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Advances in Bromine Speciation in Volcanic Plumes

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Volcanoes are a significant halogen source to the atmosphere. After water, carbon dioxide and sulfur compounds, halogens are often the most abundant gases in volcanic plumes. In the past, less attention was given to the heavy halogens bromine and iodine. However, the discovery of bromine monoxide (BrO) in volcanic plumes led to new interest especially in volcanic bromine chemistry and its impact on atmospheric processes. The BrO detection came along with advances in volcanic remote sensing techniques, in particular, robust DOAS applications and the possibility of continuous measurements by automated instruments located at safe distances from the volcano. As one of the consequences, the volcanic community developed an increased interest to use BrO/SO₂ ratios as a potential tracer of volcanic activity. BrO is a secondary volcanic gas, but the only bromine species in volcanic plumes, which has been measured by remote sensing techniques today. For a better understanding on bromine chemistry in volcanic plumes and to gain information on the original amount of emitted bromine by only measuring BrO, additional techniques were developed (alkaline traps, diffusion denuders) and adapted for drone-based sampling to determine further gaseous bromine species (i.e. Br₂, HBr, HOBr, interhalogens) at various plume ages. Additionally models of plume-atmospheric chemistry were developed to help the interpretation of field-measurements. Model studies simulating plume conditions indicated that a complex atmospheric chemistry mechanism transforms emitted HBr into BrO and other reactive bromine species such as BrOH, Br₂, BrCl, BrONO₂ or BrNO₂. To reproduce the very rapid formation of BrO observed in volcanic plumes, the volcanic emission input to the (low-temperature) plume chemistry models also needs to consider the high-temperature near-vent plume conditions, as represented by thermodynamic models. The formation of BrO and other reactive bromine species depend not only on the amount of bromine emitted but also on plume mixing processes, relative humidity, and aerosol particle acidity. However, uncertainties remain in the validation of the plume chemistry models by a lack of field-measurements. This review provides a comprehensive summary on

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volcanic bromine data of the last 15 years achieved from established and cutting edge measurement techniques as well as their treatment and interpretation in recent model experiments. It points out controversially discussed relation of bromine degassing to volcanic activity and puts a light on remaining uncertainties.

Keywords: volcanic halogen emissions, volcanic plumes, plume chemistry, bromine explosion, bromine speciation, gas monitoring, troposphere

INTRODUCTION

Volcanic activity is responsible for the formation of a permanent atmosphere on Earth, and volcanic emissions continue to influence atmospheric composition and thereby climate. Thus, even though volcanic activity has acted to promote the development of life it also induces risks and challenges for the terrestrial biosphere (Shaw, 2008). Measurements of volcanic emissions (gas flux, composition) made at the surface provide an indirect means to explore the inaccessible interior of volcanoes and their complex magmatic systems. They can thereby contribute to volcano hazard monitoring and eruption forecasting.

Knowledge about the chemistry occurring in the volcanic plume is required when making inferences about volcanic activity from volcanic gas measurements, particularly in the more chemically-aged downwind plume. Volcanic sulfur emissions become oxidized in the atmosphere to radiatively-active sulfate particles that can cause climate cooling in particular following large eruptions that injected gas into the stratosphere, e.g., Pinatubo 1991. Volcanic sulfate particles in the stratosphere also act as surfaces to activate halogens causing ozone layer depletion (Brasseur and Granier, 1992). More recently, the potentially chemistry-climate importance of volcanic halogen emissions has started to be of increased interest again (von Glasow et al., 2009; Kutterolf et al., 2013, 2015; Cadoux et al., 2015, 2018; Vidal et al., 2016; Klobas et al., 2017). Some eruptions to the stratosphere have been observed to co-inject halogens alongside sulfur (Theys et al., 2009; Hörmann et al., 2013; Carn et al., 2016), leading to volcanic halogen impact on stratospheric ozone and NO_X (Lurton et al., 2018). Besides remote sensing observations have been done via in situ sampling (Rose et al., 2006) and analysis of ice cores (Zdanowicz et al., 1999; De Angelis et al., 2003). Volcanic halogen emissions to the troposphere, particularly bromine (but also chlorine and iodine) can become activated through plume atmospheric chemistry that results in tropospheric ozone, NO_X, HO_X depletion and may convert mercury to a more easily deposited form (Bobrowski et al., 2007; Roberts et al., 2009; von Glasow, 2010; Jourdain et al., 2016). The plume chemical activation of volcanic halogen emissions, especially bromine, is the focus of this review.

In general, volcanic gas emissions consist of: 50-90% H₂O, 1-40% CO₂, 1-25% SO₂, and trace species such as ca. 0.5% H₂ and 0.03% CO (Gerlach, 2004; Textor et al., 2004). For arc volcanoes, halogen emissions are represented by about 0.84% HCl, 0.061% HF, 0.0025% HBr (Gerlach, 2004), with trace-level HI emissions. Global average bromine degassing fluxes from arc

volcanoes are estimated to be 5 ± 15 Gg/year HBr (Pyle and Mather, 2009), based on collated data on Br/S ratios in volcanic emissions combined with assuming an annual arc-related SO₂ flux of 15 Tg/a. However, this may underestimate total volcanic bromine emissions, given the most recent estimate of the global SO₂ flux from passively degassing volcanoes is 23 ± 2 Tg/a (Carn et al., 2017), and the new identification of bromine-rich volcanic sources (e.g., Nyiragongo, Democratic Republic of the Congo, Bobrowski et al., 2015).

Previous reviews on volcanic halogens have focused on petrological approaches (e.g., Devine et al., 1984; Wallace, 2005) and volcanic degassing fluxes of halogens (Pyle and Mather, 2009) or their general cycling and distribution in the earth system (Webster et al., 2018). Platt and Bobrowski (2015) summarized observations of reactive bromine species mainly by spectroscopic technologies. Several reviews have also been published about the origin and impact of volcanic halocarbons as a minor halogen emission (Schwandner et al., 2004, 2013; Frische et al., 2006). Lately developments of new measurement techniques, collection and interpretation of data sets (by both, new and old techniques), and approach by model applications moved forward the knowledge related to bromine transformations in volcanic plumes. To our knowledge no comprehensive summary and overview of this still relatively new and fast advancing field has been done today. Here we conflate all recent approaches to draw an overall image about the state of the art by pointing out commonly confirmed statements, inconsistencies and general knowledge gaps suggesting further work. We focus on inorganic bromine compounds within passively degassing volcanic plumes entering the troposphere, for which the datasets and plume model studies are most comprehensive, but including a brief discussion of other halogens (Cl, I).

This review starts with an overview of existing measurement techniques for the determination of both speciated and total bromine in volcanic plumes. This is followed by a discussion of numerical modeling of plume bromine chemistry. Then a summary of volcanic bromine measurements is presented. An overview is given of the chemical reaction mechanisms that activate bromine (as well as chlorine and iodine) in volcanic plumes, detailing the roles of both low and high-temperature chemistry, photochemical and multi-phase reactions, aerosol/humidity, and impacts and feedbacks of ozone, NO_X, and mercury. This conceptual framework is then used to discuss reported field-studies and trends in the collated volcanic bromine datasets. The review concludes by describing the current knowledge gaps, particularly in relation to the use of bromine as monitoring parameter for forecasting eruptions, and highlights

recent developments that provide new opportunities in this field.

METHODS USED TO INVESTIGATE BROMINE IN VOLCANIC PLUMES

Measurement Techniques

Volcanoes present a challenging context for both instruments and scientists seeking to measure volcanic plume gases in remote, inaccessible areas and during hazardous volcanic eruptions. For continuous and safe observations remote sensing techniques are often the instruments of choice. However, not all species can be measured by remote sensing. Also, *in situ* and direct sampling techniques can enable measurements closer to the emission source and thus capture degassed species at an earlier point of plume age. Making measurements very near-to-source is essential for the case of highly reactive species such as bromine: it allows to investigate as "pure" volcanic signal as possible, opening up the possibility to study the chemical transformation processes as a function of distance (and travel time) downwind, as the volcanic emissions progressively mix with the surrounding atmosphere.

In Situ – Application of Instruments in the Plume

Earlier determinations of volcanic halogens were realized by condensate sampling in volcanic fumaroles (Oana, 1962; Sugiura et al., 1963; Goff and McMurtry, 2000; Amachi et al., 2001) and by the use of passive alkaline traps (Noguchi and Kamiya, 1963; Goff et al., 1998; Witter et al., 2004). Today, actively pumped alkaline traps are typically applied. The main idea is to retain and enrich acid compounds within strong alkaline solutions (e.g., 1-4 M NaOH) when air is pumped through the substrate. Due to their light weight and easy handling, "filter-packs" (FP) are still the most commonly applied alkaline traps to sample volcanic plume gases. Different kinds of filters (cellulose, Nylon, Teflon) have been used with several mainly alkaline coatings (e.g., 1M NaHCO₃) and glycerol to improve the absorption efficiency. Nevertheless, saturation of filters or evaporation can readily occur. Limitations of the sampling efficiency were pointed out especially for saturated filter packs leading to elevated halogen/sulfur ratios due to distinct acidity (e.g., SO₂ and HF) and therefore different retention in the alkaline solution (Martin et al., 2010; Sawyer et al., 2011; Wittmer et al., 2014). To avoid or at least be aware of saturation effects, several filters are usually mounted in series. If the last filter of a pack yields <10% of the total amount of the species of interest, a quantitative trapping of the compounds is assumed (Lazrus et al., 1976; Sedlacek et al., 1984). If particle filters in front of filter packs are installed (i.e., to avoid uptake of aerosol and ash particles), possible interactions of gases with the particle-loaded filter have also to be considered. For volcanic bromine, HBr is assumed to be efficiently collected, but uncertainties remain whether species such as HOBr, Br₂, BrCl, BrNO₂, and BrONO₂ are trapped quantitatively by alkaline impregnated filters (Kitto et al., 1988).

Using these techniques extensive data sets of volcanic plume composition were established, however, in the case of halogens most of them focus on chlorine and fluorine and do not take bromine and iodine into account. One of the early plume studies on heavy halogens was carried out by Aiuppa et al. (2005) who assumed to determine the total molar amount of emitted bromine species (Br_{total}) on alkaline filters since bromine on particulate filters were negligible and no BrO was measurable with DOAS at the same location.

Uncoated filters can be used for the determination of particulate bromine, although few observations have been reported (Sturges and Shaw, 1993; Zelenski et al., 2013). Size-resolved analysis is enabled when cascade impactors are used. Here, the particles are deposited on several filter membranes according to their size. In comparison to alkaline filter sampling packs (sampling volume about 120 L; Aiuppa, 2009), particle analysis typically requires much higher sampling volumes ranging from 2,000 up to 10,000 L (Erebus, Antarctica: Ilyinskaya et al., 2010; about cascade impactors: Marple et al., 1991; Ma and Kim, 2008).

For avoiding saturation during sampling, active liquid alkaline traps were developed, which solve acid gas species in alkaline solutions by bubbling volcanic gases through the solution. To enhance the efficiency in Drechsel bottles (based on gas washing bottles named after Edmund Drechsel, DB) frits are used to decrease bubble size and therefore increase the relative surface of introduced gas samples to the alkaline solution (Liotta et al., 2012; Wittmer et al., 2014). The same aim of increasing the surface between the solution and introduced gas is used by filling a glass tube with glass rings (so-called Raschig rings; Raschig, 1914) and adding alkaline solution which is able to cover all the surface by rotating the tube (the so-called Raschig tube; RT, Levin et al., 1980). The enhanced efficiency allows a higher volume flow-rate that yields in increased sample concentrations and facilitates analyses close to detection limits for a specified sampling time or alternatively sampling time in the field can be reduced, in cases where a reduced detection limit is not necessary. Since it is assumed that the dominant fraction of bromine compounds are acidic and are therefore dissolved in the alkaline sampling solution the results are assumed to represent the Br_{total}. Today, all those alkaline trap samples are usually analyzed by IC and ICP-MS (see overview in Wittmer et al., 2014). An alternative are neutron activation analysis, but often much more expensive and comparison studies showed similar results between the methods (Strellis et al., 1996; Wittmer, 2012). Wittmer et al. (2014) characterized the sampling efficiency of various alkaline traps. In general, good agreements for filter packs, Drechsel bottles, and Raschig Tubes were found.

Alkaline sampling techniques do not allow speciation (i.e., the determination of individual halogen species). Moreover, alkaline traps cannot distinguish between gaseous and particulate phase. To provide a more in-depth halogen-speciation, gas diffusion denuders take advantage of the different diffusion coefficients of gaseous compounds and particles. The tube-shaped denuder systems are applied with a coated inner surface. Gas molecules diffuse to coated denuder walls, which represent an ideal sink for the gas molecules of interest and thus are retained and enriched in the coating. In contrast, due to the much smaller diffusion coefficient, the aerosol particles pass the denuder. Using various coatings, which undergo species specific derivatization of analytes, enables selective analysis of bromine species (Huang and Hoffmann, 2008, 2009; Rüdiger et al., 2017).

In situ techniques applicable for bromine determination are summarized in **Table 1**. Beside ground-based applications, drone-based sampling units proved to be a valuable tool for *in situ* volcanic gas measurements of difficult-to-access crater emissions as well as aged plumes downwind from the emission source that have become lofted away from the surface (e.g., McGonigle et al., 2008; Rüdiger et al., 2018b).

Remote Sensing

Especially during explosive episodes, remote sensing techniques become an indispensable tool for safe monitoring of volcanoes. Optical remote sensing techniques, using spectroscopic approaches from ultra-violet to thermal infrared, measure integrated gas concentrations through cross-sections of the plume. An overview of typical applied remote sensing techniques is given e.g., in McGonigle and Oppenheimer (2003) and Platt et al. (2018).

Restrictions for remote sensing techniques might occur due to high and/or variable atmospheric backgrounds for some gases (mainly for H_2O and CO_2) and since remote sensing techniques are sensitive for species with differential absorption structures, accessible compounds are limited to the wavelength range of the particular spectrometer and light source. Fourier transform infrared (FTIR) technique has been used for spectroscopic halogen observations of HCl and HF (e.g., Mori and Notsu, 1997; Oppenheimer et al., 1998). However, due to "small" absorption cross-sections, low abundances and limited spectral resolution, portable FTIR instruments have not been able to detect HBr.

In volcanic environments, BrO is the only bromine species that has been determined by remote sensing. Satellites, portable and continuously installed Differential Optical Absorption Spectroscopy (DOAS) instruments, are used to investigate BrO in volcanic plumes. DOAS instruments collect light (from scattered skylight, direct solar light or artificial lamps) that has passed through the volcanic plume. Light collected by a telescope is coupled into a spectrometer often by using an optical fiber and is then dispersed by a grating inside the

TABLE 1 | Overview *in situ* plume bromine sampling techniques.

Sampling techniqueSpeciesPhaseReferences e.g.Alkaline TrapsDrechsel bottleAcidic speciesGas + particleWittmer et al., 2014Raschig TubeAcidic speciesWittmer et al., 2014Coated FilterAcidic speciesAiuppa et al., 2005DenuderCoating: 1,3,5- trimethoxybenzeneHalogen atom with the oxidations state +1 or 0 (e.g., Br2 or BrCl)GasRüdiger et al., 2017ParticleFilterParticulateParticleZelenski et al., 2013Cascade impactorParticulateParticleIlyinskaya et al 2010					
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DenuderCoating: 1,3,5- trimethoxybenzeneHalogen atom with the oxidations state 		Coated Filter	Acidic species		Aiuppa et al., 2005
Particle Filter Particulate Particle Zelenski et al., 2013 Cascade impactor Particulate Particle Ilyinskaya et al 2010	Denuder	Coating: 1,3,5- trimethoxybenzene	Halogen atom with the oxidations state +1 or 0 (e.g., Br ₂ or BrCl)	Gas	Rüdiger et al., 2017
Cascade impactor Particulate Particle Ilyinskaya et al 2010	Particle	Filter	Particulate	Particle	Zelenski et al., 2013
		Cascade impactor	Particulate	Particle	llyinskaya et al. 2010

spectrometer. Signal intensity is quantified via a CCD array or photomultiplier device. The most commonly used instrument are the so-called passive DOAS instruments, which use scattered skylight as a light source. The plume absorption spectrum is compared with an additional spectrum taken outside the plume under nearly identical conditions to correct influences of atmospheric background absorptions and the solar spectral structure (Fraunhofer lines). Today portable Multi-Axis-DOAS (MAX-DOAS) instruments are a widespread instrument in volcanic plume investigations yielding many observations of BrO alongside SO₂. Details are described in e.g., Hönninger et al. (2004), Bobrowski et al. (2007), Bobrowski and Platt (2007), and Gliß et al. (2015). Zenith-sky pointing DOAS instruments can be deployed whilst traversing underneath the plume to yield total column amounts across a transect, which in combination with wind-speed estimates provides total SO₂ and BrO fluxes. Traverses have been undertaken by foot, car, ship, and even by UAVs in inaccessible areas (e.g., McGonigle et al., 2002; Galle et al., 2003; López et al., 2013; Rüdiger et al., 2018b).

Imaging of plumes with SO₂-cameras (see e.g., Mori and Burton, 2006; Kern et al., 2015; McGonigle et al., 2017) enables to spatially map complex plumes at high time resolution (<1 s). Imaging of volcanic plume BrO has been achieved using Imaging DOAS (IDOAS). Spatial resolution is generated by spatial and spectrally resolved vertical columns applying a two dimensional detector and by horizontally scanning gaining the second spatial dimension (Lohberger et al., 2004; Louban et al., 2009). However, the temporal resolution to record a single image by this method is in the order of minutes, much slower than the fast (seconds) plume chemical and physical (dilution) processes. The BrO image obtained is only an average. Ongoing development of Fabry Perot Interferometers (FPI; Kern et al., 2010; Kuhn et al., 2018) may enable fast imaging of BrO in the future with a temporal resolution similar to SO₂ cameras (<1 s).

Beside portable instruments, instrument networks for permanent and automatic monitoring of several volcanoes by DOAS instruments have been created [NOVAC, (Galle et al., 2010); FLAME, (Burton et al., 2009), FLYSPEC, (Businger et al., 2015)]. Typically instruments are installed 5-10 km downwind of the crater observing plumes with an age of about 10-30 min (Galle et al., 2010). Further satellite ultraviolet instruments like GOME-2 (Global Ozone Monitoring Experiment), OMI (Ozone Monitoring Instrument) and the recently launched TROPOMI (Tropospheric Monitoring Instrument) enable BrO (and SO_2) monitoring in the troposphere and stratosphere on a global scale. Main differences between satellite instruments are spatial resolution and overpass time (for details see e.g., Theys et al., 2011; Carn et al., 2013; Hörmann et al., 2013). Systematic analysis of GOME-2 satellite observations of volcanic BrO are presented from volcanic eruptions entering the troposphere (Heue et al., 2011; Rix et al., 2012), and upper troposphere-lower stratosphere region (Theys et al., 2011; Hörmann et al., 2013). Due to the spatial resolution (100's km, reaching 10's km for the new TROPOMI instrument), observations are typically available for somewhat aged eruption plumes. Interpretation of such data is made more complicated because total bromine measurements are often not available and for commonly investigated plume ages >day, SO₂ might not be a reliable conservative plume tracer anymore. We focus our discussion of volcanic bromine speciation on ground based and airborne measurements made on the volcano flank (relatively close to the emission source) by remote sensing (DOAS), *in situ* methods, and associated numerical modeling studies.

Model Application

The volcanic plume is a diverse environment with dynamic variability in temperature, humidity, aerosol loading, and chemistry. Because of this, modelers typically employ separate techniques to describe phenomena at the vent (high-temperatures) and at ambient atmospheric conditions downwind (low temperatures). Numerical models of volcanic plume chemistry have been developed and applied at both high-temperature and low-temperatures. Comparison of model simulations to plume observations provides a wider context to improve our understanding of bromine evolution in the plume and to forecast impacts of bromine chemistry. Calculations identified that volcanic gases undergo high-temperature interactions followed by low-temperature reactions in the downwind plume that has been mixed with ambient air and cooled to ambient temperatures (Gerlach, 2004; Bobrowski et al., 2007; Roberts et al., 2009).

Model Initialization – High-Temperature Near-Vent Conditions, Dilution, and Aerosols

Following the discovery of BrO in a volcanic plume (Bobrowski et al., 2003) thermodynamic equilibrium models were applied at high-temperature to forecast bromine speciation. However, these were not able to simulate bromine monoxide to sulfur dioxide (BrO/SO₂) ratios (10^{-4} mol/mol) observed downwind from the crater (Gerlach, 2004). Rather, the predicted BrO/SO₂ ratios at the vent (10^{-8}) are consistent with the observation that BrO is often not detectable at the crater rim (see also section Dependence on Plume Age–Temporal Evolution).

Numerical models of atmospheric chemistry were then adapted to volcanic plume environments and/or newly developed and applied to forecast the plume halogen chemistry at ambient-temperatures. These model simulations were able to reproduce measurement results (e.g., BrO/SO₂ downwind from the crater). Three key aspects to the initialization of atmospheric chemistry models of volcanic plume halogen chemistry are: (1) the high-temperature near-vent conditions, (2) dilution (that controls entrainment of background air), and (3) volcanic aerosols.

For near vent conditions, it is assumed that near the emission source volcanic gases mix with atmospheric gases and undergo high-temperature chemistry. If reaction kinetics proceed faster than plume dilution a thermodynamic equilibrium may be assumed. This equilibrium systems depend on pressure, temperature (magmatic and ambient) and oxygen fugacity (P-T-fO₂) conditions and have been applied to calculate the composition of the "effective source region" (Gerlach and Nordlie, 1975; Gerlach, 2004; Martin et al., 2006; Bobrowski et al., 2007). One of the most common used thermodynamic equilibrium models is HSC Chemistry (Outokumpu Tech, Finland). A free-energy minimization algorithm is used to

calculate molecular equilibrium compositions of gas mixtures. The estimated model input data are based on crater rim or fumarole measurements and include temperature and pressure as well as a starting elemental plume composition. This is based on mixing the observed composition of volcanic gases with air (N₂, O₂, Ar), for which the proportion of mixing between atmospheric and magmatic gases (VA:VM) must be assumed. Thermodynamic modeling of the hot plume-air mixing predicts that a $V_A:V_M > 0$ ambient air oxidizes gases like H₂S and H₂ and produces small quantities of reactive halogen species such as Br, BrO, Cl and ClO, as well as relatively high quantities of HO_X (hydrogen oxide radicals) and NO_X (NO, NO₂ and combinations) (Gerlach, 2004; Martin et al., 2006). Thermodynamic modeling also predicts the high-temperature formation of SO₃ that is a precursor to sulfate aerosols (see also section Bromine Explosion). The output from the thermodynamic models is then used as input ("the effective source region") to atmospheric chemistry models that simulate halogen processes in the downwind plume at ambient temperatures. The high-temperature composition of the near vent plume is crucial for the initialization of low-temperature kinetic models: the radicals generated accelerating the onset ("kick-start") of halogen cycling at ambient temperatures, hence can affect plume evolution downwind from the crater.

However, the high-temperature near-vent conditions ("effective source region") remain a source of uncertainty in the initialization of low-temperature models of the plume halogen chemistry. The high-temperature modeling involves several unverified assumptions since high-temperature conditions are hard to constrain by direct measurements. For example, VA:VM in the near-vent plume exerts a strong influence on the predicted composition but is somewhat arbitrarily chosen: early studies used 40:60, and later on 15:85, 10:90 and 5:95. For some processes assumptions of the thermodynamic equilibrium seem to be inappropriate [e.g., re-equilibration of H₂S (Martin et al., 2009), H₂ (Aiuppa et al., 2011) or formation of NO_X from background N2 that is likely kinetics limited due to the high bond strength for N₂ (Martin et al., 2012a)]. At $V_A:V_M = 40:60$, high levels of Cl radicals are predicted to form, that subsequently act to rapidly destroy H₂S in the atmospheric chemistry models. However, measurements by Aiuppa et al. (2007) did not show a decrease of H₂S as a function of distance to the emission source (0.1-1 km). Measurements of HO₂ and OH are very rare even though they play an important role in the atmospheric chemistry/halogen reaction cycles. OH radicals can be measured by very high-resolution active DOAS instruments (Platt et al., 1988) or by laser-induced fluorescence (LIF). Indirect detection of HO₂ is possible by titrating HO₂ with NO to form OH, which can be detected by LIF. Subtraction of the measured OH background then yields HO₂ (Hard et al., 1984; Brune et al., 1995). However, the power requirements and large size of such measurement techniques for OH and HO2 generally limit their use on the volcano, restricting their potential applicability to aircraft measurements of the (typically more aged) plume. One of those examples is the airplane measurements of the eruptive plume of Mt Hekla, Iceland in 2000 (Hunton et al., 2005). No measurements are available, to the authors' knowledge, in the proximity of volcanic emission sources.

When the plume reaches the crater rim, more ambient air is entrained and the plume cools to ambient temperatures. The background air contains oxidants (HO_X, NO_X, O₃) whose entrainment into the plume affects the plume chemistry. Therefore, important controls on the halogen evolution in the downwind plume are the rate or extent of dilution and nature of the background atmosphere. Beside latitude, longitude and altitude dependent parameters (e.g., temperature, time of the day) more polluted and NO_X-rich atmosphere may need to be considered for some locations. In contrast to emissions entering cleaner free tropospheric conditions like for many high altitude volcanoes in the Andes, impacts of other environmental emissions on the plume chemistry have to be considered close to large cities or for instance, for volcanic emissions entering the planetary boundary layer like it is the case for Stromboli in the Mediterranean, or Masaya in Nicaragua. Dilution of the downwind plume can be represented by simple dispersion parameterizations such as Pasquill-Gifford or can be adapted to reproduce reported plume dilution rates.

Aerosol particles in volcanic plumes are highly acidic, often sulfate-rich, and include both primary aerosol (directly emitted) and secondary aerosols (e.g., formed from the atmospheric oxidation of volcanic SO₂). Volcanic aerosols can be measured by in situ offline time-averaged sampling using filter packs (Allen et al., 2002; Mather et al., 2003) or in situ real-time sampling by optical particle counters (Allen et al., 2006) and by remote sensing with sun-photometers (Watson and Oppenheimer, 2000; Roberts et al., 2018). The role of aerosol in partitioning between the gas phase and particles is considered by equilibrium models AIM (Aerosol Inorganic Model; Wexler and Clegg, 2002; Martin et al., 2012b) or MOCCA (Model of Chemistry in Clouds and Aerosols in MISTRA; Sander and Crutzen, 1996; Vogt et al., 1996; von Glasow, 2000). Aerosols catalyze heterogeneous reactions that have a crucial impact on plume halogen chemistry (see section Bromine Explosion). This is demonstrated by model simulations that can only reproduce downwind observations of BrO/SO2 when volcanic aerosol is included. Reactions of halogens on volcanic aerosol are thus a key element in the atmospheric models of volcanic plume halogen chemistry outlined below.

Modeling Plume Chemistry – Low-Temperature Kinetic Models

In the downwind plume low-temperature kinetic models investigate plume chemistry. Mainly two low-temperature kinetic 0/1D models have been developed: MISTRA (simulates an advected column of air) and PlumeChem (in expanding box or multi-grid box modes). Recently also the regional 3D model CCAT-BRAMS was applied to study volcanic halogen chemistry. All these models take into account detailed atmospheric chemistry schemes including reactive halogens (bromine and chlorine) with gas-phase photolytic and gas-aerosol reactions. These kinetic models of volcanic plume chemistry at ambient temperatures used an "effective emissions source" (Bobrowski et al., 2007) in their inputs based on thermodynamic equilibrium models (HSC) representations of the near-vent high-temperature chemistry.

The numerical model MISTRA simulates a column of atmosphere that is divided into multiple layers. This column is moving over a volcano where the plume is emitted as a puff into few adjacent model layers at a given height. This column of air is then advected downwind according to an assumed wind speed (Bobrowski et al., 2007), whilst the plume puff disperses vertically in the column. Horizontal dilution is taken into account with a simple parameterization, assuming that the horizontal evolution of the plume corresponds to a Gaussian plume. MISTRA was originally developed for microphysics in the marine boundary layer including a hydrodynamic part, a radiation code and a parameterization of the turbulence by means of a first-order. The turbulence closure was based on the model of Mellor and Yamada (1982) and the meteorological core was developed by Bott et al. (1996). von Glasow (2000) extended MISTRA with a module that describes chemical reactions of the gas phase, aerosol particles and cloud droplets (this extension was based on MOCCA). So far, collision and coalescence of aerosol and new particle formation are not considered in the model. The onedimensional MISTRA has been applied in volcanic plumes with focus on halogen mechanisms by Aiuppa et al. (2007), Bobrowski et al. (2007), von Glasow (2010), Bobrowski et al. (2015), and Surl et al. (2015).

PlumeChem is a 0D/1D box model that also uses a Lagrangian-type approach to follow a volcanic plume puff with time or distance downwind according to an assumed wind-speed (Roberts et al., 2009, 2014b, 2018; Kelly et al., 2013). The model simulates chemical reactions in the plume as it disperses and entrains background air. This approach is similar to that applied to modeling the chemistry of shipplumes (Song, 2003). In the single grid box mode, plume dispersion is simulated by a single grid box (an ellipse) that expands as background atmospheric air is entrained into it. In the multiple grid box mode, the plume is described by a series of chemical boxes in a row, i.e., plume chemistry is spatially resolved horizontally. Plume dispersion is represented in the vertical direction by box-growth, whereas it is simulated in the horizontal direction by mixing between adjacent boxes. The plume dispersion is modeled assuming a Gaussian plume dispersion and parameters are given by Pasquill-Gifford parameterizations, or can be user-defined to approximate actual observed dispersion of plume tracers (Kelly et al., 2013). The PlumeChem model includes detailed volcanic plume halogen atmospheric chemistry similar to MISTRA, but the gas-aerosol reactions as treated using reactive uptake coefficients on an aerosol surface area rather than directly simulating the aqueousphase chemistry. This reduces the computational requirements of the model, and is the approach also used in 3D models as discussed below. Considered model parameters are the chemical composition of model initialization, the aerosol loading and plume dispersion.

Recently a non-hydrostatic mesoscale atmospheric chemistry model, CCATT-BRAMS, was developed to simulate volcanic plume halogen chemistry and regional impacts in a Eulerian-type study (Jourdain et al., 2016). The BRAMS model has been used for studies of planetary boundary layer, operational weather forecasting and climate and includes parameterizations of physical processes such as surface–air exchanges, turbulence, convection, radiation and cloud microphysics that are particularly well adapted for tropics. Coupling to CCATT enables the transport, chemical evolution, emission and deposition of molecules and aerosol particles (Freitas et al., 2009; Longo et al., 2013). The CCATT preprocessor was adapted to include volcanic emissions, and the chemistry scheme developed to include volcanic halogens. Two-way nested grids are used to capture plume processes at the small-scale (<1 km) with larger grids simulating the tropospheric impacts of the dispersed plume at the regional scale (>1,000 km). A similar development has also been undertaken to include volcanic halogens in the WRFChem model (Surl et al., 2014).

First steps have also been made to introduce volcanic halogens and their tropospheric plume chemistry into global models. A major issue is the coarse resolution of global models (typically 100–200 km). This causes the volcanic emissions to be too rapidly diluted compared to reality. Global models, therefore, cannot resolve the volcanic plume halogen processes that occur at much smaller scales (km), particularly in young plumes/near-to-source. Grellier et al. (2014) proposed a sub-grid-scale parametrization approach to represent the near-source halogen chemistry plume processes in a global tropospheric model. Future work should further develop the volcanic plume halogen chemistry in such approaches.

The comparison of the currently available model results with one another and with measured observations is usually difficult because starting conditions vary between models due to consideration of different data arrays and volcanoes in model applications. In general, total emissions and their composition may vary immensely over just a few days or between different volcanoes. Furthermore, measurement data sets at different distances downwind are usually not taken simultaneously. The model parameter space is vast. Sensitivity studies show how variability in initial plume compositions (e.g., variable HBr emissions, plume composition, mixing or high-temperature interactions of the young plume in the source region) affect downwind BrO formation (e.g., Roberts et al., 2014b).

CURRENT UNDERSTANDING

Overview of BrO/SO₂ and Br/S Field-Measurements

Upon the discovery of BrO in the volcanic plume of Soufriere Hills, Montserrat (Bobrowski et al., 2003) detection of BrO in many volcanic plumes globally shows that BrO formation is a common process in volcanic plumes including different settings of arc, rift and hotspot volcanoes as shown below.

Nevertheless, data sets are still very limited regarding the geographical distribution and temporal variability of bromine emissions. **Figure 1** and the **Supplementary Table 1** summarize all available volcanic plume bromine data since 2003. Earlier results are already summarized elsewhere (e.g., Gerlach, 2004). **Figure 1** gives an overview on particulate and gaseous bromine to sulfur (Br/S) and BrO/SO₂ ratios, across different volcanological settings (arc, hotspot and rift).

Bromine species are typically reported relative to sulfur, to enable distinction between plume chemistry and dilution effects. Assuming slow in-plume oxidation, SO₂ will act as a plume tracer on the relatively short observation timescales (typically minutes to hours) of measurements on the volcano flank. Indeed, volcanic plume SO₂ can often be considered to be conserved over the short distance between "source" and "downwind" observation points on the volcano flank because the acidity of the volcanic aerosol in the plume, prevents the SO₂ uptake, and because SO₂ concentrations are in excess of most oxidants (McGonigle, 2004; von Glasow et al., 2009; Galeazzo et al., 2018). Variations in bromine/sulfur observed at just one fixed measurement point might be a result from changes in emission composition. But the assumption of SO₂ as a plume tracer may not always be valid for measurements of very dilute or aged plumes (in the order of days), e.g., as may be detected by satellites.

The Br/S emissions in alkaline-trap samples are assumed to represent total bromine, Br_{total} . Figure 1 shows that Br/S values (displayed as blue circles) are generally higher than BrO/SO₂ ratios (red crosses) measured by DOAS instruments. This is consistent with the partial conversion of emitted bromine into BrO, Particulate measurements tend to report higher Br/S values (green dots) in comparison to gas ratios or gas-particle mixtures. Increased Br/S ratios in particles may demonstrate enhanced bromine partitioning into particles compared to sulfur, or unusually low sulfate for the few volcanoes studied e.g., Erebus, Antarctica (Ilyinskaya et al., 2010).

Several factors may explain the variability in **Figure 1**, including:

- Degassing derived from different magma types and volcanic (tectonic) environments e.g., deep or shallower sources associated with melt generation and degassing systems or crustal fluids (Pyle and Mather, 2009; Mather et al., 2012) as discussed below.
- Different states of volcanic activity such as eruptive or quiescent degassing (section Forecasting–Correlating Volcanic Plume Composition to Volcanic Activity).
- Different distances to the source and therefore different extents to which bromine has been chemically transformed into BrO as a function of plume age (section BrO Evolution in Volcanic Plumes).
- Also, processes such as scrubbing of water-soluble compounds may impact Br/S (Symonds et al., 2001; Textor, 2003).
- Uncertainties in individual measuring techniques (section Measurement Techniques), as discussed below.

Considering subaerial actively degassing volcanoes, arc volcanoes are the most abundant tectonic setting and therefore provide the majority of data. The volcanic Br/S shows a very large variation over five orders of magnitude. This is probably primarily due to variation in bromine abundance of the arc volcano plumes.

Hotspot volcanoes have typically been described as halogenpoor (e.g., Pyle and Mather, 2009). However, we find that the available hotspot volcanoes fall in the same range of the data available for arc zones (**Figure 1**). Note that the representation of the samples formerly and still up to now is not statistically relevant. Hawaii falls at the very low end (about 0.3×10^{-4})



measurement uncertainties (see section Overview of BrO/SO₂ and Br/S Field-Measurements).

but Mt Etna, Sicily, Italy is at the high end of Br/S ratios (up to 0.05). There may be influence of recycled oceanic crust as a halogens source in some hotspot cases, e.g., Eyjafjallajökull, Iceland (Moune et al., 2012).

Intriguingly, the few samples taken from rift volcanoes tend to show lower BrO/SO₂ ratios, but no clear differences for Br/S ratios in comparison to arc and hotspot volcanoes.

Within the arc volcano category, the bromine abundances of southern American volcanoes seem to lay at the lower end of the Br/S range. Maximal BrO/SO₂ values for South American volcanoes are below the mean of other available arc volcanic values, with the exception of Tungurahua (max. 1.7×10^{-4}). Lowest global Br/S values since 2003 were found for Copahue's plume (0.20 $\times 10^{-4}$ and 0.29 $\times 10^{-4}$). Although for Japanese arc volcanoes Gerlach (2004) cited also low Br/S ratios (Tokachi 0.25 $\times 10^{-4}$, Kudryavy 7.8 $\times 10^{-4}$), BrO/SO₂ obervations by Lee et al. (2005) at Sakurajima are a bit in contrast to those earlier

determined values in Japan. However, some caution must be applied to the comparisons regarding the methods, because most of the observations of the South American volcanoes are groundbased, in contrast, many observations of the Kamchatka arc (Kizimen, Benzymianny, Kliuchevskoi) were gained by satellite measurements of potentially more aged plume.

Uncertainties in individual measuring techniques were discussed in section Measurement Techniques. Measurements of trace bromine emissions can be particularly challenging close to the detection limit. Also, high gas concentrations can limit the accuracy of the results e.g., due to saturation of alkaline filter packs (see section *In Situ*–Application of Instruments in the Plume). In **Figure 1** the high Br/S ratios of up to 475 × 10^{-4} measured on Etna, Sicily, Italy in 2005 (Martin et al., 2008) are significantly greater than Br/S observations from January to October 2004 where all data points are below 100 × 10^{-4} (Aiuppa et al., 2005), as well as Br/S on Etna reported

by Oppenheimer et al. (2006) and Wittmer et al. (2014). Whilst change in volcanic activity could be one explanation, oversaturated filter samples might be a conceivable explanation of the exceptional high Br/S ratios by Martin et al. (2008).

Plume Chemistry and the Formation of Reactive Bromine Bromine Explosion

Since observations show BrO is not directly emitted from the volcano (see section Dependence on Plume Age-Temporal Evolution), model studies were undertaken to explain the

measured BrO content and its formation process. Thermodynamic studies predict that halogens are mainly degassed as hydrogen halides, e.g., HBr (Symonds et al., 1994). Gerlach (2004) calculated that BrO in volcanic plumes must originate from the enhancement of reactive halogen species by oxidation of HBr and HCl through autocatalytic cycles involving heterogeneous chemistry. Numerical models of volcanic plume atmospheric chemistry were then developed to explain the measured BrO content and its formation process. Volcanic BrO formation is currently assumed to occur through a bromine reaction cycle, the so-called "bromine explosion," that has been previously identified in the polar and marine boundary layers and above salt plains (Barrie et al., 1988; von Glasow and Crutzen, 2003; Saiz-Lopez and von Glasow, 2012).

The most relevant gas phase reaction of HBr is the formation of Br radicals with OH radicals (R1; von Glasow, 2010).

$$HBr + OH \rightarrow Br + H_2O$$
 (R1)

High-temperature chemistry is suggested to accelerate the start of low-temperature cycles by elevated concentrations of HO_X derived from high-temperature chemistry in the near-vent plume that enhance the initial formation of Br radicals via R1 (Oppenheimer et al., 2006; Martin et al., 2009; Roberts et al., 2009; von Glasow, 2010). Therefore, small amounts of BrO (R2) and subsequent formation of HOBr (R3) are already expected to be formed in the high temperature zone as calculated by thermodynamical equilibrium models.

$$Br + O_3 \rightarrow BrO + O_2$$
 (R2)

$$BrO + HO_2 \rightarrow HOBr + O_2$$
 (R3)

However, those gas phase reactions alone cannot explain the measured abundance of volcanic BrO. Calculations predict that heterogeneous reactions enhance BrO formation compared to gas phase oxidation. Indeed, plume model simulations that do not include volcanic aerosols cannot reproduce the downwind BrO. The key reaction is the reactive uptake of HOBr (reacting with aqueous HBr in the acidic aerosol) to form the reactive halogen Br_2 (R4).

$$HOBr + H_{(aq)}^{+} + Br_{(aq)}^{-} \xrightarrow{aerosol-surface} Br_2 + H_2O_{(aq)}$$
(R4)

Br₂ has low solubility in the aqueous acidic sulfate particles and thus partitions into the gas phase where it is photolyzed into Br radicals (R5).

$$Br_{2(g)} + h\nu \rightarrow 2Br$$
 (R5)

Overall, this repeated sequence R2–R5 each time doubles BrO concentrations and is therefore known as the "bromine explosion" (**Figure 2**). Due to its autocatalytic nature, it can produce a very rapid increase in BrO concentrations.

The sequence can be terminated by the loss of Br due to its conversion back to HBr (R6–R7; von Glasow and Crutzen, 2003; Roberts et al., 2009).

$$Br + HCHO \rightarrow HBr + HCO$$
 (R6)

$$Br + HO_2 \rightarrow HBr + O_2$$
 (R7)

In the following, we discuss the importance of photochemistry, the role of aerosols, the production of HNO₂ (von Glasow, 2010) or HNO₃ (Roberts et al., 2009) predicted by models, the depletion of tropospheric ozone (section Causes and Effects of O₃ Depletion), interhalogen, chlorine and iodine plume chemistry (section Interhalogen, Chlorine, and Iodine Plume Chemistry), and conversion of mercury into a more toxic and easily deposited form (von Glasow, 2010; von Glasow et al., 2015) in relation to reactive bromine formation in volcanic plumes (section Mercury in Volcanic Plumes).

Importance of photochemistry

Photochemistry plays a key role for doubling Br radicals in the autocatalytic cycle of the bromine explosion (R5). BrO measurements made round-the-clock at Masaya volcano, Nicaragua, confirmed that photolytic reactions are necessary for the formation of BrO (Kern et al., 2009). Kern et al. (2009) observed significant BrO during daytime whereas BrO was below the detection limit during night. Other measurements were performed at Mt Etna, Sicily, Italy in the early morning during and shortly after sunrise (Gliß et al., 2015; Butz et al., 2017). The data shows an increase of the BrO/SO₂ ratio within the first hour of increasing sunlight and a constant ratio afterward.

Influence of aerosol or humidity

Regarding volcanic halogen emission estimates, the non-reactive uptake of HCl and HBr onto sulfate particles should be considered as it may overprint magmatic HX/SO₂ gas ratios. Significant uptake could lead to underestimation of volcanic halogen emissions if the particle phase is not taken into account alongside the gas phase (Pyle and Mather, 2009).

Aerosols provide surface area promoting heterogeneous reactions within or on the particles and therefore have a key role in volcanic BrO formation (R4), the cycling, and speciation of bromine in volcanic plumes. The volcanic plume environment is abundant in particles, including both primary and secondary aerosols (see also section Model Initialization–High-Temperature Near-Vent Conditions, Dilution, and Aerosols). Primary sulfate is emitted at up to about 1% (mol/mol) of the volcanic SO₂ emission. One mechanism for primary sulfate formation is the high-temperature formation of SO₃ radicals in the near-vent plume, that upon cooling are converted into H_2SO_4 in the presence of water vapor. Secondary sulfate aerosol particles are formed from the oxidation of SO₂ to H_2SO_4 as the plume disperses into the background



atmosphere. Oxidation can occur by gas-phase reaction with OH, and aqueous-phase reaction with ozone, H_2O_2 , and O_2 with transition metal ions. In volcanic plumes, catalysis by metallic ions might significantly enhance oxidation of SO_2 with O_2 (Galeazzo et al., 2018). H_2SO_4 is highly hygroscopic and can create aqueous $H_2SO_{4(aq)}$ droplets via homogeneous nucleation or condense onto existing accumulation mode particles (Seinfeld et al., 1998; Mather et al., 2006; Martin et al., 2012b; Galeazzo et al., 2018).

The surface area of hygroscopic sulfate-rich particles is a strong function of humidity and temperature conditions. However, as the abundant amounts of aerosol provide a substantial surface area for reactions, the formation of BrO may not be limited by aerosol availability, particularly neardownwind. Bobrowski and Giuffrida's studies (Bobrowski and Giuffrida, 2012) at 6 km downwind did not find influences on the BrO/SO₂ ratio due to seasons, relative humidity, and wind velocities. Although higher relative humidity and aerosol with high sulfuric acid concentrations are proposed to increase the probabilities and rates of halogen reactions, they seem not to be a real limiting factor in volcanic plumes. However, Dinger et al. (2018) identified correlations between observed BrO/SO2 molar ratios and relative humidity (correlation coefficient 33%) by investigating periodic patterns in the BrO/SO₂ ratio of Cotopaxi's volcanic plume.

Nitrogen compounds

There is still a great deal of uncertainty as to whether nitrogen compounds NO_Y (NO_X and oxidation products including organic nitrogen compounds) are emitted or formed in volcanic

emissions and which products are built by the interaction of bromine and nitrogen compounds in volcanic plumes. In polluted environments, NO_X might be present in the atmosphere and enter volcanic plumes upon dilution. Mather et al. (2004a) observed nitric acid at the volcano crater rim but proposed not a direct emission of nitric acid. Rather NO is predicted to be formed thermally by equilibrium models at near vent conditions from atmospheric N₂ and O₂ (Gerlach, 2004; Martin et al., 2006). It is assumed that in the volcanic plume at ambient temperature NO will be rapidly oxidized with O₃ to NO₂ (von Glasow, 2010). Further oxidation pathways (R10–R11 below), or reaction of NO₂ with OH may then produce HNO₃.

von Glasow (2000) suggested a reaction of Br radicals with volcanic NO₂ immediately after plume release of BrNO₂ (R8) and due to photolysis a lifetime of about 150 s (R9). Since large amounts of NO₂ are proposed in the plume the continuous reformation and accumulation of BrNO₂ is assumed. This reduces the amount of bromine available to form BrO.

$$Br + NO_2(+M) \rightarrow BrNO_2(+M)$$
 (R8)

M is a chemically inert collisional partner.

$$BrNO_2 + h\nu \rightarrow Br + NO_2$$
 (R9)

In contrast, Roberts et al. (2014b) suggest the volcanic plume $BrNO_2$ lifetime will be short. Roberts et al. (2009) suggest rather a formation of $BrONO_2$ due to the reaction of BrO with NO_2 (R10). $BrONO_2$ rapidly reacts heterogeneous on volcanic aerosols producing HOBr that may immediately react via R4,

forming Br₂ and nitric acid (R11). Under high aerosol conditions, BrONO₂ can thereby promote the "bromine explosion,"

$$BrO + NO_2(+M) \rightarrow BrONO_2(+M)$$
 (R10)

$$BrONO_2 + H^+_{(aq)} + Br^-_{(aq)} \rightarrow Br_2 + HNO_{3(aq)}$$
 (R11)

However, the importance of these chemical pathways depends on the abundance of NO_X in volcanic plumes. As mentioned above a key question is whether NO_X is generated from volcanoes or simply being entrained into the plume by dilution. Although the formation of NO_X is predicted by equilibrium models, a hightemperature kinetic model study found that the time-scale to produce significant quantities of thermal NO_X formation is too slow in the near-vent plumes (Martin et al., 2012a). However, several observations of elevated NO_X and NO_Y in volcanic plume have been reported: an overview is given in Martin et al. (2012a). For example, aircraft measurements were able to identify volcanogenic HNO3 and HO2NO2 (Oppenheimer et al., 2010). However, this study did not observe any NO_X in the downwind plume. They assume that near-source NO_X was quickly oxidized to HNO3 and HO2NO2. HNO3 has been reported in several volcanic plumes (e.g., Mt Etna, Sicily, Italy; Voigt et al., 2014), Masaya, Nicaragua; Mather et al., 2004b). However, Martin et al. (2010) did not detect any HNO3 at Masaya in 2009. For some eruptions (particularly explosive), volcanicinduced lightning could be a source of NO_X in parts of the plume. For volcanic plumes entering polluted background atmospheres, the entrained air can be a source of NO_X (and NO_Y) as the plume dilutes. Overall there remain uncertainties in the sources of volcanic plume NO_Y, and hence the role of coupled nitrogenbromine chemistry in volcanic plumes and its importance for Br-speciation.

Mercury in volcanic plumes

Mercury is emitted from volcanoes, but is also present in the background atmosphere, predominantly in the inert form Hg(0). It is believed that reactive bromine (as well as chlorine, see section Interhalogen, Chlorine, and Iodine Plume Chemistry) species might promote conversion from relatively inert Hg(0) (emitted as well as atmospheric mercury) into more reactive and toxic oxidized mercury [Hg(II), R12–R13], although observations of volcanic mercury transformations are few and somewhat inconclusive to date. Oxidized mercury is soluble and partitions into aerosol particles so can be more easily deposited near the volcano (R14).

$$Hg^0 + Br (+M) \rightarrow HgBr (+M)$$
 (R12)

$$HgBr + Br (+M) \rightarrow HgBr_2 (+M)$$
 (R13)

$$HgBr_{2(gas)} \leftrightarrow HgBr_{2(aq)}$$
 (R14)

There are uncertainties in mercury chemistry in volcanic plume reactions, notably whether a back-reaction can reform inert Hg(0) from oxidized Hg(II), possibly involving photolysis, SO_2 and particles. Mercury concentrations are generally thought to be at sufficiently trace-levels that the mercury chemistry does not impact overall Br-speciation (Bagnato et al., 2007; von Glasow, 2010).

Causes and Effects of O₃ Depletion

Models predict the depletion of tropospheric ozone destroyed by the reaction sequence R2–R5 each cycle (e.g. Bobrowski et al., 2007; Roberts et al., 2009; von Glasow et al., 2009; Kelly et al., 2013; Surl et al., 2015). In addition, ozone loss occurs through a second cycle between the self-reactions of BrO (R15, R16). This rapid interconversion causes ozone loss in the plume at high halogen concentrations, and so can be particularly important in destroying ozone in the near-downwind (concentrated) volcanic plume.

$$BrO + BrO \rightarrow 2Br + O_2$$
 (R15)

$$BrO + BrO \rightarrow Br_2 + O_2$$
 (R16)

Note also that the reactions R15 and R16 will tend to reduce the abundance of BrO at high halogen concentrations and therefore can bring additional non-linearities to the chemistry of volcanic BrO formation. Determination of ozone in volcanic plumes by UV absorption is difficult due to high SO₂ concentrations (see Table 2). As both O₃ and SO₂ broadly absorb UV radiation, two or more simultaneous observations at different wavelengths are required to deconvolute SO₂ from O₃ by UV absorbance alone. Because many UV absorbance instruments employ a UV line source (frequently Hg vapor, 253.7 nm), additional treatments are needed to correct for SO₂ coabsorption. In those cases to avoid SO₂ interferences a CrO₃ scrubber is necessary to remove SO₂ from the inflow to the instrument (e.g., Schumann et al., 2011; Surl et al., 2015). Dual channel ozone instruments also yield accurate results in the presence of SO₂. These detect ozone by taking the difference between a "measurement" cell and "background" cell (with in-line ozone scrubber), therefore any SO₂ interference cancels out (Vance et al., 2010). In cases of large ozone depletion at low SO₂ the interference may be estimated using field-observations of SO₂ and subtracted in data post-processing (Kelly et al., 2013). Electrochemical concentration cells (ECCs) sense ozone as it reacts with a diluted solution of potassium iodide to produce a weak electrical current proportional to the ozone concentration of the sampled air. A negative cross-sensitivity (observed ozone destruction) on mole-per-mole-basis may be caused due to SO2 (Vömel and Diaz, 2010). However, no interferences in volcanic applications are known with chemiluminescence-based techniques that measure O₃ by sensing light output from the reaction of O3 with ethylene, although an overread of O3 (few %) at high humidity may occur (Vance et al., 2010). Recently an Aeroqual instrument, WO3 semiconductor, was used for ozone measurements, for which the most important cross-sensitivity in volcanic plumes is that of H₂S (Roberts, 2018). Time-averaged ozone measurements have also been made using diffusion tube passive sampling devices (Vance et al., 2010). These are filled with an adsorbing cartridge that has a highly specific irreversible chemical affinity to ozone (e.g., the "ozonolysis" of double bonds to form an aldehyde). In the lab, hydrazine is added to form an acid that can be detected via absorption.

Ozone loss due to heterogeneous reactions in the presence of halogens has already been observed in several tropospheric

TABLE 2 Ozone measurement de	vices and the	ir advantages a	and disadvantages
for application in volcanic plumes.			

instrumentation	Advantage	Disadvantage
UV absorption		High positive SO ₂ cross-sensitivity
UV absorption + CrO ₃ scrubber	No SO ₂ cross-sensitivity	
Dual UV absorption + O ₃ scrubber	No SO ₂ cross-sensitivity	
Electrochemical concentration cell		Negative SO ₂ cross-sensitivity
Chemiluminescence	No known interferences Response time <5s	High humidity causes \mbox{O}_3 overread
Diffusion Tubes	Highly specific	Poor time resolution (passive sampling)
WO ₃ sensor	Small, low cost	H ₂ S cross-sensitivity

TABLE 3 | Overview of sampling campaigns on O₃ depletion in volcanic plumes.

Volcano	Instrumentation	% O ₃ loss	Reference
Sakurajima	Dual UV absorption	n.a.	Lee et al., 2005; Surl et al., 2015
Augustine	Chemiluminescence	0–100, av 60	Vance et al., 2010
Etna	Diffusion Tubes	<70%	
	UV absorption $+$ CrO ₃	15–40	
Eyjafjallajökull	Dual UV absorption $+$ O ₃ scrubber	4–84, av 37	
Erebus	UV absorption	max 35	Oppenheimer et al., 2010
Redoubt	Dual UV absorption $+$ O ₃ scrubber	max 90	Kelly et al., 2013
Etna	$UV + CrO_3$ scrubber	15–45	Surl et al., 2015
Kilauea	WO ₃ sensor	<5%	Roberts, 2018

n.a., not available.

volcanic plumes (see Table 3, for a discussion of halogen impacts on ozone in the stratosphere see section Introduction). Ozone in Kilauea's plumes was reported close to ambient levels in craterrim emissions (Halema "uma"u) and only slightly depleted in the 10 km downwind plume (Pu'u 'O'ō) (Roberts, 2018). This is consistent with Kilauea being a low halogen emitter. Large ozone depletions (several 10's%) were observed in the plumes of higher halogen emitters such as Mt Etna, Sicily, Italy and Mt Redoubt, Alaska (see Table 3). In general, the largest O₃ depletion was observed in the core of the plume and was correlated with volcanic gases (e.g. Kelly et al., 2013), with a maximum deficit of -30 ppbv and an average O₃ deficit reaching -12 ppbv at 20-40 min plume age. That is consistent with model simulations that reproduced ozone depletion in the core of the plume (Kelly et al., 2013). Observational and modeling results suggest that the conversion of volcanic Br emissions into reactive bromine are key controls in the depletion of tropospheric ozone downwind from the volcano. If ozone becomes entirely depleted in the plume this might in turn limit the partitioning of reactive bromine into BrO in the near-source (concentrated) plume, e.g., as suggested for conditions of extreme emissions and/or low background ozone concentrations (Jourdain et al., 2016). However, full depletion of ozone in a volcanic plume has only been observed in two measurements at Augustine in 1976 (Vance et al., 2010). Note that humidity might have an influence and is not considered here and that Hegg et al. (1976) reported a calibration accuracy of $\pm 5\%$ when the chemiluminscence technique was compared to ozone detection via potassium iodide. Reactive bromine is the principal cause of ozone depletion in volcanic plumes, but models predict that the interaction of BrO with reactive chlorine (or iodine) can augment the ozone loss.

Interhalogen, Chlorine, and Iodine Plume Chemistry *Chlorine*

The bromine explosion can ultimately also lead to the production of reactive chlorine. First, studies of Roberts et al. (2009) using the AIM model (Wexler and Clegg, 2002) on gas-particlepartitioning for mean arc volcanic compositions (Gerlach, 2004) reported that Br_2 is the favored product in heterogeneous reactions (R4) for typical volcanic Br/Cl emission compositions, according to aqueous-phase equilibria (R18). However, depletion of plume HBr through the bromine explosion can significantly reduce HBr/HCl ratios (100–1,000 times lower than arc mean composition). Under these conditions, the reactive uptake of HOBr can lead to the formation of BrCl (R17) instead of Br_2 in R4.

$$\begin{split} HOBr + H^{+}_{(aq)} + Cl^{-}_{(aq)} &\xrightarrow{aerosol-surface} BrCl + H_2O_{(aq)} (R17) \\ Br^{-}_{(aq)} + BrCl_{(aq)} &\rightleftharpoons Br_2Cl^{-}_{(aq)} \rightleftharpoons Br_{2(aq)} + Cl^{-}_{(aq)} (R18) \end{split}$$

Subsequent partitioning of BrCl (and Br₂) into the gas phase and photolysis produces Cl alongside Br radicals (R19) that can react with ozone (R20) and contribute to ozone loss (see also R5 and R2).

$$BrCl + h\nu \rightarrow Br + Cl$$
 (R19)

$$Cl + O_3 \rightarrow ClO + O_2$$
 (R20)

The reaction of ClO with BrO can lead to OClO (R21, Bobrowski and Platt, 2007; Gliß et al., 2015; Roberts et al., 2018).

$$BrO + ClO \rightarrow OClO + Br$$
 (R21)

OClO/SO₂ has been reported to increase from below detection limit at the vent up to $(3-6) \times 10^{-5}$ at Mt. Etna, Sicily, Italy [BrO/SO₂: $(1-5) \times 10^{-4}$; Bobrowski et al., 2007; General et al., 2015; Gliß et al., 2015] and $(4-6) \times 10^{-4}$ at Soufriere Hills, Montserrat [BrO/SO₂: $(1-4) \times 10^{-4}$; Donovan et al., 2014]. Compared to bromine activation <1% HCl is converted into reactive chlorine, according to both models and observations (Gliß et al., 2015). The few field-data are too limited to determine further controls on OClO formation, although the increasing OClO/SO₂ is consistent with an in-plume formation pathway.

Bromine Speciation in Volcanic Plumes

Recently, models have been able to reproduce volcanic OClO observations (Roberts et al., 2018), according to the HBr depletion and BrCl formation pathway. Models predict that bromine emissions and humidity can both impact OClO formation rates. For higher bromine emissions HBr may not become depleted immediately by the bromine explosion, leading to a later onset of BrCl formation and thus delayed rise in OClO/SO₂. Modeled OClO/SO₂ formation near downwind is enhanced at greater relative humidity because this enhances the aerosol surface that supports the heterogeneous reactions, leading to a more rapid bromine cycling and faster depletion of HBr and enabling BrCl formation (Roberts et al., 2018).

Kern and Lyons (2018) report observations of OClO (with BrO) at the plume edge, whilst only BrO is observed at the plume center. The measurements are very consistent with the horizontal spatial distribution of BrO-OClO predicted by Roberts et al. (2018). Kern and Lyons (2018) propose that measurements of OClO alongside BrO can help determine the degree of depletion of Br_(aq) (and HBr) and therefore guide interpretation of BrO observations in terms of total bromine emission.

Importantly, the fact that Br_2 (or BrCl when HBr is depleted) are the preferential products released from the reactive uptake of HO_X (X = Br, Cl) and the fast reaction of Cl with methane (R22) prevents a self-amplifying cycle for chlorine oxides. Thus, HCl cannot become depleted by plume reactive halogen chemistry (unlike HBr).

$$CH_4 + Cl \rightarrow CH_3 + HCl$$
 (R22)

Iodine

Only a few studies are available that report about iodine in tropospheric volcanic plumes, although iodine chemistry is wellstudied in marine environments (Saiz-Lopez and von Glasow, 2012 and references therein). Iodine is released as a trace emission in tropospheric volcanic plumes. For instance a total iodine to sulfur ratio (I/S) has been observed between (4-7) \times 10⁻⁶ detected with filter packs at Etna, Sicily, Italy in 2004 (Aiuppa et al., 2005), $(7-22) \times 10^{-6}$ and $(4-90) \times$ 10^{-6} at Etna and Stromboli, Mediterranean with alkaline traps in 2010-2012 (Wittmer et al., 2014), $(3-5) \times 10^{-6}$ with the Raschig Tube at Nyamuragira, in the Virunga Mountains of the Democratic Republic of the Congo in 2015 (Bobrowski et al., 2017) and 2×10^{-5} and 5.8 $\times 10^{-6}$ at Masaya and Telica, both Nicaragua respectively with filter packs in 2006 (Witt et al., 2008). Bobrowski et al. (2017) reported that the total iodine already comprises 8-18% non-hydrogen iodide measured at the crater rim of the pit crater at Nyamuragira. A major eruption of Kasatochi, Aleutian Islands 2008 enabled rare detection of IO in the volcanic plume in the stratosphere by satellite (Schönhardt et al., 2017). We highlight that the ozone depletion potential of iodine may have significant ozonedepletion impacts even as a trace emission. Iodine has an alpha factor (chemical effectiveness relative to chlorine for global ozone destruction, globally and annually averaged) of 150-300. Compared to bromine (alpha factor ca. 60), iodine can affect ozone levels even more effectively (World Meteorological Organization, 2014 and references therein).

Like the other halogens volcanic iodine is thought to be emitted mainly as hydrogen iodide with subsequent radical formation near the vent in a similar manner to bromine (R1, see section Bromine Explosion). IO is formed out of iodine radicals reacting with ozone (R23).

$$I + O_3 \rightarrow IO + O_2 \tag{R23}$$

The reaction of IO with another halogen oxide (XO, with X = Br, I) leads to OIO formation (R24). OIO can form higher iodine oxides which are precursors in particle formation and thus affect the atmospheric radiation balance (Hoffmann et al., 2001).

$$IO + XO \rightarrow OIO + X$$
 (R24)

Reactions with HOI (formed analogously to R3) form interhalogens (R25, XI with X = Br, Cl) that are released to the gas phase. Interhalogens can be photolyzed (analogous to R10) leading to halogen radicals that are involved in the catalytic cycles of halogens through plume chemistry as outlined above.

$$HOI + H^{+}_{(aq)} + X^{-}_{(aq)} \xrightarrow{aerosol-surface} XI + H_2O_{(aq)}$$
(R25)

Further details are given in von Glasow and Crutzen (2003) and literature therein.

BrO Evolution in Volcanic Plumes Dependence on Plume Age – Temporal Evolution

Observations of BrO/SO₂ ratios as a function of distance from the emission source allow to trace chemical transformations of volcanic bromine (under the condition that SO₂ is conserved). BrO was reported to be not detectable at crater rims but is formed inside the atmosphere-volcanic gas plume mixture over very short time-scales (minutes). The development of BrO/SO2 with distance downwind is shown in Figure 3, based on results of Bobrowski and Giuffrida (2012) and Platt and Bobrowski (2015) complemented by further measurement series (derived from the Supplementary Table 1). Each data series is normalized to the maximum BrO/SO2 observed, and averages were taken for measurements at the same distance within a given data series. All data arrays confirm a general rising trend. Close to the emission source BrO/SO2-ratios are low. Excluding Ambrym, Vanuatu BrO/SO₂-ratios seem to reach a relatively stable level within 5-10 km from the source. The near-constant levels then appear to decrease in some cases, for example Etna, Sicily, Italy 2004, Nyiragongo, in the Virunga Mountains of the Democratic Republic of the Congo 2004 but not for other data series e.g., Etna 2005 and Ambrym 2007.

The development of BrO is a function of plume age (and therefore depends as well on the distance); however, the distancetime interconversion depends on wind-speeds that are specific to each data-series. This means BrO formation, even if taking place at the same plume age, might be measured at a different distance. Mixing between the plume and air is also a function of distance or time downwind, and is suggested by models to strongly influence halogen speciation, hence BrO/SO₂. Indeed, whilst all volcanic plume model studies predict the initial rise



in BrO/SO₂, the occurrence of a subsequent plateau or decline depends strongly on the downwind plume-air mixing (Roberts et al., 2014b). For large gas emission fluxes like at Ambrym a lower proportion of air may initially mix into the plume. This could reduce BrO/SO₂ close to the source, as is found for both models and observations (Bani et al., 2009; Jourdain et al., 2016). However, this aspect still needs to be further investigated using measurement data.

Spatial Distribution in the Cross-Section of the Plume

Beside distribution of BrO along the plume-direction (as a function of plume age, dependent on distance from the emission source), measurement and model studies have also investigated the vertical or horizontal distribution of BrO and OCIO in the cross-section of the plume.

Vertical profiles were calculated from two-dimensional BrO and SO₂ distributions measured by imaging DOAS instruments (Bobrowski et al., 2007; Louban et al., 2009; General et al., 2015). Gliß et al. (2015) and recently Kern and Lyons (2018) investigated the vertical BrO/SO₂ distribution by using a 1 dimensional scanner unit. Bobrowski et al. (2007) found highest BrO/SO₂ values at the edges of the plume (3×10^{-4}) compared to lower values in the center (2×10^{-4}) and reproduced this trend with their model data.

Jourdain et al. (2016) discussed favored BrO formation for the edges of a volcanic plume more quantitatively, finding general agreement between model and measurement data in the trend in BrO/SO₂. While at the edges BrO reached up to 60% of the total bromine species, in the core BrO remains below 30%. Modeling predicts that Br radicals are the abundant species in the core. An explanation might be the entrainment of O₃-rich ambient air at

the plume edges to promote BrO formation, but limited transport of tropospheric O_3 and HO_X radicals toward the plume center (see section Causes and Effects of O_3 Depletion). However model studies show that plumes without ozone depletion in the core can also exhibit greater BrO/SO₂ (and OCIO/SO₂) at the plume edge (Roberts et al., 2018). This is due to the interconversion reactions between reactive bromine species, in particular Br, BrO (section Causes and Effects of O_3 Depletion), that enhances BrO/Br in the more dilute edge compared to the core of the near-downwind plume.

BrO as a Fraction of Total Bromine Under Different Plume Conditions

The conversion of HBr and the formation of BrO compared with total SO₂ flux, as well as the depletion of ozone have been reported in few measurement observations (see section Causes and Effects of O₃ Depletion) and have been investigated with model studies. Roberts et al. (2014b) showed in studies of Etna's plume HBr fully converted within tens of minutes with HBr/SO₂ ratios of 7.4×10^{-4} . While HBr/SO₂ ratios of 24×10^{-4} a conversion of only about 50% was determined within an hour of simulation (both with 20 kg/s SO₂). Bani et al. (2009) observed an increased and faster BrO/SO₂ formation dependent on the distance by comparing data with SO₂ flux below 100 kg/s and above 100 kg/s at Ambrym, Vanuatu in 2005 and 2007. Although it has to be noted that no influence of bromine emissions can be investigated in this case due to the lack of simultaneous HBr measurements.

The impact of total (Br flux) and relative (Br/S) bromine emissions on BrO development can be investigated using simultaneous measurements of Br/S and BrO/SO₂, which enables the calculation of BrO as a fraction of total bromine (BrO%). Near-simultaneous Br/S and BrO/SO₂ data sets from **Supplementary Table 1** have been used to calculate the BrO% fraction as a function of the distance from the emission source in **Figure 4**. The dependence of BrO% with distance shows a positive dependency (similar as for **Figure 3**) further supporting the assumption that increased BrO/SO₂ is caused by the formation of BrO. The particularly low BrO% for Redoubt, Alaska might be explained by the fact that the measurements of Br/S and BrO/SO₂ were not quite simultaneous (which could introduce non-linearities if the bromine emission changed).

The highest BrO% between 40 and 65% was observed at Ambrym, Vanuatu in 2007 (SO₂ flux of 47 kg/s). These datasets were also obtained at the furthest distance downwind (12–14 km) i.e., greater time-evolution to form BrO. Note that BrO/SO₂ were measured in August 2007 (Bani et al., 2009) while Br/S results derive from October 2007 (Allard et al., 2016). Jourdain et al. (2016) performed model simulations of the Ambrym plume, comparing to BrO/SO₂ measurements in January 2005 (Bani et al., 2009). The SO₂ flux was significant higher in 2005 (average 218 kg/s) than in 2007. The model predicted up to 60% BrO% at 200 km distance to the emission source at the edge of the plume (HBr/SO₂ 6.87 × 10⁻⁴). In the core only below 30% BrO% is observed.

A high BrO fraction of about 40% was observed at Etna, Sicily, Italy in 2004 with Br/S 4.8×10^{-4} and an SO₂ flux of 11 kg/s (Oppenheimer et al., 2006). Note that measurements on Etna in 2012 (Wittmer et al., 2014) resulting in 3 and 6% at similar Br/S ratios of 8.8×10^{-4} and 4.2×10^{-4} respectively occurred at plume ages <1 min. While 40% were reached by observations of Oppenheimer et al. (2006) at 2–3 km (3–4 min estimated plume age), the model results by Roberts et al. (2014b) mentioned above with HBr/SO₂ 7.4 × 10⁻⁴ (SO₂ 20 kg/s) resulted in approximately 25% at 5 min plume age. This comparison lacks of the fact that we do not know if the model set-up would agree with Oppenheimer et al.'s plume measurement parameters. It is worth highlighting that the value of 40% BrO% found for the Etna 2004 measurements is rather high compared to other datasets around this distance downwind (**Figure 4**).

Model studies of Bobrowski et al. (2015) compared Etna's plume (bromine flux 22 g/s, maximum BrO/SO₂ 2.1×10^{-4}) with slightly greater bromine flux of Nyiragongo's plume (in the Virunga Mountains of the Democratic Republic of the Congo, bromine flux 30 g/s, maximum BrO/SO₂ 0.7×10^{-4}). Simulations predicted similar BrO columns in both cases but differ in relative BrO formation. Even though Nyiragongo has greater Br/S ratios in its volcanic plume than Etna, observations of the relative conversion to BrO downwind show smaller values (BrO%: 0–2%) in the aged plume. For Nyiragongo 2007 model predictions fit with measurements, if a plume age of 1–2 min is assumed.

Comparison of Nyiragongo with studies by Kelly et al. (2013) at Mt Redoubt results in a slightly higher conversion of HBr (although low compared to Etna) in Mt Redoubt's plume. Even though Br/S emissions of 41×10^{-4} are smaller than in some cases of Nyiragongo (note also the SO₂ flux of 4.3 kg/s is lower, resulting in also lower quantities of bromine

compared to Nyiragongo as well as Etna). Model simulations that reproduced the observed ozone loss in the plume of Mt. Redoubt predicted only 30% of emitted HBr to be converted into reactive forms. This may be due to the rather dilute Redoubt plume because conversion of HBr into reactive bromine via the autocatalytic bromine explosion strongly depends on abundances of bromine and aerosols. BrO formation in this case seems to be rather low compared to other data sets mentioned above (see Figure 4). Note that model parameters and Br/S results are based on measurements at Mt Redoubt in June 2010 but BrO/SO₂ observations were made in August 2010. The observed BrO/SO_2 of 1 × 10⁻⁴ in August is rather low (BrO% of 2.4%) compared to the June model results (maximum BrO/SO₂: 6 \times 10⁻⁴, BrO%: 14%). For comparison, GOME-2 observations of Mt. Redoubt plume in the end of May found 2.5×10^{-4} BrO/SO₂ (Hörmann et al., 2013, see Supplementary Table 1). Also, the ozone depletion of August (up to 70% below ambient air at 19 min estimated plume age) is lower than observed in June (up to 90% below ambient air at 38 min estimated plume age). Since the rapid conversion of HBr to form BrO leads to ozone consumption (see section Causes and Effects of O₃ Depletion), the differences in observed ozone loss over June-August imply that BrO/SO₂ was likely higher in June 2010 than reported in August 2010. A possible explanation is that the bromine emission decreased over these months, or that changes in the plume chemistry led to a reduced conversion of emitted bromine into BrO. Unfortunately, this hypothesis cannot easily be tested as Br/S was not monitored throughout.

To study the relationship between emission fluxes and BrO fractions there are mainly three mandatory tasks: (1) Simultaneous measurements of Br/S and BrO/SO2. (2) The determination of bromine fluxes or at least SO2 fluxes so Br flux can be calculated from Br/S ratios. (3) Observations of BrO should be made at a certain distance/plume age from the emission source. As discussed before (section Dependence on Plume Age-Temporal Evolution) for comparison of BrO results in plumes measurements should be used, which reached the BrO/SO₂ "plateau" after \sim 5–10 min (2,5–10 km, depending on wind speed and plume condition). All three tasks are more or less realized by only two data sets for Etna in 2004 (Oppenheimer et al., 2006) and Gorely in 2011 (Aiuppa et al., 2012; Bobrowski et al., 2012; Platt and Bobrowski, 2015). Remarkable is the fact that Etna's plume (Br/S: 4.8×10^{-4} , SO₂: 11 kg/s) results in a significant higher HBr to BrO conversion of \sim 40% (BrO% 40%) than in Gorely's plume that had a slightly lower SO₂ flux (9.3 kg/s) and slightly higher Br/S ratio (5.3 \times 10⁻⁴, i.e., resulting in similar Br flux) resulting in ~13% BrO%.

All low-temperature kinetic models pointed out that the absolute amount of total gas emission, the total emitted bromine (also chlorine) and the initial formation of HO_X or halogen radicals in the effective source region, and the role of plume-air mixing all seem to be exert strong controls on BrO formation. However, these parameters are not well constrained in the models, nor for the intercompared field-observations, due to lack of datasets. For the intercomparison of volcanoes, the overall atmospheric setting where volcanoes are located at



should also be noted. Etna is in a more polluted region, while Nyiragongo and Redoubt release plumes in more remote, clean atmospheres. Therefore, there might be a difference in the ambient ozone and NO_X levels. It is also possible that volcanic plume halogen chemistry might be influenced by marine sources of halogens and aerosols. Finally, temporal variations due to meteorological or/and volcanological conditions may introduce variability in the datasets used for intercomparisons (noting Br/S and BrO/SO₂ measurements are difficult to perform exactly simultaneously as they are made at different locations on the volcano).

The challenge to interpreting BrO observations is highlighted by Cadoux et al. (2018). GOME-2 observations of Merapi's eruption in Indonesia 2010 resulted in extremely low BrO/SO2 ratios (maximum 8 \times 10⁻⁶; Hörmann et al., 2013), much lower than expected based on petrological estimates of the halogen/sulfur emission. Cadoux et al. (2018) discussed that BrO formation could be limited by the restriction of photochemical driven reactions caused by the ash-rich paroxysmal phase and the absence of UV light during the eruption in the night (the BrO was detected by GOME-2 in the following day). According to Cadoux et al. (2018) other possibilities include: (a) scavenging of bromine in the troposphere (e.g., adsorption of bromine on ash), (b) kinetic factors lead to favored S degassing from the magma, (c) brine saturation leading to Br uptake when magma rises, and (d) changes in Br partitioning caused by other volatile species in the magma. To this, we add the important role of plume chemistry in converting emitted HBr into BrO, as emphasized in this review.

Forecasting – Correlating Volcanic Plume Composition to Volcanic Activity

The characterization of volcanic gases can give insights about processes inside volcanoes. Volcanic gases are solved in magma at depth under high pressure. When magma is rising, decompression, diffusion, and coalescence lead to bubble formation and growth and therefore the release of gases. The depth at which gases start to escape can be different with different types of magma. The release of gases has an influence on the rising magma due to their influence on density, compressibility and flow properties. Therefore, magma degassing is a crucial control of eruption dynamics (Malinconico, 1979; Carroll and Holloway, 1994; Sparks, 2003; Moussallam et al., 2016). A strong influence is already known for the most abundant emitted gases like H₂O and CO₂. Nevertheless, even trace gases like fluorine are thought to control the viscosity of magmas with effects on crystallization-fractionation paths, controlling degassing paths and therefore the eruptive behavior (Giordano et al., 2004).

 H_2O is the most abundant emitted species, but water vapor emissions are difficult to determine due to high and very variable atmospheric background. Further, emitted water vapor is difficult to interpret because it does not always purely originate from magma but might derive also from hydrothermal systems or in some cases even from incorporated local seawater. Nevertheless, recent studies show the potential to determine water vapor in volcanic plumes and renew the hope to use them as eruption precursors in the future (Moor et al., 2016a,b; Kern et al., 2017).

Monitoring of CO_2 by remote sensing instruments is difficult as well. Non-volcanic CO_2 sources and high and variable atmospheric background concentrations may lead to variability in volcanic CO_2/SO_2 measurements (Burton et al., 2000). Today, CO_2/SO_2 ratios need usually to be determined by *in situ* sampling, however these are spatially limited and sampling points are not always representative and are frequently destroyed during eruptive periods. CO_2 is exsolved at relatively large depth and SO_2 is released at much shallower levels. Degassing of the halogens Cl and F occurs often at even lower pressure, closer to the Earth's surface (**Figure 5**; Spilliaert et al., 2006; Burton et al., 2007; Aiuppa, 2009). Observations of decreasing CO_2/SO_2 ratios and increasing Cl/SO_2 ratios could therefore be interpreted as gases derived from magma moving to shallower depth or magma from already exhausted reservoirs.

Although bromine is not known to have a large influence on eruption dynamics, bromine degassing could be nevertheless a good indicator for rising magma. In contrast to CO_2 and H_2O , SO_2 and bromine species have negligible background concentrations and clearly derive from magma degassing. The fact that SO_2 and BrO are measurable simultaneously by remote sensing instruments facilitates its applicability for monitoring volcanic activity.

Aiuppa et al. (2005) correlated Br/S ratios from filter pack samples to Etna's activity in 2004 observing increasing Br/S as well as Cl(F)/S prior to an eruption, followed by a rapid decrease. In contrast to the Br/S observation of Aiuppa et al. (2005), several authors observed lower BrO/SO₂ ratios during or also prior to eruptive activities in comparison to observations made during quiescent degassing or after eruptive activity phases. For instance as reported by Bobrowski and Giuffrida (2012) during eruptive and non-eruptive periods of Etna, Sicily, Italy from 2006 to 2009, Lübcke et al. (2014) at Nevado del Ruiz, Colombia from 2009 to 2013, Dinger et al. (2018) and Hidalgo et al. (2018) at Cotopaxi, Ecuador 2008, Warnach et al. (in review) at Tungurahua, Ecuador 2008–2014. The time lag between gas composition change and start of eruptive activities differs between the various studies and volcanoes; however, the general trend seems to be similar. Conversely, the discussion of BrO-mediated ozone depletion at Mt Redoubt, Alaska (see section BrO as a Fraction of Total Bromine Under Different Plume Conditions) points toward a possible decrease in bromine emission with assured activity.

In contrast to most chlorine and fluorine studies, recent investigations observe a correlation between BrO/SO_2 ratios and CO_2/SO_2 ratios (Bobrowski et al., 2015, 2017) leading to the assumption that bromine exsolution might be deeper than sulfur. Only recently studies on partitioning and/or knowledge of the magma movement through other geophysical or geochemical data start to deliver a basis for the interpretation of available field data (Cadoux et al., 2018). With increased degassing a change of the gas composition from S-rich to Cl-Br-rich is predicted, favoring the hypothesis of a relatively shallow degassing of bromine in comparison to sulfur that goes along with measurement results of Aiuppa et al. (2005). However, both possibilities are discussed–shallow and deep degassing of bromine (**Figure 5**).

SUMMARY OF MAIN FINDINGS

For the interpretation of BrO data regarding volcanic activity, it is essential to have a detailed understanding of the ongoing chemistry, to be able to recalculate total bromine emissions. There is a temporal and spatial heterogeneous distribution of the BrO concentration in volcanic plumes (see sections Dependence on Plume Age–Temporal Evolution and Spatial Distribution in the Cross-Section of the Plume). Current model and field





studies demonstrate that most of the BrO is not directly emitted and describe its formation through low-temperature, kinetically controlled photochemical and multiphase reactions and mixing processes with the surrounding atmosphere in the downwind plume (see section Plume Chemistry and the Formation of Reactive Bromine, summary in **Figure 2**).

Common main findings are:

- BrO/Br_{total} rises rapidly within the first minutes of plume age but usually stabilizes and then decreases again further downwind (section Dependence on Plume Age–Temporal Evolution).
- Low-temperature "bromine explosion" chemistry mechanism is able to explain BrO abundances in the downwind plume. But elevated concentrations of radicals e.g., HO_X and/or NO_X derived from high-temperature near-vent chemistry in the initial plume seem necessary to accelerate the start to enhance low-temperature BrO formation downwind in order to reach observed magnitudes of BrO (section Plume Chemistry and the Formation of Reactive Bromine).
- The autocatalytic cycles forming BrO are driven by reaction cycles involving aerosol- and gas-phase and photolysis reactions and affected by O₃ entrainment, leading to spatial heterogeneity in plume BrO/SO₂ as seen in vertical plume profiles (sections Spatial Distribution in the Cross-Section of the Plume).
- The products from heterogeneous halogen reactions are controlled by aqueous-phase equilibria that produce Br₂ (hence BrO) in volcanic plume environment. Formation of BrCl (hence OClO) occurs upon very low levels of HBr as a consequence of the bromine explosion (section Interhalogen, Chlorine, and Iodine Plume Chemistry).
- Studies of the near-source plume chemistry point to nonlinearities in HBr conversion into BrO depending on the bromine content of the emissions, plume-air mixing and other factors.

LIMITATIONS

Today, the knowledge on bromine plume chemistry is limited mainly by measurement restrictions. Model studies that have been performed in relation to particular data arrays can generally reproduce the available (limited) observations but operate in a vast parameter space that is currently not well constrained. Discrepancies exist in the predicted bromine speciation between the two available model (MISTRA and PlumeChem) probably as a result from differences in simulation algorithms, but also the wide volcanological and meteorological parameter space to which they have been applied. A general takeaway of obtained findings on one volcano cannot necessarily be applied to other volcanoes (e.g., comparison of Etna, Sicily, Italy and Nyiragongo, in the Virunga Mountains of the Democratic Republic of the Congo in Bobrowski et al., 2017). The hightemperature near-vent plume compositions are crucial for initializing low-temperature models that simulate the downwind plume chemistry. Near-source chemistry is usually investigated with high-temperature thermodynamic models. Since the hot near-vent plume cannot easily be observed by measurements, there are still uncertainties in parameterizations. The often assumed thermodynamically equilibrium conditions might not hold true (see section Model Initialization-High-Temperature Near-Vent Conditions, Dilution and Aerosols), therefore, a better characterization of the near-source volcanic plume chemistry is an essential step for understanding the evolution of volcanic plumes in the atmosphere. Even though the "bromine explosion" chemistry mechanism is relatively well-understood although still not fully characterized for volcanic plume environments there remain also some uncertainties in the low-temperature modeling for example in the parameterization of the gas-aerosol reactions (Roberts et al., 2014a), and the role of NO_X in volcanic plumes. A major issue, however, is that available datasets are typically too limited to fully constrain the model initialization such that some parameters have to be assumed, and only very limited data exist for model evaluation.

The growing amount of data sets and detailed model studies help to improve hypotheses. However, most collected data sets are incomplete toward bromine speciation and development in the plume. Future data sampling is recommended to be taken as comprehensive as possible. Therefore, three main points need to be considered: (1) Simultaneous measurements of the emissions-plume chemistry and physics are needed as both emissions and plume dynamics can vary rapidly with time. (2) Measurements of BrO for volcanic activity monitoring should be systematically undertaken at a certain plume age also because the influence of plume age on bromine chemistry needs to be taken into account to enable the interpretation of the data in the context of volcanic activity. (3) Additional parameters like ozone, aerosol particle composition, and size distribution as well as meteorological conditions, and constraints on gas flux and plume dilution/mixing into the background atmosphere are mandatory. Evidently, many restrictions reduce the availability of those data sets. For most of the bromine species as well as key chemical compounds such as OH and HO₂ no applicable methods are available for definite speciation. Recent developments have started to provide complement measurements for reactive bromine species, as outlined below. However, data sets are still rare. This activity demands a large pool of instruments, involved people, and logistical resources. Application of UAV systems is revealed as a useful tool for plume investigation but is still limited in its scope due to the often high altitude of volcanoes or other environmental effects.

Recent Developments – Opportunities and Challenges

Development of measuring techniques to add further knowledge on bromine speciation and quantification of other involved species such as OH and bromine speciation is needed to evaluate model predictions of plume chemistry. To pursue this target Rüdiger et al. (2017) presented a new *in situ* method for the determination of gaseous reactive bromine species in volcanic plumes (see section *In Situ*-Application of Instruments in the Plume). Applying active alkaline traps (Raschig Tubes) can simultaneously determine Br_{total}. So far, data sets are very

rare. The method has been applied at Etna, Sicily, Italy 2015 (Rüdiger et al., 2017), Nyamuragira, in the Virunga Mountains of the Democratic Republic of the Congo 2015 (Bobrowski et al., 2017), Stromboli, Mediterranean 2016 (Rüdiger et al., 2018b) and Masaya, Nicaragua 2016 (Rüdiger et al., in preparation). The proof of principle of ground-based sampling sets in 2015's campaigns resulted in averaged reactive bromine to Br_{total} ratios of 29 \pm 17% at Etna (plume age <1 min) and 25% \pm 9% at Nyamuragira. In 2016 applicability of UAV-based sampling at Stromboli was undertaken and resulted in an average reactive bromine to SO₂ ratio of $(5.1 \pm 4.2) \times 10^{-4}$ for the very young plume. Measurement results regarding reactive bromine do not allow an ideal and entire description of bromine plume evolution yet, because spatial information in the plume is still very limited (distances mainly at 100-200 m from emission sources and at several craters in the case of Etna). Additional methods for determination of other bromine species such as HBr in the gas phase are under development (Gutmann et al., 2018) and may complement bromine speciation soon.

Development of Fabry Perot Interferometers (FPI, Kern et al., 2010; Kuhn et al., 2018) may allow faster imaging of BrO in the future, than current DOAS application. This can be particularly valuable to separate plume chemistry (as a function of dilution, plume age etc.) from temporal variability in bromine emissions. Permanently installed DOAS instruments (e.g., NOVAC network) now provide the potential for longterm BrO/SO2 time series on a daily basis, at many volcanoes worldwide. Whilst these datasets are regularly being exploited for SO₂, so far only a few of these datasets have been analyzed for BrO, but the number is increasing. As our understanding of volcanic plume bromine chemistry improves, we may be able to better interpret these observations in terms of volcanological and meteorological conditions, to yield a valuable resource for daily bromine emissions monitoring. Recently, Kern and Lyons (2018) demonstrate the use of co-measured OClO to help better interpret volcanic BrO.

Atmospheric simulation chamber studies show promise to discover further details of volcanic plume chemistry by mimicking the plume gas and particle conditions in a controlled environment. In those experiments chemical reactions are continuously observed in real-time by a multi-instrument setup, measuring BrO, O_3 , NO_X and further compounds as well as meteorological parameters to constrain their effect on the bromine activation (Rüdiger et al., 2018a).

To provide better constraints for the initial composition of the emissions-plume ("effective source region") there are efforts to improve the understanding of high-temperature chemistry in the transient near-vent plume. A major limitation to existing thermodynamic models is the assumption of chemical equilibrium. Development of new chemical-kinetics approaches is being undertaken to more accurately reproduce the transient (hot, rapidly diluting and cooling) near-vent plume (Roberts et al., in review). This effort will improve the representation of the high-temperature near-vent chemistry as needed for initialization of the low-temperature atmospheric models of volcanic BrO.

Recent advances introducing the mechanisms developed from 1D/box models into 3D models deliver a more comprehensive

understanding of volcanic halogen chemistry as plumes disperse more widely into the atmosphere, enabling the quantification of impacts on tropospheric composition and the potential for halogens to be transported into the stratosphere. Mesoscale models with nested grids enable the simulation of plume processes and impacts from local up to regional scales (Surl et al., 2014; Jourdain et al., 2016). Advances have also been made toward developing parameterizations to include volcanic plume halogen processes in global models (Grellier et al., 2014). Whilst the box/1D models are an essential tool to probe the chemistry of volcanic plumes at the plume-scale and for comparison and to aid interpretation of near-downwind plume observations, regional and global models are needed to assess more fully impacts and to compare to satellite observations of BrO in more aged/evolved plumes. In particular, the recently launched TROPOMI satellite instrument will enable to measure BrO globally at higher spatial resolutions than previously possible.

Advances in *in situ* and remote sensing field-observations from the crater to flank, and by satellite, and development of a chain of numerical models from local (0D/1D) to regional and global (3D) scales, as well as laboratory experiments, will together address current knowledge gaps in volcanic plume bromine chemistry. Anticipated future benefits include the potential for widespread use of bromine monitoring as part of volcano hazard forecasting and a quantitative assessment of volcanic bromine impacts on the atmosphere over regionalglobal scales.

AUTHOR CONTRIBUTIONS

AG and NB collected research results. AG processed data collections. AG, NB, and TR wrote the paper with input from all authors. JR and TH contributed to the final version of the manuscript. All authors provided critical feedback and helped shape the research, analysis and manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/feart. 2018.00213/full#supplementary-material

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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