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https://doi.org/10.1038/s43247-024-01210-3

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# Use of cold waters geochemistry as a geothermal prospecting tool for hidden hydrothermal systems in Réunion Island

Bhavani Bénard o 1,2,3 <sup>™</sup>, Vincent Famin 1,2, Pierre Agrinier 1, Pascale Louvat 1,4, Geneviève Lebeau 1,2 & Pierre Burckel 1

Most untapped high-enthalpy geothermal resources are blind, meaning lacking surface evidence of their existence. The first step in their discovery is to find evidence of hydrothermal activity. Here we apply an approach based on the geochemistry of cold waters, which allowed us to identify evidence of the existence of a hydrothermal system at Piton de la Fournaise volcano (Réunion Island), and constrain its location. This approach uses the concentrations in B, Li,  $SO_4$ , F, Mo, P, V, As and  $HCO_3$  and the isotopic ratios  $\delta^{13}C$  and  $\delta^{11}B$  as geochemical markers of hydrothermal activity that can be used even in waters with extremely low ion content (Electrical conductivity <80  $\mu$ S/cm), and even when their geochemical composition is mainly controlled by other processes. This noninvasive approach is easy to implement and can be applied wherever the presence of a blind geothermal system is suspected.

<sup>&</sup>lt;sup>1</sup> Université de Paris, Institut de Physique du Globe de Paris, CNRS, F-75005 Paris, France. <sup>2</sup> Université de La Réunion, Laboratoire GéoSciences Réunion, F-97744 Saint Denis, France. <sup>3</sup> Observatoire Volcanologique du Piton de la Fournaise, Institut de Physique du Globe de Paris, F-97418 La Plaine des Cafres, France. <sup>4</sup> Université de Pau et des Pays de l'Adour, E2S UPPA, CNRS, IPREM, UMR, 5254 Pau, France. <sup>⊠</sup>email: benard@ipgp.fr

eothermal development consists in harvesting the energy naturally generated by hydrothermal activity. Geothermal exploration is therefore greatly facilitated by the presence of obvious surface markers of hydrothermal activity such as hot springs or fumaroles. While there are plenty of efficient methods for exploring geothermal systems displaying such evidence, less visible systems prove hard to uncover, thus restraining the development of geothermal energy worldwide. Yet, with the acceleration of climate change from the use of fossil fuels, geothermal energy represents a necessary renewable and local alternative. To address this issue, the scientific community in all Earth-sciences related disciplines has been developing exploration strategies for almost two decades, specifically targeting "hidden" or "blind" geothermal systems, i.e., with no obvious surface evidence of hydrothermal activity<sup>1-7</sup>. For high-enthalpy systems and conventional geothermal use, the first milestone is to determine whether or not there is hydrothermal activity, as it represents the base condition. So far, few approaches have been proposed using water geochemistry. Indeed, the widely used geochemical tools in non-blind systems rely on the study of hot springs and fumaroles, absent in blind systems. Yet, small hydrothermal components are often highlighted in groundwater studies in volcanic environments<sup>8–22</sup> and could therefore theoretically constitute an exploration tool. In most cases the quantification of the hydrothermal contribution is made using anionic mass budgets (Cl, SO<sub>4</sub>, HNO<sub>3</sub>) but also trace elements concentrations and isotopic ratios that are typically enriched (B, Li, ...) or depleted (Se, Ge) or fractionated ( $\delta^7 \text{Li}$ ,  $\delta^{11} \text{B}$ ,  $\delta^{34} \text{S}$ ,  $\delta^{13} \text{C}$ ,  $\delta^{30} \text{Si}$ ) in the hydrothermal fluids compared to the "cold" waters. Only in the recent exploration campaigns in the islands of Hawai'i, unconventional geochemical tools were used on cold groundwaters, whose compositions result mostly from non-hydrothermal processes, which along with other methods proved efficient to evidence the existence of hidden geothermal systems<sup>3</sup>. In these ocean island volcanoes, it is assumed that strong lateral groundwater flows conceal the rise of hot fluids and that to some extent, these hot fluids are diluted in a wider hydrogeological network. While bulk geochemical data for these "mixed" groundwaters fall into the range of expected values for "pure" groundwaters, by comparing the temperatures, Cl/Mg concentration ratios and SiO2 concentrations between groundwaters samples harvested at the scale of the islands, Lautze et al.3 were able to decipher geochemical anomalies associated with hydrothermal activities in some samples. Ferguson<sup>23</sup> showed that these waters also displayed anomalous He content and R/Ra ratios. These studies touch upon the possibilities of using geochemical markers of hydrothermal activity in cold groundwaters as an exploration tool for the first stage of geothermal exploration in blind systems, which is the approach we develop here on a larger scale.

We chose the Piton de la Fournaise volcano (Réunion Island) as our natural laboratory. This ocean island volcano displays no visible surface indicators of hydrothermal activity despite an ongoing frequent volcanic activity. As in other blind systems, geophysical studies as well as studies on the geochemistry of soil gas have been conducted for geothermal exploration, which concluded to the possible presence of hydrothermal systems at different locations in the volcanic edifice<sup>24</sup>. Additionally, in unrelated studies, observed peculiarities in the Radon content and the isotopic composition ( $\delta^{13}$ C and  $\delta^{11}$ B) of several cold groundwater samples of the massif have tentatively been attributed to the presence of magmatic fluids<sup>25–27</sup>. Therefore, to date, the presence of a hidden geothermal system on this volcano, although strongly suggested, is still questioned, making it an ideal candidate for the development of new exploration strategies such as the study of the chemistry of cold waters, as a complementary approach to the other methods. Here, we acquired a large dataset

both in terms of samples and analyzed parameters to try and find markers of a hydrothermal signature and fit them in a hydrogeological circulation model. We sampled 80 cold springs, groundwater wells, rivers and streams on the volcanic massif (Fig. 1A) and analyzed them for major and trace elements concentrations and  $\delta^{11}B$  and  $\delta^{13}C$  isotopic ratios.

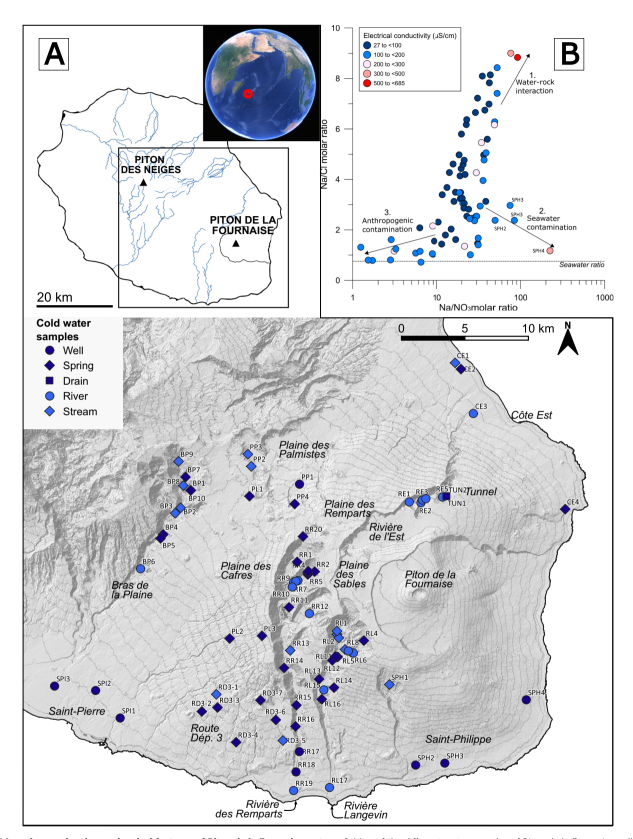
We show that high temperature water-rock interaction and/or dissolution of magmatic gases can be evidenced in Piton de la Fournaise waters by the relative enrichments in As, Rb, P, V, Mo, Sb, F, Li,  $SO_4$ ,  $HCO_3$  and B, as well as by the isotopic ratios  $\delta^{11}B$  and  $\delta^{13}C$ . With these geochemical markers of hydrothermal activity, we built a unique index that measures the relative probability of the presence of a hydrothermal component in waters. The spatial distribution of waters with the most evidence of hydrothermal activity indicates that a caldera fault system surrounding the Plaine des Sables region is a preferential pathway for rising hydrothermal fluids. These results are consistent with previous geophysical exploration studies which identified a dense body and a conductive layer as a potential geothermal resource in this region  $^{24,28,29}$ .

### **Results and discussion**

Identifying geochemical markers of hydrothermal activity. At Piton de la Fournaise, the aquifer system is unconfined and constituted by stacked lava flows<sup>30</sup>. Lavas are transitional between the tholeitic and alkaline fields and have a range of compositions from picrite to trachyandesite, with a large majority of aphyric basalt<sup>31-33</sup>. At a regional scale, permeability is decreasing from the surface to the depth of the edifice, the deeper units being older and thus more affected by secondary mineralization processes. These oldest units act as a wall for interconnected groundwater flows of regional extent with waters going from the inner to the outer part of the edifice. Preferential pathways or isolated circulations can result from local discontinuities in the stacked lava flows, such as permeable paleo-valleys and less permeable paleosoils<sup>30,34,35</sup>. Because the aquifer system is interconnected and of regional extent, we expect that the presence of a hydrothermal system would result in a dilution of thermal waters into the aquifer system, as in Hawai'i<sup>3</sup>.

There are no resurgences of waters displaying a strong thermal component at Piton de la Fournaise, which was confirmed by our extensive fieldwork. From their interaction with hot rocks, hot fluids, and magmatic gases, thermal waters are hot and/or acidic and/or with a high total dissolved solid content. Piton de la Fournaise waters wave low temperatures, ranging between 8.5 and 21.1 °C, and neutral pH values ranging between 6.5 and 8.6. Most samples also display low electrical conductivities, although five (TUN1, TUN2, TUN3, SPH3, SPI3) display values superior to the others (266 to 685  $\mu$ S/cm against a mean of 129  $\pm$  100  $\mu$ S/cm, Supplementary Data 1 and Fig. 1B).

The mineralization of rainwaters, which infiltrate the volcanic edifice, is mostly influenced by sea spray and thus have a Na/Cl ratio close to that of seawater, with however low Na and Cl concentrations<sup>36</sup>. From this initial composition, as water circulates through the ground, its composition evolves according to different mineralization processes. In Fig. 1B, the comparison of Na/Cl and Na/NO<sub>3</sub> molar ratios gives three trends corresponding to three common mineralization processes: water-rock interaction, seawater contamination, and anthropogenic contamination. As the water residence time in the aquifers increases, water-rock interaction increases the Na concentration through the dissolution of silicate bearing minerals<sup>36</sup>, giving the first trend. Additionally, seawater may contaminate samples collected at river mouths and in wells close to the coast (SPH2, SPH3, SPH4), increasing both Na and Cl concentrations<sup>36</sup>, giving the



**Fig. 1 Locations and main geochemical features of Piton de la Fournaise waters. A** Map of the different water samples of Piton de la Fournaise collected for this study and (**B**) Na/Cl vs. Na/NO<sub>3</sub> molar ratios and electrical conductivity (μS/cm) of Piton de la Fournaise waters, showing that water rock interaction, anthropogenic contamination and seawater contamination are the main mineralization processes. Insert: ©2023 Google Earth.

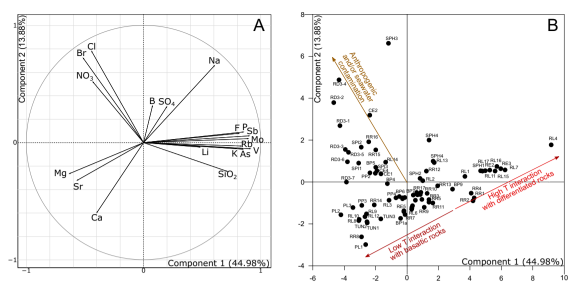


Fig. 2 Principal component analysis of the major and trace elements composition of Piton de la Fournaise waters. Distribution of (A) variables and (B) indices for components 1 and 2, representing 58.86% of the total variance.

second trend. Finally, the presence of urban areas and agricultural lands located on the catchment areas of several sampled watersheds can lead to the infiltration of artificially chlorinated waters from the domestic water supply, of organochlorine compounds and muckspreading used for agriculture and of wastewaters. These anthropogenic pollutions result increased Cl and NO<sub>3</sub> concentrations in waters, giving the third trend. The electrical conductivity of Piton de la Fournaise waters (blue to red scale, Fig. 1B), which is a proxy for their total dissolved mineral content, increases along each trend. This means that those three processes contribute the most to the geochemical composition of the waters. Here, we are searching for evidence of mineralization associated with hydrothermal processes. Seawater contamination and anthropogenic contamination are not hydrothermal processes. Water-rock interaction can occur at low temperature, but it is facilitated by heat and acidity, and can thus be increased by hydrothermal activity. However, from the data presented in Fig. 1B alone, it is not possible to decipher whether water-rock interaction evidenced here is associated with hydrothermal activity.

For a broader understanding of the geochemical composition of our samples, and in order to highlight a possible hydrothermal contribution, we performed a principal component analysis (PCA) on a large geochemical dataset of major and trace element compositions (variables) of our water samples from Piton de la Fournaise (individuals).

Among all of the analyzed elements, the variables retained for interpretation are:

Major elements (except HCO<sub>3</sub> for which not all samples had available data),

Trace elements that can originate from hydrothermal activity Li, B, Br, F, P, Sb, Mo, Rb, V, As, Sr,

SiO<sub>2</sub> as a maker of water rock interaction due to the weathering of silicate minerals<sup>30,37</sup>,

NO<sub>3</sub> as a marker of anthropogenic contamination.

We did not include  $HCO_3$  in the PCA because it requires a homogeneous dataset and  $HCO_3$  concentrations had not been measured for some samples.

Using elemental concentrations for the PCA gives larger weights to the elements with the highest concentrations. Since in our case the hydrothermal component may not constitute the main geochemical feature, and since most elements associated with hydrothermal activity are trace elements, we chose to rather study the enrichment in each element compared to the others for each sample. To obtain this enrichment, we normalized the data by dividing each elemental concentration by the sum of the concentrations of the major cations (Ca, Mg, Na, K).

The results of the analysis are presented in Fig. 2A, B, which display the contribution of each chemical element to the two principal components, and the distribution of water samples on these components, respectively.

The first component explains 44.98% of the total geochemical variability. The elements contributing the most to this component are F, P, Sb, Mo, Rb, V, K, As, and to some extent SiO2, Na and Li. F, P, Sb, Mo, Rb, V, K, As plot close together and define a pole. These chemical elements are found in magmatic gases, their condensates and associated deposits<sup>38-41</sup>, as observed at Piton de la Fournaise<sup>42,43</sup>, which can dissolve in waters. But in our case, F, P, Sb, Mo, Rb, V, K and As plot close to SiO<sub>2</sub>, which is a marker of water-rock interaction, so we preferably explain their presence by a high temperature interaction with differentiated rocks<sup>44–48</sup>, and thus by hydrothermal activity. By comparison, at the other volcano of La Réunion, Piton des Neiges, thermal springs display high concentrations in F, Rb, K, As and SiO<sub>2</sub> but not in V and P, and there is no available data for Mo and Sb49. Elements that can form oxyanions such as P, V, Mo and Sb have a higher solubility in oxidizing environments<sup>47,50–52</sup>. Therefore, they are more likely to be present in an unconfined aquifer such as in Piton de la Fournaise, especially at the top of the water column, where the conditions are more oxic.

The second component explains 13.88% of the total geochemical variability. The elements contributing the most to this component are Br, Cl, NO<sub>3</sub>, Na and to some extent, B and SO<sub>4</sub>. Br, Cl and NO<sub>3</sub> plot close together and define a second pole which we infer to be anthropogenic and/or seawater contamination.

A third pole can be identified on the negative end of both components, which corresponds to an enrichment in Mg, Ca and Sr. In the aquifers of Réunion Island, Ca and Mg are inferred to come from low temperature interaction with basaltic rocks<sup>30</sup>, which we infer to be the main mineralization process for this third pole. However, an enrichment in magmatic gases can cause the pH to decrease and facilitate water-rock interaction.

Therefore, an enrichment in Mg, Ca and Sr could also be a knockon effect of hydrothermal activity.

Na plots close to the circle and in-between the components 1 and 2, meaning it is contributing to both.

Elements Li, B and SO<sub>4</sub> are poorly represented on the two main components resulting from the PCA analysis (Fig. 2A), indicating that they originate from multiple sources. Sulfur species, Li and B are found in magmatic gases at Piton de la Fournaise<sup>53,54</sup> that could contribute to a hydrothermal system. Li and B can also come from high temperature interaction with differentiated rocks<sup>45,55–57</sup>. At Piton des Neiges volcano, previous studies show that several thermal springs and rivers streams contain high concentrations of Li, B, and SO<sub>4</sub> (up to 137.1 µmol/l, 1545.4  $\mu$ mol/l and 8.28 mmol/l, respectively) with low  $\delta^7$ Li and δ<sup>11</sup>B values, which can be associated with hydrothermal activity<sup>49,58,59</sup>. At Piton de la Fournaise, as expected, Li, B, and SO<sub>4</sub> concentrations are several orders of magnitude lower than those of Piton des Neiges thermal springs. However, some samples display higher concentrations in these elements (up to 0.22 µmol/l, 14.92 µmol/l and 0.09 mmol/l, respectively) compared to all samples (median values: 0.02 µmol/l, 0.48 µmol/l and 0.01 mmol/l, respectively), which could correspond to a small hydrothermal contribution. Regarding the Li content of Piton de la Fournaise waters, it should be noted that Henchiri et al.<sup>58</sup> analyzed the composition in Li at two locations, the mouth of Rivière Langevin and the mouth of Rivière de l'Est. Samples were taken at the same locations for our study. The two studies show that the Li concentration is low: 0.032 and 0.009 µmol/l, respectively, in Henchiri et al.<sup>58</sup>, and 0.021 and 0.025 µmol, respectively, in our study (Supplementary Data 1). Therefore, we do not consider these datapoints to be representative of the enriched Li pole with a possible hydrothermal input which we identified. Indeed, their  $\delta^7$ Li values do not indicate a contribution from a hydrothermal source.

The hydrothermal contribution in boron can come from the dissolution of the host rock, of a magmatic gas phase or of fumarolic condensates or sublimates. Little fractionation is expected during these processes<sup>60</sup>, and therefore, in any case the  $\delta^{11}B$  value should be close to that of ocean island basalts (OIB). There are no available  $\delta^{11}B$  values for Réunion basalts, but other OIB unaltered basalts display  $\delta^{11}B$  values between -12 and  $-3\%^{61}$ . Hydrothermal alteration or contamination by subducted slabs can increase these values. At Piton des Neiges volcano, where hydrothermal alteration is evidenced in the host rock<sup>62</sup>, thermal springs have an isotopic ratio  $\delta^{11}B$  comprised between 0.2 and 3.9% (this study<sup>26</sup>). At Piton de la Fournaise, where there is less hydrothermally altered rocks<sup>62–64</sup>, the  $\delta^{11}B$  value of the hydrothermal contribution should be lower, close to the basalt value. Additionally to a possible hydrothermal contribution, in groundwaters and river waters, boron is controlled by water-rock interaction at low temperature, by processes fractionating boron isotopes (mainly the sorption onto mineral and organic surfaces and coprecipitation of boron in secondary minerals, all favoring incorporation of isotope <sup>10</sup>B, and leaving the waters enriched in <sup>11</sup>B, so with higher  $\delta^{11}B^{65}$ ), but also by the biological productivity, by the influence of sea spray in the infiltrated rainwater and by mixing with seawater. All of these other contributions are expected to result in natural waters displaying a higher  $\delta^{11}$ B isotopic ratio than that of thermal waters (>12\%^65). Therefore, the isotopic ratio  $\delta^{11}B$  can be used to discriminate the hydrothermal contribution from these other sources of B in groundwaters and river waters<sup>66-68</sup>. Only anthropogenic contamination would possibly bring B with low  $\delta^{11}$ B signatures, either issued from urban/industrial uses or from fertilizers<sup>69–71</sup>. Measured  $\delta^{11}$ B values for Piton de la Fournaise waters range from

-5.0 to 42.7‰ in agreement with previous values by Louvat et al.<sup>58</sup> (from 3.7 to 44.2‰). Nine samples out of the 28 for which  $\delta^{11}$ B was analyzed display low  $\delta^{11}$ B values (typically <12‰): TUN1, TUN2, TUN3, RL4, RL1, RE4, RR3, RL5 and RL3. These samples come from preserved natural environments, and do not show anthropogenic contamination according to their position in the principal component analysis plot (Fig. 2). Therefore, these waters are interpreted as being strongly influenced by a hydrothermal system.

CO<sub>2</sub> is the most abundant magmatic gas<sup>72</sup> and can dissolve in waters. Considering the pH  $(7.6 \pm 0.4)$  and temperature  $(16.2 \pm 2.6 \,^{\circ}\text{C})$  of the waters of Piton de la Fournaise, this dissolved carbon should be mostly in the form of HCO3. The HCO<sub>3</sub> content can thus be an indicator of the presence of magmatic degassing. HCO<sub>3</sub> concentrations of Piton de la Fournaise waters range between 0.20 and 4.77 mmol/l. This upper value is in the range of Piton des Neiges thermal waters  $(3.1 to 42.7 mmol/l^{49})$ , indicating that there is a probable magmatic source of carbon at Piton de la Fournaise. However, part of the dissolved carbon can also come from a biogenic source. The isotopic ratio  $\delta^{13}$ C of the total dissolved inorganic carbon (TDIC) can be used to discriminate between a biogenic and a magmatic source of carbon. At Piton de la Fournaise, magmatic gaseous  $CO_2$  is expected to have a  $\delta^{13}C$ value comprised between -0.5 and  $-8\%^{73-75}$ . Dissolution of this gaseous carbon in waters does not affect the isotopic ratio, but there is an isotopic fractionation if this CO<sub>2</sub> is lost through degassing from the water, resulting in higher values of  $\delta^{13}$ C in the residual waters<sup>49,74,76</sup>. A biogenic contribution (through the respiration of micro-organisms in the soils and vegetation C use) results in lower values of  $\delta^{13}$ C in waters ( $\delta^{13}$ C  $\leq -20$  in Réunion<sup>77</sup>). For Piton de la Fournaise waters, we were only able to get  $\delta^{13}$ C isotopic ratios for 54 out of 82 samples, the remaining samples having too low TDIC (see methods section).  $\delta^{13}$ C values range between -21.4 and 0.6% (Supplementary Data 1). Fourteen samples display  $\delta^{13}$ C values in the range of that of magmatic CO<sub>2</sub> and 3 have a clear biogenic signature. For the others, we infer that the carbon content is a mix between both sources. In Fig. 3, we represented the  $\delta^{11}$ B and  $\delta^{13}$ C values for the 25 samples for which both parameters were analyzed. Three samples display  $\delta^{11}$ B and  $\delta^{13}$ C in the range of the expected values for B and C originating from magmatic or hydrothermal activity, and several appear to tend toward this pole.

Piton de la Fournaise hydrothermal system revealed. From our geochemical study, we can retain several geochemical markers of hydrothermal activity: the relative enrichments in As, Rb, P, V, Mo, Sb, F, Li, SO<sub>4</sub> and B compared to other elements, the concentration in HCO<sub>3</sub> and the isotopic ratios  $\delta^{11}B$  and  $\delta^{13}C$ . These markers are found in several samples and in different proportions. In this section, we use these markers to compare the hydrothermal contributions in the different samples.

Each of the markers (Y) can be converted to a linear scale from 0 to 1 (index  $Y_i$ ) measuring the relative likelihood of the presence of a hydrothermal component (Table 1). Depending on the elements, additional operations can be made to account for the influence of the dilution of the hydrothermal fluid and of seawater and anthropogenic contaminations.

The enrichments in As, Rb, P, V, Mo, Sb and F, Li, SO<sub>4</sub> and B are obtained by dividing the elemental concentrations by the sum of the major cations, which approximates the total mineral content. This corrects for both dilution of hydrothermal fluids and seawater contamination, as both affect the global geochemical composition of waters. SO<sub>4</sub> and B concentrations can be affected also by anthropogenic contamination, which only partially affect

the global geochemical composition. Therefore, here, we correct for anthropogenic contamination by also dividing the  $SO_4$  and B concentrations by the  $NO_3$  concentration (Table 1).

 $\rm HCO_3$  concentration can be affected by dilution. However, this compound is inferred to originate from  $\rm CO_2$  dissolution, which promotes mineral dissolution and in turn increases the total mineral content. Hence,  $\rm HCO_3$  concentration cannot be corrected for dilution.

While elemental concentrations of B and HCO<sub>3</sub> give a relative probability of the presence of a hydrothermal component of each water sample compared to the others, isotopic ratios  $\delta^{11}$ B and

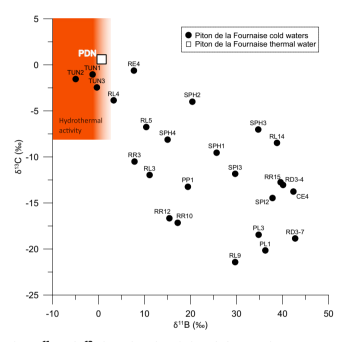


Fig. 3  $\delta^{11}B$  and  $\delta^{13}C$  isotopic ratios of Piton de la Fournaise waters compared to a thermal water sample from Piton des Neiges. The expected ranges for boron and carbon originating from hydrothermal activity are also represented by the red field 14,34,49,53,55-57.

 $\delta^{13}C$  can be considered as absolute linear scales. The maximum likelihood of the presence of a hydrothermal component corresponds to the highest value for  $\delta^{13}C$  and to the lowest value for  $\delta^{11}B$ , resulting in an inverse scale (Table 1).

We use these indexes  $Y_i$  to build a global hydrothermal index (HI = Hydrothermal Index, Eq. (1)). All indexes  $Y_i$  are normalized between 0 and 1, weighted and summed (Table 1). From the result of the PCA, we know that As, Rb, P, V, Mo, Sb and F are representative of a unique hydrothermal process. Therefore, their 7 indexes  $Y_i$  are summed, and each is given a weight of 1/7. Other indexes are each given a weight of 1. The hydrothermal index is then computed with the following equation:

$$HI = \frac{\frac{As_{i} + Rb_{i} + P_{i} + V_{i} + Mo_{i} + Sb_{i} + F_{i}}{7} + SO_{4,i} + B_{i} + Li_{i} + HCO_{3,i} + \delta^{11}B_{i} + \delta^{13}C_{i}}{n}$$
(1)

where n corresponds to the number of terms of the numerator.  $\delta^{11}$ B and/or  $\delta^{13}$ C were not analyzed for all samples. As a result, n equals at least 5 and at most 7. We then normalized the hydrothermal index between 0 and 1, giving the number HI<sub>norm</sub>.

The spatial distribution of the values of the normalized hydrothermal index HI<sub>norm</sub> in Piton de la Fournaise waters is represented in Fig. 4, along with major geological features that can constitute preferential pathways for hydrothermal fluids or heat sources: caldera faults<sup>78</sup>, rift zones<sup>79</sup> dense bodies and conductive layers<sup>9</sup>. The waters most likely to have a hydrothermal component (HI<sub>norm</sub> > 0.5) are all located on or close to the caldera faults surrounding the Plaine des Sables region, except for four of them (wells SPH2, SPH3, SPH4 and spring RL15). However, these four samples drain an aquifer which can be considered continuous from the Summit/Plaine des Sables region down to the shore<sup>30,80</sup>, meaning that their geochemical composition could be inherited from upstream. This spatial distribution indicates that the caldera faults act as preferential pathways for rising hydrothermal fluids. In La Plaine des Sables region, located at the center of this caldera fault system, a dense body and a conductive layer have already been identified as a possible geothermal resource (Fig. 4). Our study provides additional evidence in favor of this interpretation. Therefore, la Plaine des Sables can be considered the best target for geothermal exploration in terms of geothermal potential. This

Marker Y	Index Y <sub>i</sub>	Corrections	Weight
[As]	$As_i = \frac{[As]}{[Ca] + [Mg] + [Na] + [K]'}$ normalized between 0 and 1	Dilution, Seawater contamination	1/7
[Rb]	$Rb_i = \frac{[Rb]}{[Ca] + [Mg] + [Na] + [K]'}$ , normalized between 0 and 1		1/7
[P]	$P_i = \frac{[P]}{[Ca]+[Mg]+[Na]+[K]'}$ normalized between 0 and 1		1/7
[V]	$V_i = \frac{[V]}{[Ca] + [Mg] + [Na] + [K]'}$ , normalized between 0 and 1		1/7
[Mo]	$Mo_i = \frac{[Mo]}{[Ca]+[Mg]+[Na]+[K]'}$ normalized between 0 and 1		1/7
[Sb]	$Sb_i = \frac{[Sb]}{[Ca] + [Mg] + [Na] + [K]'}$ normalized between 0 and 1		1/7
[F]	$F_i = \frac{[F]}{[Ca] + [Mg] + [Na] + [K]'}$ normalized between 0 and 1		1/7
[Li]	$Li_i = \frac{[Li]}{[Ca] + [Mg] + [Na] + [K]'}$ , normalized between 0 and 1		1
$[SO_4]$	$SO_{4,i} = \frac{[SO_4]/[NO_3]}{[Ca]+[Mg]+[Na]+[K]}$ , normalized between 0 and 1	Dilution, Seawater contamination, Anthropogenic contamination	1
[B]	$B_i = \frac{[B]/[NO_3]}{[Ca]+[Mg]+[Na]+[K]'}$ normalized between 0 and 1		1
[HCO <sub>3</sub> ]	$HCO_{3,i} = [HCO_3]$ , normalized between 0 and 1		1
$\delta^{11}B$	$\delta^{11}B_i = 1/\delta^{11}B$ , normalized between 0 and 1		1
$\delta^{13}C$	$\delta^{13}C_i = \delta^{13}C$ , normalized between 0 and 1		1

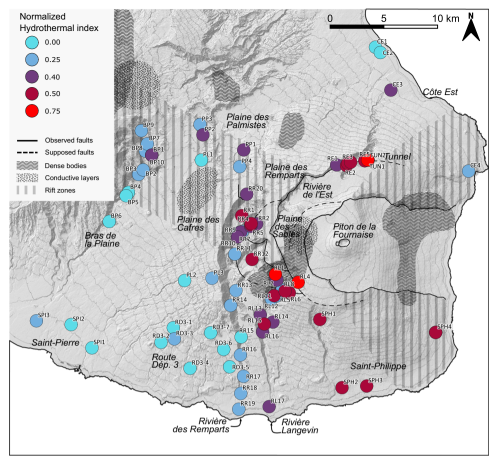


Fig. 4 Spatial distribution of the normalized hydrothermal index, indicating the relative probability of the presence of a hydrothermal component in the waters of Piton de la Fournaise. Geological, structural and geophysical features are also represented: caldera faults<sup>78</sup>, rift zones<sup>79</sup> dense bodies and conductive lavers<sup>24,28,29</sup>.

region has exceptional landscapes and biodiversity, and as such is part of a national park and is a UNESCO World heritage site, protected by strict environmental regulations, which any geothermal project will need to account for.

Our study demonstrates that the geochemistry of cold waters is an efficient way to address the first stage of geothermal exploration in blind systems, which is to determine whether or not there is hydrothermal activity generating energy at depth, and it proves to be complementary to other approaches. Regardless of the extent of the contribution of hydrothermal activity to the geochemical composition of waters, as long as it exists, the study of relevant elemental and isotopic ratios can highlight it. Spatializing this diffuse hydrothermal signal points to geographical and structural targets for geothermal exploration.

## Methods

The groundwaters and surface waters of Piton de la Fournaise were sampled during the dry season in 2020 and 2021, when the water tables are at a low level, in an effort to minimize mixing with newly infiltrated rainfall. These campaigns yielded a total of 82 water samples (from 80 sampling points) to be analyzed for major and trace elements concentrations and  $\delta^{13}C$  and  $\delta^{11}B$  isotopic ratios. The pH, temperature, and electrical conductivity (EC, expressed at 25 °C) of each sample were measured in situ. All water samples were filtered at 0.2  $\mu$ m on site and the containers were rinsed three times with filtered water. Samples for the analysis of major, trace elements and B isotopes were collected in LDPE bottles. Samples for the analysis of major cations and trace elements were acidified with nitric acid (HNO<sub>3</sub> 0.6 N) on site.

Samples for the analysis of total dissolved inorganic carbon content (TDIC) and C isotopes were collected in Labco 12 ml Exetainer vials with no headspace. All samples were stored at a temperature of 6 °C before shipment.

All analyses were performed at the Institut de Physique du Globe de Paris (IPGP). Major anions, fluoride and nitrate concentrations were determined by ion exchange chromatography. Major cations and trace element concentrations were determined using inductively coupled plasma mass spectrometry (ICP-MS). The accuracy for major cations and anions concentrations is <5%, and <20% for trace elements, as inferred from replicate measurement of solutions of known compositions. The TDIC and carbon isotope ratios of the TDIC were determined together by gas chromatography and isotope ratio mass spectrometry (GC-IRMS), after releasing the TDIC as CO<sub>2</sub> by H<sub>3</sub>PO<sub>4</sub> acidification in a vial from an aliquot of 5 ml<sup>81</sup>. For the analysis with the GC-IRMS, a finite amount of gas is pumped inside the vial and sent into the analysis circuit. This step is repeated several times and the TDIC and  $\delta^{13}$ C are computed from the results of several runs. This method proved not adapted for the analysis of our samples due to a too low content of TDIC. With an aliquot of just 5 ml, the available gaseous CO<sub>2</sub> in the vial sampled by the GC-IRMS was too low and the analysis did not yield results for several runs and for several samples. We obtained reliable results for only 51 out of 82 samples. For these samples, HCO<sub>3</sub> concentrations were computed from TDIC, pH and temperature, using the temperature dependent acid dissociation constants from Mook and Koene,  $(1975)^{82}$ . For the other samples, HCO<sub>3</sub> concentrations were estimated for an ionic balance equal to 0%.  $\delta^{13}$ C are

reported relative to PDB standard with a 1 SD reproducibility,  $\pm 0.05\%$ . Boron isotope ratios were determined by MC-ICP-MS with a direct injection nebulizer (d-DIHEN83), after boron extraction through ion chromatography84.  $\delta^{11}B$  are reported relative to NIST SRM951 standard with a 2 SD reproducibility, between 0.1 and 0.4‰.

### **Data availability**

The data used in this manuscript are available as Supplementary Data and also through IPGP Research Collection (research-collection.ipgp.fr) at the following https://doi.org/10.18715/IPGP.2023.lq6ndlzz<sup>85</sup>.

Received: 15 May 2023; Accepted: 9 January 2024; Published online: 29 January 2024

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# **Acknowledgements**

We thank Alix Marianne, Chris Payet and Manaarii Salmon for their participation in the fieldwork. The Communauté d'Agglomération du Sud de La Réunion (CASUD) funded the postdoctoral fellowship of Bhavani Bénard and this research. Chemical analyses were also supported by IPGP multidisciplinary program PARI and Paris–IdF region SESAME (Grant no. 12015908).

### **Author contributions**

Bhavani Bénard contributed to all aspects of the project and wrote the article. Vincent Famin supervised the project, took part in the field work and data interpretation, and contributed to writing the article. Pierre Agrinier and Pascale Louvat conducted some of the analytical work of this project and contributed to interpreting the data and writing the article. Geneviève Lebeau contributed to the conceptualization and execution of the field work. Pierre Burckel conducted some of the analytical work of this project.

### **Competing interests**

The authors declare no competing interests.

### **Additional information**

**Supplementary information** The online version contains supplementary material available at https://doi.org/10.1038/s43247-024-01210-3.

Correspondence and requests for materials should be addressed to Bhavani Bénard.

Peer review information *Communications Earth & Environment* thanks the anonymous reviewers for their contribution to the peer review of this work. Primary Handling Editors: Lucia Pappalardo and Joe Aslin. A peer review file is available.

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