The caves of the Irazú volcano (Costa Rica), became accessible after the partial collapse of the NW sector of the Irazú volcano in 1994, offering the opportunity to investigate active minerogenetic processes in volcanic cave environments. We performed a detailed mineralogical and geochemical study of speleothems in the caves Cueva los Minerales and Cueva Los Mucolitos, both located in the northwest foothills of the main crater. Mineralogical analyses included X-ray diffraction (XRD) and Raman spectroscopy, while geochemical characterization used Energy Dispersive X-ray spectroscopy (EDX) coupled to Scanning Electron Microscopy (SEM). In addition, measurements of environmental parameters in the caves, cave drip water and compilation of geochemical analyses of the Irazú volcanic lake (~150 m above the cave level) and fumarole analyses were conducted between 1991 and 2014. We identified forty-eight different mineral phases, mostly rare hydrated sulfates of the alunite, halotrichite, copiapite, kieserite and rozenite groups, thirteen of which are described here as cave minerals for the first time. This includes the first occurrence in cave environments of aplowite, bieberite, boyleite, dietrichite, ferricopiapite, ferrinatrite, lausenite, lishizhenite, magnesiocopiapite, marinellite, pentahydrite, szomolnokite, and wupatkiite. The presence of other new cave minerals such as tolbachite, mercallite, rhomboclase, cyanochroite, and retgersite, is likely but could not be confirmed by various mineralogical techniques. Uplifting of sulfurous gases, water seepage from the Irazú volcanic lake and hydrothermal interactions with the volcanic host rock are responsible for such extreme mineralogical diversity. These findings make the caves of the Irazú volcano a world-type-reference locality for investigations on the formation and assemblage of sulfate minerals and the biogeochemical cycle of sulfur, with potential implications for Astrobiology and Planetary science.

Extremely high diversity of sulfate minerals in caves of the Irazú Volcano (Costa Rica) related to crater lake and fumarolic activity

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Abstract: The caves of the Irazú volcano (Costa Rica), became accessible after the partial collapse of the NW sector of the Irazú volcano in 1994, offering the opportunity to investigate active minerogenetic processes in volcanic cave environments. We performed a detailed mineralogical and geochemical study of speleothems in the caves Cueva los Minerales and Cueva Los Mucolitos, both located in the northwest foothills of the main crater. Mineralogical analyses included X-ray diffraction (XRD) and Raman spectroscopy, while geochemical characterization used Energy Dispersive X-ray spectroscopy (EDX) coupled to Scanning Electron Microscopy (SEM). In addition, measurements of environmental parameters in the caves, cave drip water and compilation of geochemical analyses of the Irazú volcanic lake (~150 m above the cave level) and fumarole analyses were conducted between 1991 and 2014. We identified forty-eight different mineral phases, mostly rare hydrated sulfates of the alunite, halotrichite, copiapite, kieserite and rozenite groups, thirteen of which are described here as cave minerals for the first time. This includes the first occurrence in cave environments of aplowite, bieberite, boyleite, dietrichite, ferricopiapite, ferrinatrite, lausenite, lishizhenite, magnesiocopiapite, marinellite, pentahydrite, szomolnokite, and wupatkiite. The presence of other new cave minerals such as tolbachite, mercallite, rhomboclase, cyanochroite, and retgersite, is likely but could not be confirmed by various mineralogical techniques. Uplifting of sulfurous gases, water seepage from the Irazú volcanic lake and hydrothermal interactions with the volcanic host rock are responsible for such extreme mineralogical diversity. These findings make the caves of the Irazú volcano a world-type-reference locality for investigations on the formation and assemblage of sulfate minerals and the biogeochemical cycle of sulfur, with potential implications for Astrobiology and Planetary science.

Keywords: hydrated sulfates, sulfate speleothems, volcanic caves, crater lake, cave minerogenesis

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INTRODUCTION

Although still poorly studied compared to karstic carbonate environments, lava tubes and volcanic caves have drawn the interest of many speleological and mineralogical investigations over the past 20 years (Forti, 1994, 2005, Forti et al., 2003, 2004; Daza & Bustillo, 2014; Miller et al., 2014, among others). Increasing attention is being paid to minerogenesis and weathering processes in terrestrial magmatic materials, and in particular to subterranean volcanic environments, as analogues of mineral-forming mechanisms on Mars and other planets (e.g., Morse et al., 2011; Miller et al., 2014; Lalla et al., 2015, 2016). Indeed, volcanic caves host over 40% of the total number of cave minerals reported to date, whereas...
10% of the mineral species found in caves worldwide are limited to subterranean volcanic environments (Hill & Forti, 1997; Forti, 2005; Onac & Forti, 2011a). Silicate minerals (e.g., opal and clay minerals) and carbonate speleothems (predominantly made of calcite and/or aragonite), sometimes derived from biological processes, are relatively common in volcanic caves (Daza & Bustillo, 2014; Miller et al., 2014). However, sulfates are the main group in terms of mineralogical diversity, representing over 70% of the different minerals reported from volcanic caves (Hill & Forti, 1997; Forti, 2005). The five main processes leading to sulfate speleothem precipitation in volcanic caves include: (1) deposition from aerosols, (2) evaporation of SO$_4^{2-}$ rich waters, (3) alteration and weathering (e.g., oxidation of reduced sulfur species), (4) biomediated mineral precipitation, and (5) bat guano mineralization.

Sulfur-rich caves commonly present sulfate speleothems (secondary mineral deposits formed in caves) such as stalactites, stalagmites, and crusts. Less common are geysermites, vent-shaped speleothems with a central hole that continues in the cave floor and are formed by rising thermal fluids (vapor) (Hill & Forti, 1997 and references therein); rims and blisters, formed in fractures with fumarole vapors influence (Onac & Forti, 2011b). In some of these sulfur-rich caves [e.g., Cueva de Villa Luz in Mexico, Hose et al. (2000); Frasassi cave in Italy, Jones et al. (2016)] white mucous-like soda straw biofilms hanging from cave walls and known as “snottites” (Hose & Pisarowicz, 1999) have been reported.

We conducted a detailed mineralogical and geospeleological study of Cueva Los Minerales (CMI) and Cueva Los Mucolitos (CMU), in the Irazú Volcano, Costa Rica (Fig. 1A). This area is characterized by its intense volcanic influence, with relevant CO$_2$ emissions (Galindo et al., 2004; Epiard et al., 2017) and evidence of passive fumarole activity and hydrothermalism (Alvarado et al., 2006). Here, we discuss factors related to minerogenesis, the formation of the caves themselves and their linkage with the hydrothermal system. In addition, we provide measurements of environmental parameters in the cave, hydrochemical analyses of cave infiltration waters, those present in the Irazú volcanic lake and in fumaroles (from 1991 to 2014). Abundant sulfur gases and dissolved sulfur species in the cave water are proposed as being responsible for the precipitation of an unusual diversity of sulfate minerals, many of which are reported here as cave minerals for the first time. The exact ages of the investigated caves are unknown, although access from the surface to these caves exist since December 8th, 1994, when a large landslide on the volcano flank occurred. Thus, minerogenetic processes in the caves of the Irazú Volcano can be considered as a recent and ongoing example of sulfate mineral precipitation in the volcanic subterranean environment.

**STUDY SITES**

The Costa Rica Volcanic Front is developed parallel to the Middle American Trench that separates the Coco and Caribbean plates. The convergence rate
between the subducting Coco Plate and the Caribbean Plate in Costa Rica is close to ~90 mm/yr (DeMets, 2001) (Fig. 1B). Upper plate deformation is affected by differences in subducting plate morphology, age, and dip, resulting in different deformation styles in northern and southern Costa Rica (Protti & Güendel, 1995). The Central Costa Rica Deformed Belt is the onshore expression of the transition of these two deformation styles (Marshall et al., 2000; Montero et al., 2013). It is characterized by shallow seismicity (<15 km) and a broad array of conjugate northwest-striking dextral faults and northeast-striking sinistral faults (Montero et al., 2013).

Irazú Volcano is located on an area of structural weakness dominated by the Atirro-Río Sucio Fault System. It consists of a 150 km long northwest-trending zone of dextral strike-slip faults with strong morphotectonic expression (Montero et al., 2013), it presents active seismicity, and it is composed of different active faults (Montero & Alvarado, 1995; Montero, 2001; Montero et al., 2013).

The two main craters of the Irazú Volcano, the highest volcano in Costa Rica (altitude 3,432 m a.s.l.), are located in the 35 by 10 km wide pull-apart basin comprised between traces of the Eastern and Western Río Sucio Faults (Montero et al., 2013). The Irazú Volcano evolved from Early Pleistocene to Holocene in age. The Irazú summit hosts the Main Crater “Crater Activo” (active 1962-1965). The older crater “Diego de la Haya” is located to the east of the summit. The remaining structure of a composite oldest crater called “Playa Hermosa” together with a prehistoric scoria cone and tuff rings can also still be recognized. Evidence of a debris avalanche can be seen to the northwest of the main crater (Alvarado et al., 2013). An intermittent volcanic lake has been present on “Crater Activo” after the last eruptive period (1965). This lake lasted until March 2013 when it disappeared, but it started forming again in 2017. According to Ramirez et al. (2013), the lake had a variable temperature (16-35°C), pH (3.0-5.85) and color over time (red, green, turquoise, mustard). The permeability due to fractures in the “Crater Activo” probably contributed significantly to the volume changes (draining) of the volcanic lake, and the lake-hydrothermal system is probably hydrogeologically connected to the Río Sucio springs, NW of the volcano (Ramírez et al., 2013; Pierre et al., 2015). Intra-crater fumaroles were reported several times for the period 1998-2001 on the Crater Activo (Ramírez et al., 2013).

To the northwest of the “Crater Activo”, there is an unstable area with strong hydrothermal alteration associated with the Central Río Sucio Fault. On this area three volcanic caves have been found (Fig. 1A): CMI, CMU, and Pizote Espantado (Ulloa et al., 2013, 2015). The caves were exposed after a debris avalanche in 1994. CMI consists of a big chamber (~50 m wide and~27 m long) with a wide entrance that enables sunlight to reach most parts of the cave. It shows the highest mineralogical and speleothem diversity and thermal anomalies. CMU is a narrow, completely dark cave, approximately 38 m long and 15 m deep, in which microbial communities (e.g., snottites, microbial mats) can easily be recognized. Pizote Espantado Cave has not been mapped and sampled, thus was not included in this study.

METHODS

Survey and sampling

Since 2011, four surveys for sampling and measuring environmental variables in the caves have been carried out. The caves were surveyed using a laser-distance meter, compass, and clinometer. Cave temperature, humidity (wet bulb/dry bulb), and light intensity were measured with portable thermometer (Kestrel 4500, ± 1°C), humidity probe (Kestrel 4500, ± 3%), and photometer equipment (CEM DT-8820 4-in-1 digital Multifunction Environment Meter). Thermal images were obtained using a thermal camera FLIRSC660. In CMI and CMU the CO₂ and H₂S concentrations at the floor level and in the cave atmosphere were measured with a West System WS0820 flux meter.

Mineral and water sampling in CMI and CMU were conducted in 2011, 2013, and 2014. Representative mineral samples were collected based on their visual appearances (e.g., color, texture) and location in the caves. Different types of speleothems (e.g., stalactites, geysermites, crusts on the walls) were collected, preferably broken specimens to minimize impact on the cave. Water samples for trace element analyses were taken in amber glass 50 ml bottles. Samples for analyses of heavy metals were acidified with HNO₃ to a pH < 2 to prevent precipitation during storage.

Hydrogeochemical and fumarole gas analyses

Analyses were carried out following certified procedures (INTE-ISO/IEC17025:2005) at Agrotec Analytical Laboratory (San Jose, Costa Rica). Trace element analyses were measured by inductively coupled plasma optical emission spectrometry (Thermo Trace Jarrel-ash 51-ij) (Eaton et al., 1998, Standard Method 3120b). Calibration was conducted using synthetic standards with a relative standard deviation (RSD) of 2% or better.

In this paper, we used cation and anion analyses of 25 water samples collected by the Instituto Costarricense de Electricidad (ICE) from June 1991 to February 1993 and 17 samples collected by the Observatorio Vulcanológico y Sismológico de Costa Rica (OVSICORI) since March 1999 to December 2010 (Supplementary Table 1). Water samples were collected in two 500 ml bottles. Cation samples were preserved with 5 N HNO₃ (one ml per 100 ml). The water samples were analysed for cations by Atomic Absorption Spectrophotometer and for anions by titration methods (bicarbonate), spectrophotometric methods and ion chromatography.
Errors in the determination of the different ions are lower than 10% in all cases.

The compiled eight fumarole gas analyses for 1991, 2003, 2006m and 2007 (Table 1), were sampled by ICE and OVSICORI using bottles under vacuum and partially filled with 4N NaOH (Giggenbach, 1975). The caustic solution was used for the analyses of F and Cl by ion chromatography and SO₂ as SO₄²⁻ after oxidation with H₂O₂ by ion chromatography.

Table 1. Historical fumarole analyses in Irazú Volcano.

| Date       | pH  | Temp (°C) | Conductivity (µS/cm) | Salinity (g/L) | F (mg/l) | Cl (mg/l) | SO₄²⁻ (mg/l) |
|------------|-----|-----------|----------------------|---------------|----------|-----------|--------------|
| 7/17/1991  | 2.35| 88        | 2.41                 | 1.08          | 1.06     | 503       |              |
| 5/27/2003  |     | 88        | 2.86                 | 0.048         | 2.673    | 42.116    |              |
| 6/12/2003  |     | 88        | 57.76                | 0.39          | 8.535    | 496.798   |              |
| 10/13/2006 |     |           | 648                  | n.a.          | 1.791    | n.a.      |              |
| 12/13/2006 | 2.88| 87        | 648                  | 0.1           |          |           |              |
| 2/4/2007   |     |           | 2.35                 | 2.235         | 2.173    | 99.224    |              |
| 5/5/2007   |     |           | 5/5/2007             | 2.863         | 86.589   | 57.761    |              |
| 5/15/2007  | 5.8  | 83        | 185.4                | 0.8           | 5.525    | 15.214    | 190.662      |
| Maximum    | 2.88| 88        | 1854                 | 0.8           | 5.525    | 86.589    | 503          |
| Mean       | 2.55| 86.5      | 1251                 | 0.45          | 2.023    | 16.862    | 231.594      |
| Minimum    | 2.35| 83        | 648                  | 0.1           | 0.048    | 1.06      | 42.116       |

source was a Laser Research Electro-Optics (REO) working at 632.8 nm. The KOSI HoloSpec f/1.8i spectrometer from Kaiser Optical covered a spectral range of 150–3800 cm⁻¹ and a spectral resolution of 5 cm⁻¹, while the CCD (charge coupled device) employed was a DV420A-CE-130 model from Andor. The Raman head used was KOSI MKII, HFPF-FC-S-632.8 model from Kaiser Optical Systems coupled by optical fiber to a Nikon Eclipse E600 microscope, which in turn was attached to a JVC TK-C1381EG videocamera for visual analysis and precise control of the measured spots. Two objectives were used, 50x and 100x allowing microanalyses of 37 and 15 μm diameter spots, respectively. The laser power on the sample was maintained around 2 mW to ensure no thermal damage occurred to the samples. Typical integration time for spectral acquisition was 10 s and 10 accumulations were done. The sample was manually scanned, while the height of focus was varied in order to optimize the intensity of the spectra signals.

In addition, 53 powdered bulk samples were analyzed using a FT-Raman Bruker instrument (model RFS100/S). The coherent laser was a Compass 1064-500 model (Nd:YAG, 1064 nm, 500 mW). The CCD Bruker used was a D418-T model, operating in 20 kV. A total of 96 micro-analyses were performed at the X-Ray diffraction (XRD) analyses were performed at the Servicios Centrales de Investigación of the University of Almería (Spain). SEM microphotographs of 26 samples were taken using a Hitachi S-3500 instrument in variable-pressure mode. The elemental chemistry was determined by EDX microprobe at different points with different typology over several grains of the samples, using an Oxford INCA 7210 diffractometer, with a Cu anode (CuKα, λ = 0.154 nm) and graphite monochromator. A Ni filter and Al sample holders were utilized. Tension and current produced by the generator were 40 kV and 30 mA respectively, for all analyses. The analysis used the 20 scanning method, using 0.400 seconds per step and within the angular limits of 5 to 70°.

Elemental composition and microphotographies

Energy Dispersive X-ray spectroscopy (EDX) coupled to Scanning Electron Microscopy (SEM) analyses were performed at the Servicios Centrales de Investigación of the University of Almería (Spain). SEM microphotographs of 26 samples were taken using a Hitachi S-3500 instrument in variable-pressure mode. The elemental chemistry was determined by EDX microprobe at different points with different typology over several grains of the samples, using an Oxford INCA 7210 X-ray detector, with a voltage of 20 kV. A total of 96 micro-analyses were performed (Supplementary Table 5). The diameter of the beam was approximately 1 μm. The limit of detection of this technique enables major elements such as Fe, Mn, O, Si, Al, Ca, Pb, Zn, and Ba to be analyzed.

RESULTS

Geological observations

The host rock of the caves consists of interbedded pyroclastic rocks, with centimetric to decimetric...
stratification and poorly selected volcanic breccia with angular volcanic clasts associated with pyroclastic density current deposits. Pyroclastic rocks are light grey, white to yellowish color, and in some areas present evidence of silicification. These rocks are normally poorly to moderately compacted or welded and weathering is intense due to harsh volcanic conditions (e.g., high temperature and volcanic gases). The dip direction (318/26°) is well defined by the strata where the caves are located, and this plane served as rupture surface of the 1994 debris avalanche. Pyroclastic rocks close to the CMU entrance, displays synthetic and antithetic Ridel shears with sub-horizontal striation and dextral motion. This fault corresponds to the Central Río Sucio Fault (Montero et al., 2013) and extends to CMI and Crater Principal (Fig. 1A and 2). To the south of the main entrance of CMI, there is a sector with evidence of runoff watermarks and mud deposits.

**Hydrogeochemistry**

Evidence for multiple aquifers has been reported in Irazú Summit (Alvarado, 1993). An ephemeral upper water level (~3,170 m a.s.l.) is associated with the geological contact between recent volcanoclastic deposits and hydrothermally altered rocks. The ephemeral main crater volcanic lake is located between ~3,080 and <3,050 m a.s.l. We provide geochemical evidence for multiple aquifers has been reported in Irazú Summit (Alvarado, 1993). An ephemeral upper water level (~3,170 m a.s.l.) is associated with the geological contact between recent volcanoclastic deposits and hydrothermally altered rocks. The ephemeral main crater volcanic lake is located between ~3,080 and <3,050 m a.s.l. We provide geochemical

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**Chemical fumarole gas composition**

Fumaroles are located ~300 m northeast of the cave area (Fig. 1) and ranged in temperature from 83 to 88°C and pH from 2.35 to 2.88. Table 1 displays the chemical composition of main fumarole components from volcanic lake seepage waters, was performed in 2013 for selected elements (Fig. 3C). The existence of a deeper aquifer is suggested (Alvarado et al., 2006) associated with phreatomagmatic eruptions and the hydrothermal system in the area. This deeper aquifer could be the source of sulfate-rich water and also one brownish-yellow drain (possibly acidic) close to the base of the cliff (at ~2,825 m a.s.l.) could be associated with the upwelling of these deeper waters.
Environmental parameters

CMI presents a wide entrance ~50 m (Fig. 4A), with a main chamber developing 27 m from the cave entrance to the end of the cave and approximately 40 m wide, whereas Sala de los Cristales Verdes is 10 m wide and presents green stalactites (Fig. 4B). Due to the cave morphology (Fig. 5A), the cave is exposed to sunlight (>1.32 lux during the day), because of the very large cave entrance. Temperature in the cave ranges from 9 to 16 °C in the main chamber. Thermal map (Fig. 5B) was drawn showing the highest temperatures (28-30 °C) close to the north wall fissures (where important sulfate crusts are located) and some unmapped areas close to Sala de los Cristales Verdes show temperatures over 40 °C. The southern part of the main chamber presents the highest elevation and acts as a hot air trap, with temperatures between 30-32 °C. Lower mineral diversity was found in this section of the cave. Based on 2013 field trip measurements, the relative humidity in CMI varies from 74.2% (close to entrance) to 92.4% (inner area of the main chamber). CO₂ concentrations up to 950 ppm were measured; H₂S was 29 ppm and SO₄ and H₂ were below detection limits.

CMU is smaller in volume, the entrance is small (~50 cm) and is developed in a collapsed area, 38.2 m long and 14.6 m deep. This cave showed temperatures ranging between 16 and 18 °C and relative humidity of 96.8% in 2013. Unlike CMI, this cave is totally dark.

Infiltration water is evident in both caves, and drip points are generally associated with active microbial mats (snottites). The pH of these drips ranged from 1.00 to 2.61.

Mineralogy, speleothems and elemental composition

We identified 48 different minerals in CMI (45) and CMU (6) (Table 2). This includes 19 new cave minerals, of which 14 are referred to as “confirmed.”
Sulfates were the most common mineral group in the caves of the Irazú Volcano (91.5% of the total diversity). Most of the identified sulfates correspond to hydrated acid and sulfates (33). Also, sulfates belonging to hydrated sulfates containing hydroxyl or halogen (4), anhydrous acid and sulfates (3), anhydrous sulfates containing hydroxyl or halogen (4), and halides (1) have also been detected (Fig. 6). Most of the minerals reported in this study by Raman techniques present a percentage of match with the reference spectra from RUFF Database generally >85%.

**Sulfates**

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Table 2. List of the minerals identified in CMI and CMU. The numbers correspond to how many times the mineral species were detected using different techniques.

| Mineral                  | Chemical formula | New cave Mineral | Status | Cave | Cave location | Color | Habit | Crust crystal | Stalactites | Geysermite | 2015 XRD (%) | 2011 XRD (%) | micro-Raman (%) | FT Raman (%) | Total |
|--------------------------|------------------|------------------|--------|------|--------------|-------|-------|---------------|-------------|------------|---------------|---------------|----------------|--------------|-------|
| **NATIVE ELEMENTS (1)**  |                  |                  |        |      |              |       |       |               |             |            |               |               |                |              |       |
| Sulfur                   | S₈               |                  |        |      |              | Y, O  | MA    | X             |             |            |               |               |                |              |       |
| Tolbachite               | CuCl₂            | X                | P      | CMI  | R            | LG    | X     |               |             |            |               |               |                |              |       |
| **OXIDE (1)**            |                  |                  |        |      |              |       |       |               |             |            |               |               |                |              |       |
| Goethite                 | FeO(OH)          |                  | C      | CMU  | W            | DA    | X     |               |             |            |               |               |                |              |       |
| **PHYLLOSILICATE (1)**   |                  |                  |        |      |              |       |       |               |             |            |               |               |                |              |       |
| Kaolinite                | Al₂Si₂O₅(OH)₄    |                  | C      | CMU  | W            | Y     |       |               |             |            |               |               |                |              |       |
| **SILICATE (1)**         |                  |                  |        |      |              |       |       |               |             |            |               |               |                |              |       |
| Marinellite              | Na₆Ca₂Al₂Si₂O₁₄₂⁺⁺SO₄Cl₂·6H₂O | X | C | CMI | W, F | Y, O | HE | X |               |             |            |               |               |                |              |       |
| **SULFATES (43)**        |                  |                  |        |      |              |       |       |               |             |            |               |               |                |              |       |
| Aluminocopiapite         | [Al,Mg]Fe²⁺⁺⁺(SO₄)₉(OH,OH)₂·20H₂O | C | CMI | F | W, Y |       | X | 1 |               |             |            |               |               |                |              |       |
| Alunogen                 | Al₃(SO₄)₃(H₂O)₁₇·5H₂O | C | CMI | W, R, F | W, Y | T, GR, AC | X | 10 | 2 | 6 | 6 | 24 |           | |
| Anglesite                | Pb(SO₄) | C | CMI | W, R, F | LB, G, DG, TG | BO, AC | X | 10 | 10 | 4  | 4 | 24 |           | |
| Anhydrite                | Ca(SO₄) | C | CMI | W, R | O |       | X | 2 |               |             |            |               |               |                |              |       |
| Apjohnite                | Mn²⁺⁺⁺⁺Al₃(SO₄)₁⁺₂·22H₂O | C | CMI | W | W, TR | AC | X | X | 3 | 3 | 6 | 12 |           | |
| Aplowite                 | CoSO₄·4H₂O | X | C | CMI | W | W |       | X | 1 |               |             |            |               |               |                |              |       |
| Birberite                | CoSO₄·7H₂O | X | C | CMI | R | G |       | X | 1 |               |             |            |               |               |                |              |       |
| Blödite                  | Na₂Mg(SO₄)₂·4H₂O | P | CMI | W | Y, R |       | X | 1 |               |             |            |               |               |                |              |       |
| Boyleite                 | Zn[S'O₄]·4H₂O | X | C | CMI | R | G |       | X | 1 |               |             |            |               |               |                |              |       |
| Chalcanthite             | Cu(SO₄)·5H₂O | P | CMI | R | TG | GR |       | X | 1 |               |             |            |               |               |                |              |       |
| Copiapite                | CaFe₃⁺⁺⁺(SO₄)₂(OH)₂·20H₂O | C | CMI | W, R, F | Y |       | X | X | 16 | 11 |               | 27 |           |         | |
| Coquimbite               | (Fe²⁺⁺⁺⁺SO₄)·9H₂O | C | CMI | W, R, F | P, Y, W | X | X | 20 | 3 | 6 | 29 |           |         |         | |
| Cyanochroite             | K₂Cu(SO₄)·6H₂O | X | P | CMI | F | TR |       | X | 1 |               |             |            |               |               |                |              |       |
| Dietrichite              | ZnAl₂(SO₄)₂·22H₂O | X | C | CMI | W, F | W, TR | AC | X | X | 1 | 1 | 1 | 8 |           |         |         |
| Mineral Name          | Formula                                      | Color | Habit | Location | Reference                      |
|-----------------------|----------------------------------------------|-------|-------|----------|---------------------------------|
| Epsomite              | MgSO₄·7H₂O                                    | C     |       |          |                                 |
| Ferricopiapite         | Fe⁺³⁺Fe³⁺(SO₄)₂·2OH₂O                        | X     |       |          |                                 |
| Ferrinatrite           | Na,Fe⁺²⁺(SO₄)·2H₂O                           | X     |       |          |                                 |
| Gypsum                | Ca(SO₄)·2H₂O                                 | C     |       |          |                                 |
| Halotrichite          | Fe⁺²⁺Al(SO₄)·22H₂O                           | C     |       |          |                                 |
| Hexahydrite           | MgSO₄·6H₂O                                   | C     |       |          |                                 |
| Jarosite              | KFe⁺³⁺SO₄·6H₂O                               | C     |       |          |                                 |
| Kalinite              | KAl(SO₄)·11H₂O                               | C     |       |          |                                 |
| Lausenite             | Fe⁺³⁺SO₄·2H₂O                                | X     |       |          |                                 |
| Lithihzenite          | ZnFe⁺²⁺SO₄·1H₂O                              | X     |       |          |                                 |
| Magnesiocopiapite      | MgFe⁺²⁺SO₄·3H₂O                              | X     |       |          |                                 |
| Melanterite           | FeSO₄·7H₂O                                   | C     |       |          |                                 |
| Mendozite             | NaAl(SO₄)·11H₂O                              | C     |       |          |                                 |
| Mercallite            | KH(SO₄)·2H₂O                                 | X     |       |          |                                 |
| Metavolinite          | K₂Na₅Fe⁺²⁺Fe⁺³⁺SO₄·6H₂O                     | C     |       |          |                                 |
| Natroalunite          | NaAl(SO₄)·3H₂O                               | C     |       |          |                                 |
| Nabrojarosite         | NaFe⁺³⁺(SO₄)·6H₂O                           | C     |       |          |                                 |
| Pentahydrite          | MgSO₄·5H₂O                                   | X     |       |          |                                 |
| Pickeringite          | MgAl₂(SO₄)·12H₂O                             | C     |       |          |                                 |
| Retgersite            | Ni(SO₄)·6H₂O                                 | X     |       |          |                                 |
| Rhomboclase           | (H₂O)Fe⁺³⁺SO₄·1H₂O                           | X     |       |          |                                 |
| Römerite              | Fe⁺⁺Fe⁺³⁺SO₄·14H₂O                           | C     |       |          |                                 |
| Rozenite              | Fe⁺⁺SO₄·4H₂O                                 | C     |       |          |                                 |
| Siderotil             | Fe⁺⁺Cu(SO₄)·5H₂O                            | C     |       |          |                                 |
| Starkeyite            | MgSO₄·4H₂O                                   | C     |       |          |                                 |
| Szomolokite           | Fe(SO₄)·1H₂O                                 | X     |       |          |                                 |
| Tamarnugite           | NaAl(SO₄)·6H₂O                               | C     |       |          |                                 |
| Voltaiite             | K₃Fe⁺³⁺Fe⁺²⁺Al(SO₄)·18H₂O                    | C     |       |          |                                 |
| Wupatkiite            | CaAl(SO₄)·22H₂O                              | X     |       |          |                                 |

NOTE: Status: C: Confirmed, P: Possible. Location: W: Wall, R: Roof, F: Floor. Color: B: Blue, BL: Black, BR: Brown, CL: Colorless, DG: Dark Grey, G: Green, GA: Garnet, GY: Golden Yellow, LB: Light blue, LG: Light green, O: Orange, DO: Dark Orange, P: Purple, Pi: Pink, R: Red, TG: Turquoise green, TR: Translucent, W: White, Y: Yellow. Habit: AC: Acicular, BO: Botryoidal, DR: Drusy, GR: Granular, HE: Hexagonal, MA: Massive, T: Tabular. (#) Corresponds to number of identifications by different analyses, and does not refer to abundance.
CMU too. Hydrated acid sulfates are sub-classified according to Dana’s classification (Mindat, 2018).

Hydrated acid and sulfates with the general formula \((A)\ _2(B)(XO_4)\ _4\ _x(H_2O)\) are represented by alunogen, coquimbite (Fig. 8), and lausenite in crusts on the walls, roof, and floor of the cave. Coquimbite was also found in geysermites. Alunogen and coquimbite have been identified by Raman techniques and XRD, lausenite only by FT and micro-Raman. The presence of rhomboclase is suggested by micro-Raman (Fig. 9) and the presence of Fe in EDX analyses.

Hydrated acid and sulfates belonging to the group with general formula \((A')\ _2(B)(XO_4)\ _4\ _x(H_2O)\) are represented by the possible presence of blödite (identified by XRD) and cyanochroite (identified by micro-Raman, Fig. 9), whereas ferrinatrite and metavoltine, have been found on walls and roof of the cave.

Some hydrated acid and sulfates with formula \(A(B)(XO_4)\ _4\ _x(H_2O)\) are difficult to differentiate by Raman because of the similarity in the spectra (e.g., apjohnite, dietrichite, halotrichite, pickeringite), with peaks usually at 994 and 975 cm\(^{-1}\), and because they commonly appear in mineral mixtures. However, the combination of XRD-Raman and EDX has made identification possible. Some samples also show a minor peak at 1,025 cm\(^{-1}\) (CR-1C-A and MIM-9A) possible associated with the presence of coquimbite. Minerals from this group, have been found forming part of crusts on the walls and geysermites (Fig. 10A), in some cases associated with ferricopiapite (Fig. 10B), and generally showing acicular habit (Fig. 10C, D). Kalinite, mendozite, and tamarugite, have been detected in several samples.

Hydrated acid and sulfates with formula \(A\ _XO_4\ _x(H_2O)\) are especially common in these caves. Thirteen species have been found as crust on the walls. Melanterite and rozenite are also part of geysermites, szomolnokite is part of stalactites (Fig. 11A, B, and C). Gypsum is a very abundant mineral in CMI, covering the north walls of the cave and often associated with some microbial mats. Other new cave minerals from this sub-group are pentahydrute and szomolnokite, both identified by Raman and XRD techniques. Chalcantite is another possible mineral but was identified only from one micro-Raman analysis; however, it matches 98% with the reference Raman spectrum (982, 609, and 455 cm\(^{-1}\)).

Retgersite is the only nickel sulfate identified by micro-Raman (984, 609 cm\(^{-1}\)) (Fig. 9) that belongs to...
the sub-group with the formula A\(\text{XO}_4\cdot x(\text{H}_2\text{O})\). Finally, voltaite displays black pyramidal shaped crystals under the SEM (Fig. 8D) and is the only member of Hydrated Acid and Sulfates with miscellaneous formula. Generally, it is present in the cave as crusts and crystals on the walls, floor, and roof. The presence of voltaite has been confirmed by XRD and micro-Raman techniques.

**Hydrated sulfates containing hydroxyl or halogen**

This group comprises minerals of the copiapite group, represented in CMI by aluminocopiapite, copiapite, ferricopiapite, and magnesiocopiapite. These minerals are common forming part of geysermites and normally show yellow colors, and sometimes a tabular habit (Fig. 10B). Ferricopiapite and magnesiocopiapite are new cave minerals confirmed by XRD.

**Other minerals (native elements, halides, oxides, and silicates)**

Native sulfur was detected in 11 samples from CMI by FT-Raman, micro-Raman, and XRD. Native sulfur deposits in this cave appear close to volcanic vents as crusts made of crystals <5 cm large, in high temperature areas, acid conditions (pH ~2) and sometimes forming stalactites in association with anhydrite (Fig. 7). The possible presence of the halide tolbachite is suggested by XRD as accessory mineral associated with römerite, gypsum, melaniterite, rozenite, and szomolnokite, detected in the same stalactite. This mineral was forming part of a light green stalactite (>8 cm length) in CMI. Tolbachite was not reported from cave environments yet.

The only oxide detected, goethite, was found in CMU as dark orange and reddish masses filling cracks in the walls. It was identified by XRD in only one sample. The only phyllosilicate identified is kaolinite, found in a single muddy yellow sample from CMU. It was associated with natrojarosite and minor gypsum in snottite-like speleothems hanging from the roof of the cave, probably related to extremophile microorganism activity (pH 1–2). Finally, marinellite was identified by micro-Raman spectrometry in two different samples, but it was not confirmed or detected by XRD.

**Mineral distribution and speleothems**

Mineral distribution in CMI is represented in Fig. 5, however due to the high diversity and possible spatiotemporal variable distribution of minerals (e.g., different temperature and relative humidity affecting the hydration state of some hydrous minerals), this study did not aimed at quantitatively estimating this spatial diversity. Other techniques, including photogrammetry or LIDAR scanning may be necessary.

Most of the reported sulfate minerals are very soluble, therefore, the only reason why they exist in the tropical weather of Costa Rica (rainy season from May to November, with 2,387 mm/yr) is because of the particular cave climate conditions. Most minerals in CMI occur as crusts on the wall and roof and are concentrated on the northern side and also covering walls and roof of the Sala de los Cristales Verdes (where most of the stalactites are located). Other speleothems such as stalactites and stalagmites are associated with \(\text{SO}_4^{2-}\) and Fe-rich infiltration waters (Fig. 11 and 4B), or Ca-rich solutions as a particular anhydrite-sulfur stalactite (Fig. 7). Geysermites are associated with \(\text{SO}_4^{2-}\) rising thermal waters, and present high-mineral diversity with more than 11 mineral species in a single sample. Most of the identified mineral species present yellow, white, and translucent color followed by green. The most common mineral habits are acicular followed by massive. Most of the crust minerals are centimeters in size, while stalactites and geysermites are normally decimeters.

**Elemental analyses and microphotography**

The results of 93 EDX analyses on samples from both caves allowed the detection of 17 chemical ions and their relative abundance (Supplementary Table 5). Presence of iron is related with abundant iron sulfate mineral phases (e.g., coquimbite, copiapite, ferricopiapite, rozenite). Other metals such as aluminum have been detected in several samples associated with sulfates containing Al (e.g., marinellite, alunogen, dietrichite, halotrichite,
pickeringite, aluminocopiapite). The presence of Si can be associated with the silicates (e.g., kaolinite or marinellite) and diatom skeletons found in the sample MU-170514-4 next to snottites (at pH ~2). Alkaline elements, such as calcium correspond to gypsum but can also be related to other minerals such as marinellite, anhydrite, and copiapite. The other alkaline cations detected by EDX (Na, Mg, K) are quite common in many samples. V, Br, and Sr were also reported but no mineral containing those elements has been identified in this work. This suggests that there could be more minerals present in the samples. SEM photographs of the samples show complex associations between these minerals and are useful to distinguish the different habits (Fig. 7, 8, 10, and 11).

DISCUSSION

Speleogenesis
The caves of the Irazú Volcano have a complex origin, which differs considerably from the speleogenetic mechanisms associated with normal carbonic and sulfuric acid caves worldwide (Hose et al., 2000b; Klimchouk, 2009; Onac et al., 2009). Ulloa et al. (2013) analyzed different possible mechanisms for the origin of CMI (i.e., landslide, phreatic explosion, or water runoff). These authors concluded that the landslide model with a stepped fault plane generated these caves. However, preliminary results of Baldoni et al. (pers. comm.) estimated a rock removal rate of ~3,000 m$^3$/yr in the Irazú-Turrialba system by chemical leaching from the springs. Also, based on an infrared survey in 1964, thermal anomalies were also identified (Gawarecki et al., 1980), which coincide with the Río Sucio fault traces and some of them were located close (~250 m NW) to the actual position of the caves (covered by the material that slipped in 1994). Therefore, the possibility exists that dissolution weathering and underground removal of volcanic pyroclastic rocks by very acid waters along active faults in the Irazú caves area may have caused the formation of the underground voids.

This pseudokarstic mechanism is evidenced by the presence of a cylindrical hole ~1 m in diameter in the roof of the cave, similar to holes formed in dissolution caves. Linked to this, in Llano Grande mine quarry (6.5 km SW of the Crater Activo), there are channels ~1 to 2 m wide in similar pyroclastic hydrothermally altered rocks, completely filled with native sulfur. Likewise, these presumed proto-channels were found during a drilling campaign, when a borehole intersected a possible hypogenic cave, ~1 to ~5 m depth. Therefore, a combination of tectonic, weathering, high temperature degasification, leaching, and erosion could be involved in the genesis of the caves on Irazú Volcano.

Sulfur in the caves
Sulfur is common in sulfuric acid caves [e.g., Cueva Villa de la Luz (Mexico; Hose et al., 2000); Carlsbad and Lechuguilla caves (New Mexico; Hill, 1995); Lower Kane Cave (Wyoming; Egemeier, 1981); Diana Cave (Romania; Onac et al., 2009); Santa Cesarea Terme (Italy; D’Angeli et al., 2017)] as wall crusts and stalactites, in places associated with gypsum (Hill & Forti, 1997). On Irazú Volcano, elemental sulfur has been reported in fumaroles (Stoiber & Rose, 1974) and as sulfur flow in the main crater in 1982 (McClelland et al., 1989). In the Irazú caves, elemental sulfur is present in some volcanic vents on the eastern wall of the cave (pH 1-2) and crystals can be up to 6 cm in length. Furthermore, elemental sulfur was found in a ~10 cm

Fig. 9. Micro-Raman spectra of the new cave minerals reported only by this technique.

Fig. 10. An example of geysermite (Photography taken by Victor Carvajal in 2011) in CMI (A); SEM microphotograph correspond to ferricopiapite (B and D); and pickeringite minerals (C and D).
long stalactite (Fig. 7A). Similar sulfur speleothems have been reported in strongly acid environments as a result of H2S oxidation (Forti & Mocchiutti, 2004; D’Angeli et al., 2017). Sulfur is present in all the EDX analyses (Supplementary Table 5), ranging from 5.2 to 70.8 wt%. The abundant presence of sulfur in these minerals, is related to the typically sulfur-rich volcanic environments. Sulfur is present in a variety of valence states; the most reduced form as hydrogen sulfide (H2S) in volcanic gases is -2, elemental sulfur (valence 0) and the most oxidized form as sulfate (SO4^2-) is +6. Some of the changes in valence are attributed to the biogeochemical reactions, in which microorganisms frequently play a main role as catalysts, taking advantage of the associated electron fluxes to satisfy their metabolic needs (Morse et al., 1987).

**Minerogenetic mechanisms in the Irazú caves**

The transport of ions and chemical compounds in the caves and the consequent mineral precipitation can occur from mineralized solutions and this may happen both in subaerial or subaqueous settings. Each of the transport mechanisms can trigger different minerogenetic processes. Mineralized solutions play a very important role in the transportation of chemical elements and compounds into the caves. Water geochemistry analyses indicate considerable concentrations of sulfates in the main crater lake (Supplementary Table 1), and infiltration water analyses indicated the presence of Se, Fe, Al, Ca, Mg, Na, Mn, K, Zn, Ni, and Co ions (Fig. 3C). Selenium is the only element that is not reflected in the elemental composition of the reported minerals. Enrichment in selenium (993.1 mg/l) might be associated with the input of geothermal waters (Floor & Román-Ross, 2012). Iron concentration is 440.2 ppm in the infiltration waters and is present in the chemical composition of 20 minerals, followed by calcium present in 14 different minerals. Stalactites and stalagmites (usually green to blue in color) are associated with dissolution-precipitation processes of Fe-sulfate-rich infiltration waters. They are found in “Sala de los Cristales Verdes”, formed by iron hydrated sulfates including melanterite, szomolnokite, römerite, rozenite, and volaita with presence of gypsum (probably in lower quantities) and possible tolbachite. In the case of iron sulfate minerals as szomolnokite-rozenite-melanterite-römerite, they differ from each other by their content in water molecules, with hydration and dehydration processes possibly causing their transformation from one species to another. The growth of the blue-green stalactite (Fig. 4B) is relatively fast: based on field observations (growth and collapse of green-stalactites) they can grow in the order of ~30 cm/a. Similar melanterite stalactites and stalagmites have been reported in caves in California, Nevada, and Tennessee (USA), Sicily (Italy), and mine caves (Hill & Forti, 1997).

At least 7 different geysermites have been identified in CMI distributed in the main chamber and ranging from ~15 to ~40 cm height and ~8 to ~15 cm width (Fig. 5 and 10). Field observations revealed no thermal anomalies in these speleothems suggesting that the process that formed them, is not currently active. The minerogologival composition of the analyzed geysermites consists of hydrated acid sulfates (apjohnite, coquimbite, dietrichite, halotrichite, kalinite, pickeringite, and wupatiikite) and hydrated sulfates containing hydroxyl or halogen (aluminocopiapite, copiapite, ferricopiapite, magnesiocopiapite). The genesis of these geysermites is related to upwelling sulfate-rich waters of the Irazú hydrothermal system. The cooling of the rising thermo-mineral fluids induced supersaturation and mineral precipitation (Onac & Forti, 2011b) and may explain the high mineralogical diversity of such speleothems. The presence of geysermites suggests the existence of lower thermal aquifers which can be responsible of acid drainage in the Río Sucio River basin (Arce-Rodriguez et al., 2017) and some thermal springs in the surroundings.

Aerosols (solid or colloidal particles, 10^7 to 10^2 cm) and hydrosperosols (liquid droplets) are another way to transport chemical compounds, which is a feasible mechanism where there is a mixture of air at different temperatures in caves (Cigna & Hill, 1997). Speleothems associated with aerosols are related to thermal caves (preferable associated with fumarole vapors), where temperature and humidity gradients are present (Onac & Forti, 2011b). Speleothems associated with aerosols as transport-media of chemical compounds is strongly suggested for CMI because: (1) presence of minerals close to hydrothermal vents, higher density and radial distributions around them; (2) elevated concentrations of SO4^2- in fumarolic analyses (Table 1) and presence of H2S in the cave atmosphere (29 ppm) being evidence of sulfur-compounds in the air; (3) some minerals present morphologies similar to “rims and blisters” (according to the description of Onac & Forti, 2011b), which are associated with speleothems developed by aerosols and vapors. CMU did not present considerable thermal anomalies and shows less mineral diversity, which could be suggesting that aerosols are probably one of the most...
active mechanisms in the higher mineral diversity of CMI.

The presence of elemental sulfur crystals in CMI could be associated with cooling down of fumarole gases (sublimation at temperatures >100°C), which is suggested as minerogenetic mechanism (Forti, 2005; Onac & Forti, 2011b). The atmosphere of CMI is rather complex, and shows significant variations in temperature (9 to 30°C) and relative humidity (74.2 to 92.4%). These variations can influence precipitation of aerosol minerals and also rule the hydration-dehydration processes for those mineral species that only differ in their water content (e.g., aplowite-bieberite, epsomite-hexahydrite and melanterite-rozenite-szomolnokite series).

Finally, another possible mineral-forming mechanism in the cave is microbial activity. This mechanism is suggested based on field observations where gypsum-stalactite are intrinsically related to snottites in some parts of the cave.

Relative abundance of elements and minerals

CMI may be the cave (or one of the caves) in the world with the greatest mineralogical diversity. Because of the particular conditions in an active volcano environment (SO₄²⁻ rich waters and influence of volcanic gases), 48 minerals species have been identified, 41 are confirmed and 7 mineral species are suggested. According to Onac & Forti (2011a) in the past 20 years, on average, three new cave minerals per year have been discovered. This study adds 18 minerals to the list Onac & Forti (2011a), including 12 confirmed sulfates (aplowite, bieberite, boyleite, dietrichite, ferrinatrite, ferricopiapite, lausenite, lishizhenite, pentahydrite, magnesiocopiapatite, szomolnokite, and wupatkiite), one confirmed silicate (marinellite), four other possible sulfates (cyanochroite, merrallite, retgersite, and rhomboclase, Fig. 9), and also the possible presence of an halide (tolbachite).

Five of these new cave minerals (aplowite, ferrinatrite, magnesiocopiapatite, wupatkiite, and boyleite) in CMI were previously reported by Ulloa et al. (2013) and apjohnite is not in the cave minerals list of Onac & Forti (2011a), but was reported by Onac et al. (2009). However, it must be considered that the number of minerals present in these caves is still underestimated because it was not possible to associate some of the Raman spectra with any specific mineral species of the Raman database, as well as the presence of more elements (e.g., Br, Sr, V) identified by EDX analyses (Supplementary Table 5) which are not reflected in the composition of the minerals listed in Table 2.

Apart from selenium, the majority of elements in infiltration waters (Fig. 3C) corresponds to the most abundant elements present in the identified sulfates (Fig. 6D). Fe, Al, Ca, Na, and Mg are the most abundant cation species in sulfates and also the major elements in infiltration waters. Thus, the abundance of accessory elements in sulfate minerals is proportional to the abundance of elements in the infiltration waters.

Gypsum is probably the most abundant mineral in the cave. The north walls of CMI are totally covered with gypsum crystals, mainly along the external edge of the cave, and this mineral is also present in CMU. Gypsum tends to be idiomorphic and some of the crystals presents an interesting association with microbial mats.

Summit volcanic crater lake and hydrothermal system

The last eruptive activity of the main Irazú crater stopped after 1965 and in July of the same year the volcanic lake was formed in the main crater. The last eruptive activity of the main Irazú crater stopped after 1965 and in July of the same year the volcanic lake was formed in the main crater. This lake has an ephemeral behavior, being repeatedly dry (years 1977, 1979, 1982-83, 1987, 1990, and 2013-17) and precipitation has not been the main factor in its change in volume. Fumaroles and sub-aquatic fumaroles have also been reported at different time-intervals, during 1973, 1977-1978, 1982, 1985-1986, 1991-2000, and 2002-2008 (Ramírez et al., 2013 and references therein). The lake of the main crater showed important changes in pH, temperature, chemical composition (Fig. 3A and B; Supplementary Table 1) and water level since 1991, making this lake a complex geochemical system. An acidification trend in the lake was measured in the years 1991-1993 and was associated with enrichment in SO₄²⁻ and Cl⁻. After 2000, lower concentrations of SO₄²⁻ and Cl⁻ were detected and an alkaline trend predominated. The average temperature of the lake is 29.5°C, although it showed temperatures over 50°C in 1991 and 2003.

These variations in water levels and volcanic gas emissions are strongly associated with the active faulting and secondary permeability in the volcanic conduit. Based on historical reports it is suggested that rising of hot fluids and volcanic gases from the hydrothermal system has changed over time, sometimes reaching the volcanic crater-lake level. Hydraulic communication between the main crater lake, and the caves and hydrothermal system is suggested. The main supporting signs are: the presence of diatom skeletons in a sample in a dripping zone at CMU, which may come from lake; the enrichment in sulfates in historical water analyses of the main crater lake associated with infiltration waters in the caves and presence of sulfate minerals in the cave. The hydraulic connection lake-caves coinciding with the model proposed by Ramírez et al. (2013) is associated with high-permeability pathways within the faults. According to this hypothesis, variations of regional tectonic forces, seismic activity, clays on cracks, and raining may be factors controlling the ephemeral behavior of Irazú Volcanic Lake in the Crater Principal.

For periods for which water geochemical analyses are available (Supplementary Table 1) in Irazú Volcanic Lake, evidence of sub-aquatic fumarole activity is reported for years 1991, 1992, and 2003 to 2008 (Ramírez et al., 2013). These ephemeral volcanic emissions are connected with different pulses of the hydrothermal system activation and probably directly linked with variations in the geochemical lake system. According to this model, cave inputs of sulfate-rich
waters can present two origins, one related with volcanic lake infiltration, which was presently active during field trips. A second sulfate-rich water source is suggested, related with the rising of geothermal fluids through faults, which can provide permeable paths. Infiltration sulfate-rich solutions are responsible for speleothem precipitation, such as stalactites and stalagmites, whereas geothermal rising sulfate-rich waters probably caused the formation of geysermites.

**Fumaroles and diffuse gas at Irazú System**

Sulfur and gypsum minerals have been reported as fumarole minerals in the volcano (Stoiber & Rose, 1974). Fumarole locations have changed over the last decades, but actually at least two still active fumaroles are present in the northwest part of Irazú Volcano (Fig. 1A). In the area affected by the 1994 landslide another fumarole was present, maybe associated with the actual location of the caves. The only published data from Irazú fumaroles shows concentrations of CO$_2$ from 98.0% to 99.8%, SO$_2$ from 0.106 % to 0.007 % and H$_2$S from 0.015 % to 0 % in 1981 and 1982, respectively (Global Volcanism Program, 1983) and CO$_2$ 98.9% and H$_2$S 0.8% in the year 1993 (Global Volcanism Program, 1993). The compiled historical analyses of Irazú fumaroles (Table 1) shows an important content of SO$_2^-$ ranging from 42 to 503 mg/l, being SO$_2^-$ the most abundant compound, making aerosols mineralogenesis a potential mechanism.

Irazú Volcano also presents an important diffuse CO$_2$ flux: Galindo et al. (2004) established a CO$_2$ base flux at ~20 gm$^{-2}$d$^{-1}$ and maximum values of 316 g m$^{-2}$d$^{-1}$ for sites studied in 2001, while Epiard et al. (2017) reported maximum values of 2.71 x 10$^4$ gm$^{-2}$d$^{-1}$ for the northern flank and 2.17 x 10$^3$ gm$^{-2}$d$^{-1}$ north of the main crater in 2015. Those diffuse CO$_2$ flux anomalies are located in areas close to active fault traces, being the structural control very important for the escape of gases and associated rock alteration, but probably with less relevance for the formation of speleothems in the caves of the Irazú Volcano.

**Implications for planetary exploration**

Caves are considered extreme environments for life because they are resource-limited due to the absence of light and implicitly of photosynthesis; however, chemolithoautotrophic organisms can metabolize inorganicelements such as iron, sulfur, and manganese in caves (Northup et al., 2011, 2012; Riquelme & Northup, 2013). Thus, terrestrial subterranean environments and secondary cave minerals (speleothems) have drawn attention as potential analogues of life niches for planetary exploration, especially Mars (Boston et al., 2001; Bost et al., 2013; Gázquez et al., 2014). During the past decades important efforts have been done by the scientific community to understand primary composition of Mars and alteration products that are associated with paleo-environmental conditions of the planet. Chemical analyses suggest than the Martian crust is basaltic, where mineralogy is dominated by primary silicates (e.g., olivine, clinopyroxene, orthopyroxene, plagioclase, alkalifeldspars) and secondary minerals groups (e.g., oxides, clay minerals, hydrated silicates, carbonates, sulfates, and chlorides). These secondary minerals are associated with alteration related to presence of liquid water in some stage of the Martian evolution (Mustard et al., 2008; Ehlmann & Edwards, 2014). The discovery on the Martian surface of soil sulfur-enrichment (Chevrier & Mathé, 2007), presence of jarosite suggestive of existence of acidic waters (pH < 4) (Ehlmann & Edwards, 2014; Thollot et al., 2012), Ca, Fe, and Mg-sulfates (Ehlmann & Edwards, 2014) and CO$_2$-rich atmosphere, are factors which coincide with the mineralogy and some physical data presented in this article for the Irazú volcanic caves. Therefore, we consider that our investigated caves can be used as analogue for the Mars environment. Future geomicrobiological analyses on these caves could give us clues in understanding the potential life-forms and geo-environmental interactions from an astrobiological perspective.

**CONCLUSIONS**

The northwest flank of the Irazú Volcano accommodates three volcanic caves, of which CMI and CMU have been studied in detail. The speleogenetic process that generated these caves is complex and involved tectonic (fault zone), weathering, high temperature degasification, leaching and erosion. The caves are developed in interbedded pyroclastic rocks, affected by hydrothermal alteration.

The studied caves show an extremely high mineralogical diversity, with forty-eight identified mineral species. The secondary mineral assembles of the studied caves are constituted mainly by sulfates, most of them belonging to hydrated acid and sulfates sub-group. Other mineral groups are also present, including native elements and minor halides, oxides and silicates. Thirteen of the mineral species are confirmed as new cave minerals, whereas five others are not confirmed yet by various techniques.

The particular environmental conditions in the studied caves, including location in an active volcano, active faulting, presence of thermal anomalies, interaction of sulfate-rich and acid water from volcanic lake infiltration, and probably rising of geothermal waters somewhere in the history of the caves, are the main factors that control this extreme mineralogical diversity.

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