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The tropospheric processing of acidic gases and hydrogen sulphide in volcanic gas plumes as inferred from field and model investigations

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ACPD

6, 11653–11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◀ ▶I

■ Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

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Abstract

Improving the constraints on the atmospheric fate and depletion rates of acidic compounds persistently emitted by non-erupting (quiescent) volcanoes is important for quantitatively predicting the environmental impact of volcanic gas plumes. Here, we present new experimental data coupled with modelling studies to investigate the chemical processing of acidic volcanogenic species during tropospheric dispersion. Diffusive tube samplers were deployed at Mount Etna, a very active open-conduit basaltic volcano in eastern Sicily, and Vulcano Island, a closed-conduit guiescent volcano in the Aeolian Islands (northern Sicily). Sulphur dioxide (SO₂), hydrogen sulphide (H₂S), hydrogen chloride (HCI) and hydrogen fluoride (HF) concentrations in the volcanic plumes (typically several minutes to a few hours old) were repeatedly determined at distances from the summit vents ranging from 0.1 to ~10 km, and under different environmental conditions. At both volcanoes, acidic gas concentrations were found to decrease exponentially with distance from the summit vents (e.g., SO₂ decreases from $\sim 10\,000\,\mu\text{g/m}^3$ at 0.1 km from Etna's vents down to $\sim 7\,\mu\text{g/m}^3$ at $\sim 10\,\text{km}$ distance), reflecting the atmospheric dilution of the plume within the acid gas-free background troposphere. Conversely, SO₂/HCI, SO₂/HF, and SO₂/H₂S ratios in the plume showed no systematic changes with plume aging, and fit source compositions within analytical error. Assuming that SO₂ losses by reaction are small during short-range atmospheric transport within quiescent (ash-free) volcanic plumes, our observations suggest that, for these short transport distances, atmospheric reactions for H₂S and halogens are also negligible. The one-dimensional model MISTRA was used to simulate quantitatively the evolution of halogen and sulphur compounds in the plume of Mt. Etna. Model predictions support the hypothesis of minor HCl chemical processing during plume transport, at least in cloud-free conditions. Larger variations in the modelled SO₂/HCl ratios were predicted under cloudy conditions, due to heterogeneous chlorine cycling in the aerosol phase. The modelled evolution of the SO₂/H₂S ratios is found to be substantially dependent on whether or not the interactions of H₂S with halogens are

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

included in the model. In the former case, H_2S is assumed to be oxidized in the atmosphere mainly by OH, which results in minor chemical loss for H_2S during plume aging and produces a fair match between modelled and measured SO_2/H_2S ratios. In the latter case, fast oxidation of H_2S by CI leads to H_2S chemical lifetimes in the early plume of a few seconds, and thus SO_2 to H_2S ratios that increase sharply during plume transport. This disagreement between modelled and observed plume compositions suggests that more in-detail kinetic investigations are required for a proper evaluation of H_2S chemical processing in volcanic plumes.

1 Introduction

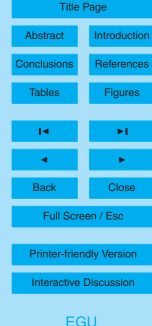
Volcanic plumes are peculiar chemical/physical systems formed by the atmospheric dispersion of gases and particles released from both eruptive and quiescent volcanoes. While eruptive stratospheric plumes and their atmospheric impacts have been extensively studied in the past (e.g., Robock, 2000; Robock and Oppenheimer, 2003), the impacts of tropospheric emissions from quiescent volcanoes are far less well characterised. However it is becoming clear that these less dramatic but persistent emissions can have local to regional environmental impacts which may be at least as important as those of short-lived volcanic eruptions in the long-term (e.g. Delmelle, 2003; Mather et al., 2003a).

A matter of particular interest is to assess the paths and rates of chemical reactions in tropospheric volcanic plumes; these may be significantly different to those in the background atmosphere, reflecting higher temperatures, humidity, aerosol content and possibly radical production (Gerlach, 2004). Besides CO₂, which is likely to be relatively inert during atmospheric dilution, volcanic degassing releases large amounts of acidic gaseous volatile species such SO₂, H₂S and hydrogen halides (HCI, HF, HBr) (Symonds et al., 1994). As they are emitted from the vent, cooled and mixed in with atmospheric gases, these reactive species are available to take part in both homogeneous (gas-only) and heterogeneous (gas-liquid-solid) reactions. Although these reac-

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases



tions have been studied in detail in the laboratory, uncertainties remain when applying such results to the conditions in volcanic plumes. Recently, evidence for rapid reaction following emission, or during transport, has been found for HBr, which is at least partially converted to BrO following emission (Bobrowski et al., 2006; Oppenheimer et al., 2006).

The recent detection of CIO and OCIO in volcanic plumes (Lee et al., 2005; Bobrowski et al., 2006) suggests that similar reaction mechanisms (Martin et al., 2006) may also apply to emitted HCI.

A proper evaluation of the reaction schemes involved is of fundamental importance for assessing the residence time and thus deposition rates of these acid compounds, and for quantitatively predicting the environmental impact of volcanic gas plumes. In addition, since the determination of volcanic gas composition is increasingly accomplished by remote-sensing spectroscopic methods, knowledge of atmospheric processes within volcanic plumes is required to verify the extent to which the composition of distal plumes, sampled many km downwind from a volcanic vent, is representative of the source composition.

The ultimate fate of much of the sulphur dioxide in volcanic plumes is thought to be conversion into sulphate aerosols (its other sinks are dry and wet deposition), which can occur via gas phase reactions (mostly via reaction with OH radical) or more probably on the surface of solid/liquid particles in the plume (Penkett et al., 1979; Ravinshankara, 1997). Although the rates of such reactions can vary significantly depending on temperature, humidity, insulation, and aerosol loading (Martin et al., 1986; Oppenheimer et al., 1998), first order sulphur dioxide loss rates have recently been estimated at only ~4% per hour in ash-free non-condensing volcanic plumes (McGonigle et al., 2004). Hydrogen chloride and hydrogen fluoride are more soluble in water than sulphur dioxide, and hence wet deposition may cause major depletion of these species from the plume (especially during the night or in wet periods). Still, Horrocks et al. (2003) failed to detect any change in the plume SO₂/HCl ratio at Masaya volcano (Nicaragua), taking FTIR measurements simultaneously at the vents and downwind of the volcano

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



(up to 40 km from source, with plume ages of up to ~1.5 h), suggesting that there is no differential influence of tropospheric processes on plume SO₂ and HCl, at least under the conditions prevalent at a low altitude (600 m a.s.l.), tropical volcano (11.98° N) such as Masaya. However, their measurements were limited to the drier season (December to April) in Nicaragua and during day time, when relative humidity would have been at a minimum.

Here, we test further Horrocks et al.'s (2003) suggestion that the SO₂/HCl ratio in volcanic plumes is not altered by tropospheric processing over distances of tens of kms by presenting results from a series of experiments carried out on Mount Etna volcano (Sicily, Southern Italy) in the 2002-2005 period. Sulphur dioxide, HCl and HF timeaveraged concentrations in the plume were determined repeatedly by the use of a network of diffusive tubes (Aiuppa et al., 2004) at distances from the summit vents ranging from 0.1 to ~10 km. Measurements were made during different seasons, ranging from cold and rainy periods (October-November 2003) to hot and dry periods (e.g., August 2004). By so doing, we attempt to evaluate the extent to which SO₂/HCl and SO₂/HF ratios are altered during atmospheric transport in a tropospheric volcanic plume under a wide range of environmental conditions. We also present the first distal "diffusive" determinations of H₂S concentrations in plumes from Mount Etna and Vulcano Island, allowing us to assess its residence time compared to SO₂. Finally, the comparison of plume data with the model results from the MISTRA one-dimensional atmospheric reaction code (von Glasow et al., 2002) is used to better evaluate the chemical processing of halogen and sulphur compounds in Mt. Etna's plume.

1.1 Introduction to field sites

Mount Etna (Fig. 1) is the largest (summit 3.3 km a.s.l.; base 60×40 km) and most-active strato-volcano in Europe (Bonaccorso et al., 2004). It is an excellent test-site for plume-chemistry investigations, because of the un-interrupted open-conduit passive degassing from four summit craters and the massive release of SO_2 ($\sim 5500 \, \text{T d}^{-1}$; Caltabiano et al., 2004), halogens (Francis et al., 1998; Aiuppa et al., 2005a) and H_2S

11657

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



(Aiuppa et al., 2005b). Although Etna's plume is generally dispersed by winds at the same altitude as the emission point, it is sometimes forced by the prevailing westerly winds (Fig. 1) to flow at ground level down the volcano's slope, leading to fumigation of its upper eastern flank (Aiuppa et al., 2004; Allen et al., 2006).

Vulcano is a closed-conduit quiescent volcano in the Aeolian Islands (Fig. 2). The volcanic plume is fed by degassing from many fumaroles ($T_{max} \sim 450^{\circ}C$), with mean SO_2 and H_2S emissions of ~ 15 and $\sim 6 T d^{-1}$, respectively (Aiuppa et al., 2005b, c).

2 Methods

2.1 Measurement techniques

Diffusive tubes have recently been shown to provide valuable information on the plume dispersion of volcanic acidic gases (Delmelle et al., 2001, 2002; Allen et al., 2002; Aiuppa et al., 2004). Diffusive tubes are passive air sampling devices that rely on the molecular diffusion of the species of interest through an entrapped air volume. A chemically-specific sorbent (an impregnated filter designed to selectively react with and capture a certain gaseous species) is placed in the closed end of an open tube, and an inert filter positioned at the inlet end to prevent turbulent transport of gases within the device, or ingress of particles. Uptake of the species of interest by the sorbent creates a concentration gradient inside the tube and the net flux of the sorbed gas is proportional to the ambient concentration. The average ambient concentration can hence be calculated from the amount of the gaseous species that is captured and the exposure time using Fick's first law of diffusion (Ferm, 2001).

The diffusive tubes used in this study for the determination of SO_2 , HCl and HF were polypropylene tubes using a paper filter impregnated with NaHCO $_3$ in glycerol as sorbent (see Aiuppa et al., 2004 for more details). After exposure, the filter was leached in water and analyzed by ion chromatography to determine the amounts of F^- , Cl^- and SO_4^{2-} accumulated. The average HF, HCl, and SO_2 concentrations in

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



the air (precisions, ±20%, ±10% and ±5%, respectively) during the sampling periods were then calculated (see Ayers et al., 1998 for further details). The diffusive tubes used for H₂S determination (Radiello® from Fondazione Maugeri) were of the radial sampling geometry and used a porous polyethylene cartridge impregnated with zincacetate as a sorbent (for details, see http://www.radiello.it/english/h2s_en.htm). The sorbed sulphide was later recovered by extraction with water, mixed with N,N-dimethyl-p-phenylenediammonium ion to yield methylene blue (measured by visible spectro-photometry).

Four field campaigns were carried out on Etna, in May–June 2002, October–November 2003, July–August 2004 and July–August 2005. In each campaign, about 20 diffusive tubes were simultaneously exposed for about 1 month around the volcano (Fig. 1). One sampling campaign was carried out during June–July 2005 at Vulcano Island (Fig. 2). Hydrogen sulphide concentrations were measured at a sub-set of sampling sites during the 2004 and 2005 campaigns only (Fig. 1).

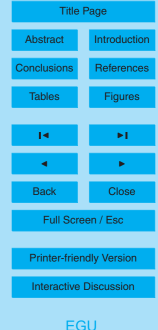
2.2 Model description

We used the one-dimensional model MISTRA to study the evolution of halogen and sulphur compounds in the plume of Mt. Etna. The model version that we used has been described in more detail in Bobrowski et al. (2006) and is an extension of the model presented in von Glasow et al. (2002). In brief, the model includes gas phase, aerosol and cloud droplet chemistry with a focus on halogen and sulphur chemistry. Photolysis frequencies are calculated online. Vertical mixing is treated in detail in the model, whereas horizontal mixing and entrainment of ambient air into the volcanic plume is parameterized. In this paper we discuss model runs that are based on the setup that showed best agreement with measurements at Mt. Etna of BrO and SO₂ and BrO/SO₂ (run "60–40", see Bobrowski et al., 2006). The initial composition of volcanic volatiles has been calculated with a thermodynamic equilibrium model under the assumption of a mixture of 60% volcanic volatiles and 40% ambient air at about T=600°C. Model start is at local noon and the duration of the runs is 6 h. For more details on the model and

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases



the basic setup of the runs for the modelling of volcanic plumes the reader is referred to Bobrowski et al. (2006).

In Sect. 3.2 we will discuss results from a model run where the ambient humidity has been set to 60% so that no cloud forms in the model after release of the volcanic volatiles ("cloud-free") and a second run where the ambient humidity was set to 95% resulting in cloud formation after plume release ("cloudy"). This cloud has a vertical extent of 10–50 m and is present for 70 min after plume release, which corresponds to 42 km at a wind speed of 10 m/s (see Fig. 6).

We have also conducted model runs where we included the oxidation of H₂S.

3 Results and discussion

3.1 Plume measurements

Figure 3 is a plot of SO_2 concentrations in air at the locations shown in Fig. 1 against the relative distances from Etna's summit craters. This clearly demonstrates the volcanic origin of SO_2 , since the concentrations decrease exponentially with distance from Etna's summit vents, from $\sim 10\,000\,\mu\text{g/m}^3$ at $0.1\,\text{km}$ to $\sim 7\,\mu\text{g/m}^3$ at $\sim 10\,\text{km}$ distance. Assuming that SO_2 is virtually inert during short-term plume dispersal (McGonigle et al., 2004), this trend reflects an increasing atmospheric dilution of the plume within the SO_2 -free background troposphere, and further supports the idea of recurrent plume fumigation in an area $\sim 10\,\text{km}$ in diameter centred east of the volcano's summit (Fig. 1), as first proposed by Aiuppa et al. (2004). The ~ 100 -fold concentration contrast between "upwind" and "downwind" sites on Etna's flanks, confirms that plume fumigation occurs less often around Etna's western flank.

Hydrogen chloride and HF concentrations display similar decreasing (e.g., dilution) trends over the same distance interval, ranging from $\sim\!800$ and $\sim\!900\,\mu\text{g/m}^3$, respectively, at 0.1 km from the vents, to below detection limit (0.6 and 0.35 $\mu\text{g/m}^3$, respectively) at 10 km distance.

ACPD

6, 11653–11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



Molar SO₂/HCl and SO₂/HF ratios in air at the diffusive tube locations of Fig. 1 are shown in Fig. 4. Large variations of the ratios are observed between one campaign and another, reflecting important compositional changes at the source over time: the higher ratios in the 2002 campaign are consistent with the larger SO₂/HCl and SO₂/HF 5 ratios in the near-vent summit craters' plume at that time (4.7 and 85, respectively; Table 1). SO₂/HCl and SO₂/HF in the source plume were lower (and more homogeneous) during the 2003, 2004 and 2005 campaigns (Table 1). Equally, data from each campaign consistently show that there are no systematic changes (within the measurement errors) in the SO₂/HCl ratios with distance (i.e. with plume aging) from the vents (Fig. 4). The mean SO₂/HCl ratios (Table 1) fit source compositions within analytical uncertainty for each of the four campaigns, and suggest that, for these short transport distances, atmospheric reactions during plume transport for both species are either negligible or of the same rate. The scatter of SO₂/HF ratios for each campaign is larger than for SO₂/HCl ratios, as is the discrepancy from the source's composition (Table 1). However, there is still no apparent systematic distance-ratio dependence, and the scatter in the dataset seems to be more likely related to analytical uncertainties in fluoride determination, with concentrations often being very close to the detection limit. These observations are consistent with the usefulness of diffusive samplers for volcanic monitoring (Aiuppa et al., 2004).

Hydrogen sulphide concentrations, measured in a sub-set of Etna's locations during the 2004 and 2005 campaigns, are plotted against corresponding SO_2 concentrations in Fig. 5. They range between $0.3-281\,\mu\text{g/m}^3$ and are positively correlated with SO_2 concentrations. SO_2/H_2S ratios are highly scattered (Table 1), reflecting a combination of poor analytical accuracy of H_2S determinations (8.7% at 2σ ; http://www.radiello.it/english/h2s_en.htm), multiple H_2S sources (fumarolic vs main vents, or natural vs anthropogenic), or complex atmospheric chemistry. Nevertheless, mean SO_2/H_2S ratios from diffusive tubes for both campaigns (54 and 60, respectively) are in reasonable agreement with the source plume composition (\sim 60; Table 1). Data from the Vulcano Island campaign (Fig. 5), showing consistent SO_2/H_2S ratios

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Title Page



in source (2.2±1.1; Aiuppa et al., 2005c) and distal (2.2±1.2, this study) plumes, also point to only minor alterations of sulphur speciation during medium-term plume transport (for plume ages of ~30 min, assuming a plume travelling speed of ~10 km/h). These results extend the earlier conclusions by Aiuppa et al. (2005b), who demon-5 strated that hydrogen sulfide is essentially inert in volcanic plumes on timescales of seconds to minutes.

Model results

3.2.1 Halogen and SO₂ chemistry

The basic features of the plume differ in the cloud-free and cloudy model runs, as can best be seen in the evolution of SO₂. The evolution of liquid water content of the cloud developing at the crater in model run "cloudy" is shown in Fig. 6. In the cloudfree run (Fig. 7, top panel), the plume rises due to the initially high temperature of the plume (~600°C) and radiative heating due to absorption by the volcanic aerosols. In the cloudy run (Fig. 8, top panel), the plume descends due to long-wave cooling of the cloud-top and by additional cooling caused by the evaporation of cloud droplets. For numerical reasons we were not able to prescribe the temperature of the fresh plume in the cloudy run to ~600°C as in the cloud-free run, but only to ambient temperatures, therefore the plume descent is somewhat stronger than it would be if thermal buoyancy effects had been included.

Due to plume dispersal, the concentration of SO₂ in the core of the plume is higher than at the edges. The smaller concentrations at the plume base in the model are in the range measured with the diffusive tubes, which is consistent with the fact that the plume edges will be more likely to be in contact with the ground (where the diffusion tubes are located) than the core of the plume. At the altitude of Mt. Etna's summit a concentration of $1 \mu g(SO_2)/m^3$ roughly corresponds to a mixing ratio of 5×10^{-10} mol/mol.

In order to compare the model results with the diffusive tube measurements presented in this paper, we show the evolution of the SO₂/HCl ratio as a function of time

11662

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Introduction

References

Figures



and altitude for both the cloud-free and the cloudy run (see bottom panels of Figs. 7 and 8). The SO₂/HCl ratio in the cloud-free runs is constant for approximately the first 130 min after plume release and it averages at about 2, very close to the measured ratios. This matching between modelled and measured ratios further supports 5 only minor hydrogen chloride partitioning into the liquid/solid aerosol phase, at least in cloud-free conditions and in the short-term (minutes). Although fluorine chemistry was not included in the modelling, we propose that similar considerations also apply to even less-reactive HF. Starting ~130 min after plume release, the SO₂/HCl ratio rapidly increases in the model (note the strongly non-linear increments of the contour lines in Fig. 7), which is due to heterogeneous cycling of HCl on aerosols: after this time, plume dilution has decreased the gas phase concentrations enough to make uptake onto the aerosol phase an important sink for HCl. It should be noted, however, that model CIO vertical columns underestimate the measured columns by about a factor of 40 in the early plume (see Bobrowski et al., 2006), so that some uncertainty remains as to whether chlorine chemistry is properly reproduced in the model. In the case of HCl, however, it is likely that the model captures its chemistry with some accuracy, as it is fairly un-reactive in the gas phase and its aqueous fraction is small due to the high acidity of the aerosol particles until the gas phase concentration is reduced by dilution. This is also supported by field measurements of night- (cloudy plume with RH ≥88%) and daytime (transparent) plume SO₂/HCl ratios in the undiluted plume at the crater rim of Masaya volcano, Nicaragua (Mather et al., 2003b).

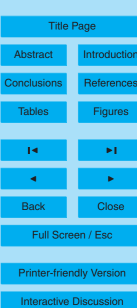
The evolution of the SO₂ to HCl ratio in the cloudy run (see Fig. 8) is different from that in the cloud-free run. The ratio is fairly high in the cloud layers due to uptake of HCl in the cloud droplets, which contain a lot more water than aerosol particles. The lower, cloud-free part of the plume in these runs would seem more likely to ground though, and in this region the modelled ratio is consistent with the measurements. Even if the diffusion tubes are within a "grounded cloud", during part of the sampling time one can expect a mixture of the modelled "cloud-free" and "cloudy" conditions during sampling with the diffusion tubes which would still be consistent with the field

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



data. These modelling results are in agreement with the relatively minor chlorine bulk deposition fluxes measured over the volcano perimeter, accounting for \leq 1% of total emissions from the volcano (Aiuppa et al., 2006), which argue against a substantial HCl removal from the plume.

5 3.2.2 H₂S chemistry

We already demonstrated elsewhere (Aiuppa et al, 2005b) that the measured SO_2/H_2S ratios in the short-lived Etna's plume (plume ages of seconds to a few minutes) reflect thermodynamic equilibrium at magmatic conditions ($T\sim1100^{\circ}C$; $P\sim1-10$ bars) and redox conditions buffered by the coexisting silicate melt at about the Nickel-Nickel oxide (NNO) buffer ($fO_2\sim10^{-9}$ bar at 1150°C; Metrich and Clocchiatti, 1996). Conversely, our gas-only thermodynamic equilibrium calculations with the software HSC (Outokumpu Tech., Finland) show that under "60-40, T=600 K" conditions, the concentration of H_2S in the early plume should be negligible, due to much higher oxygen fugacity ($fO_2\sim0.08$ bar) in the air-volcanic gas mixture. This is in stark contrast to our measurements, and implies that the emitted H_2S is not fast-oxidized upon cooling and dilution in the high-temperature volcanic gas-air mixing region. Thus the assumption of thermodynamic equilibrium in the volcanic volatiles-ambient air mixture does not hold for H_2S . We therefore assume that the H_2S concentration equals $SO_2/60$ in the fresh plume, and this was used as input data for further modelling with MISTRA code.

Typically, H_2S is assumed to be oxidized in the atmosphere mainly by OH. Including only this reaction as chemical loss for H_2S our model results very closely reproduced the measured ratio of SO_2/H_2S (see Fig. 9, top panel). As model ozone concentrations are strongly reduced in the early plume, the resulting OH concentrations are very small (see also discussion in Bobrowski et al., 2006), therefore one would expect H_2S to have a long chemical lifetime in early volcanic plumes. The halogen radicals Cl and Br, however, react very rapidly with H_2S as well, leading to the production of the hydrogen sulphide radical, SH, as in the case of H_2S + OH. The rate coefficient for the reaction H_2S + Cl is fairly well known (Sander et al., 2006) but only one measurement has

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◆ I

Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

been reported for H_2S + Br (Nicovich et al., 1992). In line with recent measurements of halogen oxides (CIO, OCIO, BrO) in volcanic plumes (Bobrowski et al., 2003, 2006; Lee et al, 2005), we also included the interactions of H_2S and its break-down products with halogens in the model.

SH, the first break-down product of H_2S , can be reduced back to H_2S by self reaction and by reaction with HBr (Nicovich et al., 1992) and according to the model in the first few minutes after plume release the reduction by HBr is indeed more important than oxidation of H_2S by Br. The oxidation of H_2S by CI, however, is very fast leading to H_2S chemical lifetimes in the early plume of a few seconds and a drastically changing SO_2 to H_2S ratios (see Fig. 9, bottom panel). This is an obvious contradiction to our measurements (Fig. 5). According to thermodynamic equilibrium calculations of the composition of the initial plume, there are significant amounts of atomic halogens in the mixture of volcanic volatiles and ambient air (see also Gerlach, 2004; Martin et al., 2006), so that the fast oxidation of H_2S by CI should also proceed at night.

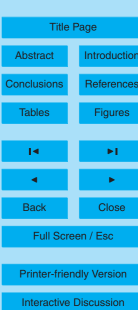
Possible explanations for the mismatch between modelled and field-measured SO_2/H_2S ratios could be that our model strongly overestimates the CI concentrations in the plume. However, again it should be noted that a comparison of model results with observed column densities of CIO, a species closely linked to CI, showed an underestimate of the CIO columns in the model. Even though we cannot exclude the possibility that we have neglected important chlorine reactions in the model that might lead to smaller chlorine atom concentrations, this is unlikely to yield a sufficiently different CI/CIO ratio to be consistent with the measurements of high CIO columns (assuming that the spatial gradients are not extraordinarily high). Another possible explanation would be rapid reduction of SH by HCl back to H_2S , as is the case for bromine. For chlorine, however, this reaction is endothermic. Another complication with regard to H_2S – halogen links is the fact that the hydrogen sulphide radical is known to react rapidly with dihalogens (CI_2 , BrCI, Br_2) producing XSH (Sander et al., 2006). The fate of XSH, however, remains unknown and one can only speculate about reaction pathways and rate constants that would lead to a possible reformation of H_2S .

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



There is currently a significant lack of information regarding the possible reactions that may be involved in halogen $-H_2S$ links in volcanic plumes so we unfortunately cannot use the model to explore our observations in more detail. Further study of the possible reactions that might be involved and their kinetics is to be encouraged.

4 Conclusions

Our combined experimental and modelling investigations support volcanogenic HCl and HF being essentially inert (their chemical processing being negligible compared to their reservoir within the plume) during short-term plume transport, as already hypothesised for SO₂ (McGonigle et al., 2004). We suggest that HCl heterogeneous cycling into the aerosol phase, which is expected to be the main atmospheric sink for chlorine under plume conditions, is relatively minor under low relative humidity conditions, when chlorine partitioning into the liquid phase is limited by the high acidity of the aerosol particles (at least until the gas phase concentration is reduced by dilution). Chlorine heterogeneous cycling may be more effective under cloudy conditions, due to HCl uptake in cloud droplets. However, even under these conditions, the virtually-constant SO₂/HCl plume ratios measured during rainy periods (e.g., October 2003), and the relatively minor chlorine bulk deposition fluxes measured over the volcano perimeter, accounting for ≤1% of total emissions from the volcano (Aiuppa et al., 2006), argue against a substantial HCl removal from the plume. Our measurements also show that SO₂/H₂S ratios are constant (within analytical uncertainty) for plume travel distances of up to several kilometres from the summit vents, suggesting that H₂S is also conserved during short-range transport (i.e. for plume travel times of tens of minutes). This suggestion is supported by model calculations only when H₂S-halogens interactions are neglected. At this stage we can only pose open questions with regard to H₂S in the plume but are unable to answer them without further information about the possible reactions involved: (1) Which reactions are able to preserve H₂S during cooling and dilution within the high-temperature (>600°C) volcanic gas-air mixing region? (2) What

ACPD

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Title Page



is the fate of XSH? (3) What other possibly halogen related reactions could be able (like HBr) to reduce SH to H₂S?

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ACPD

6, 11653–11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Title Page Introduction Abstract Conclusions References **Tables Figures** I⋖ Back Close Full Screen / Esc **Printer-friendly Version** Interactive Discussion

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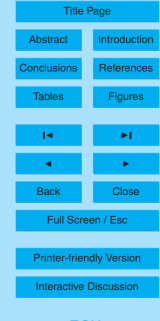
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6, 11653–11680, 2006

The tropospheric processing of acidic gases



Table 1. Mean SO_2/HCI , SO_2/HF and SO_2/H_2S molar ratios from diffusive tubes for the four Etna campaigns. Standard deviations are given in parentheses. For comparison, the characteristic molar ratios in the near-vent source plume are given for each campaign (as averages of the periodic measurements performed on Etna's vents during each exposure period).

Campaign	SO ₂ /HCI		SO ₂ /HF		SO ₂ /H ₂ S	
	Source	Diffusive	sourcea	Diffusive tubes	Source	Diffusive
	plume ^a	tubes		plume ^b	tubes	
2002	4.7	4.9 (0.9)	90	90 (26)	_	_
2003	1.0	1.0 (0.2)	21	16 (5)	_	_
2004	1.5	1.7 (1.0)	20	16 (6)	~60	60 (55)
2005	2.2	2.2 (0.5)	10	12 (3)	~60	54 (20)

 $^{^{\}rm a}$ Measurements of acidic gases in the crater's plume by filter packs (Aiuppa et al., 2005a) and this study.

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Title Page

Abstract	Introduction			
Conclusions	References			
Tables	Figures			
I◀	►I			
4	•			
Back	Close			
Full Screen / Esc				
Printer-friendly Version				
Interactive Discussion				

^b Measured by specific electrochemical sensors according to the method described in Aiuppa et al. (2005c), and assuming equal (1:1) sulphur contributions from North-east crater (SO_2/H_2S ~20) and Voragine crater (SO_2/H_2S ~100).

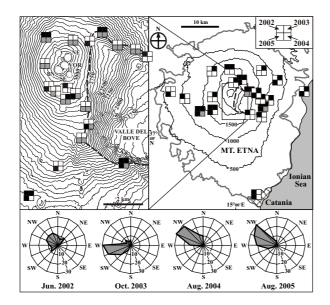


Fig. 1. Location of the sites where diffusive tubes were exposed on Etna during the different measurement campaigns. The bold grey line limits the outcrop area of Etnean volcanic products, while the dashed line shows the rim of the Valle del Bove. The summit craters are indicated with different initials: NE – North-East Crater; SE – South-East Crater; BN – Bocca Nuova; VOR – Voragine. Diffusive tubes were exposed for about one month in four campaigns on May–June 2002, October–November 2003, July–August 2004 and July–August 2005. For each sampling site, the symbol is subdivided into four sectors, each referring to a different survey. A white sector indicates no sampling; a black sector indicates sampling of SO₂ and halogens; a grey sector sampling of SO₂, halogens and H₂S. The wind rose diagrams showing the frequency of wind provenance (directions from which the wind originated are marked) at 700 hPa during the four surveys are also shown – as based on Trapani Birgi atmospheric sounding data (http://www.weather.uwyo.edu/upperair/sounding.html).

6, 11653–11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



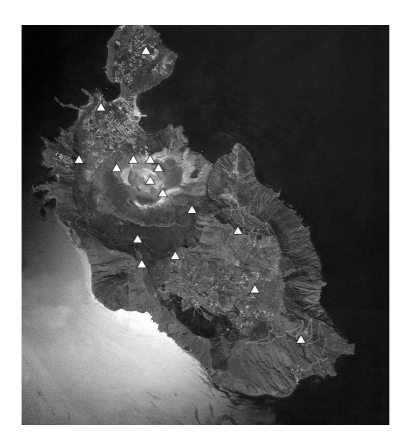


Fig. 2. Location of the sites of exposure of diffusive tubes on Vulcano Island during June–July 2005.

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ← ►I

← ► Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

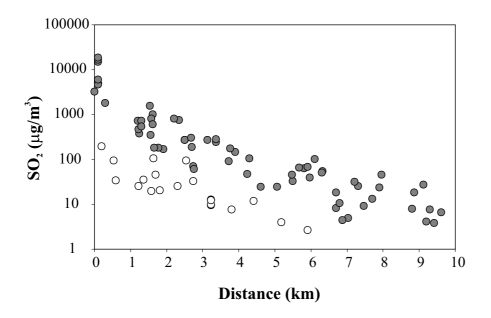
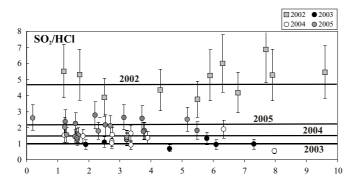


Fig. 3. SO_2 concentrations in air (in μ g/m³) versus the corresponding distance from Etna's summit craters. Data for the four Etna campaigns are integrated. White circles, sites upwind of Etna's summit vents; grey circles, sites downwind of Etna's summit vents. Each site was assigned as upwind or downwind based on the rose diagrams of Fig. 1.

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



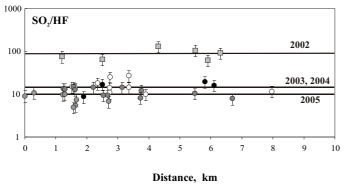
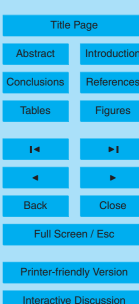


Fig. 4. SO_2/HCl and SO_2/HF molar ratios in air at the diffusive tube locations of Fig. 1, shown against the corresponding distance from Etna's summit vents. The horizontal solid lines indicate - for each campaign – the mean SO_2/HCl and SO_2/HF molar ratios in the near-vent source plume (Table 1). The diagram demonstrates contrasting compositions between the 2002 campaign (where higher SO_2/HCl and SO_2/HF were measured) and 2003–2005 campaigns. It also points to minor variations (within analytical uncertainty) of ratios with distance (e.g., with increasing plume travel times) in each campaign.

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



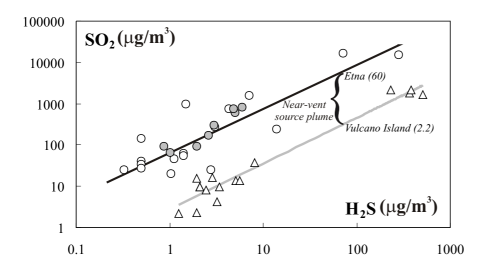


Fig. 5. Scatter diagram of SO_2 versus H_2S concentrations (in $\mu g/m^3$), measured at a sub-set of Etna's locations (Fig. 1) in 2004 and 2005 campaigns (symbols as in Fig. 4). Data for the 2005 Vulcano Island campaign are also shown (triangles). For both Etna and Vulcano Island, the characteristic SO_2/H_2S ratios of the near-vent source plumes (numbers in parentheses; see Table 1) are indicated by the solid diagonal lines.

6, 11653-11680, 2006

The tropospheric processing of acidic gases



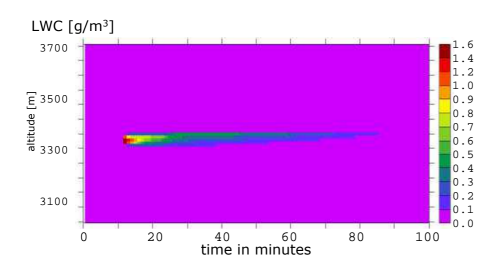
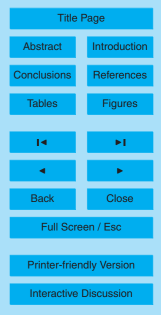


Fig. 6. Contour plot of the liquid water content of the cloud developing at the crater in model run "cloudy". Time is in minutes since model start (plume release is 11 min after model start) and altitude in metres.

6, 11653–11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



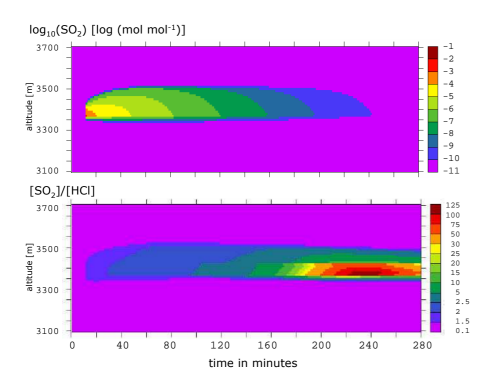
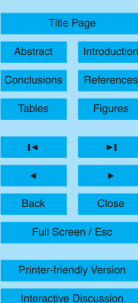


Fig. 7. Contour plot of the evolution of the SO_2 and the ratio SO_2 to HCl for the "cloud-free" model run. Time is in minutes since model start (plume release is 11 min after model start) and altitude in metres. Note the strongly non-linear contours for $[SO_2]$:[HCl].

6, 11653–11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



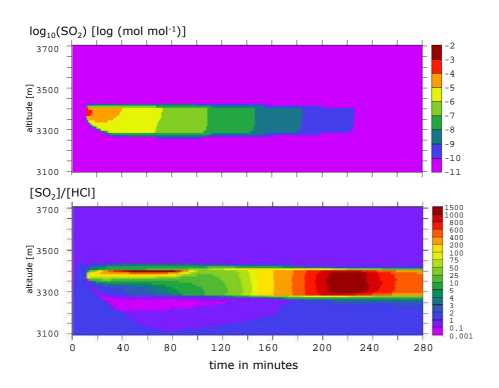
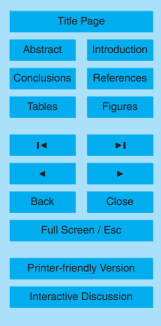


Fig. 8. Same as Fig. 8 but for the "cloudy" model run.

6, 11653-11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



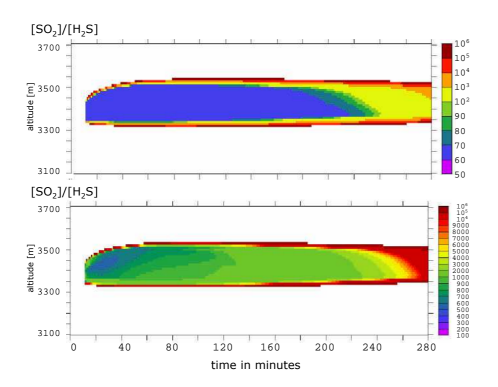


Fig. 9. Contour plot of the SO_2 to H_2S ratio for a cloud-free model run without (top) and with halogen $-H_2S$ interactions (bottom), respectively. Time is in minutes since model start (plume release is 11 min after model start) and altitude in metres.

6, 11653–11680, 2006

The tropospheric processing of acidic gases

A. Aiuppa et al.



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Interactive Discussion