



Space-based observation of volcanic iodine monoxide

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Abstract. Volcanic eruptions inject substantial amounts of halogens into the atmosphere. Chlorine and bromine oxides have frequently been observed in volcanic plumes from different instrumental platforms such as from ground, aircraft and satellites. The present study is the first observational evidence that iodine oxides are also emitted into the atmosphere during volcanic eruptions. Large column amounts of iodine monoxide, IO, are observed in satellite measurements following the major eruption of the Kasatochi volcano, Alaska, in 2008. The IO signal is detected in measurements made both by SCIAMACHY (Scanning Imaging Absorption Spectrometer for Atmospheric CHartography) on ENVISAT (Environmental Satellite) and GOME-2 (Global Ozone Monitoring Experiment-2) on MetOp-A (Meteorological Operational Satellite A). Following the eruption on 7 August 2008, strongly elevated levels of IO slant columns of more than 4×10^{13} molec cm⁻² are retrieved along the volcanic plume trajectories for several days. The retrieved IO columns from the different instruments are consistent, and the spatial distribution of the IO plume is similar to that of bromine monoxide, BrO. Details in the spatial distribution, however, differ between IO, BrO and sulfur dioxide, SO₂. The column amounts of IO are approximately 1 order of magnitude smaller than those of BrO. Using the GOME-2A observations, the total mass of IO in the volcanic plume injected into the atmosphere from the eruption of Kasatochi on 7 August 2008, is determined to be on the order of 10 Mg.

1 Introduction

Halogen oxides strongly influence atmospheric composition. Catalytic reaction cycles involving chlorine, bromine or iodine, lead to ozone depletion in the troposphere. In the stratosphere, the role of chlorine and bromine, which are released predominantly as a consequence of anthropogenic emissions of chlorofluorocarbon compounds, is well established (World Meteorological Organization, 2014). The potential importance of iodine reactions for stratospheric ozone depletion is discussed in Solomon et al. (1994), Hossaini et al. (2015), and Saiz-Lopez et al. (2015a).

Stratospheric concentrations of iodine species are much lower than those of chlorine and bromine compounds (Bösch et al., 2003; Butz et al., 2009). From balloon-borne observations, an upper limit for stratospheric iodine monoxide, IO, of 0.1 parts per trillion by volume (pptv) was determined in the tropics (Butz et al., 2009), while upper limits for IO of 0.2 pptv at 20 km, or 0.1 pptv at 15 km, were derived in the mid- and high latitudes (Pundt et al., 1998). Butz et al. (2009) estimate upper limits of total gaseous iodine of about 0.09 to 0.16 pptv in the tropical lower stratosphere (21.0 to 16.5 km) and 0.17 to 0.35 pptv in the tropical upper troposphere (16.5 to 13.5 km). A recent study by Saiz-Lopez et al. (2015a) estimates that stratospheric iodine may range between 0.25–0.7 pptv. This is based on, for example, new aircraft observations in the tropics from which volume mixing ratios (VMRs) of IO between 0.1–0.2 pptv at altitudes up to 14 km were retrieved (Volkamer et al., 2015).

The ozone destruction potential of stratospheric iodine is significantly higher than that of the other halogens. Bromine is about 60 times more effective in destroying ozone than chlorine and, for iodine, the factor is about 150 to 300

(World Meteorological Organization, 2014). The effective chain length of the catalytic cycles involving iodine and IO is larger than those involving the other halogens. This is in part because the temporary reservoir species containing iodine are photolysed and/or react more rapidly with stratospheric free radicals than their chlorine or bromine analogues. As a result, reactive iodine may impact stratospheric ozone chemistry (Solomon et al., 1994; Hossaini et al., 2015) even at sub-pptv levels.

IO is formed by the reaction of iodine radicals with ozone, O₃. Catalytic cycles including IO, by which tropospheric O₃ is effectively destroyed, were already proposed in the 1980s (Chameides and Davis, 1980). As a result of self reactions, iodine oxides may lead to particle formation and thereby affect atmospheric radiation balance (Burkholder et al., 2004; O'Dowd and Hoffmann, 2005; Saunders et al., 2010). These effects of iodine motivate the scientific interest in the assessment of sources, amounts and distributions of iodine species in the atmosphere.

Atmospheric iodine is of organic as well as inorganic origin, e.g. from emissions of I₂ and of halogenated organic compounds such as CH₃I and CH₂I₂ (Saiz-Lopez et al., 2012, and references therein). The largest iodine sources in general are the world's oceans. Iodine compounds are emitted into the marine boundary layer, e.g. from algae (Schall et al., 1994; Aliche et al., 1999; Carpenter, 2003) or via inorganic pathways involving the ocean surface (Garland and Curtis, 1981; Carpenter et al., 2013). In the polar troposphere, bromine and iodine oxides are both observed predominantly during spring time. Release mechanisms of iodine and bromine above sea ice areas, however, are considerably different. Bromine monoxide, BrO, is released following an autocatalytic Br activation (Vogt et al., 1999), also known as the bromine explosion mechanism. Iodine most probably takes different pathways involving the release of organoiodine compounds (Saiz-Lopez et al., 2015b), while inorganic reactions cannot be excluded.

Volcanic eruptions are an important source of halogens in the atmosphere, especially for the free and upper troposphere and the lower stratosphere (von Glasow et al., 2009). Volcanic plumes are known to contain halogen species initially in acidic form, e.g. HF, HCl, HBr and HI (Aiuppa et al., 2009). Bromine oxides as well as chlorine oxides have been previously observed in volcanic plumes. Volcanic BrO was first detected by Bobrowski et al. (2003), who applied the well-established Differential Optical Absorption Spectroscopy (DOAS) technique (Platt and Stutz, 2008) with a ground-based Multi-Axis DOAS (MAX-DOAS) system. Volcanic chlorine oxides, ClO and OClO, were also measured, e.g. by Lee et al. (2005) and Bobrowski et al. (2007), using ground-based DOAS instruments. From space, volcanic BrO was detected for the first time from the Kasatochi eruption in 2008 (Theys et al., 2009), followed by volcanic OClO from the Puyehue eruption in 2011 (Theys et al., 2014). Several further observations using ground-based mea-

surements (Bobrowski et al., 2006; Bobrowski and Platt, 2007; Kern et al., 2009), airborne instrumentation (General et al., 2015) as well as satellites (Hörmann et al., 2013) have confirmed and further quantified the abundances of bromine oxides injected into the atmosphere following volcanic eruptions. The release mechanism of volcanic BrO is believed to be similar to polar tropospheric BrO and is based on an autocatalytic reaction cycle involving volcanic aerosols (Bobrowski et al., 2007). Ozone depletion was observed within volcanic plumes and is attributed to reactive halogen chemistry (Lee et al., 2005; Surl et al., 2015, and references therein).

Using filter techniques, measurements at Mt Etna in Italy (Aiuppa et al., 2005) and at the Masaya and Telica volcanos in Nicaragua (Witt et al., 2008), for example, showed that gaseous HI, I and HBr are relevant constituents in the degassing of these specific volcanos. Only a few studies are available that report on the analysed iodine content in samples of volcanic gases or volcanic fluids. Snyder and Fehn (2002) investigated the ¹²⁹I/I ratio in volcanic fluids in order to determine the ages of iodine species. The determined iodine ages are in agreement with the expected age of subducted sediments. An iodine accumulation takes place here, because marine sediments contain concentrated amounts of organic iodine. Iodine oxides have not been previously detected in the emission plumes of volcanos. Gliß et al. (2015) report an upper limit for IO slant columns of 7.6 to 8.6 × 10¹² molec cm⁻² based on the detection limit of their ground-based DOAS observations at Mt Etna, Italy, during a stable quiescent degassing phase in September 2012.

The composition of volcanic gases is in general strongly variable, with individual characteristics changing from volcano to volcano as well as between eruption and degassing phases (Witt et al., 2008; Aiuppa et al., 2009, and references therein). While the gas-phase composition is individual for each volcanic eruption, there is also a general difference between iodine and other halogens in volcanic gases at high temperatures. At around 1000 °C, the main constituents are HF, HCl and HBr for the other halogens. For iodine, however, HI and atomic I may be present in equal amounts (Aiuppa et al., 2005).

Up to the present, no detection of gaseous iodine oxides of volcanic origin has been reported, either by in-situ measurements or by remote sensing from ground or satellite. Iodine monoxide is retrieved from satellite measurements of backscattered solar radiation by the DOAS technique and is observed, for example, in the south polar region (Saiz-Lopez et al., 2007; Schönhardt et al., 2008, 2012). In most cases, atmospheric amounts of IO are fairly small, so that usually temporal averages of the satellite data of at least 1 month are created in order to improve signal-to-noise ratio.

In August 2008, the eruption of the Kasatochi volcano took place (Waythomas et al., 2010). Kasatochi belongs to the volcanic arc of the Aleutian Islands, Alaska. The violent explosions started on 7 August 2008, in the afternoon.

The Volcanic Explosivity Index (VEI) (Newhall and Self, 1982), which classifies the eruptive volume and eruption cloud height, was VEI 3–4. Large amounts of ash and sulfur dioxide, SO₂, were released into the atmosphere, reaching the lower stratosphere (Waythomas et al., 2010). In total about 1.7 Tg of SO₂ were emitted and spread over large parts of the globe.

In the following, the detection of IO from the eruption of the Kasatochi volcano using observations of the SCIAMACHY (Scanning Imaging Absorption Spectrometer for Atmospheric CHartographY) and GOME-2A (Global Ozone Monitoring Experiment-2A) satellite instruments is presented and discussed. The applied instruments and retrieval settings are briefly described, and the IO spectral fit quality is investigated. The IO results are analysed in terms of spatial distribution, temporal evolution and integral amount. In addition, IO and BrO distributions in the volcanic plume are compared among each other and to those of SO₂.

2 Instruments and Measurements

The only satellite-borne spectrometer for which an IO product has been reported so far is the SCIAMACHY instrument onboard the European Environmental Satellite (ENVISAT) (Saiz-Lopez et al., 2007; Schönhardt et al., 2008, 2012). The mission operated between March 2002 and April 2012. In the present study, data from the GOME-2A onboard MetOp-A (Meteorological Operational Satellite A) have also been successfully analysed, and the signatures of IO and BrO absorption have been retrieved. IO detection by GOME-2A is reported here for the first time. The DOAS method is used for the retrieval of trace gas amounts.

2.1 Satellite instruments and data

SCIAMACHY is a spectrometer measuring direct, scattered and reflected sunlight in the UV, visible and near-IR spectral regions. The spectra are measured contiguously from 214 to 1773 nm and in two spectral bands within the ranges of 1934–2044 and 2259–2386 nm. The operation modes include nadir, limb and occultation geometries (Burrows et al., 1995; Bovensmann et al., 1999; Gottwald and Bovensmann, 2011). The present study uses the nadir observations. ENVISAT has a Sun-synchronous, near-polar orbit with a local equator crossing time of 10:00 a.m. in descending node. Individual SCIAMACHY ground pixels in the spectral range used here have a typical size of 30 × 60 km². For the IO retrieval in general, spatial averaging over four ground pixels is applied to reduce noise. A further reduction in spatial resolution occurs for some parts of each orbit as a result of using the SCIAMACHY read-out from cluster 14 in channel 3 (404–424 nm) in addition to the more commonly used cluster 15 (424–527 nm). Cluster 14 has a partly longer integration time than cluster 15, and the integration time is adapted for the

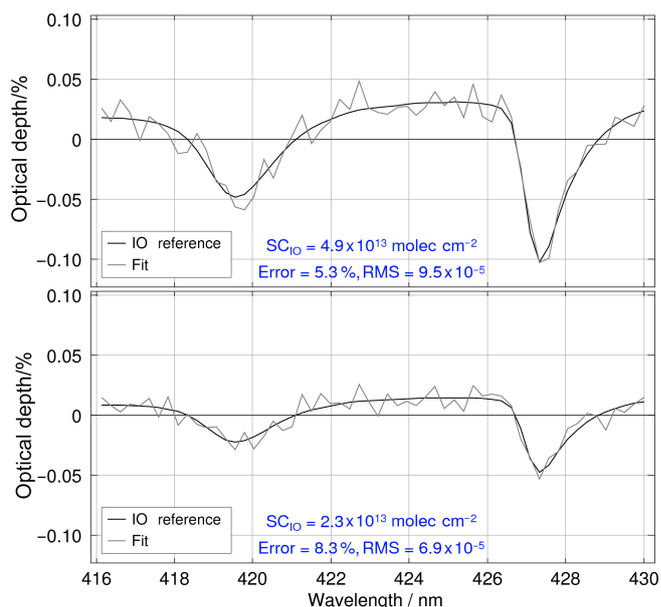


Figure 1. Example fitting results from SCIAMACHY on day 11 August 2008, with two different IO slant column amounts recorded at 55.34° N, 220.21° E (top) and 54.90° N, 215.92° E (bottom). The fit (grey) shows the measurement optical depth after all other features except for the IO absorption (black) have been subtracted. The optical depth RMS values are 9.5 and 6.9 × 10^{−5}, respectively.

entire spectral region to achieve smoothed spectra across the cluster border. The largest across-track ground-scene pixel size is 240 km.

The GOME-2A instrument observes in the UV and visible spectral regions from 240 to 790 nm and performs measurements in nadir viewing geometry. Launched in July 2006 onboard MetOp-A, GOME-2A is the first of three nearly identical instruments. The mission officially started in October 2006, and data is available since January 2007. The equator crossing time of MetOp-A is 09:30 a.m. As for SCIAMACHY, spatial averaging is applied for the GOME-2A data in order to achieve noise reduction. The typical ground pixel size of 40 × 80 km² is thus increased to 80 × 160 km². For some direct comparisons with BrO, however, the IO results without spatial averaging are used.

2.2 DOAS retrievals of IO and BrO

The DOAS method is applied to the satellite measurements in order to retrieve IO and BrO column amounts. For the SCIAMACHY IO product, the standard retrieval settings as published in Schönhardt et al. (2008) and summarized in Table 1 are used. Two example fitting results from day 11 August 2008 are displayed in Fig. 1, showing the spectral fits for IO columns of 4.9 × 10¹³ molec cm^{−2} (top) and 2.3 × 10¹³ molec cm^{−2} (bottom). The comparably large IO column amounts are detected with rather small relative fitting errors of 5.3 and 8.3 %, respectively.

Table 1. Retrieval settings for IO from SCIAMACHY and GOME-2A observations.

Retrieval settings	SCIAMACHY	GOME-2A 2T	GOME-2A 3T
Fitting window	416–430 nm	416–430 nm	418–438 nm
Polynomial degree	2 (quadratic)	2 (quadratic)	3 (cubic)
Trace gases	NO ₂ , O ₃ IO	NO ₂ , O ₃ IO	NO ₂ , O ₃ , O ₄ IO
Other features	Ring effect: SCIATRAN calculation (Roazanov et al., 2014; Vountas et al., 1998) Linear intensity offset correction		
Background	Daily earthshine, Siberia (60–70° N, 80–120° E)		

For the analysis of GOME-2A data, two alternative retrievals are used and are also listed in Table 1. The 2T retrieval corresponds to the standard SCIAMACHY IO retrieval and therefore covers the same two IO transition bands. The GOME-2A data show higher noise levels than the respective SCIAMACHY measurements. Consequently, for the analysis of GOME-2A data, the use of more spectral information from a larger fitting window is investigated. The resulting 3T retrieval covers three transition bands of the IO absorption spectrum. For SCIAMACHY, the 3T retrieval was not successful (Schönhardt et al., 2008) due to instrument-related spectral features above 430 nm. It leads, however, to an improved quality of the IO retrievals from GOME-2A measurements. If not specified otherwise, in the following, GOME-2A IO results from the 3T retrieval are used. In terms of IO amounts, the results are consistent within the uncertainties between both GOME-2A retrievals as well as between GOME-2A and SCIAMACHY, cf. Sect. 3.2.

For all IO retrievals, a daily averaged earthshine spectrum is used as reference background. This background spectrum is generated from a reference area 60 to 70° north and 80 to 120° east – a continental region which is likely to contain only small column amounts of IO. Consequently, a differential slant column between the specific location and this reference region is retrieved.

The cross sections used for SCIAMACHY retrievals are NO₂ (223 K) (Bogumil et al., 2003) and O₃ (223 K) (Bogumil et al., 2003). For GOME-2A retrievals, NO₂ (223 K) and O₃ (223 K) from measurements with the GOME-2 flight model are used (Chehade et al., 2013, and P. Spietz, personal communication, 2005) as well as O₄ (Greenblatt et al., 1990), with the latter only for the GOME-2A 3T retrieval. The absorption structures in the O₄ spectrum are small in the spectral range of the IO fitting window and, in addition, these small structures differ rather strongly between the three available O₄ cross sections in the literature (Greenblatt et al., 1990; Thalman and Volkamer, 2013; Hermans, unpublished data, <http://spectrolab.aeronomie.be/o2.htm>). However, the inclusion of any of the three O₄ cross sections or completely omitting O₄ from the IO retrieval has no significant influence on the resulting IO slant columns in the volcanic plume. For all retrievals, the IO (298 K) cross section measured by

Gómez Martín et al. (2007) is applied, convolved with the slit function of the respective instrument.

BrO columns are retrieved from GOME-2A in a fitting window from 336 to 347 nm, taking into account absorption features of O₃ (223 and 273 K), NO₂ (223 K) and BrO (Be-goin et al., 2010). A cubic polynomial with four coefficients is fitted for the broadband spectral effects.

SCIATRAN calculations (Roazanov et al., 2014) are used to determine reference spectra for rotational Raman scattering (Ring effect), which is taken into account in all retrievals. An additional additive intensity offset compensates for effects such as stray light or different types of inelastic scattering, e.g. not fully compensated Ring structures, the influence of vibrational Raman scattering, VRS, in air (Lampel et al., 2015) and VRS on liquid water or liquid water absorption (Peters et al., 2014). Including VRS spectra of N₂ and O₂ explicitly in the IO retrieval does not change the resulting IO slant columns significantly.

The DOAS analysis yields the differential trace gas slant column amounts, which are the differences between two spectra in absorber concentrations integrated along the mean light path. In order to convert these slant column amounts into vertical column amounts, the air mass factor (AMF), i.e. the ratio between the slant and vertical column, is computed. For both IO and BrO, a geometric AMF is applied here which is suitable for a stratospheric absorber. For the current study, assuming a geometric AMF is adequate since the volcanic plume is located at fairly high altitudes (Theys et al., 2009) and the relevant solar zenith angle is below 50°. The influence of aerosols on light scattering and thus on the AMF is not considered in this work. Aerosols can increase or decrease visibility of trace gases depending on several aspects such as aerosol characteristics and the relative altitude distributions. Here we concentrate on a more qualitative discussion of the observed halogen amounts and distributions.

3 Results

3.1 Observation of volcanic IO

After the eruption of Kasatochi, enhanced IO column amounts are detected within the volcanic plume for several

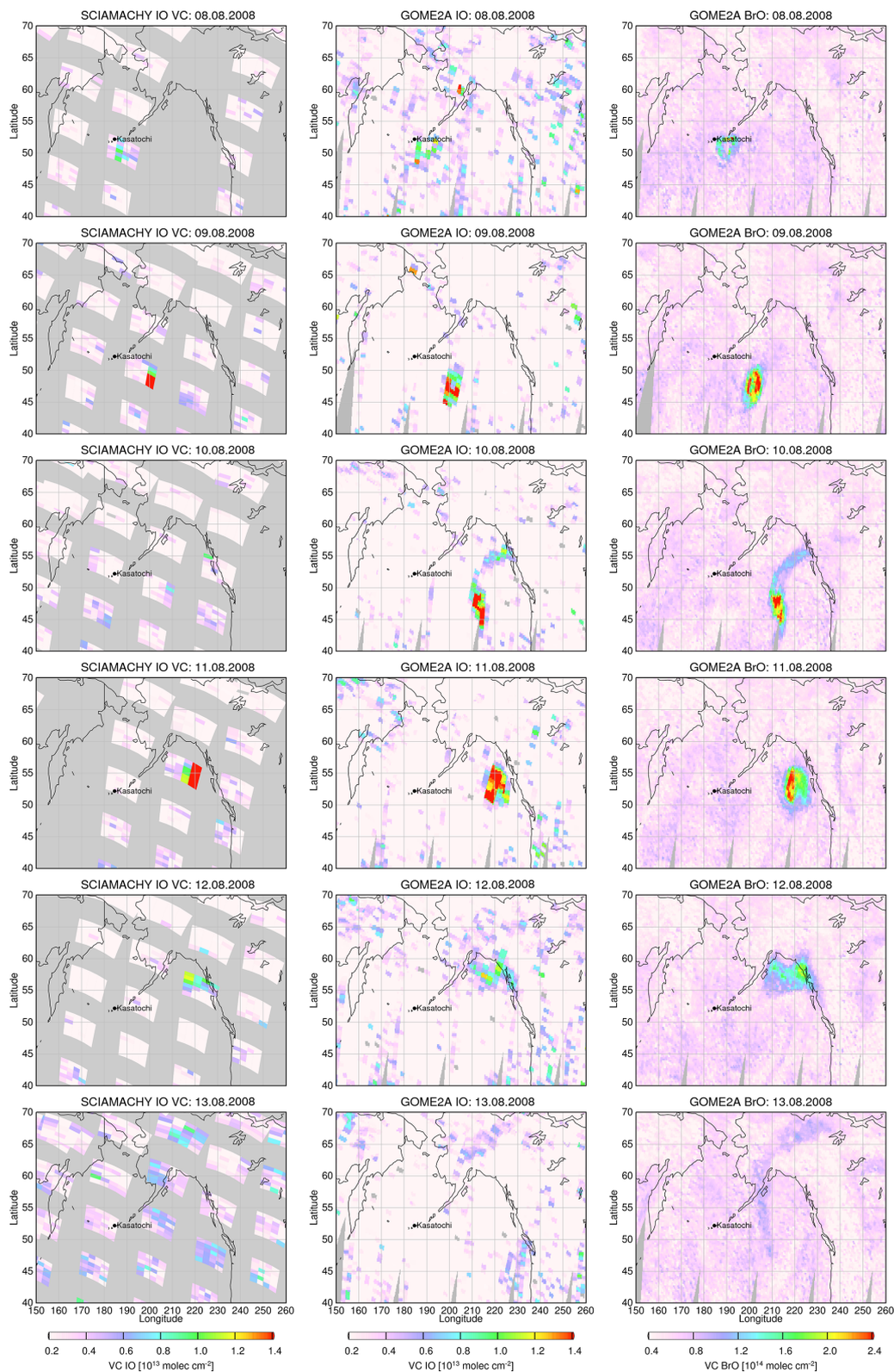


Figure 2. Retrievals of IO from SCIAMACHY (left) and GOME-2A (middle) together with observations of BrO from GOME-2A (right) for 6 days following the eruption of the Kasatochi volcano on 8 August 2008. Regions without data coverage are shaded in grey.

days. As a consequence of the morning overpass times of the satellite instruments, the eruption, which started in the afternoon of 7 August 2008, can be observed from 8 August 2008 onwards. In Fig. 2, left column, the observational results from the SCIAMACHY IO retrieval are shown for 6 days from 8 to 13 August 2008. IO enhancements are detected on all 6 days as well as enhancements of BrO (not shown, see http://www.iup.uni-bremen.de/doas/scia_data_browser.htm?gas=bro&column=strat&view=nh&year=2008&month=8&day=8). On 8 August, a loop-shaped area with enhanced IO is visible, and maximum slant column amounts are around 2.3×10^{13} molec cm⁻². In the same area, BrO reaches slant column values up to 4.2×10^{14} molec cm⁻². The slant column amounts on 9 August are higher, with 3.4×10^{13} and 5.6×10^{14} molec cm⁻² for IO and BrO, respectively. While on day 10 August the volcanic plume is situated just in between two SCIAMACHY orbits, and only slightly enhanced amounts are seen at the edges of the plume in the adjacent orbits (at 50° N, 210° E, and 55° N, 225° E), the SCIAMACHY IO column amounts are largest on day 11 August. Slant columns reach up to 4.9×10^{13} molec cm⁻² for IO and 5.6×10^{14} molec cm⁻² for BrO. These values correspond to vertical columns of 2.1×10^{13} molec cm⁻² for IO and 2.5×10^{14} molec cm⁻² for BrO. While these large column amounts of BrO from volcanic emission have been reported before (Theys et al., 2009), IO produced by volcanic activity is observed for the first time.

The IO column amounts in the Kasatochi emission plume are larger than the upper limit for IO slant columns of 7.6 to 8.6×10^{12} molec cm⁻² reported by Gliß et al. (2015) for the degassing of Mt Etna in September 2012. These results are not in contradiction with the satellite observations in the present study, as different volcanos show individual gas-phase compositions, and degassing phases may differ strongly from eruptive periods.

3.2 IO detection with GOME-2A

Maps of IO retrieved from GOME-2A data for the 6 days after the eruption are shown in Fig. 2 (middle), next to the SCIAMACHY results for direct comparison. BrO amounts retrieved from GOME-2A are shown in the right column. Due to the much better spatial coverage of the GOME-2A instrument as compared to the SCIAMACHY instrument, the IO plume from the volcanic emission is clearly visible on all 6 days. The spatial shape of the IO enhancement agrees well with the area where higher BrO is observed.

As a comparison of the IO results retrieved from the two different sensors, an example collocation case from 11 August 2008 has been chosen. The comparison includes the IO from the SCIAMACHY retrieval as well as IO from both GOME-2A retrievals, and the results are summarized in Table 2. The IO retrieval settings are discussed in Sect. 2.2, and the corresponding spectral fits are shown in Fig. 3.

The IO results of the three retrievals are consistent within their uncertainties. The GOME-2A spectral retrievals are of good quality, with relative fitting errors of around 14 %. The fitting error is thus larger than for the SCIAMACHY retrieval. The IO detection limit for GOME-2A observations is on the order of 5×10^{12} molec cm⁻² in terms of vertical columns and around 1×10^{13} molec cm⁻² for the retrieved slant columns, depending on several factors, such as the received radiance and solar zenith angle. For the discussed examples, the GOME-2A instrument detects slightly less IO than the SCIAMACHY instrument. On other collocation events the relation is, however, reversed. The ground scenes of the two instruments are not identical, and the measurement times typically differ by half an hour. For rapidly moving volcanic plumes, differences in the detected IO column amounts by the two instruments are expected, either as result of changing IO concentrations due to relatively fast and complex multiphase photochemical reactions, the size of the ground scene, or changing ground or cloud albedo.

3.3 Analysis of IO and BrO amounts

The spatial sampling of spectra by the GOME-2A instrument is intrinsically better than that of the SCIAMACHY instrument, and the full volcanic plume is observed on several days. Consequently, GOME-2A IO results provide a more accurate analysis and representation of the total iodine amount and mass emitted from the Kasatochi eruption than the IO results retrieved from SCIAMACHY. Integration over the IO amount inside the plume is performed. For this purpose, the plume itself needs to be defined first. Here, the plume is defined as the area enclosing those satellite pixels with an IO column amount above a certain threshold. This threshold is defined as $VC_{IO,thr} = \langle VC_{IO} \rangle + 2\sigma_{IO}$, where $\langle VC_{IO} \rangle$ is the mean IO vertical column and σ_{IO} is its standard deviation. Both parameters are derived from measurements on the days before the eruption as explained below. For BrO, the procedure is almost the same, but the threshold is set at $3\sigma_{BrO}$ above the mean. For IO, the weaker criterion of 2σ is necessary in order to capture the plume well. The reason for this is the larger noise as compared to that of the BrO data, i.e. enhanced IO amounts are closer to the detection limit than is the case for BrO. Mean and standard deviation values for IO and BrO are calculated using the data from 3 consecutive days of satellite coverage prior to the eruption (5 to 7 August 2008) and from within a wide area around the volcano (40–62.5° N, 183.5–231° E) enclosing all main plumes on the following days. Threshold values are 5.3×10^{12} and 9.7×10^{13} molec cm⁻² for IO and BrO, respectively. Only a small background IO slant column is found prior to the eruption (around 0.4×10^{12} molec cm⁻² and below the detection limit), while the BrO column has a substantial stratospheric as well as free-tropospheric contribution of around 6.1×10^{13} molec cm⁻² in this area.

Table 2. IO retrieval results from the collocation case between SCIAMACHY and GOME-2A on 11 August 2008. The three results agree within their fitting errors.

Retrieval results	SCIAMACHY	GOME-2A 2T	GOME-2A 3T
Fitting window	416–430 nm	416–430 nm	418–438 nm
$SC_{IO}/10^{13}$ molec cm^{-2}	4.58 ± 0.28	4.25 ± 0.60	4.14 ± 0.59

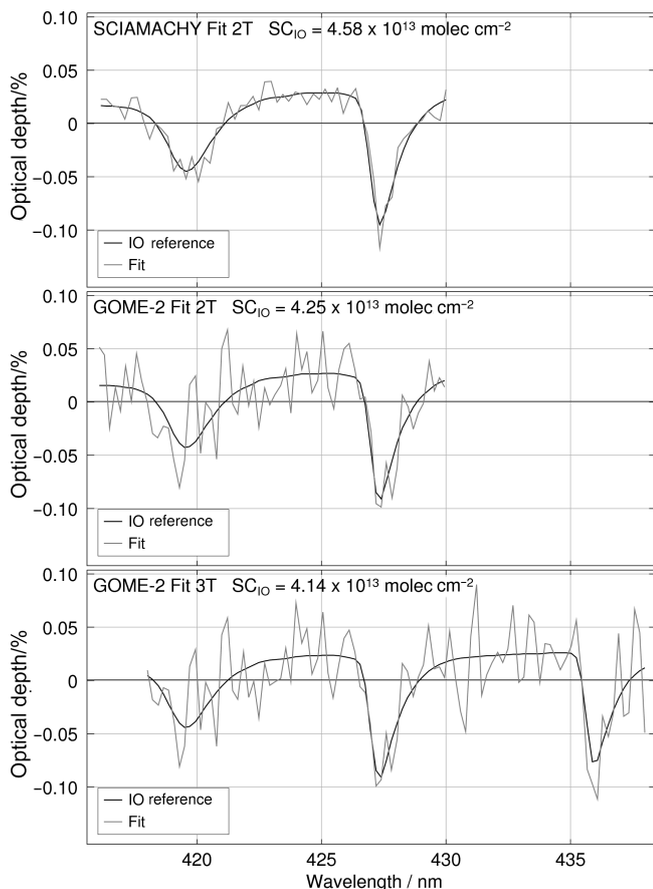


Figure 3. Spectral fitting results from day 11 August 2008 for a collocation between SCIAMACHY (top) and GOME-2A around 55° N and 220° E. For GOME-2A, the results from two different fitting windows are shown, using the SCIAMACHY standard IO fitting window (centre) and using a larger spectral window covering three spectral absorption bands of IO (bottom).

In an alternative approach, the observed SO_2 column amount (cf. Sect. 3.4) is used in order to select the BrO and IO in the volcanic plume. For this SO_2 mask approach, the plume is defined by applying a 10 DU limit to the SO_2 distribution. Following the plume definition and background subtraction, the IO and BrO column amounts are integrated over the selected plume area. This yields an integrated number of molecules originating from the volcanic eruption. For the days 8 to 12 August 2008, the results of this procedure are shown as a time series for IO and BrO in Fig. 4. Results from

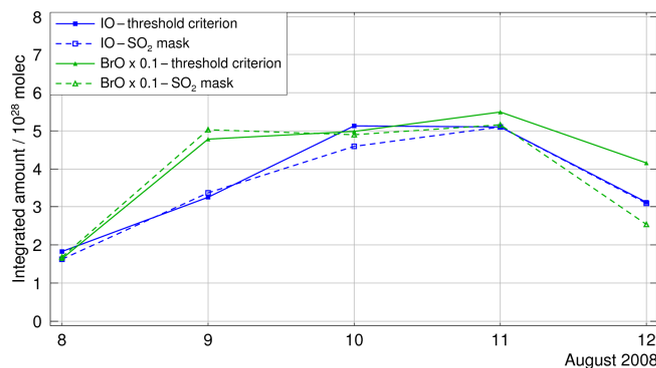


Figure 4. Time series of integrated IO (blue) and BrO (green) amounts. BrO data is scaled by a factor of 0.1. For both trace gases, two methods for the plume definition are applied, the threshold criterion and the SO_2 mask.

the threshold criterion are displayed and compared to those obtained using the SO_2 mask approach.

On days 8 to 11 August 2008, the two methods agree within a few percent, while the actual plume shape differs slightly at the edges. On 12 August 2008, the difference of the plume positions between SO_2 on one hand and BrO and IO on the other is larger. For IO, the difference does not affect the integrated value much, while for BrO, the results from the two different selection routines differ by 60%. Using the SO_2 mask, part of the BrO plume is missed. For IO, the values in that region are lower. Consequently, the influence of the precise plume shape on the calculation of the integrated amount is less pronounced. In general, the two methods are in agreement but, due to the latter finding, the method using the σ -level criterion is preferred.

On day 8 August, just after the start of the eruption, approximately 1.8×10^{28} molecules of IO are observed in the plume, corresponding to a mass of 4.3 Mg or metric tons, t, of IO. The amount of IO increases to 7.7 t on 9 August, reaches up to 12.2 and 12.1 t (i.e. around 5.1×10^{28} molecules of IO) on the peak days 10 and 11 August, respectively, and decreases back to 7.4 t on 12 August. The integrated mass of IO hence ranges between 4.3 and 12.2 t. Using the molar masses of iodine and oxygen, this amount of IO contains an integrated mass of reactive iodine between 3.9 and 10.8 t.

The integrated mass of BrO within the plume increases from 26 t on 8 August to 76 t and 79 t on 9 and 10 August and reaches a maximum of 87 t on 11 August. On 12 August

2008, an integrated mass of 66 t of BrO remains in the volcanic plume. Directly converting the integrated mass of BrO between 26 and 87 t to the corresponding integrated mass of reactive Br, a range between 22 and 73 t is derived using the molar masses of bromine and oxygen. These integrated BrO amounts are larger but in broad agreement with calculations by Theys et al. (2009), who use the FLEXPART dispersion model and derive the total amount of BrO within the volcanic plume to be around 30 to 42 t. In addition to BrO, other bromine compounds contribute to the total bromine mass. In the relevant altitude between 8 and 12 km, 30–50% of the total inorganic bromine exist in the form of BrO (Theys et al., 2009). Using this relation, the integrated BrO amount corresponds to a total mass of 50 to 290 t of reactive bromine.

Although knowledge on iodine chemistry in a volcanic plume is limited, other iodine compounds such as I₂, I, HI, HOI, OIO and higher iodine oxides are presumably present in the emission plume as well. Consequently, the emitted mass of iodine (3.9 to 10.8 t) can be regarded as a lower limit for the iodine content in the Kasatochi emission plume because this range is derived directly from the IO observations. Detailed chemical modelling would be needed to derive the total amount of reactive iodine in the volcanic plume from the observed IO column amounts by taking into account the other iodine species and all known chemical reactions that are taking place in the hot exhaust of the individual volcano. Such a modelling exercise is, however, out of the scope of the current study. In addition to the presence of other iodine species, iodine oxides may polymerize into particles, while there is no evidence that bromine oxides do under atmospheric conditions. This might lead to an underestimation of the iodine to bromine ratio if only gas-phase species are considered.

The emitted mass of iodine inferred for the Kasatochi eruption in August 2008 is of the same order of magnitude as previously determined for the annually integrated flux for degassing volcanos, e.g. 10 t yr⁻¹ of iodine at Mt Etna, Italy, (Aiuppa et al., 2005) or 12 t yr⁻¹ at Satsuma-Iwojima, Japan (Snyder and Fehn, 2002). This is in line with observations for bromine, where for one given volcano the Br flux from an individual eruption can be of the same order of magnitude as the annual Br flux from degassing (Aiuppa et al., 2005). The temporal evolution of the integrated amounts is discussed in Sect. 4.

3.4 Spatial distributions of IO, BrO and SO₂

In order to investigate the spatial plume structure more closely, Fig. 5 gives an expanded view of the volcanic plume. The retrieved column amounts of IO (left) and BrO (centre) are shown together with those of SO₂ (right) for the days 9 August 2008 (top) and 11 August 2008 (bottom). SO₂ column amounts are derived in the spectral window between 312.5 and 327 nm using an iterative retrieval approach (Richter, 2009).

Previous satellite studies reported that BrO is often enhanced around the plume centre (Hörmann et al., 2013). For the two depicted cases, the IO column amount is also lower in the plume centre than in some areas around the centre. In general, the IO and BrO plumes have a similar spatial extent and shape. It is however interesting to note, that maximum IO and BrO column amounts are not observed in the same satellite pixels and that the details of the spatial patterns differ. On 9 August 2008, the largest BrO enhancements are detected in the west and east of the plume, while IO is also enhanced there but even more so in the south of the plume. On 11 August 2008, BrO maxima are seen in the west, and IO maxima are split into two regions in the north and south of the volcanic plume.

The IO and BrO vertical column amounts that are observed within a rectangular latitude–longitude box which encloses the entire volcanic plume are investigated for each individual day between 8 and 12 August 2008. The correlation coefficient between IO and BrO considering the data from the respective area lies between $R = 0.62$ and $R = 0.84$ on the days from 9 to 12 August 2008. On the first day, 8 August, the correlation is lower, at $R = 0.42$. These results with relatively large and positive values of R indicate that iodine and bromine compounds are emitted together into the volcanic plume but also that there are factors influencing the temporal evolution of the two gases differently as R is clearly below unity ($R < 0.85$).

The IO and BrO distributions are again similar to those of SO₂, but even larger differences occur than between the distributions of the two halogen compounds. For SO₂, no occurrence of lower values in the plume centre is observed. On some days, such as the example day 11 August, SO₂ is at maximum in the plume centre. On day 9 August, two SO₂ maxima are seen, with one part crossing the plume centre and one part situated more to the southern edge of the plume. The three different trace gases observed by satellite hence show several individual aspects in their spatial distribution within the volcanic plume.

4 Discussion

When comparing the integrated numbers of IO and BrO molecules in the volcanic plume, one important and interesting point is that the amount of iodine is only about 1 order of magnitude smaller than that of bromine. For the individual days from 9 to 12 August 2008, the ratio for the integrated number of BrO to IO molecules lies between 6.7 and 10.0 and amounts to 4.2 on 8 August 2008. The corresponding mass ratio for BrO to IO ranges between 4.0 and 6.7 and amounts to 2.8 on 8 August 2008, using data from Fig. 4.

Figure 6 shows a scatterplot between IO and BrO column amounts from the individual satellite observations. Data from the 4-day period, 9 to 12 August 2008, is included in the comparison. As in the correlation analysis described in Sect. 3.4,

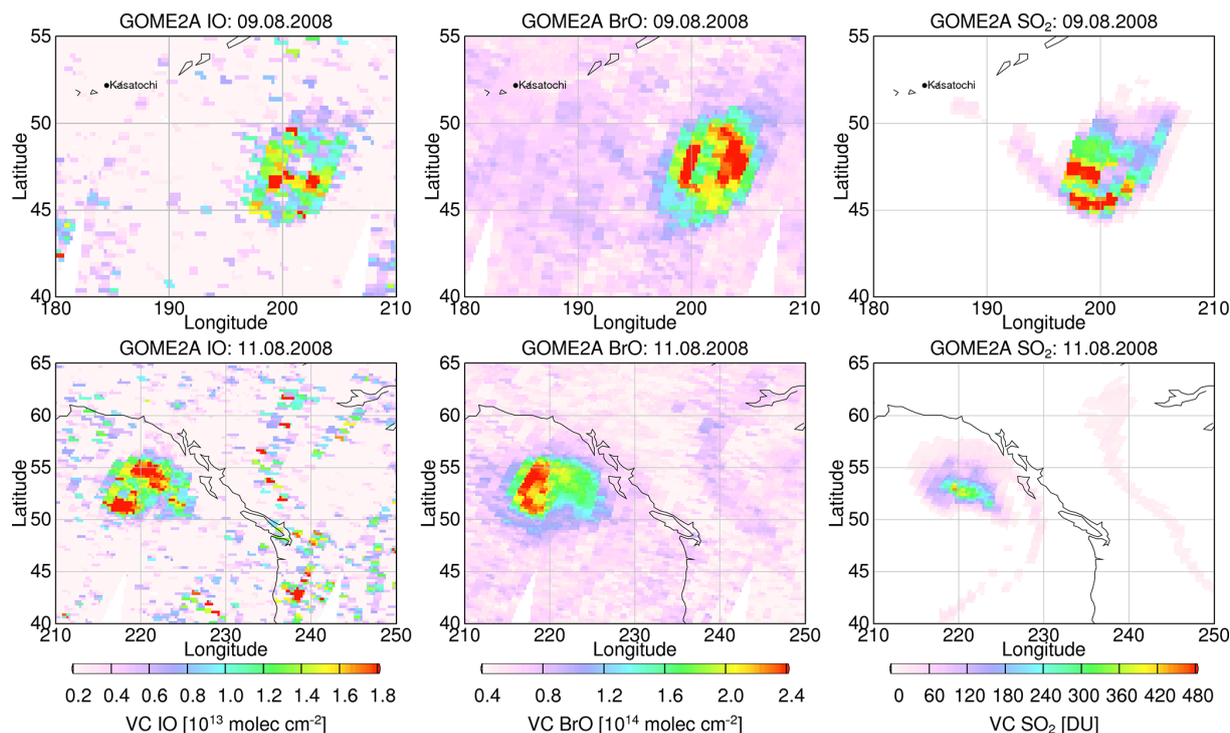


Figure 5. Close-up of the volcanic plumes of IO (left), BrO (centre) and SO₂ (right) on the days 9 August (top) and 11 August (bottom), 2008. While the plume extent and shape are similar, differences in the spatial distribution patterns are visible.

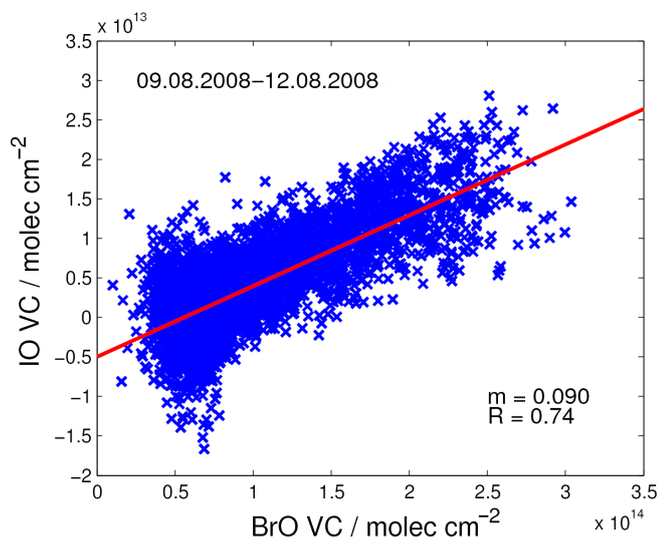


Figure 6. Scatterplot of IO vs. BrO column amounts. Data from the 4 days from 9 to 12 August 2008 is included.

for each day those measurements that fall into a rectangular area enclosing the volcanic plume are used. The slope for all data of IO vs. BrO columns is 0.09, with a correlation coefficient of 0.74. This observation is consistent with findings by Aiuppa et al. (2009), who estimate a 1 order of magnitude difference between the volcanic abundances of the two

halogen species iodine and bromine. In addition, Pyle and Mather (2009) estimate the annual fluxes of volcanic HBr and HI to be on the order of 5–15 Gg yr⁻¹ for HBr and 0.5–2 Gg yr⁻¹ for HI, respectively. The uncertainties in emission fluxes given by the latter study are rather large, but the results for the halogen flux ratios agree with the present satellite observations within their error bars.

The interesting point is that the seawater abundance yields a ratio of Br / I of 15 000, i.e. a 4 order of magnitude difference between I and Br. By considering singly the IO and BrO observations, the number ratio between iodine and bromine atoms is thus enhanced by about 3 orders of magnitude in the volcanic plume as compared to seawater. Modelling of the halogen chemistry within the volcanic plume would be required to calculate the iodine and bromine amounts from the IO and BrO column observations. These estimates are performed under the given restrictions, and the present observations of volcanic emissions of IO shall encourage including the chemistry of iodine and iodine oxides in volcanic halogen chemistry modelling in the future. Other studies have shown that while the Cl vs. Br ratio for volcanic condensates is in agreement with the seawater ratio of around 650 (Gerlach, 2004; Aiuppa et al., 2005), the ratio of Cl vs. I is about 2 orders of magnitude lower in volcanic plumes than in seawater (Honda et al., 1966; Honda, 1970; Snyder and Fehnl, 2002; Aiuppa et al., 2005). Consequently, an enhancement of io-

dine species takes place in the processes which determine the release of halogens from volcanic activity.

Explanations for the observed enhancement of iodine in volcanic emissions relative to seawater are connected to the magma composition of the specific volcano. As Kasatochi is an oceanic arc volcano, marine sediments which are carried into the Earth mantle at the subduction zone directly influence the composition of the volcanic material. Marine sediments in turn are enriched in iodine compounds from organic material (Muramatsu and Wedepohl, 1998). In addition, volcanic emissions are influenced by the composition of the melts and fluids in the volcanic chamber. Results of hydrothermal experiments were used to analyse the compositions of hydrous fluids and silicate melts with respect to the different halogens (Bureau et al., 2000). It was found that the partition coefficient between fluid and melt is clearly larger for iodine than for bromine and chlorine. The partitioning into the fluid phase is therefore stronger for iodine than for bromine, which is again stronger than for chlorine. Consequently, volcanic emissions to the atmosphere are expected to be enhanced in iodine relative to the other halogens.

It is also of interest to study the temporal evolution of the observed IO and BrO column amounts within the plume. The observations on the first day after the eruption, however, may be influenced by dust and clouds accompanying the eruption, especially close to the volcano (Theys et al., 2009). Consequently, trace gas amounts could be larger than quantified by the spectroscopic observations. Comparison of the temporal behaviour of IO and BrO shows that their evolution is similar, with maximum integral amounts detected 1 to 3 days after the eruption (cf. Fig. 4). BrO reaches its highest values (around 5×10^{29} molecules) earlier than IO. The different chemical pathways and time constants for IO and BrO production and destruction also influence the temporal variation of the I/Br ratio. However, the temporal changes between 9 and 12 August 2008 are close to the limit of being significant. Considering only the 1σ standard deviation of IO on the order of 2.5×10^{12} molec cm^{-2} , the uncertainty on the integrated IO molecule number within the volcanic plume lies between 0.8 and 1.2×10^{28} , using the plume areas from 9 and 11 August, respectively. As a result, details in the temporal evolution need to be interpreted with care. Overall, it is interesting to note that the ratio of observed IO and BrO (Fig. 4) shows little change during the aging of the plume within the 5 analysed days. This observation may imply that higher iodine oxides, which are formed more rapidly at larger IO concentrations (cf. estimation of IO mixing ratios below), are photochemically labile inside the volcanic plume. Thereby, the IO may persist in the plume for a longer time period than what would be expected from the atmospheric lifetime of IO. The evolution of iodine species in the volcanic plume may be further affected by particle formation and heterogeneous reactions. Murphy and Thomson (2000) measured enhanced iodine content in aerosols in the upper troposphere and lower stratosphere (UTLS) region. This finding has two further im-

plications. Particles may serve as a sink for iodine, reducing the availability of reactive iodine, and, on the other hand, they may provide pathways for heterogeneous reactions from which reactive iodine compounds may be released again.

The spatial distributions of IO, BrO and SO₂ are described in Sect. 3.4, and some differences between the three species are observed. The chemical pathways of iodine and bromine within the plume are probably not independent from each other. Formation and loss processes may interfere with each other. Although the rate coefficients for the reactions of I and Br with O₃ are similar, the smaller expected concentrations of I than Br imply that the time constant for IO production is larger than that for BrO. As a consequence, large amounts of Br that react with O₃, thereby strongly reducing the O₃ abundance, may prevent the build-up of IO. This results in spatially separated maximum values for the two halogen oxides. The reactions between IO and BrO, as well as self reactions of IO, also impact the spatial distributions and maximum amounts. Furthermore, the time of emission of the precursor substances may differ to some degree. Iodine and bromine have different solubility in volcanic fluids (Aiuppa et al., 2009). For the two halogen species, degassing from the magma may therefore take place at different pressures, i.e. at different depth of the volcanic abyss. In addition, some clear differences between the spatial distributions of the halogen oxides and SO₂ are found. In general, the comparison between the trace gas spatial distributions is interesting because it potentially yields information on the eruption process and chronology. Details of the plume composition and evolution need to be analysed in the future by chemical transport modelling to provide better insight into the complex reactions taking place within the plume.

For an estimate of the impact of volcanic iodine on atmospheric chemistry, the volume mixing ratio is a more relevant quantity than the column amount. For a rough estimate, the vertical plume extent derived by Theys et al. (2009) is used. They determine the major part of the plume to reside between 8 and 12 km altitude. The retrieved integrated number of IO molecules of about 5×10^{28} molecules, for 10 and 11 August 2008, is used as lower limit of the emitted iodine amount. On both days, the plume extends horizontally over 5×10^5 km². Spreading the observed IO homogeneously within the 4 km thick layer and over the entire plume extent, the average VMR would be around 3 pptv at 10 km altitude using US standard atmosphere pressure and temperature values. Certainly, local VMR values will exceed this average VMR due to an inhomogeneous distribution within the plume. Iodine mixing ratios of 3 pptv may have a strong impact on ozone concentrations (Bösch et al., 2003; Saiz-Lopez et al., 2015a) and constitute a large perturbation of stratospheric iodine, which is measured and estimated to be on the sub-pptv level.

Iodine from volcanic eruptions has several possible implications for atmospheric composition. The upper part of the Kasatochi plume may have reached into the lower stratosphere. Consequently, the presented satellite-based observa-

tions of iodine monoxide indicate that volcanic eruptions may have an impact on the iodine concentrations in the upper troposphere and lower stratosphere, at least regionally.

The above estimated IO VMR of 3 pptv in the Kasatochi plume will be diluted with time. Spreading the released trace gas amount over the area of the entire globe decreases the VMR at the given altitude by 3 orders of magnitude as compared to the plume area. Consequently, strong implications for ozone depletion through iodine from a single volcanic eruption are probably mainly regional and restricted in time. Primarily, the lower stratosphere or UTLS region is affected. However, the region impacted by the emitted iodine may be dislocated from the erupting volcano due to the quickly moving volcanic plume covering distances of typically around several hundred km per day.

Due to the larger chain length for the removal of O₃ by BrO_x and IO_x than by ClO_x, loss of O₃ in the stratosphere can be significantly impacted by the BrO and IO in addition to ClO released from volcanic eruptions. In this case the lower stratosphere may become most affected. This could impact on ozone hole chemistry when volcanic eruptions enter the polar vortices, an issue recently raised by Solomon et al. (2016).

Background iodine amounts between 0.1 and 0.4 pptv in the free troposphere as observed recently (Puentedura et al., 2012; Dix et al., 2013) are possibly also influenced by volcanic activity. Following a volcanic eruption, the iodine amount will directly influence the local and regional chemistry by reducing the ozone levels. The impact of the ability of volcanic IO to form aerosol condensation nuclei requires further study. In addition, volcanic plumes may be subject to long-range transport and therefore lead to effects also at larger distances.

The Kasatochi eruption was in some respect special as it was a major eruption; the plume altitude was relatively large and bromine amounts were also larger than for other investigated volcanic plumes (Hörmann et al., 2013). IO has not yet been detected for any other eruptions investigated, at least not at the Kasatochi levels. Scaling with the observed bromine amounts, iodine levels for the other eruptions could just be below or around the detection limit of current space-based instruments. Future satellite instruments with finer spatial resolution and improved signal-to-noise ratio may allow the observation and detailed investigation of iodine species in volcanic plumes more frequently.

It is interesting to speculate on the amount of halogens emitted to the atmosphere from past major eruptions which have severely impacted atmospheric composition prior to halogen observations from space. For the Pinatubo eruption in 1991, for example, a total mass of about 20 Tg of SO₂ was emitted. The eruption injected gases and aerosols up to 25–30 km altitude, i.e. around the maximum stratospheric ozone mixing ratio. In relative terms, the IO VMR will be increased at these high altitudes due to much lower air density as compared to the Kasatochi estimates. Assuming a similar magma

composition as that of Kasatochi, i.e. similar halogen to sulfur ratios, an amount of around 100 t of IO as well as 1 kt of BrO could have been emitted into the stratosphere from Pinatubo, with corresponding impact on stratospheric chemistry over extended horizontal distances and periods. A detailed assessment again requires better knowledge and studies of the loss of iodine and bromine into the stratospheric aerosol.

5 Summary and conclusions

Following the major eruption of the Kasatochi volcano in August 2008, iodine monoxide is observed by satellite in the volcanic plume for several days. This is the first experimental evidence of IO emitted from a volcanic eruption. The satellite sensors SCIAMACHY and GOME-2A both detect slant column amounts of IO above 4×10^{13} molec cm⁻² in the volcanic plume for several days following the Kasatochi eruption. Maximum vertical columns above 2×10^{13} molec cm⁻² are derived. The presented observations also represent the first reported retrievals of IO from measurements of the GOME-2A instrument. In comparison to tropospheric IO observations in polar and mid-latitudinal regions, the observed column amounts are large, reducing the uncertainties and facilitating analysis of individual measurements. The IO data in the plume shows good fitting quality with fitting errors around 6 % for SCIAMACHY and below 15 % for GOME-2A retrievals.

Overall, the IO enhancements coincide in space with previously published observations of BrO and SO₂. While the plumes of IO, BrO and SO₂ are roughly found in the same area with similar shape, the maximum amounts of the individual species, however, do not always coincide. Differences between IO and BrO are smaller than those between the halogens and SO₂. The emission chronology as well as chemical conversions are presumably individual for the three compounds and could probably lead to the observed differences in spatial distributions.

Correlating all observations of IO and BrO between 9 to 12 August 2008 yields a slope of 0.09, i.e. IO amounts are about 1 order of magnitude smaller than those of BrO. Judging from the IO and BrO column amounts alone, this volcanic ratio indicates a 3 order of magnitude difference with respect to the seawater ratio between iodine and bromine, in agreement with previous filter measurements of volcanic samples at arc volcanos. For this relative enhancement of iodine, two reasons play a role. Iodine shows a stronger preference than bromine for partitioning into volcanic fluid than melt in the volcanic chamber located underneath the volcano. This relative partitioning between fluid and melt determines the gas-phase composition of an eruption plume. In addition, iodine-enriched marine sediments are carried into the Earth's mantle in the subduction zone and directly influence the composition of the magma.

An integration of the observed IO amount within the emission plume results in a large mass of around 10 t (4 to 12 t) of IO emitted from the volcano. By comparing the integrated numbers of IO and BrO molecules found within the volcanic plume, the BrO / IO number ratio ranges between 6.7 and 10.0, while the BrO / IO mass ratio lies between 4.0 and 6.7. Together with the knowledge that the Kasatochi BrO plume predominantly reached the altitude between 8 and 12 km, it can be concluded that a substantial input of iodine to the lower stratosphere, UTLS and free troposphere has taken place following the Kasatochi eruption. If the IO amount is homogeneously spread over the plume area and within the main 4 km thick vertical layer, a VMR of 3 pptv at an altitude of 10 km results. The local VMR can be even higher due to inhomogeneous distribution in the volcanic plume. Iodine volume mixing ratios of around 3 pptv may have substantial impact on atmospheric composition, e.g. through regionally reducing the ozone concentrations.

The investigation of past and future volcanic eruptions with respect to their IO content and impact on tropospheric and stratospheric chemistry is subject to further work and will be facilitated by improved satellite instrumentation in the future.

Data availability. Satellite trace gas column data from SCIAMACHY and GOME-2A observations can be obtained from the authors upon request.

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

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