Research Article

Geothermal Fluid Variation Recorded by Banded Ca-Carbonate Veins in a Fault-Related, Fissure Ridge-Type Travertine Depositional System (Iano, southern Tuscany, Italy)

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Banded Ca-carbonate veins in travertine deposits are efficient recorders of the compositional fluctuations of geothermal fluids flowing (or flowed) from deep reservoirs up to the surface, within fault zones. In this view, these veins represent key tools for decoding those factors that influenced the geochemical variations. We have analyzed veins developed in fractures channeling geothermal fluids forming travertine deposits. The studied veins cut a fossil travertine fissure ridge, near the Larderello geothermal area (Iano area, southern Tuscany) where geothermal fluid circulation is favored by NE-trending strike-to-oblique-slip faults and their intersections with NW-trending normal ones. U-Th dating indicates that fluid circulation occurred from (at least) 172 ka to 21 ka. In this time span, the geothermal fluid changed in composition, and the banded Ca-carbonate veins recorded these variations in terms of mineralogical and stable isotope composition and temperature (T) of deposition. We also documented for the first time the occurrence of Mn-rich black tree-shaped structures within the veins. Mineralogy coupled with stable and clumped isotope measurements allows the reconstruction of some features (i.e., crystal texture, temperature, and CO₂ origin) and the inference of the processes (i.e., pH, T, and pCO₂ variations) that have controlled the fluid evolution through time. Multiple-stage and one-stage deposition processes have played an important role in modifying the stable isotope composition of banded Ca-carbonate veins; temperature coupled with pCO₂ also influenced their mineralogical composition. Interpreted in the context of the tectonic setting, the data show that the NW-trending faults have mainly controlled travertine deposition. Their intersection with NE-trending faults, interpreted as transfer faults, highlights the important role of transfer zones in channeling the geothermal fluids.
1. Introduction

Faults and associated fractures are discontinuities of the upper crust able to control the circulation of geothermal fluids, thus favoring combined convective-advective heat propagation processes [1]. If geothermal fluids are stored in carbonate reservoirs, or interact with carbonate rocks, fluid composition is enriched in Ca$^{2+}$ and CO$_3^-$, while the saline content depends on the amount of dissolved CO$_2$, fluid temperature, and pH [2–8]. If fluids reach the surface through permeable faults and their damaged volumes, thermal springs develop [9], depositing masses of CaCO$_3$ (i.e., travertine deposits; [5]) mainly due to the CO$_2$ leakage [5, 8, 10, 11]. In these settings, the rise of geothermal fluids and travertine deposition are assumed to be contemporaneous with the activity of faults channeling the fluids [3, 12–15]. Thus, travertine deposits can act as tectonic markers in geothermal areas [16–20] and potentially provide key geochronologic constraints since carbonate deposits can be dated using different radiometric methods [7, 13, 15, 21].

The interplay between fault activity and travertine deposition was addressed by several authors in the last decades [3, 16, 18, 19, 22–25]. Travertine fissure ridges [26] are considered the most useful deposits for neotectonic investigations in geothermal areas. A fissure ridge-type travertine deposit consists of a ridge made of bedded travertine, from gently to steeply dipping away from the apical part, where the central fissure is located, corresponding to the fault trace intersecting the substratum. From such a fissure, the geothermal fluids flow out, and travertine deposition takes place, growing the ridge.

The relationships among fluid flow, travertine deposits, and tectonic activity have been discussed by several authors, especially focusing on fissure ridges (e.g., [3, 12, 18, 19, 27–31]). Variations in the chemistry and isotope composition of the travertine forming the ridge and, especially, of the banded Ca-carbonate veins (i.e., banded travertine in [22]) filling the fissure can be linked to different processes influencing hydrogeochemistry, including climate [8, 32], volcanism [33], fluid-rock interaction, and tectonics [3, 7, 34, 35], with particular concern on the palaeoseismicity (cf. [8, 13, 15]). This attempt encounters several difficulties, because there are many parameters that influence the geochemical characteristics of this kind of carbonate deposit. In particular, the tectonic activity controls the permeability of the fault rocks and, as a consequence, the hydrothermal fluid pathways within the crust. Such tectonic effects likely exert stronger control than climatic effects on hydrothermal fluid property variations within hypogene conduits, such as faults and fractures. In order to constrain the possible impact of the tectonic activity on the geochemical and isotopic composition of the geothermal fluids, we analyzed banded Ca-carbonate veins formed in a late Pleistocene fissure ridge-type travertine depositional system cropping out nearby the Larderello geothermal area (southern Tuscany, Italy; [36]). This is an ideal region for this kind of study, since it records a long evolution of the geothermal fluid flow, veins, and related travertine deposits that are exposed in the area (Iano area, Figure 1) located in the shoulder of a broad Neogene structural depression (known as the Volterra Basin, [37, 38]). Here, late Quaternary tectonics gave rise to a shallow hydrothermal system and travertine deposition and still ensures active thermal springs. The analyzed veins formed during several episodes show different fabrics and colors suggesting compositional changes through time. We discuss the variation of the mineralogy (i.e., aragonite and calcite) and stable isotopic composition in relation with the tectonic activity. Furthermore, we document the occurrence of Mn-rich black three-shaped structures forming some bands within the banded Ca-carbonate veins. We interpret these observations as evidence of change in geothermal fluid properties, possibly induced by the tectonic activity, promoting a variation in the fluid-rock interaction.

2. Geological Setting

The study area is located in a sector of the inner Northern Apennines, about 20 km North of the Larderello geothermal area (Figure 1(b)) and in the eastern shoulder of a wide, NNW-SSE oriented Neogene structural depression (Volterra Basin, [37, 38]). The Northern Apennines originated from the convergence and collision (late Cretaceous–early Miocene) between the Adria promontory of the African plate and the European plate represented by the Sardinia-Corsica massif (i.e., [39], and references therein). This process gave rise to the stacking of several tectonic units deriving from different palaeogeographic domains [40]. From the top, these units are [41] (a) the Ligurian units, derived from the Ligurian-Piedmont Domain, and consisting of remnants of Jurassic oceanic crust and its late Jurassic-Cretaceous, mainly clayey and sedimentary cover; (b) the sub-Ligurian units (sub-Ligurian domain), made up of Cretaceous-Oligocene turbidites; and (c) the Tuscan units forming a duplex system and composed of HP metamorphic and sedimentary units ranging from Palaeozoic to early Miocene in age [41–45]. After nappe stacking, an eastward migrating extension (e.g., [39, 46–53]) affected the inner Northern Apennines (i.e., northern Tyrrhenian Sea and Tuscany), since early–middle Miocene [48, 54–58]. Extensional tectonics thinned the previously thickened lithosphere and continental crust, up to 50 and 20-25 km, respectively [59]. This geological process was a consequence of a mainly eastward dipping normal fault system [48, 60–62] that produced (a) the lateral segmentation of the more competent levels within the previously stacked tectonic units [63], (b) the direct superimposition of the Ligurian units on the late Triassic evaporite and/or on the Palaeozoic phyllite, both representing regional detachment levels [61, 62], and (c) the development of bowl-shaped tectonic depressions where Miocene continental and marine sedimentation occurred [60, 61, 64] and superimposed Pliocene-Pleistocene basins bounded by faults which dissected the previous extensional detachments [41, 46, 62, 65–68].

These Neogene-Quaternary basins, including the Volterra Basin, were filled by Pliocene-Quaternary marine and continental sediments [37, 52]. Their deposition was controlled by the activity of NNW-SSE-striking normal faults and coeval NE-trending transfer zones [69, 70], along which
most of the late Miocene-Pleistocene magmatic activity is concentrated [24, 68, 71–75]. Geothermal fluid circulation in the whole inner Northern Apennines also appears to be controlled by transfer zones [19, 75–78]. The Iano area is located within a transfer zone (Figure 1(b)) dissecting a NNW-SSE trending normal fault system that bounds the Volterra Basin to the east. The interplay between the two fault systems (i.e., transfer and normal faults) gave rise to favorable conditions for hydrothermal circulation and thermal springs occurrence (Figure 2). Travertine deposits, some of which have been quarried, developed near the thermal springs. Deposition is still active in the surroundings of the

Figure 1: (a) Structural map of the Northern Apennines highlighting the main transfer zones from [19] and (b) geological map of the Larderello geothermal area and Volterra Basin. The study area is also indicated. Redrawn after Brogi et al.[134] and Liotta and Brogi [19].
Iano village where low temperature thermal springs \((T = 22^\circ C)\) are still discharging Ca(Mg)-HCO_3 fluids with high content of Sr and SO_4 [11].

The Iano area was investigated during Hg and coal mining exploration since eighteenth and nineteenth centuries, and detailed information about its geological setting can be found in Barberi [79] and Costantini et al. [80].

3. Methods

We conducted detailed description and analysis of the banded Ca-carbonate veins crossing travertine deposits, to investigate their (i) geometrical setting in relation to the main structures, (ii) mineralogy and textural features, (iii) stable- and clumped isotopic compositions related to the fluid geochemistry and temperature, and (iv) depositional ages as inferred by U-Th geochronology.

3.1. Geometrical Setting, Mineralogy and Textural Features. A detailed study of travertine stratigraphic setting and its structural features was carried out applying the classical methods of sedimentary and structural geology. The collected data were interpreted in the context of the area’s tectonic setting, which we have represented in a new geological map at 1:10 000 scale (Figure 2). This map was produced via integration of preexisting geological maps (geological map of Tuscany 1:10.000 scale and geological map of [80]) and new field data, covering an area of about 11 km^2. Structural and kinematic data were collected in favorable exposures through structural stations, as indicated in Figure 2. The attitude of
banded Ca-carbonate veins was also analyzed and compared with the fault striking.

To visualize textural and mineralogical changes, a petrographic investigation of the samples was performed by polarized light optical microscopy on thin sections from several samples of banded Ca-carbonate veins using a Zeiss Axiopline polarizing microscope. Based on this investigation, three key samples, labelled PAM1, PAM13 and ET3, were selected for SEM-EDS, X-ray powder diffraction, micro-Raman analyses, stable isotope measurements, and U-Th dating. Several aliquots of one sample (IA1) were selected for clumped isotope thermometry.

Scanning electron microscope (SEM) observations and microanalysis (EDS) of selected bands were performed at the Department of Earth and GeoEnvironmental Sciences (University of Bari), using a ZEISS EVO 50XVP instrument, coupled with energy dispersive X-ray Spectroscopy (EDS, Oxford-Link). Selected areas were observed on thin sections of samples PAM1 and PAM13 after graphite sputtering coating (30 nm thick). Elemental distributions of Mn were also mapped.

Mineralogical identification of PAM1 and PAM13 samples was determined by means of X-ray powder diffraction using an Empyrean, Malvern Panalytical Diffractometer at the Department of Earth and Geo-Environmental Sciences (University of Bari). X-ray powder diffractogram (XRPD) data were collected in the 6-80 2θ range using Ni-filtered CuKα radiation obtained at 40 kV and 40 mA. For this purpose, polished sample fragments and randomly oriented powder mounts were used. The analysis of the powder XRPD patterns was carried out using Highscore Plus software. The lattice parameter determination was carried out by means of Rietveld refinement using the software BGMN [81]. In addition to XRPD analysis, confocal Raman microscopy was used for a more detailed mineralogical characterization. Micro-Raman spectra were recorded with a LabRAM HR Evolution Horiba spectrometer at the Department of Chemistry (University of Bari). This spectrometer is equipped with an Ultra-Low Frequency (ULF) notch filter and an electrically cooled CCD detector. Raman spectra were excited by a He/Ne (λ = 632.8 nm) laser source using a 50× long working distance objective. The instrument was calibrated using the 520 cm⁻¹ line of a Si wafer. Raman spectra were collected on polished thin section of PAM1 and PAM13 samples by using LabSpec6-Horiba software and reworked by using Origin software. Repeated acquisitions were accumulated in the 100-3800 cm⁻¹ range to improve the signal-to-noise ratio. Micro-Raman spectra were recorded on different colored bands for each sample. Spectral scanning was also performed in the sample areas where black tree-shaped structures were clearly identified.

3.2. Vein Isotopic Composition and Geochronology. Stable carbon and oxygen isotope measurements were performed on single bands composing each vein (PAM1, PAM13, and ET3). Analyses on microdrilled powders were carried out at the Institute for Geological and Geochemical Research, Research Centre for Astronomy and Earth Sciences (Budapest, Hungary). Powdered sample analyses were carried out using the continuous flow technique with the H₃PO₄ digestion method [82]. ¹³C/¹²C and ¹⁸O/¹⁶O ratios were measured in CO₂ gases using a Finnigan Thermo delta plus XP mass spectrometer equipped with an automated GasBench II. The results are expressed in δ notation [δ = (R1/R2 - 1) × 1000, where R1 is the ¹³C/¹²C or ¹⁸O/¹⁶O ratio of the sample, and R2 is the corresponding ratio of the Vienna Pee Dee Bellemnite (V-PDB) standard] in parts per thousand. Isotopic compositions are expressed in the standard δ notation in parts per thousand (%) relative to V-PDB (δ¹³C and δ¹⁸O) and Vienna Standard Mean Ocean Water (V-SMOW; δ¹⁸O). Reproducibility was better than ±0.1‰ for carbonate δ¹³C and δ¹⁸O.

Clumped isotope thermometry [83–85] was applied to determine the precipitation temperature of carbonate minerals in order to reconstruct the palaeotemperature of the hydrothermal fluids channeled through the faults. The powdered sample (IA1) was analyzed at the University of Washington’s IsoLab, Seattle, WA, USA, using the procedures of Burgener et al. [86]. For each replicate analysis, 6-8 mg of carbonate powder was digested in a common acid bath at 90°C, and the resulting sample CO₂ was purified cryogenically using an automated system. Measurements were performed using a Thermo Scientific MAT 253 mass spectrometer with dual inlet configured to measure m/z 44, 45, 46, 47, 48, and 49. Data were corrected following Schauer et al. [87], using the ¹⁷O interference correction parameters recommended by Brand et al. [88] as well as a pressure baseline correction [89] to calculate δ¹³C, δ¹⁸O, and Δε. The carbonate calibration data of Petersen et al. [90], which is the most up-to-date available for carbonates of different mineralogies and sample types (synthetic, natural, or biogenic). This calibration includes the calibration dataset of Kele et al. [92], which is travertine/tufa-specific, and the synthetic carbonate calibration data of Kelson et al. [93], which was produced in IsoLab using the same methods as the sample data presented here.

The carbonate clumped isotope values measured at ATOMKI (Institute for Nuclear Research, Debrecen, Hungary) were performed using Thermo Scientific Kiel IV Automatic Carbonate Device coupled to a Thermo Scientific MAT253 Plus isotope ratio mass spectrometer. The measurement system was installed in 2018 in Isotope Climatology and Environmental Research Center (ICER). Each carbonate sample consisted of 10-12 replicate analyses of 100 µg aliquots. ETH1, ETH2, and ETH3 were used as normalization
4. Results

In the study area, several patches of fossil travertine deposits are exposed. In a few cases, deposition is still active. These deposits unconformably overlie Neogene-Quaternary and pre-Neogene units (i.e., Ligurian Units).

In the following sections, we illustrate the main stratigraphic and structural features of the travertine substratum. Then, the main stratigraphic and sedimentological features of the travertine deposits are reported, followed by a detailed description of the textural, mineralogical, geochemical and isotopic features of the banded Ca-carbonate veins, crossing the travertine deposits and filling fractures.

4.1. The Travertine Substratum. The travertine substratum (Figure 2) is formed by units stacked during the Northern Apennines evolution and orogenic belt building.

These are from the bottom (i) Palaeozoic-Mesozoic metamorphic rocks (Monticiano-Roccastrada Unit, CP, VER, and TCC in Figure 2); (ii) late Triassic-Oligocene sedimentary rocks (Tuscan Nappe, Ev, Di, Sc, and Mac in Figure 2); (iii) Cretaceous sandy-marl turbiditic succession (external Ligurian Unit); and (iv) Jurassic-Cretaceous ophiolite succession overlain by the marine shale and limestone cover (inner Ligurian Unit, Jc, and APA, respectively, in Figure 2).

The contractional tectonics that gave rise to the orogenic belt produced superimposed structures already described in several papers [80, 99, 100]. For the goals of this work, we focus on the Neogene-Quaternary extensional tectonics, which overprinted all the previously developed structures. Thus, low-angle normal faults (LANFs) caused the direct superimposition of the successions belonging to the uppermost unit (inner Ligurian Unit) either above the metamorphic rocks of the lowermost unit (Monticiano-Roccastrada Unit) or above the evaporite level forming the basal part of the Tuscan Nappe (Figure 2). This setting implies the lateral segmentation of several geological bodies as well as the partial or complete tectonic omission of the previously stacked units. Later, high-angle normal to oblique-slip faults, NW- and NE-striking, dissected the LANFs and determined the eastern shoulder of the Volterra Basin, where the study area is located (Figure 2).

The Neogene sediments are mostly made up of sand, clay and sandy-clay with some conglomerate embedded at different levels. These sediments were deposited in the eastern margin of the Volterra Basin, recording its tectonic evolution, as documented in detail by Brogi and Liotta [61].

Travertine deposits are mainly located along the NE- and NW-striking faults, overall at their intersection, which juxtaposes the Neogene sediments with the pre-Neogene units (e.g., nearby Iano village, Figure 2). In the southern part of the study area (e.g., near Il Palagio locality, Figure 2), the travertine deposits are in part still active and appear to be mainly related to NW-striking faults. NE- and NW-striking faults dissect Zanclean-Piacenzian sediments and, in a few cases, the travertine deposits, suggesting Quaternary activity, as discussed below.

The major faults are never exposed but their occurrence can be inferred by NE- and NW-striking scarpata recognized across the whole area. Nevertheless, secondary faults are exposed in abandoned quarries or along road fresh cuts. These fault zones are characterized by apparent offsets of few meters and consist of damaged rock volumes exceeding 30 m in thickness with pervasive subvertical fractures and minor faults (Figures 3(a) and 3(b)). In some cases, hydrothermal alteration and/or localized mineralization (Figure 3) characterize fault planes and their damage zones. Fault slip planes are surrounded by fault cores, up to 10 cm thick, formed by brecciated rocks which, in some cases, show yellowish coatings as an effect of the circulation of previous geothermal fluids (Figure 3(a)).

Structural and kinematic data were collected in structural stations (Figure 2), in damaged rock volumes exposed in abandoned quarries where exposures and lithotypes are favorable to record and preserve striation on the slip surfaces.
The results (Figure 4) indicate that the orientation of the analyzed mesofaults is in agreement with the regional-scale structures (Figure 2), thus displaying dominant NW- and NE-striking fault systems. Kinematic indicators mainly consist of mechanical striations (Figures 3(c) and 3(d)), chatter marks, lunate structures, tool marker and, in some cases, quartz and calcite slickensides, often accompanied by Fe-hydroxides crusts and patinas (Figures 3(e) and 3(f)). Kinematic indicators on the NE-striking faults show right- and left-lateral strike to oblique-slip movements, as well as normal ones (Figure 4). Their chronological relation was not defined, although it is likely that the normal movements are probably the last ones, based on chronologies that have been already described in the surrounding areas (cf. [68]). The NW-striking faults show dominant normal component, though left- and right-lateral oblique-slip movements have been locally observed (Figure 4).

These measurements from faults were used to derive the palaeostress field, assuming that the measured slickenlines are parallel to the main shear stress. Figure 4 shows the fault plane solutions obtained for each fault trend and their dominant kinematics. For the NE-striking faults, the maximum (compressional) and minimum axes are close to horizontal, while the intermediate axis is close to vertical, during the
right and left-lateral movement, which we hypothesize that occurred first. In contrast, the maximum axis is vertical, and the intermediate and minimum axes are subhorizontal during the later movement. Concerning the NW-striking faults, all axes are oblique.

4.2. The Travertine Depositional System. Travertine deposits consist of several bodies less than 1 km² each, distributed in two different areas: (i) the Iano village area and its northern sector and (ii) Il Palagio area and its western sector (Figure 2).

The travertine patches of the Iano area are fossil deposits. These bodies are partly dismantled by human activity and covered by thick vegetation, and their original morphology cannot be reconstructed with complete confidence. Here, the maximum thickness of the travertine does not exceed 15 m. According to the classification proposed by Capezzuoli et al. [101], travertine beds consist of laminated abiotic crystalline crusts (Figure 5(a)), microbial mats, and micritic beds. Deposits often encrust bushes and other plants (Figures 5(b) and 5(c)), suggesting a slope depositional system that passed to distal zones by subhorizontal morphology and low-energy
environments. Facies distribution and bedding attitudes suggest cone-shaped mounds, indicating the location of the palaeothermal springs at their top. In this scenario, the travertine deposits were fed by thermal springs which deposited laminated abiotic crystalline crusts, interrupted by small ponds of microbial mats, and formed where fluids run along slopes, close to the palaeothermal springs. In the distal parts, fluids fed palustrine environments, where travertine deposited in vegetated areas suggest the cooling of the flowing waters.

The travertine patches in the II Palagio area consist of two main bodies (Figures 2 and 6(a)). The smaller is only a few hundred square meters wide and consists of a partly dismantled fossil deposit, developed in the surroundings of a thermal water discharge area. The larger body is characterized by fossil deposits (Figure 6(b)) as well as an active thermal spring, causing present travertine deposition (Figure 5(d)) in its southern portion. From the active springs, a $22^\circ C$ water is flowing, with a flow rate minor than 1 liter per second [102]. The water is characterized by $\mathrm{pH} = 6.30$ and high TDS values (6665 mg/L), Ca(Mg)-HCO$_3^-$, and high Sr and SO$_4^{2-}$ contents, accompanied by CO$_2$ and H$_2$S leakage [11].

The fossil deposits were quarried and today, the abandoned saw cut walls are covered by thick vegetation. The fossil travertine body is defined by NW-trending fissure ridge completely filled by banded Ca-carbonate veins, showing the locations of samples PAM13 and IA1. (c) Schematic reconstruction of the travertine fissure ridge with the location of the collected samples.
4.3. Banded Ca-Carbonate Veins. Veins consist of onyx-like mm-sized crystalline bands in white, black, and brown. Bands are vertical or subvertical, with local dm-sized botryoids (Figure 7(a)). The biggest vein (up to 2.10 m thick) is NW-trending and fills the central part of the ridge (Figures 6(b) and 6(c)), representing the main feeder conduit. Thinner veins (up to 10 cm thick) laterally cross the flank of the ridge (Figures 6(c) and 7(b)) mostly in the NW direction, accounting for secondary conduits. The veins are arranged in parallel bands, locally with internal unconformities cross-cutting the previously formed bands; the unconformable contact relationships range from parallel to angular, thus recording an interruption in the crystallization. Some veins include black bands that look like tree-shaped structures (Figure 7(d)) characterized by groups of branches in a white matrix. Their shape is very similar to the black “shrubs” identified by previous authors in travertine deposits in several localities such as Morocco (Africa), Bagni di Tivoli (Italy), Mammoth Hot Springs (Yellowstone National Park, USA) and Hierapolis (Turkey), [103–105]. Their thickness ranges from submillimeter to a few centimeters, forming thin laminae between vein bands (Figure 8(a)).

Each band is thought to correspond to a crack-and-seal event ([17], and references therein). Crack episodes mostly caused reopening of previously formed fractures. The new fracture developed parallel to slightly oblique with respect to the previously deposited bands, thus giving rise to a neoformational mm- to cm-thick, geometrical irregular, syn-taxial veins (Figure 7(c)). In some cases, the geothermal fluid permeated the mechanical discontinuities of the bedded travertine, thus producing cm-thick syntaxial vein packages extending for meters, parallel to the bedded travertine strata (Figure 5(b)). These strata, mainly formed by abiogenic crystalline crusts and made up of feather-like crystals and bedded porous laminae, are gently to steeply dipping (from 5° to 50°).

We selected for further analysis key samples representing the main features of two different vein systems: (i) banded veins formed within the main fissure ridge feeder conduit (samples PAM13 and IA1 Figures 6(c), 7(a), and 9(a)) and (ii) banded veins crossing the flank (samples PAM1, ET3, Figures 6(c), 7(b), 8(a), and 10(d)). These latter recorded at least three superposed depositional events (V1, V2, V3), represented by crosscutting veins (Figure 8(a)). In particular, sample PAM1 recorded all three events (Figure 8(a)) and also includes black tree-shaped structures; in contrast, sample ET3 recorded V1 and V2 events (Figure 10(d)).

The sample location is indicated in Figures 6(a) and 6(c) and described in Table 1.

4.3.1. Mineralogy of Banded Ca-Carbonate Veins. Vein V1 of PAM1 (Figure 8(a)) and PAM13 (Figure 9(a)) samples was analyzed by XRD to identify the carbonate minerals. XRPD analysis reveals that the V1 diffractogram (Figure 8(b)) can be indexed to two carbonate phases: (i) aragonite, with the unit cell parameters of $a = 4.9579(1), b = 7.9778(2), c =$
Figure 8: Continued.
and low-intensity reflections located at 26.6° and 44.6° (indicated with the "?" symbol in Figure 8(b)), likely due to a poorly crystalline impurity phase that could not be indexed with calcite or aragonite phases. The cell parameters of calcite are slightly smaller than those present in the Joint Committee on Powder Diffraction Standards (JCPDS 47-1443), while the aragonite unit cell parameters deviate slightly from those for pure aragonite causing a small enlargement of cell volume (JCPDS 41-1475). Although calcite has a rhombohedral unit cell and aragonite is orthorhombic, both structures show similar Ca lattice but different CO$_3$$_2$ ion arrangements. Carbonate structures can accommodate the trace element by exchange of Ca with Mg, Mn, Fe, or Sr, which is governed by the flexibility of the Ca site surrounding structure. The substitution causes dilation or contraction of the cell volume. Therefore, the small variation of cell volumes is associated with small amounts of larger and smaller sized ions substituting for Ca in the structure of aragonite and calcite, respectively [106–108]. To substantiate XRPD results, SEM/EDX analysis of calcite particles detected Mg, Ca, C, and O, whilst SEM/EDX of aragonite crystals detected Sr, Ca, and O. Confo cal Raman analyses also support XRPD results. Indeed, the Raman spectra (Figure 8(i), points 2 and 3) of vein V1 (PAM1 sample) showed the typical vibration modes $\nu_1$ (1085 cm$^{-1}$ calcite and 1086 cm$^{-1}$ aragonite) and $\nu_4$ (711 cm$^{-1}$ calcite and 705 cm$^{-1}$ aragonite) for the CO$_3$$_2$ group in a carbonate crystal lattice. Additional lattice mode peaks were located at 156 cm$^{-1}$, 282 cm$^{-1}$ (calcite), and 207 cm$^{-1}$ (aragonite). In addition, we used EDS to determine various elements at spots forming the black tree-shaped phase. The elemental mapping (Figure 8(f)) shows high intensities for manganese oxide, varying between 77.87% and 83.22%. Micro-Raman spectra were also acquired in the sample PAM1, where black spots were clearly identified. All the Raman spectra show a strong wide band at about 620 cm$^{-1}$ with shoulders at about 492 and 530 cm$^{-1}$ (Figure 8(j), point 5). The spectra show the characteristic bands ascribed to natural manganese dioxide minerals [109]. Based on SEM/EDX and Raman analyses, the black material appears to consist of poorly defined manganese minerals (Figures 8(c) and 8(e)). Minor percentages of MgO, FeO, and NiO have been measured in the black deposits, with values around 2%, 1% (with some values higher than 5%), and 5%, respectively (see Table 2 for data).

For sample PAM13 (Figure 9(a)), the X-ray diffraction pattern (Figure 9(c)) reveals that the carbonate bands are made up predominantly of aragonite. Also, in this case, the cell parameters calculated from diffraction peaks ($a = 4.9637(4)$, $b = 7.9715(7)$, $c = 5.7449(5)$ Å, and space group $Pmcn$; and (ii) calcite, with the unit cell parameters $a = 4.9713(1)$, $c = 17.017(3)$ Å, and space group $R-3c$. The diffraction pattern also shows broad and low-intensity reflections located at 26.6° and 44.6° (indicated with the "?" symbol in Figure 8(b)), likely due to a poorly crystalline impurity phase that could not be indexed with calcite or aragonite phases. The cell parameters of calcite are slightly smaller than those present in the Joint Committee on Powder Diffraction Standards (JCPDS 47-1443), while the aragonite unit cell parameters deviate slightly from those for pure aragonite causing a small enlargement of cell volume (JCPDS 41-1475). Although calcite has a rhombohedral unit cell and aragonite is orthorhombic, both structures show similar Ca lattice but different CO$_3$$_2$ ion arrangements. Carbonate structures can accommodate the trace element by exchange of Ca with Mg, Mn, Fe, or Sr, which is governed by the flexibility of the Ca site surrounding structure. The substitution causes dilation or contraction of the cell volume. Therefore, the small variation of cell volumes is associated with small amounts of larger and smaller sized ions substituting for Ca in the structure of aragonite and calcite, respectively [106–108]. To substantiate XRPD results, SEM/EDX analysis of calcite particles detected Mg, Ca, C, and O, whilst SEM/EDX of aragonite crystals detected Sr, Ca, and O. Confo cal Raman analyses also support XRPD results. Indeed, the Raman spectra (Figure 8(i), points 2 and 3) of vein V1 (PAM1 sample) showed the typical vibration modes $\nu_1$ (1085 cm$^{-1}$ calcite and 1086 cm$^{-1}$ aragonite) and $\nu_4$ (711 cm$^{-1}$ calcite and 705 cm$^{-1}$ aragonite) for the CO$_3$$_2$ group in a carbonate crystal lattice. Additional lattice mode peaks were located at 156 cm$^{-1}$, 282 cm$^{-1}$ (calcite), and 207 cm$^{-1}$ (aragonite). In addition, we used EDS to determine various elements at spots forming the black tree-shaped phase. The elemental mapping (Figure 8(f)) shows high intensities for manganese oxide, varying between 77.87% and 83.22%. Micro-Raman spectra were also acquired in the sample PAM1, where black spots were clearly identified. All the Raman spectra show a strong wide band at about 620 cm$^{-1}$ with shoulders at about 492 and 530 cm$^{-1}$ (Figure 8(j), point 5). The spectra show the characteristic bands ascribed to natural manganese dioxide minerals [109]. Based on SEM/EDX and Raman analyses, the black material appears to consist of poorly defined manganese minerals (Figures 8(c) and 8(e)). Minor percentages of MgO, FeO, and NiO have been measured in the black deposits, with values around 2%, 1% (with some values higher than 5%), and 5%, respectively (see Table 2 for data).

For sample PAM13 (Figure 9(a)), the X-ray diffraction pattern (Figure 9(c)) reveals that the carbonate bands are made up predominantly of aragonite. Also, in this case, the cell parameters calculated from diffraction peaks ($a = 4.9637(4)$, $b = 7.9715(7)$, $c = 5.7449(5)$) indicate a slight dilation of the cell volume with respect to the ideal pure aragonite phase. Raman analyses (Figure 9(b)) confirm that the sample PAM13 is made up of aragonite, showing the typical aragonite peaks of the lattice modes located at 154 cm$^{-1}$, 207 cm$^{-1}$, and 1086 cm$^{-1}$.
4.3.2. Textural Features and Age of the Banded Ca-Carbonate Veins. Vein V1 from sample PAM1 (Figure 8(a)) is mostly composed of mm-thick bands formed by elongate-blocky crystals of calcite or aragonite. The most external band (i.e., the primary formed band) is characterized by the presence of a 5 mm-thick band, made up of black material simulating tree-shaped structures (Figures 7(d) and 8(a)), in a white CaCO₃ matrix with scattered Mn-oxides (cfr. 4.3.1 section). In some cases, the black material is organized in micrometer-thick laminae (Figures 8(a) and 8(d)) made up of multilayered, rounded or subrounded shapes (Figure 8(e)).

In the polished hand specimen of sample PAM1, calcite bands appear translucent and defined by different shades of gray. Calcite crystals consist of elongate-blocky crystals (Figure 10(a)), 150-300 µm wide, and from few millimeters to 5 mm long. Crystals enlarge in the growth direction, and their terminations are characterized by pointed curved faces (Figure 10(a)), suggesting that the crystals had the time to develop. Under crossed polars, these crystals show wavy extinction. Some calcite bands include black points, aligned perpendicularly to the growth direction (Figure 10(b)). These are made up of Mn-oxides, determining a dark color to the band, at the specimen scale (see Figure 8(a)).

Aragonite crystals confer a white, near opaque aspect to the bands in the polished hand specimen. The acicular aragonite crystals are made up of needle-like crystals extending along their c-axis direction (Figure 8(g)). Calcite bands appear slightly darker gray on the backscattered images compared to the aragonite bands (Figure 10(c)). This is because calcite bands contain small percentages of Mg (3.25%). In contrast, aragonite bands do not contain Mg, but small percentages of Sr (3.08%), as also confirmed by the mineralogical analyses (cf. 4.3.1 section).

Vein V2 shows a syntaxial fabric developed perpendicular to the V1 band. This crystal texture is well represented in sample ET3 (Figure 10(d)). Millimeter-sized voids are in the central part of the vein (Figure 10(e)); the lateral bands are made of small elongate-blocky calcite crystals (about 10 µm in length) growing toward the central part of the vein (i.e., voids, in Figure 10(e)); dark hiatuses trace the edges of the elongate-blocky crystals (Figure 10(e)).

Vein V3 developed parallel to the V1 and interrupted the continuity of the V2 (Figure 8(a)). The bands forming the
vein V3 do not exceed 2 mm in thickness and are composed of acicular crystals of aragonite, showing a well-ordered texture (Figure 10(f)). The crystals are densely packed forming thin fans of ray crystals that develop from an apical point. Their ending faces form pointed terminations, highlighting that some crystals grew faster than others. Mn-oxide-rich lamina retrace the ending crystal faces (Figure 8(h)), separating vein V1 from vein V3.

Similar features are observed in sample PAM13, formed by parallel whitish bands (Figures 7(a) and 9(a)). This vein is completely made of aragonite crystals. Aragonite bands are made up of elongated polycrystals ranging in length from...
0.2 to 1.3 cm and resulting from the packing of needle thin crystals (Figure 10(g)). The latter shows wavy extinction caused by the systematic variation of the c-axis orientation with respect to the substratum of needle-like crystals [110] and very high interference colors (Figure 10(g)).

Two samples were selected for U-Th dating [95]: (i) V1 and V3 from sample PAM1, in order to get information on the time interval between the different veining events and (ii) sample PAM13. All samples are representative of a single band. The resulting data are summarized in Table 3. These highlight that V3 and V1 developed in a restricted time span encompassed between $26 \pm 1$ and $172 \pm 3$ thousand years ago (ka, before 1950 AD). In contrast, a radiometric age ranging between $21_{-1}^{+1}$ and $27_{-1}^{+1}$ ka was computed for the sample PAM13.

### 4.3.3. Stable and Clumped Isotope Analyses

The results of the $\delta^{13}C$ and $\delta^{18}O$ analyses of the veins (i.e., PAM1, ET3, and PAM13) indicate values ranging from -1.2‰ to 4.2‰ (V-PDB) and from 24.3‰ to 26.8‰ (V-SMOW), respectively. These ranges of values can be ascribed to the group of thermogenic travertine (see [5]).

The dataset is reported in Table 4 and displayed in the diagram of Figure 11(a).

Vein 1 from PAM1 ($n = 12$) and ET3 ($n = 10$) samples (Figures 8(a) and 10(d)) is characterized by $\delta^{13}C$ and $\delta^{18}O$ values ranging from -1.2‰ to 1.5‰ (V-PDB) and 24.3‰ to 26.3‰ (V-SMOW), respectively. $\delta^{13}C$ and $\delta^{18}O$ values of vein V2 ($n = 3$) range from -0.1‰ to 1.4‰ (V-PDB) and 25.6‰ to 26.0‰ (V-SMOW), respectively. V1 data are consistent with those from vein V2, suggesting constant fluid properties during fracturing. However, although the $\delta^{18}O$ data of vein V3 ($n = 3$) are similar to V1 and V2, $\delta^{13}C$ data show different values, ranging from 2.5‰ to 4.2‰ (V-PDB).

Similarly, the vein from the main feeder conduit of the fissure ridge (PAM13, $n = 3$) is characterized by $\delta^{13}C$ and $\delta^{18}O$ values ranging from 2.9‰ to 4.2‰ (V-PDB) and from 25.7‰ to 26.8‰ (V-SMOW), respectively. On the basis of $\delta^{13}C$ values, data can be separated in two distinct groups. The veins crossing the flank of the fissure ridge (Figure 6(c)) are characterized by $\delta^{13}C$ values with respect to those defining the vein from the feeder conduit. $\delta^{18}O$ values are very similar for the two vein types, although PAM13 and V3 samples show slightly higher values than those from V1 and V2 veins. Stable isotope values of vein V3 fall between the two groups (Figure 11(a)).

Clumped isotope composition ($\Delta_{47}$) was measured on a band of aragonitic sample (IA1) from the central part of the
### Table 3: Uranium isotopic composition and U-Th ages for banded Ca-carbonate veins.

| Sample ID | Weight, g | $^{238}$U $10^{-9}$ g/g | $^{232}$Th $10^{-12}$ g/g | $\delta^{234}$U measured$^a$ | $^{230}$Th/$^{238}$U activity$^c$ | Age (yr ago) uncorrected | Age (yr BP) relative to 1950 AD$^d$ | $\delta^{234}$U$_{initial}$ Corrected$^b$ |
|-----------|-----------|------------------------|------------------------|------------------|-----------------------|------------------|------------------|------------------|
| PAM1-1    | 0.0511    | 678.59 ± 0.73          | 64066 ± 407            | 318.2 ± 1.8      | 0.3004 ± 0.0040       | 52.46 ± 0.77     | 27.902 ± 421    | 25.957 ± 1033    | 342.5 ± 2.2      |
| PAM1-2    | 0.0514    | 39.489 ± 0.084         | 2246 ± 10              | 373.1 ± 5.3      | 1.1557 ± 0.0076       | 335.1 ± 2.6      | 173.372 ± 3063  | 172.362 ± 3073   | 607 ± 10         |
| PAM13-2   | 0.0490    | 681.25 ± 0.79          | 4253 ± 15              | 342.4 ± 1.9      | 0.2814 ± 0.0016       | 743.1 ± 4.9      | 25.395 ± 165    | 25.205 ± 176     | 367.7 ± 1.6      |
| PAM13-4   | 0.0463    | 342.10 ± 0.35          | 5864 ± 18              | 324.5 ± 1.7      | 0.2397 ± 0.0022       | 230.5 ± 2.2      | 21.591 ± 220    | 21.184 ± 277     | 344.5 ± 2.4      |
| PAM13-1   | 0.0397    | 436.63 ± 0.46          | 930 ± 12               | 353.8 ± 1.6      | 0.30169 ± 0.00090    | 2336 ± 30        | 27.188 ± 99     | 27.077 ± 101     | 382.0 ± 1.7      |

Analytical errors are 2$\sigma$ of the mean. $^a$$^{238}$U = $^{235}$U × 137.818 (±0.6%) (Hiess et al., 2012); $\delta^{234}$U = ($^{234}$U/$^{238}$U$_{activity}$ − 1) × 1000. $^b$$\delta^{234}$U$_{initial}$ corrected was calculated based on $^{230}$Th age (T), i.e., $\delta^{234}$U$_{initial} = \delta^{234}$U$_{measured} \times e^{234 \times \lambda_{234} \times T}$, and T is corrected age. $^c$$^{230}$Th/$^{238}$U$_{activity} = 1 - e^{-\lambda_{230}T} + (\delta^{234}$U$_{measured}/1000)[\lambda_{234}/(\lambda_{234} - \lambda_{230})][(1 - e^{-(\lambda_{230} - \lambda_{234})T})]$, where T is the age. Decay constants are 9.1705 × 10$^{-8}$ yr$^{-1}$ for $^{230}$Th, 2.8221 × 10$^{-9}$ yr$^{-1}$ for $^{234}$U (Hiess et al., 2012). $^d$Age corrections relative to 1950 AD, were calculated using an estimated atomic $^{230}$Th/$^{232}$Th ratio of 4 (±2) × 10$^{-6}$. 

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### Table 4: Results of the stable isotopes analyses.

| Sample  | Mineralogy                  | δ¹³C (‰, V-PDB) | δ¹⁸O (‰, V-PDB) | δ¹⁸O (‰, V-SMOW) |
|---------|-----------------------------|-----------------|-----------------|------------------|
| PAM1_16 | Aragonite                   | 4.2             | -4.4            | 26.3             |
| PAM1_15 | Aragonite                   | 2.5             | -5.9            | 24.8             |
| PAM1_14 | Aragonite                   | 3.6             | -5.1            | 25.7             |
| PAM1_13 | Mn-oxide lamina             | -1.2            | -5.6            | 25.2             |
| PAM1_12 | Calcite                     | 0.0             | -5.8            | 24.9             |
| PAM1_11 | Calcite                     | 0.0             | -5.8            | 24.9             |
| PAM1_10 | Aragonite                   | 1.5             | -6.0            | 24.7             |
| PAM1_9  | Aragonite                   | 1.2             | -6.1            | 24.6             |
| PAM1_8  | Calcite                     | -0.1            | -6.4            | 24.3             |
| PAM1_7  | Blocky/chaotic              | 0.9             | -6.0            | 24.7             |
| PAM1_6  | Calcite                     | 0.1             | -6.0            | 24.7             |
| PAM1_5  | Calcite                     | -0.2            | -5.7            | 25.0             |
| PAM1_4  | Aragonite+calcite           | 1.5             | -5.8            | 24.9             |
| PAM1_3  | Calcite                     | 0.1             | -5.6            | 25.1             |
| PAM1_2  | Calcite                     | -0.3            | -5.7            | 25.0             |
| PAM1_1  | Tree-shaped structures      | -0.4            | -5.4            | 25.3             |
| PAM13_1_8 | Aragonite                   | 3.2             | -5.0            | 25.7             |
| PAM13_1_7 | Aragonite                   | 3.6             | -5.0            | 25.8             |
| PAM13_1_6 | Aragonite                   | 3.4             | -4.7            | 26.1             |
| PAM13_1_5 | Aragonite                   | 3.8             | -4.9            | 25.9             |
| PAM13_1_4 | Aragonite                   | 3.3             | -4.8            | 25.9             |
| PAM13_1_3 | Aragonite                   | 3.2             | -4.6            | 26.2             |
| PAM13_1_2 | Aragonite                   | 3.8             | -4.2            | 26.5             |
| PAM13_1_1 | Aragonite                   | 3.6             | -4.6            | 26.2             |
| PAM13_2_1 | Aragonite                   | 3.2             | -4.7            | 26.0             |
| PAM13_2_2 | Aragonite                   | 3.1             | -4.5            | 26.2             |
| PAM13_2_3 | Aragonite                   | 3.5             | -4.4            | 26.4             |
| PAM13_2_4 | Aragonite                   | 4.0             | -4.0            | 26.8             |
| PAM13_2_5 | Aragonite                   | 3.9             | -4.0            | 26.8             |
| PAM13_2_6 | Aragonite                   | 4.2             | -4.2            | 26.6             |
| PAM13_2_7 | Aragonite                   | 3.8             | -4.3            | 26.4             |
| PAM13_4_1 | Aragonite                   | 3.6             | -4.7            | 26.1             |
| PAM13_4_2 | Aragonite                   | 3.0             | -5.1            | 25.7             |
| PAM13_4_3 | Aragonite                   | 3.2             | -5.0            | 25.7             |
| PAM13_4_4 | Aragonite                   | 3.8             | -4.5            | 26.3             |
| PAM13_4_5 | Aragonite                   | 3.4             | -4.8            | 26.0             |
| PAM13_4_6 | Aragonite                   | 3.5             | -4.3            | 26.5             |
| PAM13_4_7 | Aragonite                   | 3.6             | -4.7            | 26.1             |
| PAM13_4_8 | Aragonite                   | 3.6             | -4.8            | 25.9             |
| PAM13_4_9 | Aragonite                   | 3.9             | -4.4            | 26.3             |
| PAM13_4_10 | Aragonite                  | 4.1             | -4.3            | 26.5             |
| PAM13_4_11 | Aragonite                  | 4.1             | -4.6            | 26.1             |
| PAM13_4_12 | Aragonite                  | 3.8             | -4.7            | 26.1             |
| PAM13_4_13 | Aragonite                  | 4.0             | -4.3            | 26.5             |
| PAM13_4_14 | Aragonite                  | 4.0             | -4.6            | 26.2             |
| ET-3_1 | Tree-shaped structures      | -0.3            | -5.6            | 25.1             |
| ET-3_2 | Chaotic                     | 0.1             | -5.7            | 25.0             |
| ET-3_3 | Aragonite                   | -0.2            | -5.6            | 25.1             |
ridge (in the proximity of PAM13 location), in order to reconstruct the palaeotemperature of the fluids that deposited the Ca-carbonate veins. The calculation gave a temperature of 67°C ± 6°C. Clumped isotope composition was determined at ATOMKI in order to calculate palaeotemperature on two other bands of the IA1 sample, which gave values of 84°C ± 5°C and 111°C ± 5°C, respectively. The data are reported in Table 5. In addition, we also report the temperature values calculated using Kele et al.’s [92] calibration for these last two values, for the reasons explained in Section 3.2. The calculation gave temperatures of 68° ± 4°C and 85° ± 3°C, which are more suitable for the geothermal system of

| Sample | Mineralogy | δ¹³C (‰, V-PDB) | δ¹⁸O (‰, V-PDB) | δ¹⁸O (‰, V-SMOW) |
|--------|------------|----------------|----------------|-----------------|
| ET-3_4 | Chaotic    | -0.2           | -5.8           | 24.9            |
| ET-3_5 | Chatic     | -0.2           | -6.2           | 24.5            |
| ET-3_6 | Calcite    | 0.0            | -6.4           | 24.4            |
| ET-3_7 | Calcite    | 0.0            | -5.8           | 24.9            |
| ET-3_8 | Aragonite  | 0.9            | -5.4           | 25.3            |
| ET-3_9 | Calcite    | 1.2            | -5.1           | 25.6            |
| ET-3_10| Calcite    | 1.4            | -5.0           | 25.8            |
| ET-3_11| Calcite    | -0.1           | -4.8           | 26.0            |
| ET-3_12| Aragonite  | -0.5           | -4.5           | 26.3            |
| ET-3_13| Calcite    | -0.6           | -5.7           | 25.0            |
| IA1_W  | Aragonite  | 2.5            | -5.9           | 24.9            |
| IA2_D  | Aragonite  | 3.2            | -5.3           | 25.5            |

**Figure 11:** (a) Graph showing the distribution of stable isotope data for the analyzed samples. Blue rhombus refers to the PAM13 data; yellow square refers to the vein V1 data from PAM1 and ET3; the symbol “x” refers to the vein V2 from ET3; pale blue square refers to the vein V3 data from the PAM1 sample. (b) PAM1 isotope data distinguished on the basis of their mineralogy. See the good correlation ($R^2 = 0.8527$) between stable isotopes of aragonite crystals.

**Table 5: Results of the clumped isotopes analyses.**

| Sample | Replicates | $\delta^{13}C$ (VPDB, ‰) | $\delta^{13}C$ std dev | $\delta^{18}O$ (VPDB, ‰) | $\delta^{18}O$ std dev | $\Delta_{47}$ (average, CDES25 ‰) | $\Delta_{47}$ std error (‰) | $\Delta_{47}$ std error based on long-term std dev of carbonate standards (‰) | $T$ (°C) cal. with Petersen et al. [91] ±C |
|--------|------------|----------------|----------------|----------------|----------------|----------------|----------------|--------------------------------------------------------|---------------------------------|
| IA 1   | 2          | 2.819          | 0.050          | -5.231         | 0.017          | 0.589          | 0.003          | 0.012                                                 | 67 ± 6                          |
| IA1_d  | 17         | 3.230          | 0.050          | -5.260         | 0.090          | 0.558          | 0.008          | 0.006                                                 | 84 ± 5                          |
| IA1_w  | 14         | 2.660          | 0.050          | -5.750         | 0.090          | 0.518          | 0.007          | 0.006                                                 | 111 ± 5                         |
the study area. These values partially confirm the temperature value computed at the University of Washington. For this reason, we consider a fluid temperature range between 67°C ± 6°C and 85°C ± 3°C.

5. Discussion

The discussion focuses on three main points: (i) fluid origin, (ii) tectonic control on the fluid flow, and (iii) role of geochemistry in reconstructing the geothermal fluid property variations and the possible triggering factors.

5.1. Fluid Origin. Geothermal fluids that formed the travertine deposits interacted with carbonate and sulfate rocks, as suggested by their high potential to deposit Ca-carbonate and by their Sr content [11]. This implies that the geothermal fluids flowed within the late Triassic evaporite and Jurassic carbonate located at the base of the Tuscan Nappe [36, 111]. It is known that in the neighboring Larderello area, the geothermal fluids are mostly of meteoric origin (>95%) with a minor inflow of thermometamorphic and magmatic fluids [112–114]. It is therefore reasonable to assume that the parent fluids of the travertine would be related to meteoric waters, too.

Regarding the presence of CO₂, this can be the result of different processes [115]: (i) mantle degassing [116, 117], (ii) limestone dissolution [118], or (iii) hydrothermal metamorphism of limestone [119]. On the basis of δ¹³C CO₂, the potential source for CO₂ can be discerned: (a) -7‰ < δ¹³C CO₂ < -4‰ (V-PDB) suggests mantle degassing [120], (b) δ¹³C CO₂ = -7‰ (V-PDB) indicates dissolved CO₂ in rainwater in equilibrium with atmospheric CO₂ [121], (c) δ¹³C CO₂ < -20‰ (V-PDB) drives at biogenic CO₂ [122], and (d) -1‰ < δ¹³C CO₂ < +2‰ (V-PDB) suggests CO₂ from hydrothermal processes [123].

In our case, we computed δ¹³C CO₂ values from the carbonate δ¹³C values of the study veins, using the empirical equation (δ¹³C CO₂ = 1.2δ¹³C trav - 10.5) of Panichi and Tongiorgi [118], resulting from the comparison between the Italian thermal springs and associated travertine deposits. The δ¹³C CO₂ calculations for V1 and V2 veins give values ranging from -12‰ to -9‰ (V-PDB). In contrast, the computed δ¹³C CO₂ values for V3 vein and for banded veins from the central fissure of the ridge are slightly higher, ranging from -7‰ to -5‰ (V-PDB). Interestingly, the computed δ¹³C CO₂ values are significantly lower than most of the CO₂ gas emissions in central Italy, which vary from -2‰ to +2‰ (V-PDB, [124]).

As previously said, banded Ca-carbonate veins are the results of crack-and-seal events, likely triggered by the tectonic activity. This process could account for the variation of pCO₂ (probably due to tectonic activity) during the multistep precipitation of banded Ca-carbonate veins, from fluids that migrated from depth to the surface. According to Minissale et al. [115], this process could produce a large isotopic fractionation, causing the deposition of banded Ca-carbonate veins that are enriched in ¹³C at depth. As a consequence, the rising fluid, depleted in ¹³C, could deposit banded Ca-carbonate veins close to the surface, resulting in lower δ¹³C values, as we observe in V1 and V2 from PAM1 and ET3 samples. In contrast, δ¹³C CO₂ calculations for V3 and PAM13 overlap with the δ¹³C CO₂ values recorded in the northern part of the Larderello geothermal area, where the study area is located (Figure 1). Here, most of the gas emissions are characterized by δ¹³C CO₂ ranging between -7‰ and -4‰ (V-PDB), with few values lower than -7‰ (V-PDB), [115]. These values have been interpreted as the results of primary CO₂ rising up from the mantle [116] or, alternatively, a mixing between isotopically heavy and light CO₂ coming from the geothermal reservoir and organic sediments produced in the Neogene basins, respectively [124]. Therefore, V3 and PAM13 veins could be the result of a one-stage deposition from a fluid directly coming from depth.

Temperature of the hydrothermal fluid circulated within the main fissure of the travertine ridge was determined by clumped isotope thermometry to be 67°C ± 6°C for the band analyzed at the University of Washington and 68°C ± 4°C and 85°C ± 3°C for the bands analyzed at ATOMKI. These values exceed the temperature of the modern fluids (22°C, [11, 125]) of the active thermal spring located in the Il Palagio area (Figure 2), confirming that the temperature of the fluid has changed through time.

We calculated the δ¹⁸O values from the measured carbonate δ¹³C O values, using the clumped isotope derived temperature for IA1 and the Kim et al. [126] and Chacko and Deines [127] equations for the temperature dependence of aragonite-water oxygen isotope fractionation, assuming equilibrium conditions during aragonite deposition. The results using the two equations are indistinguishable (2.6‰ V-SMOW). The δ¹⁸Ow values are out of the range of present-day regional rainwater δ¹⁸Ow values (δ¹⁸Ow = -7 to -6‰) that feed the shallow hydrothermal system of southern Tuscany [11, 124]. This result could suggest that (a) the fluids were possibly influenced by magmatic source water (in Larderello geothermal field δ¹⁸O of magmatic waters ranges from +3‰ to +8‰, in [128]), causing a heavier composition of the O isotopes, or that (b) fossil meteoric water extensively interacted with the Tuscan Nappe carbonate rocks that are characterized by a positive signature of δ¹⁸O (±25‰, [129]).

5.2. The Tectonic Control on the Fluid Flow. Mesostructural data indicate that the fault-slip surfaces and damage zones of the two main NE- and NW-trending fault systems are often characterized by Fe-hydroxides and banded Ca-carbonate veins (Figure 3). This observation implies that both systems channeled geothermal fluids. Comparable evidence comes from the field mapping (Figure 2), which indicates that both the fossil and active travertine deposition is mainly located along the NW-striking faults, at the intersection with the NE-striking major faults. Thus, we conclude that in the study area, permeability and hydraulic conductivity of deep fluids are controlled by fractures.

In this view, it is noteworthy to underline that the main fissure ridge (from which the analyzed samples were collected) displays a NW trend, parallel to the main fault zones characterizing the whole study area. Furthermore, in the proximity of Iano village, travertine bodies are associated
with the interplay between NE- and NW-trending faults. Similar trends, parallel to the main NW-trending faults, characterize the minor banded Ca-carbonate veins cutting both the travertine bodies and units forming the travertine substratum. The main banded Ca-carbonate vein developed in the central part of the NW-trending fissure ridge (Figure 6(a)) suggests that this travertine deposit developed along a permeable sector of a NW-striking fault. At least part of the fault activity (from 172 ± 3 to 21.2 ± 0.1 ka) is recorded by the banded Ca-carbonate veins development. Similarly, a small number of thin veins are NE-oriented, such as vein V2 from PAM1 and ET3 samples. This superposed veining (V1, V2, and V3) is indicative of repeated deformation episodes, followed by progressive fracture sealing by Ca-carbonate deposition.

Kinematic indicators on the NE-trending faults indicate strike-slip movements. This is significant since, in agreement with Sibson [130], the intermediate axis attitude indicates the preferential direction of fluid flow. Thus, an oblique, near vertical, intermediate axis can explain the upflow migration of deep fluids and therefore the occurrence of CO2 of mantle origin (Figure 4). This suggests that the NE-striking fault-zones play a role in channeling upward fluid migration, as already suggested for the Larderello area [68].

Interaction with NW-trending faults is then significant for two related reasons: (a) the intersection of NW- and NE-trending faults increases fracture distribution, promoting hydraulic conductivity and (b) the intermediate axis of the NW-trending faults being close to horizontal enables, geo-hydraulic conductivity and (b) the intermediate axis of the NE-trending faults increases fracture distribution, promoting for two related reasons: (a) the intersection of NW- and with the interplay between NE- and NW-trending faults. Similar trends, parallel to the main NW-trending faults, characterize the minor banded Ca-carbonate veins cutting both the travertine bodies and units forming the travertine substratum. The main banded Ca-carbonate vein developed in the central part of the NW-trending fissure ridge (Figure 6(a)) suggests that this travertine deposit developed along a permeable sector of a NW-striking fault. At least part of the fault activity (from 172 ± 3 to 21.2 ± 0.1 ka) is recorded by the banded Ca-carbonate veins development. Similarly, a small number of thin veins are NE-oriented, such as vein V2 from PAM1 and ET3 samples. This superposed veining (V1, V2, and V3) is indicative of repeated deformation episodes, followed by progressive fracture sealing by Ca-carbonate deposition.

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Interaction with NW-trending faults is then significant for two related reasons: (a) the intersection of NW- and NE-trending faults increases fracture distribution, promoting hydraulic conductivity and (b) the intermediate axis of the NW-trending faults being close to horizontal enables, geothermal fluids to flow laterally, interacting with the host rocks (hence, enriching their content in 18O, for instance), and arriving to the surface when the dilatation component of the normal fault is efficient.

This framework of crosscutting relationships was also envisaged in similar geological contexts [19, 75, 131–133] and proposed to determine alignment of thermal springs by the tectonic activity [15, 29, 142, 152] and/or climatic variations [8, 32, 153]. Climate can control the amount of CO2 discharge and CO2 degassing. In our opinion, pulses of the tectonic activity are reasonable factors. In particular, for samples PAM1 and ET3, it can be assumed that the high CO2 degassing rate could have caused mineralogical differentiation between calcite and aragonite. This implies that the CO2 partial pressure has changed through time. This is supported by the stable isotope data of our banded Ca-carbonate veins (Figure 11(a)) which display larger δ18O variation than δ13C. In particular, aragonite bands are characterized by the highest δ13C values, suggesting that fast CO2 degassing has strongly influenced the deposition of these bands, causing disequilibrium due to the kinetic effect. In addition, δ18O and δ13C values correlate (R² = 0.85, Figure 11(b)), implying a deposition under disequilibrium conditions. In fact, during degassing, isotopically light carbon dioxide preferentially escapes and, as a consequence, the 13C and 18O contents in depositing CaCO3 increase in the same direction [5, 7, 8, 34, 115, 154–156]. For this reason, a positive
correlation is recorded. Furthermore, the texture of these crystals shows ending faces with pointed terminations (cfr. 4.3.3), implying that some crystals grew faster than others. Therefore, fast degassing could have increased the depositional rate of aragonite crystals.

In contrast, isotope values of calcite crystals of sample PAM1 do not show any correlation and δ\(^{13}\)C values are all around 0‰, with some values slightly negative. This evidence implies that the isotopic composition of fluids remained constant during calcite crystals formation, and (possibly) crystals formed under isotopic equilibrium. The texture of calcite crystals validate this hypothesis because crystals enlarge in the growth direction and their terminations are characterized by pointed curved faces, suggesting that the crystals grew slowly and had the time to develop.

Therefore, if CO\(_2\) degassing rate was a factor, it would imply that the veins record CO\(_2\) partial pressure changes through time.

In contrast, sample PAM13 is completely made up of aragonite (Figure 9(a)), and stable isotope values do not show any correlation (Figure 11(a)). Clumped isotopes highlight a depositional temperature between 67° and 85°C (Table 5), suggesting that temperature played a more important role than pCO\(_2\) in depositing aragonite, an interpretation that is also supported by the lack of correlation between stable isotope values.

Finally, we discuss the tree-shaped black structures described for the vein V1 (Figures 7(d) and 8(a)). These structures have never been described in banded Ca-carbonate veins; they have only been documented for travertine deposits [103–105, 157, 158] and speleothems [159].

From a mineralogical point of view, the black tree-shaped structures contain manganese dioxides, which are based on [MnO\(_6\)] octahedra units linked in chains to form a framework structure containing monodimensional tunnels with specific sizes. Tunnels can accommodate different chemical species, such as Mg, Cu, Ni, Co, and Fe in their ionic state [105]. For this reason, in some cases, high percentages of Ni (5.01% on average) have been measured. The Raman results suggest that the black tree-shaped structures consist of todorokite, although the possible presence of cryptomelane or hollandite cannot be ruled out. Indeed, our data are in good agreement with Bernardini et al. [109] who report for todorokite a broad and strong band at 630 cm\(^{-1}\) with shoulders at ~507 and 575 cm\(^{-1}\). However, the possible occurrence of cryptomelane or hollandite cannot be ruled out. Although spectroscopic data from the literature are conflicting, some recent studies [109] report for cryptomelane a position of the strong band centered between 623 and 655 cm\(^{-1}\) and a shoulder at ~580 cm\(^{-1}\). Similar positions are reported for hollandite.

In speleothems and travertine deposits, tree-shaped structures are strictly associated with bacterial activity inducing the precipitation of Mn-oxides. Precipitation produce aggregates with high Mn concentration (Mn-oxides), even from water with low Mn\(^{2+}\) concentration (parts per million, [103]). This process was very likely promoted by specific environmental parameters: in surficial flowing waters, pH changes are favored by temperature and CO\(_2\) degassing rate variations. In contrast, these factors cannot abruptly change in deep conduits. As a consequence, the occurrence of Mn-rich deposits is indicative of changes of the...

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**Figure 12:** Not to scale cartoon showing the conceptual model of the hydrothermal fluid circulation in the Iano area. Meteoric waters infiltrated at depth through dolostone and limestone forming the Late Triassic evaporite level that is exposed in the eastern basin shoulder. The heated fluids interacted with the evaporite rocks, increase their δ\(^{18}\)O, and mixed with mantle derived fluids channeled along the main fault zones. The same fault zones (mainly N120° striking) channeled the fluids up to the surface, where thermal springs, active since late Pleistocene, are depositing travertine bodies. Travertine fissure ridges and mounds are the main tectonometamorphic features occurring in the Iano area.
hydrothermal fluid conditions. Changes in pH within the circulating waters suggest fluid compositional variation. Values of pH above 5 [103] and oxidizing environments promote the precipitation of Mn-oxides [160]. In the case of Iano, these tree-shaped black structures are characterized by our lowest measured δ13C values (-1.2‰ and -0.4‰ V-PDB), suggesting significant mixing between hydrothermal and meteoric waters. In these conditions, microorganisms could thrive, locally modifying the pH conditions and discharging metabolic products able to oxidize Mn2+ [105, 161]. As long as these conditions persist, Mn-oxides continue to deposit due to microbial metabolic processes. As a consequence, tree-shaped black structures are indicative of a low-temperature fluid depleted in CO2. These conditions can develop after rapid CO2 degassing in waters that remain in surficial levels, favoring mixing with meteoric waters. These structures are absent in the banded Ca-carbonate veins filling the central fissure of the ridge, suggesting that the fluids in the central fissure maintained high temperature through time, favoring aragonite deposition and did not interact with surficial waters.

In this view, the tree-shaped black structures can suggest a phase of transition after a fast CO2 degassing event, during which hydrothermal waters mix with more surficial ones, causing temperature decrease. Thus, tree-shaped black structures can be useful indicators for a stasis phase of the hydrothermal fluid circulation and a mixing of hydrothermal and meteoric waters.

6. Conclusions

Banded Ca-carbonate veins represent key tools for decoding variations in geothermal fluid properties though time. Veins forming in the Iano travertine depositional system preserve compositional fluctuations of the geothermal fluids (red arrows in Figure 12) that flowed from the deep reservoir up to the surface, along fault zones (Figure 12). These fault zones are characterized by changes in their kinematics, which combined with their orthogonal arrays, and gave rise to localized permeable volumes able to channel deep fluids up to the surface, therefore mixing with shallow (i.e., meteoric) origin fluids (blue arrows in Figure 12). Fluids move following the intermediate axis along the transfer faults indicated by orange arrows in Figure 12. Our record started in the middle Pleistocene, where the fault permeability induced the geothermal fluids to flow at the surface. From 172 ± 3 ka until (at least) 27.1 ± 0.1 ka, the circulating fluids deposited banded Ca-carbonate veins following a multistep process. During its rise along faults and fractures, the fluid interacted with carbonate deposits, becoming enriched in δ18O (Figure 12); as a consequence, the stable isotopic composition of the fluid changed. Banded Ca-carbonate veins at different depths developed within faults and fractures, probably due to pulses of tectonic activity. Starting by at least 27.1 ± 0.1 ka, the fluid preserved the initial stable isotope imprint, from depth to the surface, following a one-stage deposition process.

Travertine deposits mostly formed from thermal springs aligned along NW-trending faults and at their intersection with NE-trending faults. Most of the banded-Ca carbonate veins follow the NW direction, even if in some cases, some very thin veins cut the previously deposited veins in the NE direction.

Our observations from the Iano area represent the first documented tree-shaped black structures, made up by Mn-oxides, in banded Ca-carbonate veins. The mineralogy is mostly likely todorokite and their deposition can be indicative of a stasis phase following a rapid CO2 degassing that favors the mixing of deep origin hydrothermal fluids with shallow meteoric waters. As a consequence, fluid temperature and CO2 content decrease, allowing the development of microbial communities in oxidizing and acid environment (pH > 5). Thus, Mn-rich deposits could have a key role in decoding periods of low fluid flux and mixing with more surficial waters.

Data Availability

The data support the results of this study and are reported in the Tables.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this paper.

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Supplementary Materials

The carbonate clumped isotope values measured at ATOMKI (Institute for Nuclear Research, Debrecen, Hungary) were performed using Thermo Scientific Kiel IV Automatic Carbonate Device coupled to a Thermo Scientific MAT253 Plus isotope ratio mass spectrometer (as previously said in Section 3.2). The normalization standards used were ETH1, ETH2, and
ETH3 coupled with IAEA–C2, used as quality control sample with the values reported in Table S1 [91]. $\Delta_{\text{PDB}}$ is reported in the CDES (carbon–dioxoide equivalent scale) [91], using an acid fractionation factor of -0.066‰ to set down data generated with 70°C acid digestion on the 25°C digestion scale. Easotope application [95] was used for data evaluation, using CO$_2$ with a fractionation factor of +0.066.

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