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https://doi.org/10.1098/rsta.2005.1668

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Volcano remote sensing with ground-based spectroscopy

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The chemical compositions and emission rates of volcanic gases carry important information about underground magmatic and hydrothermal conditions, with application in eruption forecasting. Volcanic plumes are also studied because of their impacts upon the atmosphere, climate and human health. Remote sensing techniques are being increasingly used in this field because they provide real-time data and can be applied at safe distances from the target, even throughout violent eruptive episodes. However, notwithstanding the many scientific insights into volcanic behaviour already achieved with these approaches, technological limitations have placed firm restrictions upon the utility of the acquired data. For instance, volcanic SO\textsubscript{2} emission rate measurements are typically inaccurate (errors can be greater than 100\%) and have poor time resolution (\textit{ca} once per week). Volcanic gas geochemistry is currently being revolutionized by the recent implementation of a new generation of remote sensing tools, which are overcoming the above limitations and are providing degassing data of unprecedented quality. In this article, I review this field at this exciting point of transition, covering the techniques used and the insights thereby obtained, and I speculate upon the breakthroughs that are now tantalizingly close.

Keywords: volcanology; remote sensing; spectroscopy

1. Introduction

In June 1991, Mt Pinatubo in the Philippines exploded violently, propelling material some 35 km high into the stratosphere (e.g. Newhall \textit{et al.} 2002). This was the world’s largest eruption in 80 years and ten times greater than Mt St Helens. Yet, in spite of the enormous loss of life that could have followed, it is estimated that only a few hundred people died as a result. In what is regarded to be the most successful volcanic-hazard mitigation in history, some 80 000 people were safely evacuated shortly before the beginning of this eruption, saving many thousands of lives. Scientists had been carefully monitoring the volcano’s behaviour, since activity commenced some months previously, and correctly forecasted this climax to the eruption. One of their key lines of evidence was a rapid tenfold increase in the volcano’s SO\textsubscript{2} emission rate (flux), observed with ground-based spectroscopy (Daag \textit{et al.} 1996).

One contribution of 17 to a Triennial Issue ‘Astronomy and earth science’.
Volcanoes emit gases to the atmosphere both during and between eruptions. These can be exsolved from magma and released through major craters and vents: the so-called primary emissions (see figure 1), which are the focus of most remotely sensed volcanic gas observations. Additionally, volcanoes may release fluids from, or interact with, groundwaters (hydrothermal systems; Symonds et al. 1994), and gases can be diffusely emitted through volcanic flanks. The most abundant chemical species in volcanic plumes are $\text{H}_2\text{O}$ and $\text{CO}_2$, and other significant components include $\text{SO}_2$, $\text{HCl}$, $\text{HF}$ and $\text{H}_2\text{S}$. These gases carry important information about underground conditions such as magmatic composition, volume, transport and degassing dynamics. As such, gas chemical composition and flux are routinely measured at a number of volcanoes worldwide to provide fundamental scientific insights. As the concentration of gas in magma and the dynamics of its escape to the atmosphere are arguably the primary drivers of volcanic activity, strongly influencing whether an eruption is explosive or effusive and triggering rapid switches between these regimes (Sparks 2003), these data are also valuable for forecasting eruptions.

Volcanic gases have important impacts upon the atmosphere and climate (Robock 2000), which must be correctly accounted for in climate change models. In particular, volcanic $\text{SO}_2$ is converted to radiatively active sulphate aerosol, which reflects sunlight back to space (Graf et al. 1997). Some scientists think this effect may have been responsible for prehistoric mass extinction events. More recently, aerosol from the Mt Pinatubo eruption cooled the troposphere, entirely offsetting the anthropogenic greenhouse effect for a limited duration. This eruption also caused stratospheric heating, an alteration in global stratospheric circulation patterns and an approximately 20% reduction in Antarctic ozone levels (McCormick et al. 1995).

A third motivation for interest in these emissions is the serious environmental and medical impacts they can have (Delmelle et al. 2002). For instance, at Masaya volcano, Nicaragua, atmospheric $\text{SO}_2$ concentrations are elevated over a downwind area greater than 1000 km$^2$, resulting in loss of revenues from coffee plantations and exposing approximately 50 000 people to levels exceeding World Health Organization air quality standards. One of the world’s worst environmental disasters was precipitated by the 1783–1784 eruption of Laki in Iceland, which released over a hundred million tonnes of $\text{SO}_2$ (Thordarson et al. 1996).

Figure 1. Volcanic gas emissions from Mt Etna. (a) Crater degassing. (b) Individual fumaroles on a crater rim.
This gas formed a blanket of haze that covered much of Europe, causing destruction of agricultural crops, contamination of pasture and human morbidity and mortality. Benjamin Franklin, then acting as ambassador to France, was one of the first to note the correlation between this volcanic pollution and the anomalously cold weather experienced in Europe and North America at that time.

The conventional method of measuring volcanic gases is to collect a gas sample directly, then transport it to the laboratory for subsequent analysis. Collection methodologies include bubbling gases from vents through alkaline solutions or pumping the gases through base-treated filters, which act as acid traps (Symonds et al. 1994). While direct sampling is the most detailed and accurate source of degassing data, this approach suffers a number of limitations: first, routine surveillance may be difficult or impossible owing to the risks involved, which can restrict observations to low-temperature vents that are not representative of the volcano as a whole; second, the samples’ chemical compositions may be altered in storage; and third, analysis may involve delays (during volcanic crises data are often required immediately). In comparison, the geophysical techniques used to monitor volcanoes: seismology and geodesy (ground-deformation measurements), enable data to be acquired each second or so from autonomous installed units.

Over the last 30 years, ground-based remote sensing techniques have been increasingly used for volcanic plume measurements, significantly broadening the surveillance capability of volcanic gas geochemistry. The primary advantage of these approaches is safety, as monitoring can take place multiple kilometres from the targets, allowing, in principle, the acquired data to resemble more closely seismic and geodetic data, in respect of continuous monitoring (particularly throughout violent eruptive phases), high time resolution (seconds to minutes) and near-real-time data evaluation. Remote sensing is also non-invasive, thereby avoiding sample-contamination issues; it involves integrated measurements across the whole plume, thus capturing the volcano’s bulk degassing behaviour, and it can provide fluxes, which are difficult to extrapolate from directly sampled data.

Spectroscopic plume observations can also be made from space. This began rather serendipitously, when an anomalously high ‘ozone’ signal was observed over Mexico by the Total Ozone Mapping Spectrometer, following the 1982 eruption of El Chichón. It was soon recognized that this corresponded to absorption of ultraviolet (UV) radiation by volcanic SO$_2$ (Krueger 1983), and satellite sensors have been used for volcanic plume monitoring ever since (e.g. Bluth et al. 1993). However, in view of the inferior spatial and temporal resolution of these instruments, and their higher detection limits relative to ground-based remote sensing devices, they are of limited value in eruption monitoring at present. Rather, their primary utility is in measuring SO$_2$ burdens of clouds from large explosive eruptions, which cannot be constrained from the ground; a number of significant advances in our understanding of volcanic impacts upon the atmosphere have thus been obtained. As the satellite sensors are being improved constantly (e.g. Carn & Bluth 2003; Carn et al. 2005; the newest generation can image far lower altitude plumes than possible hitherto), an
exciting future prospect will be increased synchronous applications of space-
borne and ground-based volcano remote sensing.

In this article, I outline the two most applied ground-based volcanic plume 
remote sensing techniques: UV and Fourier transform infrared (FTIR) 
spectroscopy. A description of some of the observations made, and volcanological 
insights gleaned, will then be presented, culminating in a discussion of future 
research prospects.

2. Methodologies

(a) Generic background

The techniques described here are based upon measurements of the spectra of 
electromagnetic radiation that has been attenuated by passing through a volcanic 
plume (figure 2). Depending upon the methodology used and the volcano studied, 
the radiation source may be natural, e.g. direct sunlight, scattered skylight, hot 
rocks or active lava bodies, or artificial, e.g. a lamp. As each of the volcanic gas 
species has a unique molecular structure, they have characteristic absorption 
features, e.g. a particular wavelength will be attenuated to a different extent by 
each gas present. Therefore, by examining an appropriate spectral window, the 
presence of these species can be inferred from the distinctive ‘bar-code’ spectral 
imprints they impress upon the incident radiation. This principle is used to 
determine gas concentrations quantitatively, using the Beer–Lambert Law:

\[
I(\lambda) = I_0(\lambda) \exp[-\sigma(\lambda)NL].
\]

\(I(\lambda)\) and \(I_0(\lambda)\) are the intensities of the radiation measured by the instrument at 
wavelength \(\lambda\), when the plume is present and absent from the optical path, 
respectively; \(\sigma(\lambda)\) is the absorption cross-section of the absorbing molecule, a 
quantitative measure of how strongly it attenuates light. The product of the 
mean concentration of the attenuating species \((N)\) and the width \((L)\) of the 
plume viewed (the column amount) can thus be found. In practice, these are

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Figure 2. Transmission of radiation (in this case from an artificial lamp) through a volcanic plume, 
before detection by a spectrometer. The emitted source spectrum \(I_0(\lambda)\) and the measured spectrum 
\(I(\lambda)\) differ depending upon the composition of absorbing species in the plume.
simultaneously obtained for all volcanic species of interest, using computational fitting routines, which scale $\sigma(\lambda)$ for each gas, across a spectral window.

(b) Ultraviolet spectroscopy

The most commonly used ground-based volcanic plume remote sensing technique is UV spectroscopic measurement of volcanic SO$_2$ fluxes. Attention has been focused on SO$_2$ because of its strong UV absorption features, low background atmospheric concentrations and relatively high plume abundances: typically it is the third most ubiquitous species, behind H$_2$O and CO$_2$. In contrast, it is extremely difficult to sense volcanic H$_2$O and CO$_2$ remotely because of their high ambient concentrations. SO$_2$ flux measurements have been made for 30 years with correlation spectrometers (e.g. Stoiber et al. 1983), which were recently upgraded with laptop PC controlled USB-powered spectrometers (in particular Ocean Optics Inc. USB2000s; McGonigle et al. 2002; Galle et al. 2003), which are considerably smaller, lighter, cheaper and less power consuming. During flux measurements, the USB2000 is optical fibre coupled to a vertically pointing telescope, then the plume is traversed from below, by foot, boat, aircraft or road vehicle, and overhead scattered skylight spectra are continuously acquired and saved, as are positional data from a handheld GPS receiver (figure 3a).

The spectra are then processed to determine overhead SO$_2$ column amounts. First, all spectra collected underneath the plume are divided by a spectrum acquired outside it, according to the Beer–Lambert law, thereby isolating the attenuation of the skylight UV radiation by the plume. As a slight departure from the generic methodology outlined above, the resulting spectra are then high-pass filtered, before being fitted with the SO$_2$ absorption cross-section, itself identically filtered. This methodology, termed differential optical absorption spectroscopy, removes the possible interference of spectrally broadband attenuation from the plume aerosols. The GPS data are then used to resolve the component of motion perpendicular to the plume transport direction, corresponding to each spectrum acquired underneath the plume; a cross-sectional profile of the plume SO$_2$ column amounts is thus generated. By integrating across this, then multiplying by the plume transport speed, the SO$_2$ flux value is obtained.

While most measurements are made by manually traversing the plume (data typically have a time resolution of a few days and are often sustained only for the
duration of a field campaign, e.g. one to two months), a new monitoring configuration, based on scanning the plume from below with a USB2000, has recently been developed (Edmonds et al. 2003a; McGonigle et al. 2003; figures 3b and 4). With knowledge of the plume’s height, which can be determined, for instance by tomographic analysis of the plume profiles observed with two or more scanners, fluxes can be computed in a similar manner to the traverse case. Such a system was automated at the Soufrière Hills volcano, Montserrat, and has been collecting data autonomously since 2002. This is an outstanding example of the value added to volcanological monitoring by rapid capitalization upon new technological opportunities (Edmonds et al. 2003a).

While SO$_2$ fluxes are the most ubiquitous of all ground-based remotely sensed volcanic data, and have provided many valuable scientific insights, they are limited in their utility by large errors (potentially greater than 100%). These primarily originate in the standard practice of using readings from ground-based anemometers as proxies for wind speeds at the plume altitude, typically hundreds of metres above the surface. In addition, the SO$_2$ flux measurements use scattered skylight, and, while this makes alignment trivial, the flux calculation relies upon the somewhat naïve assumption that the collected light has been scattered into the spectrometer’s field of view from above the plume and has thence travelled in a straight vertical line downwards. In reality, the sampled radiation can be scattered above, within (perhaps multiple times from aerosols) and/or below the plume; there has yet to be a rigorous characterization of errors arising from this. (c) Fourier transform infrared spectroscopy

FTIR spectrometers, operating in the near- to mid-infrared wavelength range, are capable of measuring column amounts of gases such as HCl, H$_2$O, SO$_2$, HF, CO$_2$, SiF$_4$, OCS and CO in volcanic plumes (e.g. Notsu et al. 1993; Francis et al.

Figure 4. Device for automated scanning SO$_2$ flux measurements. Skylight is coupled into the USB2000 (a) using a telescope (b) and optical fibre (c). A rotating mirror, located in the protective housing (d), pans the instrument’s field of view across the scan plane (see figure 3b) — the stepper control board is contained inside (e). The measurement position, required for the flux calculation, is found using the handheld GPS (f). Spectral acquisition and mirror rotation are controlled using a laptop PC.
With the miniaturization of FTIR spectrometers for military applications, commercial field-portable devices became available in the 1990s. In many cases, SO$_2$ flux data are open to highly divergent explanations; interpretation can be clarified with FTIR-derived multiple gas column amount data, because of the species’ different physico-chemical properties. Each acquired FTIR spectrum is analysed by using a radiative transfer model to generate a synthetic spectrum, which simulates the absorption along the optical path length by those atmospheric and volcanic species present. By iteratively varying the latter column amounts until the best fit to the observed spectrum is achieved, the plume composition is determined.

A variety of radiation sources can be used with FTIR spectroscopy: direct sunlight, artificial lamps, hot rocks or active lava bodies. While this provides design flexibility and avoids the scattering problems of the UV skylight measurements, alignment in the latter case is, generally, more straightforward. Likewise, FTIR spectrometers are considerably bulkier and more expensive than the USB2000s, and the retrievals are more complex. For these reasons, this approach has not yet become as widespread as UV spectroscopy among volcanologists. Correlation spectroscopy has also been used, albeit to a far lesser extent than FTIR spectroscopy, for remotely sensing multiple volcanic gas species concentrations.

3. Observations

UV spectroscopy was initially applied in a volcanological context to determine whether observed SO$_2$ fluxes correlate with changes in activity. The simplest mechanism proposed to affect these rates in advance of an eruption is magma rising to shallow depths, where pressures are insufficient to confine the dissolved gases, such that they exsolve and escape to the atmosphere. This interpretation has been ascribed to the behaviour of a number of volcanic events in the past, such as the increase in flux observed before the 1991 Mt Pinatubo eruption (Daag et al. 1996). Likewise, increasing SO$_2$ emissions from Mt Etna have been noted prior to a number of eruptions (Caltabiano et al. 1994).

While predicting the onset of eruptions is a vital task for volcanologists, identifying the end or decline of activity can be equally important, informing resettlement strategies for surrounding areas. Remotely sensed data can also provide useful insights here. For instance, spectroscopic measurements following the cataclysmic 1980 eruption of Mt St Helens revealed a general exponential decline in SO$_2$ flux over the following 8 years (McGee 1992), suggesting a gradual cessation of activity, as the intruded magma, responsible for the main explosion, was steadily losing its dissolved gases and not being recharged. Smaller eruptions during this period were preceded by elevated SO$_2$ fluxes, indicating intermittent phases of rising magma (McGee & Sutton 1994).

During eruptions volcanic SO$_2$ fluxes are elevated as gas is erupted with lava at the surface; however, emission rates can also be modulated by non-magmatic trends, such as solid Earth tides. One of the most surprising insights arising from the UV spectroscopic measurements is that volcanoes can also sustain very large SO$_2$ emissions with little or no corresponding lava eruption (e.g. Kazahaya et al. 2004). An extreme example of this was the passive emission of up to
60 000 tonnes day$^{-1}$ from Popocatépetl volcano in Mexico during 1997, roughly double the time-averaged SO$_2$ release from all the volcanoes in the world. A model, whereby gas is released from a bubbly layer at the top of magma chambers, has been proposed to account for such behaviour (Wallace 2001).

SO$_2$ flux measurements are relatively straightforward to interpret where the gases are free to escape to the atmosphere. However, in many cases the situation is more complicated, e.g. the magmatic gases might bubble through a hydrothermal system, which dissolves each species to a different extent, or fracture networks above the magma chamber can seal, inhibiting gas escape. For these reasons, SO$_2$ flux data alone have been less successful when applied to predicting the vent-clearing cataclysmic explosions that often begin eruptions. For instance, prior to the eruption of Mt St Helens, and in spite of the presence of magma at shallow depth, measured SO$_2$ fluxes were very low, owing to a rock carapace, which severely restricted gas release.

The ongoing activity of the Soufrière Hills volcano has been monitored using both UV and FTIR spectroscopy. A degassing model has thus been developed, whereby the emissions are derived from two distinct sources: SO$_2$ is released from a ‘mafic’ magma, intruded from depth into the chamber of an ‘andesite’ magma, itself rich in HCl (Edmonds et al. 2001, 2003b). The latter magma is relatively viscous; therefore, when extruded through the volcano’s conduit, it forms a lava dome at the volcano summit, rather than the lava flows typical of other eruptive styles. These episodes are characterized by elevated HCl fluxes, with exsolution of this gas upon ascent; conversely, HCl fluxes are low when dome growth does not take place. In contrast, SO$_2$ is emitted discontinuously, depending on the permeability of the system. One proposed sealing mechanism is closure of interconnecting bubble networks in the andesite.

Activity at the Soufrière Hills volcano had apparently stopped during 1998, with cessation of dome growth and seismic signals; yet SO$_2$ emissions did not decline, but rather continued with higher rates than observed during previous phases of lava extrusion. This provided key evidence that the eruption was not over, and indeed activity was later resumed, highlighting the importance of remote sensing in this context. Gas-measurement-based criteria for identifying the end of this eruption have also been highlighted: persistently low HCl and SO$_2$ fluxes would indicate that mafic charging of the magma chamber has ended, as has the resulting dome growth.

While FTIR spectroscopy is normally based upon monitoring volcanic gases’ absorption of radiation from a hotter background source, Love et al. (1998) used FTIR spectroscopy to observe the plume’s emitted spectra, in contrast to that of the cold sky background. At Popocatépetl volcano, Mexico, they observed a steady increase in SiF$_4$/SO$_2$ ratio over a few days prior to an explosive eruption during February 1997, followed by a rapid tenfold decrease to the pre-eruptive value. Owing to temperature-dependent equilibration between the volcanic species present, this ratio is a proxy for gas temperature and indicated cooling prior to the eruption. The interpretation given was that the gases adiabatically expanded as the conduit plug began to give way in advance of the eruption.

Different gases have different pressure-dependent solubilities in magma, so the ratios of volcanic gas concentrations are also important indicators of the depth of exsolution from the melt. For instance, SO$_2$ is removed at greater depths than HCl; therefore, a high SO$_2$/HCl ratio is indicative of deep exsolution, and a low...
ratio implies shallower degassing. On occasions, volcanoes can manifest ‘lava
fountaining’ behaviour, where gas jets propel lava fragments to heights of several
hundred metres. However, whether this gas is exsolved during these episodes, or
they are driven by a gas layer previously accumulated at depth, has been a
matter of debate. FTIR spectroscopic observations of relatively high \( \text{SO}_2/\text{HCl} \)
gas ratios for a lava fountain on Mt Etna during 2000 have clarified this matter
by indicating the latter origin of gas in this case (Allard et al. 2005). Constraints
upon exsolution depths have also been used to map the underground branched
magmatic system, which feeds the active vents on Mt Etna (Burton et al. 2003).

Even greater scope for interpretation is provided by combining parallel
geophysical and remotely sensed volcanic gas observations. Galeras volcano in
Colombia has been the focus of considerable scientific attention, following a
resurgence of activity in 1988. During this period, a number of explosive
eruptions occurred, one of which resulted in the tragic deaths of six scientists and
three tourists. Prior to such events reduced \( \text{SO}_2 \) fluxes were observed, in parallel
with increasing ‘long-period’ seismic signals, analogous to the resonances formed
in organ pipes. In retrospect, this behaviour was interpreted to indicate sealing
and pressurization of gas within the system, followed by explosive release (Stix
et al. 1993; Fischer et al. 1994). One proposed mechanism for this was
precipitation of minerals, arising from a hydrothermal system, in gas pathways
to the surface. In cases where hydrothermal interactions are suspected, FTIR
spectroscopic data act as valuable adjuncts to UV spectroscopic \( \text{SO}_2 \) fluxes in
clarifying interpretation, given the different water solubilities of the gases (e.g.
\( \text{HCl}, \text{SO}_2 \) and \( \text{HF} \)) accessible to the former approach (e.g. Symonds et al. 2001).

The UV flux measurements, made at many targets worldwide, have been used
to derive a total global volcanic \( \text{SO}_2 \) emission of approximately 20 Mton yr\(^{-1} \)
(e.g. Stoiber et al. 1987); from this the total fluxes of other volcanic species have
been calculated. While this volcanic flux is only around one-fifth of the total
anthropogenic release, sulphate loading of the atmosphere may be comparable
from the two sources, as the former emitters are, generally, at higher altitudes,
where aerosol lifetimes are longer. This is highly significant, as anthropogenic
sulphate is presently considered to be the most important factor offsetting the
anthropogenic greenhouse effect. Recently, USB2000s have been adapted for
remotely sensed measurements of two other volcanogenic gases: \( \text{NO}_2 \) and \( \text{BrO} \)
(Bobrowski et al. 2003). The former observation supports a recent report that
volcanoes may have played a significant role in generating biologically available
forms of nitrogen, via thermal fixation, during the evolution of the early Earth
(Mather et al. 2004). The \( \text{BrO} \) data indicate that volcanoes may constitute a
significant (yet hitherto largely overlooked) supply of bromine to the lower
atmosphere, with potentially important impacts upon tropospheric ozone
chemistry.

4. The future

Ground-based remote sensing has provided invaluable insights into volcanic
behaviour, which in many cases could not have been obtained in any other way,
owing to the hazards of working closer to the source. These data can be used as
proxies for underground magma mass and migration, changes in temperature,
sealing or opening of systems and depth of degassing, and they have assisted eruption forecasting measures. However, until recently, they have suffered from a number of significant limitations, such that their full potential has yet to be realized: first, flux measurements have large errors (potentially greater than 100%), primarily owing to the anemometric proxies used for plume speeds; second, they normally have low time resolution (ca one week) and are typically sustained for a few months at most; and third, measurements of multiple species (often vital for accurate geochemical interpretation) are not widespread, owing to the complexity and expense of FTIR spectroscopy. For these reasons, while degassing is arguably the primary driver of volcanic activity, determining much observed seismicity and ground deformation, measurements of the latter phenomena are presently considered to be of greater utility in volcano monitoring.

The volcanological application of the USB spectrometers heralds a new era for volcanic gas geochemistry. Automated scanning versions of these devices have been developed over the last few years and enable acquisition of high time resolution (a few minutes) sustained SO$_2$ fluxes. In addition, it has been demonstrated very recently that accurate plume transport speeds can be measured using two vertically pointing USB2000s, located underneath the plume at different distances from the source (McGonigle et al. 2005; Williams-Jones et al. in press). By cross-correlating the overhead SO$_2$ column amount time-series from these devices, the time of transit of individual plume ‘parcels’ can be obtained, allowing accurate flux measurements (errors less than 10%). By combining these two approaches, truly unprecedented sustained flux data will become available, in respect of time resolution and accuracy. In addition to the gases already measured, the USB2000 shows potential for remote measurements of other volcanogenic species, such as ClO. While these devices are unlikely to replace FTIR spectroscopy as the standard remote sensing technique for multiple volcanic gas column amount measurements, at least not in the near future, they could prove to be a useful and relatively inexpensive adjunct in this regard.

As a proxy for eruptive activity, SO$_2$ suffers the disadvantage of being readily removed by hydrothermal systems. CO$_2$, which exsolves from magma deeper, and is comparatively water insoluble, is a far more useful magmatic tracer. Unfortunately, it is notoriously difficult to measure volcanic CO$_2$ using remote sensing, owing to its high ambient concentrations. While FTIR spectroscopic observation of this species has been achieved in volcanic plumes (Burton et al. 2000), this has been performed only with an artificial lamp source, a configuration that is not widely applicable and requires relatively close access to the source. A number of research groups have tried, without success to date, to develop instruments for spectroscopic CO$_2$ measurements, using scattered skylight as the radiation source. One possible solution is pressure modulation radiometry, which measures vertically resolved gas profiles and may be able to spatially filter out much of the background CO$_2$ signal. While LIDAR are capable of making volcanic CO$_2$ flux measurements, these devices are power consuming and not field portable at present. A new radiometric technique for monitoring H$_2$O fluxes has recently been developed and applied at a number of Japanese volcanoes.

O’Dwyer et al. (2003) have measured SO$_2$/H$_2$S ratios with a USB spectrometer and deuterium bulb source over open optical paths (figure 5) and by pumping
fumarolic gas into an optical cell. With ruggedization, the latter configuration shows potential for sustained autonomous operation; and by adding further spectrometer channels it could potentially monitor HCl, CO$_2$ and H$_2$O, in addition. More complex variations of this, based on laser spectroscopy, have been developed, which measure isotope ratios and are thus capable of providing additional geochemical information.

With the development and implementation of the above technologies, spectacular breakthroughs in volcanology are foreseen during the coming decade, in two particular ways. First, the great improvements in accuracy and time resolution of SO$_2$ flux time-series will enable rigorous corroboration of degassing data with geophysical datasets (seismic, geodetic, thermal and acoustic). Such inter-parametric analyses were hindered by the poor degassing data previously available, yet are a key to increasing our volcanological understanding, given growing evidence for the interconnectedness of these phenomena (e.g. Watson et al. 2000). Second, these technological improvements will allow the first detailed flux-data-based testing and parameterization of numerical volcanic conduit models (e.g. Melnik & Sparks 1999). The gas data will thus progress from being primarily a useful means of cataloguing volcanic behaviour in retrospect to playing a more vital role in forecasting eruptions, thereby attaining a status more commensurate with the great importance of degassing in driving volcanic activity.

While our understanding of the effects of plumes from explosive eruptions upon the stratosphere is relatively advanced, less is known about the impacts of smaller plumes upon the troposphere. With better and more widely distributed ground-based remote sensing technologies, in parallel with novel satellite sensors, this situation will improve as global volcanic fluxes become better constrained, both spatially and temporally. Further investigations into the conversion of SO$_2$ to sulphate aerosol will provide additional insights here; however, it remains to be seen whether USB2000 observations alone can provide sufficient accuracy to

Figure 5. Spectroscopic measurements of H$_2$S/SO$_2$ ratios, using a USB spectrometer and deuterium bulb, at Solfatara, Campi Flegrei, Italy. Dashed line denotes optical path. Photograph courtesy of Miles Padgett.
improve our present understanding of this phenomenon. Better characterization of the important radiative forcing role played by tropospheric volcanic plumes would help improve climate change models, at a time when accurate climate forecasting has never been so important.

The author is supported by a NERC post-doctoral fellowship. Pierre Delmelle and David Pyle are thanked for their conscientious reviews, which have greatly improved the quality of this paper.

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Andrew J. S. McGonigle was born in 1973. He grew up in Edinburgh, graduating from St Andrews University with an M. Sci. degree in theoretical physics in 1995. After spending a year working for a charity in Zambia, he returned to university to study for a D.Phil. in laser physics at the University of Oxford. He then worked for a start-up company, before joining the Department of Geography at the University of Cambridge in 2001 as a research associate. From 2002 to 2005 he was a NERC Post-doctoral Research Fellow at Cambridge, developing and applying spectroscopic techniques for monitoring emissions from power stations, volcanoes and biomass burning. He is presently a DTI Academic Research Fellow at the University of Sheffield. Photograph: helicopter-borne UV spectroscopic measurements of SO$_2$ flux at Stromboli volcano, courtesy of Saibbo Inguaggiato.