

# The acid-sulfate zone and the mineral alteration styles of the Roman Puteolis (Neapolitan area, Italy): clues on fluid fracturing progression at the Campi Flegrei volcano.

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**Abstract.** Active fumarolic solfataric zones represent important structures of dormant volcanoes, but unlike emitted fluids, their mineralization are omitted in the usual monitoring activity. This is the case of the Campi Flegrei caldera in Italy, among the most hazardous and best-monitored explosive volcanoes in the world, where the landscape of Puteolis is characterized by an acid sulfate alteration that is active at least since Roman time. This paper provides temperature, mineralogical, textural, compositional and stable isotope data for those solfataric terrains sampled at the crater and Pisciarelli slope of the Solfatara volcano between 2013 and 2019. Temperatures vary between 40 ° and 95 °C. Minerals include alunite with grain sizes generally larger than 20 µm, alunogen, native sulfur, well-ordered kaolinite, and, common at Pisciarelli, pyrite, illite and NH<sub>4</sub>-sulfates. Sulfate terrains have higher contents of Ti, Ba, Au, As, Hg and Tl relative to their parent substrate. The Pisciarelli slope is anomalous in terms of the presence of NH<sub>4</sub>. δ<sup>34</sup>S values for sulfides and native S range between -3.00 and 0.49 ‰ and from -4.42 to 0.80 ‰, respectively. Sulfates show δ<sup>34</sup>S and δ<sup>18</sup>O values in the range of -3.35 to 3.80 ‰ and between 0.3 and 31.33 ‰, respectively. The style of mineralization and the stable isotope geochemistry do produce complex and not completely consistent classifications and genetic constraints. We merge our data with volcanological information, data from exploration drillings and geophysical results. With the conceptual model, we suggest a series of shallow and deep aquifers interconnected like “communicating vessels” through a main fault system that downthrows Solfatara with respect to Pisciarelli. Fluid outflow from the different discrete aquifers hosted in sediments – and possibly bearing organic imprints – is the main dataset that allows determination of the steam-heated environment with a supergene settings superimposed. Supergene conditions and high-sulfidation relicts, together with the narrow sulfate alteration zone buried under the youngest volcanic deposits, point to the existence of an evolving paleo-conduit. The data will contribute to monitor and evaluate the volcanic hazards.

## 1 Introduction

Active solfataric landscapes are among the most peculiar and fascinating environments on the Earth that may be considered as planetary analogues (e.g., White and Hedenquist, 1990; Rye et al., 1992; Lowe et al. 1993; Zillig et al., 1996; Ciniglia et al., 2005; Rye, 2005; Glamoclija et al., 2004; Sgavetti et al., 2008). Their peculiarity arises from the stringent interaction between inorganic (mineral assemblages and geochemistry) and organic (biota) substances under extreme ambient conditions (pH, temperature, salinity, oxygen deficiency, etc.) associated with endogenous degassing (i.e., H<sub>2</sub>O, CO<sub>2</sub>, CH<sub>4</sub>, H<sub>2</sub>S, SO<sub>2</sub>, HCl, HF, etc.) and hot water fluid circulation (hydrothermal/geothermal systems) on dormant volcanoes. They allow investigating a variety of processes in the field of geology (i.e., magma and volcano dynamics), biology (i.e., physiological adaptation to environmental stresses and the origin of the life), medicine, astrology and archaeology (i.e., thermal bath and antibacterial applications), with possible future medical and biotechnological applications.

The Solfatara volcano (Campi Flegrei, CF, Italy; Fig. 1a) is perhaps the most famous and hazardous geothermal solfataric setting in the world (e.g., Rittmann, 1950; Rosi and Sbrana, 1987; De Vivo et al., 1989; Barberi et al., 1991; Piochi et al., 2014)

with exploration since Greek times up to Medieval age (e.g., Photos-Jones et al., 2016). The generation of new minerals (herein after referred to as neogenesis) received limited discussion in the recent literature (Cortecchi et al., 1978; Valentino et al., 1999; Piochi et al., 2015; Russo et al., 2017). By contrasts several studies relate to bradyseism phenomena that concern with seismicity, ground deformation and outgassing (e.g., Corrado et al., 1976; Barberi et al., 1984; Chiodini et al., 2016; Cardellini et al., 2017; Moretti et al., 2017), life in these environments (e.g., Zillig et al., 1996; Glamoclija et al., 2004; Sgavetti et al., 2008), and a continuous interest into the thermal bath and medical care (e.g., Photos-Jones et al., 2016; Giacomelli and Scandone, 2012).

This paper focuses on the solfataric mineral assemblages updating our previous research (Piochi et al., 2015) and presenting the result of our progressing work on the CF solfataric volcano. Results derive from temperatures determination contextually to sampling, investigations by Optical Microscope (OM), X-Ray Powder Diffraction (XRDP), Electron Microscopy (EDS-BSEM), Diffuse Fourier Infrared Spectroscopy (DRIFT-FTIR), whole-rock geochemistry (WRG) and stable isotope geochemistry (SIG) of sulphur and oxygen. By merging new and published information (Celico, 1986; Guglielminetti, 1987; Rosi and Sbrana, 1987; Chiodini et al., 1988; Celico et al., 1992; Aiuppa et al., 2006; Caliro et al., 2007; Piochi et al., 2014; Di Giuseppe et al., 2017; Moretti et al., 2017), we reflect on the significance of the sulfate alteration zone and related volcanological implications.

## 2 Background

### 2.1 Geological setting

The Solfatara volcano (Fig. 1a,b,c) displays impressive and powerful hydrothermal activities with hot fumaroles, thermal springs, mud pools and diffuse degassing (Allard et al., 1991; Valentino et al., 1999; Chiodini et al., 2001; Valentino and Stanzone, 2003; 2004; Chiodini et al., 2010; Piochi et al., 2015; Cardellini et al., 2017; and references therein). The hydrothermalism intensely altered the faulted volcano slopes (Rosi and Sbrana, 1987) and the solfataric landscapes (Fig. 1a-e) have locally replaced the original pyroclastic sequences (e.g., Agnano Monte Spina, Astroni and Solfatara tephra) and lavas (Monte Olibano, Solfatara cryptodome) younger than 5 ka (e.g., Di Vito et al., 1999; Piochi et al., 2005).

The study area is located at Puteolis, the area of maximum ground uplift (in excess of 3 m) and seismicity (more than 16,000 low-magnitude earthquakes), activated during the unrest episodes in 1970 ÷ 1972 (Corrado et al., 1976) and in 1982 ÷ 1984, namely “bradyseisms” (Barberi et al., 1984), that is slowly on-going (e.g., Bodnar et al., 2007; Chiodini et al., 2016; Moretti et al., 2017).

The solfataric area has been exploited for centuries for its alum occurrences (Photos-Jones et al., 2016 and references therein). The intense mining in Roman and Medieval times modified their original context (Photos-Jones et al., 2016): the Pisciarelli gorge-valley was a quarry, while caving activity exposed the eastern (the Monte Olibano inner wall) and northern flanks of the Solfatara volcano, and rework deposits in the crater floor.

Old pictures and descriptions (Sicardi, 1959) suggest that the most evident manifestations along the SE and NE rim remain roughly the same: 1) the main Bocca Grande fumarole (Fig. 1d) with various exhalative branches northward along the morphological heights; 2) the minor fumarolic vents around the old thermal baths (Sst site; Fig. 1d) and 3) the mud pools (Fig. 1b,c,d,e). Also, the thermal spring in Pisciarelli (Fig. 1a,b,e), known as the “Bulla”, i.e., the bubbling one, was known at least since Medieval periods (Photos-Jones et al., 2016). Moreover, the same descriptions indicate the presence of a lake in the Agnano Plain (Fig. 1a). According to Ventriglia (1942), the lake extended up to the slope base of the Solfatara volcano and had a maximum depth of 15 m; drillings recovered related sediments (de Vita et al., 1999). Ventriglia (1942) also indicated high temperatures in the lake preventing fish from living. Today, the area shows several mud pools and thermal springs, while some (“de Pisis” and “Sprudel” springs in the Terme of Agnano; Fig. 1a) disappeared, although high temperatures can be still detected.

At present, ground waters nearby Solfatara are rich in  $\text{Cl}^-$  and  $\text{SO}_4^-$  (Aiuppa et al., 2006). Temperatures at the fumaroles exceed 160 °C (e.g., Cardellini et al., 2017; Gresse et al., 2017), in agreement (or locally lower respect) with measurements reported by Sicardi (1959) in the range of 141-215 °C at Bocca Grande and of 99-110 °C at other sites, between 1921 and 1951. Pisciarelli waters exhibited temperatures mostly around 95 °C with minimum values of 84 °C between 1978 and 1999 (Celico, 5 1992; Valentino and Stanzione, 2004). A geyser-like vent at Pisciarelli has a temperature of up to 116 °C (Chiodini et al., 2016). Only, the mud pool (“La Fangaia”; Fig. 1d) was hotter in the past with values up to 100 °C (Sicardi, 1959). This author describes a mud vent in the southeastern area of the crater that is no longer present.

Emitted gases include  $\text{H}_2\text{S}$ ,  $\text{CH}_4$ ,  $\text{N}_2$ ,  $\text{H}_2$ ,  $\text{CO}$ , in addition to the dominant water vapour and the secondary abundance of  $\text{CO}_2$  with reaching a flux of at least 1500 tonnes/day and a maximum value of 3000 tonnes/day (Allard et al., 1991; Aiuppa et al., 10 2013; Chiodini et al., 2016). Previous studies further report a Hg flux between 0.9 and 4.5 g/day (Ferrara et al. 1994; Bagnato et al., 2014), a detectable (very low) abundance of gaseous  $\text{SO}_2$  (Ferrara et al. 1994; Aiuppa et al., 2013), the occurrence of light hydrocarbons (Capaccioni and Mangani, 2001), and the presence of As and Hg (17 - 5200  $\mu\text{l}$  and 1 - 30  $\mu\text{l}$ , respectively; Valentino and Stanzione, 2003) in the pools and waters. High  $\text{NH}_4$  concentrations are described for waters at Pisciarelli (Martini et al., 1991; Celico et al., 1992; Valentino and Stanzione, 2003), but understanding of the nitrogen source and cycling 15 at the Phlegraean area, showing a unique isotopic composition with  $\delta^{15}\text{N}$  at  $6.3 \pm 0.3 \%$  (Chiodini et al., 2010), remains elusive.  $\delta^{18}\text{O}$  and  $\delta^2\text{H}$  of emitted fluids are in the range -5 to -0.5 ‰ and between -30 and -20 ‰, respectively (Caliro et al., 2007). The average  $\delta^{34}\text{S}$  value of  $\text{H}_2\text{S}$  gases is  $-0.3 \pm 0.3\text{‰}$  (Allard et al., 1991).  $\delta^{34}\text{S}$  values determined for shallow subsurface sulphur-bearing minerals range between -5.5 to 0.0‰, while the deep-seated pyrite has values from 3.3 to 7.4‰ (Piochi et al., 2015).  $\delta^{18}\text{O}$  values for alunite are from 4.2 to 7.0‰ (Piochi et al., 2015). The pH of water pools and soils is neutral to acid, with 20 values <2 around the pools (Valentino and Stanzione, 2003; 2004; Gresse et al., 2017).

## 2.2 Sampling, sample preparation and analytical techniques

Sampling was conducted within the Solfatara crater and in the Pisciarelli and Cinofilo areas (Fig. 1a,d,e) with additional sites compared to Piochi et al. (2015); the crater floor, except the pool, was intentionally avoided because the reworking in historical 25 time (Photos-Jones et al., 2016) and the possible anthropogenic contamination. This study intends to enlarge the dataset on the acid sulfate alteration zone of the Phlegraean area, in order to understand the quiescent dynamics of the volcano. Similar observations and data are also available for Ischia island (Piochi et al., 2019) that belongs to the Phlegraean Volcanic District (Piochi et al., 2005).

Our new collection is, therefore, widening the observation period on the Puteolis sulfate area that now spans between January 30 2013 and April 2019 (Table S1). Selection of sampling sites (herein after referred using the acronyms in Fig. 1d,e) is based on variable macroscopic features including degassing “magnitude”, tectonics and fracturing evidences, mineral occurrences and exhalative vents locations, as visible in the field and described in the literature (Allard et al., 1991; Ferrara et al. 1994; Valentino and Stanzione, 2003; Aiuppa et al., 2013; Bagnato et al., 2014; Chiodini et al., 2016). A thermo-couple digital probe 51/52 II by Fluke with precision of  $\pm 0.3 \text{ °C}$  was used to measure temperature on the field, contextually to sample collection. 35 Samples were air-dried for several days to one week. Subsequently, these were studied under the Optical Microscope (OM) in order to define their general mineral assemblages. Where possible, the various S-bearing phases (or enriched portions) were handpicked for subsequent isotopic analyses. Figures 2 and 3 show the appearance of most representative samples.

Both bulk-rocks and separated phases were crush in an agate mortar for X-Ray Powder Diffraction (XRDP), Diffuse Fourier Infrared Spectroscopy (DRIFT-FT-IR) and Whole-Rock Geochemistry (WRG). Scansion Electron Microscope (BSEM), 40 Electron Diffuse System (EDS) and Stable Isotope Geochemistry (SIG) used aliquots of bulk materials and isolated mineral phases. Appendix A reports details for analytical techniques. Details on XRDP and DRIT-FT-IR are in the Supplement (S1) together with representative spectra (Fig. S1, S2).

### 3 Results

#### 3.1 Mineral data by OM, XRDP, BSEM, DRIFT-FT-IR results

New and previously published (Piochi et al., 2015) mineralogical data for the Solfatara-Pisciarelli area (Table S1, Table 1) provide information on a yearly to monthly basis since 2013 along with measurements of temperature. The mineral assemblage dataset derives from XRDP analyses (Supplement) corroborated by textural and chemical information obtained at the EDS-BSEM. DRIFT-FT-IR spectra determined on representative samples, display characteristic bands of minerals they include (see below and Supplement), and help in material characterization.

Through time, K, Al sulphates (alunite) and native S (Fig. 3a,b,c,d,e,f) are the main and widely distributed secondary mineral phases associated to surface degassing. Alunogen and pyrite (Fig. 3a,b) are second in abundance. All these mineral phases can form single phase concretion or coexist in up to mm-sized grains. Alunogen often – if not generally – associates with alunite and occurs in two distinct morphologies (Fig. 3a,d,e). Most commonly, it consists of fibrous tangled masses of white crystals. Where coexisting, alunogen fibres grow from the edges of alunite crystals (e.g., sample L100 zucc in Table 1; Fig. 3d). This appearance seems usual along the fault scarp, north of the pool (L1 site, Fig. 1e). Secondly, alunogen has thin, platy crystal habits (Fig. 3b,d,e). Many of these crystal groups show rounded to corroded edges suggesting alteration after crystallization (Fig. 3e). Dendritic and/or sometimes bi-pyramidal crystallites (Figs 2a,3a,3c) are ubiquitous habits for native sulfur (typically sampled at L1, SMO, some places along ASA, Sst in Fig. 1d and Table 1) that mostly cluster within the alunitic surface and the rock voids (Fig. 2c). Along the fracture, S may form a yellow ductile patina (L1 vent, BG, BN in Fig. 1d,e and Table 1). Locally (PINT, PEXT, L19, L20, L60 in Fig. 1e and Table 1), S produces encrustations with pale yellowish fibrous-like texture (Fig. 2b).

Pyrite (Fig. 2d) occurs as smaller ( $\leq$ mm-sized) rhombododecahedral grains (Fig. 3a) or as fine-grained mineral uniformly coating other components (i.e., feldspars, lava, etc.; Figs 2d,3i). It has a particular high abundance within the Pisciarelli muds (Figs 1e,2d), i.e., both within the main pool where it could reach cm-sizes and in the geyser (G site in Fig. 1e) emission. The blackish colour of the muds should also derive from the pyrite abundance. Pyrite also occurs around the degassing areas. Barite can be further detected at Pisciarelli.

Clays have a low relative abundance in the studied samples (Supplement). They are mostly smectite, and most common are illite/montmorillonite, while kaolinite seems least abundant (Table S1, Table 1), as derived by the XRDP spectra (see Fig. S1c,d,e) and supported by DRIFT-FT-IR study (see below; Fig. S2 and Supplement). In particular, the infrared technique is suitable to detect the kaolinite and the related bands in the OH-region, in agreement with Madejová et al. (2002). Illite/montmorillonite usually occurs in the muds at Pisciarelli (from geyser and around other emissive vents) and occasionally at Solfatara (Table 1, S1). Kaolinite characterizes the newly formed pool within the Solfatara crater and occurs locally at Pisciarelli (Fig. 1c,d,e and Table 1). Figure 4 illustrates the platy particles of kaolinite with typical widths of  $< 10 \mu\text{m}$  that assembly in packages and associated with alunite crystals.

Other efflorescent phases (Fig. 3c,i) occur randomly. Rarely, Al, Fe sulphates (halotrichite) have been identified nearby the Pisciarelli geyser as crust-like aggregates. Na and  $\text{NH}_4$  sulphates induce the pale orange painting on efflorescences and encrustations, and generally of soils. Sulphates, bearing Na, Ca, Mg are least common, and represent a typical occurrence in the new Solfatara mud. Alum has been detected at Pisciarelli.

Air-dried evaporation of water sampled at the Pisciarelli pool resulted in the precipitation of mascagnite, tschermigite, and letovicite (Fig. 2e,f, S1a and Table S1, Table 1). Figure 3i shows the euhedral tschermigite that coexists with native S in the sample L30 eff-blocchetto (Table S1, Table 1). Instead, evaporation of Solfatara mud pool water produced alum, as documented already in the Medieval and Roman times (Photo-Jones et al., 2016). Water from the Stufe di Nerone (west side not shown in figure) crystallized halite.

Realgar (detected at the EDS-BSEM and not listed in Table S1) and ammonium chloride (Fig. 3g,h) appears as peculiar precipitates at the Bocca Grande and Bocca Nuova sites (Fig. 1d).

Accessory minerals include hematite, quartz, and, possibly, Fe-hydroxides and phlogopite.

Furthermore, amorphous phases are largely present at various sites (Table S1, Table 1), particularly, in muds and in the samples from Bocca Grande and L1 vent (Fig. 1d,e). The widespread amorphous phases could correspond to material from the both original volcanic rock and alteration. General assumptions (Piochi et al., 2015; Montanaro et al., 2017) indicate amorphous silica, although it merits a more rigorous examination.

Finally, Fe-oxide, and fresh to variably altered feldspar and biotite are the most common primary volcanic mineral phases.

DRIFT-FTIR spectra collected on selected samples (Fig. S2) produce data consistent with XRDP results (Fig. S1) and furthermore allow useful details on structure and eventual minor phases or impurities (Supplement). Table S2 lists the relevant vibration modes of spectra and the proposed mineral assignments. The crystals formed by evaporation of water in the Pisciarelli pool (Fig. S2a; Supplement) show a sharp band at 1422-1411  $\text{cm}^{-1}$  that is in the region of the  $\nu_4(\text{NH}_4^+)$ , as described in the literature (e.g., Weis and Ewing, 1996; Parafiniuk and Kruszewky, 2010) and in agreement with XRDP mineral data (Table S1; Fig S1).

Native S from two different samples (PINT S tozzo 18/10/17 and PINT S 18/1/18 in Table 1; Fig. S2b) is evident in the DRIFT-FTIR spectra at  $< 2950 \text{ cm}^{-1}$ , with the strongest bands at 843 and 468  $\text{cm}^{-1}$  that coincide with those of sulfur in the both the USGS (see Sulfur GDS94; Clark et al., 2007) and RRUFF (<http://rruff.info/>) databases. The spectra differ in the OH stretching region, likely indicating the occurrence of impurities, although water absorption by the KBr solution can be a further possibility.

As expected (Clark et al., 1990), alunite can be determined through its major band at 3483  $\text{cm}^{-1}$  coupled with a smaller one at 3513  $\text{cm}^{-1}$  in the OH-stretching region (Fig. S2c,d,e). Accordingly, the spectra show a small band at ca. 4605  $\text{cm}^{-1}$  from the Al-OH bend also ascribed to alunite. Furthermore, it is possible recognizing the doublet mode at 1088  $\div$  1099  $\text{cm}^{-1}$  and at 1028  $\div$  1025  $\text{cm}^{-1}$  (Toumi and Tlili, 2008) and the mode at 3971  $\div$  3978  $\text{cm}^{-1}$  (see USGS database; Clark et al., 2007) from alunite. However, in the OH-stretching region there are some other vibrations. Based on Madejová and Komadel (2001), illite is likely producing the vibration at 3622  $\div$  3629  $\text{cm}^{-1}$ .

Notably, the DRIFT-FTIR spectra of muds from Pisciarelli (Fig. S2d) show a vibration in the region of 1430  $\text{cm}^{-1}$ . Because the muds were separated from water, as before, the band can be related to the  $\text{NH}_4$  (Weis and Ewing, 1996; Parafiniuk and Kruszewski, 2010) in tschermigite, mascagnite and letovicite (Table S1, Fig. S2, Fig. S1, Supplement).

The new pool at Solfatara characterizes for peculiar DRIFT-FTIR spectra in the OH-stretching region (Fig. S2e; note the inset) due the presence of kaolinite, in addition to alunite, and minor (or occasional) S, feldspar, pyrite and amorphous phases. Specifically, these are i) alunite (Clark et al., 1990) with a major band at 3483  $\text{cm}^{-1}$  coupled with a smaller one at 3510  $\text{cm}^{-1}$ , and ii) kaolinite (Madejova, 2003) with two minor bands at 3667 and 3651  $\text{cm}^{-1}$  between two major vibrational modes at 3695 and 3620  $\text{cm}^{-1}$ . Accordingly, the spectra show vibrations at ca. 4605  $\text{cm}^{-1}$  ascribed to the Al-OH stretching contribution in alunite; furthermore, it is possible recognizing the OH-deformation at ca. 915 and 938  $\text{cm}^{-1}$ , the Si-O stretch at 1008 and 1026  $\text{cm}^{-1}$ , and the Al-OH bending contribution at 4523  $\text{cm}^{-1}$  from kaolinite.

The four vibration modes of kaolinite in Fig. S2e point for a well-ordered mineral structure (Madejová, 2003; Fitos et al., 2015), giving a strong support to the XRDP results (Fig. S1e, Supplement), also in multiphase samples (Madejová, et al., 2002).

The IR spectra of samples for which XRDP confirmed illite/montmorillonite (Fig. S1, Supplement) lack of overtones at 4255 and 4081  $\text{cm}^{-1}$ . Following the literature (Clark et al., 1990; Madejová; 2003), these overtones are characteristics of illite and their absence can support the presence of montmorillonite.

### 3.2 Stable isotopes of sulfur and oxygen (SGI)

A new set of  $\delta^{34}\text{S}$  and  $\delta^{18}\text{O}$  values for sulfur-bearing minerals is listed in Table 2.  $\delta^{34}\text{S}$  values of sulfides and native S range between -3.00 and 0.49 ‰ and between -4.42 and 0.80 ‰, respectively. Sulfates are characterized by  $\delta^{34}\text{S}$  and  $\delta^{18}\text{O}$  values ranging from -3.35 to 3.80 and from 0.3 to 31.33‰, respectively. Temporal variations in  $\delta^{34}\text{S}$  for different sulphur-bearing phases at the different locations, reveal dominantly negative signature, regardless of their mineralogy, with native S showing the most negative values (Fig. 5).

The new sulfur isotope results are generally comparable with literature values for Campi Flegrei (Piochi et al., 2015), although studies earlier than 2000 (Cortecci et al., 1978; Valentino et al., 1999) also show positive  $\delta^{34}\text{S}$  values (Fig. 5a,b). In comparison to previous studies:

- 1) our new S-isotope data for Pisciarelli also include few positive values (Fig. 5b,c).
  - 2) the new O isotope values for sulfate are the highest obtained until now (Fig. 5d). To note, the muds generally have the least heavy oxygen, with except for the 2013 - 2014 data for which O-isotope determinations are lacking. The diagram also indicates a lowering in  $\delta^{34}\text{S}$  for neoformed minerals reported later than 2000.
  - 3) the sulfides at Pisciarelli show  $\delta^{34}\text{S}$  values mostly at 0 ‰ (Fig. 6).
  - 4) the different sites display a homogeneous range in  $\delta^{34}\text{S}$  (Fig. 5a,b,c); however, the variability for the ASA and L1 sites reflects different sample heights along slope (Fig. 1d) and wall (Fig. 1e), respectively.
  - 5) a likely appearance of positive correlation between S isotopes from pyrite and sulfate phases coexisting at Solfatara, with two from the 1994 data outside the trend (Fig. 6a).
  - 6) new isotope data for sulfate reveal a difference compared to studies older than 1990 and the most recent one (Fig. 7).
- In addition, the  $\delta^{18}\text{O}$  values for sulfates at Pisciarelli show a slight tendency to increase with topographic elevation at the sampling site. Specifically, the highest values generally occur at Solfatara that is at > 96 m asl while Pisciarelli lies at 66 - 74 m asl.

### 3.3 Content of major and trace elements (WRG)

Table S3 reports the whole-rock geochemical compositions of selected samples. As expected, samples are highly hydrated and sulfur-rich, due the OH-group and/or S in the crystalline network (see ideal formula in Table 1), and/or native S in the analysed sample. LOI can be up to 80 wt% (sample L20 camino 18.9.17), although most commonly at 20-30 wt%, and the S content is up to more than 50 wt%. Carbon is generally low (< 1.25 %), and always < 22 %. Notably,  $\text{SiO}_2$  content is highly variable. Depending on the sample mineralogy, it can be as high as 70-80 wt%. Also, MnO is always < 0.1 wt%.

With respect to the local volcanic rock substrate (Table S3; Fig. 8a), some lithophile elements (Si, Al, P, Sc, Ti, V, Zr, Ba, Yb, Th, Hf) are comparable or depleted (for examples, Be at < 1-5 ppm vs > 4 ppm, Ga < 23.8 ppm, Rb < 95.6 ppm, Y < 8.1 ppm). Depletion concerns with Na, Mg, K, Ca, Rb, Eu, U content. Only Ba displays a significant level of enrichment reaching very high values up to thousands of ppm. Cs shows concentrations of up to tens of ppm. Siderophiles and chalcophiles ratios (Fig. 8b) mostly lies at 1 or slightly above, with depletions for Fe, Zn, Ga, and enrichments for S, Au, As, Tl, Hg (Table S3). Sb can be higher than the primary rock composition.

## 4 Discussion

### 4.1 Environmental realms: stationary phenomena and runoff processes

Throughout the years, the various Solfatara and Pisciarelli sampling sites display a nearly constant mineral alteration assemblage (Table 1). Commonly, the mineral neogenesis variably develops on sub-mm- to dm- to m- scales, in relation to the outgassing dynamics, runoff, weather conditions, outcropping substrate, and anthropogenic activity.

However, the various sites further display reproducible rock geochemistry and stable isotope compositions at the timescale of survey, and with respect to oldest data (e.g., Valentino et al., 1999) as well; they can be considered reference points for future investigations.

Based on the presented dataset, we propose the existence of major realms, in which some (minor/peculiar) mineral phases can appear or disappear, in response to changing physical-chemical conditions mainly associated to weather circumstances, mostly humidity and water abundance. The realms are the geographical zones discriminated by their dominant and repetitive mineralogy, rock chemistry, and isotopic compositions, and characterized by temperature variations in a narrow range. Such a constancy is revealed when comparing results reported by Sicardi (1959) (see Geological setting at 2.1 section) with the present results, corroborating the existence of “stationary” realms that are presented in the following. The only exception is the mud pool in the crater.

The Pisciarelli and Solfatara pools (Fig. 1d,e) are the two major and distinct realms. They display persistent differences in dynamics, temperature and mud (solute plus water) mineralogy. The main pool at Pisciarelli shows vigorous boiling (Fig. 1b,e), with temperatures ranging from 63.9 to 94.3 °C (Tables S1 and 2) and a relative dominance of water vapour. The mud is typically grey in colour, mostly ash to sand up to mm-sized grains, with generally rounded/smoothed shapes as a function of the boiling vigour and material supply from the nearby slopes. On the other hand, the main Solfatara pool is characterized by degassing with a temperature at around 49 °C (Tables S1 and 2). The mud is beige and fine, always with a fetid odour. We recurrently detected an enrichment in pyrite, illite/montmorillonite and feldspar at Pisciarelli and in native sulfur at Solfatara. The latter is enriched in As, Hg, Nb, W, Zr, Sb and depleted in Sr, Ba, Co compared to the original deposits (Fig. 8). Sulfides and sulfates display nearly similar  $\delta^{34}\text{S}$  values at Solfatara, whereas they are different at the Pisciarelli mud pool (Fig. 6a, inset). The DRIFT-FTIR spectra of muds from Pisciarelli, in contrast to those from Solfatara, always gave the vibration at 1430  $\text{cm}^{-1}$  related to the  $\text{NH}_4$  (see section 3.1). Notably, the La Fangaia mud pool (Fig. 1d) is likely cooling as Sicardi (1959) reported up 100 °C, several tens of degrees higher than at the present (Tables S1 and 2). A slightly cooling can be also supposed by our measurements (Table S1) with respect those in the literature (Martini et al., 1991; Celico et al., 1992; Valentino and Stanzione, 2003). In contrast, the Pisciarelli area should be hotter, although only by a few degrees Celsius (Tables S1 and 2) if compared with increasing temperature values at the geyser-vent (Chiodini et al., 2016).

The PINT-PEXT realm (Fig. 1e) - an isolated morphological height - is composed of an alunitic-rich low-cohesive reddish terrain with a temperature around 95 °C, typically comprising kaolinite (Tables S1 and 2). The kaolinite is easily discriminated also by DRIFT-FTIR spectra (not shown) that, in agreement with XRD traces, point for a well-ordered structure. This terrain bears variably sized (up to few dm) clasts and is subject to slumping and sinking.

A hole up to 2-3 m deep represents a distinct realm that we emphasize because it opened 180 m North from the main pool within the crater in May 2017, by surface collapse. A grey viscously boiling mud fills the hole (Fig. 1c,d), with a minimum temperature of 70 °C; 91 °C is the temperature on April 2019. The mud is dominated by alunite with subordinate native sulfur, showing vague similarity to Solfatara mud (Table S1), although its grey colour and the occasional pyrite are reminiscent of the Pisciarelli mud. However, kaolinite is the main clay mineral in the new pool (Fig. S1e; Fig. S2e, note the inset). Mud geochemistry reveals peculiarly high concentrations of Sr, Ga, Co, Th, V, REE, Sb (Fig. 9).

Finally, a rather broad realm includes the other various sampling sites that are characterized by encrustations of alunite with a well-defined, although relative large, range of  $\delta^{34}\text{S}$  values between ca. -4 and 1 ‰ (Tables S1,2; Fig. 5c). These occurrences reflect the nearby presence of vapour degassing. We suspect that those at the ASA, M. Olibano and SSt sampling sites along the slopes of the Solfatara crater (Fig. 1a,d) are ascribable to long-lived encrustations; further investigations are useful to ascertain this suspect. The minor fumarolic vents around the old baths (Sst site) seem unchanged in respect to the description by Sicardi (1959), particularly showing the occurrence of native S and a comparable temperature at around 95 °C. These show variable bulk-rock geochemical compositions.

Widespread alunite formation reflects the potassium and feldspar-rich rock substrate on which they develop (see Piochi et al., 2014 and references therein).

Vapour effluents around the various geysers/vents at Pisciarelli are the most important factors that influence the neogenesis of the alunite-dominated realms. Pisciarelli is a decametres-deep incision on the NE Solfatara slope and the degassing vents are constrained in a gorge-like morphology. This setting favours the stagnation of the hydrothermal steam that impregnates the rock substratum and supplies elements to the formation of a variety of Na and NH<sub>4</sub> sulphates. We detected high abundance of those phases around the pool as desiccation during the summer season. The NH<sub>4</sub><sup>+</sup> ions were present in the solution and, possibly, as droplets in the humid air, as revealed by experimental desiccation tests of the water. Aerosol particles from inside and nearby the Solfatara crater that bear NH<sub>4</sub><sup>+</sup> (and Cl<sup>-</sup>, possibly in the form of NH<sub>4</sub>Cl) as major ion (Mather et al., 2004), as well as the NH<sub>4</sub>Cl inside the BG and BN orange-yellow encrustations (Fig. 3h,i), furthermore support the widespread presence of ammonium species. Notably, N<sub>2</sub> is emitted at the main intra-crater fumarole of Bocca Grande (see Chiodini et al., 2010). DRIFT-FTIR and XRDP spectra (Fig. S2d, S1a,d), however, point to the presence of NH<sub>4</sub> in both the mud and water pools at Pisciarelli, but not at Solfatara (except for BG and BN). Alum – (K) has also been found in relation to the wet conditions at Pisciarelli.

Vapour emissions outflow and the conditions of hydrothermal steam stagnation are dependent on atmospheric pressure and wind conditions.

So that, it is likely the meteoric weather the main cause for appearance to disappearance (and vice versa) of some phases.

This is also particularly evident for the PINT, PEXT, L19, L20 (Fig. 1e) and the SMO, ASA, SSt (Fig. 1d) realms that may typically present bipyramid and/or fine dendritic sulfur crystallites (Fig. 2a). Their crystallization seems to be favoured by relatively strong exhalations and porous terrain (PINT, PEXT, L19, L20; Fig. 1e) or conditions where gases remain briefly trapped (SMO, some places along ASA, Sst; Fig. 1d). Respective conditions also prevail in close proximity (< dm scales) to the main fumaroles (L1 vent, BG, BN; Fig. 1d), where sulfur forms a cream-like patina reflecting the condensing gas flow along the host fractures and fissures.

However, native S disappears during run off and we have macroscopically tested at several places that re-crystallization needs one-to-two months, if not longer (i.e., sample L20 camino; Fig. 1e).

Furthermore, periods of intense rainfall determine the areal extent and depth of the mud pools, the generation of secondary mud vents and the erosion in Pisciarelli and its periodic water puddle. Sicardi (1959) also described the occurrence of mud vents and black mud pools following rainy periods. Notably, pools at Pisciarelli are supported by anthropogenic embankment. Meteoric and surface waters can dilute the aggressive endogenous fluids determining alteration degree conditions low enough for the generation of illite/montmorillonite (Pirajno, 2008) at Pisciarelli. Further studies need to be performed in order to better characterize clays as they can bear information useful to further constrain the hydrothermal setting.

Al, Fe sulphates (halotrichite) have been rarely found nearby the Pisciarelli geyser (see G in Fig. 1e; Tables 1,S1).

The distribution of sulfates appears irregular, and this should be an object of future investigations.

#### **4.2 Classification of alteration and genetic environments: the contradictory data**

The style of mineralization (Arribas, 1995; Sillitoe, 1993; Pirajno, 2008; Ercan et al.; 2016) and the stable isotope results (Rye et al., 1992) allow classification of alteration and differentiation of genetic environments. Table 3 summarizes characteristic mineralogical, lithological and isotopic features of these environments, in comparison to observations made at the study sites. Several contrasting interpretations can result from the data.

Alunite plus kaolinite and montmorillonite form in steam-heated environments at 100 to 160 °C where fumarolic vapour condenses above the boiling zone of nearly neutral-pH, H<sub>2</sub>S-rich fluids representing a low sulfidation environment.

Nevertheless, alunite shows grain sizes in the range of 50 to 100 μm (Fig. 3a,e,f), unlike the finest (<20 μm) ones detected in high-temperature (>40-50 °C) steam-heated (Simón et al., 2005) or supergene (Arribas, 1995) environments. Those coarser

sizes usually occur in low-temperature steam-heated (and hypogene, as well) environments (Hedenquist et al., 2000), or could directly crystallize from a SO<sub>2</sub>-rich magmatic vapour that rapidly ascent through fractures (namely high sulfidation setting; Rye et al., 1992; Stoffregen and Alpers, 1992). The occurrence of kaolinite and alunite at several Solfatara and Pisciarelli realms (particularly, new pool and PINT, Table 1) fits with the high sulfidation environment; in fact, the two phases usually coexist in the advanced argillic alteration zones proxy to ascent plumes (e.g., Pirajno, 2008). The evidence of K-feldspar replacement by alunite (Piochi et al., 2015), the disseminated pyrite and the presence of native S at Pisciarelli apparently support the high sulfidation - magmatic hydrothermal environment (Rye et al., 1992). Nevertheless, illites/montmorillonites are most common in intermediate argillic alteration zones (Pirajno, 2008) and their widespread occurrence in the various studied realms is among the results that contrasts this high sulfidation environment. Indeed, sampling temperatures are higher than 40 °C (Tables 2 and S1) and SO<sub>2</sub> is rare or absent (Aiuppa et al., 2013) at Campi Flegrei. Also, the lack (or rare detection) of the lower temperature (< 40 °C), disordered polymorphs (i.e., halloysite) point to limited supergene alteration.

Only, the alunite coexisting with kaolinite in the new hole pool has the finest grain sizes. Accordingly, the XRDP and DRIFT-FTIR analyses of CF samples point to slightly ordered kaolinite forms that usually occurs at temperatures < 150 °C in epithermal systems (Sillitoe, 1993), but could also represent a metastable form in hotter settings (Zotov et al., 1998). In this later case, the new pool realm with kaolinite and alunite can represent a local, well-circumscribed advanced argillic alteration condition indicative of a proxy plume.

However, when considering litho-geochemical parameters, schematic diagrams further produce contrasting visions. For example, following Ercan et al. (2016), the clay-bearing muds can be ascribed to various supergene to hypogene alteration field in the binary diagram of immobile Zr vs. TiO<sub>2</sub> (Fig. 9a). They also have high (above 1,000 ppm) Ba+Sr and low (<200 ppm) Ce+Y+La concentrations (Fig. 9b).

The stable isotope geochemistry of minerals supports an interpretation of steam-heated to supergene environments (Fig. 7). S-isotope equilibrium occurs between sulfides and sulfates, with reliable re-calculated temperatures in high-sulfidation environments (Arribas, 1986). In contrast, this equilibrium cannot be accounted at Campi Flegrei and any reliable temperatures result because of S-isotope fractionation between sulfates and H<sub>2</sub>S. In fact, δ<sup>34</sup>S values of alunite and pyrite roughly overlap. Instead, sulfur-H<sub>2</sub>S and sulfides-H<sub>2</sub>S at the sampling temperature reflect equilibrium S-isotope fractionation: the theoretical δ<sup>34</sup>S value of dissolved H<sub>2</sub>S fluid is between -3.84 and -0.84 ‰ (Allard et al., 1991). This also implies that the sulphate altered rocks are not magmatic hydrothermal in origin, in agreement with the lack of typical mineral phases that show high oxidation state of S (SO<sub>2</sub>-rich, high-sulfidation according to Henley and Ellis, 1983). Based on Rye et al. (1992), SO<sub>2</sub> disproportionation results in the formation of <sup>34</sup>S-enriched H<sub>2</sub>SO<sub>4</sub> and <sup>34</sup>S-depleted H<sub>2</sub>S. In addition, the intense Al-leaching in a high sulfidation system is not typical for the Campi Flegrei setting (Fig. 8; Table 3).

Actually, Campi Flegrei lacks of the occurrence of enargite and luzonite, both diagnostic for high-sulfidation environments, and instead shows minor occurrences of realgar (AsS) as well as cinnabar (HgS) (Tables 1,S1), and orpiment has been described (Russo et al., 2017).

Significantly lower δ<sup>34</sup>S values (< 15 ‰) for alunite can derive from: (i) the light sulfur isotopic composition of H<sub>2</sub>S during boiling (steam-heated or low-sulfidation setting), (ii) δ<sup>34</sup>S of sulfides (supergene) or (iii) the bulk sulfur isotope composition of magmatic steam (Rye et al., 1992). The first possibility could partly account for the isotopic composition of alunite-pyrite and alunite-H<sub>2</sub>O pairs (Fig. 10), although contrasting with conclusions based on texture, mineral assemblage and bulk-rock geochemistry (Table 3). The presence of kaolinite in the subsurface, under an alunitic cover and the occurrence of argillic alteration at depth suggest a vertical zonation at the Solfatara crater and, following Rye et al. (1992), a steam heated setting.

Furthermore, δ<sup>34</sup>S values for the pyrite-H<sub>2</sub>S pairs further support a supergene setting (Fig. 10).

Finally, we are not able to directly identify any microbial sulphur cycling, although FT-IR and rock geochemistry corroborate the absence or limited biota contribution. The analysed samples do not produce bands attributable to C=H ligands (Supplement)

and the carbon content is <1.25 wt% (most common < 0.2 wt%; Table S3). Yet, some higher  $\delta^{34}\text{S}$  and  $\delta^{18}\text{O}$  values for sulfates could be indicative, particularly considering the dry-wet alternating conditions.

#### 4.3 Merging information and unravelling the setting

Merging all available information, it appears that observations concerning the both realms' "stationarity" and contradictory classification environment reflect the evolving conditions that have followed the last magma intrusion and eruption and that probably are overlapping through time.

The solfataric alteration zone has a strongly limited extent within the central sector of the Campi Flegrei caldera. It coincides with the area of eruptive vents (e.g., Mt. Olibano, Accademia, Solfatara; Fig. 1a) and uplift of the most recent period of volcanism (Di Vito et al., 1999). The zone appears to be limited under the later Fossa Lupara and Astroni vents, while degassing and thermal aquifers occur within the caldera. However, there is an indication for their discrete, more than their continuous distribution, both across the caldera and through depth (Guglielminetti, 1986).

The studied deposits are young and nearly coeval (<5000 years) with the altered volcanic basement deposits (i.e., Monte Olibano, Solfatara).

The alteration zone locally presents high Ti, Ba, Au, As, Hg, Tl, S concentrations relative to the above parent basement lithology (Fig. 8).

The zone also appears anomalous in terms of ammonium content. Therefore, we here adopt ammonium as a possible tracer, but we have no information yet about the various contributing sources for the N-species and the cycling of nitrogen at the local scale. The presence of  $\text{NH}_4$ -bearing sulfates is particularly abundant at Pisciarelli; those sulfates systematically form by drying water collected at the various pools of the area, in relation to the abundance of nitrogen species (0.2-1 g/l) in those waters (Martini et al., 1991; Celico, 1992; Valentino et al., 1999; Holloway and Dahlgren, 2002; Valentino and Stanzione, 2003; 2004; Aiuppa et al., 2006). Actually, the  $\text{NH}_4^+$  content in the shallowest Phlegraean ground waters is generally low (<0.03 g/l; Martini et al., 1991; Celico, 1992; Valentino et al., 1999; Valentino and Stanzione, 2004; Aiuppa et al., 2006). Yet, Mather et al. (2004) measured a significant abundance of ammonium chloride particles/aerosols at the Solfatara crater. Some realgar encrustations sampled at BG and BN also associate with  $\text{CINH}_4$  (Table S1; Fig. 3h,i).

The concentration of some metals and metalloids requires sources different from the parent basement. Anthropogenic contributions are obviously possible (Alloway, 2012), e.g. when considering that  $\text{NH}_4$  is generally attributed to agricultural (i.e., fertilizer) and urban soils. However, Hg and As have been detected at the main fumaroles with similar concentrations today and in the last century (Ferrara et al. 1994; Bagnato et al., 2014). These fumaroles continuously emit  $\text{H}_2\text{S}$  and  $\text{CO}_2$  (Allard et al., 1991; Aiuppa et al., 2013; Chiodini et al., 2016) and are the preferred location of crystallization of native S and alunite.  $\text{NH}_4^+$  emissions are also present at Solfatara (Chiodini et al., 2010), although ammonium is of limited importance inside the crater waters literature data <0.001 g/l (Aiuppa et al. 2006) and crystallization of alum instead than  $\text{NH}_4^+$ -sulfates in the water from Solfatara pool. In agreement with these authors, the plausible source is the conventional geothermal reservoir. High concentrations (20-100 g/l) of ammonium are reported in the deeper (>500 m; Carlino et al., 2012) aquifers at the Mofete wells, located on the western side of the Campi Flegrei (Chiodini et al., 1988). Those deeper aquifers are located within a sequence of tuffs and marine sediments also drilled by the CF23 well (1000-1200 m depth; Piochi et al., 2014; 2015), i.e. nearby the solfataric area.

In summary, we propose an environmental setting that merges all collected information (Fig. 11). Fluid outflows from discrete aquifers hosted in sediments – and bearing organic imprints – feed the Pisciarelli site giving its ammonium peculiarity. Our purpose does not exclude the possible biological contribution that has been ascertained in the studied sites (e.g., Ciniglia et al., 2005; Glamoclija et al., 2004). However, marine strata and volcano-clastic sequence intercepted by deep drillings (San Vito1, Mofete and CF23 wells; Rosi and Sbrana, 1987; Piochi et al., 2014) are the key sediments of the  $\text{NH}_4$  species. An additional supply can be the swampy sediments encountered in shallowest boreholes (de Vita et al., 1999) located in the central sector of

the caldera (Fig. 11), nearest to the study area. Results from cluster analysis of resistivity, P-wave velocity and density parameters, from Pisciarelli across the Solfatara crater (Di Giuseppe et al., 2017) lithologically constrain the model section. These authors highlight a sudden upraise up to ca. 1500 m depth of buried rocks through a tectonic structure just beneath Pisciarelli. The dislocated rocks are the fossiliferous marine and volcano-clastic sequences drilled across the caldera. The deep aquifer is represented by a confined body with a high electrical conductivity ( $\log\rho = 0.97 \Omega\text{m}$ ). At the deepest position in the model, we correlate thermo-metamorphic rocks with the brines characterized by  $\log\rho$  at  $2.7 \Omega\text{m}$ ,  $V_p = 3800 \text{ m/s}$  and  $\Delta\sigma = 38.8\text{kg/m}^3$ . These physical features are indicative for the occurrence of voids/fracturing and the migration of gases. Therefore, we infer a deep source of gases emitted at the surface, which likely also indicates the location of the heat source.

Shallow and deeper aquifers are interconnected via a network of “communicating vessels” through a fault system, allowing deeper and shallower water to mix and being expelled at Pisciarelli. This justifies an apparent persistence of thermal springs around the Agnano Plain also in presence of the desiccating lake described by Ventriglia (1942). It also supports the depth of the water table, being at a higher topographic position in the Solfatara area in respect to the surroundings (Astroni, Bruno et al., 2007).

In the model, we further speculate that the acid-sulfate alteration zone at the Campi Flegrei is actually evidence of a paleo-conduit. This idea comes from field observations that indicate the alteration deposits locally underlying the most recent eruptive units (e.g., Astroni) that are unaltered. Therefore, the texture of the mineral assemblage, the enrichment in some metals and the lithogeochemical parameters are relict of a “high-sulfidation system”. The evolutionary dynamics within the conduit and, in particular, the water overflows from the aquifers alternating with runoff processes, explains the contradictory mineral environments with superimposed intermediate and advanced argillic alteration.

At present, a steam-heated (or low-sulfidation) environment (as derived by most isotope data on alunites; see previous section) is developing in relation to the presence of aquifers and their chemical compositions. This is in agreement with previous studies (e.g., Aiuppa et al., 2017; Piochi et al., 2015; Gresse et al., 2017). Following Hedenquist and Lowensten (1994), this is also in agreement with the shift in  $\delta^{18}\text{O}$  at constant  $\delta^2\text{H}$  values of the emitting fluids (Caliro et al., 2007). Based on the  $\delta^{18}\text{O}$  values of alunite, the recalculated environmental temperature is  $\leq 200 \text{ }^\circ\text{C}$  (Fig. 8, inset a).

Furthermore, the presence of  $\text{NH}_4^+$  is considered strictly connected to the surface environment and likely to organic/biological processes (Jaffe, 2000), which is consistent with S-isotope values of pyrite- $\text{H}_2\text{S}$  pairs, the heavier  $\delta^{34}\text{S}$  values suggesting microbial sulfur cycling and the supergene environment that is locally developing.

## 5 Conclusions and Outlook

The acid sulfate alteration zone at Pisciarelli and Solfatara is located in the sector of the Campi Flegrei caldera that was the most volcanically active area in the last 5 ka. The alteration zone includes discrete realms with very constant mineralogy, temperature and chemistry, considering the studied time interval. Outgassing dynamics, weather conditions, and runoff are the most important factors affecting the generation of new mineral phases at the variable sub-mm- to dm- to m- scales.

The new minerals include alunite, alunogen, native sulfur, pyrite, kaolinite and subordinately mascagnite.

The limited areal extent of the alteration zone underlying the most recent unaltered volcanic units, its mineralization texture and style, the  $\delta^{18}\text{O}$  and  $\delta^{34}\text{S}$  values of S-bearing minerals, and the enrichment in Ti, Ba, Au, As, Hg and Tl, are possibly attributes of the evolution of a paleo-conduit. Our mineralogical and isotopical results overlap with those in Valentino et al. (1999), favouring a stability in the hydrothermal dynamics over the past 20 years. The zone is anomalous in terms of the presence of  $\text{NH}_4^+$ . These features result from the mixing between waters that overflow through a fault system intercepting discrete aquifers supplied by surface water and deep fluids. Most of the alunite forms above the water table at a temperature  $\leq 200 \text{ }^\circ\text{C}$ . At present, the dominant steam-heated environment coexists with local supergenic conditions.

Based on presently available data, several key aspects await further investigations.

In particular, a detailed survey of the distribution of aquifers in the subsurface will foster our understanding of caldera dynamics and contributes to the debate existing between as “hydrothermal” (Moretti et al., 2017) vs. “magmatic” (Cardellini et al., 2017) unrest. Assessing the composition and spatial extent of aquifers - also including the contribution from rain fall - is crucial in solving the non-magmatic role into processes at the surface. Soluble acid components (SO<sub>2</sub>, HCl and HF) sourced at depth are  
5 condensing in the shallower aquifer system (Valentino and Stanzione, 2003; Aiuppa et al., 2007; Caliro et al., 2007; Vaselli et al., 2011; Piochi et al., 2014; 2015; Chiodini et al., 2016). However, the ability to buffer these magmatic fluids clearly depends on the water availability in relation to the volume of fluid, with implications on the sourced magma volume(s) evolving/degassing at depth. On the other hand, the circulation of fluids in the subsurface, sourced from both the downward surface infiltration and the ascent of deep fluids, would contribute to the pressurization that is evident through shallow  
10 seismicity as previously suggested (Saccorotti et al., 2007; D’Auria et al., 2011; Di Luccio et al., 2015). Most important, knowing the water availability in the subsurface is crucial for evaluating the volcanic hazard in an area dominated by phreatomagmatic events, such as the Campi Flegrei caldera (Rosi and Sbrana, 1987; Di Vito et al., 1999).

What cause the presence of NH<sub>4</sub><sup>+</sup> is still rather elusive. Based on Moretti et al. (2017), it appears useful for evaluating the temperature-dependence of N<sub>2</sub> vs. ammonia production and the relative role of hydrothermal vs. magmatic systems. We add  
15 the possible role of organic materials and bacteria, and atmospheric sources to the terrestrial cycle. Furthermore, ammonia NH<sub>3</sub> is toxic (Fromm and Gillete, 1968) and this requires ascertaining its concentration level in an inhabited environment.

Finally, the Pisciarelli site appears suitable for the biota and the life evolution studies. Here, the water-dominance, nitrogen richness, ≤ 200 °C temperatures and supergenic conditions are all needed ingredients for the growth of the organic substance and organisms (Jaffe, 2000). Consequently, this site could become a natural laboratory for investigating the complex organic-  
20 inorganic interface/relations through multidisciplinary collaborations among mineralogists, geochemists, petrologists, volcanologists, and biologists.

## 6 Appendix

### Appendix A

XRDP and DRIFT-FTIR spectra were acquired at the Osservatorio Vesuviano (Istituto Nazionale di Geofisica e Vulcanologia,  
25 Naples, Italy).

The XRDP instrument was a PANalytical X’Pert equipped with a high speed PIXcel detector (Mormone et al., 2014). The configuration includes Ni-filter, CuK $\alpha$  radiation, pyrolytic graphite crystal monochromator, 40 kV and 40 mA current, 3–70°, 2 $\theta$  range, 0.02° steps and 8 s/step. X’Pert HIGH Score Plus computer program and JCPDS PDF-2 database allowed the interpretation of diffraction patterns.

30 DRIFT was mounted on a Nicolet 670 Nexus<sup>TM</sup> both by ThermoFisher Scientific S.p.a.. The FTIR comprises a heated ceramic (Globar) source, a 670 Laser unit, a KBr beamsplitter, and an MCT detector, constantly purged from a high-pressure Nitrox dry air and CO<sub>2</sub>-trapping 280 generator by Domnick Hunter. The OMNIC Data Collector 5.2© allows data collection and interpretation in the investigated range of 5000 - 400 cm<sup>-1</sup> (resolution: <0.1 cm<sup>-1</sup>). Data collection was conducted on KBr mixed with 5 – to – 10 % of sample by grounding in an agate mortar, following the background acquisition for the KBr powder.

35 Additional acquisition on no diluted samples allowed checking for possible hygroscopic effects, obtaining similar results.

The appearance, morphology and chemical composition of minerals were determined on selected samples prepared as opaque mounts coated by cord and rod graphite, by JEOL and ZEISS electron microscope (EDS-BSEM) facilities. The JEOL-JSM 5310, equipped with a Link EDS and a Inca 4.08 software (CISAG Laboratory University of Napoli Federico II), has operating conditions of 15 kV accelerating voltage, 50–100 mA filament current, variable spot size and 50 s net acquisition time. ZEISS  
40 instrument is a SIGMA field emission scanning electron microscopy (Osservatorio Vesuviano, department of Istituto

Nazionale di Geofisica e Vulcanologia, Naples, Italy), equipped with XMAN micro-analysis system by Oxford, controlled by a SMARTSEM and AZTEC softwares. Operating conditions for SIGMA were 15 kV accelerating voltage, 50–100 mA filament current, 5-10 nm spot size and variable time acquisition time (several to tens of seconds).

Rock geochemistry (WRG) were carried out at Bureau Laboratories Ltd. (Vancouver, Canada). Major elements were analysed by Inductively Coupled Plasma Emission Spectrometer (ICP-ES) using  $\text{LiBO}_2/\text{Li}_2\text{B}_4\text{O}_7$  fusion, minor and trace elements were determined by inductively coupled plasma-mass spectrometry (ICP-MS) using a four acid ( $\text{HNO}_3\text{-HClO}_4\text{-HF-HCl}$ ) digestion. The uncertainty is generally <1% for major/minor oxides, <5–20% for trace elements. LECO was used for determining the C and S abundances. Loss on ignition (LOI) was calculated by weight loss after ignition at 1000°C.

Sulfur and oxygen isotope measurements were performed directly on pure mineral separates without and with further chemical preparation in the stable isotope laboratory at the Institut für Geologie und Paläontologie (University of Münster). Chemical preparation was different depending on sample type: i.e., sulfates+elemental S+sulfides, elemental S +sulfides, or muds. Oxidized S-bearing and multi-phases samples first required the extraction of sulfate by sample leaching in a 10% NaCl solution, filtration through a 0.45 micron cellulose acetate filter, and followed by the addition of 8.5% barium chloride solution to precipitate dissolved sulfate as barium sulfate for isotope measurements. Elemental S and pyrite extraction was performed on sulfate-free powders. This extraction consisted of a wet chemical treatment (acidic chromous chloride solution at sub-boiling conditions) that liberates sulfur as hydrogen sulfide which will ultimately be precipitated as silver sulfide, ready for isotope measurements. Elemental sulphur was liberated from each sample via acetone leaching and subsequently converted to silver sulfide applying the acidic chromous chloride treatment (Canfield et al., 1986). Again, resulting hydrogen sulfide was precipitated as silver sulfide ( $\text{Ag}_2\text{S}$ ). For S-isotope measurements, mineral separates as well as silver sulfide precipitates were homogenously mixed with vanadium pentoxide in tin capsules and combusted in a Carlo Erba elemental analyzer interfaced to a ThermoFinnigan Delta Plus mass spectrometer (EAIRMS: Elemental Analyzer-Isotope Ratio Mass Spectrometry). Results are reported in the standard delta notation ( $\delta^{34}\text{S}$ ) as per mil difference to the Vienna Canyon Diablo Troilite (VCDT) standard. Reproducibility as determined through replicate measurements was generally better than  $\pm 0.3$  ‰. Sulfates were also measured for O-isotopes by using a ThermoFinnigan TC/EA interfaced with a ThermoFinnigan Delta Plus XL. Results are reported in the standard delta notation ( $\delta^{18}\text{O}$ ) as per mil difference to the Vienna Standard Mean Ocean Water (VSMOW) standard. Reproducibility as determined through replicate measurements was generally better than  $\pm 0.5$  ‰.

## 7 Supplement

Three supplementary tables:

Table S1 – List of samples, collection date, temperature and mineralogical associations as resulting by XRDP analyses and corroborated by FTIR and EDS-BSEM study. The sampling includes water spring sampled at Stufe di Nerone. In the temperature column: tc, thermo couple (see chapter 2.2 Sampling, sample preparation and analytical techniques), infr, infrared gun. In the mineralogy column: ?, for minerals to be validated; minerals in red are approximate attribution based on XRDP patterns. The orange cells evidence water samples. Representative XRDP spectra are in Fig. S1. Further details in this supplement.

Table S2 – Vibration modes and related tentative assignment of functional groups, and mineral attribution for selected samples by DRIFT-FTIR investigations. Alu = alunite, Clay = illite/montmorillonite, Masc =  $\text{NH}_4$ -bearing sulfates, am = amorphous, Kao = kaolinite, KAl = alum - (K). ?, uncertain attribution. Note: assignments and attributions are based on mineralogy derived by XRDP study and corroborated by EDS-BSEM analyses. Further details in this supplement.

Table S3 – Selected whole-rock geochemistry of multi-phases materials sampled at different locations (i.e., sample name as in Fig. 1) within the Pisciarelli and Solfatara areas and at different times. MDL indicates the detection limit for major, trace, C and S contents.

- 5 Figure S1 – Representative XRDP spectra of NH<sub>4</sub>- sulfates dominating the assemblage formed from drying the Pisciarelli water (a, b) and of various muds from Solfatara (c,e) and Pisciarelli (d,f). Each panel reports the sample name in Table S1. The muds show the large hump between 18 ° and 30 ° 2θ degree attributed to the amorphous phase. The smaller panels evidence the reflection intensity in the most significant range useful to discriminate illite in d) and kaolinites in e). Some XRDP spectrum has a corresponding infrared spectrum in Fig. S2: the sample with mascagnite is the same of Fig. S2a, the Solfatara muds in c) produced the FT-IR spectra in Fig. S2b, the Pisciarelli mud MP 6\_16 is in Fig. S2d, the samples in e) are the same of Fig. S2e. Abbreviations (c, f): S = Sulfur; Al = Alunite; Kfd =Alkali feldspar.
- 10

Figure S2 - FT-IR spectra of NH<sub>4</sub> - sulfates (a), native sulfur (b) and various muds from Solfatara (c), Pisciarelli (d) and the new Solfatara hole (e). See Table S2 for vibrational modes and relative assignments.

## 15 8 Team list

Suspendisse a elit ut leo pharetra cursus sed quis diam. Nullam dapibus, ante vitae congue egestas, sem ex semper orci, vel sodales sapien nibh sed lectus. Etiam vehicula lectus quis orci ultricies dapibus. In sit amet lorem egestas, pretium sem sed, tempus lorem.

## 9 Author contribution

- 20 MP and AM conducted sampling campaigns and prepared samples for analyses. GB participated in some of the sampling campaigns. AM conducted the XRPD analyses and interpreted the spectra. MP acquired, elaborated and interpreted the DRIFT-FT-IR spectra and, in collaboration with AM and GB, performed the EDS-BSEM investigations. HS determined the stable isotope values and contributed to data elaboration; MP did data representations and stable isotope data modelling. MP prepared the manuscript. All authors contributed into the final manuscript.

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- 30 Angarano, Tennis Hotel and Stufe di Nerone allowed the free access at the sampling sites.

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### Caption of Figures and Tables

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Figure 3 – BSEM image showing sample texture and occurrences of S-bearing phases identified by EDS and XRPD analyses at the Puteolis sulfate lands: (a) orthorhombic baryte (Ba) front of pentagonal pyrite (Py) from the Pisciarelli mud (L3); (b) irregular platy alunogene (Alu) oriented parallel to the fracture axis; (c) rhombic native sulfur (S) with bladed tschermigite crystals (Ts); (d) acicular alunogene (Alu) crystals developed above early tabular alunogen species; (e) euhedral alunite grains (Al) showing resorbed surfaces and coexisting with tabular alunogen (Alu) and acicular gypsum (Gy) crystals; (f) massive alunite (Al) encrustation; (g) pseudo-cubic ammonium chloride ( $\text{CINH}_4$ ) crystals within encrustation sampled at Bocca Grande (BG); (h) monoclinic realgar (Rlg) and ammonium chloride ( $\text{CINH}_4$ ) individuals at Bocca Grande (BG); (i) grains coated by pyrite and massive letovicite crystals. Alu = alunogene; Al = alunite; Ba = barite;  $\text{CINH}_4$  = ammonium chloride (salammoniac);

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Table 3 – Summary of the mineralogical and isotopical features at the acid sulfate area following Rye et al. (1992); Hedenquist and Lowerstern (1994). \*highest  $^{34}\text{S}$  and  $^{18}\text{O}$  during bacteriogenic reduction of sulfates with maximum fractionation in dry-wet alternating conditions.  $\delta^{34}\text{S}$  reflects the  $\text{H}_2\text{S}/\text{SO}_2$  and temperature of fluid. 1 always present, 2 may be associated. 3 from Valentino and Stanzione (2003; 2004), Gresse et al. (2017). ‘halloysite is indicated in Montanaro et al. (2017) and included here for completeness.

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## Figures and Tables

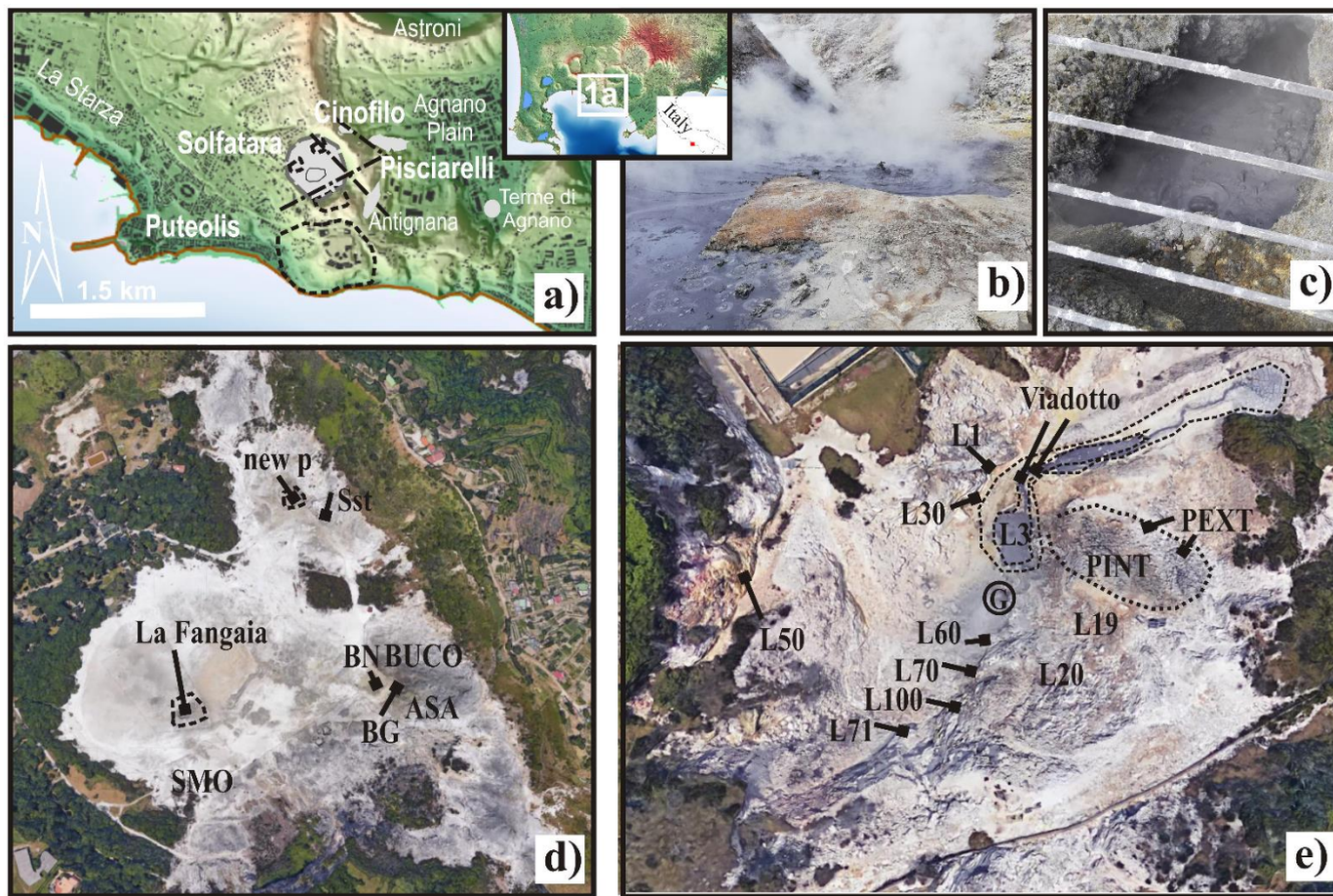


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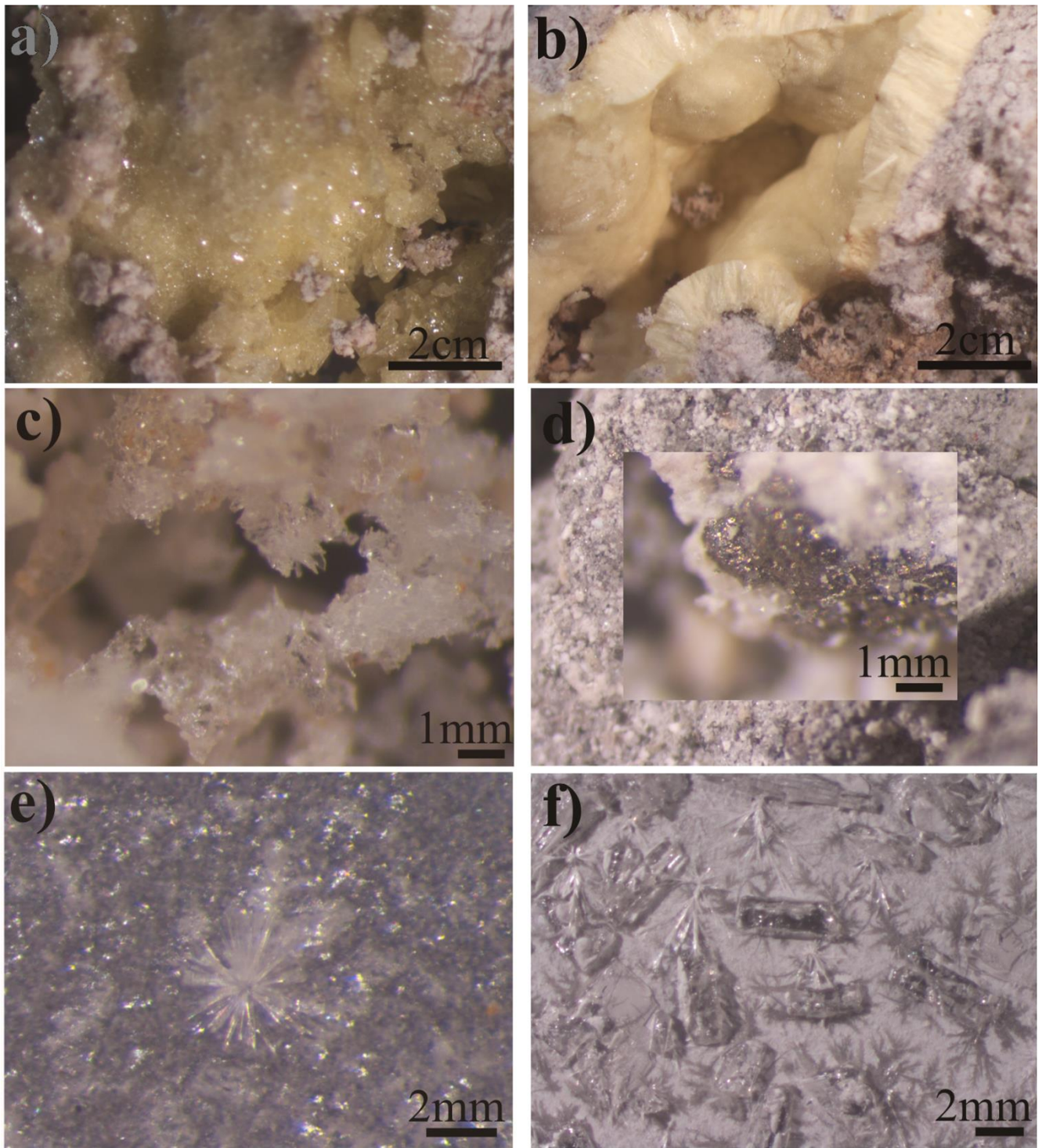


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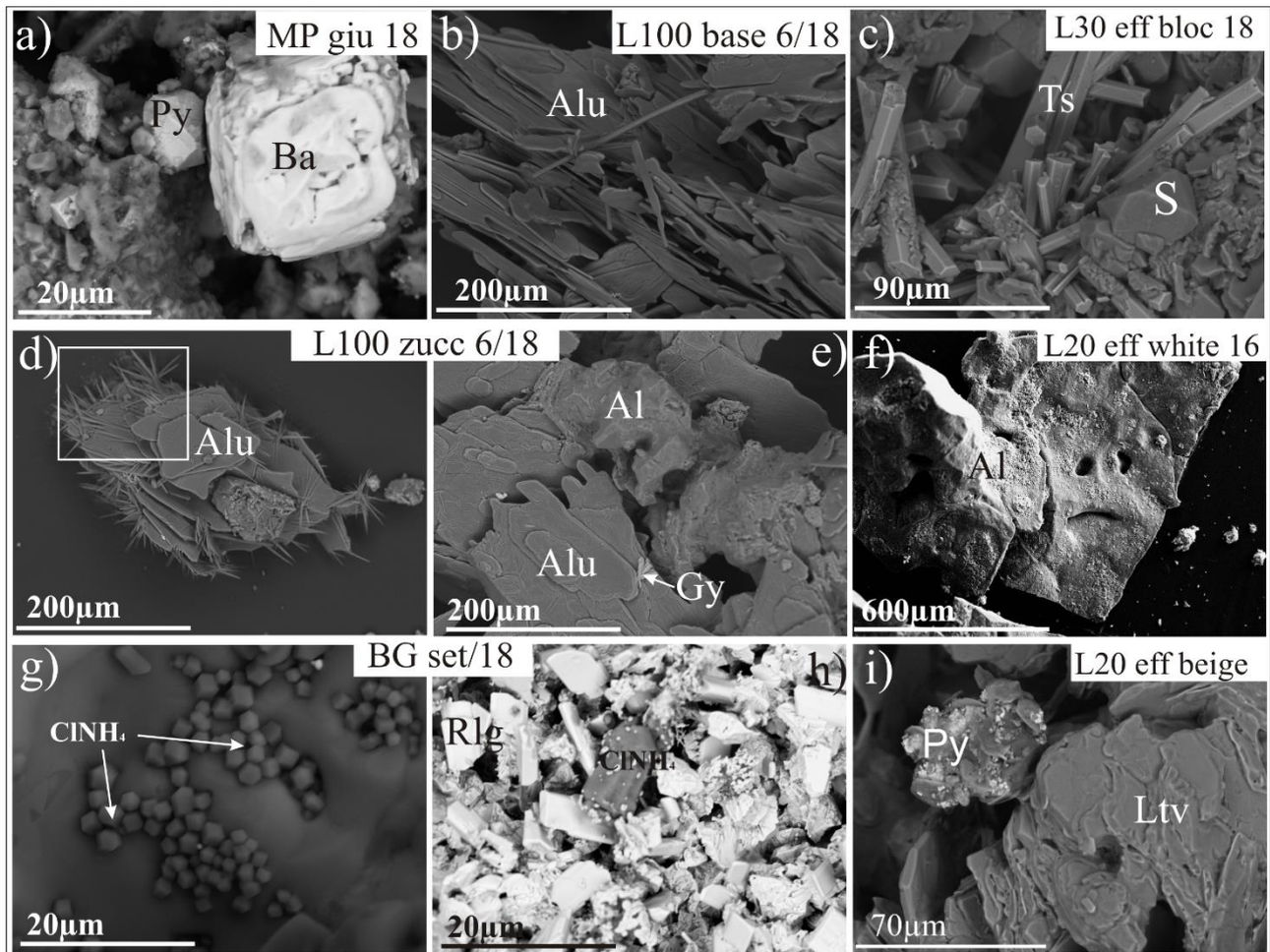


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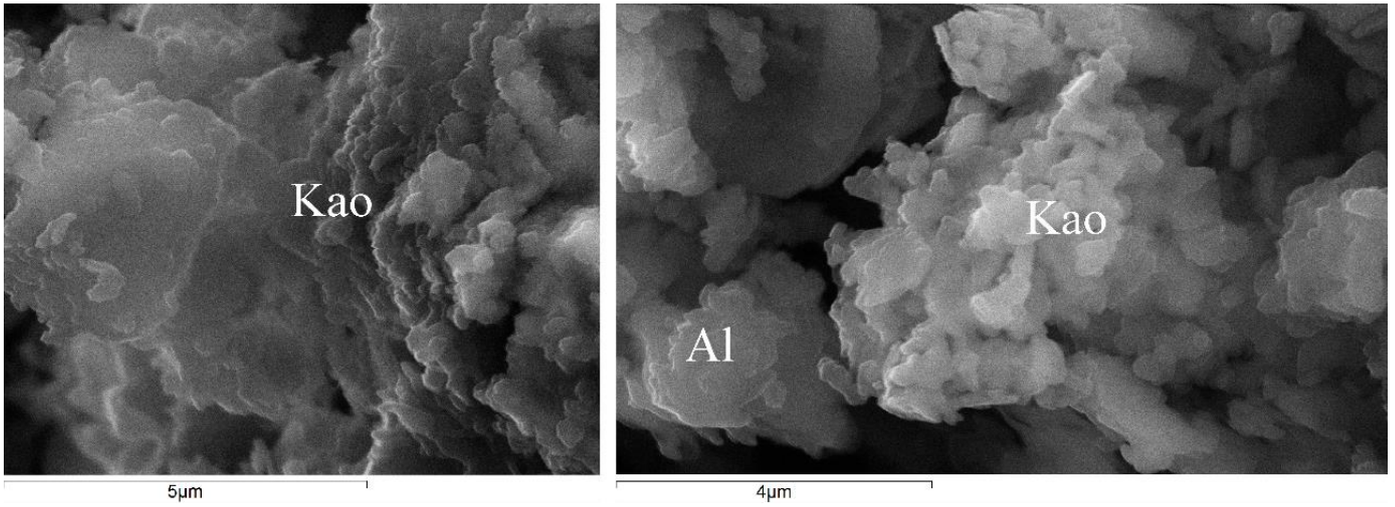


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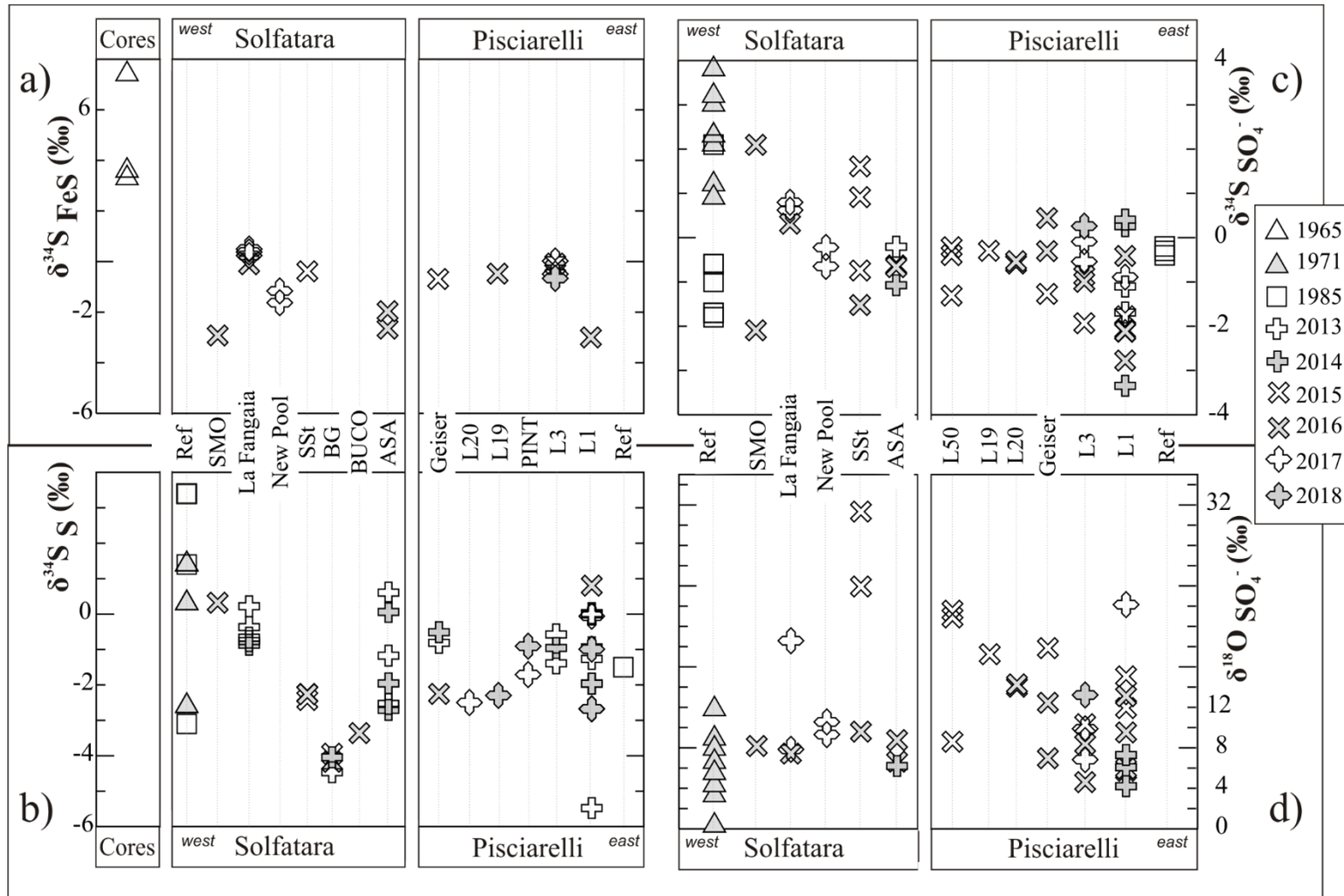


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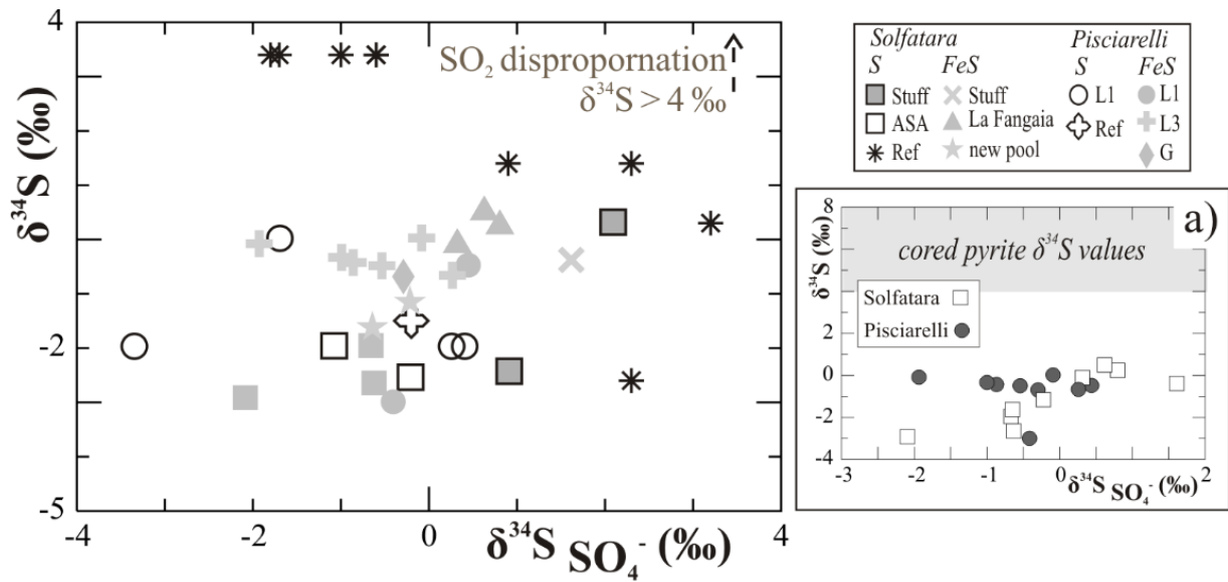


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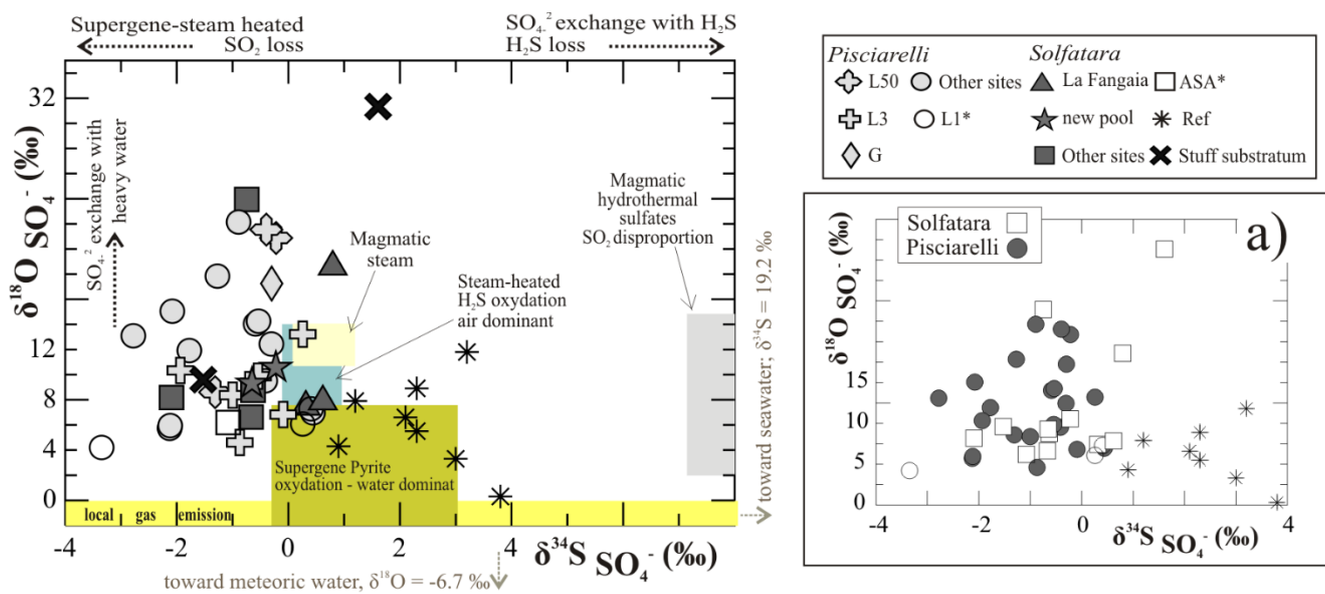


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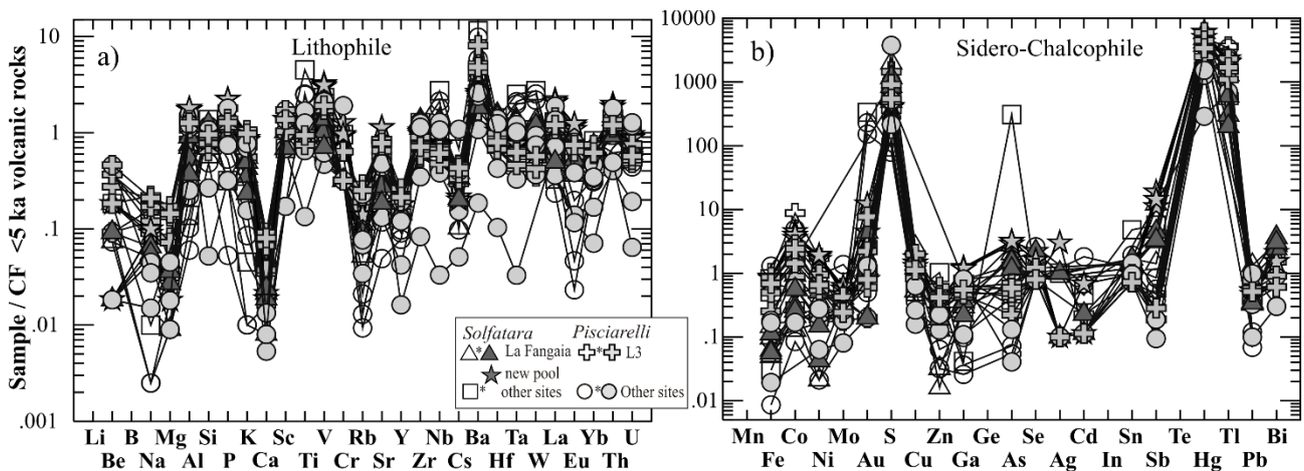


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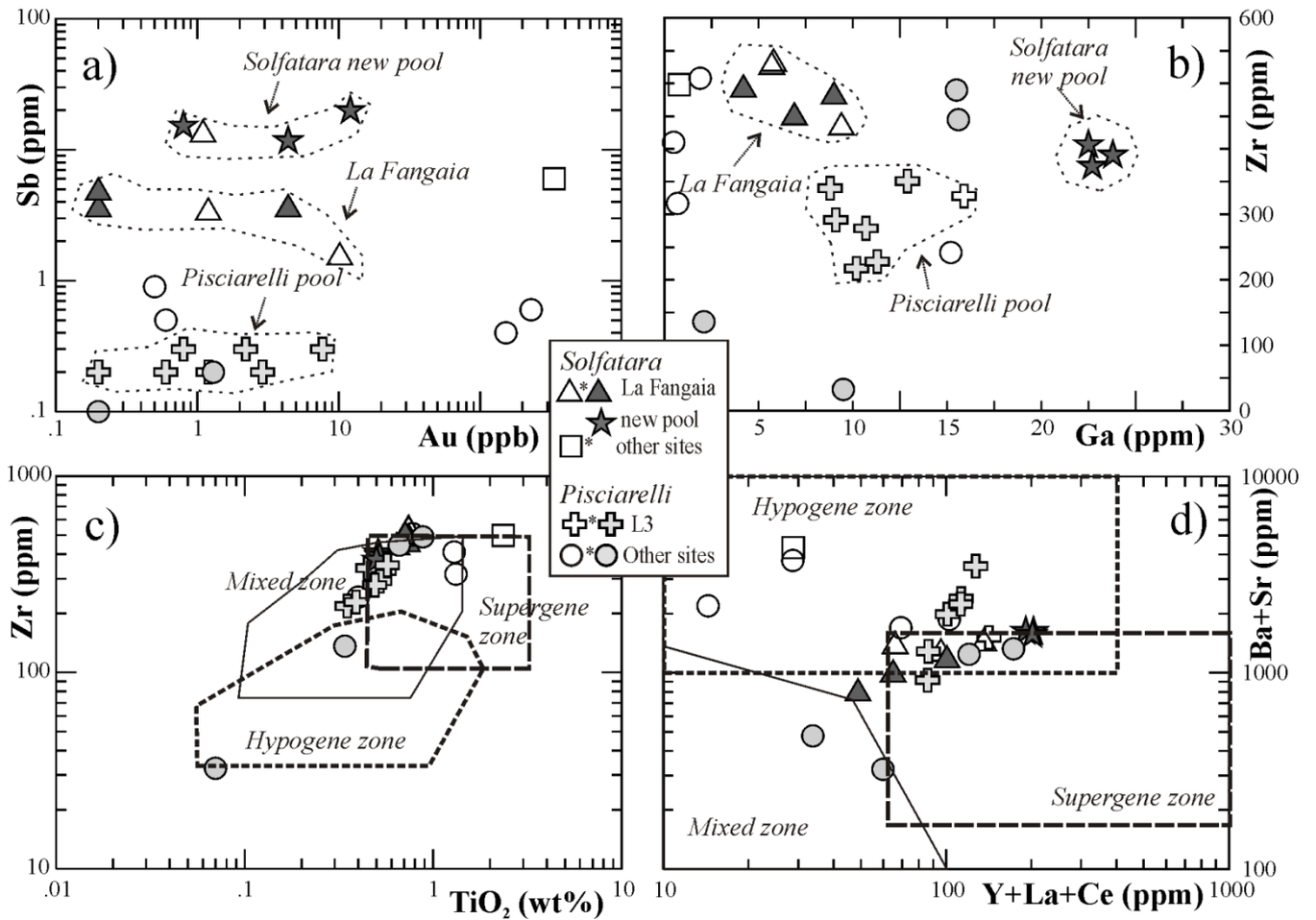


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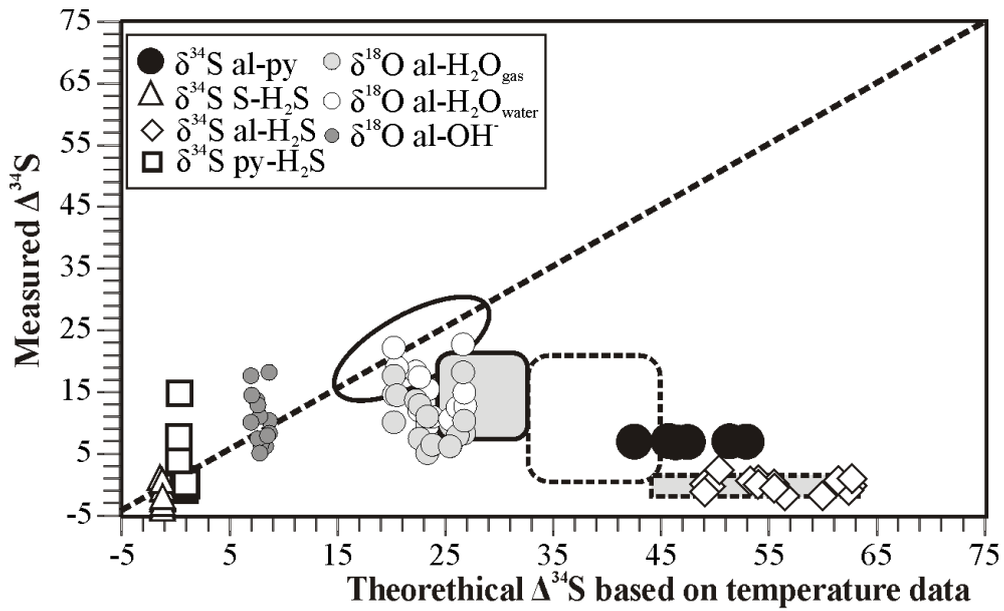


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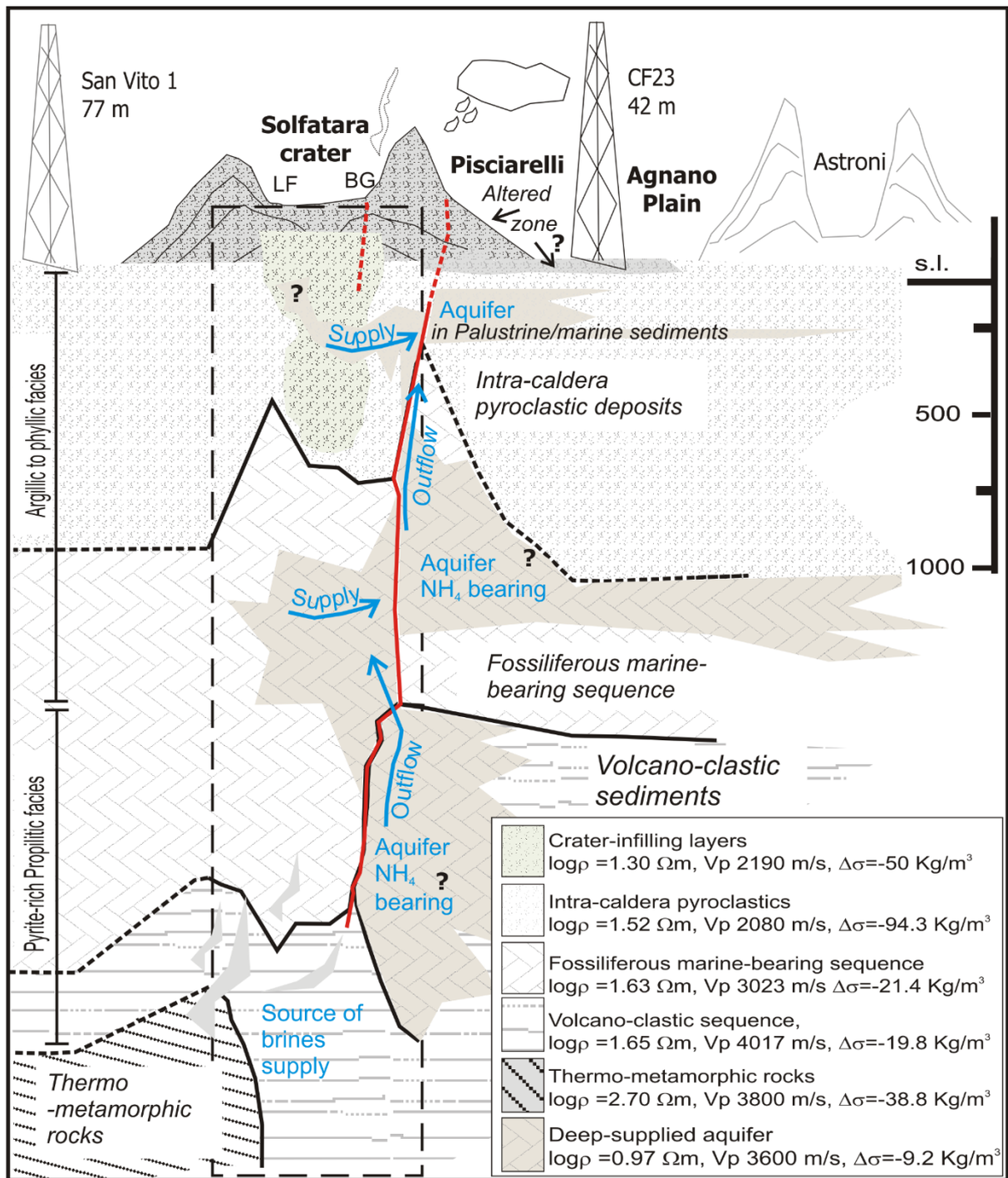


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Phase	Composition	Location
Alunite	$KAl_3(SO_4)_2(OH)_6$	Ubiquitarius
Alunogen	$Al_2(SO_4)_3 \cdot 17(H_2O)$	L1, SSt, L30, CIN, L100, L3, L70, L60, G
Alum-(K)	$KAl(SO_4)_2 \cdot 12(H_2O)$	L1, L20, PP1
Alum-(Na)	$NaAl(SO_4)_2 \cdot 12(H_2O)$	L50, New P, L20
Amarillite	$NaFe^{3+}(SO_4)_2 \cdot 6(H_2O)$	L1
Biotite	$K(Mg, Fe^{++})_3[AlSi_3O_{10}(OH, F)_2]$	MS, L3
Chabazite	$(Ca_{0.5}, Na, K)_4[Al_4Si_8O_{24}] \cdot 12H_2O$	L50, L30
Clairite	$(NH_4)_2Fe^{3+}_3(SO_4)_4(OH)_3 \cdot 3(H_2O)$	G
Coquimbite	$Fe^{+++}_2(SO_4)_3 \cdot 9(H_2O)$	L50, L3, L1
Gypsum	$CaSO_4 \cdot 2(H_2O)$	L50, L20, L3, L60, CIN, L100, SSt
Halotrichite	$Fe^{++}Al_2(SO_4)_4 \cdot 22(H_2O)$	G
Hexahydrate	$Mg(SO_4) \cdot 6(H_2O)$	CIN
Hematite	$Fe_2O_3$	PINT, L3
Kaolinite	$Al_2Si_2O_5(OH)_4$	New P, PINT, L19, L20
Koktaite	$(NH_4)_2Ca(SO_4)_2 \cdot (H_2O)$	L3, G
Illite	$(K, H_3O)(Al, Mg, Fe)_2(Si, Al)_4O_{10}[(OH)_2, (H_2O)]$	widespread at Pisciarelli (L20, L3, L50, G), CIN, rare at Solfatara
Jarosite	$KFe^{+++}_3(SO_4)_2(OH)_6$	L50, G, CIN, L30
Letovicite	$(NH_4)_3H(SO_4)_2$	L3, L1, L20
Marialite	$Na_4Al_3Si_9O_{24}Cl$	L3
Mascagnite	$(NH_4)_2SO_4$	L1, G, L20, L3, L60, New P, BG
Melanterite	$Fe^{++}SO_4 \cdot 7(H_2O)$	L3
Mereiterite	$K_2Fe^{2+}(SO_4)_2 \cdot 4(H_2O)$	L1
Minamiite	$Na_{0.6}Ca_{0.3}K_{0.1}Al_3(SO_4)_2(OH)_6$	New P
Mohrite	$(NH_4)_2Fe_2^+(SO_4)_2 \cdot 6(H_2O)$	L3
Montmorillonite	$(Na, Ca)_{0.3}(Al, Mg)_2Si_4O_{10}(OH)_2 \cdot n(H_2O)$	L50, L3, L20, CIN, SSt, G
Natroalunite	$NaAl_3(SO_4)_2(OH)_6$	L60
Periclase	$MgO$	SSt
Pickeringite	$MgAl_2(SO_4)_4 \cdot 22(H_2O)$	L1
Picropharmacolite	$Ca_4Mg(AsO_3OH)_2(AsO_4)_2 \cdot 11(H_2O)$	CIN
Pyrite	$FeS_2$	L3, BG, ASA, L1, G, New P, MS, L20, viadotto
Realgar	$As_4S_4$	BG, BN
Rostite	$AlSO_4(OH, F) \cdot 5(H_2O)$	SSt,
Salammoniac	$(NH_4)Cl$	BG, BN
Sulfur	$S^0, S_8, \text{beta}$	Ubiquitarius, subordinated in L3
Silica essentially as opal and quartz	$SiO_2$	SSt, L1, BG, CIN, L20, L50
Tamarugite	$NaAl(SO_4)_2 \cdot 6(H_2O)$	L20, CIN
Tschermigite	$(NH_4)Al(SO_4)_2 \cdot 12(H_2O)$	L20, L3, L60, L70, G
Vermiculite	$(Mg, Fe^{++}, Al)_3(Al, Si)_4O_{10}(OH)_2 \cdot 4(H_2O)$	L3
Voltaite	$K_2Fe^{2+}_5Fe^{3+}_3Al(SO_4)_{12} \cdot 18(H_2O)$	L1vent, L100, G
Zaherite	$Al_{12}(SO_4)_5(OH)_{26} \cdot 20(H_2O)$	G, CIN, L100

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Type	Sample	$\delta^{34}\text{S}$ [‰]	st. dev.	$\delta^{18}\text{O}$ [‰]	st. dev.	Date	Location*	T (°C)	Height (m)'
<b>Pisciarelli</b>									
<u>Sulfate</u>	L1d1 al/11-15	-1.78	0.06	11.93	0.23	1.11.15	L1	-	67
	L1d2 white/11-15	-2.08	0.05	15.05	0.44	1.11.15	L1	-	67
	<i>MP/11-15</i>	-1.93	0.01	10.33	0.00	1.11.15	L3	-	66
	P PP1/11-15	-0.30	0.21	17.25	0.08	1.11.15	G	92	68
	P PPb/11-15	-1.27	0.13	17.85	0.41	1.11.15	G	50.2	69
	P L50 yellow	-0.22	0.15	20.88	0.16	1.11.15	L50	-	84
	P L50 white	-0.40	0.02	21.54	0.27	1.11.15	L50	-	84
	P L50 red	-1.31	0.02	8.59	0.40	1.11.15	L50	-	84
	L1v PGw/6-16	-2.78	0.07	13.09	0.43	1.6.16	L50	-	66.5
	L1 Pwh/6-16	-2.13	0.00	5.72	0.30	1.6.16	L1	56.9	67
	L1 Psalt/6-16	-2.12	0.13	5.95	0.19	1.6.16	L1	67	67
	L1 Pblack/6-16	-0.41	0.04	9.52	0.21	1.6.16	L1	-	67
	<i>MP/6-16</i>	-0.87	0.01	4.60	0.05	1.6.16	L3	70	66
	GnvW bl/6-16	0.44	0.01	6.96	0.51	1.6.16	L19	74.7	68
	Gnv W be/6-16	-0.31	0.23	12.44	0.22	1.6.16	L19	74.7	69
	PL20 v1/6-16	-0.59	0.06	14.04	0.15	1.6.16	L20	90	71
	<i>Ps 7-16</i>	-1.00	0.08	8.37	0.06	1.7.16	L3	-	66
	PL 20V1 7-16	-0.53	0.02	14.26	0.03	1.7.16	L20	-	71
	<i>MP 29.6.17</i>	-0.09	0.03	6.82	0.2	29.6.17	L3	-	66
	L1 beije 18.9.17	-0.89	0.02	22.14	0.51	18.9.17	L1	-	67
	<i>MP L3 bocchetta 1.9.17</i>	-0.54	0.02	9.87	0.5	1.9.17	L3	49.5	66
	<i>MP 18.1.18</i>	0.26	0.06	13.21	0.2	18.1.18	L3	77.1	66
<u>Sulfide</u>	<i>MP/11-15</i>	-0.08	0.05			1.11.15	L3	-	66
	P PP1/11-15	-0.69	0.07			1.11.15	G	92	68
	L1 Pblack/6-16	-3.00	0.07			1.6.16	L1	-	67
	<i>MP/6-16</i>	-0.43	0.13			1.6.16	L3	70	66
	GnvW bl/6-16	-0.48	0.10			1.6.16	L19	74.7	68
	<i>Ps 7-16</i>	-0.34	0.01			1.7.16	L3	-	66
	<i>MP 29.6.17</i>	0.02	0.02			29.6.17	L3	-	66
	<i>MP L3 bocchetta 1.9.17</i>	-0.49	0.02			1.9.17	L3	49.5	66
	<i>MP 18.1.18</i>	-0.67	0.04			18.1.18	L3	77.1	66
<u>Sulfur</u>	L1 Pv/6-16	0.80	0.14			1.6.16	L1	56.9	66.5
	Geiser mud	-2.26	0.04			1.6.16	G	-	68
	PINT S 18.9.17	-1.71	0.06			18.9.17	PINT	-	74
	L1 S 18.9.17	-0.06	0.00			18.9.17	L1	-	67
	L 20 camino 18.9.17	-2.50	0.02			18.9.17	L20	-	76
	L1 vent S 14.12.17	-2.67	0.01			14.12.17	L1	94.7	67
	PINT S 18.1.18	-0.91	0.12			18.1.18	PINT	93.4	74
	L1 vent S 18.1.18	-2.68	0.00			18.1.18	L1	94	67
	L1 vent parete S 18.1.18	-0.99	0.05			18.1.18	L1	89.9	67.5
	L19 Geiser S 18.1.18	-2.30	0.01			18.1.18	L19	95.8	70
<b>Solfatara</b>									
<u>Sulfate</u>	SStb/11-15	-0.74	0.02	23.93	0.68	1.11.15	SSt	-	100
	SSt wr/11-15	1.61	0.11	31.33	0.47	1.11.15	SSt	-	100
	SStgc/11-15	0.91	nd			1.11.15	SSt	-	100
	SMO S/6-16	2.09	nd			1.6.16	SMO	87	109
	SMO ASA/6-16	-2.09	0.10	8.18	0.11	1.6.16	SMO	-	109
	ASA m/16-6	-0.63	0.09	8.75	0.12	1.6.16	ASA	-	124
	ASA h/16-6	-0.67	0.05	6.62	0.25	1.6.16	ASA	-	124
	SSt sub/16-6	-1.52	0.11	9.59	0.30	1.6.16	SSt	92	101
	<i>PS/6-16</i>	0.31	0.23	7.43	0.24	1.6.16	LaFangaia	52.9	94

	<i>MS 29.6.17</i>	0.80	0.00	18.58	0.3	29.6.17	<i>LaFangaia</i>	-	94
	<i>MS new 29.06.17</i>	-0.65	0.13	9.31	0.4	29.6.17	<i>new pool</i>	-	94
	<i>MS new 1.9.17</i>	-0.22	0.07	10.56	0.7	1.9.17	<i>new pool</i>	70	94
	<i>MS2 1.9.17</i>	0.62	0.09	7.84	0.1	1.9.17	<i>LaFangaia</i>	49.5	94
<u>Sulfide</u>	<i>SSt wr/11-15</i>	-0.38	0.08			1.11.15	<i>SSt</i>	-	100
	<i>SMO ASA/6-16</i>	-2.92	0.08			1.6.16	<i>SMO</i>	-	109
	<i>ASA m/16-6</i>	-2.65	0.08			1.6.16	<i>ASA</i>	-	124
	<i>ASA h/16-6</i>	-1.96	0.09			1.6.16	<i>ASA</i>	-	124
	<i>PS/6-16</i>	-0.11	0.07			1.6.16	<i>LaFangaia</i>	-	94
	<i>MS 29.6.17</i>	0.24	0.11			29.6.17	<i>LaFangaia</i>	-	94
	<i>MS new 29.06.17</i>	-1.63	0.02			29.6.17	<i>new pool</i>	-	94
	<i>MS new 1.9.17</i>	-1.16	0.10			1.9.17	<i>new pool</i>	70	94
	<i>MS1 1.9.17</i>	0.38	0.00			1.9.17	<i>LaFangaia</i>	49.5	94
	<i>MS2 1.9.17</i>	0.49	0.00			1.9.17	<i>LaFangaia</i>	49.5	94
<u>Sulfur</u>	<i>BG pg/11-15</i>	-3.95	0.03			1.11.15	<i>BG</i>	93.1	103
	<i>SStgf/11-15</i>	-2.23	0.21			1.11.15	<i>SSt</i>	-	100
		-2.13	0.28			1.11.15	<i>SSt</i>	-	100
		-2.40	0.20			1.11.15	<i>SSt</i>	-	100
	<i>SStgc/11-15</i>	-3.34	0.38			1.11.15	<i>SSt</i>	-	100
		-1.52	0.00			1.11.15	<i>SSt</i>	-	100
		-2.43	0.09			1.11.15	<i>SSt</i>	-	100
	<i>BG S/6-16</i>	-4.42	0.29			1.6.16	<i>BG</i>	93.2	103
		-3.89	0.28			1.6.16	<i>BG</i>	93.2	103
		-4.21	0.21			1.6.16	<i>BG</i>	93.2	103
	<i>up BG S/6-16</i>	-3.84	0.06			1.6.16	<i>BUCO</i>	93.2	106
		-2.78	0.11			1.6.16	<i>BUCO</i>	93.2	106
		-3.46	0.15			1.6.16	<i>BUCO</i>	93.2	106
	<i>SMO S/6-16</i>	0.29	0.28			1.6.16	<i>SMO</i>	87	109
		0.44	0.24			1.6.16	<i>SMO</i>	87	109
		0.22	0.04			1.6.16	<i>SMO</i>	87	109
	<i>SSt Sf/16-6</i>	-2.08	0.17			1.6.16	<i>SSt</i>	-	124
		-2.31	0.06			1.6.16	<i>SSt</i>	-	124
		-2.42	0.01			1.6.16	<i>SSt</i>	-	124

Table 3 – Summary of the mineralogical and isotopical features at the acid sulfate area following Rye et al. (1992); Hedenquist and Lowerstern (1994). \*highest  $^{34}\text{S}$  and  $^{18}\text{O}$  during bacteriogenic reduction of sulfates with maximum fractionation in dry-wet alternating conditions.  $\delta^{34}\text{S}$  reflects the  $\text{H}_2\text{S}/\text{SO}_2$  and temperature of fluid. 1 always present, 2 may be associated. 3 from Valentino and Stanzione (2003; 2004), Gresse et al. (2017). ‘halloysite is indicated in Montanaro et al. (2017) and included here for completeness.

	Supergene $\text{H}_2\text{SO}_4$ production by sulphides oxidation	Steam-heated $\text{H}_2\text{SO}_4$ production by $\text{H}_2\text{S}$ oxidation	Magmatic hydrothermal $\text{H}_2\text{SO}_4$ production by $\text{SO}_2$	Magmatic steam	Solfatara crater	Pisciarelli
		LOW SULFIDATION	HIGH SULFIDATION			
Alunite texture age vs original rocks	Very thin-cryptocrystalline, younger	Powdery coeval	up to 250 $\mu\text{m}$ coeval	rare up to 250 $\mu\text{m}$ in vein coeval	thin to sub mm-sized	thin to sub mm-sized
Kaolinite	disordered	well-formed	-	-	well-crystallized	well-crystallized
Halloysite	yes	possible	-	-	odd'	-
Montmorillonite	not indicate	present	in medial zones	not present	rare	present
Illite	not indicated	diffuse	in medial and outer zones	not present	rare	widespread
Gossan	at the top	-	-	-	-	possibly locally
Sinter	-	widely	-	-	widely	widely
Vuggy silica	-	-	at the core	-	not present	not present
Al solubility	-	-	highly leached	-	possibly not intense to absent	possibly not intense to absent
Sulfides - Pyrites	widespread	may be	disseminated pyrites	rare	rare	abundant pyrites
Pyrophyllite, diaspore, covellite, enargite, lauzonite	-	-	may and must be	rare	not detected	not detected
$\text{PO}_4$ in alunite	-	-	yes	unknown	not detected	not detected
S isotope fractionation	nearly absent equilibria fractionation* rarely present in sulphide rich-rocks	similar to $\text{H}_2\text{S}$	equilibria fractionation, $\text{H}_2\text{S}$ -Sulfates and Sulfate-pyrite equilibria <sup>^</sup>	equilibria fractionation with $\text{SO}_2$	Al- $\text{H}_2\text{S}$ no equilibrium (Fig. 11)	Al- $\text{H}_2\text{S}$ no equilibrium Py-Al no equilibrium Py- $\text{H}_2\text{S}$ nearly equilibrium (Fig. 11)
O isotope in Alunite	nearly absent equilibria fractionation*	equilibria fractionation	equilibria fractionation <sup>^</sup>	equilibria fractionation with $\text{SO}_2$	nearly equilibria fractionation (Fig. 11)	equilibria fractionation (Fig. 11)
pH	-	Nearly neutral	<2	-	Generally acid, locally at very low acid, locally neutral <sup>3</sup>	Locally/timely acid <sup>3</sup>
Fumarole T	-	100-to-160°C	>200°C	>200°C	Locally >160°C	Generally $\leq 110^\circ\text{C}$
Metal	-	$\text{Au}^1$ (Ag, Pb-Zn) <sup>2</sup>	(Au, Ag, Cu) <sup>1</sup> (Hg, W, Bi, Pb, Zn) <sup>2</sup>		Locally As, Hg (Fig. 9)	Hg (Fig. 9)
$\text{H}_2\text{S}/\text{SO}_4$	-	-	4±2	<1	<1	1 or higher
S state	-	Low, $\text{S}^2$	High, $\text{S}^{+4}$	High, $\text{S}^{+4}$	<1	1 or higher