

1 **Reactive bromine chemistry in Mt Etna's volcanic plume: the**
2 **influence of total Br, high temperature processing, aerosol loading**
3 **and plume-air mixing**

4

5 **T. J. Roberts¹, R. S. Martin², and L. Jourdain¹**

6

7 [1] {LPC2E, UMR 7328, CNRS-Université d'Orléans, 3A Avenue de la Recherche
8 Scientifique, 45071 Orleans, Cedex 2, France}

9 [2] {Department of Geography, University of Cambridge, Downing Place, CB2 3EN, UK}

10

11 Correspondence to: T. J. Roberts (Tjarda.Roberts@cnsr-orleans.fr)

12

1 **Abstract**

2 Volcanic emissions present a source of reactive halogens to the troposphere, through rapid
3 plume chemistry that converts the emitted HBr to more reactive forms such as BrO. The
4 nature of this process is poorly quantified, yet is of interest to understand volcanic impacts on
5 the troposphere, and infer volcanic activity from volcanic gas measurements (i.e. BrO/SO₂
6 ratios). Recent observations from Etna report an initial increase and subsequent plateau or
7 decline in BrO/SO₂ ratios with distance downwind.

8 We present daytime *PlumeChem* model simulations that reproduce and explain the reported
9 trend in BrO/SO₂ at Etna including the initial rise and subsequent plateau. Suites of model
10 simulations also investigate the influences of volcanic aerosol loading, bromine emission, and
11 plume-air mixing rate on the downwind plume chemistry. Emitted volcanic HBr is converted
12 into reactive bromine by autocatalytic bromine chemistry cycles whose onset is accelerated
13 by the model high-temperature initialisation. These rapid chemistry cycles also impact the
14 reactive bromine speciation through inter-conversion of Br, Br₂, BrO, BrONO₂, BrCl, HOBr.

15 We predict a new evolution of Br-speciation in the plume. BrO, Br₂, Br and HBr are the main
16 plume species near downwind whilst BrO and HOBr are present further downwind (where
17 BrONO₂ and BrCl also make up a minor fraction). BrNO₂ is predicted to be only a relatively
18 minor plume component.

19 The initial rise in BrO/SO₂ occurs as ozone is entrained into the plume whose reaction with
20 Br promotes net formation of BrO. Aerosol has a modest impact on BrO/SO₂ near-downwind
21 (< ~6 km, ~10 min) at the relatively high loadings considered. The subsequent decline in
22 BrO/SO₂ occurs as entrainment of oxidants HO₂ and NO₂ promotes net formation of HOBr
23 and BrONO₂, whilst the plume dispersion dilutes volcanic aerosol so slows the heterogeneous
24 loss rates of these species. A higher volcanic aerosol loading enhances BrO/SO₂ in the (>
25 6km) downwind plume.

26 Simulations assuming low/medium and high Etna bromine emissions scenarios show the
27 bromine emission has a greater influence on BrO/SO₂ further downwind and a modest impact
28 near downwind, and show either complete or partial conversion of HBr into reactive bromine,
29 respectively, yielding BrO contents that reach up to ~50% or ~20% of total bromine (over a
30 timescale of a few 10's of minutes).

1 Plume-air mixing non-linearly impacts the downwind BrO/SO₂, as shown by simulations
2 with varying plume dispersion, wind-speed and volcanic emission flux. Greater volcanic
3 emission flux leads to lower BrO/SO₂ ratios near downwind, but also delays the subsequent
4 decline in BrO/SO₂, thus yields higher BrO/SO₂ ratios further downwind. We highlight the
5 important role of plume chemistry models for the interpretation of observed changes in
6 BrO/SO₂ during/prior to volcanic eruptions, as well as for quantifying volcanic plume
7 impacts on atmospheric chemistry. Simulated plume impacts include ozone, HO_x and NO_x
8 depletion, the latter converted into HNO₃. Partial recovery of ozone occurs with distance
9 downwind, although cumulative ozone loss is ongoing over the three hour simulations.
10

1 **1. Introduction**

2 The discovery of volcanic BrO (Bobrowski et al., 2003), and its subsequent observation in
3 many volcanic plumes globally (e.g. Oppenheimer et al., 2006, Bobrowski et al., 2007a;b,
4 Kern et al., 2009, Bani et al., 2009, Louban et al., 2009, Theys et al., 2009, Boichu et al.,
5 2011, Heue et al., 2011, Bobrowski and Giuffrida 2012, Rix et al., 2012, Hörmann et al.,
6 2013, Kelly et al., 2013, Lübcke et al., 2013), demonstrates the reactivity of volcanic halogen
7 emissions in the troposphere. Volcanoes release H₂O, CO₂ and SO₂, but also a range of
8 hydrogen halides to the atmosphere including HF, HCl, and HBr (in descending order of
9 abundance in the emission, see e.g. Aiuppa et al. 2005). HF is too strong an acid for reactive
10 halogen cycling, but for HBr and HCl, observational evidence shows these are not simply just
11 washed-out from the atmosphere, but can undergo transformation into reactive halogen
12 species.

13 Notably, DOAS (Differential Optical Absorption Spectroscopy) measurements show BrO
14 forms at 100's pmol/mol to nmol/mol mixing ratios just minutes downwind, an order of
15 magnitude higher than that found in the Arctic, where BrO episodes of up to 10's pmol/mol
16 cause significant ozone depletion and mercury deposition events (Simpson et al., 2007).
17 Additionally, there is potential to use long-term BrO monitoring at volcanoes as an indicator
18 of volcanic activity (Bobrowski and Giuffrida, 2012). Thus there is strong interest in
19 developing models to simulate the formation of reactive bromine (and chlorine) in volcanic
20 plumes, and to predict the downwind impacts from both quiescently degassing volcanoes and
21 episodic eruptions to the troposphere. Studies to date usually use equilibrium models to
22 predict the high-temperature chemistry of the near vent plume, which is then used to initialise
23 kinetic atmospheric chemistry models of the downwind reactive halogen chemistry
24 (Bobrowski et al., 2007a, Roberts et al., 2009, von Glasow 2010, Kelly et al., 2013). See von
25 Glasow et al. (2009) for an overview.

26 This study uses a purpose-built kinetic model, *PlumeChem* (Roberts et al., 2009), to
27 investigate the volcanic plume reactive halogen chemistry, focusing here on bromine in a
28 case study for Mt Etna. We include a revised methodology (Martin et al., 2009) for
29 equilibrium calculations used to represent the near-vent high-temperature chemistry, and
30 discuss uncertainties in the use of thermodynamic equilibrium models. Below, we outline the
31 progression of recent research on using equilibrium models for high-temperature near-vent
32 plume chemistry and the development of kinetic models for volcanic plume reactive halogen

1 (BrO) chemistry. We then describe the new findings of this study specifically regarding the
2 in-plume reactive bromine evolution presented by the model, and to highlight uncertainties in
3 model high-temperature initialisation and the influence of total bromine, aerosol and plume-
4 air mixing on the plume chemistry.

5

6 **1.1 Application of the HSC Equilibrium model to the near-vent plume**

7 HSC is a commercially available model (Outokumpo, Finland) that predicts the
8 thermodynamic equilibrium composition of a gas mixture at a defined temperature, pressure
9 and chemical composition. Such models are used to represent the composition of the near-
10 vent volcano plume (e.g. Gerlach, 2004, Martin et al., 2006), predicting a vast array (≥ 100) of
11 chemical species. An overview of the input and outputs to HSC is provided in Table 1. The
12 chemical composition of the mixture is determined by combining magmatic (comprising of
13 H₂O, CO₂, CO, SO₂, H₂S, H₂, HF, HCl, HBr, HI, Hg, typically at around 800-1100°C) and air
14 (N₂, O₂, Ar, typically around 0-20°C) components. The magmatic gas composition varies
15 between volcanoes and may be estimated from crater-rim measurements. It is also possible to
16 predict the abundance of gases that are missing from measurements as the magmatic gas
17 H₂O-H₂, CO₂-CO, SO₂-H₂S equilibria are functions of oxygen fugacity, pressure of degassing
18 and temperature (e.g. Giggenbach, 1987). The resulting HSC output composition depends
19 critically on the assumed ratio of air to magmatic gases in the near-vent plume, $V_A:V_M$.
20 However, this ratio is poorly defined, an issue we examine further in this study.

21 The HSC output is then used to initialise low-temperature kinetic models (such as
22 *PlumeChem*, Roberts et al., 2009, Kelly et al., 2013, *MISTRA*, Bobrowski et al., 2007a, von
23 Glasow, 2010) of the volcanic plume reactive halogen chemistry including formation of BrO.
24 These models show that elevated radicals in the HSC output accelerate the onset of
25 autocatalytic BrO chemistry, leading to very rapid BrO formation. BrO formation occurs
26 more slowly in kinetic models that are not initialised with high temperature chemistry. For
27 the interest of atmospheric modellers, we simplify the complex HSC output (≥ 100 species) in
28 Table 1, following Roberts et al. (2009) who identified impacts of HO_x, NO_x, Br_x and Cl_x on
29 the downwind plume halogen chemistry. The key species are further identified to be OH, NO,
30 Br, Cl and Cl₂, noting NO₂ \ll NO and HO₂ \ll OH, Br₂ \ll Br in the HSC output. These
31 species act to accelerate autocatalytic reactive bromine formation (see Figure 4 of Roberts et
32 al., 2009). High-temperature near-vent formation of SO₃ (a precursor to H₂SO₄) also

1 influences the volcanic plume halogen chemistry by providing a source of aerosol surface
2 area.

3 However, the thermodynamic assumption behind equilibrium models such as HSC may not
4 always be appropriate for volcanic plume applications: Martin et al. (2009) noted that the
5 near-complete re-equilibration (i.e., oxidation) of H₂S within HSC is in disagreement with the
6 widespread observed presence of H₂S in volcanic plumes (exception: Erebus), and suggested
7 a revised operation of HSC in which H₂S is removed prior to re-equilibration. Furthermore,
8 recent measurements confirming volcanic H₂ (Aiuppa et al., 2011, Roberts et al., 2012)
9 indicate this argument also applies to H₂, as well as CO (although CO is typically present in
10 low concentrations, with some exceptions e.g. Mt. Erebus. See e.g. Gerlach, 2004 for various
11 collated emission compositions. Uncertainties and limitations in the use of HSC to represent
12 the near-vent plume composition are discussed further in this study in the context of
13 downwind BrO chemistry.

14

15 **1.2 Kinetic models of downwind volcanic plume reactive halogen chemistry**

16 Atmospheric chemistry models have been developed in an effort to simulate the reactive
17 halogen chemistry of volcanic plumes, explain observed BrO formation and predict impacts
18 of reactive volcanic halogens on atmospheric chemistry. To date, two models: *MISTRA* (that
19 simulates an advected column of air, Bobrowski et al., 2007a) and *PlumeChem* (in an
20 expanding box or multi-grid box modes, Roberts et al., 2009) have been developed for this
21 purpose. Initialisation of these models includes the high-temperature chemistry of the near-
22 vent plume, as represented by HSC. Calculations by Oppenheimer et al. (2006) showed BrO
23 formation to be too slow if high-temperature near-vent radical formation is ignored.
24 Bobrowski et al. (2007a) performed the first *MISTRA* kinetic model simulations of volcanic
25 plume reactive halogen chemistry, using a model initialised with HSC at V_A:V_M of 0:100,
26 15:85, 40:60 finding the 40:60 simulation yielded highest downwind BrO/SO₂. Roberts et al.,
27 (2009) queried the use of such high V_A:V_M of 40:60 which yields rather high SO₃:SO₂ ratios,
28 that implies volcanic sulfate emissions would exceed volcanic SO₂. Roberts et al. (2009)
29 presented model simulations initialised with HSC at V_A:V_M of 10:90 that reproduced the
30 rapid formation of BrO/SO₂ at a range of Arc (subduction zone) volcanoes for the first time
31 (including Etna, Soufriere Hills, Villarrica), and suggested the higher BrO/SO₂ observed in
32 the Soufriere Hills volcano plume may be fundamentally due to higher Br/S in the emission.

1 A model study by von Glasow (2010) with simulations initialised at $V_A:V_M$ of 15:85
2 demonstrated good agreement to both reported column abundances of SO_2 and BrO/SO_2
3 ratios downwind of Etna.

4 All of the abovementioned simulations (Bobrowski et al., 2007a, Roberts et al., 2009, von
5 Glasow 2010) predict substantial in-plume depletion of oxidants, including ozone, although
6 to varying extents, and predict contrasting plume halogen evolution. Roberts et al., (2009)
7 also demonstrated conversion of NO_x into nitric acid via BrONO_2 , and proposed this
8 mechanism to explain reported elevated HNO_3 in volcano plumes. Von Glasow (2010)
9 simulated the impacts of volcanic reactive halogen chemistry on mercury speciation
10 predicting significant conversion to Hg^{II} in the plume.

11 A number of observations of ozone abundance in volcanic plumes have recently been
12 reported: Vance et al. (2010) observed ozone depletion in the Eyjafjallajökull plume, and at
13 ground-level on Etna's flanks (by passive sampling). Schumann et al. (2012) presented
14 multiple measurements of the downwind plume of Eyjafjallajökull that observed ozone
15 depletion to variable degrees. There exist also observations of depleted ozone in Mt St
16 Helen's plume (Hobbs et al., 1982) that are now believed to be likely due to BrO chemistry.
17 Ozone depletion of up to ~35 % was reported in an aircraft study of Mt Erebus plume in
18 Antarctica (Oppenheimer et al., 2010), where BrO has also been observed (Boichu et al.,
19 2011). A systematic instrumented aircraft investigation of ozone depletion in a volcano
20 plume (where emissions are also quantified) is presented by Kelly et al. (2013), and
21 compared to *PlumeChem* model simulations over 2 hours of plume evolution, finding good
22 spatial agreement in the modelled and observed ozone mixing ratios. At higher altitudes,
23 ozone depletion in a volcanic plume is reported in the UTLS (upper troposphere, lower
24 stratosphere) region observed by Rose et al. (2006), and investigated and attributed to
25 reactive halogen chemistry by Millard et al. (2006).

26 However, ozone depletion has not been universally observed: Baker et al. (2010) did not
27 detect an ozone depletion signal relative to the (somewhat variable) background level during
28 an aircraft transect through Eyjafjallajökull plume. An instrumented aircraft study found no
29 evidence for O_3 depletion in the plume of Nevado del Huila (Colombia) and found ozone
30 levels 70-80 % of ambient in the plume of Tungurahua, (Ecuador), which could not be
31 conclusively attributed to BrO chemistry (Carn et al., 2011).

1 A number of modelling discrepancies also exist. For example, the model studies of Roberts et
2 al. (2009), von Glasow (2010), and Kelly et al. (2013), predict contrasting Br-speciation
3 and contrast in predicted impacts on ozone and other oxidants. These may reflect differences
4 in the model representations and modelling uncertainties or demonstrate volcano-specific
5 differences in the plume chemistry. Navigating the vast model parameter space of volcanic
6 plume chemistry is challenging due to the non-linear controls on the plume chemistry of
7 multiple inter-dependent parameters including volcanic aerosol, rate of horizontal dispersion,
8 rate of vertical dispersion, wind-speed, volcanic gas flux, bromine in the emission, and high-
9 temperature radical formation. Limited observational datasets are available to compare to the
10 models, and the available data do not fully constrain the high- and low-temperature plume
11 chemistry. To provide further insight, this study presents new *PlumeChem* model simulations
12 to compare to recently reported trends in BrO/SO₂ ratios, and illustrates several of the major
13 controls and uncertainties in the reactive halogen chemistry of volcanic plumes.

14

1 **2. Methods**

2 **2.1 HSC: equilibrium modelling of near-vent plume chemistry**

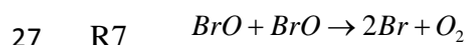
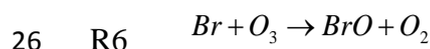
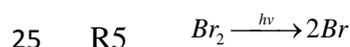
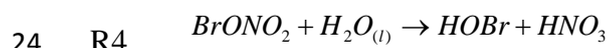
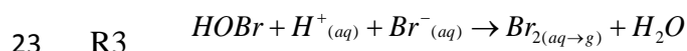
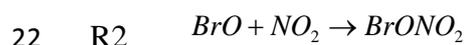
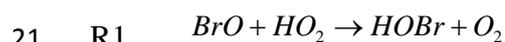
3 The use of HSC for calculating the composition of the near-vent plume is described by
4 Gerlach (2004) and Martin et al. (2006). This study uses HSC thermodynamic model version
5 7.1, and applying the modifications which were proposed by Martin et al. (2009). A simple
6 background atmosphere of N₂ (78 %), O₂ (21 %) and Ar (1%) is assumed for the HSC
7 calculations. The magmatic composition used for Etna follows that of Bagnato et al. (2007),
8 with gas mixing ratios for H₂O, CO₂, SO₂, H₂, HCl, H₂S, CO, of 0.86, $9.6 \cdot 10^{-2}$, $2.9 \cdot 10^{-2}$, $5 \cdot 10^{-3}$,
9 $1.4 \cdot 10^{-2}$, $1.5 \cdot 10^{-3}$ and $3.5 \cdot 10^{-4}$ respectively. Hg and CO are excluded for the purposes of this
10 study due to their low abundances in the volcanic emission. The bromine content as HBr, was
11 set to be either medium, high or low: ‘Medium’ bromine (molar mixing ratio of 2.16×10^{-5} ,
12 equivalent to a total bromine to SO₂ ratio (Br_{tot}/SO₂) in the emission of $7.4 \cdot 10^{-4}$) corresponds
13 to the average Br/S molar ratio at Etna NEC crater determined from filter-pack measurements
14 over 2004, Aiuppa et al. (2005). ‘High’ bromine (mixing ratio of 7.03×10^{-5} , equivalent to
15 Br_{tot}/SO₂ in the emission of $2.4 \cdot 10^{-3}$) corresponds to that assumed in a previous model study
16 of Etna (von Glasow, 2010), and is in the upper range (within one standard deviation) of the
17 observations of Aiuppa et al. (2005). Simulations are also performed at a ‘lower’ Br_{tot}/SO₂ =
18 $4.8 \cdot 10^{-4}$ which corresponds to a filterpack Br/S measurement at Voragine crater reported by
19 Oppenheimer et al. (2006). These are summarized in Table 2.

20 The magmatic temperature is set to 1050 °C in order to match that prescribed by von Glasow
21 (2010), although we note Metrich and Rutherford (1998) estimated Etna magmatic
22 temperature to be 1100°C. For the near-vent plume mixture input to HSC, ambient air
23 temperature was set to 20°C. This is somewhat high considering Etna’s elevation (3 km), but
24 this has a minor influence on the HSC output (especially considering 50 °C difference in the
25 magmatic temperature estimates outlined above). For the actual *PlumeChem* atmospheric
26 chemistry model runs, the atmospheric temperature was a more realistic 285 K. The
27 equilibrium composition was calculated for standard operation of HSC (in which H₂ and H₂S
28 are allowed to re-equilibrate) and in a revised (Martin et al., 2009) operation of HSC (in
29 which H₂ and H₂S are replaced by inert Ar such that they do not re-equilibrate). The HSC
30 calculations were performed over 16 different V_A:V_M ranging from 0:100 to 15:85.

31 **2.2 *PlumeChem*: kinetic model of downwind BrO chemistry**

1 The *PlumeChem* model simulates the reactive halogen chemistry of volcanic plume, as
 2 described by Roberts et al. (2009). It can be run in single-box (Roberts et al., 2009) or multi-
 3 box (Kelly et al. 2013) modes. Here we used the single-box that expands as a background
 4 atmosphere is entrained into it, representing dispersion of the plume as it is advected
 5 downwind. *PlumeChem* includes a background atmospheric chemistry scheme and bromine
 6 and chlorine reactive halogen chemistry, including photolysis, gas-phase and heterogeneous
 7 (gas-aerosol) phase reactions. Autocatalytic formation of BrO occurs through cycles
 8 involving reaction of BrO with oxidants, (HO_2 , NO_2), (R1,2), aerosol-phase heterogeneous
 9 chemistry (R3,4) to release a halogen dimer, whose photolysis generates two halogen radicals
 10 (R5), which may react with ozone (R6) to form BrO. The heterogeneous reactive uptake of
 11 HOBr and BrONO_2 on volcanic aerosol are thus key drivers of reactive halogen formation.
 12 Within the volcanic aerosol, aqueous-phase equilibria (Wang et al. 1994) control the nature
 13 of the product, which is Br_2 for a typical volcanic plume composition, thereby enabling
 14 autocatalytic formation of reactive bromine. Once aerosol $\text{Br}^-_{(\text{aq})}$ becomes depleted (as
 15 consequence of the BrO formation cycles), BrCl becomes a significant product from the
 16 heterogeneous reactions (R3,R4), leading to non-autocatalytic formation of reactive chlorine.
 17 Repeated cycling around R1-R6 can cause substantial ozone loss (orders of magnitude greater
 18 than the BrO mixing ratio). Repeated cycling between BrO and Br (R6, R7) further enhances
 19 ozone loss in concentrated plume environments.

20



28

1 The background atmosphere chemistry scheme used here is identical to that of Roberts et al.
2 (2009), assuming a somewhat polluted atmosphere. For the model simulations initialised
3 around midday, background ozone is ~60 nmol/mol, NO_x and HO_x are around 0.17 nmol/mol
4 and 30 pmol/mol respectively, with an ambient temperature of 285 K and 60% relative
5 humidity (RH). Plume dispersion is defined according to Pasquill-Gifford dispersion schemes
6 (see Supplementary Material). The base run plume dispersion parameterisation used in this
7 study is identical to that of Roberts et al., (2009), based on Pasquill-Gifford case D, with a
8 SO₂ gas flux of 10 kg/s at a wind-speed of 10 m/s. The influence of variations in wind-speed
9 (3-15 m/s), volcanic emission flux (10-20 kg/s SO₂) and dispersion rates (Pasquill-Gifford
10 case B,C,D) on downwind BrO/SO₂ ratios are also shown, as well as simulations with much
11 greater volcanic emission flux (5× or 10× the base run). Volcanic aerosol loading in the
12 model is investigated as part of the study, and for the majority of simulations is set to be 10⁻¹¹
13 μm² molec.SO₂⁻¹, a factor of 10 lower than that of Roberts et al. (2009), following the
14 *PlumeChem* model set-up used in Kelly et al. (2013).

15 The reaction of Br with BrONO₂ to form Br₂ + NO₃ (Orlando and Tyndall, 1996) was added
16 to *PlumeChem* model in this study. This reaction influences the overall rate of HBr
17 conversion into reactive bromine as follows: as a sink for BrONO₂ it slows the conversion of
18 HBr into reactive bromine as less BrONO₂ undergoes heterogeneous uptake (which converts
19 HBr into Br₂ via HOBr). However, as a sink for Br it slows the conversion of reactive
20 bromine back into HBr from the reaction Br + HCHO. Under a high volcanic aerosol loading
21 the former dominates, whilst the latter is more important at lower aerosol loadings. It is noted
22 that this reaction is neither included in the IUPAC Kinetics nor JPL Data evaluation
23 databases, thus is not necessarily included 'as standard' in all atmospheric models of reactive
24 halogen chemistry.

25 BrNO₂ was suggested by von Glasow (2010) to be an important reservoir for Br in the near-
26 downwind plume, based on assumed formation of BrNO₂ from volcanic NO_x and Br radicals
27 at a rate that exceeds BrNO₂ loss via photolysis. Formation of BrNO₂ was not included in
28 previous *PlumeChem* model studies (Roberts et al., 2009, Kelly et al., 2013). Here, the fate of
29 the products (BrNO₂ but also BrONO) from reaction of Br + NO₂ are investigated in more
30 detail to evaluate the potential of BrNO₂ to influence the plume chemistry.

31

32 **3. Results**

3.1 Model SO₂ column abundance, and variability in simulated BrO/SO₂

The formation of volcanic BrO is typically reported relative to SO₂, which, given slow in-plume oxidation, acts as a plume tracer on the observation time-scales (typically minutes to hours). Therefore, prior to comparing *PlumeChem* model output to the observed BrO/SO₂, a comparison is made between the simulated and reported SO₂ column abundances. Figure 1a shows slant SO₂ column abundance in Mt Etna's plume over 2004-2005, reported from DOAS (Differential Optical Absorption Spectroscopy) observations from Oppenheimer et al., (2006) and Bobrowski et al. (2007a). The data show a general decline with distance downwind, with the exception of two very near source measurements, which may have been underestimated in the very strong near-source plume, see discussion by Kern et al. (2012) and Bobrowski and Guiffrida (2012) for improved SO₂ evaluation. Also shown in Figure 1a is the model downwind plume SO₂ column abundance calculated for the plume in the vertical. The decline in modelled SO₂ column abundance with distance (or time) downwind is largely due to dispersion, given the slow rate of in-plume SO₂ oxidation. The rate of dispersion depends on plume depth, width, gas flux and wind-speed during each DOAS measurement, which are not fully constrained by available observations. Nevertheless, the broad agreement between model and observations indicates a suitable model parameterisation of plume-air mixing in the base run. This supports the use of further simulations to investigate the plume halogen chemistry using this plume-air mixing parameterisation scenario, for comparison to reported BrO/SO₂ observations.

Figure 1b shows formation of BrO (relative to plume tracer SO₂) for a range of model simulations presented later in this study, all using this same plume-air mixing parameterisation, but where the other parameters (volcanic aerosol loading, total plume bromine, initialisation using thermodynamic model output) are varied. Clearly, these variables can have a strong influence on the downwind plume halogen chemistry. Also shown are BrO/SO₂ ratios reported by Oppenheimer et al. (2006) and the observed trend in (mean) BrO/SO₂ with distance downwind reported by Bobrowski et al., (2007). Several, but not all of the model simulations in Figure 1b conform to the BrO/SO₂ observations, . Indeed simulations whose initialisations assume no plume-air mixing at high-temperature typically underestimate downwind BrO/SO₂ (see section 3.3 for further discussion). The remaining model runs demonstrate broad agreement to the BrO/SO₂ measurements and provide an

1 explanation for the observed rise and subsequent plateau or decline in BrO/SO₂ with distance
2 downwind reported by Bobrowski and Giuffrida (2012).

3 In order to provide further insight into the factors controlling volcano plume reactive halogen
4 chemistry, we investigate here the influence of the abovementioned variables, and
5 particularly uncertainties regarding the initialisation by HSC. To do so, suitable values for the
6 volcanic bromine and aerosol loading are first identified, as outlined below.

7

8 **3.2. The effect of aerosol and bromine content on downwind BrO/SO₂**

9 Highlighted in black in Figure 1b are four model runs that assume the ‘medium’ and ‘high’
10 bromine (Br_{tot}/SO₂) emission scenarios (see Table 2), and two contrasting aerosol surface
11 area loadings; namely ‘high’ aerosol estimated as $\sim 10^{-10}$ μm^2 molec.SO₂⁻¹ following Roberts
12 et al. (2009), and the ‘medium’ aerosol estimate, which is an order of magnitude lower, 10^{-11}
13 μm^2 molec.SO₂⁻¹ as was used by Kelly et al. (2013). Both the volcanic aerosol loading and
14 volcanic bromine content influence the downwind BrO/SO₂ evolution, as.

15 In general, a higher Br_{tot}/SO₂ in the emission leads to greater BrO/SO₂ far downwind. This is
16 in accordance with the proposed role of Br/S in the emission to explain order of magnitude
17 variation in BrO/SO₂ ratios across Arc volcanoes (Roberts et al., 2009). A higher aerosol
18 loading promotes the conversion of HBr into reactive forms, and promotes the occurrence of
19 reactive bromine as BrO in the far downwind plume to its role in the heterogeneous reactive
20 uptake of HOBr and BrONO₂. Interestingly, whilst the volcanic aerosol and bromine content
21 have a strong impact on the plateau in BrO/SO₂ far downwind (both in terms of value and
22 when it is reached), Figure 1b indicates that aerosol and bromine content exert a much more
23 limited impact on BrO/SO₂ in the very young plume during the first ~ 8 mins (~ 5 km) of
24 plume evolution, at least for the plume dispersion conditions simulated. For example, at 36
25 km downwind, the two contrasting aerosol loadings cause the model BrO/SO₂ to vary from
26 $4.2 \cdot 10^{-4}$ to $1.4 \cdot 10^{-3}$ (‘high’ bromine scenario) and from $2 \cdot 10^{-4}$ to $4 \cdot 10^{-4}$ (‘medium’ bromine
27 scenario), whereas at 6 km downwind all of these model runs predict BrO/SO₂ between
28 $2.5 \cdot 10^{-4}$ and $4 \cdot 10^{-4}$. This near-downwind similarity in BrO/SO₂ (despite varying Br_{tot}/SO₂ as
29 well as aerosol loading) is related to the role of oxidants in forming BrO, and differences in the
30 proportion of HBr converted to reactive bromine. This predicted near-downwind
31 independence of BrO/SO₂ on aerosol loading is consistent with the observations of

1 Bobrowski and Giuffrida (2012) at 6 km downwind that showed BrO/SO₂ was independent
2 of relative humidity (a key control on sulphate aerosol volume hence surface area). A model
3 explanation (see Section 3.4 for further discussion) is that near -downwind BrO/SO₂ ratios
4 are primarily controlled by Br to BrO partitioning– itself a function of in-plume ozone mixing
5 ratio - in this region where the plume is still relatively concentrated. See section 3.4 for
6 details of the plume reactive bromine speciation and section 3.6 for further discussion on the
7 plume impacts on atmospheric ozone.

8 Notably, the simulation with both ‘high’ bromine and the larger aerosol estimate predicts
9 BrO/SO₂ that exceeds reported observations far downwind, and does not reproduce the
10 plateau in BrO/SO₂ beyond ~ 5 km downwind of Etna summit recently reported by
11 Bobrowski and Giuffrida (2012). We acknowledge the Etna bromine emission may vary with
12 time therefore use both medium and high bromine emission scenarios alongside a low Br
13 scenario in the further model simulations of this study. However, for the high bromine
14 scenario, only the simulation with ‘medium’ aerosol surface area loading appears consistent
15 with the BrO/SO₂ observations by Bobrowski and Giuffrida (2012). The ‘medium’ aerosol
16 surface area loading is considered as likely being more representative and this estimate, ~10¹¹
17 μm² molec.SO₂⁻¹, is used in all further model simulations of this study. Further discussion
18 on estimates of the aerosol surface area loading is provided in Supplementary Material.

19

20 **3.3. The high-temperature near-vent plume - a source of model uncertainty**

21 An important model parameter in the *PlumeChem* model initialisation is the use of output
22 from thermodynamic model HSC to represent the composition of the high-temperature near-
23 vent plume.. Figure 2 compares the key species in the HSC output (Table 1) for the near-vent
24 plume of Etna (‘medium’ Br scenario), using the standard HSC methodology, in which (a) H₂
25 and H₂S re-equilibrate, and (b) the modified method (Martin et al., 2009) whereby H₂S and
26 H₂ do not re-equilibrate. NO, OH, Cl and Br and Cl₂ gas mixing ratios are shown for V_A:V_M
27 ranging from 0:100 to 15:85, where V_A:V_M is the ratio of air to magmatic gases in the near-
28 vent plume (plotted as a fraction in Figure 2), with the HSC temperature varied according to
29 the mixture of magmatic (1050 °C) and ambient (20 °C) temperatures.

30 Of note is a step increase in radical mixing ratios in Figure 2a (in which H₂ and H₂S
31 re-equilibrate). This is the so-called compositional discontinuity, C.D., (Gerlach, 2004), which

1 occurs at around $V_A:V_M \sim 0.02$ for Etna's magmatic composition. At the C.D., the reduced
2 magmatic gases (H_2S , H_2 , CO , etc) are essentially fully oxidised (SO_2 , H_2O , CO_2), thus
3 addition of further oxidant (increasing V_A/V_M) yields increases in the mixing ratios of the
4 radicals (Br, Cl, NO, OH). As $V_A:V_M$ increases further, the greater proportion of air relative
5 to magmatic gases yields a lower HSC temperature, leading to slight declines or a plateau in
6 the mixing ratios of NO and OH, and altering the balance between Cl_2 and Cl radicals (Br₂
7 remains low over the whole $V_A:V_M$ range). Formation of Br with increasing $V_A:V_M$ also leads
8 to a corresponding decrease in its 'parent' or 'source' species HBr (note other 'parent'
9 species e.g. HCl, H_2O are in excess relative to Cl_x and OH). However, in the revised HSC
10 methodology (in which H_2 and H_2S do not re-equilibrate) the C.D. has shifted to low $V_A:V_M$,
11 as first shown by Martin et al. (2009). Indeed, it may no longer be relevant to talk of a C.D. at
12 all, as an increase in radicals occurs immediately as $V_A:V_M$ is increased; this is because the
13 composition of the mixture is no longer buffered by magmatic H_2/H_2O and H_2S/SO_2 ratios.

14 The fact that certain species need to be 'protected' from re-equilibration within presents a
15 major limitation to the use of thermodynamic models to represent near-vent plume, as neither
16 the choice of $V_A:V_M$, nor the protection of certain species (but not others) are fully justified
17 on a physical basis. It is likely that some processes may be kinetics limited thus poorly
18 described by thermodynamic models. Studies suggest this is indeed the case for formation of
19 NO_x from background N_2 entrained into the plume (Martin et al. 2012), due to the high bond-
20 strength for N_2 (945 kJ/mol). Nevertheless, some evidence for the high-temperature
21 formation of radicals in the near-vent plume, for example in the presence of crater-rim
22 sulphate at $SO_4^{2-}:SO_2 \sim 1:100$ (e.g. Mather et al., 2003, Martin et al., 2008), from which near-
23 vent SO_3 production might be inferred. Further, a volcanic source of HO_x is suggested by
24 plume H_2O_2 observations of Carn et al. (2011), a source of HO_x and NO_x is suggested by
25 observations of HO_2NO_2 at Erebus (Oppenheimer et al. 2010), and elevated NO and NO_2 in
26 plumes of Masaya (Mather et al. 2004) and Mt St Helens (see Martin et al., 2012 and
27 references therein). Given abovementioned kinetic limitations to near-vent NO_x production
28 from entrained background air, these results imply the need for alternative explanations for
29 NO_x at volcanoes where it has been reported, and raise the possibility that volcano NO_x
30 emissions at other volcanoes (e.g. Etna) might be lower than predicted by HSC.

31 A representation of high-temperature radical formation in the near-vent plume is, however,
32 necessary for the initialisation of atmospheric chemistry models of downwind BrO chemistry.

1 The HSC model output is thus used for this purpose, despite above-mentioned limitations.
2 Figure 3 shows 1 hr *PlumeChem* model simulations for the three bromine emission scenarios
3 (low, medium, high), initialised using HSC operated at a range of $V_A:V_M$ varying from 0:100,
4 2:98, 5:95, 10:90 to 15:85, compared to reported BrO/SO₂ ratios from Oppenheimer et al.
5 (2006) and Bobrowski et al. (2007). Simulations initialised with $V_A:V_M$ of 0:100 (i.e. with no
6 air mixed into the near-vent plume) under-predict BrO/SO₂ ratios compared to the
7 observations, as has been shown previously (e.g. Bobrowski et al., 2007, Roberts et al., 2009,
8 von Glasow, 2010) using atmospheric chemistry models. This is due to the low radical
9 content at $V_A:V_M = 0:100$ as shown in Figure 2). Previous studies therefore chose HSC
10 initialisations using $V_A:V_M > 0:100$, e.g. Roberts et al. (2009) suggested $V_A:V_M = 10:90$, Von
11 Glasow (2010) suggested $V_A:V_M = 15:85$. Given the revised location of the compositional
12 discontinuity outlined above in Figure 2, even lower $V_A:V_M$ e.g. $V_A:V_M = 2:98$ or $V_A:V_M =$
13 $5:95$ (shown in red) can become suitable. Further progress will require more sophisticated
14 models to be developed e.g. to include full kinetic representations of chemical and mixing
15 processes.

16 Nevertheless, an interesting feature of Figure 3 is that whilst choice of HSC initialisation
17 affects the 1 hr downwind plume BrO/SO₂ strongly, the model runs show a degree of
18 convergence towards the end of the model run (particularly for low/medium Br cases).
19 Understanding the < 1 hr plume chemistry is, however, important for interpretation of flank
20 volcano BrO/SO₂ observations, and is investigated further with simulations initialised using
21 HSC with $V_A:V_M = 5:95$.

22

23 **3.4. Speciation of reactive bromine in Etna plume and implications for** 24 **observations of volcanic BrO**

25 The evolution of reactive bromine speciation is also illustrated in Figure 3 for the three
26 bromine emission scenarios, with simulations initialised using HSC at $V_A:V_M = 5:95$. A
27 number of interesting features are identified:

- 28 • BrO/Br_{tot} rises rapidly in the first few minutes, but then stabilises or declines further
29 downwind.
- 30 • HBr is fully converted to reactive bromine in the ‘medium’ and ‘low’ bromine
31 simulations but only partially converted in the ‘high’ bromine simulations.

- 1 • BrO is formed in the plume at up to 40-50 % ('medium' and 'low' Br emission
2 scenario') or 10-20 % (high Br emission scenario) of total bromine. This difference is
3 related to the extent of HBr conversion, as BrO reaches a similar maximum fraction
4 (~50%) of reactive bromine in the three simulations
- 5 • An increase in plume BrCl occurs when HBr becomes depleted, which is due to the
6 aqueous-phase equilibria producing substantial BrCl in place of Br₂.
- 7 • HOBr and BrONO₂ are present in all simulations, and represent an increasing
8 proportion of reactive bromine as the plume disperses downwind, whilst the
9 proportion of BrO declines.

10 The observed and modelled trend in BrO/SO₂ shown in Figure 1 and 3 is thus explained as
11 follows: HBr is converted into reactive forms by autocatalytic bromine chemistry cycles
12 involving volcanic aerosol, entrained atmospheric oxidants and sunlight. The HBr conversion
13 is accelerated by radical species present in the high-temperature initialisation. The initial rise
14 in BrO/SO₂ primarily reflects trends in reactive bromine speciation; entrainment of
15 background air containing ozone into the plume, promotes greater partitioning to BrO via the
16 reaction Br + O₃. Plume-air mixing is thus an important control on BrO/SO₂, because the
17 dilution of volcanic components and entrainment of air alter the balance between Br and BrO,
18 e.g. by reducing the rate of BrO loss by the self-reaction BrO + BrO (to form 2Br or Br₂), R7,
19 relative to the formation of BrO by Br+O₃, R6. The subsequent decline or plateau in BrO/SO₂
20 occurs due to net conversion of reactive bromine from BrO to HOBr and BrONO₂ in the
21 downwind plume (R1, R2). These species are formed at an accelerated rate in the downwind
22 plume as it disperses and entrains background air containing oxidants (HO₂, NO₂) which
23 react with BrO. Further, the heterogeneous loss pathways for these species are slowed in the
24 dispersed downwind plume where volcanic aerosol is diluted. The heterogeneous reactions of
25 HOBr and BrONO₂ with aerosol present a more rapid loss pathway than photolysis in the
26 aerosol-rich environment of a volcanic plume. As the plume disperses and dilutes further
27 downwind, net accumulation of HOBr (and BrONO₂) occurs whilst BrO declines (as a
28 fraction of Br_{tot}), although it is emphasized that plume chemistry cycling between these
29 species is ongoing throughout the simulation and is very rapid.

30 This predicted reactive bromine evolution is somewhat similar to that of Roberts et al. (2009)
31 but contrasts to the one hour simulations of von Glasow (2010) that did not predict the in-
32 plume presence of HOBr and BrONO₂. The higher proportion of total bromine as BrO in the

1 'medium' and 'low' Br emission scenarios (40-50 %) compared to the 'high' Br emission
2 scenario (10-20 %) is related to the extent (complete and partial, respectively) of HBr
3 conversion into reactive bromine species. This dependence of the HBr conversion on
4 $\text{Br}_{\text{tot}}/\text{SO}_2$ in the emission may to some extent explain differences between the model studies
5 of Roberts et al. (2009) and von Glasow (2010) that predicted complete and partial in-plume
6 conversion of HBr into reactive bromine, respectively.

7 Predicted BrO/SO_2 and $\text{BrO}/\text{Br}_{\text{tot}}$ trends for the three Br emission scenarios (initialised with
8 $V_{\text{A}}:V_{\text{M}} = 5:95$) are shown in Figure 4. The 'low' Br emission scenario simulation can be
9 compared to observations by Oppenheimer et al. (2006) who reported (using DOAS)
10 BrO/SO_2 ratios reached $\sim 2 \times 10^{-4}$ within 3-4 minutes downwind of Etna summit, and used
11 filter-packs to quantify the emitted $\text{Br}_{\text{tot}}/\text{SO}_2$ to be $4.8 \cdot 10^{-4}$, i.e. implying a $\text{BrO}/\text{Br}_{\text{tot}}$ of ~ 40
12 %. For the 'low' Br model run initialised at $\text{Br}_{\text{tot}}/\text{SO}_2 = 4.8 \cdot 10^{-4}$, which predicts complete
13 conversion of HBr into reactive forms over 4 minutes, BrO/SO_2 rises to 10^{-4} within 4
14 minutes, reaching a maximum of $2.5 \cdot 10^{-4}$ at about 18 minutes downwind (i.e. earlier than the
15 'medium' and 'high' bromine cases of this study) after which BrO/SO_2 declines, Figure 4a.
16 $\text{BrO}/\text{Br}_{\text{tot}}$ reaches 25 % within 4 minutes, and 40 % by about 8 minutes (Figure 4b), thus
17 converging towards the observations of Oppenheimer et al. (2006): the agreement is
18 relatively good considering the predicted $\text{BrO}/\text{Br}_{\text{tot}}$ can also be affected by other model
19 parameters kept constant here e.g. in HSC initialisation, rate of plume-air mixing, aerosol
20 loading, whilst observations of $\text{BrO}/\text{Br}_{\text{tot}}$ are subject to measurement uncertainties e.g. in
21 filterpack Br/S, DOAS measurement of BrO/SO_2 .

22 The non-linearity of HBr conversion to BrO shown in Figure 4 yields the following
23 implications for volcanology: BrO/SO_2 ratios for these simulations (initialised at $V_{\text{A}}:V_{\text{M}} =$
24 $5:95$) reach maxima of $3.6 \cdot 10^{-4}$ and $4.6 \cdot 10^{-4}$ and $2.5 \cdot 10^{-4}$ for the medium, high and low Br
25 scenarios respectively in the downwind plume. Thus, whilst the modelled bromine emission
26 has varied by a factor of three between the 'medium' and 'high' bromine scenarios, the
27 simulated BrO/SO_2 ratio has varied by less than 30 %. This result for small-scale bromine
28 variations contrasts to the earlier *PlumeChem* simulations (Roberts et al., 2009) that
29 suggested order of magnitude differences in BrO/SO_2 between Souffrière Hills volcano
30 ($\text{BrO}/\text{SO}_2 \sim 10^{-3}$) and other Arc volcanoes like Etna ($\text{BrO}/\text{SO}_2 \sim 10^{-4}$) could be attributed to
31 order of magnitude differences in the ratio of total bromine to SO_2 in their emissions.
32 However, the non-linear relationship between BrO and emitted HBr, as identified in Figure 4

1 for small-scale bromine variations, presents a complexity to efforts to quantify volcanic
2 bromine emissions using DOAS observations of plume BrO/SO₂ ratios within volcano
3 monitoring programmes, and to modelling efforts to quantify impacts from volcanic halogen
4 emissions to the troposphere. Nevertheless, DOAS observations (e.g. Bobrowski et al. 2003;
5 2007b) do suggest a positive correlation between BrO/SO₂ and volcanic HBr emissions. For
6 Soufrière Hills volcano, where high Br/S in the emission was proposed to lead to high plume
7 BrO/SO₂, further aspects to consider include the low altitude emission where ambient
8 humidity and background aerosol might be high, potentially promoting both BrO chemistry
9 and SO₂ oxidation rates.

10 Further understanding of the extent to which volcanic bromine is rapidly converted into
11 reactive forms in the near-downwind plume is needed as part of efforts to evaluate global
12 impacts from volcanic halogen degassing. Further studies of the wider model parameter space
13 can contribute to this aim, although more observations are also needed to constrain model
14 uncertainty. Overall, the model suggests HBr conversion into reactive bromine depends on a
15 balance between the autocatalytic “bromine explosion” cycles in the near-downwind plume
16 (accelerated by radicals produced in the high-temperature near-vent plume), and the
17 conversion of reactive bromine back into HBr (e.g. via the reaction Br + HCHO).

18

19 **3.5 Low in-plume prevalence of BrNO₂**

20 Formation of BrNO₂ from Br + NO₂ was excluded from the 1hr simulations presented in
21 Figure 3 and 4. However, the plume chemistry modelling study of von Glasow (2010)
22 predicted high in-plume prevalence of BrNO₂, due to reaction of Br with NO₂, given high Br
23 and NO_x mixing ratios are assumed in the (HSC) model initialisation. In the Etna simulations
24 of von Glasow (2010) formation of BrNO₂ exceeds its photolytic loss rate in the young
25 plume, leading to a significant partitioning (> 30 %) of plume bromine as BrNO₂. To further
26 evaluate this model difference, a similar two-reaction scheme for BrNO₂ was introduced into
27 the *PlumeChem* model, with BrNO₂ the assumed (sole) product of the reaction Br + NO₂.
28 With this two-reaction scheme, model runs for the three bromine scenarios also show rapid
29 formation of BrNO₂, Figure 5 (acd). The in-plume BrNO₂ prevalence (< 30 % of plume
30 bromine declining to just a few percent after 30 minutes), is still somewhat less than that of
31 von Glasow (2010), and model differences remain in Br-speciation regarding presence of
32 HOBr and BrONO₂, potentially due to differences between the models’ aerosol loading or

1 dispersion schemes. Figure 5 (acd) highlights that the rapid formation of BrNO₂ in these
2 model runs causes a slight delay to the formation of BrO downwind compared to the standard
3 model runs of Figure 3.

4 However, we do not recommend use of the two-reaction BrNO₂ scheme, because the
5 chemistry is in fact more complex. Firstly, the reaction Br + NO₂ primarily produces BrONO
6 (~92%) rather than BrNO₂ (~8%), Bröske and Zabel (1998), Orlando and Burkholder (2000).
7 Secondly, BrONO undergoes a more rapid thermal dissociation ($\tau \sim 1$ s at room temperature),
8 and photolytic loss ($\tau \sim$ seconds) than BrNO₂, Burkholder and Orlando (2000). BrONO and
9 BrNO₂ also react with NO₂ (Bröske and Zabel, 1998). BrONO (and possibly also BrNO₂)
10 also react with Br radicals. The reactions are summarized in Table 3. *PlumeChem* simulations
11 using a more detailed reaction scheme for BrNO₂-BrONO-BrNO, incorporating the
12 quantified reactions of Table 3, are illustrated in Figure 5 (bdf). With this revised BrNO₂-
13 BrONO-BrNO model scheme, these species account for only < 12 % of reactive bromine
14 (with BrONO and BrNO at only < 1%). The impact of this scheme on Br-speciation is rather
15 modest but some differences can be seen in comparison to the ‘standard’ simulations of
16 Figure 3; for example a slightly faster rate of HBr conversion to reactive bromine. However,
17 this more detailed reaction scheme is itself limited in that it does not include reaction of
18 BrNO₂ with Br (rate constant unknown), and assumes the two possible BrONO photolysis
19 pathways occur equally (as products are unknown). Further, the scheme does not include
20 potential heterogeneous reactions relevant for BrNO₂. Heterogeneous reactive uptake of N₂O₅
21 might produce BrNO₂ or ClNO₂, however, these products might react further within the
22 aerosol to form Br₂ or BrCl (Frenzel et al. 1998). Proper investigation of such heterogeneous
23 chemistry on volcanic aerosol would require detailed consideration of the underlying rate
24 constants for all the aqueous-phase reactions (e.g. in a manner similar to that recently
25 attempted for HOBr reactive uptake, Roberts et al., 2014). In addition to uncertainty in the
26 model chemistry, the model findings are also subject to uncertainty in the HSC initialisation
27 (which determines the volcanic Br and NO₂ radical source), see Section 3.3. Nevertheless, the
28 more detailed reaction BrNO₂-BrONO-BrNO scheme findings suggest the influence of
29 BrNO₂ on the plume chemistry is much lower than that proposed by von Glasow (2010).
30 Further simulations of this study therefore do not include BrNO₂.

31

32 **3.6 Influence of plume-air mixing on BrO formation and ozone depletion**

1 Here we investigate the role of plume-air mixing on the (low-temperature) halogen chemistry
2 evolution of the downwind plume. A first study investigates small variations as might be
3 expected on a day-to-day basis at Etna. A second study investigates how large variations in
4 the volcanic emission flux (e.g. due to an eruption) influence the plume chemistry, albeit
5 within the limitations of an idealised model scenario.

6

7 **3.6.1 Influence of plume dispersion parameters, volcanic emission flux and** 8 **wind-speed on BrO/SO₂**

9 As already discussed in Section 3.4, BrO formation is initially promoted by the entrainment
10 of background air (containing ozone, HO_x and NO_x), due to the balance between the reaction
11 Br + O₃ (R6) and the self-reaction of BrO (R7), but as the plume becomes more diluted the
12 entrainment of air acts to reduce BrO/SO₂ due to the reaction of BrO with HO₂ and NO₂ (R1,
13 R2). Thus, the proportion of background air that has been entrained into the plume acts as a
14 key control on BrO/SO₂. In the single-box Gaussian plume dispersion model used here, the
15 extent of mixing of the background air into the plume is controlled by the Pasquill-Gifford
16 dispersion parameters as a function of distance downwind, and whose choice depends on
17 atmospheric turbulence (a function of wind-speed and atmospheric stability). Further, for a
18 given dispersion parameterisation, the extent of mixing depends inversely on the volcanic
19 emission flux, and also depends on wind-speed (through dilution along the plume). Here the
20 effects of these three variables are illustrated for a range of plausible volcanic and
21 meteorological conditions at Etna.

22 For the base run simulations (Figure 3-4), a Pasquill-Gifford (PG) dispersion case D was
23 used, that is for a relatively neutral atmosphere, with a wind-speed of 10 m/s and volcanic gas
24 flux of 10 kg/s SO₂ (with the emission of all other volcanic gas and aerosol components
25 scaled accordingly). This SO₂ flux estimate is close to the ~13 kg/s reported by McGonigle et
26 al. (2005) for 30 July 2004. The model 10 kg/s SO₂ flux is, however, a somewhat low
27 representation for Mt Etna during 2004-5 in general. Aiuppa et al. (2005) report gas flux data
28 that show summer-time variations between 800-2000, equivalent to 9-23 kg/s SO₂, with even
29 greater SO₂ flux during eruption periods. Burton et al. (2005) report 7-day average SO₂ fluxes
30 of 1000-2500 t/d (12-25 kg/s). To illustrate the influence of variation gas flux and plume
31 dispersion, simulations were also performed at 20 kg/s SO₂ flux, and for a range of dispersion
32 and wind-speed cases. Cases C and B are introduced for more unstable atmospheric

1 conditions involving enhanced plume-air mixing, which occur more readily at lower wind-
2 speed (< 6 m/s), see Supplementary Material.

3 Simulations performed at wind-speeds of 10 m/s (case D and C), 15 m/s (case D and C), 5
4 m/s (case D and C), and 3 m/s (case C and B) are shown in Figure 6 (a ‘medium’ bromine
5 scenario is assumed for all these simulations, with $V_A:V_M = 5:95$ in the initialisation). The
6 model runs illustrate how plume-air mixing may cause variation in the downwind BrO/SO₂.
7 The variation is of a similar magnitude to that identified in the model runs with the three
8 bromine scenarios, Figure 4 (which themselves encompass only a portion of the reported
9 variability in Br/S in the emission, see Aiuppa et al., 2005). The model runs suggest that a
10 combination of variations in plume-air mixing and bromine emission could provide – at least
11 theoretically – a variability in BrO/SO₂ similar to the observed variability in BrO/SO₂ ($5 \cdot 10^{-5}$
12 - $3.9 \cdot 10^{-4}$) reported by Bobrowski and Guiffrida (2012) at 6 km downwind. Variability in the
13 volcanic aerosol emission could potentially add further to this.

14 Plume dispersion causes a transition between the two chemical regimes outlined above and
15 an intermediate maximum in BrO/SO₂. The magnitude and location of the downwind
16 maximum in BrO/SO₂ depends on the extent of plume-air mixing, as determined by the gas
17 flux, rate of dispersion and wind-speed, as well as on the volcanic aerosol loading and
18 bromine content, and the HSC initialisation. Variations in background atmospheric
19 composition (e.g. ozone, HO_x, NO_x, aerosol) could further modify the results. Finally, if
20 applying these results to volcanoes elsewhere, the summit altitude is also a relevant
21 consideration, as the greater atmospheric density at lower altitude will yield a higher in-
22 plume ratio of background oxidants to bromine, for a given volcanic SO₂ flux.

23 Nevertheless, large increases in the volcanic emission flux tend to maintain for longer the
24 more ‘concentrated’ regime where BrO/SO₂ is limited by the balance between R6 and R7, as
25 discussed further below.

26

27 **3.6.2 Effect of a large increase in volcanic flux on BrO/SO₂**

28 The sensitivity study is continued to for high emission scenarios by keeping the plume
29 dispersion case and bromine emission scenario constant (case D, ‘medium’ Br_{tot}/SO₂), but
30 increasing the volcanic gas and aerosol emission (by a factor of $\times 5$ and $\times 10$ of the base run
31 10 kg/s SO₂ flux). Such an increased volcanic emission maintains higher concentrations of

1 volcanic gases thus reduces the extent of plume-air mixing, hence entrainment of background
2 oxidants into the plume. We caution that in a real volcanic environment, such a large change
3 in degassing rate may also be accompanied by a change in composition of the volcanic
4 emission (including halogen content or aerosol loading) or act to alter the plume dimensions
5 somewhat (e.g. by the dynamics of explosive eruptions). The model results here focus solely
6 on the effect of (substantially) enhanced gas flux with all other variables held constant.

7 Simulations of three hours duration (equivalent to 108 km downwind plume propagation
8 assuming 10 m/s windspeed) with volcanic emission flux increased from the base run to $\times 5$
9 and $\times 10$ are shown in Figure 7, for both the ‘medium’ and ‘high’ bromine emission scenarios
10 (initialised with HSC using $V_A:V_M = 5:95$). The enhanced volcanic emission flux linearly
11 enhances in-plume SO_2 abundance, as expected, but exerts a non-linear effect on the plume
12 chemistry and impacts.

13 In particular, the greater volcanic emission (lower plume-air mixing) leads to a slower rise,
14 and a later onset and slower decline in BrO/SO_2 . At distances far downwind (> 2 hr for the
15 specific simulation conditions), high BrO/SO_2 is sustained for longer in plumes with high gas
16 flux. Conversely, in the near downwind (several 10’s of mins), plumes with lowest gas flux
17 exhibit the fastest initial rise and highest BrO/SO_2 ratios. As described above, these model
18 findings are readily explained by the model chemistry that partitions reactive bromine
19 between Br and BrO (during the initial rise), and BrO, and HOBr, BrONO_2 (during the
20 subsequent decline) as the plume disperses. The onset and magnitude of the decline is
21 greatest for low flux plumes that are more dilute and where a higher proportion of
22 background air has been mixed into the plume. Conversely, high flux thus more concentrated
23 plumes have a slower initial increase in BrO/SO_2 , with a delayed maximum. In the relatively
24 near-downwind plume (0-30 minutes), the model predicts lower BrO/SO_2 at greater volcanic
25 gas fluxes, as shown by the arrows in Figure 6. Implications for the interpretation of volcano
26 plume observations are discussed in Section 3.7.

27

28 **3.6.3 Atmospheric impacts of volcanic reactive halogen chemistry**

29 BrO chemistry causes ozone, HO_x and NO_x to become depleted in the downwind plume,
30 Figure 8. For HO_x and NO_x the near-downwind plume abundances are initially elevated as
31 the HSC initialisations used assumed a volcanic source of these species (Figure 2), but

1 become depleted within a few to 10's minutes downwind. The maximum depletion reaches is
2 near 100 % and > 70 % depletion relative to background values of around 30 pptv and 0.17
3 ppbv for HO_x and NO_x respectively. HO_x is converted into H₂O_(l) via HOBr chemistry (R1,
4 R3). HO_x abundances are also reduced by the gas-phase reaction of OH with SO₂, and by
5 ozone depletion in the plume (see below). The volcanic NO_x source is converted into HNO₃
6 by BrONO₂ chemistry (R2, R4), causing a rapid increase in-plume HNO₃, particularly in the
7 concentrated near-downwind plume, where HNO₃ reaches up to 60 nmol/mol (exceeding the
8 background NO_y of ~6 nmol/mol). This mechanism was proposed by Roberts et al. (2009) as
9 an explanation for observations of high HNO₃ in volcanic plumes. See collated observations
10 by Martin et al. (2012) reporting plume HNO₃/SO₂ that can reach up to 10⁻¹. For Etna in
11 particular, reported crater-rim HNO₃/SO₂ ratios are somewhat inconsistent and show large
12 variability (-2.3·10⁻⁴, 7.8·10⁻⁶, 4.2·10⁻³), which in itself might be indicative of a role of plume
13 chemistry processing. Recently Voigt et al. (2014) also observed elevated HNO₃ in the
14 downwind Etna plume, with HNO₃ as the dominant form of NO_y. Importantly, elevated
15 'volcanic' HNO₃ produced by the BrONO₂ mechanism can originate from both NO_x of
16 volcanic origin, and from NO_x from background air entrained into the plume. As
17 consequence, the in-plume NO_x declines from initially elevated abundance (due to the
18 assumed high temperature volcanic NO_x source) to become depleted relative to the
19 background abundance downwind. Finally, it is noted that simple acidification of nitrate
20 aerosol from background air entrained into the plume could also lead to gas partitioning
21 therefore enhance the 'volcanic' HNO_{3(g)} signature. Such acid-displacement of HNO_{3(g)} by
22 H₂SO_{4(aq)} has been observed by Satsumabayashi et al. (2004). The observations of volcanic
23 HNO₃ collated by Martin et al. (2012) and Voigt et al. (2014) thus require consideration in
24 the context of these two mechanisms.

25

26 Ozone is also depleted in the plume and reaches a maximum depletion (up to 100%) around
27 10 minutes downwind, coincident with the highest in-situ BrO abundances that reach ~ 1
28 nmol/mol (Figure 7). For the base run, the maximum plume ozone depletion is 30 or 45
29 nmol/mol for the medium and high bromine emission scenarios respectively. Greater in-
30 plume ozone loss occurs at higher emissions flux (lower relative plume-air mixing), however
31 for these runs the maximum ozone loss is constrained by the fact it cannot exceed ~60
32 nmol/mol (the background ozone mixing ratio). Thereafter ozone begins to recover as the

1 plume disperses (Figure 8), entraining background air, and BrO declines (Figure 7), albeit at
 2 a slower rate than the SO₂ plume tracer. Ozone recovery is greater for the base run than the
 3 higher volcanic flux cases due to both physical and chemical consequences of enhanced
 4 plume-air mixing. Thus presence of a detectable ozone depletion signature at distances far
 5 downwind depends on the emission flux and plume-dispersion. Further, the single box
 6 simulations presented here that predict the downwind trend do not simulate the ozone
 7 distribution across the plume cross-section. Ozone loss is typically greater in the plume centre
 8 than near the edges, see for example the spatially resolved model simulations for Redoubt
 9 plume that assumes similar Gaussian plume dispersion (Kelly et al. 2013). The single-box
 10 simulations should be interpreted in this context, e.g. a predicted loss of 45 nmol/mol implies
 11 greater loss at the plume centre (likely close to 60 nmol/mol or 100 %) declining to near-
 12 ambient ozone at the plume edges. For example spatially resolved model simulations for
 13 Redoubt plume predicted greater loss in the plume centre than the edges (Kelly et al. 2013).
 14 The ozone mixing ratio starts to increase when the entrainment of ambient air containing O₃
 15 is faster than the local O₃ destruction. Nevertheless, ongoing occurrence of ozone depleting
 16 BrO chemistry is demonstrated by the continuing negative trend in the cumulative ozone loss:
 17 the ozone difference (plume-background) integrated across the plume cross-sectional area
 18 declines along the 3 hr simulations to reach ~1, 4, and 7 g/cm⁻¹ for the three flux scenarios
 19 (SO₂ flux = 10, 50, 100 kg/s) respectively with greater ozone loss for the high Br compared to
 20 the medium Br scenario, as expected. These Lagrangian simulations of plume ‘puff’ ozone
 21 evolution over 3 hr can also be viewed in a Eulerian context: the 3 hr impact of continuous
 22 volcano emissions is calculated by integrating the cross-sectional impact (g/cm⁻¹) over the
 23 distance downwind. This yields ozone losses of 35·10³ (38·10³), 26·10³ (23·10³), and 6·10³
 24 (4·10³) kg for the ×10 flux, ×5 flux and base run (10 kg/s SO₂ flux) scenarios respectively the
 25 assuming the medium Br scenario (numbers in brackets refer to high Br scenario). Whilst
 26 there is some linearity in ozone loss per Br emitted (e.g. in comparing the base run to ×5 flux
 27 cases), the constraint that ozone loss cannot exceed 100% of the background abundance
 28 introduces some non-linearity for the ×10 flux case, thereby reducing its overall ozone loss.
 29 Note that the plume cross-sectional area after 3 hrs is $\pi \cdot \sqrt{2} \cdot \sigma_h \cdot \sqrt{2} \cdot \sigma_z = 2 \cdot \pi \cdot 4470 \cdot 485 = 1.4 \cdot 10^7$
 30 m². The volcanic plume cone thus resides within a cylinder of volume $1.4 \cdot 10^7 \cdot 108 \cdot 10^3$
 31 = $1.5 \cdot 10^{12}$ m³, containing approx. $110 \cdot 10^3$ kg ozone.

32 Figure 8 indicates that the plume atmospheric impacts extend beyond the one to three hour
 33 simulations presented in this study. Simulations over the lifetime of volcanic plumes under

1 different volcanological and meteorological conditions are therefore required to quantify the
2 global tropospheric impact from volcanic halogen emissions.

3

4 **3.7 Implications for modelling and observations of volcanic BrO**

5 The parameter space governing volcanic plume reactive halogen chemistry is vast, and is not
6 fully constrained by available observations. Of particular importance in controlling the
7 reactive bromine formation and downwind plume bromine speciation are: $\text{Br}_{\text{tot}}/\text{SO}_2$ in the
8 emission, the volcanic aerosol loading, and the extent of background air mixing into the
9 plume (itself a function of the plume dispersion parameterisation, volcanic emission flux and
10 wind-speed). These factors exert non-linear influences on the conversion of emitted HBr into
11 plume reactive bromine, and its speciation through interconversion of BrO, Br, Br₂, BrCl,
12 HOBr, BrONO₂.

13 The onset of the autocatalytic reactive bromine formation is also accelerated in the model by
14 radicals in the high-temperature model initialisation (Br, Cl, NO_x, HO_x). A major area of
15 uncertainty is, however, the representation of this high-temperature near-vent plume
16 environment using thermodynamic models such as HSC. Development of high-temperature
17 kinetic models of the near-vent plume is encouraged for progress in this area.

18 Further uncertainty to the downwind plume chemistry is contributed by uncertainty in the
19 volcanic bromine emission, and in aerosol surface area, that sustains halogen cycling
20 downwind. Crater-rim filter-pack measurements (e.g. Aiuppa et al., 2005) provide estimates
21 of volcanic Br/S emissions for model initialisation (see Table 2) but also highlight temporal
22 variability in this parameter. The volcanic aerosol emission is poorly constrained by
23 observations at Etna, and from volcanoes globally. A surface area loading of $\sim 10^{-11} \mu\text{m}^2$
24 molec SO₂⁻¹, i.e. an order of magnitude lower than that used by Roberts et al. (2009) yields
25 simulated (0-20 km) downwind BrO/SO₂ more consistent with that observed in the Etna
26 plume. Volcanic aerosol has a small influence on BrO/SO₂ ratio near source, but is an
27 important control in the more dispersed plume downwind. Uncertainties in the volcanic
28 aerosol emission magnitude, and its size distribution (which for sulfate varies as a function
29 temperature and humidity) thus contribute to uncertainties in models of the plume halogen
30 chemistry. Plume aerosol may be augmented by in-plume oxidation of volcanic SO₂ to
31 H₂SO₄, and the entrainment and acidification of background aerosol may also promote

1 halogen cycling. Future model evaluation of volcanic reactive halogen impacts in the wider
2 troposphere will require development of regional and global models, with detailed treatment
3 of aerosol processes as well as plume dispersion (shown to be a key control on the downwind
4 chemistry). An improved quantification of the kinetics of HOBr reactive uptake on volcanic
5 aerosol is also needed according to Roberts et al. (2014). Global models may need to include
6 a representation of the sub-grid scale volcanic plume processes, particularly as this study
7 highlighted how the proportion of emitted HBr converted into reactive forms is non-linearly
8 dependent on the degassing scenario.

9

10 We emphasize the complex role of plume chemistry in the interpretation of volcano flank
11 DOAS measurements of BrO/SO₂. Bobrowski and Giuffrida (2012) recently reported
12 variation in BrO/SO₂ ratios at Etna related to the onset of eruption activity, for example with
13 increasing BrO/SO₂ shortly prior to an eruptive event, and lower BrO/SO₂ during the eruption
14 event, according to DOAS measurements 6 km downwind from the summit. These
15 observations have been interpreted in the context of variable bromine and SO₂ emissions,
16 related to subsurface magmatic processes. Lübcke et al. (2013) identified a decrease in
17 BrO/SO₂ observed using a DOAS instrument prior to an eruption event at Nevado del Ruiz,
18 Colombia (in a period whilst SO₂ emissions were increasing). However, we emphasize that a
19 variation in plume BrO/SO₂ can also result from differences in the plume chemistry for
20 varying volcanic emission flux magnitudes. Figure 6 shows that changes in volcanic gas flux
21 (for a fixed plume dimension) can yield substantial changes in plume BrO/SO₂ ratio, even for
22 a fixed Br_{tot}:SO₂ ratio in the emission. In the near-downwind plume, a key control on BrO
23 formation is the entrainment of oxidants. A substantial increase in volcanic emission flux
24 leads to greater plume strength and reduced ratio of background oxidants to bromine in the
25 model. Thus, on the <60 min timescale of volcano flank DOAS observations, a substantially
26 enhanced rate of volcanic degassing generally leads to lower plume BrO/SO₂ ratios in more
27 concentrated plumes. Potentially, the variations in BrO/SO₂ identified by Bobrowski and
28 Giuffrida (2012), and Lübcke et al. (2013) may result from a combination of volcanological
29 and plume chemistry factors. This example highlights the complexity surrounding
30 interpretation of volcanic BrO and shows the role of plume chemistry modelling in the effort
31 to use volcanic BrO observations to monitor and predict volcanic activity.

1 We also highlight that the plume chemical evolution causes a decline in BrO/SO₂ ratios in the
2 dispersed plume further downwind through net conversion of BrO into reservoirs such as
3 HOBr and BrONO₂. This plume chemical evolution acts to reduce the BrO column
4 abundance, contributing additional limitations to its possible detection in dispersed plumes,
5 and is the model explanation for the plateau in BrO/SO₂ downwind of Etna reported by
6 Bobrowski and Guiffrida (2012). Detection of volcanic BrO by satellite is primarily
7 constrained to large volcanic emissions (Theys et al., 2009, Rix et al. 2012, Hörmann et al.,
8 2013). Smaller volcanic emissions that generate high but localised BrO at lower altitudes are
9 less readily detected particularly due to dilution effects across the satellite measurement pixel
10 (Afe et al., 2004). The modelled plume chemical evolution adds to this limitation for satellite
11 detection of BrO in dispersed volcanic plumes (even at higher resolution). Importantly,
12 however, the model Br-speciation shows that a declining trend in BrO abundance as the
13 volcanic plume disperses does not preclude the occurrence of continued in-plume reactive
14 bromine chemistry as predicted by the model.

15

16 **4. Conclusion**

17 We present a *PlumeChem* model study of the reactive halogen chemistry of Mt Etna volcano
18 plume that reproduces the recently reported trends in BrO/SO₂; namely a rapid increase in the
19 near-downwind followed by stability or decline in the far-downwind. A new in-plume
20 evolution of Br-speciation is predicted: BrO Br₂, Br and HBr are the main plume species in
21 the near downwind plume whilst BrO, HOBr (and BrONO₂, BrCl) are present in significant
22 quantities further downwind. An evaluation of the (quantifiable) chemistry surrounding
23 BrNO₂ suggests a rather low prevalence in volcanic plumes, although uncertainties in model
24 chemistry and initialisation are highlighted.

25 Emitted volcanic HBr is converted into reactive bromine by autocatalytic bromine chemistry
26 cycles whose onset is accelerated by the model high-temperature initialisation. The initial rise
27 in BrO/SO₂ is primarily due to entrainment of ozone through plume dispersion that promotes
28 BrO formation from Br radicals. A subsequent decline or plateau in BrO/SO₂ occurs upon
29 plume dispersion, which both dilutes the volcanic aerosol (slowing HOBr and BrONO₂
30 heterogeneous loss rates) and entrains HO₂ and NO₂ from the background atmosphere
31 (promoting HOBr and BrONO₂ formation from BrO). This promotes net accumulation of
32 reservoirs HOBr and BrONO₂ and a reduction in BrO in the dispersed downwind plume.

1 Thus the model can explain the reported BrO/SO₂ trend at Etna. We demonstrate the role of
2 plume chemistry models to interpret volcanic BrO/SO₂ observations as well as quantify
3 atmospheric impacts on HO_x, NO_x, HNO₃ and ozone. A number of volcanological and
4 meteorological factors can influence plume BrO/SO₂ ratios, and we illustrate simulations
5 with contrasting total bromine content and volcanic aerosol loading. The influence of plume-
6 air mixing is shown by simulations with varying dispersion rate, as well as wind-speed and
7 volcanic gas flux.

8 BrO contents reach up to 20% and ~50% of total bromine (over a timescale of a few 10's of
9 minutes), for the high and medium/low bromine emission scenarios, respectively. The latter
10 agrees well with observations that report BrO (at 3-5 min downwind) can reach up to 40% of
11 the total bromine emission at Etna (Oppenheimer et al., 2006). Partial (up to ~50%) or
12 complete (100%) conversion of HBr to reactive forms is predicted over the one hour
13 simulations, depending on bromine content (high, medium or low, respectively) as well as
14 other the plume conditions (e.g. aerosol, dispersion, HSC initialisation). Simulations using
15 the two volcanic aerosol loadings significantly differ in the downwind plume chemistry but
16 result in a similar initial rise in BrO/SO₂ near-downwind (up to 6 km), a finding that is in
17 agreement with the reported low relative humidity dependence of BrO/SO₂ (Bobrowski and
18 Giuffrida, 2012).

19 Simulations with a fixed dispersion rate but enhanced volcanic emission flux are presented.
20 For higher emission fluxes, the stronger plume and reduced ratio of background
21 oxidants:bromine causes a slower rise in BrO/SO₂ in the near downwind plume (< 40 min)
22 and a slower and delayed onset of the decrease in BrO/SO₂ in the far downwind plume (> 2
23 hr, for the volcanic conditions simulated). This simulated dependence of BrO/SO₂ on
24 volcanic emission flux (albeit in an idealised model scenario) is particularly relevant towards
25 the interpretation of changes in BrO/SO₂ during/prior to eruptive events (e.g. Bobrowski and
26 Giuffrida, 2012, Lübcke et al., 2013).

27 Impacts of the plume halogen chemistry include downwind depletion of HO_x, NO_x and
28 ozone, and formation of HNO₃. Partial recovery of ozone is predicted, particularly for low
29 gas flux emissions. However cumulative impacts on ozone are ongoing over the 3 h
30 simulations.

31

1

2 **Acknowledgements**

3 This study was financed by LABEX VOLTAIRE (VOLatils- Terre Atmosphère Interactions -
4 Ressources et Environnement) ANR-10-LABX-100-01 (2011-20) and an NSINK career
5 development allowance that enabled HSC software purchase. RSM acknowledges Christ's
6 College, Cambridge for a research fellowship. Contributions of authors: TJR designed and
7 performed the *PlumeChem* model experiments and HSC calculations and wrote the
8 manuscript. RSM advised on HSC methodology and contributed to manuscript writing. LJ
9 advised on scientific scope and contributed to manuscript writing.

10

11 **References**

12

13 Afe O.T., Richter A., Sierk B., Wittrock F., and Burrows J.P.: BrO emission from volcanoes:
14 A survey using GOME and SCIAMACHY measurements, *Geophysical Research Letters*, 31,
15 L24113, doi:10.1029/2004GL020994, 2004.

16

17 Aiuppa A., Federico C., Franco A., Giudice G., Guierri S., Inguaggiato, Liuzzo M.,
18 McGonigle A.J.S., Valenza M.: Emission of bromine and iodine from Mount Etna volcano,
19 *Geochemistry, Geophysics, Geosystems*, 6,8, Q08008, doi:10.1029/2005GC000965, 2005.

20

21 Aiuppa, A., Shinohara H.,³ Tamburello G., Giudice G., Liuzzo M., Moretti R.: Hydrogen in
22 the gas plume of an open-vent volcano, Mount Etna, Italy, *Journal of Geophysical Research*,
23 116, B10204, doi:10.1029/2011JB008461, 2011.

24

25 Bagnato E., Aiuppa A., Parello F., Calabrese S., D'Alessandro W., Mather T.A., McGonigle
26 A.J.S., Pyle D.M., Wangberg I.: Degassing of gaseous (elemental and reactive) and
27 particulate mercury from Mount Etna volcano (Southern Italy), *Atmospheric Environment*,
28 41,7377–7388, 2007.

29

1 Baker, A. K., Rauther- Schöch A., Schuck T. J., Brenninkmeijer C. A. M., van Velthoven P.
2 F. J. Wisner A., Oram D. E., Investigation of chlorine radical chemistry in the
3 Eyjafjallajökull volcanic plume using observed depletions in non-methane hydrocarbons,
4 Geophysical Research Letters, 38, L13801, doi:10.1029/2011GL047571, 2011.
5
6 Bani P., Oppenheimer C., Tsanev V. I., Carn S. A., Cronin S. J., Crimp R., Calkins, J. A.,
7 Charley D., Lardy M., and Roberts T.J.: Surge in sulphur and halogen degassing from
8 Ambrym volcano, Vanuatu, *B. Volcanol.*, 71, 10, 1159–1168, DOI 10.1007/s00445-009-
9 0293-7, 2009.
10
11 Bobrowski, N., Honniger, G., Galle, B. and Platt, U.: Detection of bromine monoxide in a
12 volcanic plume. *Nature*, 423, 273-276, doi:10.1038/nature01625, 2003.
13
14 Bobrowski, N., von Glasow, R., Aiuppa, A., Inguaggiato, S., Louban, I., Ibrahim, O. W.
15 and Platt, U.: Reactive halogen chemistry in volcanic plumes, *J. Geophys. Res.*, 112, D06311,
16 doi:10.1029/2006JD007206, 2007a.
17
18 Bobrowski, N. and Platt, U.: SO₂/BrO ratios studied in five volcanic plumes. *J. Volcanol.*
19 *Geoth. Res.*, 166, 3-4, 147-160, 10.1016/j.jvolgeores.2007.07.003, 2007b.
20
21 Bobrowski N. and G. Giuffrida: Bromine monoxide/sulphur dioxide ratios in relation to
22 volcanological observations at Mt. Etna 2006–2009, *Solid Earth*, 3, 433–445, doi:10.5194/se-
23 3-433-2012, 2012.
24
25 Boichu, M., Oppenheimer C., Roberts T. J., Tsanev V., Kyle P. R.: On bromine, nitrogen
26 oxides and ozone depletion in the tropospheric plume of Erebus volcano (Antarctica), *Atmos.*
27 *Environ.*, 45, 23, 3856-3866, 2011.
28

1 Bröske R. and Zabel F.: Kinetics of the Gas-Phase Reaction of BrNO₂ with NO, J. Phys.
2 Chem. A 1998, 102, 8626-8631
3

4 Burkholder J. B. and Orlando J. J., UV absorption cross-sections of cis-BrONO, Chemical
5 Physics Letters, 317, 6, 603–608, 2000.
6

7 Burton, M. R., Neri M., Andronico D., Branca S., Caltabiano T., Calvari S., Corsaro R. A.,
8 Del Carlo P., Lanzafame G., Lodato L., Miraglia L., Salerno G., and Spampinato L.: Etna
9 2004–2005: An archetype for geodynamically-controlled effusive eruptions, Geophys. Res.
10 Lett., 32, L09303, doi:10.1029/2005GL022527, 2005.
11

12 Carn S. A, Froyd K. D., Anderson B. E., Wennberg P., Crouse J., Spencer K., Dibb J. E.,
13 Krotkov N. A., Browell E. V., Hair J. W., Diskin G., Sachse G., and Vay S. A., In situ
14 measurements of tropospheric volcanic plumes in Ecuador and Colombia during TC, Journal
15 of Geophysical Research, 116, D00J24, doi:10.1029/2010JD014718, 2011.
16

17 Frenzel A., Scheer V., Sikorski R., George Ch., Behnke W., Zetzsch C.: Heterogeneous
18 Interconversion Reactions of BrNO₂, ClNO₂, Br₂, and Cl₂, J. Phys. Chem. A, 102, 1329-
19 1337, 1998.
20

21 Gerlach, T.M.: Volcanic sources of tropospheric ozone-depleting trace gases, Geochemistry.
22 Geophysics. Geosystems (G³) 5, Q09007, doi:10.1029/2004GC000747, 2004.
23

24 Giggenbach, W.F.: Redox processes governing the chemistry of fumarolic gas discharges
25 from White Island, New Zeland, Appl. Geochem., 2, 143-161, 1987.
26

27 Grimley, A. J.; Houston, P. L. The photochemistry of nitrosyl halides: The X+NOX--
28 >X₂+NO(v) reaction (X=Cl, Br), J. Chem. Phys. 72, 1471, 1980.

1

2 Heue, K.-P., Brenninkmeijer, C.A.M. , Baker, A. K. , Rauthe-Schöch, A. , Walter, D. ,
3 Wagner, T. , Hörmann, C. , Sihler, H. , Dix, B. , Frieß, U., Platt, U. , Martinsson, B. G. ,
4 van Velthoven, P. F. J. , Zahn, A. and Ebinghaus, R.: SO₂ and BrO observation in the plume
5 of the Eyjafjallajökull volcano 2010: CARIBIC and GOME-2 retrievals, *Atmos. Chem.*
6 *Phys.*, 11, 2973-2989, doi:10.5194/acp-11-2973-2011, 2011.

7

8 Hippler, H.; Luu, S. H.; Teitelbaum, H. and Troe, J.: Flash photolysis study of the NO-
9 catalyzed recombination of bromine atoms, *Int. J. Chem. Kinet.* 10, 155. 1978.

10

11 Hobbs, P. V., Tuell, J. P., Hegg, D. A., Radke, L. F. and Eltgroth, M. W.: Particles and
12 gases in the emissions from the 1980-1981 volcanic eruptions of Mt. St. Helens., *J. of*
13 *Geophys. Res.*, 87, C13, 11062-11086, 1982.

14

15 Hörmann C., Siöler H., Bobrowski N., Beirle S., Penning de Vries M., Platt U., and Wagner
16 T.: Systematic investigation of bromine monoxide in volcanic plumes from space by using
17 the GOME-2 instrument, *Atmos. Chem. Phys.*, 13, 4749–4781, 2013.

18

19 Kelly P.J., Kern C., Roberts T.J., Lopez T., Werner C., and Aiuppa A.: Rapid chemical
20 evolution of tropospheric volcanic emissions from Redoubt Volcano, Alaska, based on
21 observations of ozone and halogen-containing gases, *Journal of Volcanology and Geothermal*
22 *Research*, *Journal of Volcanology and Geothermal Research*, 259, 317–333, 2013.

23

24 Kern, C., Sihler, H., Vogel, L., Rivera, C., Herrera, M. and Platt, U.: Halogen oxide
25 measurements at Masaya Volcano, Nicaragua using active long path differential optical
26 absorption spectroscopy, *B. Volcanol.*, 71, 6, 659-670, 2009.

27

28 Kern, C., T. Deutschmann, C. Werner, A. J. Sutton, T. Elias, and P. J. Kelly (2012),
29 Improving the accuracy of SO₂ column densities and emission rates obtained from upward-

1 looking UV-spectroscopic measurements of volcanic plumes by taking realistic radiative
2 transfer into account, *J. Geophys. Res.*, 117, D20302, doi:10.1029/2012JD017936.

3

4 Louban I., Bobrowski N. , Rouwet D., Inguaggiato S., and Platt U.: Imaging DOAS for
5 volcanological applications, *Bulletin of Volcanology*, 71, 753–765, 2009.

6

7 Lübcke P., Bobrowski, N., Arellano, S., Galle, B., Garzón, G., Vogel, L., Platt U., BrO/SO₂
8 molar ratios from scanning DOAS measurements in the NOVAC network, *Solid Earth*
9 *Discuss.*, 5, 1845–1870, 2013.

10

11 Martin, R. S., Mather, T. A. and Pyle, D. M.: High-temperature mixtures of magmatic and
12 atmospheric gases, *Geochemistry, Geophysics, Geosystems (G³)*, 7, Q04006,
13 doi:10.1029/2005GC001186, 2006.

14

15 Martin, R. S., Roberts, T. J., Mather, T. A. and Pyle, D. M.: The implications of H₂S and H₂
16 stability in high-T mixtures of magmatic and atmospheric gases for the production of
17 oxidized trace species (e.g., BrO and NO_x). *Chem. Geol.* 263, 143-150, 2009.

18

19 Martin, R. S., Mather, T. A., Pyle, D. M., Power, M. Allen, A. G., Aiuppa, A., Horwell, C. J.
20 and Ward E. P. W.: Composition-resolved size distributions of volcanic aerosols in the Mt.
21 Etna plumes, *Journal of Geophysical Research*, 113, D17211, doi:10.1029/2007JD009648,
22 2008.

23

24 Martin R.S., Ilyinskaya E., Oppenheimer C., The enigma of reactive nitrogen in volcanic
25 emissions, *Geochimica et Cosmochimica Acta* 95, 93–105, 2012.

26

27 Mather T.A., Allen A. G., Oppenheimer C., Pyle D. M., McGonigle A. J. S., Size-Resolved
28 Characterisation of Soluble Ions in the Particles in the Tropospheric Plume of Masaya

1 Volcano, Nicaragua: Origins and Plume Processing, *Journal of Atmospheric Chemistry* 46:
2 207–237, 2003.

3

4

5 Mather T.A., Pyle D.M. and Allen A.G.: Volcanic source for fixed nitrogen in the early
6 Earth's atmosphere, *Geology*; October 2004; v. 32; no. 10; p. 905–908; doi:
7 10.1130/G20679.1.

8

9 McGonigle A. J. S., Inguaggiato S., Aiuppa A., Hayes A. R. and Oppenheimer C., Accurate
10 measurement of volcanic SO₂ flux: Determination of plume transport speed and integrated
11 SO₂ concentration with a single device, *Geochem. Geophys. Geosyst.*, 6, Q02003,
12 doi:10.1029/2004GC000845.

13

14 Mellouki, A.; Laverdet, G.; Jourdain, J. L.; Poulet, G., Kinetics of the reactions Br + NO₂ +
15 M and I + NO₂ + M, *Int. J. Chem. Kinet.* 1989, 21, 1161.

16

17 Metrich, N; Rutherford, M J. Low pressure crystallization paths of H₂O-saturated basaltic-
18 hawaiitic melts from Mt Etna: Implications for open-system degassing of basaltic volcanoes.
19 *Geochimica et Cosmochimica Acta*, 62, 7: 1195-1205. 1998.

20

21 Millard G. A., Mather T. A., Pyle D. M., Rose, W. I. and Thornton B.: Halogen emissions
22 from a small volcanic eruption: Modeling the peak concentrations, dispersion, and
23 volcanically induced ozone loss in the stratosphere, *Geophysical Research Letters*, 33,
24 L19815, doi:10.1029/2006GL026959, 2006.

25

26 Oppenheimer, C., Tsanev, V. I., Braban, C. F., Cox, R. A., Adams, J. W., Aiuppa, A.,
27 Bobrowski, N., Delmelle, P., Barclay, J. and McGonigle, A. J. S.: BrO formation in volcanic
28 plumes. *Geochim. Cosmochim. Ac.*, 70, 2935-2941, 2006.

29

1 Oppenheimer C., Kyle P. Eisele F., Crawford J., Huey G., Tanner D., Saewung K., Mauldin,
2 L., Blake, D., Beyersdorf A., Buhr M. and Davis D.: Atmospheric chemistry of an Antarctic
3 volcanic plume, *J. of Geophys. Res.*, 115, D04303, doi:10.1029/2009JD011910, 2010.
4

5 Orlando J.J., Burkholder J.B.: Identification of BrONO as the Major Product in the Gas-
6 Phase Reaction of Br with NO₂, *J. Phys. Chem. A*, 104, 10, 2048–2053, DOI:
7 10.1021/jp993713g, 2000.
8

9 Orlando, J. J., and Tyndall, G. S.: Rate coefficients for the thermal decomposition of BrONO₂
10 and the heat of formation of BrONO₂, *Journal of Physical Chemistry*, 100, 19398-19405,
11 1996.

12 Rix M., Valks P. Hao N., Loyola D., Schlager H. Huntrieser H., Flemming J., Koehler U.,
13 Schumann U. and Inness A.: Volcanic SO₂, BrO and plume height estimations using GOME-
14 2 satellite measurements during the eruption of Eyjafjallajökull in May 2010, *Journal of*
15 *Geophysical Research*, 117, D00U19, 19 PP., 1984-2012, DOI: 10.1029/2011JD016718,
16 2012.
17

18 Roberts, T. J., Braban, C. F., Martin, R. S., Oppenheimer, C., Adams, J. W., Cox, R. A.,
19 Jones R. L. and Griffiths., P. T, Modelling reactive halogen formation and ozone depletion in
20 volcanic plumes. *Chem. Geol.*, 263,151-163, 2009.
21

22 Roberts T.J., Braban C.F., Martin R.S., Oppenheimer C., Dawson D. H., Griffiths P. T. G,
23 Cox R.A., Saffell J.R. Jones R.L.: Electrochemical Sensing of Volcanic Plumes, *Chem.*
24 *Geol., Chemical Geology* 332-333, 74–91, 2012.
25

26 Roberts T. J., Jourdain L., Griffiths P. T., Pirre M., Re-evaluating the reactive uptake of
27 HOBr in the troposphere with implications for the marine boundary layer and volcanic
28 plumes, in review for *ACPD*, 2014.
29

1 Rose, W. I., Millard G. A., Mather T. A., Hunton D. E., Anderson B., Oppenheimer C.,
2 Thornton B. F., Gerlach T. M., Viggiano A. A., Kondon Y., Miller T. M., and Ballenthin J.
3 O.: Atmospheric chemistry of a 33–34 hour old volcanic cloud from Hekla Volcano
4 (Iceland): Insights from direct sampling and the application of chemical box modeling, *J.*
5 *Geophys. Res.*, 111, D20206, doi:10.1029/2005JD006872, 2006.

6

7 Satsumabayashi, H., Kawamura M., Katsuno T., Futaki K., Murano K., Carmichael G. R.,
8 Kajino M., Horiguchi M., and Ueda H.: Effects of Miyake volcanic effluents on airborne
9 particles and precipitation in central Japan, *J. Geophys. Res.*, 109, D19202,
10 doi:10.1029/2003JD004204, 2004.

11

12 Scheffler, D., Grothe, H., Willner, A., Frenzel, A., and Zetzsch, C.: Properties of Pure Nitryl
13 Bromide. Thermal Behavior, UV/Vis and FTIR Spectra, and Photoisomerization to trans-
14 BrONO in an Argon Matrix, *Inorg. Chem.* 36, 335-338, 1997.

15

16 Schumann U., Weinzierl B., Reitebuch, O. Schlager, H., Minikin A., Forster, C., Baumann
17 R., Sailer, T., Graf, K., Mannstein, H., Voigt, C., Rahm S., Simmet R., Scheibe, M.,
18 Lichtenstern, M., Stock, P., Rüba H., Schäuble, D., Tafferner, A., Rautenhaus, M., Gerz T.,
19 Ziereis, H., Krautstrunk, M., Mallaun, C., Gayet J.-F., Lieke K., Kandler K., Ebert M.,
20 Weinbruch S., Stohl A., Gasteiger J., Groß S., Freudenthaler V., Wiegner M., Ansmann A.,
21 Tesche M., Olafsson H., and Sturm K.: Airborne observations of the Eyjafjalla volcano ash
22 cloud over Europe during air space closure in April and May 2010, *Atmos. Chem.*
23 *Phys.*, 11, 2245-2279, 2011.

24

25 Simpson W. R., von Glasow R. Riedel K., Anderson P., Ariya P., Bottenheim J., Burrows J.,
26 Carpenter L. J., Frieß U., Goodsite M. E., Heard D., Hutterli M., Jacobi H.-W., Kaleschke L.,
27 Neff B., Plane J., Platt, Richter A., Roscoe H., Sander R., Shepson P., Sodeau J., Steffen A.,
28 Wagner T., and Wolff E., Halogens and their role in polar boundary-layer ozone depletion,
29 *Atmos. Chem. Phys.*, 7, 4375–4418, 2007.

30

1 Theys N., Van Roozendael M., Dils B., Hendrick, F., Hao, N., and De Mazière M. First
2 satellite detection of volcanic bromine monoxide emission after the Kasatochi eruption,
3 *Geophysical Research Letters*, 36, L03809, doi:10.1029/2008GL036552, 2009.

4

5 Vance A., McGonigle, A. J. S., Aiuppa, A., Stith J. L. , Turnbull, K., and von Glasow R.,
6 Ozone depletion in tropospheric volcanic plumes, *Geophys. Res. Lett.*, 37, L22802,
7 doi:10.1029/2010GL044997, 2010.

8

9 Voigt, C., P. Jessberger, T. Jurkat, S. Kaufmann, R. Baumann, H. Schlager, N. Bobrowski, G.
10 Giuffrida, and G. Salerno, Evolution of CO₂, SO₂, HCl, and HNO₃ in the volcanic plumes
11 from Etna, *Geophys. Res. Lett.*, 41, 2196–2203, doi:10.1002/2013GL058974, 2014.

12

13 Von Glasow, R., Bobrowski, N. and Kern, C.: The effects of volcanic eruptions on
14 atmospheric chemistry, *Chem. Geol.*, 263, 131-142, 2009.

15

16 Von Glasow, R.: Atmospheric Chemistry in Volcanic Plumes, *PNAS*, 107, 15, 6594-6599,
17 2010.

18

19 Wang, T. X., Kelley, M. D., Cooper, J. N., Beckwith, R. C. and Margerum, D. W:
20 Equilibrium, kinetic and UV-spectral characteristics of aqueous bromine chloride, bromine
21 and chlorine species, *Inorg. Chem.*, 33, 5872-5878 , 1994.

- 1 Table 1. Thermodynamic modelling of the high temperature near vent plume using HSC:
- 2 Overview of Inputs and Outputs.

HSC Input: Chemical	Comments
H ₂ O, CO ₂ , SO ₂	Major Volcanic Gases
HF, HCl, HBr, HI	Halogen Emissions
H ₂ S, CO, H ₂	Reduced Gases
Hg	Trace Metals
N ₂ , O ₂ , Ar	Air

HSC Input: Physical

V _A :V _M	Atmospheric:Magmatic Gas Ratio
Temperature	Magmatic and Ambient Temperature

HSC Output:

Full Matrix of Species	(* see footnote)
------------------------	------------------

Key Reactive Species in Output:

NO, OH, Cl, Br, Cl ₂	Species that act to kick-start BrO chemistry
SO ₃	Sulfur trioxide: direct precursor to sulphuric acid H ₂ SO ₄ (or SO ₄ ²⁻ : Sulfate)

Major Volcanic Gases in Output:

SO ₂ , HCl, HBr, CO ₂ , H ₂ O	Present in plume & in HSC output
H ₂ S, H ₂ , CO	Present in plume but missing in HSC output

3

4 * Full Matrix of Species typically included in HSC output:

5 H₂O, N₂, CO₂, SO₂, H₂, HCl, O₂, H₂S, CO, Ar, S₂, SO₃, SO, NO, HBr, COS, HS, OH, Cl, Br,
 6 S₂O, H₂S₂, Cl₂, I, HOCl, S₃, HI, HF, H, H₂SO₄, BrCl, NO₂, S, ClO, O, HO₂, Br₂, HIO, H₂O₂,
 7 HNO₂, SOCl, ICl, HCOOH, CS₂, BrO, S₂Cl, N₂O, NOCl, HSO₃Cl, IBr, SCl, S₄, IO, NOBr,
 8 COOH, HNO, NH₃, ClOO, S₅, SCl₂, CH₄, HNO₃, HCO, BrOO, CS, OCIO, O₃, I₂, ClO₂,
 9 SBr₂, HClCO, SOCl₂, ClClO, ClOCl, NOI, NO₂Cl, SO₂Cl₂, SOF, IOO, HSO₃F, ClOCl, SN,

- 1 COCl, NO₃, S₂Cl₂, OBrO, S₆, F, NBr, HOCN, HNCO, BrOBr, CH₃, ClF, HCN, COCl₂,
- 2 N₂O₂, BrF, NH₂, OIO, IF, N, BrBrO, S₂Br₂, NOF, IIO, N₂O₃, NH₂OH, SO₂ClF, SF

1 Table 2. Parameters varied in *PlumeChem* sensitivity studies

Parameter	Values
HSC $V_A:V_M$	0:100 8:92 5:95 10:90 15:85
Aerosol Loading: $\mu\text{m}^2 / \text{molec SO}_2$	
High	10^{-10}
Medium	10^{-11}
$\text{Br}_{\text{tot}}/\text{SO}_2$: molar ratio	
Medium	$7.4 \cdot 10^{-4}$
High	$2.4 \cdot 10^{-3}$
Low	$4.8 \cdot 10^{-4}$
Gas Flux kg/s SO_2	
(small variations)	10, 20
(large variations)	10, 50, 100
Wind-speed, m/s	3, 5, 10, 15
Dispersion	B, C, D
Pasquill-Gifford cases	

2

3

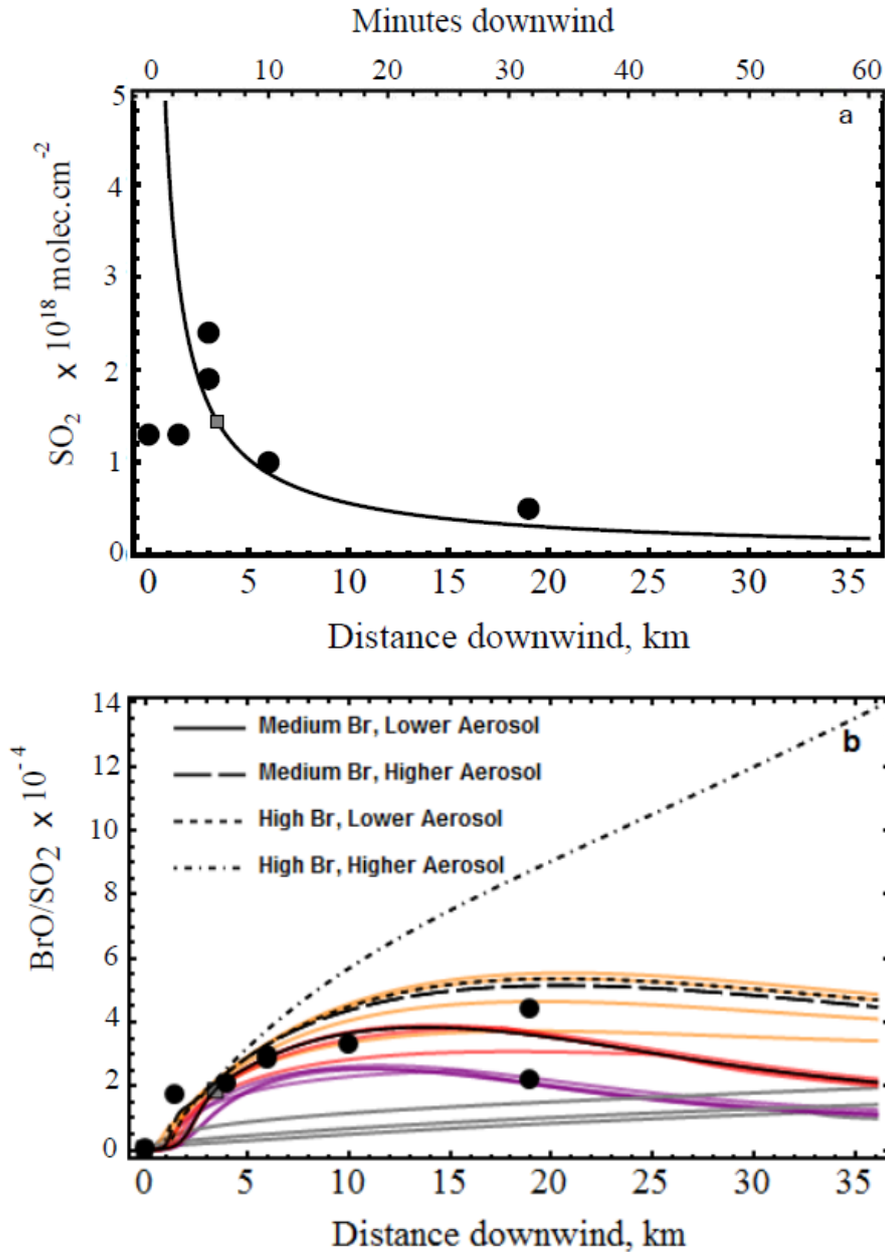
1 Table 3. List of gas-phase and photolytic reactions related to formation of BrNO₂, BrONO
 2 and BrNO. Reactions listed are used in the -BrONO-BrNO scheme. The 2-reaction BrNO₂
 3 scheme assumes BrNO₂ as the sole product from both Br + NO₂ reactions and photolysis of
 4 BrNO₂ as the only loss pathway. See text for discussion of possible additional heterogeneous
 5 pathways.

6

Reaction	Rate Coefficient	at 285 K
$Br + NO_2 \rightarrow BrNO_2$	$\sim 3.8 \times 10^{-13} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$	Brökse et al., 1998
$Br + NO_2 \rightarrow BrONO$	$\sim 4.8 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$	Brökse et al., 1998
$BrONO + Br \rightarrow Br_2 + NO_2$	$2.4 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$	Mellouki et al. 1989
$BrONO + NO_2 \rightarrow BrNO_2 + NO_2$	$\sim 2 \times 10^{-16} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ (uncertain)	Brökse et al., 1998
$BrONO \rightarrow Br + NO_2$	$\sim 1.2 \text{ s}^{-1}$ (at 298 K, 1 atm) $\tau < 1 \text{ s}$ at 298 K	Brökse et al., 1998 Orlando and Burkholder 2000
$BrONO \rightarrow BrNO_2$	unknown	-
$BrNO_2 + Br \rightarrow Br_2 + NO_2$	unknown	-
$BrNO_2 + NO \rightarrow BrNO + NO_2$	$2.3 \times 10^{-12} \text{ Exp}[-17.8/RT] \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$	Brökse et al., 1998
$BrNO_2 + NO \rightarrow BrNO + NO_2$	unknown, larger than BrNO ₂ equivalent	-
$BrNO + Br \rightarrow Br_2 + NO$	$3.7 \times 10^{-10} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ or: $5.2 \times 10^{-12} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$	Hippler et al. 1978 Grimley et al. 1980
$BrNO_2 \rightarrow Br + NO_2$	$\leq 4.0 \times 10^{-4} \text{ s}^{-1}$ $\sim 6.4 \times 10^{-5} \text{ s}^{-1}$	Brökse et al., 1998
$2BrNO_2 \rightarrow Br_2 + 2NO_2$	Unknown (slow)	Brökse et al., 1998
$BrONO \xrightarrow{h\nu} Br + NO_2$	$\tau \sim \text{s}$ (products unknown)	Burkholder and Orlando, 2000
$BrONO \xrightarrow{h\nu} BrO + NO$	or $\tau \sim \text{s}$ (products unknown)	
$BrNO_2 \xrightarrow{h\nu} Br + NO_2$	$\tau \sim \text{min}$	Scheffler et al. 1997

7

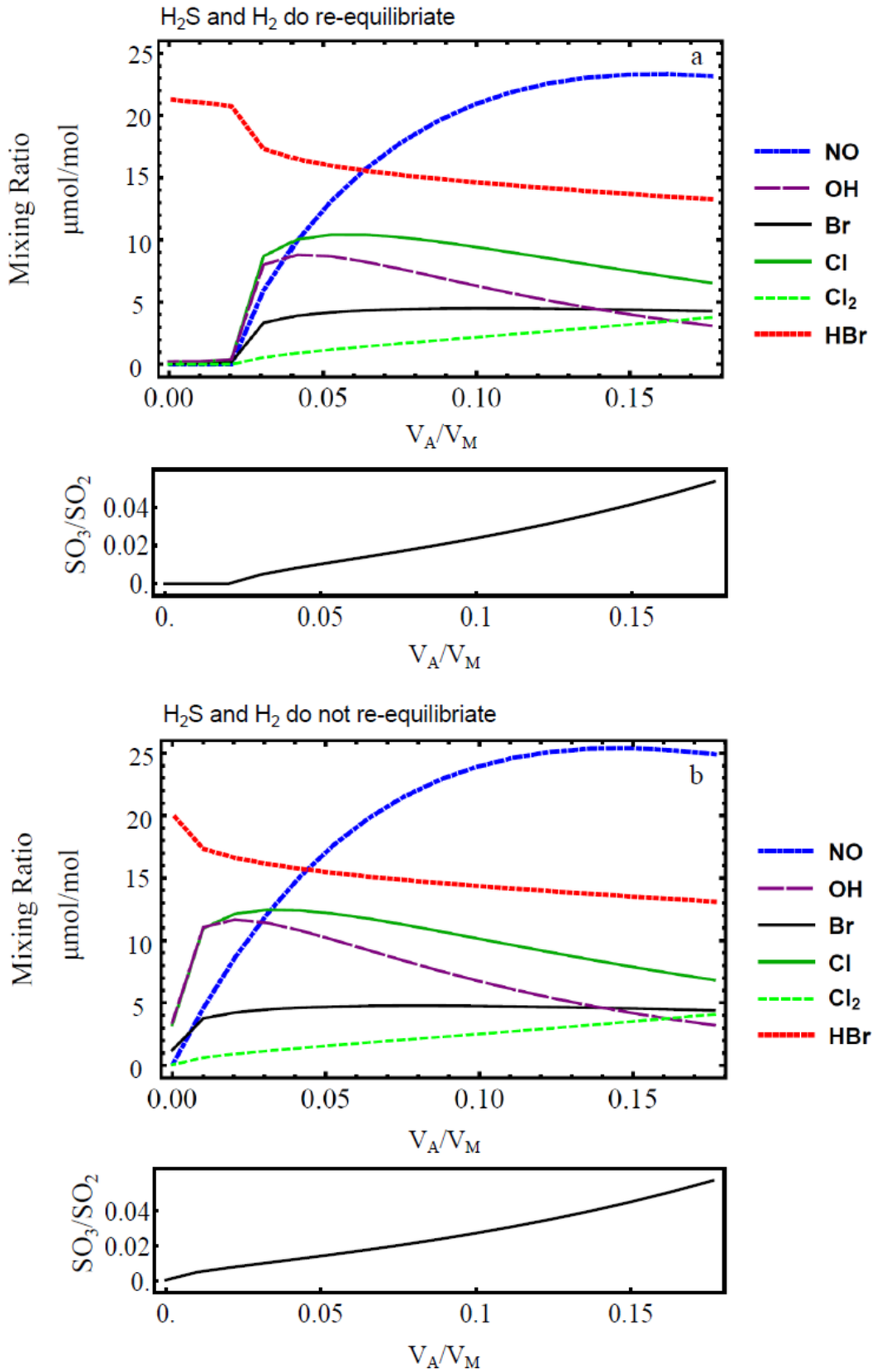
1



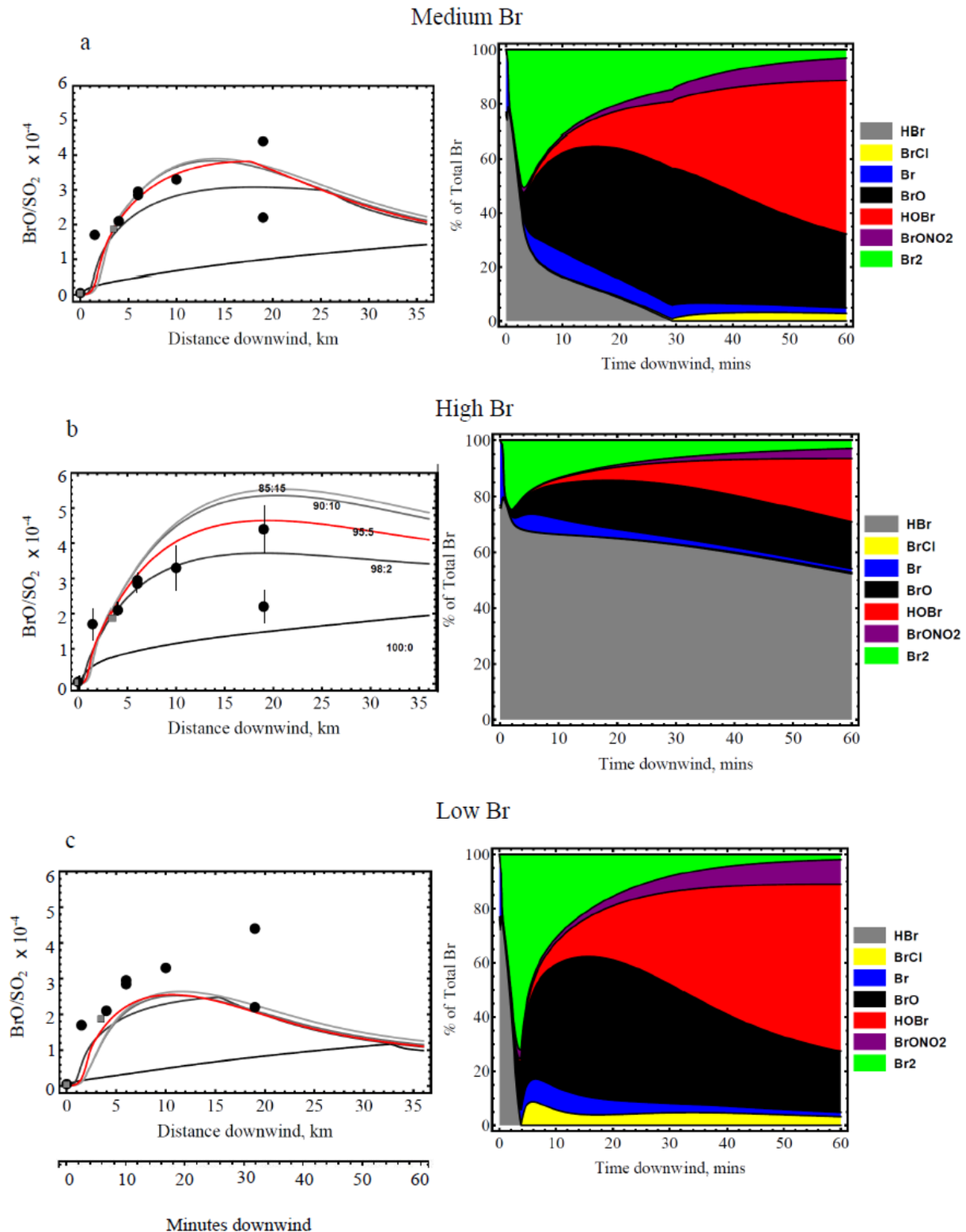
2

3 Figure 1. *PlumeChem* simulations illustrating (a) predicted SO₂ column abundance in the
 4 downwind plume (black line) according to the model dispersion parameterisation, (b)
 5 simulated downwind BrO/SO₂ ratios for model runs using this dispersion parameterisation
 6 but where: bromine in the emission ($\text{Br}_{\text{tot}}/\text{SO}_2$), volcanic aerosol loading, and the high-
 7 temperature initialisation are varied. The simulations are compared to DOAS SO₂ column
 8 abundances and (mean) BrO/SO₂ ratios reported by Oppenheimer et al., (2006), and
 9 Bobrowski et al., (2007a), gray squares and black disks, respectively. Simulations with

1 varying aerosol emission (for two bromine scenarios) are highlighted in black. Simulations
2 assuming medium aerosol loading and varying bromine emission (for a range of plausible
3 high-temperature model initialisations) are shown in red, orange and purple for medium, high
4 and low Br emission scenarios, respectively. Simulations assuming no plume-air mixing in
5 the high-temperature initialisation (VA:VM = 0:100) are shown in grey.



1 Figure 2. Mixing ratio (10^{-6} mol/mol) of key species (NO, OH, Br, Cl, Cl₂) in the HSC output
2 as a function of V_A/V_M , the assumed magmatic: atmospheric gas ratio in the near-vent plume,
3 ranging from 0 (0.00:1.00) to 0.18 (0.15:0.85). SO₃:SO₂ ratios (that prescribe the volcanic
4 sulfate/SO₂ emission) in the HSC output are also shown. (a) Standard operation of HSC in
5 which volcanic H₂S and H₂ are allowed to re-equilibrate, yielding near-zero mixing ratios of
6 these gases in the HSC output. (b) A revised operation of HSC (Martin et al., 2009) in which
7 volcanic H₂S and H₂ are removed (and temporarily replaced by inert Ar) such that they do not
8 re-equilibrate within HSC.

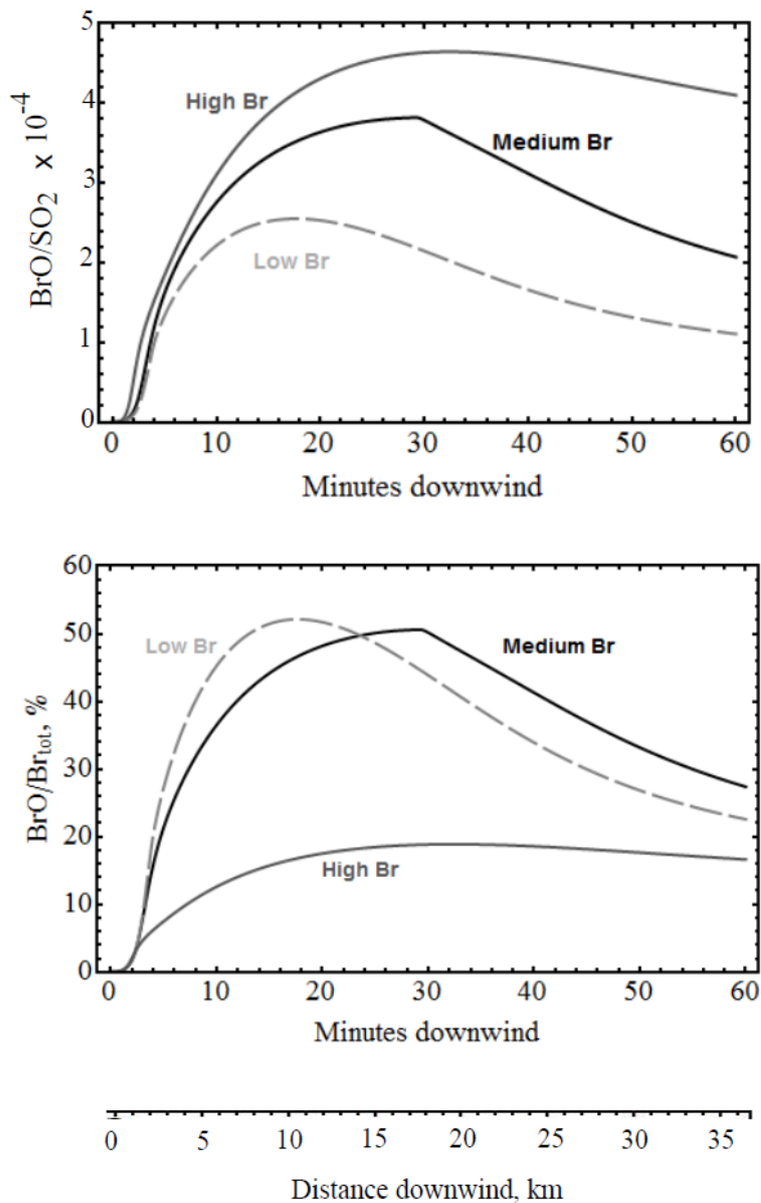


1

2 Figure 3. Left: Simulated 1 hr evolution of plume BrO/SO₂ for the three bromine emission
 3 scenarios, with varying atmospheric:magmatic gas ratio V_A:V_M (0:100, 5:95, 10:90, 15:85) in
 4 the high-temperature initialisation. Also shown are observed BrO/SO₂ ratios reported by
 5 Oppenheimer et al., (2006), and Bobrowski et al., (2007a); grey and black disks respectively,

1 with representative data error bars from Bobrowski et al., (2007a). Right: Br-speciation for
2 the three bromine emission scenarios shown for the model run initialised using HSC with
3 $V_A:V_M = 5:95$.
4

1



2

3 Figure 4. Predicted evolution in BrO/SO_2 (top) and $\text{BrO}/\text{Br}_{\text{tot}}$ ratios (bottom) over 1 hr for the
4 three different bromine emission scenarios. Model runs correspond to those shown in Figure
5 3 assuming VA:VM = 5:95 for the high-temperature initialisation.

6

2 reaction scheme (BrNO_2)

Improved scheme (BrNO_2 , BrONO , BrNO)

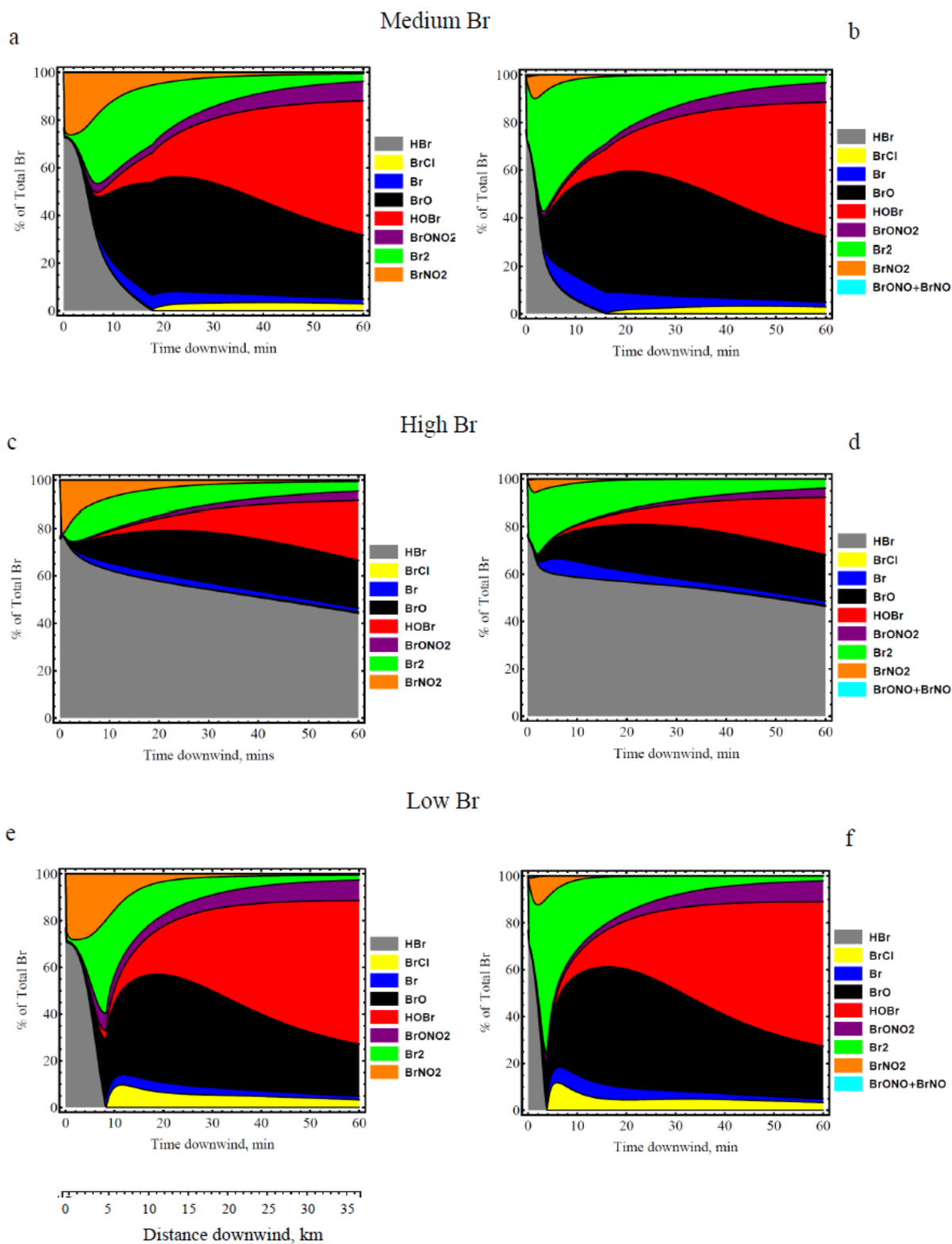
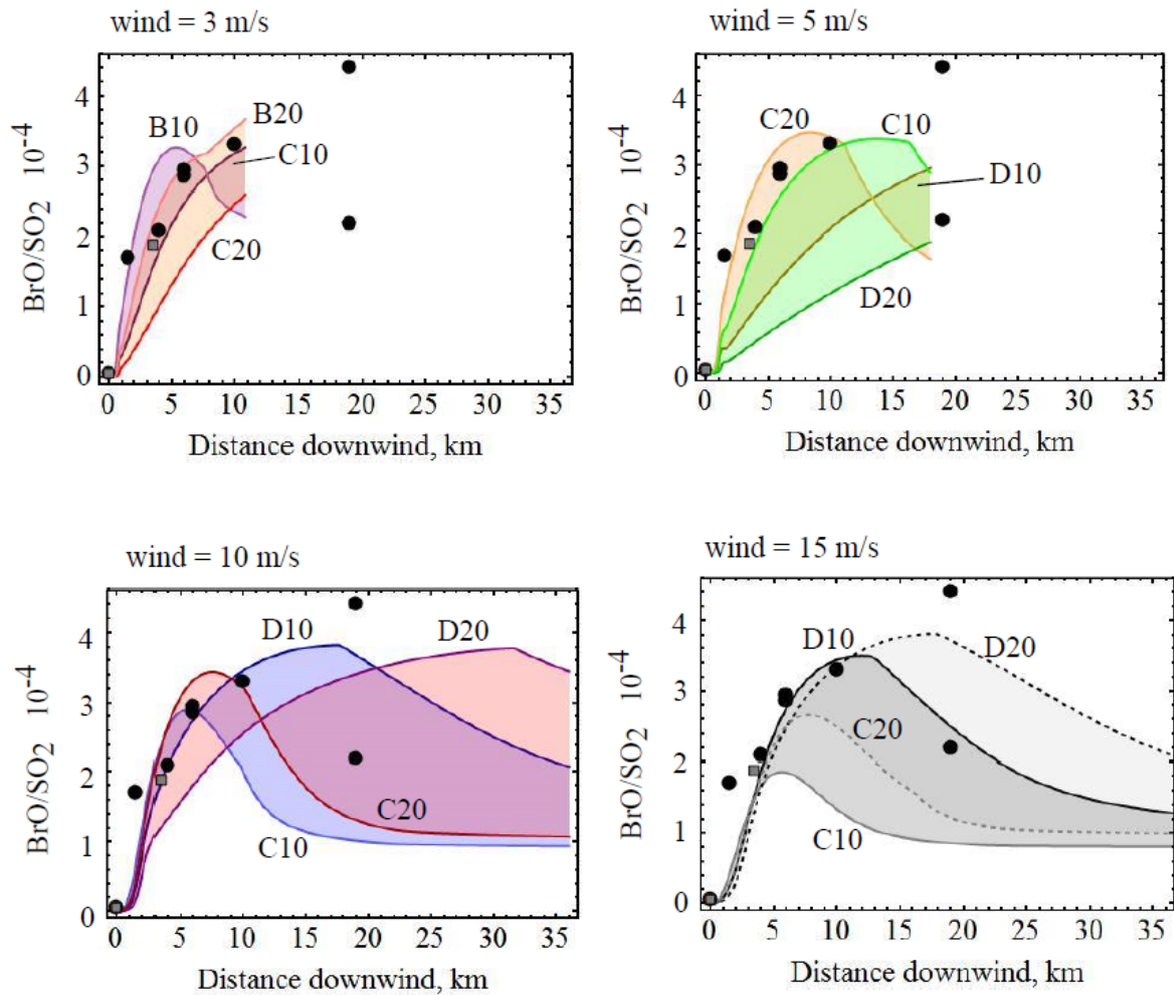


Figure 5. Br-speciation in model runs that also include formation of BrNO_2 , shown for the three bromine emission scenarios. Simulations incorporate BrNO_2 using a 2-reaction scheme

- 1 (a,c,e) or a 12-reaction scheme including BrNO_2 , BrONO and BrNO (b,d,f). See text for
- 2 details.

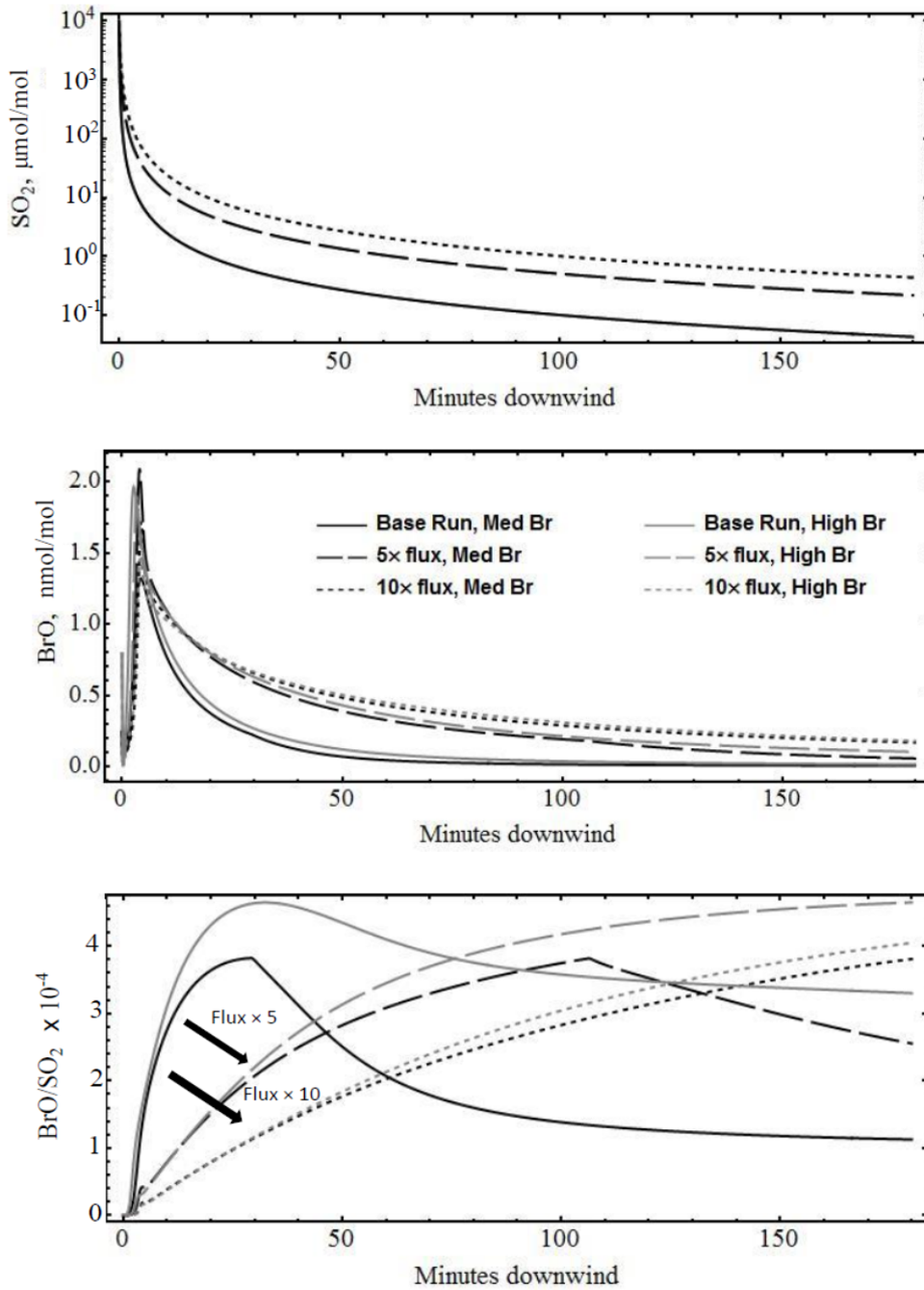


1

2 Figure 6. Simulated BrO/SO₂ over 1 h for the medium bromine emission, predicted for two
 3 emission flux scenarios (10 or 20 kg/s), and for a range of wind-speeds (3, 5, 10, 15 m/s), and
 4 Pasquill-Gifford dispersion schemes (B, C, D). See text for details of the combinations.
 5 Model runs are compared to observations from Bobrowski et al. (2007) and Oppenheimer et
 6 al. (2006), shown as black circles and grey squares, respectively.

7

1
2

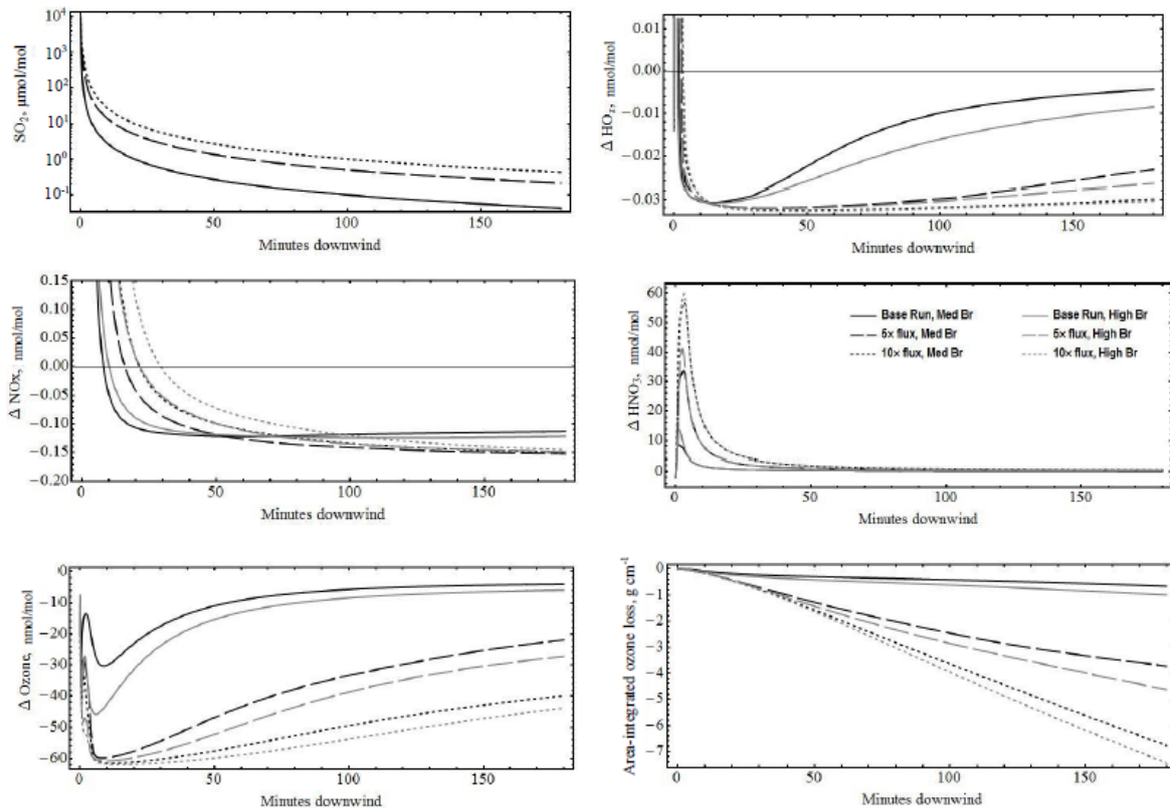


3

4 Figure 7.

5 Simulated plume SO_2 , BrO and BrO/SO_2 over 3 hours for the medium and high bromine
6 emission scenarios, and with varying volcanic emission flux (baseline run, and with volcanic
7 gas+aerosol emissions flux $\times 5$ and $\times 10$, shown by full-, long-dashed and short-dashed lines,
8 respectively), whilst keeping the same plume dispersion parameterisation, wind-speed and

- 1 initialisation (see text for model details and interpretation). Arrows highlight the reduction in
- 2 near-downwind BrO/SO₂ predicted at greater volcanic emission flux.



1
 2 Figure 8.
 3 Simulated impact of plume BrO chemistry on atmospheric oxidants, shown for the model
 4 scenarios of Figure 7. Depletion of oxidants and formation of NO_y is shown through the
 5 difference in plume – background mixing ratio for HO_x ($\text{OH}+\text{HO}_2$), NO_x ($\text{NO}+\text{NO}_2$), HNO_3 ,
 6 and ozone. Cumulative ozone loss is also calculated across the 3 hour simulations.