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A new permanent, low-cost, low-power SO₂ camera for continuous measurement of volcanic emissions

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Since its introduction to volcanology in the mid-2000 s, the SO₂ camera has become an important instrument for the acquisition of accurate and high timeresolution SO₂ emission rates, aiding in hazard assessment and volcanological research. However, with the exception of a few locations (Stromboli, Etna, Kīlauea), hitherto the majority of measurements have been made on discrete field campaigns, which provide only brief snapshots into a volcano's activity. Here, we present the development of a new, low-cost, low-power SO₂ camera for permanent deployment on volcanoes, facilitating long-term, quasi-continuous (daylight hours only) measurements. We then discuss preliminary datasets from Lascar and Kīlauea volcanoes, where instruments are now in continuous operation. Further proliferation of such instrumentation has the potential to greatly improve our understanding of the transient nature of volcanic activity, as well as aiding volcano monitoring/eruption forecasting.

KEYWORDS

volcanology, sulphur dioxide, ultraviolet camera, remote sensing, raspberry pi

1 Introduction

Volcanic emissions are the manifestation of subsurface magmatic activity, therefore providing an insight into the state of a volcanic system and potentially providing a means of hazard forecasting through their measurement (e.g., Aiuppa et al., 2009; De Moor et al., 2016; Kunrat et al., 2022). Sulphur Dioxide (SO₂) is the most common gas species detected at openvent volcanoes with remote sensing instrumentation, owing to its relative ease of detection that stems from two principal factors: Its relatively low background atmospheric concentration and its strong distinctive absorption bands at ultraviolet (UV) and infrared (IR) wavelengths (Platt and Stutz, 2008).

Since their development in the mid-2000s, ultraviolet (UV) SO_2 cameras (Mori and Burton, 2006; Bluth et al., 2007) have become extremely valuable tools for measuring volcanic emissions, due to their provision of high temporal (at times >1 Hz) and high spatial

resolution (providing two-dimensional, 2D, plume images) datasets. However, unlike the similarly popular differential optical absorption spectroscopy (DOAS) instruments, which have been installed as permanent/continuously operating instruments on a number of open-vent volcanoes worldwide (Burton et al., 2009; Galle et al., 2010; Arellano et al., 2021), SO₂ camera technology is yet to proliferate in this regard; to the best of our knowledge, previous permanent systems have been installed on only Stromboli (Burton et al., 2015), Etna (D'Aleo et al., 2016; Delle Donne et al., 2017), and Kīlauea (Kern et al., 2015), with the latter having to be removed from the field during the 2018 caldera collapse. Further installations may have been prevented, in part, by the cost of such equipment, which typically utilizes scientific grade UV cameras ($\approx1000s-10000$ s USD per camera; 2 cameras per SO₂ camera system).

Discrete field campaigns with SO₂ cameras have regularly been conducted, as they are often easier and cheaper to undertake; however, recent research has emphasized the importance of gathering long-term time series, since a volcano can exhibit significant changes in activity over a wide range of time scales (e.g., Pering et al., 2019). Such datasets form the foundation of volcano monitoring, which often relies on identifying the departure of a volcano's activity away from some baseline (Phillipson et al., 2013), and also allow more detailed/reliable volcanological research. Indeed, the permanent SO₂ cameras currently in operation have already provided important new insights into volcanic activity, as well as instrument performance. On Etna, D'Aleo et al. (2016) captured shifts in activity from one vent to another over the course of an eruptive period, inferring shallow interconnections between the vents of the volcano. Delle Donne et al. (2019) observed a mild but detectable increase in SO₂ emission rate prior to the onset of paroxysmal activity on Etna in May 2016. At Stromboli, Burton et al. (2015) found reasonable agreement of camera data with a scanning DOAS network, whilst also highlighting that the high temporal resolution allows quantification of explosive events which are too brief to be captured by the scanning DOAS network. Delle Donne et al. (2017) then found that the frequency of explosion/ puffing activity at Stromboli increased significantly prior to the onset of Stromboli's August-November 2014 effusive activity. More recently on Stromboli, Delle Donne et al. (2022) found that explosive degassing accounts for $\approx 10\%$ of the total SO₂ emission budget, over a year-long observation period. They further showed that explosive SO₂ flux correlates well with both very-long-period (VLP) seismicity and passive SO2 flux, inferring some commonality in the source processes involved in passive and explosive activity. In a somewhat different application, Elias et al. (2018) acquired highlyaccurate SO₂ emission rates by combining UV camera-derived plume velocities with SO₂ measurements made by an array of upward-looking UV spectrometers. Their approach integrates the benefit of sophisticated DOAS retrievals of SO2 column densities with the benefit of image-based plume speed estimation.

The work herein aims to contribute to the aforementioned growing need for long-term, quasi-continuous SO₂ emission rate data from active volcanoes. We present a new SO₂ camera which has significant potential to broaden the use of permanent SO₂ cameras in volcanology, owing largely to its relatively low-cost and lowpower design, as well as its associated open-source code for data processing. The instrument is based on previously developed Raspberry Pi-based UV camera technology that exploits low-cost visible imaging cameras, modifying them to improve their UV sensitivity (Wilkes et al., 2016; Wilkes et al., 2017). We discuss installations of the system on 2 volcanoes, Lascar (Chile) and Kīlauea (Hawaii, United States), presenting preliminary datasets from both. We note that the instrument has also been installed on two other volcanoes (El Reventador, Ecuador; Lastarria, Chile); however, we do not yet have data from these instruments so they are not discussed further herein.

2 Materials and methods

2.1 SO₂ camera principles

The SO₂ camera works by exploiting the significant absorption of ultraviolet (UV) light by SO₂. Images are acquired at two distinct wavelengths (bands), using bandpass filters (\approx 10 nm full width at half maximum; FWHM) inserted into the camera optics. The onband is located in a spectral region where SO₂ absorption is significant (\approx 310 nm), whilst the off-band is located in a nearby region where SO₂ absorption has diminished to be negligible in comparison (\approx 330 nm). Using the Beer-Lambert law of absorption, plume pixels in an image can be compared to the background sky radiance to calculate pixel optical depths in each band:

$$AA = \tau_{SO2} = \tau_{on} - \tau_{off} = \ln \frac{I_{0on}}{I_{on}} - \ln \frac{I_{0off}}{I_{off}}$$

where τ is the optical depth, I_0 is the background sky intensity, I is the in-plume intensity, and subscripts on and off refer to the onband (310 nm) and off-band (330 nm) images, respectively. The offband image is used to correct the on-band optical depths for radiative transfer effects that are relatively broadband in nature (thus change minimally between the on- and off-band images); this is principally scattering/absorption from aerosols in the plume.

 SO_2 optical depth can be calibrated to SO_2 column densities using either gas cells of known SO_2 amounts or a co-aligned spectrometer retrieving contemporaneous SO_2 column densities from a small region within the field of view of the camera (Lübcke et al., 2013). For permanent SO_2 camera installations, the latter method is universally employed, since gas cell calibration would require the incorporation of moving parts into the system, such as a filter wheel, to cycle through cell calibrations a number of times each day. Moving parts are likely to result in failing components over the lifetime of a camera system, as well as adding more complexity to the initial design and construction. Furthermore, contemporaneous spectroscopic measurements can allow for more sophisticated retrievals, including correction for light dilution and in-plume scattering (Kern et al., 2013; Varnam et al., 2021).

An emission rate is retrieved by multiplying a plume crosssectional mass (often referred to as integrated column amount; ICA) by the plume speed. Plume speed is primarily calculated by one of two methods: 1) cross-correlating time series from two parallel ICAs, one set downwind of the other, and therefore calculating the lag in plume motion over the series; 2) computer vision algorithms, principally optical flow, which estimate the motion of all features in an image from one frame to the next (e.g., Peters et al., 2015). The latter provides both a higher spatial (theoretically down

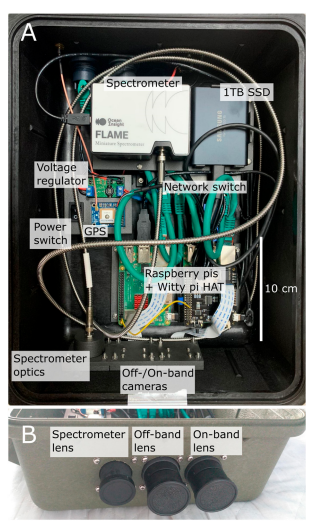


FIGURE 1

(A) Layout of permanent SO₂ camera. Most parts are mounted to the case using custom-designed three-dimensional prints. Whilst this system is housed inside a Peli Case 1,400 ($306 \times 234 \times 130$ mm), other systems have used the Peli Case 1,450 ($380 \times 266 \times 153$ mm) to provide extra space for mounting the system. (B) Front view of instrument optical systems.

to pixel-level) and temporal resolution (frame-by-frame) to plume speed estimation, relative to cross-correlation. An alternative crosscorrelation technique, which can also provide frame-by-frame temporal resolution, uses a single plume transect line, this time parallel to plume motion, and cross-correlates this spatial profile in successive images to estimate velocity (Nadeau, 2011; Nadeau et al., 2011). However, this is less commonly found in the literature to date and, thus, is not currently available in our processing software.

2.2 Instrument design

The instrument makes use of sensors from Raspberry Pi cameras, which have been modified to enhance UV sensitivity by removal of their Bayer filter (Wilkes et al., 2016). This approach provides a significant cost reduction relative to the use of scientific-

TABLE	1	Instrument	power	consumption	in	different	states
INDEL		mountent	power	consumption		unicient	Junco

Instrument state	Power consumption (Watts)			
Sleep mode	0.96			
On (not acquiring data)	6			
On (acquiring data)	12			

grade commercial UV cameras-100 s USD versus 1,000-10000 s USD for this component of the instrument (Wilkes et al., 2017). Following sensor modification, a UV transmissive optical system, designed in Zemax ray tracing software, was constructed and mounted to the sensor. A triplet design was used to improve upon the previous lower-cost plano-convex singlet (Wilkes et al., 2016; Wilkes et al., 2017), which suffered somewhat from spherical aberrations at the edges of images and had a reasonably low light throughput. Unlike our previous work, which constructed optics holders from three-dimensional (3D) prints, here the triplet mount was constructed in machined black anodised aluminium, for improved precision and longevity. The resulting instrument field of view (FOV) is 28° x 21°. The UV bandpass filters were mounted behind the triplet, to mitigate the effects of changing wavelength transmission that occurs with changing light incidence angle on the filter (Kern et al., 2010b).

The remainder of instrument design is similar to that of Kern et al. (2015), including a network switch for internal and external communications and a co-aligned spectrometer for calibration of the camera optical depths (Figure 1). The most notable difference, other than the different UV cameras used (which has a considerable influence on the overall cost of the instrument), is our omission of a thermoelectric cooler (TEC) for temperature control of the spectrometer. Whilst previous versions of the Ocean Optics (now Ocean Insight) spectrometers have shown notable temperature instabilities (pixel-wavelength calibration shift and change in instrument line shape with changing temperature), the relatively new Flame spectrometer series (c. 2015) has a much-improved thermal stability, both as quoted by the manufacturer and under external testing (see Supplementary Material S1). We therefore omitted the TEC, thereby significantly decreasing the power consumption of the instrument (the TEC consumed 3 W in Kern et al., 2015), as well as saving costs and reducing instrument complexity.

Power consumption of the system is relatively low, since the camera utilises a low-power CMOS sensor. All components of the instrument run from 5 V; however, the instrument contains a 12 to 5 V voltage regulator inside, such that the instrument accepts standard 12 V power supplies such as car batteries. Overall power consumption is summarised in Table 1. In full operation the instrument consumes a maximum of 12 W; however, in most cases, operation will occur during only a small fraction of a day, especially since the instrument cannot acquire at night due to the lack of UV radiation. Controlled by the scheduling on the WittyPi HAT, the Raspberry Pi components of the instrument can be powered down when not in operation, leaving only the network switch still running; this consumes 1 W, therefore a time-averaged power consumption (assuming 6 h of operation), may be as little as ≈ 3.75 W. This is approximately half the power of the system

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presented in Kern et al. (2015), which consumes \approx 25 W in operation and \approx 2 W in sleep mode. In locations where light conditions for solar power are poor, such as at high latitudes or regularly cloudy areas, this difference will be particularly beneficial. Lower power consumption should also mean that the instrument can be run on fewer and/or smaller solar panels and batteries, again further reducing the overall cost of the installation.

Overall, the cost of parts for a one-off instrument (bulk purchasing would reduce part costs) is ≈5000 USD, which is primarily controlled by the cost of the UV spectrometer (≈3500 USD for spectrometer and associated optics). Note, these costs do not include data transmission or powering options, which will vary from installation to installation, depending on what existing infrastructure a group may have or the availability of technology in specific countries/regions. Whilst this cost is a notable increase on our discrete field campaign instrument cost (≈500 USD), it remains considerably cheaper than the current alternatives, which employ scientific-grade UV cameras. For example, the instrument presented by Kern et al. (2015) had a part cost (not adjusted for inflation) of ≈18,000 USD (C. Kern, personal communication), not including software licenses which were also 1000 s USD; our use of open-source software also avoids this additional cost.

2.3 Data acquisition

Data acquisition is controlled by software written in Python 3, which is freely available at https://github.com/twVolc/ PyCamPermanent. With the aim of making instrument control and processing of SO_2 camera data as accessible as possible, the code provides a graphical user interface (GUI) such that users should not need significant coding knowledge to work with the instrument. The principal aim is to ensure that robust and high-quality data capture and processing is as straightforward as possible for users.

The package contains a number of scripts that control the two cameras and spectrometer through a master script ("pycam_ masterpi.py"), which also handles external communication (e.g., for adjusting settings or manual acquisition control). Once the instrument has been setup to acquire automatically, it can then run in a headless state, where connection to the GUI or an external computer is not required.

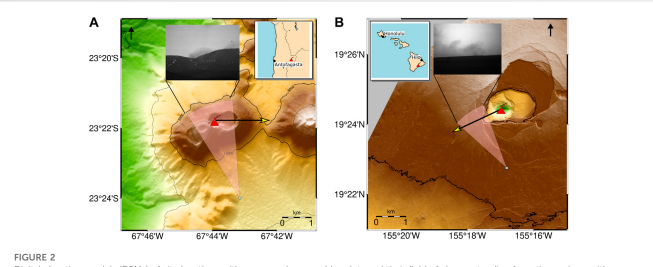
Automated image capture requires the automatic adjustment of camera and spectrometer shutter speed/integration time to avoid pixel saturation whilst ensuring sufficient light levels to obtain a good signal-to-noise ratio. To this end, image analysis performed on the instrument assesses the level of pixel saturation (relative to the maximum digital number measurable by the sensor—Defined by the sensor bit-depth) in every image/spectrum. User-defined parameters then allow the instrument to determine whether an increase or decrease in shutter speed/integration time is required for each image/spectrum. For images, the user predefines lower (*a*) and upper (β) limits for maximum pixel saturation (as a fraction), the number of pixels to average for this analysis (X), the number of image rows to use for the analysis, and whether to extract these rows from top-down or bottom-up. The option of only analysing a set number of rows allows us to only interrogate sky pixels in the analysis, omitting irrelevant pixels on the volcano flank. This is particularly important for snow-covered volcanoes, where snowcovered pixels appear very bright and would lead to an unwanted decrease in shutter speed if they were included in the analysis-It is important to optimise sky pixel intensity rather than the image as a whole. After this region of interest (ROI) has been extracted from the image, the brightest X pixels are found and the average digital number (DN) computed. Note, we choose to average a number of pixels rather than analysing only the brightest pixel, since erroneous "hot pixels" could interfere with the analysis. The saturation level of this average DN relative to the bit-depth DN is then computed. If this value is below α , the shutter speed of the next image will be increased; if the value is above β , the shutter speed of the next image will be decreased; for levels within the range α to β , the shutter speed will remain the same. For spectrometer integration time, the principle is the same; however, the user predefines a spectral window in which the saturation analysis takes place, rather than a spatial ROI. Typically this window may be 310-340 nm, ensuring that pixels do not become saturated in the spectrometer fitting window.

2.4 Data size and storage

The instrument holds a 1 TB SSD external storage device. This is the principal location for data storage. To ensure a backup, the data are also saved on the Raspberry Pi micro-SD card (128 GB), which can hold at least 100 GB of data (since it also holds the operating system of the Raspberry Pi). Images have a file size of 617 KB and spectra are 33 KB; of course, data accumulation depends on the acquisition rate and acquisition length per day. As a typical example, acquiring at a rate of 0.2 Hz for 6 h a day will generate ≈165 GB per month (2 TB per year); the storage will therefore be entirely full in 6 months, whilst back-up storage lasts less than 1 month. If frequent trips to download data are not possible, and data back-up is critical, an extra SSD external storage device could be added with relative ease; alternatively, larger SSDs are becoming more readily available. Data compression could also be explored in the future, to improve storage requirements. Where possible, telemetry of data would be desirable to allow near-realtime emission rate estimates and prevent the necessity for frequent trips to the instrument.

2.5 Data processing

Data processing can also be performed using the freely available PyCam software (https://github.com/twVolc/PyCamPermanent). Alongside in-house developed code, much of the processing makes use of the *pyplis* toolbox (Gliß et al., 2017), whilst spectrometer SO₂ retrievals are performed with *iFit* (Esse et al., 2020); light dilution correction of the spectrometer retrievals is also made possible using code associated with (Varnam et al., 2020). Information on the processing routines available can be found in those articles, and it is not within the scope of this paper to provide a detailed review of these routines. Using example datasets from Lascar and Kīlauea, Section 4 provides more details on some of the data processing algorithms available.



Digital elevation models (DEMs) of site locations with cameras shown as blue dots and their field of views extending from them, along with an example image from the on-band camera in each case. Red symbols indicate gas source locations. Typical plume motions are shown by yellow-headed arrows, but can vary significantly throughout the year. (A) Lascar (DEM from SRTM data); (B) Kilauea (DEM from LiDAR data taken from Mosbrucker et al., 2020).

We note that absolute automation of SO2 camera data processing remains troublesome, often requiring an expert user's retrospective interaction to ensure that processing routines have been optimally applied and to interrogate data quality. Others have achieved at least reasonable levels of automation (e.g., Kern et al., 2015; Delle Donne et al., 2017; Delle Donne et al., 2019), for instance, with the incorporation of a visibility index to omit data with poor visibility (Delle Donne et al., 2017) and a closed-path semi-circular integration line to capture gas emissions for all plume orientations (Kern et al., 2015). Of particular note, Delle Donne et al. (2019) investigated the difference between manual and automatic processing procedures, finding a reasonable agreement between their automatic processing routine and manual expert analysis (r^2 \approx 0.75); however, there clearly remains a notable disparity between the two, and this MATLAB-based (requiring a license) automation software is not open-source. In this respect, our work herein does not present any further steps in automation routines; however, it does provide open-source, user-friendly (with a graphical user interface) software which can promote the accessibility of SO₂ cameras to the wider volcanology community.

Data herein were processed with some manual interaction, for example, with the plume cross-sections optimized for the plume direction in each time series. Our software does, however, allow the use of multiple cross-sections, which would allow a quasi-semicircular integration line, as in Kern et al. (2015), ensuring the capture of gas emissions for all plume orientations; future software development will include the option to use true semi-circular lines. Background sky intensities also require manual derivation at times, although automated procedures, which harness the *pyplis* functionality, are available within our GUI. In general, we have found that at times of non-perfect conditions (e.g., heterogeneous cloud cover), it was optimal to manually define the background sky region for an image sequence. Other automated background sky retrieval algorithms have been proposed (Osorio et al., 2017) and may offer improved performance when moving towards a fully automated processing procedure; again, this could be the focus of future software development.

3 Field sites

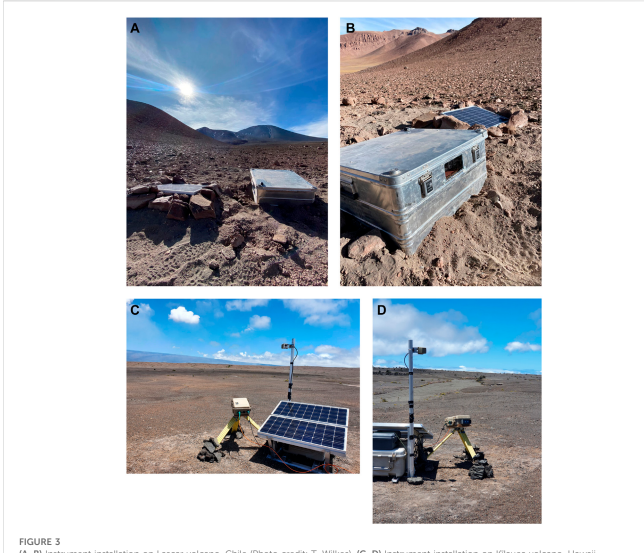
3.1 Lascar

Lascar (5,592 m; 23.37°S, 67.73°W) is a stratovolcano composed of 6 overlapping summit craters. It is predominantly andesiticdacitic in composition. In 1993 a VEI 4 eruption created a column reaching 20–22 km altitude and resulted in ash fall as far away as Buenos Aires (1,500 km SE of the volcano) (Global Volcanism Program, 1993). Current activity is primarily confined to fumarolic activity within the active crater, although during the camera installation a few small explosions were possibly heard, albeit with no clear visible associated phenomena.

The camera system was installed on Lascar on 18th May 2022, approximately 4,300 m from the crater (Figure 2A). The camera location provides a near-orthogonal viewing direction relative to the prevailing westerly winds. However, the plume has a tendency to swirl and stagnate within Lascar's crater before drifting eastwards. We also note that the front of the crater can at times obscure full view of the gas, which may lead to underestimations in emission rates at times when the plume does not loft above the crater. The installation is displayed in Figures 3A, B.

3.2 Kīlauea

Kīlauea (1,247 m; 19.41° N, 155.28° W) is a shield volcano on the Island of Hawai'i. Starting in May 2018, a series of fissure eruptions in the lower East Rift Zone accompanied drainage from



(A, B) Instrument installation on Lascar volcano, Chile (Photo credit: T. Wilkes). (C, D) Instrument installation on Kilauea volcano, Hawaii, United States (Photo credit: T. Pering).

the Halema'uma'u lava lake, which resulted in collapse of the summit caldera (Neal et al., 2019). During this period, SO_2 emission rates were estimated to have reached at least 200 kt/ day (2,300 kg/s) (Kern et al., 2020). These extreme conditions were modelled to have significantly impacted air quality hundreds of kilometres downwind of the volcano (Kern et al., 2020), whilst in the local vicinity lava flows and seismicity caused substantial damage and destruction of infrastructure (Neal et al., 2019).

The system was installed on Kilauea on 21st July 2022, approximately 2,700 m from the crater (Figure 2B). The northeasterly trade winds are relatively consistent, resulting in right to left gas motion through the image, which is almost exactly orthogonal to the viewing direction of the camera. This installation was augmented with data telemetry (up to 20 MB/s) from the instrument to a local observatory, allowing near-real-time processing of the data. This will enable rapid integration of SO₂ emission rates into hazard assessments at the volcano. The installation, along with radio antenna, is displayed in Figures 3C, D.

4 Results and discussion

4.1 Lascar

We here first present an in-depth analysis of a sample dataset retrieved shortly after camera installation on Lascar volcano, over the period 16:05:05–17:05:00 UTC on 20th May 2022. This is followed by a longer-term time series spanning 20th-21st May. Spectrometer FOV calibration is shown in Figure 4, along with the subsequent camera optical depth (τ_{SO2}) calibration. The FOV location is found through iterative calculation of the correlation coefficient between the spectrometer column density time series (Figure 4B) and τ_{SO2} time series of each pixel individually (Lübcke

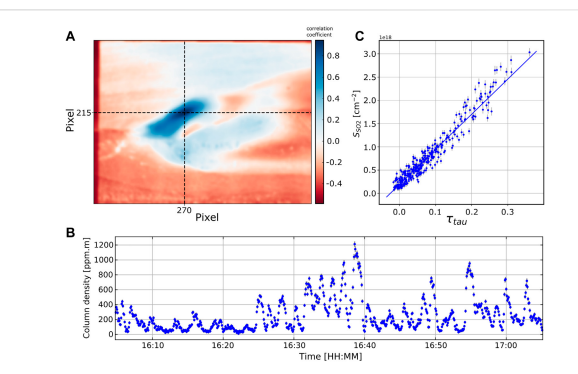
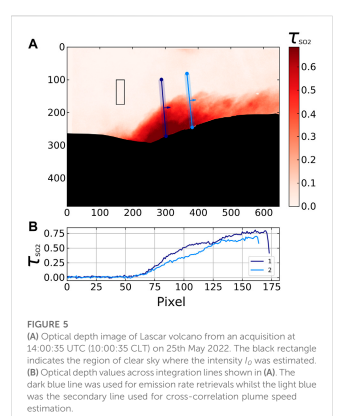


FIGURE 4

(A) Pearson correlation coefficients (colour scale) of spectrometer SO₂ retrievals with camera optical depths (τ_{SO2}), enabling determination of where the spectrometer field of view (FOV) is located within the camera image. In this case, the highest correlation coefficient is centered at pixel coordinate (270, 215). (B) Spectrometer-retrieved SO₂ slant column density retrieval time series (in UTC), highlighting that the FOV location is well-positioned to measure a wide range of column densities through time, therefore allowing a robust calibration of the camera optical depths. (C) Spectrometer SO₂ slant column density (S_{SO2}) retrievals versus camera optical depths measured in the spectrometer's FOV (as determined in A). The fitted calibration line is used to calibrate all pixel optical depths in the images.

et al., 2013; Gliß et al., 2017); Figure 4A displays the resulting correlation image and location of the spectrometer's FOV within the image. Once this is known, the spectrometer FOV can be fixed for subsequent datasets if desired. A regression between the spectrometer column densities and τ_{SO2} within the spectrometer's FOV can then be formed (Figure 4C), therefore allowing calibration of all pixel τ_{SO2} values. We note that the spectrometer's FOV is positioned just above the crater, such that it should be located within gas for most plume motions; however, there may still be periods of time where it is not located within the gas plume. In such periods it may be necessary to revert back to older calibrations, either earlier in the day or from previous days. Over time, we will build a catalog of instrument calibrations, which may then allow more robust selection of calibrations during periods where the plume is not in the spectrometer's FOV, for instance by identifying similar illumination conditions where a calibration was possible. A time series of the spectrometer SO₂ slant column density (SCD) retrievals (Figure 4C) also highlights that the spectrometer FOV is positioned such that it typically captures a wide range of SCDs in relatively short periods (<30 min). This ensures that camera optical depth calibration is unlikely to require significant extrapolation to larger values, which would result in greater measurement uncertainties. In this case the relationship between optical depth and SCDs was assumed linear, which is often reasonable for low SCDs. However, Kern et al. (2013) found that for high SO_2 burdens optical depths deviate significantly from the linear Beer-Lambert approximation, in part due to complex radiative transfer caused by in-plume aerosols and light dilution. The *pyplis* backend allows for non-linear regressions to be used in fitting camera optical depth to SCD, which can be changed within the pycam GUI.

Figure 5 shows a typical optical depth image from the instrument, also displaying the ICA line used to determine SO₂ emission rates for the time series discussed here. Emission rates can be retrieved using 4 distinct plume speed estimation algorithms; these are outlined in detail by Gliß et al. (2017), Gliß et al. (2018) and briefly summarised in Table 2. The cross-correlation technique introduced in Section 2.1 is represented by flow_glob, whilst the 3 other algorithms (flow_raw, flow_histo, flow_hybrid) utilise optical flow vectors with varying degrees of post-analysis. Figure 6 displays these effective velocities and the associated emission rate estimations; for clarity, we omit *flow_hybrid* from the plot, but these effective velocities, and thus emission rates, were extremely similar to the *flow_histo* algorithm. We note that, in this case and in our experience more generally, the optical flow algorithms typically estimate lower plume speeds than cross-correlation (flow_glob). This may be a result of cross-correlation preferentially fitting to larger pulsing events of gas (driven either by volcanic or meteorological factors), which have the most distinctive signature-notably standing out from general noise in a degassing



time series. This is supported by the higher optical flow speeds often coinciding with the larger mass loading along the ICA lines (peaks in the cross-correlation emission rate series); at these points the optical flow effective velocity is very similar to the cross-correlation plume speed (\approx 7.3 m/s in this case). In general, we therefore suggest the use of optical flow algorithms leads to more reliable emission rate estimates (Peters et al., 2015), especially the flow_histo (or flow_ hybrid) algorithm (Gliß et al., 2018), which retrieves a local average flow velocity and can potentially mitigate unreliable retrievals from the raw Farnebäck algorithm. We note, however, that there are relatively frequent periods where the *flow_histo* algorithm fails, resulting in no emission rate for these points (e.g., see Figure 6 or the "flow_histo LD corrected" sheet in Lascar spreadsheet of supplementary materials). To account for such gaps, either interpolating between recent plume speeds, or reverting to the flow_raw or cross-correlation algorithm, are solutions. More detailed discussions of plume speed algorithms and associated uncertainties can be found elsewhere (Peters et al., 2015; Gliß et al., 2017; Klein et al., 2017; Elias et al., 2018; Gliß et al., 2018).

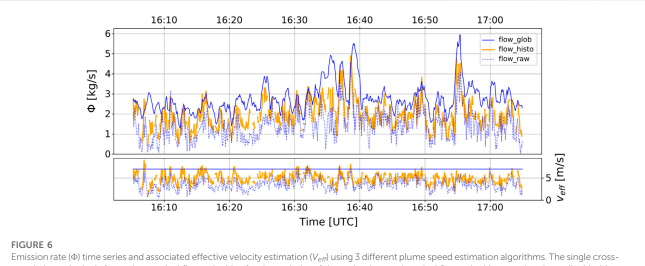
TABLE 2 Plume speed estimation algorithms, based on Gliß et al. (2017).

For the series shown in Figure 6, cross-correlation emission rates ranged between 1.3 and 6.0 kg/s, with a mean of $2.8 \pm 0.7 (\pm 1 \text{ SD}) \text{ kg/s}$, whilst *flow_histo* generated a mean emission rate of $2.0 \pm 0.7 \text{ kg/s}$, with the series ranging between 0.4 and 4.9 kg/s. The emission rate time series shows clear pulsing behaviour at Lascar, although it is difficult to definitively attribute this to volcanic activity, since the gas typically accumulates somewhat in the crater before exiting to where it can be observed by remote sensing equipment. It is therefore possible that wind turbulence at the summit is influencing the retrieved time series.

The above results were generated neglecting a light dilution (LD) correction, the phenomenon where light is scattered into the FOV of the instrument between the plume and the instrument; such light has not passed through the plume, so does not contain absorption features of SO₂, but contributes to the radiance at the detector. This can result in a significant underestimation of SO₂ emission rates (Mori et al., 2006; Kern et al., 2010a). Campion et al. (2015) proposed an image-based correction light dilution for camera optical depths, based on the changing measured intensity of the volcanic flank as it gets closer to the observer. We implement this correction using the pyplis API, finding that even in the dry Atacama air at >4,500 m altitude, there is still significant light dilution in the images (Figure 7). However, using this correction alone is only applicable to gas cell calibrated systems. Spectrometer-calibrated instruments require that the column densities measured by the spectrometer have also been corrected for associated light dilution effects; in fact, the spectrometer correction is much more critical than camera optical depth calibration, since the final calibration links column densities to somewhat arbitrary optical depths.

To correct the spectrometer-retrieved column densities we used the dual-band method proposed by Varnam et al. (2020). However, we found that the correction gave quite stochastic light-dilution factors that ranged from anywhere between 0 and 1. Correcting each spectrum individually therefore introduced large errors which resulted in a poor correlation between the spectrometer column densities and the associated optical depths in the camera images. The high correlation presented in Figure 4 suggests that the column density trend is accurate, even if absolute values are affected by light dilution. Below, we therefore present a method for stabilising the light dilution correction. Following Varnam et al. (2020), we first calculated the light dilution factor (LDF) and the associated corrected column density for each spectrum in the time series. We suggest the light dilution correction algorithm is likely to perform better on spectra with larger column densities, since the dual-band technique should see more disparity between the two bands than if fitting to low column densities. We therefore then discarded all spectra measuring column densities below 1E18 molecules cm⁻². We then took the median LDF of these

Name	Algorithm description
flow_glob	Cross-correlation
flow_raw	Raw output from Farnebäck optical flow algorithm
flow_histo	Histogram post-analysis to find a local average velocity vector for each integration line
flow_hybrid	Reliable motion vectors are used whilst unreliable ones are replaced based on histogram post-analysis



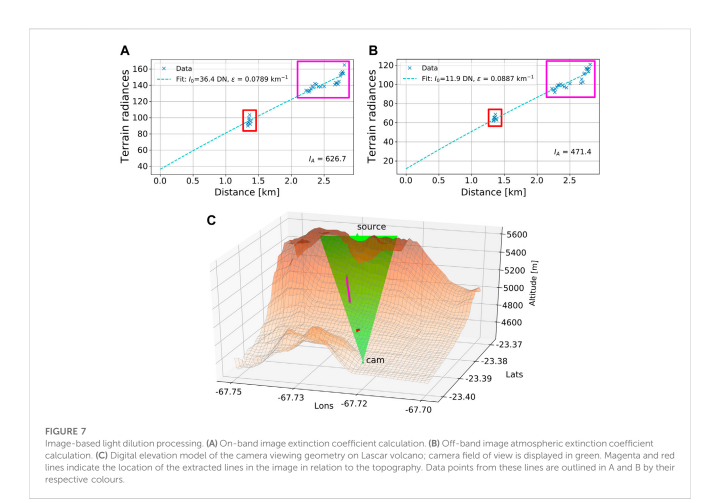
correlation velocity is faster than optical flow velocities for the majority of the series, but peak optical flow velocities match quite well with this crosscorrelation speed. We suggest *flow_histo* (bold orange) is likely to provide the most reliable data in general; however, at times this algorithm is unable to compute a reliable speed, therefore some parts of this time series are absent. Data from Lascar volcano on 20th May 2022; times are in UTC.

retrievals, in this case 0.25. Looking at the full time series again, we extracted all data points within $\pm 20\%$ of the median LDF (0.20 \leq LDF \leq 0.30). From this subset, a relationship between corrected and uncorrected column densities can be found, which is relatively linear within the range of values found here. This relationship was thus used to correct the initial column density time series. The correction increased the column densities by between ≈30% and ≈70%, depending on the initial uncorrected column density. This correction maintained the high >0.8 correlation between the spectrometer column densities and the associated optical depths within the spectrometer's FOV in the image series. We note, however, that this correction may be more complicated to implement in near-real-time when data are required rapidly for volcano monitoring purposes. Perhaps importantly, the general trends in the emission rates without a LD correction (Figure 6) are very similar to those with a LD correction (Figure 8), since the correction is relatively stable through time. This similartity suggests that emission rate time series without this correction could still have utility for rapid monitoring and response in real-time; the more accurate LD-corrected time series can then be generated post hoc.

The resulting light dilution-corrected emission rate time series increased somewhat (Figure 8A), with a mean cross-correlation emission rate of 3.8 ± 1.3 kg/s, whilst the *flow_histo* plume speed produced a mean emission rate of 2.8 ± 1.2 kg/s. From these mean values, the uncorrected retrievals therefore underestimate emission rates by 26% and 29% for *flow_glob* and *flow_histo*, respectively. These values are in the middle of the range of the examples presented by Campion et al. (2015); however, with the high altitude and dry conditions at Lascar, the level of light dilution is slightly higher than we had anticipated, highlighting that light dilution is likely to be a significant source of error for any measurements made at greater than 1-2 km from a volcano.

The longer-term series (Figure 8B), based on the same processing procedure outlined above (including light dilution correction), highlights the instrument's ability to perform shutdown/start-up sequences and adjust to changing light conditions throughout the day. Here, it is evident that the emission rates from the 2 days are significantly different. Whilst this could represent a true change in volcanic degassing, we cannot rule out the possibility that emission rates on 20th were incomplete measurements, with part of the plume being obscured by the crater as it drifted to the east. We note that plume speeds (v_{eff}) across the 2 days are relatively stable. The mean emission rate across the 2 days is 5.2 ± 3.1 kg/s, whilst the emission rate for only the 21st is 6.7 ± 3.1 kg/s. The emission rate is relatively variable, reaching a maximum of 17.5 kg/s; as mentioned above, this could be volcanogenic in nature but may also be associated with gas accumulation in the crater and subsequent large ejections driven by metrological conditions (e.g., wind turbulence).

Recent SO₂ camera measurements, in January 2019, are in good agreement with our measurements (4.7 \pm 1.4 kg/s), suggesting relative stability of the volcanic system over the last few years (Layana, 2022). However, Layana (2022) also found that, during a period where satellite-derived thermal anomalies at the volcano were absent, SO₂ emissions (measured in March and June 2018) were significantly lower-close to or below the detection limit of their SO₂ camera. This highlights the importance of more continuous monitoring, which can identify changes in activity over a broad range of timescales (from the order of minutes to sub-annual/annual and beyond); Thus, again we emphasize that discrete field campaigns cannot provide a complete picture of a volcano's state. Our measurements are somewhat higher than NOVAC scanner data between 2013-2016, which retrieved a mean of 2.6 ± 1.4 kg/s (Bucarey et al., 2020; Arellano et al., 2021). This may be a result of changing subsurface conditions in recent years, or a product of the longer-term nature of the NOVAC time series incorporating days of lower activity too. Discrete campaigns have also measured a broad range of emission rates on Lascar: 6.4 ± 2.5 kg/s in December 2012 (Tamburello et al., 2014); 27.8 kg/s in January 2003 (Mather et al., 2004). The general trend



appears to be a steady decrease in SO₂ degassing ($\approx 2003-2016$), with perhaps a new increase in emissions in the last few years; this is something which the new installation will be able to elucidate further in the coming months/years.

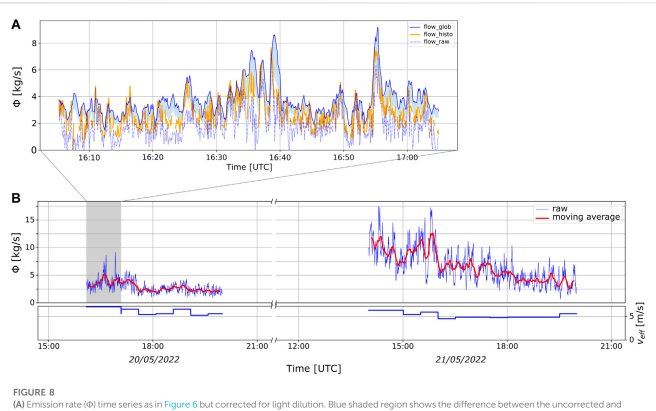
4.2 Kīlauea

We present here data acquired from Kīlauea volcano on 25th July 2022. Figure 9 displays a representative optical depth image acquired by the camera. Within the time period of this camera dataset, 9 traverse-mode DOAS measurements were also made from a vehicle transecting the plume \approx 1.5 km downwind of the Halema'uma'u Crater. Wind speeds for emission rate calculations of these traverses were retrieved from a weather station located at Sand Hill, about 2 km west of the active vent. These were then scaled by a factor of 1.2 to account for systematic underestimation of the wind speeds aloft by our ground-based measurements (Elias et al., 2018). Spectral and spatial analysis of the traverse data was performed according to standard DOAS analysis procedures (Platt and Stutz, 2008) using the DOASIS software (Kraus, 2006) in combination with a custom MATLAB code ("mDOAS") specifically developed for spatial analysis of Mobile DOAS

Using the same method as in Section 4.1, we corrected the *iFit* spectrometer retrievals for light dilution. In this case, the

image-based correction was not possible, due to the viewing geometry of the camera not providing enough volcanic flank. However, we propose that a first order light dilution correction should be possible simply by correcting the spectrometer retrievals, if we assume that image optical depths are uniformly affected by light dilution and that light dilution magnitude does not change significantly within a calibration window (typically 30–60 min). These assumptions are not strictly correct, however, since the correlation between image optical depth and column density (e.g., as in Figure 4B) remains strong following the spectrometer light dilution correction, correcting only the spectrometer retrievals may be adequate in many cases.

The traverse and camera datasets are displayed in Figure 10. Due to the significant differences in sampling frequency, and measurement location, exact comparison between the values is somewhat complicated and includes caveats; nevertheless, a basic comparison can still be useful. The 8 early DOAS traverses found an average emission rate of $18.7 \pm 8.0 (\pm 1 \text{ SD})$ kg/s, with a minimum measurement of 6.2 kg/s and maximum of 27.6 kg/s. Whilst there is a disparity, these measurements are in reasonably good agreement with the camera's contemporaneous first acquisition period (≈ 20 : 00–21:30 UTC), which has a mean emission rate of 12.7 ± 5 kg/s and a maximum of 28.0 kg/s. For the period 11:00-01:00 UTC (25/26th July), the difference in emission rates is more notable, with the single

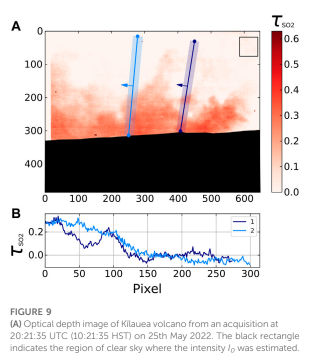


(A) Emission rate (Φ) time series as in Figure 6 but corrected for light dilution. Blue shaded region shows the difference between the uncorrected and corrected emission rates for the *flow_glob* series. (B) A longer-term series spanning 20th-21st May based on *flow_glob* plume speeds (blue) with a 10-min moving average (red). Plume speed (v_{eff}) is also displayed and is relatively consistent across the 2 days, unlike the emission rate which is significantly higher on the 21st.

traverse measurement of 23.5 kg/s, whilst the mean camera emission rate was just 7.5 \pm 3.0 kg/s.

It is not clear what has caused this large disparity in the second period (albeit comparing with only one contemporaneous traverse measurement), although some possible influences may be: differences in wind speed estimates (and the sources of these estimates); In-plume aerosol scattering of light; incomplete capture of the full gas plume by the camera, due to its viewing geometry and the crater geometry; non-orthogonal plume motion relative to camera viewing direction (e.g., Klein et al., 2017). Regarding the latter, using the plume centers based on traverse measurements, we can calculate the difference between the plume's motion and the camera's focal plane azimuth. For the first acquisition period the difference ranged between 13° and 25°, whilst for the second period this difference was 17°. Although this can lead to an underestimation in emission rates, since the second period's angular deviation falls within the bounds of the first period, it is unlikely that this is the principal cause for the larger emission rate disparities in the second period. Alternatively, incomplete capture of the gas plume by the SO₂ camera could be a notable source of error for the camera measurements. The low lying crater geometry means that at times the plume can hug the ground, therefore making it difficult to fully capture emissions close to the source with the SO2 camera. This would be particularly notable in stronger wind conditions, which would reduce the degree of plume lofting as it leaves the crater. It is difficult to quantify this error, but it certainly could contribute to a significant underestimation in emission rates. Using optical flow motion vectors, it should be possible to identify periods where plume motion is less buoyant (more horizontal motion) and flag such periods as potentially having unreliable emission rate estimates; longer-term datasets will allow deeper investigation into this.

Disparities in plume speeds, and associated emission rates, are somewhat more quantifiable; Although we note that direct comparisons may be troublesome, since the SO₂ camera images and weather stations are not collocated. Furthermore, neither technique is a direct measurement of the plume's velocity at its altitude above the traverse measurements. Details of the methodology employed for this comparison are outline in Supplementary Material S2. During the final traverse (23: 56 UTC), in the second camera acquisition period, corrected image-based horizontal plume speeds had a mean of 6.0 m/s for optical flow analysis, whilst cross-correlation measured 9.0 m/s. As discussed in Section 4.1, the discrepancy is likely due to crosscorrelation fitting to large pulses of gas exiting the crater, which are likely relate to larger gusts of wind and/or changes in volcanic degassing vigor. Indeed, much like in Figure 6, the optical flow speeds reach similar (and slightly larger) values at times; however, for the specific 10-min period of the traverse (adjusted for time for the plume to drift from crater to traverse location), the values are



20:21:35 UTC (10:21:35 HST) on 25th May 2022. The black rectangle indicates the region of clear sky where the intensity I_o was estimated. **(B)** Optical depth values across integration lines shown in **(A)**. The dark blue line was used for emission rate retrievals whilst the light blue was the secondary line used for cross-correlation plume speed estimation.

notably lower. The Sand Hill weather station measurement at this time resulted in a scaled plume speed of 10 m/s. Using our plume measurements to rescale the traverse emission rates, we find emission rates of 14.1 and 21.2 kg/s for optical flow and cross-correlation, respectively. Whilst these values remain higher than the SO₂ camera average for this period, the camera data reach 12.9 kg/s at 23:53:15 UTC and 13.8 kg/s at 23:54:30 UTC; the center of the traverse was reached at 23:56: 59 UTC and is located \approx 2–3 min downwind (1.5 km from

crater). For comparison, the difference between cameraderived and weather station-derived plume speeds for the first acquisition period (20:00–21:30 UTC) is markedly smaller; camera-derived speeds averaged 6.9 and 9.9 m/s, for optical flow and cross-correlation, respectively, whilst the weather station-derived speed was 9 m/s throughout.

Although it seems to be a contributing factor, it is very unlikely that the above plume speed difference is the only source of disparity between the traverse and SO₂ camera measurements. Whilst the academic literature currently lacks long-term comparisons between SO₂ camera measurements and traverse-/scanning-mode DOAS measurements, some previous work has found good agreement between DOAS traverses and UV camera data in discrete campaigns (Varnam et al., 2021). Conversely, however, de Moor et al. (2017) highlighted that their permanent scanning DOAS system systematically measured lower emission rates relative to contemporaneous traverse DOAS measurements. They suggest that one possible cause is having a sub-optimal scanning position that fails to capture the full plume in a large number of scans, due to variable wind direction blowing the plume away from fixed-location scanners. We note that similar issues, associated with camera location/viewing geometry, are particularly pertinent for permanent installations; during discrete field campaigns imaging location can be chosen on a daily (or sub-daily) basis depending on plume conditions. On many volcanoes it can be difficult to find a suitable viewing geometry to ensure that, given any wind direction and degassing strength, the camera will always be able to capture complete and robust emission rates. Indeed, in many cases it may be advisable to install multiple systems around a target, providing various vantage points that could facilitate more robust retrievals under a range of meteorological and volcanic conditions (e.g., Delle Donne et al., 2022). Scanning DOAS networks often do exactly this on the most comprehensively monitored volcanoes (Burton et al., 2009; Galle et al., 2010). The relatively low cost of our new permanent camera design would facilitate such ambitions. Additionally, we also suggest that longer-term comparisons

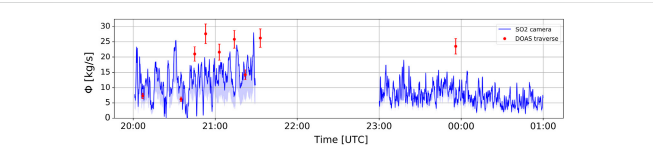


FIGURE 10

Permanent camera SO₂ emission rate (Φ) time series at Kilauea on 25th and 26th July 2022 (UTC; all measurements were 25th July HST), with contemporaneous DOAS traverse measurements (red dots). Camera retrievals were made with the cross-correlation (*flow_glob*) plume speed estimation algorithm. The magnitude of the light dilution correction (applied through the instrument's spectrometer iFit retrievals) is displayed by the shaded blue region. Agreement between the camera and traverse measurements is reasonably good for the 25th measurements, whilst on the 26th the SO₂ camera retrieves significantly lower emissions rates, albeit with only one traverse measurement for comparison in this sequence. Note that the traverse measurements were performed ≈1.5 km downwind of the source, therefore with a plume speed of ≈8–10 m/s we may expect a lag in measurements of ≈150–190 s.

between SO_2 cameras and scanning-/traverse-mode DOAS measurements, as well as satellite retrievals (e.g., TROPOMI), are critical to understanding the validity of each measurement technique under a wide range of conditions.

5 Conclusion

We have presented the design of a new permanent UV camera system for volcanic SO_2 emission rate measurements, followed by installation of the instrument on 2 active volcanoes (Lascar, Chile; Kīlauea, Hawaii, United States). The instruments make use of a suite of previously developed open-source Python packages to supplement custom software, ensuring robust/standardised and comprehensive data analysis. We have highlighted how the relatively low cost and power consumption of these instruments makes them suitable for widescale deployment on volcanoes in a range of different environments around the globe. Although we note that longer-term testing of instrument performance is still required, such instruments have the potential to provide valuable datasets for comparison with alternative ground-based remote sensing instruments (i.e., DOAS scanning networks and traverse measurements) as well as ground-truthing satellite retrievals.

We provide example datasets from Lascar (Chile) and Kilauea (Hawaii, United States). At Lascar, mean emission rates of 5.2 \pm 3.1 kg/s agree well with recent (2019) measurements but are somewhat higher than earlier (2013-2016) longer term NOVAC time series (2.6 ± 1.4 kg/s). On Kilauea we presented a comparison with contemporaneous traverse-mode DOAS measurements, which showed a reasonable agreement, albeit with the traverse emission rates $(18.7 \pm 8.0 \text{ kg/s})$ typically being higher than the camera-derived rates (12.7 \pm 5 kg/s for the first series). A later measurement series gave lower camera-derived emission rates that were typically half the value of a traverse measurement in this window. We found that differences in wind speed estimates likely contributed significantly to the disparity; however, it is likely that other effects contributed too (e.g., in-plume scattering of light, incomplete capture of the entire plume by camera measurements).

We particularly emphasise the importance of permanent/ continuous monitoring equipment in volcanic gas measurements, since volcanic activity is notably transient and variable in nature over a wide span of timescales. Whilst of course having significant value to the scientific community, the danger of discrete field campaigns is in interpreting such measurements as representative of that volcano outside of the measurement window. In some branches of volcanology (e.g., seismology), continuous monitoring is already somewhat commonplace; however, although there are a number of installations for permanent gas monitoring (e.g., most notably NOVAC), campaign-style measurements are still routine in a number of places across the globe. This is likely influenced by the cost and complexity of permanent systems. The instrument presented herein has the potential to aid the transition to more continuous geochemical monitoring of hazardous volcanoes across the globe, which in turn stands to improve our understanding of hazardous volcanic events and inform eruption forecasts (Kern et al., 2022).

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

Author contributions

TW wrote the manuscript, developed the instrument and installed the instrument on Lascar. TP contributed to the manuscript, contributed to instrument development and installed the instrument on Kīlauea. FA contributed to instrument development. SL installed the instrument on Lascar and contributed to instrument development. PN and CK installed the instrument on Kīlauea; CK also contributed to instrument development. AM contributed to instrument development. MA contributed to field work in Chile. CZ contributed to instrument development.

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Conflict of interest

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

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/feart.2023.1088992/ full#supplementary-material

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