Development of a portable active long-path differential optical absorption spectroscopy system for volcanic gas measurements

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Received: 16 April 2014 – Revised: 5 November 2014 – Accepted: 8 November 2014 – Published: 19 December 2014

Abstract. Active long-path differential optical absorption spectroscopy (LP-DOAS) has been an effective tool for measuring atmospheric trace gases for several decades. However, instruments were large, heavy and power-inefficient, making their application to remote environments extremely challenging. Recent developments in fibre-coupling telescope technology and the availability of ultraviolet light emitting diodes (UV-LEDS) have now allowed us to design and construct a lightweight, portable, low-power LP-DOAS instrument for use at remote locations and specifically for measuring degassing from active volcanic systems. The LP-DOAS was used to measure sulfur dioxide (SO₂) emissions from La Fossa crater, Vulcano, Italy, where column densities of up to 1.2 × 10¹⁷ molec cm⁻² (~500 ppmm) were detected along open paths of up to 400 m in total length. The instrument’s SO₂ detection limit was determined to be 2 × 10¹⁶ molec cm⁻² (~8 ppmm), thereby making quantitative detection of even trace amounts of SO₂ possible. The instrument is capable of measuring other volcanic volatile species as well. Though the spectral evaluation of the recorded data showed that chlorine monoxide (ClO) and carbon disulfide (CS₂) were both below the instrument’s detection limits during the experiment, the upper limits for the X/SO₂ ratio (X = ClO, CS₂) could be derived, and yielded 2 × 10⁻³ and 0.1, respectively.

The robust design and versatility of the instrument make it a promising tool for monitoring of volcanic degassing and understanding processes in a range of volcanic systems.

1 Introduction

The emission of volatile species from volcanic magmas is linked to geochemical processes occurring at depth. Volatiles are exsolved as a result of recharge, decompression, crystallisation or mixing of magmas. The composition and flux of gases being emitted from a volcanic vent can provide insights into geophysical and geochemical parameters such as magma supply, pressure, temperature and degassing depth — parameters that can used by volcanologists to forecast eruptions.

The major volatile species emitted from volcanic vents are water vapour (H₂O), carbon dioxide (CO₂), sulfur dioxide (SO₂), hydrogen sulfide (H₂S) and hydrogen halides (HX with X = Fl, Cl, Br, I). Of these, CO₂ has the lowest solubility in most magma types, and is typically emitted first as a magmatic system is recharged (Werner et al., 2013; Burton et al., 2013). Therefore, developing methods for accurately monitoring CO₂ emissions from volcanoes has been a major focus of the volcanology community in recent years.

Unfortunately, volcanic CO₂ emission rates are considerably more difficult to measure than those of some of the other species, the main reason being the significant background concentration of CO₂ in the Earth’s atmosphere. Unless measurements are made very close to the vent where high magmatic/atmospheric gas ratios can be sampled, the enhancement of CO₂ in a volcanic plume is oftentimes only a few parts per million (ppm) over the background of
approximately 400 ppm. Passive remote sensing instruments that use scattered solar radiation as a light source and therefore make an integrated measurement over the entire atmospheric column to detect an even smaller enhancement. The relative increase in the total CO$_2$ column is of the order of only $10^{-3}$ for large emitters and can be significantly less at volcanoes with moderate to small emission rates. For this reason, CO$_2$ emissions from volcanoes have not yet been detected from satellite platforms, and direct measurements of CO$_2$ emission rates from volcanic vents can only be made by laborious and costly aircraft traverse techniques (Gerlach et al., 1997).

On the other hand, SO$_2$ has a negligible atmospheric background concentration and is much more accessible to remote sensing observations, with routine measurements made both from ground-based and satellite-based instrumentation (Galle et al., 2010; Oppenheimer et al., 2011; Carn et al., 2013). In fact, arguably the most robust method currently available for measuring volcanic CO$_2$ emission rates is by determining the molecular ratio of CO$_2$ to SO$_2$ in a volcanic plume and multiplying this by the SO$_2$ molecular emission rate obtained from remote sensing instrumentation (Gerlach et al., 1998; Burton et al., 2013; Pering et al., 2014). CO$_2$/SO$_2$ molecular ratios can either be determined by collecting gas in Giggenbach bottles (Giggenbach and Goguel, 1989) for analysis in the laboratory or using in situ chemical and optical sensors in a multicomponent gas analyzer system (Multi-GAS, Aiuppa et al., 2005; Shinohara, 2005). While the sampling of fumaroles with Giggenbach bottles is limited in its applicability for continuous monitoring, these relatively novel Multi-GAS instruments now allow real-time measurements of volcanic gas compositions suitable for eruption forecasting (Aiuppa et al., 2007).

Despite their relative versatility, the Multi-GAS instruments also have some drawbacks, and determining the gas speciation representative of the bulk plume at a volcanic vent remains challenging. For one, the instruments typically rely on an optical CO$_2$ sensor and multiple electrochemical sensors for other gas species (Aiuppa et al., 2005). The different sensors tend to have significantly different response times, with the optical sensor being quicker to respond to a change in composition than the electrochemical sensors. This can lead to errors and uncertainties in the derived gas ratios if not properly taken into account (Roberts et al., 2014). Also, the electrochemical sensors have cross-sensitivities to other gases – oftentimes gases found in great abundance in volcanic plumes (Roberts et al., 2012). A typical electrochemical H$_2$S sensor, for example, can have a cross-sensitivity to SO$_2$ of up to 20% (Roberts et al., 2014). Finally, while in situ sampling lends itself to deriving the gas composition at a specific location, problems arise if the volatile speciation is heterogeneous in space in the area around the volcanic vent. Such heterogeneities are quite common, and can be caused by a complex plumbing system and interaction of volatiles with a hydrothermal system, or result from gas and fluid temperature variations in the proximity of the vent. In such cases, the gas composition measured by an in situ measurement may not be representative of the bulk plume.

In light of these challenges, improved methods for measuring plume compositions at volcanic vents are still being sought. One very sophisticated method is to use Fourier transform infrared spectroscopy (FTIR) to measure the absorption or emission of infrared (IR) radiation by volcanic gases (Oppenheimer et al., 1998; Stremme et al., 2012). FTIR is sensitive to a large number of species, including both CO$_2$ and SO$_2$, and can therefore be used in combination with SO$_2$ emission rate measurements to derive CO$_2$ emission rates (Burton et al., 2000). When used in an open-path configuration with the light path penetrating the bulk plume, representative gas compositions can be obtained. However, FTIR instruments are very costly and limited in their portability, as they rely on a moving interferometer to record spectral information. Also, deployment can be challenging in the absence of accessible hot lava, as they then require either a bulky thermal IR emitter requiring tens of watts of power to be deployed on the opposite side of the vent, or must be located such that the plume is between the instrument and the Sun, which is then used as an IR source.

Near-infrared (NIR, 800 nm $< \lambda < 2500$ nm) tuneable diode laser absorption spectroscopy (TDLAS) instruments represent a somewhat simpler alternative (Schiff et al., 1994; Carapezza et al., 2011; Pedone et al., 2014). Although typically only sensitive to a single gas species according to which the laser wavelength is chosen, today’s NIR-TDLAS units are small and portable, typically weighing $\sim 2.5$ kg, and can run for extended periods of time on battery power. When set up in an open-path configuration, these instruments only require that a retroreflector be deployed on the opposite side of the desired optical path, as both the light source and the detector are usually located in the same unit. Reflectors are considerably simpler to deploy than active light sources, as they do not require power, do not need to be carefully adjusted and can withstand harsh environmental conditions more easily. NIR-TDLAS instruments measuring CO$_2$, H$_2$S, HF or CH$_4$ alone open paths are commercially available for purchase.

Measuring SO$_2$ in the infrared absorption is also possible, but due to the lack of near-infrared (NIR) absorption bands, measurements are made in the thermal infrared regions ($3.5$ µm $< \lambda < 20$ µm). Though the technology of thermal infrared lasers is constantly evolving (Tittel et al., 2012), they remain considerably more complex than NIR diode lasers, and generally require significantly more power. Depending on which thermal IR band is chosen, additional complications can arise from interference by pressure-broadened absorption lines of other atmospheric compounds (mainly CO$_2$ and H$_2$O). Therefore, IR laser absorption measurements of SO$_2$ are usually performed in cells at reduced barometric pressures (Richter et al., 2002; Rawlins et al., 2005), and no IR laser absorption measurements of SO$_2$ along open
atmospheric paths are known to date. In the absence of an SO$_2$ measurement, emission rates of gases detected using NIR-TDLAS cannot easily be determined using the ratio method described earlier.

For this reason, implementation of a portable, active open-path measurement of SO$_2$ is highly desirable. In this study, we describe how a miniature active long-path differential optical absorption spectroscopy (LP-DOAS) instrument was designed to measure SO$_2$ and potentially also other trace gases with a high degree of accuracy and precision along well-defined open paths around volcanic vents. Emphasis was placed on ensuring portability of the instrument on foot to remote locations, a sufficient degree of environmental hardening, low power consumption that can easily be supplied by battery, and simple set-up and removal at the target site. After the instrument was constructed, tests were performed at the island of Vulcano (Aeolian arc, Italy) in February 2010, the results of which are described below.

2 Design of a portable LP-DOAS instrument

The first measurements of atmospheric trace gases by LP-DOAS were performed in the late 1970s (Platt et al., 1979). The technique has since become widely used in the atmospheric sciences community, particularly because the contact-free nature of the measurement allows the detection of highly reactive radical species (Platt and Stutz, 2008). More recently, active differential absorption measurements have also been applied to volcanic gases (O’Dwyer, 2003; Kern et al., 2008), but deployment of active LP-DOAS systems has been severely limited by the size (a typically longer than 1 m focal length telescope) and weight (typically tens to hundreds of kg) of such systems and by the power consumption of the ultra-violet light sources required (typically tens to hundreds of watts).

Two recent technological advances have now made it possible to design a lightweight portable LP-DOAS instrument. For one, the size and weight of the telescope unit can be dramatically reduced by replacing the conventional Newtonian set-up with a fibre-coupling coaxial design. In this set-up, first introduced by Merten et al. (2011), the light source is coupled into the telescope using a fibre bundle, one end of which is placed in the focal plane of the spherical primary mirror. Light from the desired light source is coupled into several of the fibres in the bundle, reflected on the primary mirror, and transmitted in a nearly parallel beam into the atmosphere to an array of retroreflectors located tens or hundreds of metres away. These reflect the light back to the primary mirror of the telescope, where, partly due to spherical aberration of the mirror, some of it enters the receiving fibres that couple it into a moderate resolution ultraviolet (UV) spectrometer for analysis. The use of this fibre-coupling design eliminates the need for any secondary mirrors and allows separation of the light source from the telescope, thus making a lighter, portable telescope design possible.

In our design, the fibre bundle consists of seven 100 µm diameter fused silica fibres, six of which are arranged in a circular pattern around a central seventh fibre. As suggested by Merten et al. (2011), we use the six outer fibres of the bundle for transmission and the central fibre to receive radiation that has passed through the optical system. However, we have modified the original design by additionally splitting the transmission fibres into three groups of two (Fig. 1). This allows the coupling of three different light sources into the optical path, a feature that is unique to our system and allows a high degree of flexibility with regard to measurement wavelength (see below).

In this coaxial set-up, the telescope primary mirror acts both as a sending and receiving unit. A three-dimensional fibre positioner is used to mount the end of the fibre bundle in the focal plane of the mirror (Fig. 1), which is protected from environmental conditions by an MgF$_2$ coating. Compared to previously described LP-DOAS systems, we use a significantly smaller primary mirror diameter $d$ of only 150 mm with a focal length $f$ of 400 mm to reduce the overall instrument size and weight. In the paraxial approximation, and considering the fibre bundle diameter of about 300 µm, this set-up leads to a beam spread angle $\alpha$ of approximately 0.04 degrees. The diameter $D$ of the retroreflector array required to reflect the entire emitted radiation back to the telescope is then given by

$$D = d + 2L \tan \left( \frac{\alpha}{2} \right) \approx d + L \tan \alpha.$$  

Here, $L$ is the distance between the telescope and reflector array, which is also exactly half of the optical path length. In our set-up, an array diameter of 19 cm is needed at a distance of 50 m. For a path length of 200 m, an array diameter of 29 cm is required to reflect all transmitted radiation. A larger telescope would reduce beam spread, but the compact design chosen here makes the instrument portable and allows measurements to be made in otherwise inaccessible remote locations. Path lengths of up to a few hundred metres are sufficient for making measurements at most volcanic vents. For the test measurements, we used an array of ten 52 mm diameter fused silica corner cubes densely packed in a triangular frame mounted on a tripod. This array had a total diameter of approximately 40 cm, thus allowing one-way paths of up to about 350 m without significant loss of radiative energy. We successfully made measurements along paths of up to 200 m with the set-up (see Sect. 3).

For one-way paths longer than 350 m, one might expect a significant drop in the measurement signal-to-noise ratio (SNR). Because the fraction of reflected light is proportional to $D^{-2}$ and therefore proportional to $L^{-2}$ for long light paths (Eq. 1), photon statistics yield an SNR of the measured spectrum that is proportional to the square root of the intensity, or $L^{-1}$. However, since the column density is proportional to
Figure 1. Photograph (a) and schematic (b) of the portable active long-path DOAS instrument. Light from three UV LEDs is coupled to a fibre bundle, collimated to a parallel beam and sent to an array of reflectors. Upon returning, it is coupled to a moderate-resolution UV spectrometer for analysis of absorption features.

the light path length (if the plume fills the optical path), the measured optical depth will be proportional to $L$, thus yielding a measurement SNR that is, in first order, independent of the light path length. However, if opaque volcanic plumes are measured, scattering of light out of the optical path by aerosols and water droplets will further decrease the SNR, particularly for longer light paths (see Sect. 4.2).

In addition to allowing for a decoupling of the light source from the telescope, the fibre design allows for fine adjustments of the beam direction without the need to adjust the pointing direction of the telescope (Merten et al., 2011). After aiming the telescope in the vicinity of the reflectors, slight adjustments to the fibre position can be used to fine-tune the pointing direction of the instrument. This technique, however, requires that the $f$/#$ of the telescope be smaller than that of the spectrometer. In other words, the area from which light is collected by the spectrometer must be smaller than the primary mirror. If this is not the case, light that has not passed along the optical path can enter the system from adjacent to the telescope primary mirror when the fibre bundle is moved out of the focal point. We avoid this problem by using an $f$/2.7 telescope and an $f$/4 spectrometer. The $f$/4 spectrometer collects light from a circular 10 cm diameter subsection of the primary mirror, so the fibre bundle can be moved up to about 2.5 cm to each side of the focal point without collecting light from beside the mirror. This allows adjustment of the telescope pointing direction by $\pm 3.6$ degrees in the left/right and up/down directions without moving the telescope itself.

The second technological advance which allowed us to design a portable LP-DOAS instrument is the availability of an efficient UV light source. Besides the weight of the instrument itself, the application of LP-DOAS systems in remote locations has also been severely limited by the high power consumption of conventional UV light sources. Typical light sources such as xenon arc and deuterium lamps only emit a fraction of their consumed energy as optical output – by far the majority of the energy is lost to wavelengths that are not relevant for the measurement, especially in the thermal IR. The advent of UV-LEDs and their applicability to LP-DOAS instruments has now revolutionised the achievable power efficiency of such systems (Kern et al., 2006; Sihler et
By utilising three UV LEDs with maximum emission wavelengths at 285, 310 and 315 nm instead of a gas discharge lamp, the power consumption of the light source was reduced from what was typically between 50 and 500 W to less than 1 W without significantly affecting the optical output power at the required wavelengths. We used Sensor Electronic Technology UVTOP285, UVTOP310 and UVTOP315 flat window LEDs with maximum emission wavelengths of 285, 310 and 315 nm, respectively. Each consuming approximately 180 mW of electrical power, these LEDs are rated as emitting between 600 and 800 µW of optical power at wavelengths within approximately ±15 nm (12 nm FWHM) around their peak emission (Sensor Electronic Technology, 2014).

As shown in Fig. 2, the UV LED wavelengths were chosen to overlap with the differential absorption features of SO₂, chlorine monoxide (ClO) and carbon disulfide (CS₂) (Bogumil et al., 2003; Vandaele et al., 2009). However, other LEDs can be chosen to target other species (e.g. bromine monoxide (BrO) at ∼340 nm, chlorine dioxide (OClO) at ∼350 nm (Kern et al., 2008) or nitrogen dioxide (NO₂) at ∼350–450 nm, Kern et al., 2006; Chan et al., 2012), or three LEDs of the same wavelength can be coupled to the system to enhance the SNR of the measurement in one specific wavelength region. The UV LEDs are each mounted on a 2-D positioner and coupled to the fibre bundle using two fused-silica plano-convex lenses (Fig. 1). The lens positioning along the optical axis is individually adjusted so as to provide maximum coupling efficiency for a given LED wavelength, thus avoiding chromatic aberration.

After the radiation has passed along the open optical path, the spectrum is analysed using an Ocean Optics QE65000 spectrometer with a cooled, back-illuminated CCD detector. The installed grating provided a wavelength range of 260 to 340 nm at a resolution of 0.5 nm. The spectrometer was placed in an insulated enclosure and the temperature of the optical bench was stabilised using a pulse-width-modulated thermoelectric cooling/heating unit to avoid fluctuations in calibration or other optical properties of the system. Excluding the notebook computer that was used for data collection in these test measurements, the power consumption of the entire system was below 25 W during continuous operation. This represents a considerable reduction in power use when compared to any previously reported LP-DOAS set-up, and allowed us to make battery-operated measurements.

The majority of the consumed power is actually used for temperature stabilisation of the spectrometer optical bench. The achieved temperature stability of better than ±0.1 °C is required when measuring weak absorption features of trace gases in the optical path of smaller than ∼0.01 optical depth (see next section). If only gas species such as SO₂ with relatively strong absorption bands are targeted, this stabilisation is not required, and the power consumption of the system can be reduced to as low as 5 W (depending on the CCD chip temperature). During our test measurements described in the next section, we used a laptop computer for data acquisition. This added an additional ∼15 W to the total power consumption, and we chose to use a 45 Ah, 12 V lead acid battery to run the measurements for the entire day. However, a set-up for making continuous measurements could either use a low-power integrated PC or make use of a USB device server (∼2 W) instead of the laptop, and include a radio telemetry link (∼5 W) to the observatory where the spectral acquisition and processing would occur. Without temperature stabilisation (e.g. if only measuring SO₂), the system could be run...
continuously in this set-up with a total power consumption of approximately 12 W. Use of a solar power system for supplying the instrument power is therefore possible.

3 Configuration of measurements at Vulcano

On 25 February 2010, the portable LP-DOAS was tested at the island of Vulcano (Aeolian arc, Italy). Measurements were performed using two different light paths. During the first measurement period from approximately 12:00 to 12:45 local time, the array of retroreflectors was positioned behind a low-temperature fumarole about 100 m from the telescope (see Fig. 3a). During this time, the gases being emitted by this fumarole were measured. Little or no disturbance from other fumaroles is expected in this data set, as favourable winds were blowing the gases from other fumaroles in the area away from the light path. Depending on the wind, the light path length through the fumarolic gases was between 0 and 15 m one way, so a total light path of between 0 and 30 m resulted.

During the second measurement period, the reflectors were moved to a position farther down in the crater (approximately 200 m from the telescope) to enable the measurement of gases being emitted from the higher-temperature fumaroles located on the edge of the crater floor (Fig. 3b). The light path length inside the fumarolic gases varied considerably due to variations in wind direction as well as variations in the degassing strength of the fumaroles themselves.

4 Evaluation of spectral data and measurement results

4.1 Spectral retrieval of SO₂

Due to its importance as a tracer for other volcanic gases (see Sect. 1), SO₂ is the primary target species for the LP-DOAS instrument. SO₂ has several distinct absorption features in the spectral region between 260 and 320 nm, the strongest of which lie around 300 nm (Vandaele et al., 2009, Fig. 2). However, we chose to measure the differential absorption of SO₂ between 280 and 290 nm, because the relatively high column densities generally encountered in the plumes of active volcanoes make absolute sensitivity a secondary concern. More importantly, the ability to measure at shorter wavelengths eliminates any interference from scattered solar radiation entering the spectrometer, as the ozone layer effectively absorbs all incident solar radiation at wavelengths shorter than 300 nm (Hartley absorption bands). A UVTOP LED with a 285 nm peak transmittance wavelength was selected for this purpose.

The SO₂ column density was derived from each measured spectrum using a DOAS retrieval. First, each spectrum was corrected for the electronic offset of the spectrometer. This offset is added to each measured spectrum by the spectrometer electronics prior to digitisation in order to avoid negative intensity values. It is best measured by co-adding a large number of exposures at very short exposure time and with no incident light. Since the electronic offset is added to each exposure, the co-added spectrum is then divided by the number of exposures, multiplied by the number of exposures in the measurement spectrum, and then subtracted from it.

In the next step, each measurement spectrum is corrected for the detector dark current. This contribution stems from thermal excitation of electrons in the CCD detector, and is proportional to the exposure time. It is therefore best
characterised by taking a long exposure of the CCD with no incident radiation. After subtracting the offset, the dark current spectrum is divided by the exposure time, multiplied by the exposure time of the measurement spectrum, and then subtracted from it. Note that both the offset and dark current are typically dependent on the temperature of the spectrometer. However, since the spectrometer and the CCD detector were both kept at a stable temperature during our measurements, characterisation of the offset and dark current could be performed before and after the measurements were made.

In order to evaluate only the differential absorption features in a measured spectrum, the next step is to derive the differential optical density \( \tau' \). \( \tau' \) is defined as the negative logarithm of the ratio of measured intensity \( I \) to transmitted intensity \( I_0 \) which, according to the Beer–Lambert–Bouguer law of absorption, is proportional to the column density \( S \) of the absorber in the light path.

\[
\tau' = -\ln \left( \frac{I}{I_0} \right) = \sigma \cdot S \quad (2)
\]

Here, \( \sigma \) is the absorption cross section of the absorbing gas. The initial intensity \( I_0 \) is the spectrum of the light sources before the radiation has passed through the atmosphere. Since \( I_0 \) can depend on the temperature of the LEDs (Kern et al., 2006; Sihler et al., 2009), it is best measured in close temporal proximity to the measurement spectra. In LP-DOAS instruments, characterisation of the light sources can be performed using a “shortcut” system which redirects light emitted from the light sources directly into the spectrometer without passing along the optical path. Since we use a fibre assembly to couple radiation into and out of the telescope, a shortcut system can be implemented by simply placing a reflective diffuser plate in front of the fibre bundle in the focal point of the mirror (Merten et al., 2011). Radiation emitted from the fibre ring is scattered on this plate and, in part, enters the central collection fibre for analysis. In between measurements, we manually inserted a sand-blasted aluminium plate a few cm from the fibre bundle and measured \( I_0 \). However, owing to the variable wind directions encountered within the La Fossa crater, we also intermittently obtained measurement spectra with no volcanic gas in the light path of the instrument. These were later found to be the best measure of \( I_0 \), because exactly the same optical path through the instrumentation is used as during the measurements, only in the absence of volcanic gas. In our evaluations, we co-added 60 such spectra and used these as an LED reference (shown in Fig. 4).

In a differential optical absorption spectroscopy retrieval (Platt and Stutz, 2008), the column density \( S \) of one or multiple absorbers \( i \) (in this initial case, just SO\(_2\)) in the instrument’s light path is derived by fitting the laboratory absorption cross sections \( \sigma_i \) to the measured optical density, along with a polynomial \( P_\lambda \) (we use a fifth order) to account for broadband spectral features. We varied this approach slightly, and instead fit the SO\(_2\) absorption cross section \( \sigma_{SO_2} \) and the logarithm of the shortcut spectrum \( I_0 \) to the logarithm of the measurement spectrum \( I \), as shown in Eq. (3).

\[
\ln I = \ln I_0 - \sum_i \sigma_i S_i + P_\lambda \quad (3)
\]

In this manner, a slight shift in wavelength calibration of the spectrometer between shortcut and measurement, should this occur, can be accounted for in the fit. Since the spectral resolution of the instrument does not match that of the laboratory reference (we use the SO\(_2\) cross section measured by Vandaele et al., 2009), the reference cross section is convolved with the instrument line shape before the fit is performed. The line shape or so-called “slit function” was characterised by recording the spectrum of a mercury vapour lamp and measuring the shape of a single atomic emission line at the spectrometer’s resolution. The convolved cross section was then used in the fit.

An example of the SO\(_2\) fit is shown in Fig. 4. In this example, an SO\(_2\) column density of 1.5 \( \times 10^{17} \) molec cm\(^{-2}\) of SO\(_2\) was retrieved. The residual, shown at the top right of the figure and representing the difference between the measured spectrum and the fit result, is about 1 % peak to peak. When compared to the SO\(_2\) absorption cross section, this noise level corresponds to an SO\(_2\) detection limit of approximately 2 \( \times 10^{16} \) molec cm\(^{-2}\) or 8 parts per million \( \times \) metres (ppmm). In other words, concentrations below 1 ppm can be detected along a 10 m total path (5 m one way).
The results of the SO$_2$ evaluation for the spectra recorded during the first measurement period at the low-temperature fumarole are shown in Fig. 5a. The time series is characterised by a strong fluctuation of SO$_2$ column densities in time which is mainly caused by wind blowing the fumarolic gas into or out of the light path of the instrument. At times, the retrieved SO$_2$ column density reached up to $2.8 \times 10^{17}$ molec cm$^{-2}$ ($\sim 110$ ppmm). Assuming a total optical path length of up to 30 m in the fumarolic gas (see Fig. 3a), this would correspond to an average SO$_2$ mixing ratio of 3.7 ppm.

The second optical path that was chosen was more favourable in that it did not pass through as dense a gas plume as in the first measurement period (Fig. 3b). This led to a lower opacity and slower fluctuations in SO$_2$ column density (Fig. 5b). Up to $1.2 \times 10^{18}$ molec cm$^{-2}$ ($\sim 480$ ppmm) of SO$_2$ were measured along this 200 m (one-way) path. We estimate that the volcanic gas filled about 1/4 of the optical path for this measurement. Under this assumption, the measured column density corresponds to an average mixing ratio of about 4.8 ppm. When further taking into account that this plume was considerably more diluted than the plume measured at the first position, this second plume appears to have a significantly higher SO$_2$ fraction. This finding is in agreement with the results of Aiuppa et al. (2005), who also found a significantly higher SO$_2$ fraction in the plume originating from the inner-crater fumaroles. The combination of slower column density fluctuations and improved integration time (typically shorter than 300 ms) also led to smaller errors in the retrieval for this time period (see next section).

4.2 Rapid variation of gas and aerosol concentrations in the optical path

The errors in the results obtained for the first measurement site (depicted in Fig. 5a) are significant, particularly for individual measurements of high column densities preceded and followed by lower values. The relatively large error in these measurements therefore appears to be associated with a change in gas and/or aerosol concentration in the instrument’s light path during a measurement. Here we discuss the instrument’s sensitivity to such fluctuations.

Two effects can be separated. First, the concentration of aerosols and water droplets in the light path can change over time. A portion of the light emitted by the LP-DOAS instrument is scattered onto particles and droplets while passing along the open optical path. In the narrow beam approximation (Platt and Stutz, 2008), light that has been scattered once will not re-enter the light path, and is therefore removed from the system. This leads to a reduction in the light intensity received by the telescope and inherently decreases the SNR of the measurement according to photon statistics.

However, the DOAS approach is not susceptible to a systematic error when the intensity changes during an exposure. Since a polynomial $P_\lambda$ is included in the fit to account for broadband spectral features such as those associated with particle scattering (see Eq. 3), the column density $S$ is retrieved solely from the differential depth of the narrowband absorption lines of the respective trace gas. If the broadband intensity of radiation returning from the atmospheric path changes without a change in the relative depth of the absorption lines, an exposure integrated over this change will still exhibit the differential optical depth associated with the encountered gas column density. This can easily be shown for two spectra with different on-band intensities $I_1$ and $I_2$, and off-band intensities $I_{0,1}$ and $I_{0,2}$. If we require that

$$\frac{I_1}{I_{0,1}} = \frac{I_2}{I_{0,2}},$$

(4)
retrieval, as is discussed in the next section. Correction is the measurement needs to be corrected during the spectral trum that appears superimposed on the spectrum of the ac-
this contribution to the instrument. This leads to a "background" solar spec-
can also scatter solar radiation into the receiving optics of the instrument. This will directly influence the atmospheric light path which can also potentially change the measured optical depth.

In other words, that the amount of gas in the light path be constant (see Eq. 2), then

$$\frac{I}{I_0} = \frac{I_1 + I_2}{I_0} = \frac{(I_1 + I_2)}{(I_0 + I_1)} \frac{(I_0 + I_2)}{(I_0 + I_1)} = \frac{I_1}{I_0} \cdot \frac{I_0 + I_2}{I_0 + I_1} \cdot \frac{I_2}{I_0} \cdot \frac{I_0}{I_0}$$

Therefore, the optical depth $\tau$ will remain an accurate measure of the column density $S$ of gas in the instrument's well-defined light path (Eq. 2) if only the aerosol concentration changes. This stands in contrast to the effect that aerosols have on passive DOAS measurements which use scattered solar radiation to measure volcanic gases. For these, any change in the aerosol concentration along the viewing direction will directly influence the atmospheric light path along which the measurement is being conducted, thus also potentially changing the measured optical depth $\tau$ and greatly impacting the accuracy of the measurement (Kern et al., 2009).

Particles and water droplets in the LP-DOAS optical path can also scatter solar radiation into the receiving optics of the instrument. This leads to a "background" solar spectrum that appears superimposed on the spectrum of the active light sources (shown in Fig. 2a). This contribution to the measurement needs to be corrected during the spectral retrieval, as is discussed in the next section. Correction is not necessary when operating at wavelengths shorter than 300 nm (as we do here), because solar radiation is negligible in this region.

The second effect that can occur during a measurement is a rapid change in the column density $S$ of the measured trace gas. This effect is significantly more problematic than a change in the aerosol concentration. When measuring volcanic plumes, the two effects will often occur simultaneously as $SO_2$ and aerosol-rich volcanic gas is blown into and out of the instrument light path. If we assume a constant off-band intensity $I_0$, then a change in trace gas column density will lead to a change in the on-band intensity of a measurement from $I_1$ to $I_2$. If these two successive measurements are co-added (or integrated during a single exposure of the CCD), the optical depth $\tau$ can be calculated from the sum of the individual intensities:

$$\tau = \ln \left( \frac{I_1 + I_2}{I_0} \right) = \ln \left( \frac{I_1}{I_0} + \frac{1}{2} \frac{I_2}{I_0} \right)$$

Equation 6 shows that, owing to the nonlinearity of the Beer–Lambert–Bouguer law (Eq. 2), $\tau$ is not equal to the average of the two optical depths $\tau_1$ and $\tau_2$ associated with the gas column densities at time 1 and time 2. Instead, there is a systematic shift towards lower optical depths. This effect will also influence the shape of the absorption lines in the measured spectra, and the fit of the absorption cross sections according to Eq. (3) will result in a systematic residual structure. Accurate measurements therefore require that the exposure time be short compared to the timescales over which the column density of the absorbers changes in the light path.

During our test measurements at the first site, individual spectra were recorded at exposure times varying between approximately 150 and 600 ms, depending on the opacity of the plume in the light path at any given time. Because the retroreflector was positioned in direct proximity to the fumaroles, high opacities were encountered fairly often, and exposure times of ~600 ms were not uncommon. As shown above, changes in plume opacity alone would not lead to systematic errors, but in this case, such changes were likely accompanied by changes in the gas column in the instrument’s light path, as the fumarole’s plume drifted in and out. This often led to a misfit between measured optical depth and the absorption cross section and, thus, resulted in the relatively large errors (Fig. 5a). During the measurements conducted at the second site, generally lower plume opacities led to shorter exposure times. Thus, the assumption of a constant $SO_2$ column density in the light path is valid, and the errors are significantly reduced (Fig. 5b). To avoid systematic errors in the future, care must be taken to ensure that the exposure time is always shorter than the timescale of variations in the trace gas column density, even if this leads to undersaturated exposures and, associated with this, slightly decreased SNRs for the measurement.
4.3 ClO and CS₂ detection limits

Besides SO₂, CS₂ and ClO both have characteristic absorption features in the 260–360 nm wavelength region (Fig. 2). The absorption cross section of ClO was originally included in the SO₂ fit described above, but no differential features associated with ClO absorption were detected during the evaluation, so it was later omitted from the SO₂ retrieval. In an attempt to improve the SNR of the measurement and thereby improve the ClO detection limit, the evaluated spectra were sorted by their SO₂ column density, binned in groups of 10 to 20 such that all spectra in a single group had comparable SO₂ columns, and co-added to improve the statistics of the measurement.

These spectra were again evaluated for ClO absorption, but none was found. Due to their high solubility in water, halogen gases are very efficiently scrubbed when passing through large-scale hydrothermal systems (Symonds et al., 2001). Considering the significant hydrothermal activity at the island of Vulcano, it is therefore not surprising that halogen emissions appear negligible. However, an upper limit could be derived from the spectral retrieval. For this, the magnitude of the fit residual was analysed.

The fit residual represents the difference between the measured optical depth and the optical depth derived from the DOAS fit. Absorption structures of gases not accounted for by the fit would appear in the residual. For a given measurement spectrum, the detection limit of a gas species not included in the DOAS model can therefore be derived by comparing the fit residual with the absorption cross section of that species multiplied by a hypothetical column density. This is demonstrated in Fig. 4. Here, the magnitude of the fit residual is compared to the absorption features associated with a hypothetical column density of $1.5 \times 10^{15}$ ClO molec cm$^{-2}$. By making the conservative assumption that any ClO absorption features larger than these could not be masked by the noise in the fit residual, a column of $1.5 \times 10^{15}$ ClO molec cm$^{-2}$ represents the upper limit for this particular spectrum. This corresponds to a path-averaged mixing ratio of about 1.5 ppb (note that the volcanic gas only fills part of the 400 m total light path) and an upper limit for the ClO/SO₂ ratio of about $5 \times 10^{-3}$ for this example. When analysing the entire collected data set, and particularly the co-added spectra containing larger amounts of SO₂ than in the given example, an upper limit for the ClO/SO₂ molar ratio of about $2 \times 10^{-3}$ was obtained for the emissions from the fumaroles on the edge of the crater floor. Considering an average SO₂ emission rate from the fumarolic area of about 12 t d$^{-1}$ (as reported by Vita et al. (2012) for the 2008–2010 period), this corresponds to an approximate upper limit for the ClO flux of 20 kg d$^{-1}$.

Finally, the recorded spectra were also evaluated with regards to the potential absorption of CS₂. CS₂ was first measured by DOAS in the urban atmosphere above Shanghai by Yu et al. (2004). Though it has never been detected by DOAS in a volcanic environment, an attempt was made here, because reduced sulfur species (particularly H₂S, Aiuppa et al., 2006) are known to be present in abundance in the fumaroles at La Fossa crater on the island of Vulcano, where these measurements were made. The differential absorption features of CS₂ in the 310–330 nm wavelength range (Fig. 2) do make it potentially accessible to our LP-DOAS instrument. A separate CS₂ evaluation was performed in which the wavelength region between 312 and 328 nm was analysed. At shorter wavelengths, strong SO₂ absorption interfered with the measurement. At longer wavelengths, the intensity of the available LED was insufficient. In addition to the cross section of CS₂, the cross section of SO₂ was included in the fit, as well as a reference spectrum compiled from 60 spectra without SO₂ absorption. A fifth-order polynomial was again used to account for any broadband structures in the retrieval. Because this retrieval was performed at wavelengths longer than 300 nm, the spectrum of solar radiation scattered into the instrument’s light path by condensed water droplets and volcanic gases was found superimposed onto the LED spectra. This is clearly demonstrated by the non-zero baseline at 350 nm in the example spectrum shown in Fig. 2b. To correct for this effect, a Fraunhofer solar reference spectrum (measured with the same geometry but with the LED switched off) and a Ring spectrum (used to correct for inelastic scattering in the atmosphere, Grainger and Ring, 1962) were also included in the fit.

As was the case for ClO, CS₂ concentrations were below the instrument’s detection limit throughout the measurement period. However, the same approach was used to derive an upper limit for the CS₂/SO₂ molecular ratio in the fumarolic emissions. CS₂ was omitted from the DOAS model and, after the fit accounted for SO₂ absorption and interference from solar radiation scattered into the optical path, the fit residual was compared to the absorption features of hypothetical columns of CS₂. Figure 6 shows an example in which an upper limit of $6 \times 10^{16}$ CS₂ molec cm$^{-2}$ (~24 ppmm) was obtained. This corresponds to an upper limit for the path-integrated average mixing ratio of about 60 ppb, a CS₂/SO₂ molecular ratio of less than 0.1 and an average CS₂ flux of less than about 1.4 t d$^{-1}$ (again assuming an average SO₂ emission rate of 12 t d$^{-1}$ as reported by Vita et al., 2012).

5 Conclusions and outlook

Recent technological advances in fibre-coupled telescopes and UV-LED technology have made design and construction of a mobile active LP-DOAS for use in remote environments possible. Here, we built an instrument specifically for measuring volcanic gas emissions. The tests conducted at La Fossa crater, Vulcano, showed that in the current configuration, the instrument is highly sensitive to SO₂, with open-path measurements of SO₂ column densities as low as about...
whether the achieved detection limit will make the LP-DOAS is only a very minor component (Halmer et al., 2002) when the altitude of CS

On the other hand, little is known about the actual magnitude of CS

emissions from volcanic systems, so it is unclear whether the achieved detection limit will make the LP-DOAS a valuable tool for assessing this trace species. Globally, CS₂ is only a very minor component (Halmer et al., 2002) when compared to other sulfur-bearing species (mainly SO₂ and H₂S), but trace amounts have been reported both in explosive eruption plumes (Rasmussen et al., 1982) and in fumarolic gases (Stoiber et al., 1971), so its exact role remains uncertain, and abundances may vary widely from system to system.

The described LP-DOAS system can also be used to measure other volcanic volatiles (Fig. 2), most notably perhaps the halogen oxide compounds OClO and BrO. The design of the fibre-coupling system allows the addition of LEDs at other wavelengths if targeting species with absorption cross sections outside the current range. Alternatively, three LEDs emitting at the same wavelength can be used to improve the light throughput and SNR of the system at that wavelength. Another envisioned modification that will likely further improve the sensitivity of the instrument is the implementation of a pulsed LED power supply. The optical output power of LEDs is limited by the heat generated by the dissipated electrical input power. Therefore, LEDs can be pulsed at significantly higher input currents than they can sustain in continuous operation. Such a pulsed operation does not significantly improve the time-integrated optical output power of the LEDs but, if synchronised with the spectrometer read-out electronics, the system could measure the background spectrum during periods when the LEDs are off, thus providing a more or less contemporaneous measurement of the background solar radiation scattered into the light path. Currently, accurate correction of the scattered solar contribution appears to be one of the main limitations of the system, particularly when applying it to opaque, optically thick plumes, and a pulsed acquisition stands to improve this correction significantly.

The flexibility of the system allows application to a wide range of volcanic system types, from active high-temperature vents to hydrothermal fumarole fields. The active light sources allow measurements to be made both during the day and at night, which can provide insights into the photochemistry taking place in volcanic plumes, as for example shown by Kern et al. (2008). The instrument’s light weight, rugged design and low power consumption allow active DOAS measurements to be made at remote locations that were previously inaccessible. It therefore has the potential to make significant improvements to our ability to monitor and understand geochemical processes in volcanic systems around the world.

Acknowledgements. The authors would like to acknowledge Tom Pering, Tamar Elias and two anonymous reviewers for their thoughtful reviews of the manuscript. Any use of trade, firm, or product names is for descriptive purposes only and does not imply endorsement by the US Government.

Edited by: A. Schütze
Reviewed by: three anonymous referees
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