The imprint of hydrothermal fluids on trace-element contents in white mica and tourmaline from the Panasqueira W–Sn–Cu deposit, Portugal

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Abstract
White mica and tourmaline are the dominant hydrothermal alteration minerals at the world-class Panasqueira W–Sn–Cu deposit in Portugal. Thus, understanding the controls on their chemical composition helps to constrain ore formation processes at this deposit and determine their usefulness as pathfinder minerals for mineralization in general. We combine whole-rock geochemistry of altered and unaltered metasedimentary host rocks with in situ LA-ICP-MS measurements of tourmaline and white mica from the alteration halo. Principal component analysis (PCA) is used to better identify geochemical patterns and trends of hydrothermal alteration in the datasets. The hydrothermally altered metasediments are enriched in As, Sn, Cs, Li, W, F, Cu, Rb, Zn, Tl, and Pb relative to unaltered samples. In situ mineral analyses show that most of these elements preferentially partition into white mica over tourmaline (Li, Rb, Cs, Tl, W, and Sn), whereas Zn is enriched in tourmaline. White mica has distinct compositions in different settings within the deposit (greisen, vein selvages, wall rock alteration zone, late fault zone), indicating a compositional evolution with time. In contrast, tourmaline from different settings overlaps in composition, which is ascribed to a stronger dependence on host rock composition and also to the effects of chemical zoning and microinclusions affecting the LA-ICP-MS analyses. Hence, in this deposit, white mica is the better recorder of the fluid composition. The calculated trace-element contents of the Panasqueira mineralizing fluid based on the mica data and estimates of mica-fluid partition coefficients are in good agreement with previous fluid-inclusion analyses. A compilation of mica and tourmaline trace-element compositions from Panasqueira and other W–Sn deposits shows that white mica has good potential as a pathfinder mineral, with characteristically high Li, Cs, Rb, Sn, and W contents. The trace-element contents of hydrothermal tourmaline are more variable. Nevertheless, the compiled data suggest that high Sn and Li contents are distinctive for tourmaline from W–Sn deposits.

Keywords Alteration geochemistry · Tourmaline · White mica · Panasqueira · Tungsten–tin deposits · Magmatic-hydrothermal systems · Trace elements

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Introduction

The alteration assemblages associated with hydrothermal deposits typically have much larger footprints than the areas containing metals at economic grades and thus constitute important exploration guides (Lowell and Guilbert 1970; Large and McGoldrick 1998; Mathieu 2018). Identifying mineral pathfinders and developing their use for exploration require studying deposits to generate a detailed understanding of what controls the mineral composition and to what extent this reflects the chemical signature of the mineralization they are associated with. In the last decades, many studies have focused on mineral exploration tools based on fertility indicators and chemical vectoring (Kerrick and Wyman 1997; Kelley et al. 2006; Jackson 2010; Cooke et al. 2014; Wilkinson et al. 2015; Champion and Huston 2016; McClennen and Layton-Matthews 2017; Soltani Dehnavi et al. 2018; Dill 2018; Codeço et al. 2018; Uribe-Mogollon and Maher 2018; Luz et al. 2019). In that context, there is an increasing interest in using multivariate statistics and machine learning methods (Cheng et al. 2011; Yang and Cheng 2014; Makvandi et al. 2016a, 2019; Gonçalves et al. 2018; Ordóñez-Calderón and Gelcich 2018; Grunsky and de Caritat 2019; Huang et al. 2019; Gonçalves and Mateus 2019) and in developing discriminant diagrams to separate different styles of mineralization (Dupuis and Beaudoin 2011; Montreuil et al. 2013; Makvandi et al. 2016b; Fresia et al. 2017; Huang et al. 2019).

The world-class Panasqueira W–Sn–Cu deposit in central Portugal has more than 100 years of activity and is currently Europe’s second largest tungsten mine, after Felbertal, Austria (USGS 2019). The deposit consists of sub-horizontal mineralized quartz veins that are hosted by metasedimentary rocks above a late-Variscan granite and its greisen cupola (Kelly and Rye 1979; Bussink 1984; Polya et al. 2000) (Fig. 1). White mica and tourmaline are the most abundant alteration minerals throughout the deposit, occurring in both the early and late mineralization stages. Thus, these minerals are of prime interest as indicators for the composition of the mineralizing fluids and may have potential as ore pathfinders applicable to similar deposits worldwide.

Under equilibrium conditions, the composition of hydrothermal mica and tourmaline is controlled by the composition of the fluid and the partition coefficients of the elements between the mineral and fluid phase. Partition coefficients are sensitive to pressure, temperature, and element speciation in the fluid (van Hinsberg et al. 2010). Fluid composition is, in turn, sensitive to the fluid–rock ratio, with a progressively larger impact of the local host-rock composition as the fluid–rock ratio decreases. Preliminary tourmaline-fluid partition coefficients have been estimated for high-pressure subduction zone fluids (van Hinsberg et al. 2017), and tourmaline-melt partitioning was studied by van Hinsberg (2011), but no data are available at hydrothermal conditions.

Experimental studies related to mica-fluid partitioning are also limited (Bos 1990; Zhu and Sverjensky 1991; Melzer and Wunder 2001; Green and Adam 2003). Chemical variations and microstructures of mica have been used to trace the degree of differentiation and magmatic-hydrothermal processes in highly fractionated granitic and associated ore systems (Roda et al. 2007; Lichtervelde et al. 2008; Johan et al. 2012; Breiter et al. 2017, 2019). While it is debated to what degree mica can be used as a fluid tracer, recent studies suggest that the trace-element contents in mica are controlled by their presence in the fluids and by fluid-mica partition coefficients which themselves vary as a function of pressure, temperature, and chemical composition conditions (Legros et al. 2016, 2018).

In this study, we first establish the bulk chemical changes caused by hydrothermal alteration based on analyses of altered and unaltered host rocks, as well as the underlying granite and greisen. We then present in situ trace-element compositions of white mica and tourmaline from the vein selvages and the alteration zones by laser ablation ICP-MS analyses. We use principal component analysis (PCA) of the compositional data from whole rocks and minerals to help identify and interpret the geochemical effects of the hydrothermal alteration and the ability of the minerals to reflect them. The mica studied is exclusively dioctahedral but shows a range of compositions with variable muscovite–celadonite–phengite components (Codeço et al. 2019). For simplicity, we will use the terms white mica and muscovite interchangeably.

The Panasqueira W–Sn–Cu deposit

The Panasqueira deposit is located in the Iberian Massif (Fig. 1), which is part of the European Variscan Belt that formed as a result of the Laurussia–Gondwana collision through the Late Devonian (ca. 370 Ma), culminating with the closure of the Rheic Ocean (ca. 300 Ma) (Ribeiro et al. 1990; Kroner et al. 2007; Martínez Catalán et al. 2007). A large number of post-collisional granites in the Variscan Belt are highly evolved and several host important Sn–W deposits, which take various forms including veins, stocks, skarns, breccia pipes, and greisen bodies (Štěmplok 1980). The Iberian Massif contains one of the largest domains of granitoids in the Variscan Belt, among which is the 290 ± 10-Ma-old (Clark 1970) Panasqueira granite, a peraluminous, S-type, two-mica granite containing quartz, microcline, plagioclase, muscovite > biotite, apatite, ilmenite, and zircon (Kelly and Rye 1979; Neiva 1987). The Panasqueira granite does not crop out at the surface but has been intersected by drilling. The granite intruded greenschist–facies biotite–chlorite–quartz schists of the Almaceda Formation, which is part of the regional Schist–Greywacke Complex—Beiras Group (SGC-BG) of Late Ediacaran to Early Cambrian age (Fig. 1). A contact
metamorphic aureole with cordierite porphyroblasts (spotted schist) is locally developed in the mine area above the granite (Conde et al. 1971; Bussink 1984). The exposed greisen cupola of the granite (Fig. 1c) is composed of medium- to coarse-grained quartz and white mica, with minor amounts of topaz, chlorite, K-feldspar, plagioclase, pyrite, arsenopyrite, and zircon (Kelly and Rye 1979; Bussink 1984; Neiva 1987; Codeço et al. 2019).

The flat-lying quartz–wolframite ± cassiterite ± sulfides veins have been exposed by underground mining over an area of 2500-m length, 400 to 2000-m width, and at least 500-m depth (Wheeler 2016). The paragenetic sequence
of the ore veins has been divided into six stages of mineralization: quartz–tourmaline alteration, muscovite selvage formation, main oxide-silicate stage, main sulfide stage, pyrrhotite alteration stage, and late carbonate stage. Tourmaline, white mica, and quartz coexist as gangue minerals from the early stages to the main sulfide stage (Kelly and Rye 1979; Polya 1989; Foxford et al. 1991, 2000; Polya et al. 2000). Hydrothermal alteration of the wall rocks produced concentric zones with progressively greater distance from the veins, consisting of a proximal tourmaline–quartz–muscovite zone, 2 to 30-cm thick, containing more than 50 vol% of tourmaline, and a distal muscovite–quartz zone including minor tourmaline or topaz (Bussink 1984). Even though muscovite and tourmaline are present in different stages at Panasqueira, we exclusively focus on the occurrences that are related to the ore-forming event in the veins and altered host rocks at the immediate vein contact.

Quartz-hosted fluid inclusions indicate that the main oxide and sulfide stages formed from fluids with low to moderate salinities (5–10 wt% NaCl eq.) at temperatures between 230 and 360 °C, while inclusions from the late carbonate stage indicate lower temperatures (<200 °C) and fluid salinities below 5 wt% NaCl eq. (Kelly and Rye 1979; Bussink 1984; Polya et al. 2000; Jaques and Pascal 2017; Lecumberri-Sanchez et al. 2017). Geothermometry based on arsenopyrite compositions, Ti-in-quartz, and muscovite–tourmaline B–isotope exchange (Jaques and Pascal 2017; Codeço et al. 2017, 2019) indicates significantly higher temperatures ranging from 420 to 490 °C for the early stages of hydrothermal alteration and vein formation and about 250 °C for late-stage mineralization hosted in cross-cutting fault zones.

The major-element composition of Panasqueira tourmaline has been studied by several workers (Neiva et al. 2007; Codeço et al. 2017; Launay et al. 2018). Tourmaline has intermediate schorl–dravite compositions with significant excess Al in octahedral Y-sites (up to 0.9 apfu), variable X-site vacancies (0.1 to 0.6 apfu), and relatively high F contents (up to 0.7 apfu). Tourmaline grains are typically zoned and show significant increases in Fe, F, and Na and decreases in Mg, Ca, and Al from core to rim. Codeço et al. (2017) noted a trend of decreasing Al and increasing Mg/Fe with distance from the vein contact, which was attributed to the influence of the host schists, where tourmaline formed at the expense of the chlorite–biotite-rich protoliths. Codeço et al. (2019) presented major-element data of white mica and documented a range in compositions from muscovite towards the celadonite and aluminoceladonite end members. The main chemical variations in terms of major elements are in the Fe/Mg ratio and in the Al and F contents. Mica in the greisen has the highest Fe/Mg ratios (up to 10), mica from altered host rocks (tourmaline–quartz–mica zone) has the lowest values (0.5–1), and vein-selvage micas have intermediate values. The composition of mica from the greisen, most vein selvages, and altered wall rocks overlap in terms of Al and F, the exception being mica in topaz-rich selvages, which has the highest F and lowest Al contents (1–1.5 apfu F and 1.5–2 apfu Al).

Samples

Whole-rock samples

Samples for whole-rock analysis were collected from three drill cores from underground galleries (D1602, M1609, and M1610) and four drill cores from the surface (SCB-1, SCB-2, SPS-1, and SMN-1). Their locations are shown in Figure S1 of the Electronic Supplementary Material 1 (ESM 1), and further details are in ESM2 (Table S1) and Codeço et al. (2020) (Table D1). Additionally, two outcrop samples (CXG1–2, CXG-2), located respectively at 5.5 and 10.5 km distance from the deposit, were taken as examples of the regional metasediments (ESM 1; Codeço et al. 2020, Table D2). The drill cores SMN-1, SPS-1, and SCB-2, located N and NE of the current underground workings, contain unaltered metasediments consisting of light green-colored phyllite and schist, intercalated with quartzite and metagreywacke layers (Fig. 2a, b). The main minerals are white mica, biotite, chlorite, and quartz. The unaltered schist in the drill cores is very similar to samples of the country rocks collected from outcrops outside of the mine area, whose main mineralogy comprises quartz, white mica, and minor biotite, with accessory zircon and feldspar.

Drill core SCB2 intersects the Panasqueira granite at a distance of about 800 m NE from the known cupola and 315 m below the surface. The contact zone consists of pelitic, granoblastic hornfels with a thickness of about 1 m, composed of quartz, white mica, biotite, and feldspars with accessory tourmaline, cassiterite, sulfides, and chlorite (Fig. 2c). The hornfels shows a gradational transition downward through greisen, from 316 to 327 m depth, into granite (Fig. 2d). The coarse-grained, two-mica granite has white mica dominating over biotite and also contains albite, K-feldspar, and quartz (Fig. 2e), as well as accessory zircon. The granite is altered in some domains by muscovitization of feldspars and chloritization of biotite (Fig. 2f).

 Petrographic studies of the metasediments in drill cores from underground galleries confirm that tourmalinization (tourmaline–quartz–muscovite zone) only occurs around quartz veins and veinlets. The abundance of tourmaline increases towards veins and veinlets (Fig. 2g), and the thickness of the tourmaline–muscovite zone is a function of the vein size and vein density (Fig. 2h). Samples more distal to the veins
lack tourmaline and contain predominantly quartz, biotite, chlorite, and white mica (Fig. 2i).

**Mineral samples**

Samples for LA-ICP-MS analysis were chosen from a larger sample set that was described and analyzed for major-element chemical and boron isotopes in tourmaline and white mica by Codeço et al. (2017, 2019). The samples containing mica and tourmaline represent different types of vein selvages, wall rock, greisen, and fault zones, and their locations are given in ESM 1—Figure S3, ESM2—Table S1, and Codeço et al. (2020) (Table D3).

**Veins and vein selvages**

In the veins, white mica is the second-most abundant silicate mineral after quartz. There is considerable variability in the vein selvages, but most veins are bordered by mica-rich selvages (MRS) (Fig. 3a). Less common are veins with mica-poor selvages (MPS) and topaz-rich selvages (TRS). The MRS are several centimeters thick and consist of white mica, quartz, arsenopyrite, wolframite ± chalcopyrite ± pyrite ± sphalerite and locally also tourmaline, topaz, and apatite (Figs. 3a–d). At the vein contact, mica typically forms fine-grained (FG) flakes intergrown with quartz and minor tourmaline. A coarse-grained (CG) generation of white mica typically develops towards the center of the vein and is commonly intergrown with wolframite (Fig. 3b, e). The MPS are thin layers (<1 cm) at the edge of the veins consisting of quartz, topaz, tourmaline, and variable amounts of sulfide minerals (Fig. 4a). The TRS are one to several centimeter-thick layers at vein margins that contain light-green, massive topaz aggregates, partly replaced by secondary fine-grained white mica (Codeço et al. 2019). Unfortunately, the mica in TRS samples is too fine grained for LA-ICP-MS measurements. Tourmaline is not common in the veins, but it locally forms grain aggregates at the selvages or overgrowths on disrupted fragments of wall rock enclosed in the vein quartz (Fig. 3a, b). Tourmaline commonly shows optical and chemical zoning (see Fig. 3d and Codeço et al. 2017, for details).

**Wall rock alteration zones**

White mica in the wall rock alteration zones typically forms fine-grained (<1 mm) flakes associated with tourmaline, quartz, and minor Ti-(Fe)-oxides (Fig. 3f, g). Mica is less common in the alteration zones of the coarser-grained spotted schists, these zones being instead dominated by tourmaline with minor quartz. Tourmaline in the alteration zones can reach up to 70 modal % of the rock. It is typically fine grained, <0.1 to 0.3 mm (Fig. 3f), but the grain size is larger in the spotted schist (0.1 to 1 mm).

**Cross-cutting fault zones, late vugs, and greisen**

Localities with more intense alteration related to syn- to post-mineralization faulting or fault reactivation (Fig. 3a, g) typically contain coarser-grained tourmaline and mica, as well as a more complex accessory mineral assemblages with Ti-(Fe, W, Sn, Cr, Sr, V)-oxide minerals. These zones commonly contain veinlets and vugs filled by apatite and variable amounts of sulfide minerals (Fig. 3a). Zoned tourmaline crystals are also present as needles in late vugs (∆= carbonates ± quartz) at the vein margins, i.e., post-mineralization cracks or fractures (Fig. 4b). Mica is absent in these late vugs.

The greisen zone contains white mica as a major phase, where it is fine- to medium-grained (up to 2–3 mm) and shows compositional zonation, which is more prominent along cleavage planes and rims (Fig. 4c). Tourmaline was not found in any samples of the greisen in our study or in previous publications.

**Methods**

**Whole-rock geochemistry**

All samples were crushed and subsequently powdered in an agate mill to a grain size of less than 62 μm. Major-element oxides (SiO₂, TiO₂, Al₂O₃, Fe₂O₃, MnO, MgO, CaO, Na₂O, K₂O, P₂O₅) and trace elements (Ba, Cr, Ga, Nb, Ni, Rb, Sr, V, Y, Zn, Zr) were analyzed in a subset of samples by inductively coupled plasma mass spectrometry (ICP-MS) at GFZ Potsdam. A Dionex ICS-1100 Ion Chromatograph from Thermo Fischer was used to analyze fluoride (F⁻) anions in aqueous solutions. The separation process is based on the distribution of the substance into a mobile fluid phase and a solid stationary phase. Sulfur was determined using an Eltra CS 2000 element analyzer. Samples were quantified using calibration curves generated from 105 reference materials.

Additional trace elements (As, Be, Bi, Cd, Co, Cs, Cu, Ge, Mo, Pb, REE, Sb, Sc, Sn, Th, Ti, U, W) were analyzed in a subset of samples by inductively coupled plasma mass spectrometry (ICP-MS) at GFZ Potsdam, using a HR-ICP-MS Thermo Element 2XR. A 125-μg sample material was digested in HF and aqua regia following Pretorius et al. (2006) and Romer and Hahne (2010). The internal standard indium was added to the solutions to monitor for drift. The full dataset, including major-, minor-, and trace-element analyses from this study, is given by Codeço et al. (2020) (Table D4). In addition to these data, we also included published data from the granite and micaite from Pinto (2014).
Laser ablation ICP-MS

For in situ analyses of white mica and tourmaline, 120-μm-thick polished sections from 10 samples (see ESM 1, Table S1) were prepared from the same billets used for microprobe and SIMS analyses in Codeço et al. (2017, 2019). Concentrations of $^7$Li, $^{11}$B, $^{23}$Na, $^{24}$Mg, $^{27}$Al, $^{33}$Cl, $^{39}$K, $^{47}$Ti, $^{51}$V, $^{55}$Mn, $^{57}$Fe, $^{63}$Cu, $^{65}$Zn, $^{85}$Rb, $^{88}$Sr, $^{89}$Y, $^{90}$Zr, $^{118}$Sn, $^{133}$Cs, $^{137}$Ba, $^{153}$Eu, $^{181}$Ta, $^{182}$W, $^{205}$TI, and $^{208}$Pb in mica and tourmaline grains were determined using a NewWave 213-nm Nd:YAG laser, coupled to a Thermo-Finnigan iCAP Qc quadrupole ICP-MS, at the Earth and Planetary Sciences Department, McGill University. Except for grains < 40 μm in size, which were mostly tourmaline, line scan mode was used to minimize the analysis of subsurface inclusions. The analytical conditions were 9 to 10 J/cm$^2$ fluence, 10-Hz laser pulse frequency, 20-μm line width, 5-μm/s line scan speed, 30-s warm-up, approximately 40-s ablation time, and 20-s washout. The ablated material was transported to the ICP-MS in an 800 mL/min He flow and mixed with Ar prior to injection into the plasma. Analyses of the unknowns were bracketed with analyses of reference NIST SRM 610 (glass). Also, at the beginning and end of each session, muscovite 98973, dravite 108796, schorl 112566 (Dyar et al. 2001), an in-house tourmaline (UoB), as well as BGR-2G and GSD-1G reference materials (GEOREIM database—Jochum et al. (2016)) were measured to assess analytical quality.

The raw data were processed using the Iolite software v. 2.5 (Paton et al. 2011). The limits of detection (LOD) and uncertainties of each element and mineral are reported in Codeço et al. (2020) (Tables D7 and D8). The abundance of the major elements (Na, K, Mg, Fe, Ti, Al, Mn, Si) was used to normalize the collected dataset to 15 cations at the Y, Z, T sites (for tourmaline) and 22 total charges (for white mica) and thereby correct for differences in ablation behavior between the samples and NIST-SRM 610. This method avoids the need for an externally determined reference element. Comparison with major elements determined by electron microprobe from Codeço et al. (2017, 2019) shows good agreement. Laser ablation analyses are provided in Codeço et al. (2020) (Tables D5 and D6).

Principal component analysis

Principal component analysis (PCA) is a multivariate statistical technique that identifies compositional vectors in a dataset by transforming the initial variables, e.g., element concentrations into a set of principal components (PCs). These PCs are defined as orthogonal linear combinations of the initial variables that align with the maximum variance in the dataset, with PC1 representing the most variance (Pawlowsky-Glahn et al. 2015). Eigenvalues and eigenvectors are used to characterize each PC. The eigenvalues represent the contribution of each principal component to the overall variance of the original data, while the eigenvectors represent the correlations between the initial data and the principal components (Pawlowsky-Glahn et al. 2015). The PCs with low eigenvalues are assumed to be dominated by “noise,” and when these are discarded, the dimensionality of the dataset is reduced, which facilitates interpretations of datasets with a large number of elements.

Geochemoal datasets typically contain “censored data,” i.e., missing values corresponding to analyses below detection limit (DL) or “structural zeros,” i.e., components not measured (Van Den Boogaart et al. 2006). These introduce limitations for the application of multivariate statistical methods that require complete datasets. For the PCA of the whole-rock data, we used all samples having the full suite of elements analyzed and added the analyses from Pinto (2014) that contained no censored data. Laser ablation ICP-MS always contain censored data because the small grain size of tourmaline forced laser ablation analysis to be conducted with a smaller beam size (20 μm), leading to relatively high detection limits for some trace elements. One solution to censored data is to exclude the data or to substitute the missing values by the DL, DL/2, or zero (Croghan and Egeghy 2003; Verbovšek 2011), but this can introduce bias (Dupuis and Beaudoin 2011; Makvandi et al. 2016a). Alternatively, censored data can be estimated by statistical methods (Aitchison 1982; Filzmoser et al. 2009; Hron et al. 2010), and the k-nearest neighbor (kNN) approach has been suggested to be the optimal imputation method for compositional datasets (Hron et al. 2010; Grunsky et al. 2013; Makvandi et al. 2016a, 2016b, 2019; Huang et al. 2019). Applying this method can have the drawback of neglecting the constraint that data fall below the DL, unless the method further enforces that the imputed values stay below the DL.

With respect to our datasets, the application of these different methods to pre-process censored data has only a minor influence on the PCA results. For this paper, we decided to substitute values below DL by DL/2 because the kNN imputation method (calculated with robCompositions package in R) resulted in values more than three times the DL, which clearly overestimated the low-concentration trace elements. We use the DL determined for the individual measurement...
wherever possible or, alternatively, the median DL value of the dataset for the respective element, when individual DLs are not available. Data were then transformed using the centered log-ratio (CLR) method in which each variable is divided by the geometric mean of all values and then log-normalized (Filzmoser et al. 2009). The PCA calculations and graphical results were made with the ioGAS™ software (REFLEX 2019). The imputation methods testing and the corresponding PCA results are described in full in the ESM 3.

Results

Whole-rock compositions

Both the tourmaline–muscovite–quartz alteration zone and unaltered metasediments show a wide compositional range for most major elements as a consequence of the varying proportions of sandstone– shale intervals in the protolith. Overall, the compositions of the unaltered schist and granite from our study compare well with analyses from the literature (Bussink 1984; Neiva 1987; Ugo et al. 2010; Villaseca et al. 2014), showing that the metasediments and intrusion at Panasqueira have compositions typical for this region. The major- and rare earth element (REE) contents of unaltered and altered metasediments overlap completely (Fig. 5a). REE patterns show relatively high middle- and heavy-REE contents (Sm, Eu, Gd, Yb, Lu) compared to the light REE. Since neither major elements nor REE are useful indicators of the alteration, we mainly focus on other minor- and trace-element contents. The element concentrations were normalized to those of the average upper continental crust (Rudnick and Gao 2014) (Fig. 5b). The altered schists show enrichments in As, Sn, Cd, Cs, Li, W, F, Bi, Cu, Rb, Zn, Tl, and Be compared to the unaltered metasediments, which is in good agreement with earlier findings of Bussink (1984). Also, most of these same elements are enriched in the greisen relative to the unaltered granite (Fig. 5c).
Trace-element contents of white mica and tourmaline

For the description of the trace-element results, we focus on the elements that are consistently enriched in the altered rocks, namely Sn, W, Cu, Zn, Cs, Li, and Rb, as well as some that were unaffected by alteration, e.g., Mg, V, and Sr. Some elements that exhibited important variations in the whole-rock data either could not be analyzed by LA-ICP-MS (F) or were below detection limits (As, Cd, Bi, Sb). Tantalum was not analyzed in the whole rock, but we measured Ta by LA-
ICP-MS in addition to Nb because the Nb/Ta ratio is useful as a marker of magmatic-hydrothermal processes (Ballouard et al. 2016). Chlorine was analyzed but not detected by LA-ICP-MS measurements and, therefore, excluded from the dataset.

White mica

White mica compositions show positive correlations for a number of element pairs with different slopes in different samples, in Li vs. Mg (Fig. 6a), Nb vs. Ta (Fig. 6b), and Cs vs. Rb (Fig. 6c). Strontium and V show a positive correlation in general, with highest V/Sr ratios in the fault (median = 30.2) and wall rock alteration zones (median = 6.4) and the lowest V/Sr ratios in micas from the greisen (median = 0.01) (Fig. 6d). The FG mica from MRS and MPS has higher V and Sr contents than CG mica in the same samples (Fig. 6d). Tin and W contents show an overall negative correlation, with the highest W/Sn ratio in greisen mica (median = 0.7) and the lowest W/Sn in mica within the fault zone (median = 0.08) (Fig. 6e). Copper and Zn are both highly variable and do not correlate with each other (Fig. 6f).

Mica from the greisen sample shows the most distinctive composition compared with other settings and generally has low Mg, high Ta and Nb contents (up to 100 and 200 ppm, respectively) and high W and Zn (up to 120 and 150 ppm, resp.), partially overlapping with the MRS micas, but low Sn contents (70–170 ppm). The Li, Cs, Rb, and Cu contents of greisen mica are variable and overlap with those of other settings. Some of this variability relates to compositional zoning, with mica rims having higher Li and Sn, but lower Cu and W values relative to cores (Fig. 6a, e, and f). In terms of Cs, Rb, Ta, and Nb, mica compositions form two distinctive clusters within individual grains (Fig. 6b, c), but this clustering cannot be related to growth zones.

Mica from the MRS shows distinct trace-element compositions, forming clusters in several of the binary plots. For example, sample PN46 stands out with high Nb and Ta contents that are comparable to the mica from the greisen. Sample PN22 has lower Li, Rb, and Zn contents compared to PN33 and PN46. Importantly, samples PN22 and PN33 contain two texturally distinctive mica varieties that could be analyzed separately, a fine-grained variety (FG) which occurs at the contact with the host rocks (Fig. 3a–c), and a coarse-grained variety (CG) that is commonly associated with wolframite (see Fig. 3a–d). In general, the fine-grained mica (“FG” in Fig. 6) has lower Rb and W contents, but higher Mg, Sn, Nb, Ta, Sr, and V than the coarser mica in these samples and in sample PN46, which has only CG mica.

White mica in wall rock alteration zones was generally too fine grained and intergrown with other phases to be measured by LA-ICP-MS. The single exception is sample PN30D, which yielded high Li and V contents compared with mica from vein selvages and greisen, but lower W and Sn contents. Wall rock mica from PN30D is similar to the FG mica from MRS in terms of Mg, Sr, Nb, Ta, Cs, Rb, and Zn contents. White mica from the fault zone (sample PN37–2) is distinctive from all others in terms of high Li, Mg, and Sn and low Nb, Ta, and W contents. For the other elements shown, the fault-zone mica and mica from vein selvages overlap.

Tourmaline

Tourmaline at Panasqueira is commonly zoned, with systematically higher Fe/Mg ratios from core to rim (Codeço et al. 2017). Where grain size allowed, core and rim compositions of tourmaline were analyzed separately, and some trace elements reflect the zoning; rims have higher Zn and Li and lower Sr and Mg contents than cores (Fig. 7). However, because of the small grain size, most LA-ICP-MS analyses consist of mixtures between rim and core composition. Overall, the tourmaline crystals from all settings show overlapping compositions and only occasionally are clusters by sample or type evident (Fig. 7d, f). The only example of distinct compositions are cores of tourmaline from the fault zone and the late vugs (Fig. 7), which have the highest Mg and Sr and the lowest Li and Zn contents in the dataset (see Fig. 7a, d, f). The lithophile element pairs Rb–Cs and Nb–Ta show strong positive correlations with each other (Fig. 7c, d), as was noted in mica. The chalcophile Zn–Cu element pair is not correlated, and Cu shows a random variation with no relationship to other elements, to crystal zoning, or to setting within the deposit (Fig. 7f). With few exceptions, both Sn and W concentrations are below 30 ppm. Like Cu and Zn, the ore elements Sn and W neither show correlation with other elements in tourmaline nor is there a systematic difference in concentrations relative to crystal zoning or setting within the deposit (Fig. 7e).

Discussion

Whole-rock chemistry

The altered host rocks show enrichments in W, Sn, Cu, and Zn as well as in As, F, Li, Rb, and Cs relative to the unaltered metasediments (Fig. 5). The principal component analysis (PCA) of all altered and unaltered schist samples was run with a set of 16 elements chosen to show different aspects of the element behavior (Fig. 8). We included two major elements (Fe, Mg), ten elements that are most consistently enriched during alteration (W, Sn, Cu, Zn, F, Rb, Cs, Li, As, and Tl), and four elements that are observed to remain relatively unchanged during alteration (V, Sr, Ba, and Nb) (Fig. 5b). Three principal components (PCs) account for 82% of the total variance, with 52.9% for PC1, 22.2% for PC2, and 6.7% for PC3. PC1 separates unaltered and altered metasediment samples
In contrast, the coarse-grained mica yields intermediate compositions between the FG mica and the greisen mica, suggesting greater influence of the hydrothermal fluid. Finally, mica from the fault zone stands out in terms of Li and Sn contents with strong positive loadings in PC2 and lower W, Ta, and Nb contents (Figs. 6 and 9a, b). The fault-zone mica also showed distinctly low B-isotopic values compared with mica from other settings (<23 to ~17‰; Cocco et al. 2019), which was interpreted as a consequence of cooling. Copper shows systematically negative correlation with Sn and is the only element with strong loadings in PC3, where it has a positive correlation with Li, opposite to the negative correlation in PC2 (Fig. 9b, d).

For tourmaline, the lack of clear correlations between elements observed in the bivariate plots (Fig. 7) is also reflected in the PCA results (Fig. 10). In contrast to the whole-rock and mica data, four PCs are needed to explain the main chemical variations in tourmaline (80% of the total variance), with 43.8% for PC1, 17.1% for PC2, 11.1% for PC3, and 7.9% for PC4 (ESM 3: Fig. S5). Iron, Zn, Mg, V, Sr, Sn, Li, and Cu are coupled and load positively in PC1, whereas the elements Cs, Rb, Ba, W, Nb, and Ta plot in the negative plane. Interestingly, the latter set of elements contains the ones where 25 to 37% of data are censored and were replaced by DL/√2 values. PC2 (Fig. 10b) is characterized by positive loadings for Rb, Cs, Ba and negative loadings for Nb and Ta, indicating that these elements are not related in tourmaline, which is consistent with their geochemical behavior (LILE vs. HFSE). Tungsten also loads in the negative plane, but like Cu, it is poorly represented by the first two components. We suggest that PC1 separates the “true” tourmaline composition (positive plane) from compositions likely affected by mineral inclusions (negative plane). Since Rb, Cs, Ta, Nb, Ba, and W have lower contents in tourmaline than in mica (see Figs. 6 and 7), these elements will be more sensitive to variance caused by inclusions of mica, rutile, and wolframite or fluid inclusions, which are visible in BSE images and under the microscope. If so, PC2 probably distinguishes between mica inclusions (Rb, Cs, and Ba) and rutile and/or wolframite (Nb, Ta, and W) and fluid inclusions (which may contain both these element groups). In PC3, Fe, Zn, V, and Li load in the positive plane, while Mg, Sr, and Sn load in the negative plane. This division is related to zoning since tourmaline rims are consistently enriched in Fe and Zn and depleted in Mg and Sr relative to cores as described above. Even though the zoning is an important feature of tourmaline, it is only captured by PC3 due to the fact that the CLR treats the data such that the PCA results are not overly weighted by the major elements (i.e., Mg and Fe). The relative importance of W and Cu increases from PC2 to PC4, with PC3 showing a positive correlation between the two and PC4 a negative one. This indicates that the variations of W and Cu are not systematic or are related to different processes, but our results cannot distinguish between these alternatives.

White mica and tourmaline

The PCA results for white mica show a strong clustering by setting and sample, as seen in Fig. 6. Three components account for 86% of the total variance, with 50.5% in PC1, 25.6% in PC2, and 9.4% in PC3 (Fig. 9; see also ESM 3). High concentrations in W (Zn), Ta, and Nb distinguish the greisen mica from mica in other settings. The fine-grained (FG) micas are distinct in terms of high Mg, Sr, Ba, and V contents and in terms of high Sr contents for sample PN22 (Fig. 9b, d). The PCA results show that the FG mica has a chemical affinity with the host rocks, suggesting a stronger rock control in their composition. In contrast, the coarse-grained mica yields
 Petrographic observations show that white mica and tourmaline are intergrown in many cases (see Fig. 3d), with sharp contacts, locally showing interpenetration, suggesting that they formed during the same event of hydrothermal alteration. Under equilibrium conditions, the composition of hydrothermal mica and tourmaline is controlled by partition coefficients of the elements between the mineral and fluid phase (van Hinsberg et al. 2010). We compared the trace-element compositions of such mineral pairs in contact to assess element partitioning between them (i.e., \( D_{\text{tur/mica}}^{\text{element}} \)). Correlation plots for selected trace elements (Fig. 11) show that Li, Rb, Cs, Tl, W, and Sn preferentially partition into mica, with apparent \( D_{\text{tur/mica}}^{\text{element}} \) values from 0.0001 to 0.5, with sample medians from 0.001 to 0.08. Copper, Ta, and Nb show considerable scatter, but they also generally favor mica over tourmaline (\( D_{\text{tur/mica}}^{\text{element}} = 0.01 \) to 30, median = 0.04 to 0.61), whereas Sr, V, and Zn partition preferentially into tourmaline.

![Fig. 7 Variations of major-, minor- and trace-element contents in tourmaline from different settings in the Panasqueira deposit.](image-url)

- **Fig. 7** Variations of major-, minor- and trace-element contents in tourmaline from different settings in the Panasqueira deposit. a Mg vs. Li, b Nb vs. Ta, c Sr vs. V, d W vs. Sn, f Cu vs. Zn. MRS mica-rich selvages, MPS mica-poor selvages, Wr-prox wall rock proximal to the vein-contact, Wr-dist wall rock distal from the vein contact; rims and cores of zoned grains in late vugs are outlined.

![Fig. 8 Principal components analysis (PCA) of the altered and unaltered whole-rock compositions (a–d).](image-url)

- **Fig. 8** Principal components analysis (PCA) of the altered and unaltered whole-rock compositions (a–d). a, c The log-transformed data plotted in PC1 vs. PC2 and PC1 vs. PC3 planes, respectively. b, d Element vectors (as variables) in the corresponding planes. The terms proximal and distal in the legend refer to distance from the greisen cupola. Dashed outlines in b and d highlight element clusters. e Vertical variations of PC1, PC2, and PC3 along drill core M1610 (drilled upward from a mine stope). Black rectangles correspond to the position of the samples studied. The kink in the PC1 curve at about 8 m marks the approximate boundary between altered and unaltered wall rocks.
These apparent $D$ values, while quite variable, are in reasonable agreement with the limited data in the literature, which are currently restricted to a study of the Mashhad granite, Iran (Zall et al. 2019), Zn–Pb–Ag mineralization at Broken Hill, Australia, and high-P/T metamorphic rocks from Syros, Greece (Klemme et al. 2011) (Fig. 12a). Importantly, in all previous studies and in our results from Panasqueira, the apparent $D$ values for individual elements are highly variable, which is contrary to expectation for equilibrium partitioning between...
the two minerals. For Panasqueira, tur/mica $D$ values vary by up to 3 orders of magnitude for some elements (e.g., Cs in Fig. 11c), even though the measured grains are in direct contact and appear to be coeval.

Possible explanations for the variation in element partitioning are that the mineral pairs did not reach chemical equilibrium, or that equilibrium was achieved but not maintained, or that the analyses are affected by microscopic mineral inclusions. Although we avoided all visible inclusions during analysis and excluded element spikes and troughs when integrating the laser ablation ICP-MS time-resolved spectra, tourmaline signals were commonly noisy, and incorporation of sub-micron inclusions cannot be excluded. There is evidence that white mica and tourmaline pairs from these samples reached B-isotope equilibrium (Codeço et al. 2019), but this does not necessarily argue for chemical equilibrium since chemical and B-isotope variations in tourmaline are commonly decoupled (Krienitz et al. 2008; Galbraith et al. 2009; Su et al. 2016; Codeço et al. 2017; Albert et al. 2018). The mineralization and alteration at Panasqueira involved dynamic processes and multiple stages, and this is reflected by the chemical zonation in tourmaline. White mica generally lacks zonation with the exception of mica from the greisen, so it may have re-equilibrated to changing conditions, while tourmaline could not due to its slow volume diffusion (e.g., van Hinsberg et al. 2011). If so, white mica may be in chemical equilibrium with the outer rim of coexisting tourmaline, but we cannot test this because the small size of the tourmaline grains makes reliable trace-element analysis of rims impossible.

On the other hand, the possibility that unseen mineral inclusions cause apparent disequilibrium element partitioning is very likely. Laser ablation ICP-MS profiles of some tourmaline grains show large oscillations, and a plot of Rb and Cs variations in tourmaline shows a correlation trend that extrapolates directly to the compositions of white mica in the same samples (Fig. 12b). Therefore, if mica inclusions affected the Rb and Cs concentrations, we expect that other elements also concentrated in mica (K, Ba, Tl) will be affected, and this could explain the near-vertical trends in mica-tourmaline correlation plots of Rb, Cs, Ti, W, and Sn (Figs. 11b–f). Inclusions are also consistent with the tourmaline PCA results mentioned above. There is a clear separation between the “true” tourmaline composition (PC1, positive plane) and mica inclusions (PC1, negative plane) (Fig. 10).

In conclusion, we suggest that the apparent lack of equilibrium between tourmaline and white mica may reflect the greater ability of white mica to adjust to changing fluid compositions during mineralization, as shown by differences in zoning of the two minerals, but there is definitely some effect of sub-microscopic inclusions in tourmaline. For tourmaline-mica partitioning, any mica inclusions in tourmaline would shift the apparent $D$ values closer to unity, so the measured values farthest from 1 could be closest to the “true” partitioning between tourmaline and mica.

### Estimating the fluid composition from white mica compositions

Given that white mica from Panasqueira is enriched in the majority of the mineralization-related trace elements, we can test its value as a recorder of fluid composition by comparing a calculated fluid composition based on our data with LA-ICP-MS analyses of fluid inclusions by Lecumberri-Sanchez et al. (2017). Unfortunately, mica-fluid element partition coefficients have only been determined for a few elements (Green and Adam 2003; Wunder and Melzer 2003), and these do not include the Panasqueira ore elements. As an alternative, we calculated mineral-fluid $D$ values for the elements of interest by combining mineral-melt and melt-fluid $D$ values following the approach of Keppler (1996). Table 1 shows apparent mica-fluid $D$ values derived from empirical mica-melt partitioning values of Kovalenko and Antipin (1980), based on white mica phenocrysts and groundmass in ongonites (volcanic equivalent to Sn-W granites), combined with experimental melt-fluid $D$ values from Zajacz et al. (2008). The $D$ values in Table 1 should be taken as a first approximation because the ongonite study and the fluid/melt experiments do not represent the same melt composition and P–T conditions; there is a significant spread in the melt/fluid $D$ values of Zajacz et al. (2008) depending on Cl contents; and none of the experiments were done for the low salinities (5–10 wt% NaCl) found in fluid inclusions from the Panasqueira veins. Given these uncertainties, the predicted fluid compositions (Table 1; Fig. 12c) are qualitative, but there is reasonable agreement with measured fluid compositions in quartz-hosted fluid inclusions from Panasqueira (Lecumberri-Sanchez et al. 2017). Note that in that study, high Sn concentrations in the host quartz (~50 ppm) meant that the detection limit for Sn in the included fluids was around 100 ppm, which is at the upper limit of Sn concentrations calculated from the mica concentrations (Fig. 12c). The fluid compositions calculated from mica compositions in different settings within the deposit suggest increasing Li, Cs, and Sn contents and decreasing W and Zn in the order from the greisen to the veins to the fault zone.

Fluorine and chlorine are important ligands in hydrothermal solutions generally, and they can play a significant role in Sn and W transport and precipitation (Manning and Henderson 1984; Schröcke et al. 1984; Heinrich 1990; Keppler and Wyllie 1991; Bai and Koster van Groos 1999; Wood and Samson 2000;
To investigate the potential of white mica and tourmaline as pathfinders for ore deposits of the W–Sn type more generally, we compared results from this study of Panasqueira with data from other Sn–W deposits as well as hydrothermal deposits with different metal associations (e.g., Cu–Au–Mo, Pb–Zn). Unfortunately, only a few studies report trace-element compositions of mica and most of those are for trioctahedral micas (Maoping, Legros et al. 2016; Piotang, Legros et al. 2018; Cinovec/Zinnwald, Breiter et al. 2017; Qiluling, Xie et al. 2015; Yashan, Li et al. 2015). All Panasqueira micas are dioctahedral, so the comparison needs to be interpreted with caution. Nevertheless, we find that mica with W–Sn association has higher Rb, Cs (Li), and Sn contents compared with that from porphyry Cu and VMS base-metal deposits, whereas the range of W contents overlaps (Fig. 13a–c). Most mica from W–Sn deposits, including Panasqueira, has > 10 ppm W, while mica from the majority of porphyry deposits has W contents between 1 and 10 ppm. Early mica from the Piaotang deposit, which is coeval with cassiterite, yielded lower W contents than later mica from that deposit, indicating that it can record changing conditions (Legros et al. 2018). There are no data for Zn and Cu in mica from the other W–Sn deposits, but our Panasqueira data overlap in terms of these elements with mica from VMS and porphyry deposits (Fig. 13d).

A similar comparison for tourmaline shows higher Sn and Li contents in that mineral from Sn–W deposits relative to tourmaline from the Salikvan Cu–Mo porphyry deposit and orogenic gold deposits (Fig. 13e), but no significant difference was found for W, except for the Salikvan deposit (Fig. 13f). The Sn–Li plot also discriminates between Sn–W deposits in the Massif Central and deposits with W only. The strong overlap of all deposits on the V and Sr plot (Fig. 13g) supports the interpretation that these elements are host rock controlled and depend on the local lithogeochemistry rather than mineralizing fluids (Kalliomäki et al. 2017). The range of Cu concentrations in tourmaline is similar and highly variable in all deposits, but Zn contents tend to be higher at Panasqueira than in all other Sn–W deposits (Fig. 13h). This may reflect the high sphalerite content in the Panasqueira veins.

Even though tourmaline compositions are less responsive to the composition of the hydrothermal fluid than white mica, our compilation suggests that high Sn and Li contents in tourmaline (i.e., > 10 ppm and 50 ppm, respectively) are distinctive for magmatic-hydrothermal Sn deposits. In any case, given that mica and tourmaline tend to occur together in hydrothermal systems and incorporate different sets of trace elements, the best approach for geochemical pathfinding using alteration minerals would be to combine information from both minerals.
Conclusions

White mica, tourmaline, and quartz are the most abundant hydrothermal minerals in the alteration assemblages at the Panasqueira W–Sn–Cu deposit, where mica forms thick vein selvages at the contacts with the host rocks, and tourmaline predominately occurs in the alteration halos. To explore the potential of hydrothermal minerals as indicators for mineralization at Panasqueira, we determined the trace-element contents of unaltered and...
altered rock samples by whole rock analyses and of hydrothermal white mica and tourmaline by in situ LA-ICP-MS. The resulting data were investigated with the help of principal component analysis (PCA).

The alteration zone is characterized by enrichments in As, Sn, Li, W, F, Cu, Rb, Zn, Tl, and Pb as delineated by PCA results from the whole-rock data. These elements strongly load on the first principal component (PC1) of the variability, whereas PC2 is interpreted to reflect the different modal abundances of minerals such as mica and tourmaline. PC3 identifies a subordinate geochemical variation as a function of distance from the greisen, which is mineralogically controlled, with proximal zones enriched in Rb, Cs, and Tl reflecting thick mica selvages and the distal zones richer in As, Zn, and Cu reflecting sulfide vein mineralization.

The in situ LA-ICP-MS data show that Rb, Cs, Ba, Li, Nb, Ta, W, and Sn preferentially partition into white mica over tourmaline while Zn, V, and Sr do the opposite. Thus, white mica better reflects the hydrothermal fluid and its evolution. Fine-grained mica at the wall rock contact zone has lower Rb and W contents (and Zn in some cases) but higher Mg, Sr, V, and Sn contents compared to mica of the main vein-selvage and the greisen body. This suggests that the micas at the contact zone partially reflect the composition of the host rock. In addition to a temporal evolution, the PCA results for white mica show a clear clustering by

Table 1  Apparent partition coefficients of mica-fluid ($D_{m-w}$) used in this study to reconstruct fluid compositions from analyses of white mica

| Element | $D_{m-w}$ | All vein mica | MPS | MRS | Fault | Greisen |
|---------|-----------|---------------|-----|-----|-------|---------|
| Na      | 0.17      | 24,012        | 24,814 | 24,690 | 6815 | 23,843 |
| K       | 8.7       | 10,060        | 9984 | 9975 | 10,115 | 10,028 |
| Rb      | 9.6$^vi$ | 201           | 177 | 207 | 237 | 200 |
| Cs      | 2.1       | 54            | 53  | 52  | 61  | 47     |
| Li      | 1.4       | 940           | 815 | 865 | 1942 | 549    |
| Pb      | 0.21      | 9.9           | 13  | 9.6 | 4.1  | 6.0    |
| Sr      | 0.56      | 24            | 50  | 19  | 1.9  | 3.1    |
| Ba      | 0.26      | 2079          | 3303 | 1741 | 1466 | 217    |
| Mg      | 2.5       | 2888          | 3960 | 2025 | 7495 | 372    |
| Zn      | 1.1       | 94            | 73  | 113 | 61  | 121    |
| Mn      | 4.0       | 85            | 72  | 89  | 131 | 156    |
| Fe      | 595       | 35            | 29  | 37  | 44  | 39     |
| Al      | 50        | 3625          | 3672 | 3636 | 3019 | 3779   |
| B       | 0.92$^vi$ | 65            | 57  | 65  | 126 | 55     |
| Sn      | 3.3       | 60            | 63  | 58  | 101 | 37     |
| Nb      | 121       | 0.05          | 0.03 | 0.06 | 0.01 | 1.0    |
| W       | 2.2       | 36            | 26  | 38  | 11  | 42     |

$^i$Calculated by combining empirical mica-melt and experimental fluid-melt partition coefficients from Kovalenko and Antipin (1980) and Zajacz et al. (2008), respectively (see text)

$^ii$Fluid composition based on the median value of mica analyses from the given setting

$^iii$Based on the median values of MRS and MPS mica analyses combined

$^iv$Mica-rich selvage

$^v$Mica-poor selvage

$^vi$Based on the maximum value of partition coefficients of mica-melt reported by Kovalenko and Antipin (1980)

Fig. 13  Trace-element concentrations in micas (a to d) and tourmaline (e to h) from Panasqueira and other hydrothermal deposits worldwide. Studies from Piaotang W–Sn (Legros et al. 2018), Yashan W–Ta–Nb (Li et al. 2015), and Qiguling rholyte (Furong Sn) (Xie et al. 2015) include trioctahedral and dioctahedral micas; Maoping W–Sn (Legros et al. 2016) and Cinovec/Zinnwald Sn–W–Li (Breiter et al. 2017) only comprise trioctahedral micas; data from other ore deposits are only dioctahedral micas: Ann-Mason porphyry Cu (Cohen 2011), Copper Cliff porphyry Cu (Uribe-Mogollon and Maher 2018), and Bathurst Mining Camp VMS (Soltani Dehnavi et al. 2018). Data sources for tourmaline are: Duchoslav et al. (2017)—Comwall Sn–W; Harlaux (2016)—Enguiles and St-Mélan—French Massif Central (FMC); Jiang et al. (2004)—Yunlong Sn; Harlaux et al. (2019)—Puy-les-Vignes, French Massif Central W; Hazarika et al. (2017)—Hutti-Maski (Au); Trumbull et al. (2019) and Cabral et al. (2012)—Passagem de Mariana (Au); Kalliomäki et al. (2017)—Archean Hattu Schist Belt (Au); and Yavuz et al. (1999)—Salikvan porphyry Cu–Mo porphyry deposit. PCu and PCu–Mo porphyry Cu and Cu–Mo deposits, respectively, VMS volcanogenic massive sulfide deposits, FMC French Massif Central
setting, distinguishing the greisen, vein selvages, and late fault zones from each other. The trace-element composition of hydrothermal fluid calculated from approximated fluid-mica partition coefficients shows reasonable agreement with measured compositions of quartz-hosted fluid inclusions and suggests an evolution of the fluid from greisen formation to vein selvage (MRS) to late-stage mineralized fault zones, with progressive increases in Li, Cs, and Sn with time and fluctuation in Sn/W ratios. The mica data and fluid modeling suggest that fluids released from the granitic intrusion were responsible for the hydrothermal alteration. Comparisons of mica data from Panasqueira and other deposits suggest that white mica has a great potential as a pathfinder for W–Sn mineralization with characteristically high Rb, Cs, Li, Sn, and W contents.

In contrast to white mica, tourmaline at Panasqueira mainly forms by mineral replacement in the host rock, and its composition is less influenced by the hydrothermal fluid. The PCA results from tourmaline data are complex, and the variability is complicated by analytical issues related to mineral and fluid inclusions, as well as by chemical zoning that cannot be spatially resolved by LA-ICP-MS analysis. Despite this, the comparison of tourmaline data from Panasqueira and other hydrothermal ore deposits suggests that high Sn and Li contents in hydrothermal tourmaline are indicative for Sn–W mineralization.

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