for the undersampling is accomplished by convolving the preflight slit functions with the highresolution solar spectrum and differencing its fully sampled and undersampled representations (Chance et al., 2005).

To process each OMI orbit, it is split into blocks of 100 swath lines. Spectral fitting is then performed for each block by processing the 60 cross-track pixels included in each swath line sequentially before advancing to the next swath line. The spectra are modeled as follows:

$$I = \left\{ \left(aI_0 + \sum_i \alpha_i A_i \right) e^{-\sum_j (\beta_j B_j)} + \sum_k \gamma_k C_k \right\} \operatorname{Poly}_{\text{scale}} + \operatorname{Poly}_{\text{baseline}},$$
(1)

where I_0 is the solar irradiance (used in our operational BrO retrieval) or radiance reference measurement, I is the Earthshine radiance (detected at satellite), a is albedo, α_i , β_i and γ_k are the coefficients to the reference spectra of A_i , B_j and C_k (for example, trace gas cross sections, Ring effect, vibrational Raman, undersampling correction, common mode) of model constituents. To improve cross-track stripe biases (Sect. 3.4), the OMI daily solar irradiance (I_0) is substituted by the first principal component of the solar irradiances measured by OMI between 2005 and 2007 (one for each crosstrack position). The principal component derived between 2005 and 2007 is used to process the entire mission. The reference spectra are derived separately for each cross-track position from original high-resolution cross sections convolved with the corresponding pre-launch OMI slit functions (Dirksen et al., 2006) after correcting for the solar I_0 effect (Aliwell et al., 2002). Figure 1 shows the trace gas cross sections and Ring spectra used in the current operational algorithm. The black lines are the original high-resolution reference spectra, and the colored lines show the corresponding spectra convolved with the OMI slit function, which are used in the fitting.

For improved numerical stability, radiances and irradiances are divided by their respective averages over the fitting window, renormalizing them to values of ~ 1 . BrO is fitted in the spectral window 319.0–347.5 nm, within the UV-2 channel of the OMI instrument. The switch from the previous fitting window of 340–357.5 nm to this shorter and wider fitting window is based on extensive sensitivity analysis following the method described by Vogel et al. (2013). This new fitting window aims at reducing the fitting uncertainty by including more BrO spectral structures as shown in Fig. 1 and reducing retrieval noise while preserving the stability of the algorithm. An analysis of the retrieval sensitivity to different windows is included in Sect. 3.5.

The rotational Raman scattering (Chance and Spurr, 1997; Chance and Kurucz, 2010) and undersampling correction spectra, A_i , are first added to the albedo-adjusted solar irradiance aI_0 , with coefficients α_i as shown in Eq. (1). Radiances I are then modeled as this quantity attenuated by absorption from BrO, O₃, NO₂, H₂CO, and SO₂, with coefficients β_i fitted to the reference spectra B_i as shown in Eq. (1). A common mode spectrum C_k , computed online, is added by fitting coefficient γ_k after the Beer–Lambert law contribution terms. For each cross-track position, an initial fit of all the pixels along the track between 30° N and 30° S is performed to determine the common mode spectra, derived as the average of the fitting residuals. The common mode spectra include any instrument effects that are uncorrelated to molecular scattering and absorption. This is done to reduce the fitting root-mean-square (rms) residuals and the overall uncertainties. These are then applied as reference spectra in fitting of the entire orbit. The fitting additionally contains additive (Polybaseline) and multiplicative closure polynomials (Poly_{scale}), parameters for spectral shift and, potentially, squeeze (not normally used). The operational parameters and the cross sections used are provided in Table 1.

As part of the development of the OMBRO retrieval algorithm, a significant amount of effort was dedicated to algorithm "tuning", i.e., the optimization of elements in the retrieval process, including interfering absorbers like O₂-O₂. The spectral region of 343 nm, where O₂-O₂ has an absorption feature larger than the BrO absorption, essentially is impossible to avoid in BrO retrievals: the fitting window would have to either terminate at shorter wavelengths or start past this feature, and both approaches yield low information content that is too unacceptable for the BrO retrievals to succeed. During the tuning process, we investigated the effects of, among many other things, including or excluding O₂-O₂, the use of different spectroscopic data sets (Greenblatt et al., 1990; Hermans et al., 1999, cross sections), shorter or longer wavelength windows for the retrieval and even extending the retrieval window beyond the O₂-O₂ absorption feature but excluding the approximate wavelength slice of the feature itself. The only approach that provided quantitatively satisfactory results - i.e., stability of the retrieval under a wide range of conditions, minimized correlation with clouds, low fitting uncertainties, consistency of OMI global total column BrO with published results, and low noise in pixel-to-pixel retrievals - was to exclude O₂-O₂ from the OMBRO V3. It is difficult to quantify O₂-O₂ atmospheric content from the absorption feature around 343 nm alone, and its correlation with absorption bands of BrO and H₂CO leads to spectral correlations in the course of the nonlinear least squares minimization process that are detrimental to the OMI BrO retrievals. Lampel et al. (2018) provide spectrally resolved O₂-O2 cross sections not only at 343 nm but also at 328 nm (see Fig. 1), which is about 20 % of the absorption at 343 nm and has not been shown in previous O₂-O₂ cross sections. Future updates to the operational OMBRO algorithm will investigate the effect of including Lampel et al. (2018) O₂-O₂ cross sections on the fitting.

3.3 Air mass factors

Due to significant variation in O_3 absorption and Rayleigh scattering in the fitting window AMFs vary with wavelength by 10 %–15 % as shown in Fig. 2. At large solar and viewing zenith angles it is difficult to identify a single representative AMF ad hoc. The wavelength-dependent AMFs are introduced to take into account for such strong variation within the BrO fitting window. They are applied prior to fitting to the BrO cross sections, and the spectral fit retrieves VCDs directly. This direct fitting approach is a major departure from the commonly employed two-step fitting procedure (OMI, 2002). It was first developed for retrievals of trace gases from SCIAMACHY radiances in the shortwave infrared (Buchwitz et al., 2000) and has been demonstrated for total O₃ and SO₂ retrievals from GOME/SCIAMACHY measurements in



Figure 1. Cross sections used in the current operational BrO algorithm except for the SO₂ cross section at 298 K which is to be used in the next version. The black lines are the original cross sections, and the colored lines show the cross sections convolved with approximate OMI slit function (which is assumed to be a Gaussian with 0.42 nm full width at half maximum). The O₂-O₂ calculation is based on Lampel et al. (2018) cross sections. The BrO cross section after multiplication with the wavelength-dependent AMFs used these parameters for the AMF calculation: albedo = 0.05, SZA = 5.0° and VZA = 2.5°. The rms of the fitting residuals is on the order of 9×10^{-4} , indicating that BrO spectral features are slightly bigger than typical fitting residuals.

| Parameter | Description/value |
|---------------------------------|--|
| Fitting window | 319.0–347.5 nm |
| Baseline polynomial | Fourth-order |
| Scaling polynomial | Fourth-order |
| Instrument slit function | Hyper-parameterization of preflight measurements; Dirksen et al. (2006) |
| Wavelength calibration | Spectral shift (no squeeze) |
| Solar reference spectrum | Chance and Kurucz (2010) |
| BrO cross sections | Wilmouth et al. (1999), 228 K |
| H_2CO cross sections | Chance and Orphal (2011), 300 K |
| O ₃ cross sections | Malicet et al. (1995), 218, 295 K |
| NO ₂ cross sections | Vandaele et al. (1998), 220 K |
| SO_2 cross sections | Vandaele et al. (1994), 295 K ^a ; Hermans et al. (2009); Vandaele et al. (2009), 295 K ^b |
| OCIO cross sections | Kromminga et al. (2003), 213 K |
| Molecular Ring cross sections | Chance and Spurr (1997) |
| Undersampling correction | Computed online; Chance et al. (2005) |
| Residual (common mode) spectrum | Computed online between 30° N and 30° S |

Table 1. Fitting window and parameters used to derive BrO vertical column densities.

^a Used in the current operational algorithm.

 $^{\rm b}$ Used for testing sensitivity to ${\rm SO}_2$ cross sections and will be used in the next version.



Figure 2. Wavelength- and albedo-dependent air mass factors calculated using a mostly stratospheric fixed BrO profile. The blue box shows the fitting window used in our previous versions, and the red box shows the fitting window used in the current operational algorithm.

the ultraviolet (Bracher et al., 2005; Coldewey-Egbers et al., 2005; Weber et al., 2005; Lee et al., 2008).

The albedo- and wavelength-dependent AMFs were precomputed with the Linearized Discrete Ordinate Radiative Transfer code (LIDORT, Spurr, 2006) using a single mostly stratospheric BrO profile (Fig. 3, left panel). The BrO profile, based on the model of Yung et al. (1980), has $\sim 30 \%$ BrO below 15 km, ~ 10 % BrO below 10 km and ~ 2 % BrO below 5 km. It should be noted that a fixed profile is inconsistent with the varying tropopause height (both with latitude and dynamically, e.g., Salawitch et al., 2010) and therefore with the profile shape in the stratosphere, but the impact on the AMF is typically small as the scattering weight does not change much in the stratosphere. For conditions with enhanced BrO in the lower troposphere, using this profile will overestimate the AMFs and therefore underestimate the BrO VCDs as discussed in Sect. 3.5. Surface albedos are based on a geographically varying monthly mean climatology derived from OMI (Kleipool et al., 2008). Although AMFs based on this BrO profile only slightly depend on surface albedo, albedo effects can be significant over highly reflective snow or ice surfaces, reducing VCDs by 5 %-10 %.

In order to provide the AMF in the data product for consistency with previous versions based on a two-step approach, a second fitting of all OMI spectra is performed with unmodified BrO cross sections, which yields SCDs. An effective AMF can then be computed as AMF = SCD/VCD.

The green line in the top right panel of Fig. 1 shows the modified BrO cross section after multiplication with the wavelength-dependent AMF (albedo = 0.05, SZA (solar zenith angle) = 5.0° , and VZA (viewing zenith angle) = 2.5°). The wavelength dependence in AMF is visible from the varying differences near BrO absorption peaks and the right wings at different wavelengths. The correlation of the unmodified BrO cross sections with the rest of the fit-



Figure 3. (a) A mostly stratospheric vertical BrO profile used for air mass factors calculations in OMBRO V3. Total BrO, BrO < 15 km, BrO < 10 km and BrO < 5 km are 2.05×10^{13} , 5.06×10^{12} , 1.55×10^{12} and 2.87×10^{11} molecules cm⁻², respectively. (b) A stratospheric tropospheric vertical BrO profile used to investigate the impact of high tropospheric BrO columns on air mass factors calculations. Total BrO, BrO < 15 km, BrO < 10 km and BrO < 5 km are 6.99×10^{13} , 5.45×10^{13} , 5.10×10^{13} and 4.97×10^{13} molecules cm⁻², respectively.

ted molecules is small (typically less than 0.12), except with H_2CO (0.43). However, it is safe to assume that in most polar regions with enhanced BrO there are no high concentrations of formaldehyde. It will be worthwhile for future studies to assess the interference of H_2CO under high H_2CO and background BrO conditions, similar to De Smedt et al. (2015). In addition, the AMF wavelength dependence increases with the increase of solar and viewing zenith angles and surface albedo, which increases the correlation between modified BrO cross sections and O₃ cross sections. However, the correlation with O₃ only becomes noticeable (~ 0.10) at SZAs above ~ 80°.

3.4 Destriping

OMI L1b data exhibit small differences with cross-track position, due to differences in the dead/bad pixel masks (crosstrack positions are mapped to physically separate areas on the CCD), dark current correction and radiometric calibration, which lead to cross-track stripes in the level 2 product (Veihelmann and Kleipool, 2006). Our destriping algorithm employs several methods to reduce cross-track striping of the BrO columns. First, we screen outliers in the fitting residuals. This method, originally developed to mitigate the effect of the South Atlantic Anomaly in SAO OMI BrO, H₂CO, and OClO data products, is now also being employed for GOME-2 (Richter et al., 2011). Screening outliers is done through computing the median, r_{med} , and the standard deviation σ of residual spectra $r(\lambda)$ and in subsequent refitting excluding any spectral points for which $r(\lambda) \ge |r_{\text{med}} \pm 3\sigma|$. This can be done repeatedly for every ground pixel, which makes the processing slow. However, we do it once for a reference swath line, recording the positions of the bad pixels and excluding them by default in each subsequent fit. Second, after the completion of the spectral fitting process for all ground pixels in the granule, a post-processing cross-track bias correction is performed: an average cross-track pattern is calculated from the along-track averages of all BrO VCDs for each crosstrack position within a $\pm 30^{\circ}$ latitude band around the Equator, to which a low-order polynomial is fitted. The differences between the cross-track pattern and the fitted polynomial are then applied as a cross-track VCD correction (or "smoothing") factor. The smoothed VCDs are provided in a separate data field, ColumnAmountDestriped. Smoothed SCDs are derived in an analogous fashion and are also included in the data product.

3.5 BrO VCD error analysis

Estimated fitting uncertainties are given as $\sigma_i = \sqrt{C_{ii}}$, where C is the covariance matrix of the standard errors. This definition is only strictly true when the errors are normally distributed. In the case in which the level 1 data product uncertainties are not reliable estimates of the actual uncertainties, spectral data are given unity weight over the fitting window, and the 1σ fitting error in parameter *i* is determined as

$$\sigma_i = \varepsilon_{\rm rms} \sqrt{\frac{c_{ii} \times n \text{ points}}{n \text{ points} - n \text{ varied}}},\tag{2}$$

where $\varepsilon_{\rm rms}$ is the root mean square of the fitting residuals, "*n* points" is the number of points in the fitting window and "*n* varied" is the number of parameters varied during the fitting.

The fitting uncertainties for single measurements of the BrO VCDs typically vary between 4×10^{12} and 7×10^{12} molecules cm⁻², consistently throughout the data record. The uncertainties vary with cross-track positions, from $\sim 7 \times 10^{12}$ at nadir positions to $\sim 4 \times 10^{12}$ at edge positions due to the increase of photon path length through the stratosphere. Relatively, the VCD uncertainties typically range between 10% and 20% of individual BrO VCDs but could be as low as 5% over BrO hotspots. This is roughly 2–3 times worse that what was achieved from GOME-1 data.

The BrO VCD retrieval uncertainties listed in the data product only include random spectral fitting errors. Error sources from AMFs (i.e., BrO climatology), atmospheric composition and state (pressure–temperature vertical profiles, total O₃ column, etc.) and other sources of VCD uncertainty are not included. We provide error estimates for these additional error sources here.

Uncertainties in the AMFs, used to convert slant to vertical columns, are estimated to be 10% or less except when there is substantially enhanced tropospheric BrO. Hence



Figure 4. The percentage of relative AMF errors as a function of the SZA and the wavelength (**a**), albedo (**b**) and VZA (**c**) when using the stratospheric-only BrO profile (Fig. 3a) in the case there is a significant tropospheric BrO column as shown in the stratospheric–tropospheric BrO profile (Fig. 3b).

the total uncertainties of the BrO vertical columns typically range within 15%–30%. To estimate the AMF error associated with enhanced tropospheric concentrations, we have studied the difference between AMFs calculated using the stratospheric-only BrO profile and a stratospherictropospheric profile as shown in Fig. 3b. Figure 4 shows the dependency of the relative AMF difference with respect to wavelength (panel a), albedo (panel b) and VZA (panel c) as a function of the SZA between calculations performed using these two profiles. The use of the stratospheric-only BrO profile can lead to AMF errors up to 50% depending on albedo and viewing geometry. On average, using the stratospheric-only BrO profile overestimates AMF and underestimates VCD by 41%.

We have performed sensitivity analysis of OMI BrO VCD with respect to various retrieval settings using orbit 26564 on 13 July 2009. Table 2 shows the median VCDs, median fitting uncertainties and the number of negative VCD pixels for each configuration. Table 3 summarizes the overall fitting error budget including the random fitting uncertainty, cross section errors (as reported in the literature) and various retrieval settings. We studied five wavelength windows including the current operational window (319.0-347.5 nm) version 2 window (323.0-353.5 nm), version 1 (340.0-357.5 nm) and two extra windows, exploring the impact of extending the window to shorter wavelengths (310.0-357.5 nm) and reducing it by limiting its extension to wavelengths above 325 nm (325.0-357.5). The choice of fitting window can cause significant differences in BrO VCDs of up to 50 %. The current window results in the most stable retrievals with the smallest number of pixels with negative VCD values.

| Description | Median VCD $(molecules cm^{-2})$ | Median uncertainty (molecules cm^{-2}) | Number of negatives |
|-------------------------------------|----------------------------------|---|---------------------|
| 319–347.5 nm Op. (V3) | 4.02×10^{13} | 7.11×10^{12} | 88 |
| 323.0-353.5 nm (V2) | 2.65×10^{13} | 9.27×10^{12} | 1604 |
| 340.0-357.5 nm (V1) | 2.86×10^{13} | 1.19×10^{13} | 3351 |
| 310.0-357.5 nm | 1.97×10^{13} | 6.18×10^{12} | 2728 |
| 325.0–357.5 nm | 3.16×10^{13} | 8.02×10^{12} | 1416 |
| With O ₂ -O ₂ | 3.54×10^{13} | 7.80×10^{12} | 319 |
| Online slit function | 5.09×10^{13} | 7.16×10^{12} | 68 |
| Without common mode | 3.89×10^{13} | 1.02×10^{13} | 116 |
| Without H ₂ CO | 2.52×10^{13} | 6.27×10^{12} | 816 |

Table 2. Error analysis studies. For reference, the total number of retrieved pixels not affected by the row anomaly is 58 112.

 Table 3. Summary of different errors sources in the BrO vertical column.

| Error source | Туре | Parameter uncertainty | Averaged uncertainty on BrO VCD | Evaluation method – reference | |
|--|------------|---|---|--|--|
| Measurement noise random | Random | S/N 500-1000 | $4-7 \times 10^{12}$ molecules cm ⁻² | Error propagation | |
| H ₂ CO O ₃ BrO NO ₂ SO ₂ OCIO Ring | Systematic | Based on literature re- ported error estimates | 5 % 2 % 8 % 3 % 5 % 5 % | Chance and Orphal (2011), 300 K Malicet et al. (1995), 218, 295 K Wilmouth et al. (1999), 228 K Vandaele et al. (1998), 220 K Vandaele et al. (1994), 295 K Kromminga et al. (2003), 213 K Chance and Spurr (1997) | |
| Order of baseline poly- | Systematic | Vary polynomial order | 10 % | | |
| nomial Order of scaling poly- nomial | Systematic | Vary polynomial order | 10 % | Sensitivity analysis | |
| Instrumental slit func- tion and wavelength calibration | Systematic | Preflight and online slit function | 27 % | | |
| Wavelength interval | Systematic | Varying fitting window | 50 % | | |

Including the interference of O_2 - O_2 leads to a decrease of the median VCD by ~ 12% and an increase of the median fitting uncertainty by ~ 10% with respect to the operational setup. Excluding H₂CO from the fitting significantly reduces the retrieved BrO columns by ~ 37%, given that the strong anticorrelation between both molecules is not taken into account. Fitting the mean residual (common mode) has a small impact on the retrieval results; the median VCD only changes ~ 3% but reduces the median fitting uncertainty by ~ 30% with respect to the exclusion of the common mode. To study the impact of the slit functions, we have performed the retrieval using both online slit functions, modeled as a Gaussian, and the preflight instrument slit functions. The median difference between these two retrievals is 27% for orbit number 26564. We have investigated the impacts of the order of scaling and baseline polynomials; it can cause uncertainties of $\sim 10\,\%$ as shown in Table 3.

To study the impact of the radiative transfer effects of the O_3 absorption in our retrieval, we have adopted the correction method described by Pukīte et al. (2010). We find that between 60° S and 60° N, the average difference is smaller than 10%, with values around 2% near the Equator. However, as we move near the poles with solar zenith angles above 60°, the differences start to be bigger, arriving at mean values around 30%.

4 Results and discussions

Comparisons of the OMI OMBRO product with GOME-2 satellite retrievals and remote sensing ground-based measurements over Harestua, Norway, as well as monthly mean

averages illustrate the quality of the retrieval on a global scale. On a local scale, recent scientific studies looking at BrO enhancements in volcanic plumes and over salt lakes are pushing the limits of the current OMBRO setups. In the following sections, we provide details of these comparisons (Sect. 4.1) and discuss OMI OMBRO global distribution (Sect. 4.2) and local enhancements over salt lakes and volcanic plumes observations (Sect. 4.3) and their applicability and strategies to correctly use the publicly available OMBRO product.

4.1 Comparisons with GOME-2 and ground-based observations

To assess the quality of the OMBRO product, we first compared OMI BrO VCDs with BIRA/GOME-2 BrO observations (Theys et al., 2011). GOME-2 has a descending orbit with a local Equator crossing time (ECT) of 09:30, and OMI has an ascending orbit with an ECT of 13:45. To minimize the effects of diurnal variation, especially under high SZAs (e.g., McLinden et al., 2006; Sioris et al., 2006), on the comparison, we conduct the comparison using simultaneous nadir overpasses (SNOs) within 2 min between GOME-2 and OMI predicted by NOAA National Calibration Center's SNO prediction tool (https://ncc.nesdis.noaa. gov/SNOPredictions, last access: 15 October 2017). Given Aura and Metop-A satellite orbits, all these SNOs occur at high latitudes around 75° S and 75° N. Figure 5 shows the time series of comparison of individual OMI/GOME-2 BrO retrievals from February 2007 through November 2008. The temporal variation of BrO at the SNO locations is captured similarly by OMI and GOME-2 BrO. The scatter plot in Fig. 6 quantifies the comparison between OMI and GOME-2 BrO. OMI BrO shows excellent agreement with GOME-2 BrO, with a correlation of 0.74 and a mean bias of $-0.216 \pm 1.13 \times 10^{13}$ molecules cm⁻² (mean relative bias of -2.6 ± 22.1 %). Considering very different retrieval algorithms including different cross sections and BrO profiles, such a good agreement is remarkable. GOME-2 retrievals use the BrO cross sections of Fleischmann et al. (2004), while our BrO retrievals use the BrO cross sections of Wilmouth et al. (1999). According to the sensitivity studies by Hendrick et al. (2009), using the Fleischmann cross section increases BrO by ~ 10 %. So, accounting for different cross sections, OMI BrO underestimates the GOME-2 BrO by ~ 10 %. In addition, the GOME-2 algorithm uses a residual technique to estimate tropospheric BrO from measured BrO SCDs by subtracting a dynamic estimate of stratospheric BrO climatology driven by O₃ and NO₂ concentrations and by using two different tropospheric BrO profiles depending on surface albedo conditions. This is very different from the approach of using a single BrO profile in the OMI BrO algorithm and can contribute to some of the BrO differences. Furthermore, additional algorithm uncertainties in both algorithms and different spatial sampling can also cause some



Figure 5. Time series comparison of SAO OMI (red) BrO and BIRA/GOME-2 (blue) BrO VCDs from February 2007 to November 2008 using simultaneous nadir overpasses occurring at high latitudes, around 75° S and 75° N, and within 2 min between OMI and GOME-2 observations.



Figure 6. Correlation and orthogonal regression of OMI and GOME-2 BrO for the data shown in Fig. 5. The legends show the mean bias and standard deviation of the differences, correlation and the orthogonal regression.

differences. Figure 7 shows the VCD monthly averages of GOME-2 data (green) and OMBRO (black) from February 2007 to December 2009, for which the seasonal variations are clearly seen. Our study shows that OMI has negative mean biases of 0.35×10^{13} molecules cm⁻² (12%), 0.33×10^{13} molecules cm⁻² (10%), 0.25×10^{13} molecules cm⁻² (17%) and 0.30×10^{13} molecules cm⁻² (10%) for Alaska, the South Pacific, Hudson Bay and Greenland, respectively.

We also used ground-based zenith-sky measurements of total column BrO at Harestua, Norway (Hendrick et al., 2007) to estimate the quality of the OMI BrO. We com-



Figure 7. VCD of GOME-2 (green) comparison to OMI (black) over four regions from February 2007 to December 2009 for four regions. Each region is defined by a square with the following latitude and longitude boundaries: Alaska ($50-70^{\circ}$ N, $165-135^{\circ}$ W), Hudson Bay ($50-65^{\circ}$ N, $95-75^{\circ}$ W), the South Pacific ($70-50^{\circ}$ S, $135-155^{\circ}$ E) and Greenland ($60-80^{\circ}$ N, $60-15^{\circ}$ W).

pared daily mean total BrO at Harestua with the mean OMI BrO from individual footprints that contain the location of Harestua site. Figure 8 shows the time series of the comparison between OMI total BrO and Harestua total BrO from February 2005 through August 2011 with the scatter plot shown in Fig. 9. Ground-based BrO shows an obvious seasonality with high values in the winter-spring and low values in the summer-fall. Such seasonality is well captured by OMI BrO. OMI BrO shows a reasonable good agreement with Harestua BrO, with a correlation of 0.46 and a mean bias of $0.12\pm0.76\times10^{13}$ molecules cm⁻² (mean relative bias of 3.18 ± 16.30 %, with respect to individual Harestua BrO). Sihler et al. (2012) compared GOME-2 BrO to ground-based observations at Utqiagvik (formerly Barrow), finding the correlation to be weaker (r = 0.3), likely due to both elevated and shallow surface layers of BrO. However, their correlation between GOME-2 BrO and ground-based measurements made from the icebreaker Amundsen, in the Canadian Arctic Ocean (r = 0.4), is closer to our correlation here. From the Harestua data, tropospheric BrO typically consists of 15 %-30 % of the total BrO, larger than what we have assumed in the troposphere. The use of a single BrO profile in the OMI BrO algorithm will likely underestimate the actual BrO. Accounting for the uncertainty due to profile shape, OMI BrO will have a larger positive bias relative to Harestua measurements, which can be caused by other algorithm uncertainties and the spatiotemporal differences between OMI and Harestua BrO.



Figure 8. Time series comparison of ground-based zenith-sky total BrO (black) at Harestua, Norway, and coincident SAO OMI BrO (red) from February 2005 through August 2011.

4.2 Global distribution of BrO VCDs

Figure 10 presents the global distribution of monthly mean BrO VCDs for selected months (March, June, September, December), showing BrO seasonality for three different years (2006, 2007 and 2012). BrO typically increases with latitude, with minimal values in the tropics ($\sim 2 \times 10^{13}$ molecules cm⁻²) and maximum values ($\sim 10^{14}$ molecules cm⁻²) around polar regions especially in the Northern Hemisphere winter–spring. In the tropics, BrO shows little seasonality, but at higher latitudes in polar re-



Figure 9. Correlation and orthogonal regression of OMI and Harestua BrO for the data in Fig. 8. The legends show the mean biases and standard deviations of the differences, correlation and the orthogonal regression.

gions, BrO displays evident seasonality. The seasonality is different between the Northern and Southern Hemisphere. In the Northern Hemisphere, BrO values are larger in spring (March) with widespread enhancement and are smaller in fall (September/December). In the Southern Hemisphere, BrO values are larger in southern hemispheric spring and summer (i.e., September and December) and smaller in the winter (June). Such global distribution and seasonal variation are generally consistent with previous satellite measurements (cf. Chance, 1998; http://bro.aeronomie.be/level3_monthly. php?cmd=map, last access: 15 October 2017). BrO in the tropics shows consistent zonal distributions, with lower values over land and in the intertropical convergence zone. This might be related to the impacts of clouds on the retrievals (e.g., BrO below thick clouds cannot be measured; there are uncertainties in the AMF calculation under cloudy conditions) and will be investigated in detail in future studies. The global distribution and seasonal variation are consistent from year to year, but the distributions from different years disclose some interannual variation. For example, BrO values in 2007 are smaller in January but are larger in March compared to those in 2006. Although OMI data since 2009 have been seriously affected by the row anomaly at certain crosstrack positions, the monthly mean data derived from good cross-track positions are hardly affected by the row anomaly as shown by the very similar global distribution and seasonality in 2012.



Figure 10. Global distributions of monthly mean BrO VCDs in March, June, September and December (in different rows) of 2006, 2007 and 2012 (different columns). Bromine release "explosions" during the polar spring months can be seen clearly.

4.3 Salt lakes and volcanic plumes enhancements of BrO

Following recent work by Hörmann et al. (2016) over the Rann of Kutch using OMI BrO retrievals from an independent research product, we have explored the capability of our OMBRO product to observe similar enhancements in other salt lakes. Figure 11 shows monthly averaged OMI BrO over the Great Salt Lake for June 2006, the corresponding surface albedo used in the retrieval, cloud cover (assuming a cloud filter of 40%), and the cloud pressure. Over the Great Salt Lake, BrO enhancement occurs predominantly over the lake bed, with enhancements of $\sim 5-10 \times 10^{12}$ molecules cm⁻² over background values $(3-4 \times 10^{13} \text{ molecules cm}^{-2})$. Despite observing these enhancements, the users of OMBRO for these kinds of studies should be aware of three limitations of the current retrieval algorithm. First, the BrO columns assume a mostly stratospheric BrO profile (Fig. 3a) for the AMF calculation. Second, the OMI-derived albedo climatology (Kleipool et al., 2008) used in OMBRO has a resolution of 0.5°. At this resolution OMBRO retrievals can have biases given the size of OMI pixels and the inherent subpixel albedo variability. Finally, high albedos inherent to salt lakes surface yield abnormally high cloud fractions and low cloud pressures over the salt lakes (Hörmann et al., 2016). All these factors should be considered in studies addressing the spatiotemporal distribution of BrO over salt lakes using OMBRO.

During our analysis of volcanic eruption scenarios, it was discovered that the currently implemented SO_2 molecular absorption cross sections (Vandaele et al., 1994) are a suboptimum choice (see Fig. 12). Compared to more recent laboratory measurements (Hermans et al., 2009; Vandaele et al.,



Figure 11. Mean June 2006 BrO VCD over the Great Salt Lake area. Averages have been calculated on a $0.2^{\circ} \times 0.2^{\circ}$ grid including only pixels with cloud fractions smaller than 0.4. The straight lines are borders of the state of Utah, and the curving lines represent the Great Salt Lake (eastern oval area) and Salt Flats (western oval area).

2009), the original SO₂ cross sections implemented in OM-BRO do not extend over the full BrO fitting window and exhibit the wrong behavior longward of 324 nm, overestimating the most recent measurement by up to a factor of 3. As the correlation between BrO and both SO₂ cross sections is very small $(-0.03 \text{ for the current } SO_2 \text{ and } 0.11 \text{ for the latest } SO_2$ cross sections) over the spectral range of SO₂ cross sections, interference by SO₂ in BrO retrievals is usually not an issue at average atmospheric SO₂ concentrations, but strong volcanic eruptions will render even small SO₂ absorption features past 333 nm significant. Around 334 nm, the Vandaele et al. (2009) data show an SO₂ feature that correlates with BrO absorption when SO₂ concentrations are significantly enhanced. As a consequence of this spectral correlation, SO₂ may be partially aliased as BrO, since the implemented SO₂ cross sections cannot account for it. Figure 13 presents an example from the 2010 Eyjafjallajökull eruption to show that the BrO retrieval can be affected by the choice of SO₂ cross sections. The next version of the OMBRO public release will be produced using the updated SO₂ absorption cross sections. Until then, caution is advised when using the OMI BrO product during elevated SO₂ conditions. We recommend using the OMBRO product together with the operational OMI SO₂ product (Li et al., 2013) to flag abnormally high BrO retrievals.

Figure 13a and b show daily average operational BrO VCDs from the eruption of the Eyjafjallajökull volcano on



Figure 12. Comparison of BrO absorption (red) and SO₂ absorptions under volcanic scenarios based on cross sections used in the operational algorithm (Vandaele et al., 1994) as shown in black and the recent laboratory cross sections (Vandaele et al., 2009) as shown in purple. For BrO, a SCD of 1.0×10^{14} molecules cm⁻² is assumed; for SO₂, a SCD of 15 Dobson units (i.e., 4.03×10^{17} molecules cm⁻²) is assumed. Cross sections have been convolved with the OMI slit function (which is assumed to be a Gaussian with 0.42 nm full width at half maximum).

5 and 17 May 2010, respectively. Enhanced BrO values in excess of 8.0×10^{13} molecules cm⁻² are detected in the vicinity of this volcano (e.g., plume extending southeastward from the volcano on 5 May and high BrO over Iceland on 17 May). Some of these enhanced BrO values correspond to the locations of enhanced SO₂ as shown on the NASA global SO₂ monitoring website (https://so2.gsfc.nasa.gov/, last access: 7 October 2017). This enhancement of BrO is not related to the seasonal variation of BrO as no such BrO enhancement is detected over Eyjafjallajökull during 5-17 May 2011 (a year after the eruption), with BrO values of only up to $\sim 5.3 \times 10^{13}$ molecules cm⁻² (not shown). Figure 13c and d show the same BrO retrievals using SO₂ cross sections by Vandaele et al. (2009). Using the improved SO₂ cross sections increases the BrO over a broader area on both days, supporting the theory that the choice of SO₂ cross sections can affect the BrO retrievals. However, BrO enhancement around the volcano can still clearly be seen with the improved SO_2 cross sections. This suggests that this BrO enhancement is not totally due to aliasing of SO₂ as BrO but potentially real BrO from the volcanic eruption.

5 Conclusions

This paper describes the current operational OMI BrO retrieval algorithm developed at SAO and the corresponding V3 OMI total BrO (OMBRO) product in detail. The OMI BrO retrieval algorithm is based on nonlinear least squares direct fitting of radiance spectra in the spectral range 319.0– 347.5 nm to obtain vertical column densities (VCDs) directly in one step. Compared to previous versions of two-step al-



Figure 13. Daily average BrO VCDs from Eyjafjallajökull on 5 May 2010 (a) and 17 May 2010 (b) produced using the operational SO_2 cross sections for the same days (c) and using the Vandaele et al. (2009) SO_2 cross sections (d).

gorithms, the fitting window was moved to shorter wavelengths, and the spectral range was increased to reduce the fitting uncertainty. Because air mass factors (AMFs) vary significantly with wavelength as a result of significant variation of O₃ absorption, the wavelength- and surface-albedodependent AMF, which is precomputed with the Linearized Discrete Ordinate Radiative Transfer (LIDORT) code using a single mostly stratospheric BrO profile, is applied prior to fitting to BrO cross sections for direct fitting of VCDs. Prior to the spectral fitting of BrO, wavelength calibration is performed for both irradiance and radiance at each cross-track position ,and reference spectra are properly prepared at the radiance wavelength grid. Then radiances are modeled from the measured solar irradiance, accounting for rotational Raman scattering, undersampling and attenuation from BrO and interfering gases and including additive and multiplicative closure polynomials and the average fitting residual spectrum. To maintain consistency with previous versions, a second fitting of all OMI spectra is performed with unmodified BrO cross sections to derive SCDs and the effective AMFs. Then a destriping step is employed to reduce the cross-trackdependent stripes.

The uncertainties of BrO VCDs included in the data product include only spectral fitting uncertainties, which typically vary between 4 and 7×10^{12} molecules cm⁻² (10 %–20 % of BrO VCDs; this could be as low as 5 % over BrO hotspots), consistent throughout the data record. The uncertainties vary with cross-track positions, from ~ 7 × 10¹² at nadir positions to ~ 4 × 10¹² at edge positions. We have investigated additional fitting uncertainties caused by interferences from O₂-O₂, H₂CO, O₃, and SO₂, the impact of the choice of fitting window, the use of common mode, the orders of closure polynomials and instrument slit functions. Uncertainties in the AMF calculations are estimated at $\sim 10\%$ unless the observation is made over a region with high tropospheric BrO columns. In this case, the use of a single stratospheric BrO profile is another source of uncertainty, overestimating AMFs (up to 50\%) and therefore underestimating BrO VCDs.

We compared OMI BrO VCDs with BIRA/GOME-2 BrO observations at locations of simultaneous nadir overpasses (SNOs), which only occur around 75° N and 75° S. OMI BrO shows excellent agreement with GOME-2 BrO, with a correlation of 0.74 and a mean bias of $-0.216\pm$ 1.13×10^{13} molecules cm⁻² (mean relative bias of $-2.6 \pm$ 22.1 %). Monthly mean OMBRO VCDs during 2007-2009 show negative biases of $0.25-0.35 \times 10^{13}$ molecules cm⁻² (10%-17%) over Alaska, the South Pacific, Hudson Bay and Greenland, respectively. We also compared OMI BrO with ground-based zenith-sky measurements of total BrO at Harestua, Norway. The BrO seasonality in Harestua is well captured by the OMI BrO and OMBRO retrieval, showing a reasonable good agreement with the ground-based measurements. The correlation between both datasets is 0.46, and the mean bias is $0.12 \pm 0.76 \times 10^{13}$ molecules cm⁻² (mean relative bias of 3.18 ± 16.30 %).

The global distribution and seasonal variation of OM-BRO are generally consistent with previous satellite measurements. There are small values in the tropics with little seasonality and large values at high latitudes with distinct seasonality. The seasonality is different between the Northern and Southern Hemisphere, with larger values in the hemispheric winter–spring (spring–summer) and smaller values in summer–fall (winter) for the Northern (Southern) Hemisphere. This spatiotemporal variation is generally consistent from year to year and is hardly affected by the row anomaly but does show some interannual variation. Finally, we have explored the feasibility of detecting enhanced BrO column over salt lakes and in volcanic plumes using OMBRO retrievals. We found enhancement of the BrO with respect to the background levels of $5-10 \times 10^{12}$ molecules cm⁻² over the US Great Salt Lake. We also observed a significant enhancement from the eruption of Eyjafjallajökull volcano although BrO retrievals under high SO₂ conditions can be affected by the current use of a sub-optimal choice of SO₂ cross sections.

Several important retrieval issues in the current operational algorithm that affect the quantitative use BrO VCDs have been raised in this paper such as the exclusion of O₂-O₂, nonoptimal SO₂ cross sections, the neglect of the radiative effect of O₃ absorption and the assumption of the stratospheric-only BrO profile. Users are advised to pay attention to these issues so that the product can be used properly. Future versions of OMBRO will include updated SO2 and O₂-O₂ cross sections, corrections for the radiative transfer effect of the O₃ absorption, and reoptimization of the spectral fitting windows to mitigate the interferences of other trace gases. We will also improve the AMF calculation accounting for clouds and O₃ and will consider the use of model-based climatological BrO profiles. These updates will increase the capabilities of the OMBRO retrieval to quantitatively estimate enhancements over salt lakes and in volcanic plumes.

Data availability. SAO OMI BrO retrievals were performed using radiance data publicly accessible at https://doi.org/10.5067/Aura/OMI/DATA1002 (Dobber, 2007). SAO OMBRO data are publicly available at https://doi.org/10.5067/Aura/OMI/DATA2006 (Chance, 2007).

Competing interests. The authors declare that they have no conflict of interest.

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