The gas membrane sensor (GMS) method: a new analytical approach for real-time gas concentration measurements in volcanic lakes

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Abstract: The primary goal of this study was to apply an innovative analytical method based on a gas membrane sensor (GMS) to quantify the concentrations of CO_2 and CH_4 gas in water columns from volcanic lakes. For this purpose, dissolved CO_2 and CH_4 from the Monticchio Grande and Piccolo (Mt Vulture, Italy) and Pavin (Massif Central, France) lakes, characterized by depths of 35, 38 and 92 m, respectively, were measured using two different approaches: GMS and SH (single hose), the latter being an independent conventional method.

The CO_2 and CH_4 concentrations recorded via the GMS and SH techniques were within the analytical errors of the two methods. These test measurements demonstrated that the GMS method can be successfully used to produce highly accurate *in situ* measurements of dissolved gas composition along the vertical profiles of lakes.

Volcanic lakes are commonly found in both dormant and recently active volcanic systems (Simkin & Sieber 1994; Delmelle & Bernard 2000). According to the classification proposed by Pasternack & Varekamp (1997), chemical and physical features of the water allow us to clearly distinguish two main volcanic lake categories: (1) highactivity lakes, consisting of acidic-to-hyperacidic lakes strongly affected by inputs of volcanic fluids; and (2) low-activity lakes, characterized by relatively low salinity and almost neutral pH, being located in systems with a low-to-moderate degassing rate of deep-originated fluids. Lakes pertaining to the latter group can be meromictic and show permanent dissolved gas reservoirs (Tassi & Rouwet 2014 and references therein). In this environment, the possible occurrence of vertical water mixing (rollover events) related to exo- or endogenic processes, e.g. landslides, earthquakes or increasing volcanic activity, may cause the sudden outburst of huge amounts of gases (limnic eruptions; Sabroux et al. 1987; Kusakabe 1996; Halbwachs et al. 2004; Kusakabe et al. 2008). These dramatic events represent a serious hazard, as demonstrated by the two disasters that occurred at Monoun and Nyos lakes (Cameroon) in 1984 and 1986, respectively (Kling et al. 1987; Sigurdsson et al. 1987). The origin of non-atmospheric gases found in volcanic lakes, basically consisting of CO₂ and CH₄, is mainly related to: (1) fluid contribution from underlying volcanic systems; and (2) microbial activity (Tassi & Rouwet 2014). The physicochemical evolution in time of dissolved gas reservoirs depends on both biogenic and geogenic inputs (Cabassi et al. 2014). The artificial degassing of bottom waters, a method successfully applied at Nyos and Monoun lakes (e.g. Kling et al. 2005), is considered particularly efficient to mitigate the risk of limnic eruption (Rouwet et al. 2014). However, this type of intervention may imply severe environmental concerns due to contamination of surface lake layers with reducing, toxic deep waters (Pasche et al. 2010), suggesting that the artificial degassing approach has to be evaluated with caution.

Scientific investigations based on direct physical-chemical measurements along vertical lake profiles are thus of fundamental importance to characterize and monitor the hazardous gas reservoirs that occur in meromictic volcanic lakes.

Different methods have been used for the analysis of dissolved gases in deep lake waters

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(Tassi & Rouwet 2014). Few of them allow data to be obtained in situ (Evans et al. 1993; Yoshida et al. 2010; Ohba et al. 2013). Most techniques also require laboratory analysis on samples collected using bottles with different geometry and equipped with various tapping systems (e.g. Niskin, Van Dorn, Ruttner samplers) or devices (single hose; Caliro et al. 2008: Tassi et al. 2009). Furthermore, the concentration of dissolved CO₂ has often been determined indirectly, based on pH (Kusakabe et al. 2000) and alkalinity. To overcome the disadvantages related to different problems, such as tedious laboratory analysis, sampling difficulties and instrumental efficiency, an improved gas sensor device, based on a membrane technique and suitable for measurements and sampling of dissolved gases, was developed. This device is easy to handle. It operates via a wire line and is free of any complex mechanical components.

This paper provides a description of the gas membrane sensor (GMS) and its application under conditions typical for meromictic lakes hosted in volcanic systems. The Monticchio Piccolo and Monticchio Grande (southern Italy) and Pavin (France) lakes were investigated to test the reliability of this new technical approach. Concentrations of dissolved CO_2 and CH_4 in different water layers measured with the GMS were compared with those obtained using the single hose (SH) method.

Theoretical background

The key component of the GMS system is a phase separating membrane often used in applied sciences (Burger *et al.* 1996), medicine and industry (Kesson 1984, 1985), in laboratories or otherwise protected environments for gas enrichment and separation purposes. Gas permeation through a membrane generally takes place in several steps according to the solution–diffusion model of Graham (1866). The penetrant condenses and dissolves at the surface of the membrane; then it diffuses in liquid form through the membrane under the influence of a concentration gradient (chemical potential) and evaporates to the gaseous state at the opposite surface (Klopffer & Flaconnèche 2001).

The relation between permeability, diffusivity and solubility is, as follows:

$$P_o = D_s \times S \tag{1}$$

where D_s is the diffusivity coefficient (cm² s⁻¹) and *S* the solubility coefficient (cm³_(STP) cm⁻³ cm Hg⁻¹). For the dissolution of small gas amounts at total pressure value near atmospheric, the simple 'Law of Henry' is valid and is linearly proportional to the partial gas pressure (Scholes *et al.* 2008). Hence, the

gas concentration inside the membrane is directly correlated to the dissolved gas concentration in the outside water. This correlation is expressed by the permeability coefficient (P_o), a measure of the flux through the membrane, expressed in units of 'barrer' (Alter 1962). The latter parameter depends on (equation 2):

- (1) the permeating gas flux through the membrane $(in \ cm_{(STP)}^3 \ s^{-1});$
- (2) the membrane surface area (in cm^2);
- (3) wall thickness (in cm);
- (4) partial pressure gradient (in cm Hg).

$$\frac{\mathrm{cm}_{(\mathrm{STP})}^{3} \times \mathrm{cm}}{\mathrm{cm}^{2} \times \mathrm{s} \times \mathrm{cm} \,\mathrm{Hg}} 10^{-10} = \mathrm{barrer} \quad (2)$$

The permeation rate through a polydimethylsiloxane (PDMS) membrane depends on temperature. As it is a thermally activated process (Michaels & Bixler 1961), the temperature dependence can be described by the Arrhenius relation, where E_a is the apparent activation energy for the permeation process (J mol⁻¹) consisting of the energies of diffusion and dissolution, as follows:

$$P_o = P_o(0) \exp\left(\frac{-E_a}{RT}\right) \tag{3}$$

where *R* is the gas constant $(J \text{ mol}^{-1} \text{ K}^{-1})$ and *T* the absolute temperature (K).

Terms of $-E_a/R$ for CO₂ and CH₄ with PDMS membranes are -118 K and 721 K, respectively (Ajhar 2011). These data reflect a negative temperature correlation of the CO₂ permeation coefficient, in contrast to the positive CH₄ permeation dependency.

Data on gas permeation rates for PDMS membranes (e.g. Robb 1968; Allen *et al.* 1977; Egli *et al.* 1984; Merkel *et al.* 2001; Praveen *et al.* 2006) are available in the literature, although real membrane material/gas systems generally show deviations from the theory due to different morphologies, polymer flexibilizers or previous history of the membrane (Staude 1992). Therefore, for the present study the permeation rates of CO₂ and CH₄ through the membrane tube used were determined experimentally in the laboratory by applying a thermostatic pressure device (Zimmer *et al.* 2011). At 7°C, the measured CO₂ and CH₄ coefficients were 945 and 153 barrer, respectively.

The partial pressure of dissolved gases in the lake waters was calculated with the transformed equation (2). The conversion of the partial gas pressure into the corresponding dissolved gas concentration at the measured temperature of the lake water (T in °C) was carried out according to the

Henry's law constants (H_i) of Sander (1999), whose dependencies on temperature are as follows:

$$H_{\rm CO_2} = 0.034 \\ \times \exp\left(2400 \times \left(\frac{1}{273.15 + T} - \frac{1}{298.15}\right)\right) \, [\rm{moll}^{-1} \, \rm{atm}^{-1} \,]$$
(4)

$$H_{\rm CH_4} = 0.0014$$

$$\times \exp\left(1700 \times \left(\frac{1}{273.15 + T} - \frac{1}{298.15}\right)\right) \text{ [moll^{-1} atm^{-1}]}$$
(5)

General features of the test sites

GMS and SH measurements were carried out at the Monticchio Piccolo and Monticchio Grande lakes (MP and MG, respectively) in Italy and at Lac Pavin (LP) in France. These volcanic lakes are characterized by permanent water stratifications, with the occurrence of an anaerobic hypolimnion overlaid by an aerobic epilimnion (Lopes *et al.* 2011; Cabassi *et al.* 2013).

MG and MP are located SW of Mt Vulture in two maars formed 140 ka ago during the last eruptive activity of Mt Vulture (Caracausi et al. 2009 and reference therein). MP has a maximum depth of 38 m and a surface area of 0.16 km², whilst MG reaches a maximum depth of 35 m with a surface area of 0.41 km². Dissolved CH₄ in both lakes has a typical biogenic isotope signature ($\delta^{13}C-CH_4 \leq -50\%$) V-PDP), whereas according to the δ^{13} C-CO₂ values (MG -4.3% and MP -1.6% V-PDP) CO2 was interpreted as a mixture of fluids from mantle and thermometamorphic reactions involving limestone (Cabassi et al. 2013). In both lakes, water rollover events have occurred several times during the last 200 years (Caracausi et al. 2009; Cabassi et al. 2013).

LP, located at an altitude of 1197 m above sea level., is a maar lake formed by a phreatomagmatic eruption about 6 ka ago (Camus *et al.* 1993) and represents the youngest volcanic activity in the Massif Central. LP has a maximum depth of 92 m and a surface area of 0.44 km². Dissolved CO₂ in the hypolimnion has a dominant magmatic origin, whereas the source for CH₄ is likely biogenic (Aeschbach-Hertig *et al.* 1999), although the lake was classified as low-productive (Restituito 1987).

Material and methods

Description of the GMS system

The GMS system basically consists of: (1) gas analytical devices at the surface; and (2) a gas collector that is lowered to the desired water depth (Fig. 1).

The gas collector is composed of a phase-separating membrane tube, a pressure and temperature transmitter (Keller, Typ PA-36XW), a water pump (Barwig, Typ 04, 100 ml min⁻¹) placed in a protective plastic housing, and a custom-made cable.

PDMS was chosen for the membrane tube due to the relatively high gas permeability of this material. Commercial PDMS membranes available for gas separation purposes have a thickness at the micrometre scale and are mostly stabilized by a supportive porous material. For the present study, a 0.8 mm thick silicon tube with a surface area of 270 cm² was used. This type of membrane was already tested and applied as a component for CO₂ analysis in deep boreholes (Zimmer *et al.* 2011). The cable embeds two stainless steel capillaries (inner ϕ of 2 mm), two strain reliefs, as well as three double core wires for the power supply of the water pump and the signal transmission of the pressure and temperature sensor.

A carrier gas (Ar), provided from a 2 l, 100 bar pressure vessel at the surface, was introduced through the first inlet capillary into the membrane tube. The Ar flux was set at 50 ml min⁻¹ at 1.2 bar using a gas flow controller (Analyt-MTC, 35829 Mass Controller) and pressure regulator (Air Liquid). As the outflow capillary is open to the atmosphere, the Ar pressure inside the membrane tube is constant and close to that of the atmosphere.

The gas pressure on the outer membrane face depends on the water depth at the measurement site. In order to resist the high-pressure gradient, the membrane tube was filled with glass spheres of 1-2 mm in diameter. The glass spheres prevented the membrane tube from collapsing, maintaining its form and allowing the gas to pass through. Adopting this solution, the membrane tube can resist pressures higher than 20 MPa, corresponding to a measurement site up to a depth of 2 km (Zimmer & Erzinger 2009). To prevent clogging, two filters were placed between the membrane tube and the stainless steel capillaries. To avoid depletion of the gas on the membrane surface, the pump continuously circulated the lake water through the GMS housing. Via the outflow capillary, Ar and gases permeating from the lake water through the membrane tube were carried back to the surface. At the surface, the collected gases were analysed dynamically in real time using an LI 820 sensor for CO₂ (LI-COR Inc.) with a measurement range between 0 and 2000 ppmv and a photoacoustic sensor (Innova 1412) for CH₄ (Air Tech Instruments) with a measurement range between 0.4 and 40 000 ppmv.

A laptop computer controlled the sensors and stored the signals. Two 12 V 55 Ah accumulators provided electric energy for eight hours of continuous measurements.

The standard deviation of the method for dissolved CH_4 and CO_2 concentration $<1 \text{ mmol } l^{-1}$ is 30%. For higher concentrations, the standard



Fig. 1. Schematic representation of the GMS with gas analytical devices at the surface (**b**) consisting of a box with the computer, pressurized argon, gas sensors and batteries. In addition, a gas collector element (**a**) comprising a tube membrane, a P, T transmitter, a water circulation pump and filter elements was applied under water.

deviation is 10%. The higher standard deviation at lower gas concentration is a result of a larger relative impact of stochastic noise effects from several variables (e.g. electronic and thermic noise) while approaching the detection limit of the GMS method.

GMS measurements

Gas measurements in the water column were performed in the two Monticchio lakes (MP at 40° 55' 56.26" N, 15° 36' 48.62" E and MG at 40°

55' 55.85" N, 15° 36' 12.47" E) Italy, on 15 and 16 October 2013, respectively, while those at Lac Pavin (45° 29' 44.84" N, 2° 53' 17.21" E) France, were carried out on 27 May 2014.

The sensor was manually lowered to the desired depth by the custom-made cable as the strain reliefs inside the cable take the complete weight. The sensor depth was controlled by a scale printed directly on the cable. The manual cable winding and the handling of the sensor were relatively easy due to the flexibility of the cable and the light total weight (about 12 kg). The computer, analytical sensors, batteries and Ar cylinder were placed inside a single waterproof box to protect them from splashing water in the boat and for easier handling on the shore.

The GMS measurements were carried out along the vertical profiles, from 5 m below the lake surface down to the lake maximum depth (MP = 38 m, MG = 35 m, LP = 92 m) at intervals of 5 m. At each measurement site, the GMS remained in position for 15 minutes, i.e. until the measured gas concentration in the Ar carrier gas was stable.

Temperature and SH measurements

Water temperature (T), along the vertical profiles of the MP, MG and LP lakes, was measured using a Hydrolab probe (Idroprobe) equipped with a data logger for data storage. The nominal precisions are, as follows:

depth, ± 0.05 m; temperature, $\pm 0.03^{\circ}$ C.

The sampling of dissolved gases using the SH method was carried out simultaneously to the GMS measurements. The SH device consisted of 10 m long Rilsan tubes (inner $\phi = 6 \text{ mm}$) connected by steel joints. Once the tube end was lowered to the chosen depth, water was pumped up to the surface by means of a 150 ml glass syringe, connected to a three-way valve, and transferred into pre-evacuated 250 ml glass vials equipped with a Teflon stopcock after the displacement of a water volume double that of the inner volume of the tube (Tassi et al. 2009). The sampling flasks were filled with water up to three-quarters of the inner volume. The free gas phase in the flask headspace was analysed by gas chromatography using Shimadzu 14a and Shimadzu 15a gas chromatographs equipped with TC- and FI-detectors, respectively. The chemical composition of dissolved gases was calculated using: (1) the gas composition; (2) the pressure; (3) the volume of the flask headspace; and (4) the solubility coefficient in water of each gas compound (Tassi et al. 2009). The analytical error for the dissolved gas analyses was $\leq 5\%$.

Results and discussion

Profiles of temperature

Water temperature vertical profiles of the investigated lakes are reported in Figure 2a-c.

MP and MG were sampled at the end of the summer season, explaining the relatively high water temperatures measured at the surface (up to 18.7°C and 18.2°C, respectively). LP was sampled in spring, thus the surface water was relatively cold (10.7°C). At MP (Fig. 2a), temperature decreased with increasing depth showing a clear thermocline at 7-13 m depth. The lowest temperature was reached at -15 m (7.1°C) and then progressively increased up to 9°C at the lake bottom. At MG (Fig. 2b), the thermocline was at 5-9 m depth. whereas the minimum temperature $(6.6^{\circ}C)$ was measured at the bottom. At LP, after the thermocline at 5-8 m depth, the temperature slightly decreased down to the minimum value $(3.7^{\circ}C)$ at 25-30 m depth, and then increased up to 5°C at the lake bottom.

Profiles of CO₂ and CH₄ concentrations

The vertical distribution of the CO_2 and CH_4 concentrations at MP and MG lakes, as well as that of



Fig. 2. Water temperature–depth profiles at (a) Monticchio Piccolo, MP, (b) Monticchio Grande, MG and (c) Pavin, LP, lakes.

Depth (m)	T (°℃)	$\begin{array}{c} \text{CO}_2 \\ (\text{mmol } l^{-1}) \\ \text{SH} \end{array}$	$\begin{array}{c} \text{CO}_2 \\ (\text{mmol } l^{-1}) \\ \text{GMS} \end{array}$	P _{CO2} (mbar) GMS	CH ₄ (mmol l ⁻¹) SH	CH ₄ (mmol l ⁻¹) GMS	P _{CH4} (mbar) GMS
Monticc	hio Piccolo						
0	18.7						
5	18.4	0.27	0.27	6.6	0.02		
10	10.2	0.93	0.98	19	0.05	0.04	21
15	7.1	6.4	6.1	108	0.66	0.69	340
20	7.3	9.2	9.4	166	1.05	1.09	544
25	7.7	10.1	10.1	181	1.17	1.19	597
30	8.2	11.4	11.1	202	1.57	1.54	784
35	8.7	14.5	14.2	262	2.21	2.10	1078
38	9.0	16.0	15.0	280	2.51	2.30	1187
Monticc	hio Grande						
0	18.2						
5	17.3	0.36	0.23	5.5	0.01	0.01	6.2
10	8.3	0.71	0.55	10	0.11	0.13	65
15	7.0	1.37	1.20	21	0.17	0.15	75
20	6.8	2.52	2.12	37	0.24	0.26	128
25	6.7	3.45	3.74	65	0.28	0.26	127
30	6.7	3.74	3.97	69	0.37	0.35	171
35	6.6	3.64	3.46	60	0.36	0.39	194

Table 1. Temperature and CO_2 and CH_4 concentrations and partial pressures measured at MP and MG using the GMS and SH methods

 CO_2 at LP measured using both the GMS and the SH methods, is reported in Tables 1 and 2. No CH_4 measurements are available for LP due to a sensor failure. The concentrations of CO_2 and CH_4 at MP

increased almost regularly with depth (Fig. 3a, b), reaching their maximum concentrations at the lake bottom (up to 15 and 2.3 mmol l^{-1} respectively; GMS method). At MG (Fig. 4a, b), the dissolved

Table 2. Temperature and CO_2 concentration and partial pressure measured at LP using the GMS and SH methods

Depth (m)	T (°℃)	$\begin{array}{c} \text{CO}_2 \\ (\text{mmol } l^{-1}) \\ \text{SH} \end{array}$	$\begin{array}{c} \text{CO}_2 \\ (\text{mmol } l^{-1}) \\ \text{GMS} \end{array}$	P _{CO2} (mbar) GMS
Lac Pavin				
0	10.7	< 0.01		
5	10.0		< 0.01	0.8
10	4.5	0.01	0.01	0.9
15	4.0		0.02	2.6
20	3.8	0.02	0.02	3.4
25	3.7		0.03	4.0
30	3.7	0.02	0.03	4.9
35	3.8		0.39	6.2
40	3.8	0.28	0.39	6.2
45	4.0		0.69	11
50	4.0	0.57	0.81	13
55	4.3		1.18	19
60	4.4	2.35	1.98	32
65	4.5		4.50	73
70	4.7	7.10	7.35	120
75	4.9		8.21	135
80	4.9	8.52	9.43	155
85	5.0		9.95	164
90	5.0	9.58	10.2	168



Fig. 3. (a) CO₂ and (b) CH₄ concentrations measured with GMS (red) and SH (blue) at MP. Error bars indicate the uncertainty of the GMS (<1 mmol 1^{-1} 30% and >1 mmol 1^{-1} 10%) and SH (5%) results. Differences between corresponding data pairs and the relative bias (*B*) of the GMS against the SH method are shown below. $B = 100\% \times \sum (Y_i/X_i - 1)n^{-1}$, where *Y* is the result of the GMS system, *X* is that of the SH method and *n* the number of measurements.



Fig. 4. (a) CO₂ and (b) CH₄ concentrations measured with GMS (red) and SH (blue) at MG. Error bars indicate the uncertainty of the GMS ($<1 \text{ mmol } 1^{-1} 30\%$, and $>1 \text{ mmol } 1^{-1} 10\%$) and SH (5%) results. Differences between corresponding data pairs and the relative bias (*B*) of the GMS against the SH method are shown below.



Fig. 5. CO₂ concentrations measured with GMS (red) and SH (blue) at LP. Error bars indicate the uncertainty of the GMS ($<1 \text{ mmol } 1^{-1} 30\%$, and $>1 \text{ mmol } 1^{-1} 10\%$) and SH (5%) results. Differences between corresponding data pairs and the relative bias (*B*) of the GMS against the SH method are shown below.

gas profiles were less regular, being characterized by a maximum for both gases at 30 m depth (CO₂ and CH₄, 3.97 and 0.37 mmol 1^{-1} , respectively; GMS method). The dissolved gas composition of MG, as well as the CO₂ profile of MP, is consistent with that measured using the SH method in October 2010 (Cabassi et al. 2013). The maximum CO_2 concentration in LP was 10.2 mmol l⁻¹ (GMS method) at the lake bottom (Fig. 5). Literature data for dissolved CO2 in LP vary considerably as they range from 1.1 (Lehours et al. 2005) to 16 mmol l^{-1} (Aeschbach-Hertig *et al.*) 1999). These data, which were consistent with those measured in the present study, seem to indicate that the stratification of LP was affected by significant perturbations, although no degassing of the lake was reported since 1999.

Statistical comparison between the GMS and SH methods

The relative biases (*B*) of the GMS v. the SH data range from -1% for CO₂ in MP to 12% for CO₂ in LP and are within the analytical error of the method (Figs 3–5). The relative root mean square deviation (RMSD) of the GMS measurements based on the possible bias and the uncertainty of the SH results is $0.37 \text{ mmol } l^{-1}$ for CO₂ and 0.07 mmol l^{-1} for CH₄.

Conclusions

this study an innovative methodological In approach, the GMS technique, was used to measure the concentrations of selected gas species (CO₂ and CH₄) dissolved in three volcanic lakes from Italy and France that are characterized by permanent or semi-permanent stratification. The compositional data obtained for the Monticchio Piccolo, Monticchio Grande and Pavin lakes by GMS were in good agreement with those measured using the SH method, indicating that the accuracy of the GMS technique is at least comparable with that of one of the most common sampling and analytical methods for the measurements of dissolved gases in a lake water column. The statistical tests have shown that the two series of data cannot be distinguished, since the corresponding data pairs overlap within the analytical uncertainties of the two methods. However, at lower dissolved gas concentrations, the consistency between the GMS and SH data decreased, possibly due to a higher standard deviation of the GMS method. To improve the sensitivity of the GMS method, the thickness of the membrane can be reduced and its surface area can be increased. These structural modifications should be able to increase both the permeation rate and the 'lake gas concentration' in the carrier gas, reducing the analytical uncertainty at low gas concentrations.

It is worth noting that the GMS method is able to deliver data on the dissolved gas concentration at any depth in a volcanic lake. Similar information is of fundamental importance for monitoring the evolution of a deep gas reservoir, which potentially represents the risk of limnic eruptions. Furthermore, by applying additional analytical devices at the surface (e.g. mass spectrometer, gas chromatograph, radon detector), the number of detectable dissolved components in the lake water can be extended to all dissolved gases. For example, GMS measurements combined with a quadrupole mass spectrometer have already provided useful data for dissolved H₂, N₂, O₂, CH₄ and CO₂ in drilling fluids of deep boreholes (Zimmer et al. 2011). Real-time production of compositional data is suitable for mapping large lake areas in a relatively short time and allows for the construction of very detailed (at depth intervals in the order of fraction of metres) vertical profiles of dissolved gas concentrations. A high number of compositional measurements at different depths and locations may be used to construct a 3D map of gas distribution, possibly helping to locate inflows of volcanic gas discharging from deep sources.

A GMS sensor could also be installed at one specific location at a certain lake depth. Maintenance of the system (replenishment of the Ar carrier gas, recalibration of the gas analytical devices) can be performed without pulling the sensor back to the lake surface.

With this method, it is also possible to obtain and collect gases from deep water layers for further geochemical and isotope studies in the laboratory. All in all, GMS can be considered a reliable and effective alternative method to traditional sampling techniques for gas concentrations in volcanic lakes.

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