First Multi-GAS based characterisation of the Boiling Lake volcanic gas (Dominica, Lesser Antilles)

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ABSTRACT

We used a Multi-component Gas Analyser System (Multi-GAS) to measure, for the very first time, the composition (H₂O, CO₂, H₂S, SO₂) of the volcanic gas plume issuing from the Boiling Lake, a vigorously degassing, hot (T ~ 80-90°C) volcanic lake in Dominica, West Indies. The Multi-GAS captured in-plume concentrations of H₂O, CO₂ and H₂S were well above those typical of ambient atmosphere, while no volcanic SO₂ was detected (<0.05 ppm). These were used to derive the Boiling Lake plume characteristic ratios of CO₂/H₂S (5.2±0.4) and H₂O/CO₂ (31.4±6). Assuming that other volcanic gas species (e.g., HCl, CO, H₂, N₂, etc.) are absent in the plume, we recalculated a (air-free) composition for the sourcing volcanic gases of ~ 96.3% H₂O, 3.1% CO₂ and 0.6% H₂S. This hydrous gas composition is within the range of published gas compositions in the Lesser Antilles region, and slightly more H₂O-rich than obtained for the fumaroles of the nearby Valley of Desolation (~94.4% H₂O, 4.7% CO₂ and 0.8% H₂S, CO₂/H₂S of ~5.7). We use our results, in tandem with the output of numerical simulations of gas scrubbing in the lake-water (performed via the EQ3/6 software), to derive new constraints on the degassing mechanisms at this poorly studied (but potentially hazardous) volcanic lake.

1. Introduction

The Lesser Antilles volcanic arc, in the West Indies, hosts a productive chain of active volcanoes [Wadge 1984] which devastation power has dramatically been brought to mind by the 1902 Mt. Pelée eruption in Martinique (30,000 casualties; Lacroix [1904]) and, more recently, by the still-ongoing Soufrière Hills (Montserrat) eruption started in 1995 [Druitt et al. 2002]. With most of its active volcanoes remaining dormant for centuries but still prone to erupt again in the future, and with a still-developing socio-economical structure with clear touristic vocation, the Lesser Antilles remain an area where a potentially high volcanic risk insists [Roobol and Smith 1989]. As dramatically evocated by the great controversies arisen during management of the 1975-76 seismo-volcanic crisis of La Soufrière in Guadeloupe (references in Komorowski et al. [2005]), a better understanding of past volcanic behaviour and present-day volcano setting is vital to correctly interpret any change in the status of historically active volcanoes, potentially heralding to an eruption.

Dominica has been, since the Miocene, the most productive (in terms of erupted volumes) volcanic island of the Lesser Antilles arc [Wadge 1984], and some of the most explosive eruptions (with depositional record) in the arc have occurred in this island [Sigurdsson 1972, Carey and Sigurdsson 1980, Sparks et al. 1980, Lindsay et al. 2003]. There is no report of magmatic eruptions in historical times in Dominica, however, and the island is today only partially covered by modern instrumental volcano monitoring networks. Whilst seismological information is largely available (Seismic Research Unit, University of the West Indies, Trinidad and Tobago; see Stasiuk et al. [2002], and Lindsay et al. [2003]), the geochemical monitoring network is far less developed, albeit hydrothermal manifestations are vast and widespread over the island [Joseph et al. 2011, and reference therein], and overall an expression of the island’s still active volcanic nature.

The Boiling Lake (BL; Figure 1), located in the SE-sector of the island (Figures 2 and 3), is the most impressive hydrothermal manifestation in Dominica. With its persistent vigorous steam discharges, its high temperature (80-90°C) and acid environment (pH, 4-6) [Fournier et al. 2009], the BL is one of the most ex-

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traordinary volcanic hot lakes in the world (the second largest after the Frying Pan Lake, New Zealand; Vandemeulebrouck et al. [2008]). The BL has also been the theatre of the most recent volcanic phenomenologies on Dominica, such as a phreatic explosions in 1880 (described by Nicholls [1880a,b]) and a lethal CO2-gas release event in 1901 [Elliot 1938, Bell 1946].

A more recent phreatic explosion is reported to have occurred in 1997 in the Boiling Lake/Valley of Desolation area (Figure 3). While there have been previous reports on the chemical and isotopic compositions of lake-waters [Pedroni et al. 1999, Joseph et al. 2011], no information is presently available on the composition of the gas being discharged at the lake’s surface, due to the inherent difficulties in direct gas sampling (gas discharges occur at the centre of a vigorously convecting hot lake).

Here, we report on the first characterisation of the BL gas manifestations, which we obtained by in-plume gas observation with a Multi-GAS [Aiuppa et al. 2005] during a field expedition in late February 2012. We demonstrate the suitability of the Multi-GAS as a technique to permanently monitor activity of the lake. We also use our results to derive new information on degassing features at this potentially hazardous lake.

2. The study area

2.1. The Lesser Antilles volcanic arc

The island of Dominica (Figure 2), located between Martinique, to the south, and Guadeloupe, to the north, belongs to the Lesser Antilles archipelago, which extends for ~850 Km from the Atlantic continental margin to the Greater Antilles.

The Lesser Antilles archipelago forms the currently active volcanic arc of a convergent margin, in which the north American Plate oceanic crust is being subducted underneath the Caribbean Plate, since the Oligocene, at a 2.3 cm/yr rate [Maury et al. 1990, Dixon et al. 1998, DeMets et al. 2000]. The Lesser Antilles islands describe a double arc, coalescent in the southern part and diverging into two branches in the northern sector, at the latitude of Dominica [Martin-Kaye 1969]. Volcanic activity has been concentrated on the eastern branch until the Miocene, when it shifted to the western side of the arc [Bouysse 1984, Bouysse et al. 1990] in response to a major deformation event in the subducting plate. Because of this, the Benioff zone underneath the Lesser Antilles archipelago actually appears irregular, dipping 50-60° to the north of Martinique, to then deepen in the southern part of the arc, to become vertical near Grenada Island [Wadge and Shepherd 1984, Feuillet et al. 2002]. The slab is supposed to be located at about 170km depth underneath Martinique and Dominica; in this depth range, clusters of deep seismic events have been detected [Ruiz et al. 2011].

Volcanic rocks in the Lesser Antilles are dominantly andesites and dacites of calc-alkaline affinity. A progression from low-K tholeiitic products, to medium and high-K calc-alkaline and alkaline rocks, has been shown to occur from northern (Montserrat to Saba) to central (from St. Lucia to Guadeloupe) and southern (Grenada) parts of the arc [Sigurdsson et al. 1973, Brown et al. 1977, Hawkesworth and Powell 1980, Smith et al. 1980, Westercamp 1988]. This trend is associated to enrichment in incompatible elements and increase in radiogenic strontium and lead [Davidson 1986, White and Dupré 1986, Pedroni et al. 1999], suggesting either (i) assimilation of terrigenous sediment by the rising magma plus fractional crystallization [Davidson 1985, 1987, Smith et al. 1996, Thirlwall et al. 1996] or (ii) mixing of partially melted subducted sediments and primitive mantle in the Benioff zone [White and Dupré 1986, Westercamp 1988].

2.2. Dominica

Dominica is almost completely made up of volcanic rocks, with the exception of some minor conglomerates and corals along its west coast. Outcropping
volcanites have been dated from Miocene to present [Demange et al. 1985, Monjaret 1985, Bellon 1988] (Figure 2). The cumulative erupted magma volume of ~40 km$^3$ [Wadge 1984] makes of Dominica the most productive volcanic island in the Lesser Antilles in the last 100 ka. Basaltic magmas were erupted in early volcanic activity stages (from Miocene to the Pliocene), while Pelean-type domes and pyroclastic deposits, ranging from andesitic to dacitic in composition, characterized volcanic activity in Pleistocene to Recent.

A key phase in the formation of present-day Dominica is thought to have occurred in the Pliocene (4-2 Ma), when the bulk of the island was built up by several coalescent volcanoes. This activity overall led to edification of a large strato-volcano (Cochrane-Mahaut centre), which eroded remains are now mainly preserved in the southern-central portion of the island [Bellon 1988]. The Cochrane-Mahaut centre consisted of a basal (less than 3.7 Ma but more than 2.83 Ma old) unit of basaltic pillow lavas and submarine volcanic breccias, overlain by subaerial andesitic lava flows and inter-bedded pyroclastic deposits in its younger sections. After a probable break in volcanism, at about 2Ma (when well developed soils were formed), volcanic activity resumed and intensified in the Pleistocene, concentrating in the south-central part of the island during the last 1 Ma. Nine major pyroclastic flow deposits, the largest of which being the Roseau Tuff and the Grand Bay Ignimbrites [Lindsay et al. 2003, 2005], have been related to the activity of volcanic centres in the southern part of Dominica [Sigurdsson 1972, Lindsay et al. 2003, 2005]. The ~30 ka old Roseau Tuff was produced by the largest explosive eruption in the last 200 ka of the Lesser Antilles [Carey and Sigurdsson 1980]. The eruption was sourced by either the Morne Trois Pitons/Micotrin lava domes [Sigurdsson 1972, Carey and Sigurdsson 1980, Sparks et al. 1980] or the Wotten Waven caldera [Demage et al. 1985]. The Roseau Tuff consists of a huge (~3 km$^3$) ignimbrite deposit, occasionally welded, filling a ~8 km long valley.
up to the city of Roseau [Sigurdsson 1972, Sparks et al. 1980]. The Grand Bay Ignimbrites outcrop in the south-eastern sector of Dominica and have recently [Lindsay et al. 2003, 2005] been related to a ~39 ka old large-scale collapse event of the Plat Pays volcano, which ultimately led to formation of Soufrière bay. The Grand Savanne Ignimbrite (22-44 ka; Sparks et al. [1980]) is a pyroclastic deposit originated by Plinian eruption(s) of Morne Diablotins, the only volcanic centre of the northern part of Dominica active during the Pleistocene.

While there is no historical report of magmatic eruptions in Dominica, seven volcanoes, active in the last 10 ka, are considered likely to erupt again. The high eruption potential of Dominica is also supported by widespread geothermal activity [Joseph et al. 2011] and shallow seismicity (a major earthquake swarm was observed in 1998-2000 on the southern sector of Dominica; Stasiuk et al. [2002], Lindsay et al. [2003]). Low-temperature (90-100°C) sulphur-rich fumaroles and hot springs are common in the Plat Pays volcanic complex area (Sulphur Springs, Galion fields and Champagne Beach) and in the Watten Wave caldera, in the SW-sector of the island [Joseph et al. 2011]. Hydrothermal fluid discharges are also abundant in the Valley of Desolation/Boiling Lake sector, where a number of hot springs, bubbling pools and fumaroles are hosted in a very small area (0.5 km²). Thermal waters in the Valley of Desolation area are generally acidic (pH=1 to 4), hot (temperatures from 74 °C to 98 °C) and with a prevalent Na-SO₄ composition. They are interpreted [Joseph et al. 2011] as the result of steam-heating of meteoric fluids by rising CO₂-H₂S rich hydrothermal vapours.

2.3. The Boiling lake

The Valley of Desolation/Boiling Lake region has been the theatre of two main phreatic explosions in 1880 [Nicholls 1880a,b] and, more recently, in 1997, and is a most likely potential source region for the next volcanic eruption in Dominica. Located at about 800m of altitude, in a hill above the Valley of Desolation area (SE-sector), the Boiling Lake (Figure 1) has a volume of ~ 104 m³ (50m wide and 15m deep; Fournier et al. [2009]), and is therefore the second largest boiling lake in the world, after the Frying Pan Lake (New Zealand) [Vandemeulebrouck et al. 2008]. The lake is hosted within a depression formed by a phreatic/phreato-magmatic explosion(s) [Joseph et al. 2011]. After the 1880 eruption [Sapper 1903, Robson and Willmore 1955, Brown 2002], the Boiling Lake has undergone prolonged phases of steady-state activity, in which the depression is filled by hot (80 to 90°C) and acidic (4-6) waters, and a vigorous gas upwelling region (object of this study) takes place in the lake’s centre (Figure 1), forcing continuous turbulence and convection. This “steady-state” lake is characterized by highly saline (TDS up to 14.000 ppm) Na-Cl(SO₄) waters, which are thought to be deeply derived brines rising from a deep-seated hydrothermal reservoir, and which are variably admixed with (and diluted by) meteoric water [Joseph et al. 2011]. Apart from direct precipitations, the lake is fed by a stream on its northern shore; whilst the only visible outflow is on the southern shore, where lake waters overflow into the Rivière Blanche river [Fournier et al. 2009].

Heat, transferred to the lake via hydrothermal steam ascending through a central feeding conduit, sustains water convection observed in the centre of the lake, and maintains its temperature constant at ~90°C [Fournier et al. 2009]. During their ascent, gas bubbles drag hot water with them and, compensating the hydrostatic pressure of the above water mass [Fournier et al. 2009], force the lake water to remain “suspended” 12-15m above the local water table [Bardou 2006, Fournier et al. 2009], at least in the steady-state conditions. Condensed steam is then thought to be recycled back through the same conduit.

Since 1876, occasional variations in lake activity (e.g., water level fluctuations and changes in the lake’s chemical features) have been reported [Ober 1880, James 1988, Fournier et al. 2009], including a small lethal (2 casualties) explosion in 1901, causing the expulsions of CO₂-rich gases into the atmosphere [Elliot 1938, Bell 1946]. Fournier et al. [2009] recently provided an extensive report on the most recent lake’s deviation from steady-state conditions. In this event, starting in December 2004, a significant drop in the water level was observed (until to nearly complete emptying of the lake), accompanied by a cessation of steam inflow into the lake and consequent temperature decrease (down to 20°C). Several cycles of emptying-refilling, one of this lasting only one day, were observed in January 2005; during these events, low temperature, neutral, low-sulphate and low-chlorine waters were observed in the lake, with recurrent stop-restarts of gas bubbling. This “crisis” eventually terminated in April 2005, when steady-state conditions in the lake (normal water level, high T - 90°C -, acid pH, and vigorous convection) were finally restored. No simultaneous change was reported in the neighboring hot springs and fumaroles of the Valley of Desolation.

Fournier et al. [2009] recently proposed a model in which the lake’s drying out events would be triggered by external perturbations such as landslides and/or regional seismicity. These would result into a
net decrease in gas flux into the lake, leading to fast downward flux of liquid water back into the conduit, and consequently (i) a significant reduction of the upward stress sustaining the water lake above the piezometric level, and (ii) drainage of the Boiling Lake down to the meteoric water table level. Afterwards, when the lake has entirely been emptied, bubbles would return to rise up through the conduit and, dragging liquid water upwards, would promote refilling of the lake.

3. Technique

The highly dynamic degassing activity of the BL prevents any attempt to directly sample the vapours being released at the centre of the lake (Figure 1). Because of this, the compositional features of this gas have remained undetermined until this study. Clearly, a technique based on measurement of the lake atmospheric gas plume, rather than the hardly accessible fumaroles, is required.

The Multi-GAS, a custom-made instrument combining commercial gas sensors, is an increasingly used technique for accurate in-plume observations of volcanic gas species [Aiuppa et al. 2005, Shinohara 2005]. This instrument has traditionally been used to measure plume compositions at open-vent volcanoes, during surveys [e.g., Aiuppa et al. 2006, 2012], permanent installations [Aiuppa et al. 2007, 2009, 2010] or airborne profiling [Shinohara et al. 2003]. Two recent applications [Roberts et al. 2012, Shinohara et al. 2013]...
have first been demonstrated the ability of the Multi-GAS to measurement of gas plumes issuing from a hot degassing volcanic lake (Yudamari crater lake, Aso volcano, Japan). This measurement approach is explored further here.

The portable Multi-GAS we used in this application allowed detection of the in-plume concentrations of $\text{H}_2\text{O}$, $\text{CO}_2$, $\text{SO}_2$ and $\text{H}_2\text{S}$, using the following instrumental configuration (same as in recent studies; e.g., Aiuppa et al. [2011]): a Licor LI-840 NDIR closed-path spectrometer for $\text{CO}_2$ and $\text{H}_2\text{O}$, and specific electrochemical sensors for $\text{SO}_2$ (3ST/F electrochemical sensor by City Technology Ltd.; 0-200 ppm range) and $\text{H}_2\text{S}$ (EZ3H electrochemical sensor by City Technology Ltd.; 0-200 ppm range). All the sensors, powered by an internal battery, were housed in a water-proof box, through which the plume gas was pumped at 1.2 l/min rate. The sensor readouts were sampled at 0.5 Hz by an onboard data-logger, and saved into an internal memory board. The acquired data were post-processed using the methodology described below (cf. Section 4).

Our measurements at the BL were taken on February 26, 2012, from a fixed position (Lat, N15.31867; Long, E61.2947; elevation 792m) on the north-northwestern outer rim of the depression hosting the lake (see Figures 1a and 3d). Weather conditions, which had remained unstable over the morning of February 26, significantly improved between mid-day and 3pm, when our observations were taken in relatively dry, stable (no precipitation, relative humidity of ~70%, $T_{\text{ambient}}$ of ~24°C) conditions. The SSE winds, dominating on that day, brought the gases issuing from the lake’s surface towards our observation site between about 1 and 1.50 pm (local time). We estimate a plume travel time, from emission at the lake’s surface to our observations site, of several seconds at most, a far too short timescale for chemical processing in the plume to significantly alter the original fumarole composition. Tests made at a number of closed-conduit volcanoes in fumarolic stage of activity (e.g., La Fossa crater, Vulcano Island; Aiuppa et al. [2005]) have demonstrated fair agreement between Multi-GAS derived compositions of near-vent fumarolic atmospheric plumes and compositions of the source fumaroles (from direct sampling), at least for brief transport distances (timescales of tenths of seconds).

In order to obtain consistent data to compare with our BL dataset, we also explored the fumarolic field of the Valley of Desolation (VoD). In this case, the technique was employed in a walking traverse mode [see Aiuppa et al. 2005], i.e., while traversing the fumarolic field with the MultiGAS transported in a backpack (Figure 3f). At each degassing vent, the traverse was stopped, and the instrument was left to acquire for a few minutes with its inlet being positioned at ~0.3m from the vent. Acquiring gas concentration data at 0.5 Hz, and with simultaneous GPS positioning (Figure 3a), the Multi-GAS had the required temporal/spatial resolution to map the chemical heterogeneity of a fumarolic field (e.g., Aiuppa et al. 2005). Unfortunately in this specific VoD survey, however, the plume was condensing (e.g., liquid $\text{H}_2\text{O}$ was stable) close to most fumaroles (condensation occurred as steam cooled upon dilution and transport in air). In addition, both the $\text{H}_2\text{S}$ and $\text{CO}_2$ sensors achieved signal saturation (at ~250ppm and 3500 ppm, respectively) in several part of the traverse. Only a subset of the acquired VoD dataset was therefore accessible to quantitative analysis.

4. Results

Figure 4 shows a 40 minute-long time-series of $\text{H}_2\text{O}$, $\text{CO}_2$, $\text{H}_2\text{S}$ and $\text{SO}_2$ concentrations in the plume of the BL. For both $\text{H}_2\text{O}$ and $\text{CO}_2$, we report in Figure 4 the air-corrected concentrations, which we calculated by subtracting from the raw measured concentrations the characteristic ambient air values of ~22,300 ppm and 385 ppm, respectively (obtained by averaging 10-minute measurements in the background site BG shown in Figure 3a).

The figure demonstrates coherent temporal variations for in-plume concentrations of $\text{H}_2\text{O}$, $\text{CO}_2$ and $\text{H}_2\text{S}$, an hint for their common derivation from lake degassing. Visual observations during the measurements clearly suggest that the large and erratic concentration variations captured by the Multi-GAS reflected changes in plume direction and altitude (due to changes in wind speed and direction above the lake) and, only to a lesser extent, the time-changing dynamics of degassing of the lake. The peak concentrations (3680, 101 and 25 ppm for respectively $\text{H}_2\text{O}$, $\text{CO}_2$ and $\text{H}_2\text{S}$, evident in the central part of our record (at measurement number 600), were obtained in a (few-minutes long) temporary phase of wind speed decrease, during which the gas was forced to accumulate at our observation site. $\text{SO}_2$, instead, was virtually absent in the plume, confirming that $\text{H}_2\text{S}$ is the only $S$ species emitted, and that no rapid oxidation process is occurring in the plume.

From the concentration data of Figure 4, the scatter plots of Figure 5 were obtained. These were then used to derive – by calculating the slopes of the best fit regression lines – the characteristic $\text{CO}_2/\text{H}_2\text{S}$ (5.2±0.4) and $\text{H}_2\text{O}/\text{CO}_2$ (31±6) volatile ratios in the BL atmospheric plume (all ratios quoted here are molar).

Figures 4 and 5 compare the compositional feature
of the BL gas plume with the compositions of the atmospheric plumes of the VoD fumaroles (Figure 3). These latter were characterised by remarkably high CO$_2$ (>3500 ppm) and H$_2$S (>250 ppm) concentrations, given the proximity (50 cm) of our measurements to the exhaling vents. Limiting our analysis to the couples of simultaneous measurements in which all sensors’ outputs were below saturation (Figure 5a), we obtain a large spread of CO$_2$/H$_2$S, from <2 to >50, in agreement with large compositional variability observed in previous work [Joseph et al. 2011]. Most of the VoD fumaroles, however, converge at high gas concentrations to a CO$_2$/H$_2$S of ~5.7, near our inferred composition for the BL. The VoD plumes were condensing close to most vents, precluding any H$_2$O retrieval. Only at a few sites we obtained a volcanic signal clearly distinguishable from the atmospheric background, these measurements being summarised in Figure 5c. From these, a representative H$_2$O/CO$_2$ ratio of ~20 is obtained for the VoD fumarolic field. The VoD fumarolic plumes also have very low but still detectable SO$_2$/SO$_2$/H$_2$S of ~0.00020).

5. Discussion

The BL gas plume, for which we provide here the first compositional report (at least to our knowledge), is manifestly H$_2$O-rich (H$_2$O/CO$_2$ of 31±6). With the objective of facilitating comparison between our observations and other volcanic gas datasets (Figure 6), we recalculated – from the plume characteristic ratios described above – an air-free composition for the BL gas. This operation, while mathematically correct, leads to a “real” volcanic gas composition only at condition that other volcanic gas species (not detected by the Multi-GAS: e.g., HCl, CO, H$_2$, N$_2$, etc.) are unimportant in the BL gas. This assumption, while not strictly true, is a good approximation, however, considering that H$_2$O, CO$_2$ and H$_2$S generally make up ~100% of the total gas phase in low-temperature hydrothermal gas manifestations [Chiodini and Marini 1998], including in Dominica [Joseph et al. 2011]. For instance, low concentrations of HCl (<24 mmol/mol), H$_2$ (<82 mmol/mol), N$_2$ (<116 mmol/mol) and CO (<0.008 mmol/mol) were reported by Joseph et al. [2011] for the VoD fumaroles, where CO$_2$ and H$_2$S are normally in the hundreds of mmol/mol range (all concentrations were reported by Joseph et al. [2011] on a dry-gas basis).

We finally obtained a recalculated composition for the BL gas of ~96.3% H$_2$O, 3.1% CO$_2$ and 0.6% H$_2$S. This composition can now be used to better constrain the mechanisms of gas feeding to the BL. In their physical modelling study, Fournier et al. [2009] interpreted the BL as a suspended meteoric water body, heated by condensation of rising steam bubbles supplied (with eventually a brine component; Joseph et al. [2011]) by a deep hydrothermal reservoir. In such a model, two po-
tential sources for our measured BL gas can be identified: (i) the hydrothermal steam feeding the lake or (ii) evaporation/degassing of lake-water itself. Contribution (i) would of course be zero if steam was to completely condense into the lake before reaching its surface. With no BL gas measurements available, Fournier et al. [2009] were not in the condition to distinguish between hypotheses (i), (ii), or their combination. They however left the possibility open that, given the fast ascent of gas bubbles through a relatively thin shallow (~15m) lake, part of the supplied gas can make the whole path through the lake, to burst at its surface. Our measurements here open the way to testing the hypothesis.

5.1. Models of gas-lake interaction and scrubbing
Marini and Gambardella [2005] first used the EQ3/6 software (7.2b version; Wolery and Daveler [1992]) to quantitatively model interaction of a volcanic gas phase with an aqueous solution, and the consequent removal (“scrubbing”) of water-soluble reactive gas species [Symonds et al. 2001]. Here, we follow the approach of Marini and Gambardella [2005] to model reaction between the BL lake-water and its feeding gas. Modelling is initialised and performed following the steps detailed below.

Step 1, Fixing feeding gas composition - The composition of the gas supplying the lake from below is unknown, but is required to initialise a simulation of gas scrubbing. Given the proximity between BL and VoD, we can make the guess that the composition of the feeding gas is approximated by a VoD-type gas. This latter is evaluated at ~94.4% H2O, 4.7% CO2, 0.8% H2S, and 0.1% HCl%, using our H2O/CO2 (20) and CO2/H2S (5.7) ratios (cf. Section 4), and assuming an average H2S/HCl ratio of 13 (the mean ratio calculated from the dataset of Joseph et al. [2011]). The temperature of the feeding gas, prior to its entrance in the lake, is taken at 200 °C (the equilibrium temperature estimated from application of the Na/K aqueous geothermometer to BL lake-water; Joseph et al. [2011]). Tests made changing this temperature range (in the 100-200 °C) demonstrated no substantial effect on the simulations.

Step 2, Fixing meteoric water composition - The composition of the meteoric water component entering the BL, and therein interacting with the feeding gas, is also required as initial solution for the simulations. No such data is reported in Fournier et al. [2009] and Joseph et al. [2011]. We therefore selected, as a proxy for the meteoric water entering the BL, the composition of a cold (T = 23°C) spring water from Soufrière volcano, in nearby Guadeloupe island (sample Beaugendre, from Brombach et al. [2000]).

Step 3, EQ3/6 runs - We then numerically simulated the gas-water interaction process by running a set of isothermal model calculations with EQ3/6. We performed 3 separate runs, having the same feeding gas and...
initial aqueous solution (steps 1-2 above), but differing for the mole amounts of feeding gas allowed to interact with 1 kg of initial solution. These were increased in the sequence 5.5, 6.4 and 7.2 moles (per kg of initial solution) from run 1 to 3. The temperature of the 3 runs were then determined by a simple enthalpy balance between the feeding gas (200 °C) and the initial solution (23 °C), and increased in the succession 84 °C (5.5 moles added), 92 °C (6.4 moles added) and 99 °C (7.2 moles added). The most “gas-rich” run we performed, therefore obtained by adding 7.2 moles of feeding gas to 1 kg of initial water, was then representative of the heating process of the initial solution, from its original temperature (23 °C) up to boiling temperature (99 °C). In each of the 3 runs, the code output consisted in the equilibrium composition of dissolved aqueous species in the model solution, including the partial pressures of individual gas species.

Step 4, degassing. At the end of each of the 3 runs, the total pressure of dissolved gases (calculated by summing together partial pressures of individual gas species) exceeded 1 bar. The obtained model solutions were then degassed using a single-step degassing model [see Marini and Gambardella 2005], until 1 bar total gas pressure was obtained. The so-derived model gas phase compositions (M1 to M3), obtained via degassing of model solutions, are shown, for the different runs, in the triangular diagram of Figure 6.

The model gases M1 to M3 (Figure 6) are ultimately representative of the gas that would form by degassing of (gas separation from) the BL at 84 °C, 92 °C and 99 °C. Model gas compositions M1-M3 are clearly distinct, for manifestly being depleted in sulphur (CO2/H2S ratios of 11-17), from the feeding gas composition (VoD gas; CO2/H2S = 5.7). This mismatch demonstrates the “chemical filtering” effect potentially played by gas-water interaction, in which more water-soluble H2S is preferentially (relative to CO2) scrubbed out of the feeding gas, and oxidised to dissolved sulphate (model solutions have dissolved sulphate contents of 3500-4950 mg/l, comparable to those reported for the BL by Joseph et al. [2011]). This gas scrubbing process also acts to remove HCl from the feeding gas: M1-M3 gas compositions are virtually identical to those reported for the BL by Joseph et al. [2011]), but differing for the mole amounts of feeding gas allowed to interact with 1 kg of initial solution. These were increased in the sequence 5.5, 6.4 and 7.2 moles (per kg of initial solution) from run 1 to 3. The temperature of the 3 runs were then determined by a simple enthalpy balance between the feeding gas (200 °C) and the initial solution (23 °C), and increased in the succession 84 °C (5.5 moles added), 92 °C (6.4 moles added) and 99 °C (7.2 moles added). The most “gas-rich” run we performed, therefore obtained by adding 7.2 moles of feeding gas to 1 kg of initial water, was then representative of the heating process of the initial solution, from its original temperature (23 °C) up to boiling temperature (99 °C). In each of the 3 runs, the code output consisted in the equilibrium composition of dissolved aqueous species in the model solution, including the partial pressures of individual gas species.

5.2. The BL gas in a regional context
If, as proposed above, the BL gas is only marginally affected by scrubbing in the lake, comparison is then possible with available volcanic gas datasets in the region. Figure 6 puts our Dominica measurements in the context of published gas analyses from active volcanoes in the Lesser Antilles.

Research on volcanic gas compositional signature in the Lesser Antilles arc has received significant impetus after the 1995-present eruption of Soufrière Hills (Montserrat); still, the available information remains fragmentary and incomplete – at least relative to other better characterised arc regions [Hilton et al. 2002, Mather et al. 2006, Fischer 2008]. The high-temperature (>700 °C) gas emissions issuing from the Soufrière Hills dome should in principle represent a source of invaluable information to constrain the truly magmatic gas composition in the region. However, given the inaccessibility of the vent area, and our current technical limitations in remotely sensing the major (H2O and CO2) gas species, it is only the SO2 volatile budget to remain well characterised [see Christopher et al. 2010, and references therein]; the C/S gas signature (and consequently the CO2 fluxes) have only recently been established [Edmonds et al. 2010], instead, and the H2O flux remains unconstrained. There has been, however, one fortunate report [Hammouya et al. 1998] of gas collection on- top Soufrière Hills dome on February 1996 (just before pyroclastic flows started); the collected gas samples, while affected by air dilution, allowed to set the recalculated (on air-free basis) magmatic gas composition at ~91% H2O, ~3% CO2, 2.7% HCl and 0.6% total sulphur (mainly present as SO2). This Hammouya’s et ali [1998] H2O-CO2-S magmatic gas composition for Soufrière Hills (e.g., with HCl not being considered) is close our above-derived composition for the Boiling Lake (Figure 6). Soufrière Hills and BL gases show, for instance, nearly identical CO2/S ratios of 5.1 and 5.2, respectively.

The ongoing (1992-present) degassing crisis of La Soufrière dome (references in Bernard et al. [2006])
makes of nearby Guadeloupe Island the second most significant volcanic gas spot in the region. Here again, however, the number of published gas reports is relatively small [e.g., Brombach et al. 2000, Bernard et al. 2006, Ruzié et al. 2012]. When this previous dataset is combined with results of our recent Multi-GAS observations in March 2012 [Allard et al. 2013] the Soufrière of Guadeloupe volcanic gases display an almost linear compositional array, from the S-poor (CO$_2$/S of 11-474) compositions reported for the early 1997 activity by Brombach et al. [2000] (samples collected just prior degassing activity increased in vigour in December 1997) to the S-rich (CO$_2$/S of ~2.5) we observed on March 2012 [Allard et al. 2013], which are representative of the ongoing degassing crisis. Notably, our BL gas sits in the central part of the Soufrière of Guadeloupe array.

The overall good match between the BL gas composition and more “truly magmatic” fluids in the region (e.g., Soufrière Hills and La Soufrière dome) once more supports the conclusion of a minor role played by secondary processes (e.g., gas scrubbing) in the lake.

6. Conclusive remarks

We have presented here the first chemical characterization of volcanic gas release at the Boiling Lake, one of the most prominent hydrothermal-volcanic manifestations in Dominica, and in the entire Lesser Antilles arc. Our results support a clearly volcanic derivation for the BL gas, and a marginal role played by interaction with lake waters (gas scrubbing). Our observations also provide additional confidence [Roberts et al. 2012, Shinohara et al. 2013] on the ability of the Multi-GAS to operate in the harsh conditions (e.g., very high relative humidity) often encountered in the plume(s) issuing from active volcanic lakes. At least in the conditions met during our survey, the BL plume was “dry” enough for volcanic H$_2$O sensing (which is hampered by liquid water if present in substantial amounts), and “dense” enough for a significant volcanic signal to be resolved from the background. We conclude that this technique could valuably assist any future plan for monitoring of the BL activity. Given the relatively frequent record of harmful volcanic events (phreatic explosions, and sudden toxic gas release events) in the recent past, and the increasing visitors inflow in Dominica, implementation of the monitoring network (including gas chemistry) should be a priority of local authorities and hazard managers in the region.

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