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Hydrogen emissions from Erebus volcano, Antarctica

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Abstract

The continuous measurement of molecular hydrogen (H₂) emissions from passively degassing volcanoes has recently been made possible using a new generation of low-cost electrochemical sensors. We have used such sensors to measure H₂, along with SO₂, H₂O and CO₂, in the gas and aerosol plume emitted from the phonolite lava lake at Erebus volcano, Antarctica. The measurements were made at the crater rim between December 2010 and January 2011. Combined with measurements of the long-term SO₂ emission rate for Erebus, they indicate a characteristic H₂ flux of 0.03 kg s⁻¹ (2.8 Mg day⁻¹). The observed H₂ content in the plume is consistent with previous estimates of redox conditions in the lava lake inferred from mineral compositions and the observed CO₂/CO ratio in the gas plume (~0.9 log units below the quartz–fayalite–magnetite buffer). These measurements suggest that H₂ does not combust at the surface of the lake, and that H₂ is kinetically inert in the gas/aerosol plume, retaining the signature of the high-temperature chemical equilibrium reached in the lava lake. We also observe a cyclical variation in the H₂/SO₂ ratio with a period of ~10 min. These cycles correspond to oscillatory patterns of surface motion of the lava lake that have been interpreted as signs of a pulsatory magma supply at the top of the magmatic conduit.

Keywords: Erebus volcano Hydrogen Magma redox conditions Lava lake Volcanic degassing

Introduction

Hydrogen is one of the most abundant trace species in volcanic emissions (e.g. Giggenbach 1987; Oppenheimer et al. 2012) and is an essential participant in key redox reactions that take place in magmatic gases, e.g.

H₂+12O₂=H₂O

and

H₂S+2H₂O=SO₂+3H₂.

At Erebus volcano, Antarctica, emissions to the atmosphere result from the sustained supply of gas via the persistently active lava lake and surrounding fumaroles. The gas composition of the plume provides valuable insights into redox conditions of the lava lake, assuming thermodynamic equilibrium is achieved between the gas phase and the melt. At Erebus volcano, the volcanic plume is mainly the result of passive degassing of the magma within the lava lake. Contrary to the conventional consideration that the redox state of a magma is buffered during decompression, recent chemical modelling of the gas phase in ascending magmas (Burgisser and Scaillet 2007; Burgisser et al. 2008) has suggested an evolution of the magma redox state during ascent, partly reflecting initial volatile contents. Measurement of redox-sensitive magmatic gas species such as molecular hydrogen will enable such models to provide better constrained "inversions" of surface geochemical signatures.

The recent development of highly portable and readily deployed multi-species gas sensing systems (Shinohara 2005; Aiuppa et al. 2005, 2006; De Vito et al. 2007) has enabled measurements of volcanic gas ratios for extended periods and in some cases operationally (Aiuppa et al. 2010a, b). Such multi-species approaches complement ultraviolet and infrared spectroscopic applications (Oppenheimer 2010) to enable measurement of abundances and fluxes of a range of gas species. Until very recently, there has not been a practical means for extended surveillance of hydrogen (H₂) abundance in dilute volcanic plumes (Aiuppa et al. 2011; Shinohara et al. 2011). Here, we use a "multi-gas" instrument incorporating a sensor to measure H₂ in the plume emitted from the lava lake of Erebus volcano. Erebus is of particular interest because of the emerging evidence for redox change associated with magma ascent, with the most reducing conditions found in the lava lake itself (Oppenheimer et al. 2011). Despite challenging conditions at the summit crater, measurements were possible for several hours per day spanning a week. Our initial aims were to assess implications of the measurements for lava lake redox conditions and to identify any rapid variability in gas composition of the plume that would corroborate previous observation of cyclical behaviour of the lava lake (Oppenheimer et al. 2009).

Methodology

The measurements were made between 6 December 2010 and 3 January 2011 using a purpose-built multi-gas instrument and a LI-COR® LI-840 CO₂ and H₂O infrared analyser. The multi-gas instrument incorporated H₂, H₂S and SO₂ electrochemical sensors, a nondispersive infrared (NDIR) sensor for CO₂, and a sensor for temperature and humidity

measurements. The sampled gas is circulated via a miniature 12-V rotary pump through the sensors (Aiuppa et al. 2011). The H₂, H₂S and SO₂ sensors produce an electrical current in response to the target gas entering the electrolyte and oxidizing or reducing the electrode. This current is proportional to the concentration of the target gas in the total gas volume. The electrochemical sensor for SO₂ (City Technology, sensor type 3ST/F) has a calibration range of 0-30 ppmv, an accuracy of ±2 %, a repeatability of 1 % and a resolution of 0.5 ppmv. The electrochemical sensor for H₂ (City Technology, sensor type 3HYT) has a calibration range of 0-500 ppmv, an accuracy of ±5 %, a repeatability of 2 % and a resolution of 2 ppmv. The NDIR CO₂ sensor (model Gascard II) is calibrated for 0–3,000 ppmv and has an accuracy ±2 % and a resolution of 0.8 ppmv. In addition to the gas sensors, temperature and relative humidity (RH) sensors (Galltec) are mounted in the instrument, providing a measuring range of 0-100 % RH and an accuracy of ±2 %. All sensors were housed inside a weatherproof box, with the ambient air sampled via Teflon tubing connected to a HEPA filter fed through an opening in the box. The sampled gas was ejected via an outlet likewise fed through a hole in the case. The sampled gas was heated to ~30 °C on its way through the first hose to prevent freezing and other problems related to the low ambient temperatures (below -25 °C).

An on-board data-logger card in the multi-gas instrument captured measurements at a rate of 0.5 Hz, while the LI 840 output was logged at 1 Hz via a netbook PC. Both instruments were always started simultaneously. The multi-gas instrument was recalibrated using standard gas mixtures in the laboratory (accurately measured by gas chromatography) before and after the campaign and showed very little drift (<5 %).

The LI-COR® LI-840 instrument was operated for intercomparison with the Gascard II and humidity sensors in the multi-gas unit. The LI-840 is a non-dispersive infrared gas analyser equipped with a dual wavelength, infrared detection system allowing measurements of CO_2 and H_2O in the gas phase in the range of 0–3,000 ppmv and 0–80 parts per thousand volume (pptv), respectively. The accuracy was better than 1.5 % for both species and cross-sensitivity is <0.1 ppmv CO_2 /pptv H_2O for H_2O and <0.0001 pptv H_2O /ppmv CO_2 for CO_2 .

Both instruments were deployed intermittently on the crater rim of Erebus with their inlet filters placed side-by-side directly sampling the plume (Fig. 1). Measurements were possible when the plume was grounded and most data were collected at the "pump site" situated on the northern rim of the crater \sim 220 m vertically above and \sim 150 m horizontally from the lava lake. This site has consistently proved to be suitable for in situ plume sampling and sensing owing to the prevailing wind direction (Zreda-Gostynska et al. *1997*; Ilyinskaya et al. *2010*). Both instruments were powered by a 12-V DC battery, which sustained 6–10 h of unattended operation.

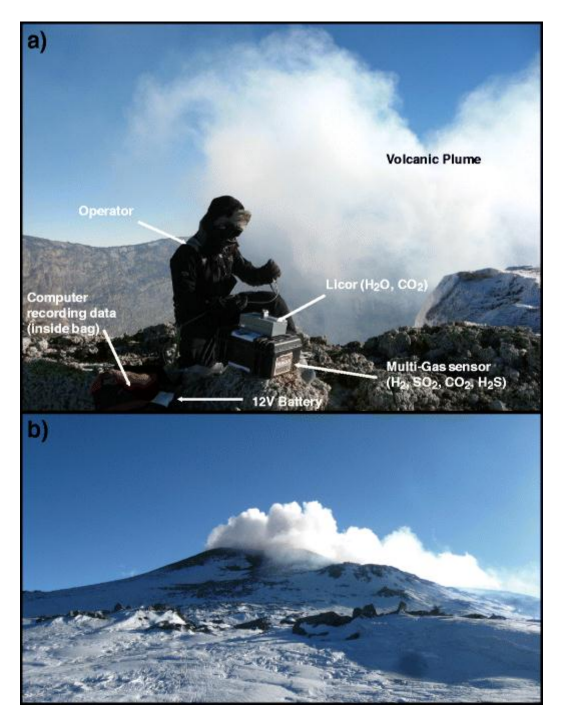


Fig. 1
a Typical field operating conditions during deployment of the LI 840 and multi-gas sensor at the crater rim. Gases are pumped through both instruments via narrow hoses connected to particle filters. b View of Erebus (looking south) during ideal plume sampling conditions on 26 December 2010. Turbulent airflow resulted in grounding of the plume at the sampling site

Data processing

Mixing ratios of SO₂, H₂O, CO₂ and H₂ in the gas phase are retrieved (in parts per million volume) directly from the laboratory-calibrated sensors using the proprietary "840–500" software for the LI-840 and in-house software (developed at INGV Palermo) for the multi-gas

instrument. The raw data collected by both instruments show a good correlation between all measured species. The good agreement between the CO_2 measurements obtained by the multi-gas instrument and those obtained by the LI-840 provides a validation of the performance of the Gascard II sensor. Response times of the different sensors vary slightly but are all rapid, requiring a few seconds to approach maximum reading. Figure 2 shows an example of a typical dataset, recorded on 27 December 2010. All species' abundances are highly correlated, and there is little or no time lag between each sensor's response to increasing gas mixing ratios. Typical abundances measured at the crater rim range from 0.3 to 1.3 ppmv for H_2 and SO_2 , 400 to 500 ppmv for CO_2 and 800 ppmv for H_2O .

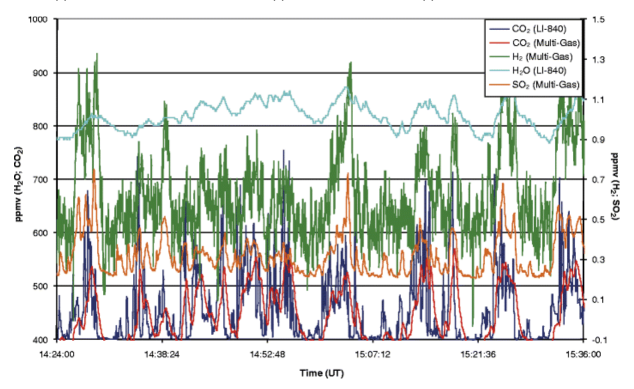


Fig. 2
Example of time series for gas mixing ratios obtained from both instruments (the multi-gas instrument and LI 840). This 1-h-long time series is an extract from a 10-h-long run acquired at the crater rim on 27 December 2010. All gas abundances are reported in in parts per million volume

In order to convert the raw abundance data into reliable measurements, several processing steps were applied. Firstly, the small difference in response times between sensors was corrected for. Secondly, the sensor signal resulting from cross-sensitivity with other gases, characterised by laboratory measurements, was subtracted. The differences in response time for the different sensors were corrected by finding the lag times from correlation analysis of the various time series. Laboratory tests were performed using a set of gas standards circulated through the multi-gas instrument in order to determine the cross-sensitivity of the hydrogen sensor to other species. Carbon monoxide was a particular concern because of its abundance in the Erebus gas/aerosol plume (Wardell et al. *2004*; Oppenheimer and Kyle *2008*). Its effect was tested for a range of CO from 7 to 500 ppmv. Mixed CO and H₂ gas calibrations were also carried out. These tests revealed a constant 4 % cross-sensitivity of the H₂ sensor due to the presence of CO. The H₂ sensor was found not to be sensitive to

 H_2O , CO_2 or SO_2 . Although there was no CO sensor in the multi-gas instrument, an estimate of CO abundances can be made, point-by-point, from measured CO_2 abundance and using a CO_2/CO molar ratio of 13 ± 1 obtained from open-path FTIR spectroscopic measurements (Oppenheimer et al. *2009*; Ilanko personal communication). At each point, 4 % of the estimated CO value was subtracted from the H_2 signal to correct for the cross-sensitivity (Fig. 3).

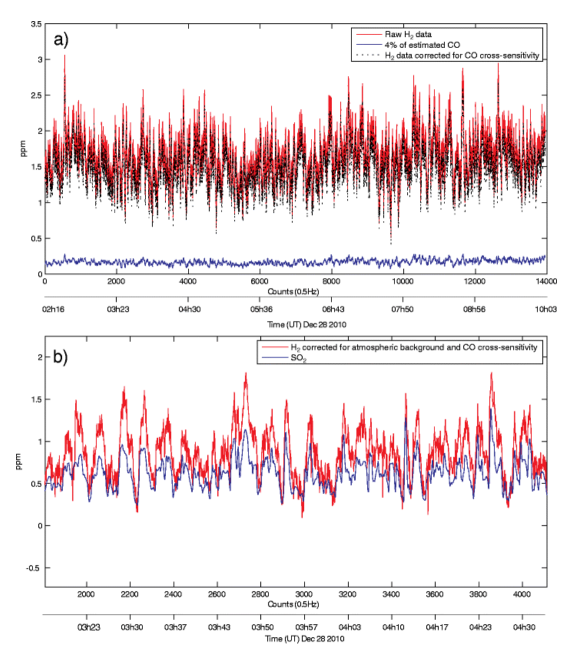
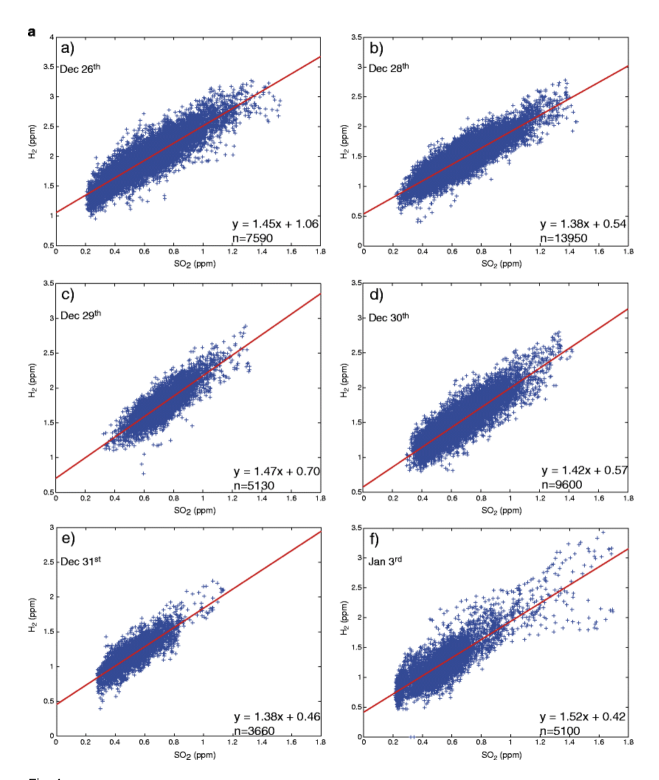


Fig. 3 a Raw (red trace) and corrected (dashed trace) time series for H_2 . The corrected time series is obtained by estimating the signal that would be due to presence of CO in the plume (blue trace). The abundance of CO has been estimated from the corresponding Gascard II CO_2 time series, from which a constant background atmospheric value of 387.5 ppmv has been subtracted, and a CO_2/CO molar ratio of 13 (from Oppenheimer et al. 2009). b H_2 and SO_2 time series after correction for atmospheric background H_2 and SO_2 cross-sensitivity and after alignment of both times series using an offset calculated from the maximum correlation factor between the time series

Results

Useful data were only acquired during favourable winds that resulted in grounding of the plume at the crater rim (Fig. 1b). We obtained 25 h of good quality observations at a sample rate of 0.5 Hz over the 180 h of data collection. As well as the variable weather conditions, delays in stabilising the internal temperature of the multi-gas instrument were also responsible for the limited collection time. It is worth noting that Strombolian eruptions, which are occasionally observed at Erebus (Aster et al. 2003; Dibble et al. 2008), did not occur during the acquisition of this dataset; only the "passive" plume emitted from the lava lake was sampled. Figure 4 shows 12 scatter plots for measurements recorded during the last week of December 2010, which offered the best conditions for plume sampling. Results from 6 days of data yield a daily mean H₂/SO₂ molar ratio between 1.38 and 1.52 (Fig. 4a); the average ratio for the whole week being 1.44. Scatter plots for the CO₂ and SO₂ measurements yield CO₂/SO₂ molar ratios varying between 36 and 45 (Fig. 4b), with the week's average being 40. We neglect the CO₂/SO₂ ratio obtained for 3 January which shows much higher variability, which we ascribe to contamination from nearby fumaroles. The intercept of first order linear regression through the scatter plots of H₂ vs. SO₂ and CO₂ vs. SO₂ should correspond to the atmospheric background H₂ and CO₂ abundances, respectively (since ambient SO₂ is very low, less than 10 pptv). In fact, we find values for ambient H₂ between 1.06 and 0.42 ppmv and background CO₂ values between 433 and 385 ppmv. These are both good approximations to expected atmospheric background abundances for the two gases. For instance, measurements from December 2010 at the South Pole weather station (available at http://www.esrl.noaa.gov/) indicate a mixing ratio of 387.5 ppmv for CO₂ and 0.54 ppmv for H₂. This station is the closest Antarctic research station routinely measuring atmospheric gas abundances at altitude (2,900 m a.s.l, c.f. the altitude of the pump site of ~3,700 m). A test run using the multi-gas instrument on 7 December carried out near Lower Erebus hut (2 km from the crater) also yielded stable H₂ readings of ~0.5 ppmv, though CO₂ readings fluctuated between 400 and 350 ppmv with temperature drift during acquisition. These estimates of the ambient mixing ratios of the two gases give further confidence in the performance of the multi-gas instrument sensors. While H₂O was being recorded simultaneously by both the LICOR and multi-gas instruments, changes in the ambient atmospheric humidity, the influence of nearby low-temperature fumaroles and conceivably absorption of H₂O on tubing and filters precluded reliable retrieval of water content of the plume.



a H₂–SO₂ and b CO₂–SO₂ scatter plots from 6 days of sampling of the Erebus plume under favourable conditions. a 11:20 to 16:33, 26 December 2010, UTC. b 02:16 to 10:01, 28 December 2010, UTC. c 11:07 to 13:58, 29 December 2010, UTC. d 06:25 to 11:45, 30 December 2010, UTC. e 04:10 to 06:12, 31 December 2010, UTC. f 22:42 to 01:32, 3 January 2011, UTC. Regression lines are shown in red and corresponding parameters are displayed on the lower right corner of each plot

Further inspection of our dataset reveals small but clear variations in the retrieved gas ratios which appear cyclical. Figure 5 shows the evolution of the H_2/SO_2 and SO_2/CO_2 ratios for 23 min on 26 December. Three complete cycles of about 7 to 8 min length can clearly be identified in both time series. The H_2/SO_2 ratio for \sim 3 h on 3 January 2011 is also reported on this figure and shows 15 cycles of about 10 min. Figure 5 also presents the corresponding pseudo-periodogram obtained for these \sim 3 h using a continuous Morlet wavelet transform analysis (Goupillaud et al. *1984*) of the time series. The pseudo-periodogram shows a strong transform modulus with a cycle of 8 to 12 min. Similar pseudo-periodograms have been produced for all the time series for which data are presented in Fig. 4, and all reveal cycles with periods of 7 to 14 min. In addition, some pseudo-periodograms show weaker signal strength at a shorter period of 3 to 5 min.

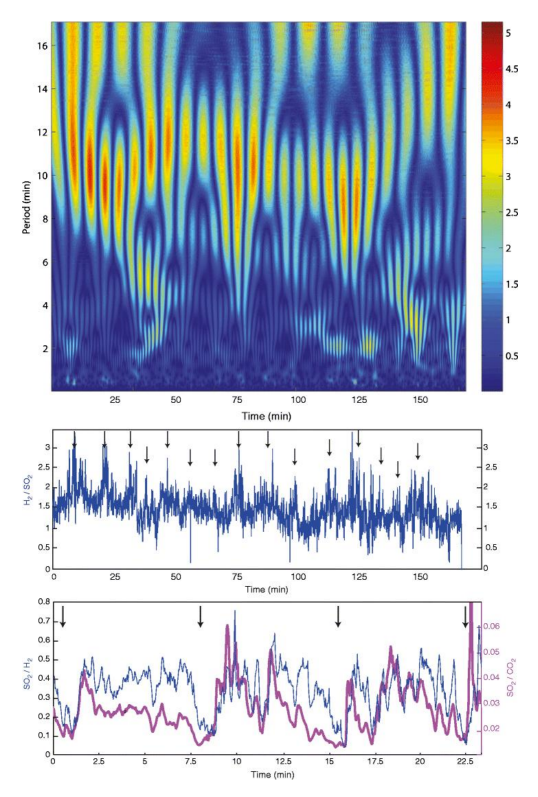


Fig. 5 Morlet wavelet transform pseudo-periodogram computed from a 3-h time series of the H_2/SO_2 ratio obtained from a 10-h-long run of the multi-gas instrument on 3 January 2011. Note the strong transform modulus emerging steadily at a period of \sim 10 min. The central panel shows the same 3-h time series of the evolution of the H_2/SO_2 ratio in which \sim 10-min cycles can be observed. Black arrows indicate the peak in H_2/SO_2 ratio of each cycle. The lower diagram shows the evolution of the SO_2/H_2 and SO_2/CO_2 ratios for the first 1,400 s (\sim 23 min) of a time series obtained from an 8-h-long run of the multi-gas instrument on 26 December 2010. This lower diagram shows three complete cycles of \sim 8 min each and shows reasonably clearly that SO_2/H_2 follows SO_2/CO_2 . Black arrows indicate peak in H_2/SO_2 ratio of each cycle

Pseudo-periodograms were produced for the CO_2/SO_2 and CO_2/H_2 ratio time series and reveal similar periodicities. The CO_2/H_2 ratios vary mostly between 20 and 35. Lower and upper extremes are approximately 15 and 50, respectively. The periodicity is noteworthy since it suggests preservation of a source signature despite the passage of the plume within the crater (and the possibility for mixing and homogenization of the time-varying signal), from the lava lake to the pump site. Time series of the SO_2/H_2 and SO_2/CO_2 ratios were constructed using background H_2 and CO_2 atmospheric values determined by the intersection of the linear regression with the H_2 or CO_2 -axis for each day (Fig. 4) except when that intercept was higher than the lowest measured H_2 or CO_2 value, in which case this lowest H_2 or CO_2 value was used as the background.

Discussion

H₂ contribution to the Antarctic atmosphere from Erebus

We have estimated the H_2 flux from Erebus volcano using the time-averaged (over approximately a decade of observations) SO_2 flux of 0.71 ± 0.3 kg s⁻¹ (Sweeney et al. 2008) and the measured H_2/SO_2 ratio. Using our mean H_2/SO_2 molar ratio of 1.44 (equivalent to a H_2/SO_2 mass ratio of 0.045), we estimate the mean H_2 flux at Erebus volcano as 0.03 kg s⁻¹ (2.8 Mg day⁻¹). Note, however, that the SO_2 emission rate itself fluctuates cyclically (from 0.17 to 0.89 ± 0.20 kg s⁻¹ in December 2005; Boichu et al. 2010). This estimated H_2 flux from Erebus is the largest recorded point source of H_2 to the Antarctic atmosphere. The Erebus source amounts to ~1 Gg a⁻¹ of H_2 corresponding to 0.004 to 0.02 % of the total global anthropogenic emission (the global anthropogenic emission of H_2 from the use of fossil fuels is estimated at 5 to 25 Tg a⁻¹) (Novelli et al. 1999). For comparison, the hydrogen flux at Mt. Etna has been estimated at ~0.00065 Tg a⁻¹ (Aiuppa et al. 2011).

Oxidation state of the Erebus lava lake

Based on a mean bulk plume SO_2/H_2O molar ratio of 0.023 obtained by FTIR spectroscopy (Oppenheimer et al. 2009), we can convert the mean multi-gas-measured H_2/SO_2 ratio to an H_2/H_2O molar ratio of 0.033. From this ratio, we can calculate the corresponding oxygen fugacity based on the redox reaction in Eq. (1).

The equilibrium constant, K, for this reaction is given by:

$$K = f_{H_2} \times (f_{O_2})_{12} f_{H_2O}$$

where
$$f_{H_2=\gamma_{H_2}\times P_{H_2}}$$
; $f_{O_2=\gamma_{O_2}\times P_{O_2}}$; $f_{H_2O=\gamma_{H_2O}\times P_{H_2O}}$

with
$$P = x \times P$$

and where f_i is the fugacity of the ith species, γ_i is the fugacity coefficient of the ith species, P_i is the partial pressure of the ith species, x_i is the mole fraction of the ith species and P is the total gas pressure.

This yields:

$$(f_{02})_{12}=K\times f_{H2}Of_{H2}=K_{V}H_{2}O\times XH_{2}O\times P_{V}H_{2}\times XH_{2}\times P$$

and therefore

$f_{O2}=(K\gamma_{H2}O\times x_{H2}O\gamma_{H2}\times x_{H2})2.$

At atmospheric pressure, the fugacity of a gas is equal to its partial pressure (assuming ideal behaviour), therefore $VH_2O/VH_2=1$. The equilibrium constant was calculated using Maier–Kelly coefficients specific for each species and obtained here from the Supcrt92 software (Johnson et al. 1992). At T = 1,273 K (the most widely accepted temperature of the lava lake), K = 3.72 × 10⁻⁸, the logfO₂ is equivalent to Δ QFM = -0.92 (using a H₂/SO₂ ratio of 1.44, where QFM refers to the quartz–fayalite–magnetite buffer, and where Δ QFM = logfO₂ – logfO₂ of QFM). Using the obtained oxygen fugacity and prior measurements (Oppenheimer et al. 2009), we can recalculate the composition of the Erebus plume to include H₂ and the expected abundance of H₂S (Table 1, first column). Note that, in the table, H₂S is estimated based on the gas redox properties calculated using the "Dcompress" software from Burgisser et al. (2008) though it has not been detected at Erebus despite multiple attempts (Oppenheimer and Kyle 2008).

Table 1
Estimated composition of the Erebus plume in mole percent and molecular ratios for given species pairs

	Mean	Top of cycle	Bottom of cycle
	mol%		
CO ₂	44.00	34.86	47.25
H ₂ O	47.84	57.62	43.37
SO ₂	1.10	1.39	0.94
CO	3.30	2.61	3.54
HCI	0.46	0.56	0.42
HF	1.16	1.39	1.05
H ₂	1.58	1.39	1.89
OCS	0.01	0.01	0.01
H ₂ S	0.55	0.16	1.53
	mol/mol		
CO ₂ /CO	13.33	13.33	13.33
SO ₂ /H ₂ O	0.02	0.02	0.02
H ₂ /SO ₂	1.44	1.00	2.00
CO ₂ /SO ₂	40.00	25.00	50.00
SO ₂ /H ₂ S	2.01	8.72	0.62
log(fO ₂) at 1,000 °C	-11.90	-11.63	-12.14
Delta QFM	-0.92	-0.65	-1.16

The CO₂/CO and SO₂/H₂O molar ratios are obtained from Oppenheimer et al. (2009) as HCl/CO, HF/CO and OCS/CO ratios are used to estimate the plume composition in mole percent. The H₂/SO₂ and CO₂/SO₂ molar ratios are obtained from the multi-gas instrument measurements and the SO₂/H₂S molar ratio is calculated using the Dcompress software (Burgisser et al. 2008). The top of cycle

composition corresponds to the mixed plume composition of Oppenheimer et al. (2009), while the bottom of cycle composition corresponds to the conduit gas composition of Oppenheimer et al. (2009)

The oxidation state of the phonolite magma in the persistent lava lake of Erebus volcano has been estimated by several techniques. Kelly et al. (2008) used mineral chemistry to estimate an oxidation state of $\Delta QFM = -0.9$ using the QUILF program (Andersen et al. 1993) and a temperature of 1,000 °C. Oppenheimer and Kyle (2008) and Oppenheimer et al. (2011) used the CO₂/CO ratio obtained using FTIR spectroscopy (and the same temperature) to estimate the oxidation state at $\Delta QFM = -0.9$ to -0.88. Both of these estimates are essentially identical to our mean $\Delta QFM = -0.92$. It should be noted however that our new estimate of the oxidation state is not entirely independent as we used the SO₂/H₂O ratio previously measured by FTIR spectroscopy in our calculation.

The presence of H_2 in the volcanic plume suggests that H_2 is not burning at the interface between the lava lake and the atmosphere as has been observed, for instance, at Kīlauea's lava lake (Cruikshank et al. 1973). The correspondence of computed redox conditions for the lava lake also indicates that the H_2 abundance at the crater rim corresponds to the high-temperature equilibrium with the lava lake as hypothesized by Martin et al. (2009) and experimentally verified at Etna by Aiuppa et al. (2011). If any H_2 is oxidizing in the plume (e.g. to form HO_x radicals), it is only in minor amounts.

Periodicity and magma supply to the lake

From the time series, the H₂/SO₂ molar ratio varies mostly between 1 and 2 for all 6 days while the CO₂/SO₂ molar ratio varies mostly between 25 and 50. These upper and lower values can be attributed to two end-member compositions associated with a cyclic dynamic behaviour of the lake (Table 1). This quasi-periodic behaviour has been recognised in the velocity field of the lava lake surface motion, the heat loss of the lava lake, the SO₂/CO₂ and HCl/CO gas ratios (Oppenheimer et al. 2009), the SO₂ flux (Boichu et al. 2010) and lidar observation of the rising and falling lava lake surface (Jones et al. 2010, 2011). All these observations reveal cycles of between roughly 8 and 18 min. These have been interpreted as consequences of an episodic arrival of magma into the lava lake, which exsolves a water-rich end-member gas composition (Oppenheimer et al. 2009; Boichu et al. 2010).

The terms "top" and "bottom" of the cycle are adopted here to echo the previous literature (Oppenheimer et al. 2009); the "top of cycle" refers to high SO_2/CO_2 ratio, faster lake motion and higher lake level, and, as shown in Fig. 5, corresponds to high SO_2/H_2 . This composition is calculated using an H_2/SO_2 molar ratio of 1 and SO_2/H_2O molar ratio of 0.0242 from the "mixed plume" composition of Oppenheimer et al. (2009), which corresponds to measurements during vigorous convection of the lava lake, while the "bottom of cycle" composition is calculated using an H_2/SO_2 ratio of 2 and SO_2/H_2O gas ratios of 0.0218 from the "conduit gas" composition of Oppenheimer et al. (2009). The difference between the two end-member compositions is quite significant in terms of redox state, representing $\Delta QFM = -0.65$ at the "top of the cycle" and $\Delta QFM = -1.16$ at the "bottom of the cycle", assuming no change in temperature.

Considering that the "tops" of the cycles are marked by increases in lake level, surface motion and SO_2 flux, Oppenheimer et al. (2009) and Boichu et al. (2010) suggested that they are associated with the arrival of foaming magma batches in the lava lake (still exsolving water at near atmospheric pressure). We now observe that the tops of the cycle are

consistently associated with significantly more oxidized conditions ($\Delta QFM = -0.65$) than the "bottoms" of the cycles ($\Delta QFM = -1.16$). This differential pattern in oxidation state is also apparent in subtle cyclic variations in CO_2/CO ratio reported in Burgisser et al. (2012). If the top of the cycles is indeed associated with the influx of rising magma batches, then these batches appear to be releasing gas whose composition is a relic of chemical equilibrium acquired at some depth. This signature may be preserved as a result of rapid ascent of the magma batch (i.e. fast with respect to the kinetics of redox reactions such as [1] and [2]). The dichotomy we identify between the oxidized top and reduced bottom of the cycles therefore provides further empirical evidence for redox stratification in the Erebus plumbing system as discussed in Oppenheimer et al. (2011) and Burgisser et al. (2012) and as hypothesized from a more general standpoint by Burgisser and Scaillet (2007).

Conclusion

In situ measurements of the gas plume emitted from the lava lake of Erebus volcano by means of a multi-gas sensing instrument indicate that the hydrogen abundance in the magmatic gas phase is around 1.6 mol%. These measurements constrain the oxidation state of the lava lake to $\Delta QFM = -0.9$ log units, consistent with previous estimates; provide strong evidence that hydrogen burning is not prevalent at the surface of the lake; and that hydrogen is at least largely kinetically inert in the gas/aerosol plume rising in the crater. The hydrogen flux to the atmosphere from the summit of Erebus is estimated at 2.8 Mg day⁻¹. A strong ~10-min cyclicity in the proportions of H_2 and other species in the plume infers corresponding redox state variations and points to a pulsatory supply of magma to the lava lake. The more oxidized signature of the magma episodically entering the lake provides strong empirical evidence of a redox stratification in the shallow plumbing system, as has been hypothesized by previous numerical models.

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