Reactive interaction between migmatite-related melt and mafic rocks: clues from the Variscan lower crust of Palmi (southwestern Calabria, Italy)

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Abstract. In the Variscan lower–intermediate crust exposed in the Palmi area (southwestern Calabria, Italy), amphibolites occur as foliated, decimeter-thick layers within migmatitic paragneiss and as a decametric main body adjacent to the migmatites. The main body is mostly fine-grained and weakly to moderately foliated; unfoliated medium-grained portions rarely occur. Amphibolites are mainly composed of plagioclase (An$_{80-91}$) frequently developing triple junctions, amphibole consisting of cummingtonite rimmed by hornblende and variable amounts of biotite. Minor quartz is present in amphibolite layers within paragneiss. Accessory allanite occurs in amphibolite layers within migmatites and in foliated, fine-grained portions from the main body. This study mainly aims to achieve information about the effects triggered by the migration of migmatite-related melts into associated mafic rocks and its role in the re-distribution of major and trace elements out of the anatectic source.

On the basis of whole-rock major- and trace-element compositions, the protolith of amphibolite is recognized as of cumulus origin, likely derived from basic melt emplaced in the framework of the late-Variscan lithospheric extension. The rocks experienced high-temperature subsolidus re-equilibration ($\sim$ 800°C) in conjunction with the development of amphibole. The origin of amphibole is attributed to a coupled dissolution–precipitation process related to the reaction between migrating SiO$_2$-rich hydrous melt and precursor orthopyroxene (± plagioclase). Reactive melt migration also caused the crystallization of biotite ± allanite ± quartz ± plagioclase. SiO$_2$-rich hydrous melt had REE (rare earth element) compositions similar to late-Variscan peraluminous granites and could have been derived by partial melting of metasediments akin to neighboring migmatitic paragneisses.

Both whole-rock and amphibole analyses reveal a decrease in Mg# (Mg/(Mg + Fe$^{2+}$)) from amphibolite layers within paragneiss to fine- and medium-grained rocks of the main body. Hornblende shows an increase in SiO$_2$ and a decrease in Al$_2$O$_3$ and K$_2$O with increasing Mg#. Amphibolites interlayered with paragneiss have higher K$_2$O, Rb, Ba, Th, U and Zr relative to those from the main body. Furthermore, amphibole from amphibolites interlayered with paragneiss is distinct for relatively high Rb, Ba, MREE (middle rare earth element) and HREE (heavy rare earth element) concentrations. Within the main body, foliated, fine-grained rock has both the whole rock and amphibole enriched in Rb and Ba and high Zr bulk-rock contents. Whole-rock and mineral chemistry heterogeneity most likely reflects (i) variation of the composition of the melt during its reactive migration, in response to dissolution of pre-existing minerals and crystallization of new phases, and (ii) variable modification of the original compositions. Foliated and fine-grained amphibolites record the strongest modification, thereby suggesting that they represent permeable pathway enabling effective interaction of the reacting melt with precursor minerals and nucleation of new mineral phases.
1 Introduction

Partial melting and felsic-melt movement from the sites of generation in the lower crust to sites of accumulation in the upper crust is indicated as a fundamental mass-transfer process responsible for the compositional differentiation of Earth’s crust (Brown et al., 2011; Sawyer et al., 2011). It is commonly envisaged that melts may preferentially enrich in most incompatible elements and transport them in the upper crust, thereby leaving a residual, depleted lower crust. Nevertheless, an opposite effect was observed in several leucosomes and granitic bodies showing lower concentrations of incompatible trace elements (i.e., U, Th, Y, Zr and REEs; rare earth elements) with respect to their residual source (e.g., Sawyer, 1991; Wolfram et al., 2017). It was also shown that fluid-absent partial melting is not associated with a depletion of U and Th (i.e., heat-producing elements) concentrations in residual metapelitic-derived granulite facies rocks; elevated U–Th contents are rather retained by basement terranes (Alessio et al., 2018). Lower-than-expected concentrations of trace elements in melts relative to their source were related to processes occurring during early stages of melt migration close to the source region (e.g., Sawyer, 1991). In particular, a strong control was imputed to accessory minerals (e.g., zircon, monazite and apatite) that are able to a incorporate large amounts of Zr, U, Th, Y and REEs (e.g., Bea, 1996). In a recent study about the anatectic turbidites exposed on Kangaroo Island, Schwindinger et al. (2020) showed that accessory minerals such as apatite, monazite, xenotime and zircon become preferentially concentrated in the melanosomes. The authors proposed that local melt–host reaction leads to crystallization of magma oversaturated in accessory minerals close to the source region in the melanosome, away from the melt channel, thereby removing U, Th and REEs from the migrating melt. Mechanisms responsible for mass transfer and distribution of most incompatible elements, in particular U, Th, Zr, Y and REEs, during anatexis at the source area and through the crust remain open to debate.

Recent studies demonstrated that melt–rock interactions and reactive porous flow are key processes in the evolution of crustal rocks. For instance, they are responsible for the redistribution of major and trace elements in the oceanic lithosphere, leading interstitial melt, migrating through a crystal framework, to change towards more evolved compositions and trace-element enrichments (e.g., Lissenberg et al., 2013; Sanfilippo et al., 2020). Reactive porous melt flow was indicated as a process able to produce the hornblende-rich rocks close to the source region in the melanosome, away from the melt channel, thereby removing U, Th and REEs from the migrating melt. Mechanisms responsible for mass transfer and distribution of most incompatible elements, in particular U, Th, Zr, Y and REEs, during anatexis at the source area and through the crust remain open to debate.

This paper aims to achieve information on the effects triggered by the reactive migration of migmatite-related melts into associated mafic rocks and its role in the distribution of major and trace elements from the source region. For this purpose, I considered a portion of the Variscan lower–intermediate crust exposed in the Palmi area (southwestern Calabria, Italy) consisting of migmatitic paragneiss showing a peak metamorphic assemblage of biotite, K-feldspar, garnet, sillimanite and cordierite. I focused on amphibolites occurring as foliated, decimeter-thick layers within migmatites and as a decametric main body adjacent to paragneiss. In this contribution, I report major and trace-element composition of major minerals (amphibole, plagioclase, garnet) combined with major- and trace-element compositions of whole rocks. The data are used to assess that reactive migmatite-related melt imparted a mineralogical and geochemical signature by migrating within associated mafic rocks, where it promoted the crystallization of amphibole and accessory phases (i.e., allanite), which played an important role in the re-distribution of REEs and most incompatible elements out of the anatectic zone.

2 Geological framework and field relations

The Calabria–Peloritani terrane is a nappe-structured belt, built up during Alpine tectogenesis and located between the NW–SE-trending southern Apennines and E–W-trending Maghrebides (Amodio-Morelli et al., 1976). The terrane drifted to the present position during the Apennine orogeny and the opening of the western Mediterranean basins (Alvarez et al., 1974). The nappe-structured edifice, from top to bottom, consists of (i) Paleozoic continental-crust units with metamorphic and igneous rocks pertaining to the southern European Variscan belt (Fig. 1), (ii) Mesozoic oceanic-crust units locally affected by Alpine high-pressure/low-temperature (HP/LT) metamorphism, and (iii) carbonate rocks of the Apennine units (Bonardi et al., 2001, and references therein).

Late-Variscan continental-crust sections, tilted and nearly completely preserved, are exposed in the Sila (Dubois, 1971; Graessner and Schenk, 2001) and Serre massifs (Schenk, 1980). They are characterized by widespread low-pressure/high-temperature (LP/HT) metamorphism, intense
crustal anatexis and granitoid magmatism, as commonly found in other segments of the southern European Hercynian belt (e.g., Pn and Vielzeuf, 1983). In the Calabria crustal sections, granulite facies gneisses of the lower crust are overlain by tabular late-Variscan granitoids and minor gabbroic bodies emplaced at intermediate to upper-crustal levels between 306 and 290 Ma ($^{40}$Ar/$^{39}$Ar on hornblende; Ayuso et al., 1994; U–Pb on monazite; Graessner et al., 2000; U–Pb on zircon, Langone et al., 2014). The contact between granitoids and high-grade metamorphic rocks is characterized by a migmatitic border zone (Rottura et al., 1990). The granitoids intruded into overlying amphibolite and in greenschist facies metamorphic rocks of the intermediate–upper crust. Granitoids mainly consist of metaluminous calc–alkaline tonalites and granodiorites and are interpreted to originate from mantle-derived magmas hybridized by either assimilation of crustal material or mixing with crustal melts (e.g., Ayuso et al., 1994). Nearly coeval, subordinate peraluminous granites also developed by lower-crust anatexis (e.g., Rottura et al., 1993; Caggianelli et al., 2003; Fiannaca et al., 2019). Minor mafic intrusions (gabbros to diorites) of tholeiitic affinity crystallized at intermediate-crustal levels (Caggianelli et al., 1994) almost simultaneously with the intrusion of the late-Variscan granitoids. Typically, these mafic intrusions can be observed in the migmatitic border zone, as relatively small bodies on the order of some tens of meters thick (Rottura, 1985; Caggianelli and Di Florio, 1989). Moreover, they occur within tonalite and quartz-diorite as syn-plutonic dikes and enclaves (Rottura et al., 1990). The late-Variscan magmatic activity in the Calabria terrane was ended by the intrusion between 295 and 277 Ma of dike swarms ranging in compositions from microgranites and microgranodiorites to minor microgabbros and microdiorites (Festa et al., 2010).

The Palmi area is located in southwestern Calabria and comprises a portion of the Variscan lower–intermediate crust, mainly consisting of migmatitic paragneisses to the north and of amphibole-bearing tonalitic gneisses to the south (Fig. 1). The overall characteristics of the Palmi Variscan basement are similar to those observed in the Serre crustal section, where the transition zone from the mid to lower crust is exposed (Schenk, 1984). The tonalitic gneisses from Palmi show a strong foliation developed mostly at the solid state as documented by widespread presence of dynamically re-
crystallized quartz (Caggianelli et al., 1997; Prosser et al., 2003). On the basis of Al-in-hornblende geobarometer, intrusion of the tonalites within migmatitic paragneiss was estimated at 610 ± 60 MPa (Caggianelli et al., 1997). A thick marble layer is comprised between the tonalitic gneisses and migmatitic paragneiss and contain abundant clasts of calc–silicate-bearing rocks interpreted as former skarns developed during the intrusion of the tonalites. Marble acted as a weak horizon during the Eocene allowing for deformation events promoting strain localization and the production of a 400 m thick mylonitic shear zone (Prosser et al., 2003). Later deformation at the brittle–ductile transition was documented by pseudotachylyte-bearing shear zones within the tonalites and related to Oligocene extensional tectonics (Grande et al., 2009). The migmatitic paragneisses show a layering defined by the alternation of quartz–feldspar felsic bands and garnet–sillimanite–cordierite–biotite restitic layers, folded and transposed by late-Variscan high-temperature deformation (Grande et al., 2009). The paragneisses frequently include layers of calc–silicate-bearing marble, orthogneiss and metagabbro (Prosser et al., 2003).

In the Palmi area, rocks largely composed of amphibole and plagioclase (hereafter amphibolites) occur as flattened microgranular enclaves within the tonalitic gneisses (Rottura, 1985; Rottura et al., 1990) and as centimeter-thick layers or decametric bodies within the migmatitic paragneisses. Amphibolites have been studied on two main outcrops along the beach, named Scoglio dell’Isola and Pietre Nere, respectively (Figs. 1 and 2a). The former covers an area of about 1400 m², with a length of about 50 m towards NNW–SSE, a width of about 35 m towards ENE–WSW and a maximum elevation of about 14 m a.s.l. It is mainly composed of migmatitic paragneisses characterized by a stromatic texture with alternating melanosomes and leucosomes. The peak mineral assemblage within the melanosomes consists of biotite, K-feldspar, garnet, cordierite and sillimanite. The latter occurs as elongated prisms within the foliation or as centimeter-sized aggregates (up to 7 cm). Leucosomes are composed of quartz, feldspars (plagioclase, K-feldspars), biotite and white mica ± garnet ± cordierite. Foliated, fine-grained amphibolites occur as layers up to 10 cm thick and are locally characterized by the presence of centimetric garnet grains. Garnet is preferentially observed at the boundary between amphibolites and migmatites and is locally surrounded by a corona of biotite (Figs. 2b and 3a). Amphibolite layers are alternated with felsic ones (up to 30 cm thick) mainly composed of centimetric feldspars, quartz, biotite and garnet.

Amphibolites also occur as decametric bodies exposed on the cliff and on the beach. The outcrop on the cliff is only accessible with difficulty, whereas on the beach amphibolites occur as an isolated body (hereafter referred to as the Pietre Nere body) without exposed contacts with host rocks (Fig. 2a). The Pietre Nere body is mainly weakly to moderately foliated and fine-grained (Fig. 2c). Medium-grained, centimeter-scaled domains rarely occur; they are unfoliated and patchily associated with the fine-grained portions. Leucocratic lenses with diffuse margins characterized by relatively high plagioclase content are locally present throughout the body. On the basis of preliminary U–Pb data obtained by sector-zoned, magmatic cores of zircons separated from a Pietre Nere foliated rock, a Carboniferous age of intrusion for the parental magma was proposed (Renna et al., 2020).

### 3 Petrographic features of selected samples

In this study, amphibolites from the Pietre Nere body and from the layers within migmatitic paragneiss of the Scoglio dell’Isola were considered. In particular, for petrographic descriptions, whole-rock and mineral analyses, I selected (Table 1) two unfoliated medium-grained rocks (PN3 and PN4), one foliated rock (PN1) and one weakly foliated fine-grained rock (PN2) from Pietre Nere and two amphibolites interlayered with migmatitic paragneiss, which include one garnet-free (SI-1) and one garnet-bearing (SI-MR) sample.

#### 3.1 Pietre Nere

Amphibolites from Pietre Nere mainly consist of plagioclase, amphibole and minor biotite. Medium-grained amphibolites typically show anhedral poikilitic amphibole enclosing plagioclase grains (Fig. 4a and b). Fine-grained amphibolites display an overall granoblastic texture and a foliation produced by the preferential orientation of biotite. In both medium- and fine-grained amphibolites, plagioclase (45 vol%–55 vol%) is subhedral and frequently shows triple junctions (Fig. 4c). Amphibole occurs in a variable modal amount; in particular medium-grained amphibole...
Poikilitic amphibole from the medium-grained amphibolites is locally composed of polygonal grains with nearly equant morphology. In fine-grained amphibolites, amphibole typically displays polygonal equant morphology (Fig. 4d); anhedral grains interstitial to plagioclase also occur. Amphibole from all Pietre Nere amphibolites is zoned with pale-brown cores and green rims (Fig. 4e), which are frequently in optical continuity. Green amphibole decreases from the modal amount of 25 vol % of the medium-grained amphibolites to the ∼15 vol % of the fine-grained samples (Table 1). Anhedral grains interstitial to plagioclase also occur. Amphibole from all Pietre Nere amphibolites is zoned with pale-brown cores and green rims (Fig. 4e), which are frequently in optical continuity. Green amphibole decreases from the modal amount of 25 vol % of the medium-grained amphibolites to the ∼15 vol % of the fine-grained samples (Table 1). Bi-

Fe–Ti oxides (up to 5 vol %) appear interstitial to plagioclase and amphibole; in medium-grained samples they were also found as inclusions or along the cleavage planes of green amphibole. Zircon and apatite are accessory phases. Accessory allanite associated with biotite and amphibole was also found in foliated fine-grained amphibolite (PN1, Fig. 4f).

### 3.2 Scoglio dell’Isola

Amphibolites interlayered with paragneiss are fine-grained and show a foliation defined by the parallel alignment of biotite and amphibole. They mainly consist of subhedral to anhedral plagioclase (40 vol %–43 vol %), variable modal abundances of anhedral amphibole (18 vol %–25 vol %) and subhedral biotite (25 vol %–30 vol %), and minor amounts of anhedral quartz (7 vol %). One sample is characterized by the oc-
Figure 4. Thin-section photomicrographs of (a, b) poikilitic amphibole enclosing plagioclase grains (samples PN4 and PN3, respectively); (c) plagioclase showing 120° triple junctions (sample PN4); (d) aggregate of polygonal amphibole grains (sample PN2); (e) amphibole showing pale-brown (cummingtonite) core and green (hornblende) rim (sample PN4); (f) allanite associated with biotite and amphibole (sample PN1); (g) amphibole showing pale-brown (cummingtonite) core and green (hornblende) rim (sample SI-1); and (h) inclusions of plagioclase, quartz and cummingtonite in garnet (sample SI-MR).

currence of a centimetric garnet grain (Fig. 3). Accessory phases are Fe–Ti oxides, zircon, apatite and allanite, which commonly occur in association with biotite and amphibole.

In the garnet-free quartz amphibolite (hereafter referred to as quartz amphibolite), plagioclase locally intersects other plagioclase grains or amphibole at 120° triple junctions and amphibole is optically zoned with pale-brown cores and thin green rims (Fig. 4g). In the quartz–garnet amphibolite (hereafter referred to as garnet amphibolite), amphibole is pale brown. In this sample, garnet occurs as a poikiloblast of ~1 cm in size, with euhedral faces and cuspat terminations. It is mantled by biotite and minor quartz and contains numerous inclusions of plagioclase, commonly showing embayed boundaries and mainly distributed in the core.
Other inclusions comprise pale-brown amphibole with elliptical morphology, irregularly shaped to subhedral biotite, anhedral quartz and Fe–Ti oxides (Figs. 3 and 4h). Only one small inclusion of green amphibole was found within the garnet. Zircons from the garnet amphibolite are 50–500 µm in size and have euhedral to subhedral morphology. The internal structure of zircons from the garnet amphibolite was investigated with backscattered electron (BSE) microscopy and cathodoluminescence (CL) using a Philips XL30 electron microscope equipped with a Centaurus CL detector. Prior to the CL imaging, the sample was carbon coated and the images were obtained using 15 kV acceleration voltage and a working distance of 26 mm. CL and BSE images revealed internal structures commonly characterized by an irregularly curved patchy pattern and convolute zoning, surrounded by thin high-luminescent rims (Fig. 5). The zircons also frequently showed fractures, pores and inclusions of xenotime.

4 Mineral chemistry

Amphibole, plagioclase, biotite and garnet were analyzed for Si, Ti, Al, Cr, Fe, Mn, Ni, Mg, Ca, Na and K (Tables 2–5 and S1–S4 in the Supplement) by electron microprobe (JEOL JXA-8200 SuperProbe) at the Dipartimento di Scienze della Terra, Università degli Studi di Milano, in wavelength-dispersive spectrometry mode. For all analyses, electron microprobe operating conditions were 15 kV accelerating voltage and 5 nA beam current, counting times were 30 s on the peaks and 10 s on the backgrounds, and natural standards were used.

Trace-element mineral compositions of amphibole, plagioclase, garnet and allanite were determined by laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) at the CNR IGG UOS of Pavia (Consiglio Nazionale delle Ricerche Istituto di Geoscienze e Georisorse Supporting Unit of Pavia; Table S5). The instrument couples a 193 nm ArF excimer laser microprobe (Geolas200QMicrolas) to a triple quadrupole ICP-MS system (ICP-QQQ, 8900 from Agilent). NIST SRM 610 was used as an external standard; 44Ca or 29Si were adopted as the internal standards, depending on the mineral composition. In each analytical run the USGS reference sample BCR2 and NIST SRM 612 were analyzed together with the unknowns for quality control.

4.1 Major elements

Plagioclase from Pietre Nere amphibolites is rich in anorthitic component ranging from 80 mol % to 86 mol %. Anorthite contents in plagioclase from the quartz amphibolite interlayered with paragneiss ranges from 86 mol % to 89 mol %. In the garnet amphibolite, plagioclase in the matrix and included in the garnet have similar anorthite contents, overall ranging from 87 mol % to 91 mol % (Tables 2 and S1).

According to the classification of Hawthorne et al. (2012), pale-brown amphiboles from all selected samples are cummingtonite and green amphiboles are hornblende. Cummingtonite from the medium-grained Pietre Nere rocks have higher Mg# (Mg/(Mg + Fe2+)) in atoms per formula unit) than that from the associated fine-grained samples (57–62 vs. 51–56; Fig. 6). In the garnet amphibolite, cummingtonite within the garnet has higher Mg# (59–61) and lower MnO (∼ 0.6 wt %) than that in the matrix (53–56 and 0.6 wt %–1.0 wt %). Cummingtonite from the quartz amphibolite has low Mg# (50–53) and MnO contents similar to those of cummingtonite from the matrix of the garnet amphibolite (Table 3, Fig. 6). MnO contents in cummingtonite from Pietre...
Nere rocks are relatively low (0.4 wt %–6.0 wt %). Estimated Fe ratios (Fe<sup>2+</sup>/(Fe<sup>2+</sup> + Fe<sup>3+</sup>)) overall range from 0.94 to 1.00 with an average of 0.99 (Table S2). The cummingtonite has 0.4 wt %–1.7 wt % of Al<sub>2</sub>O<sub>3</sub> and 0.5 wt %–1.7 wt % of CaO.

Mg# of hornblende from the medium-grained Pietre Nere amphibolites is higher than that from fine-grained samples (62–67 vs. 44–54; Fig. 7). In Pietre Nere rocks, hornblende has also variable contents of Al<sub>2</sub>O<sub>3</sub> (7.0 wt %–18.0 wt %) and CaO (7.0 wt %–12 wt %) and low amounts of K<sub>2</sub>O (0.2 wt %–0.4 wt %). Hornblende from the fine-grained amphibolites frequently shows higher Al<sub>2</sub>O<sub>3</sub> and lower SiO<sub>2</sub> than that from the medium-grained samples (Fig. 7). Hornblende from the quartz amphibolite has low Mg# (39–45) and SiO<sub>2</sub> (~43 wt %) and relatively high Al<sub>2</sub>O<sub>3</sub> (14 wt %–16 wt %) and K<sub>2</sub>O (0.4 wt %–0.5 wt %). Hornblende included within garnet has relatively high Mg# and SiO<sub>2</sub> (60 wt % and 47 wt %) and low Al<sub>2</sub>O<sub>3</sub> (11 wt %). Estimated Fe ratios (Fe<sup>2+</sup>/(Fe<sup>2+</sup> + Fe<sup>3+</sup>)) overall range from 0.89 to 1.00 with an average of 0.99 (Table S3). Hornblende has TiO<sub>2</sub> in the range of 0.1 wt % to 0.9 wt %.

Biotite from fine-grained Pietre Nere amphibolites differs from that of associated medium-grained samples for the higher Fe<sub>tot</sub><sup>2+</sup>/(Fe<sub>tot</sub><sup>2+</sup> + Mg) values, which are in the range of 0.41–0.44 and 0.33–0.39, respectively (Tables S5 and S4). In the garnet amphibolite, biotite within the poikilitic garnet has lower Fe<sub>tot</sub><sup>2+</sup>/(Fe<sub>tot</sub><sup>2+</sup> + Mg) values than that in the matrix (0.33–0.37 vs. 0.40–0.44). High Fe<sub>tot</sub><sup>2+</sup>/(Fe<sub>tot</sub><sup>2+</sup> + Mg) values were found for the biotite from the quartz amphibolite (0.47–0.53). Overall, biotite has TiO<sub>2</sub> from 1.7 wt % to 2.8 wt % and Na<sub>2</sub>O from 0.1 wt % to 0.4 wt %. Biotite from the garnet amphibolite is characterized by high K<sub>2</sub>O (9.0 wt %–10.0 wt %), which frequently exceed those of biotite from other samples (8.2 wt %–9.2 wt %).

Garnet is almandine-rich (Alm<sub>60</sub>–69, Gr<sub>7</sub>–14, Py<sub>13</sub>–19, Sp<sub>9</sub>–9) and shows (i) a rough, oscillatory zoning for Ca, which is higher along the garnet rim and in the core, slightly
Figure 7. Variation of Al₂O₃ (wt %), SiO₂ (wt %) and K₂O (wt %) vs. Mg# in hornblende.

Table 4. Electron microprobe analysis of hornblende (average values, wt %).

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<tr>
<th>Sample</th>
<th>PN4</th>
<th>PN3</th>
<th>PN2</th>
<th>PN1</th>
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<th>SI-MR</th>
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<td>n = 6</td>
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<td>SD</td>
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<td>TiO₂</td>
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<tr>
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<td>0.04</td>
<td>0.03</td>
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<tr>
<td>FeO</td>
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<td>14.0</td>
<td>0.9</td>
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<td>MnO</td>
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<tr>
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<td>0.6</td>
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<tr>
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<td>64.1</td>
<td>1.8</td>
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</tbody>
</table>

M-G amph.: medium-grained amphibolite. F-G amph.: fine-grained amphibolite. Qtz amph.: quartz amphibolite. Grt amph.: quartz–garnet amphibolite. Mg# = 100 × Mg³⁺/(Mg³⁺ + Fe²⁺).

decreasing in the intermediate zone, and (ii) slight depletion of Mg towards the rims (Fig. 3, Tables 6 and S4).

### 4.2 Trace-element compositions

#### 4.2.1 Amphibole and plagioclase

The chondrite-normalized REE pattern of cummingtonite from Pietre Nere amphibolites shows negative Eu anomaly (Eu / Eu*) ~ 0.6) and LREE (light rare earth element) depletion with respect to MREEs (middle rare earth elements) and HREEs (heavy rare earth elements), which are nearly flat at ~ 10 times chondrite (Fig. 8). Cummingtonite from the fine-grained amphibolite PN1 differs for a more marked LREE depletion (CeN / SmN = 0.2 for SmN = 12) with respect to other Pietre Nere samples (CeN / SmN = 0.6 for SmN = 13). Within each sample, the REE pattern of hornblende is parallel to that of the associated cummingtonite. The hornblende has higher absolute REE concentrations (by about an order of magnitude) with a pattern exhibiting a slight decrease from Gd to Lu (GdN / LuN = 2.0–2.7 for LuN = 35–58) and a marked negative Eu anomaly (Eu / Eu* = 0.4). With respect to cummingtonite, hornblende is also characterized by higher V, Cr and Sc concentrations (Table S5). Overall, both hornblende and cummingtonite from the fine-grained sample PN1 have higher V, Cr and Sc than those from other Pietre Nere amphibolites.

Cummingtonite and hornblende from Pietre Nere amphibolites have parallel chondrite-normalized incompatible-element patterns, which overall show Rb, Ba, Nb, Ta and HfSE (high-field-strength element) depletion with respect to MREEs–HREEs and Y (Fig. 8). Rb, Ba, Nb and Ta are also enriched relative to La–Ce. Hornblende has a parallel chondrite-normalized incompatible-element pattern to associated cummingtonite but at higher concentration levels (by about an order of magnitude). Phlogopite from Pietre Nere amphibolites (Fig. 9) have nearly similar chondrite-normalized REE patterns, which...
Figure 8. REE and incompatible-trace-element compositions normalized to chondrite (Anders and Ebihara, 1982) of cummingtonite and hornblende.

Table 6. Electron microprobe analysis of garnet (average values, wt %).

<table>
<thead>
<tr>
<th>Sample</th>
<th>SI-MR</th>
<th>SI-MR</th>
<th>SI-MR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mineral</td>
<td>Grt rim</td>
<td>Grt interim</td>
<td>Grt core</td>
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<tr>
<td>n = 4</td>
<td>SD</td>
<td>n = 8</td>
<td>SD</td>
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<tr>
<td>SiO₂</td>
<td>38.5 ± 0.4</td>
<td>38.4 ± 0.2</td>
<td>38.9 ± 0.2</td>
</tr>
<tr>
<td>TiO₂</td>
<td>&lt; 0.04 ± –</td>
<td>&lt; 0.04 ± –</td>
<td>&lt; 0.04 ± –</td>
</tr>
<tr>
<td>Al₂O₃</td>
<td>22.1 ± 0.2</td>
<td>21.9 ± 0.2</td>
<td>22.2 ± 0.1</td>
</tr>
<tr>
<td>Cr₂O₃</td>
<td>0.03 ± 0.03</td>
<td>0.03 ± 0.02</td>
<td>0.03 ± 0.02</td>
</tr>
<tr>
<td>FeO</td>
<td>29.9 ± 0.1</td>
<td>31.0 ± 0.2</td>
<td>28.9 ± 0.7</td>
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<tr>
<td>MnO</td>
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<td>3.39 ± 0.11</td>
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<tr>
<td>MgO</td>
<td>3.59 ± 0.38</td>
<td>4.00 ± 0.11</td>
<td>4.57 ± 0.15</td>
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<tr>
<td>CaO</td>
<td>5.13 ± 0.14</td>
<td>4.11 ± 0.21</td>
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<tr>
<td>Na₂O</td>
<td>0.02 ± 0.03</td>
<td>0.02 ± 0.03</td>
<td>0.02 ± 0.02</td>
</tr>
<tr>
<td>K₂O</td>
<td>&lt; 0.01 ± –</td>
<td>&lt; 0.01 ± –</td>
<td>&lt; 0.01 ± –</td>
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<tr>
<td>Sum</td>
<td>102.8 ± 0.5</td>
<td>102.9 ± 0.2</td>
<td>103.2 ± 0.3</td>
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<tr>
<td>Prp</td>
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<td>16.1 ± 0.6</td>
<td>17.9 ± 0.5</td>
</tr>
<tr>
<td>Alm</td>
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<td>67.7 ± 0.7</td>
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</tr>
<tr>
<td>Grs</td>
<td>7.96 ± 0.75</td>
<td>7.74 ± 0.32</td>
<td>7.43 ± 0.58</td>
</tr>
<tr>
<td>SpS</td>
<td>12.6 ± 1.1</td>
<td>8.50 ± 1.19</td>
<td>12.6 ± 1.2</td>
</tr>
</tbody>
</table>

Grt amph.: quartz–garnet amphibolite.
display a marked LREE enrichment relative to MREEs (LaN / SmN = 18–22 for SmN = 1.2–1.8), a prominent positive Eu anomaly (Eu / Eu* = 27–38) and a steady decrease from Gd to Er (GdN / ErN ~ 7 for ErN ~ 0.1). In the quartz amphibolite, the REE pattern of plagioclase is characterized by a flat LREE (LaN / SmN = 1.0 for SmN = 7.0) and a moderate positive Eu anomaly (Eu / Eu* = 8) and depletion of HREEs relative to MREEs (GdN / ErN = 7 for ErN = 0.5).

4.2.2 Garnet and allanite from the garnet amphibolite

The core of the poikilitic garnet has a chondrite-normalized REE pattern (Fig. 9) showing HREE enrichment with respect to MREEs (GdN / LuN = 0.21 for LuN = 490) and a marked negative Eu anomaly (Eu / Eu* = 0.1). Normalization to chondrite reveals that the rim of the garnet differs for a nearly flat HREE pattern, which are at lower concentration levels (by a factor of ~5) and a slight negative Eu anomaly (Eu / Eu* = 0.5).

The chondrite-normalized REE pattern of allanite (Fig. 9) is characterized by a marked enrichment of LREE relative to MREEs (LaN / SmN = 5.7 for SmN = 32 294), a prominent negative anomaly (Eu / Eu* = 0.2) and a steady decrease from Gd to Lu (GdN / LuN = 36 for LuN = 381). Uranium and Th contents are high (1130 and 1750 ppm, respectively). Allanite also has relatively high contents of Ti, Ba, V, Cr and Sc (Table S5).

5 Whole-rock chemistry

Whole-rock major- and trace-element analyses of the selected rocks (Table S6) were carried out at Activation Laboratories (Ancaster, Ontario) by inductively coupled plasma (ICP) optical emission spectroscopy and ICP mass spectrometry (4Lithores method of lithium metaborate and tetraborate fusion, ICP, and ICP-MS). Three blanks and five controls (three before sample group and two after) were analyzed per group of samples. Precision and accuracy are estimated to be better than 5 % for the other major elements. Precision and accuracy of trace-element analyses are typically assessed to be within 10 %.

Medium-grained amphibolites from Pietre Nere have higher Mg# (molar Mg/(Mg + Fe2+)) × 100 = 57–59) than associated fine-grained samples (50–51, Fig. 10). Fine-grained amphibolite PN1 has lower CaO and higher K2O than other Pietre Nere samples, which overall range from 10.1 wt % to 11.0 wt % and from 0.41 wt % to 0.56 wt %, respectively. The TiO2 concentrations are higher in medium-grained amphibolite PN3 than in other Pietre Nere samples, which reflects its higher modal content of Fe–Ti oxides (Table 1), as also confirmed by normative calculations revealing a significant amount of normative ilmenite in this sample (Table S6). The garnet amphibolite has higher Mg# than quartz amphibolite (54 and 49, respectively). The samples interlayered with the paragneiss are also characterized by relatively low CaO (7.0 wt %–8.7 wt %) and Na2O (0.6 wt %–0.8 wt %) and high K2O (2.3 wt %–4.0 wt %). Amphibolites have overall Al2O3 in the range of 19 wt % to 21 wt % (Fig. 10) and relatively low SiO2 / Al2O3 values (2.2 to 2.5, Table S6).

Medium-grained amphibolites from Pietre Nere have the highest contents of V (280–487 ppm), which reflects the higher modal content of (i) hornblende, which is characterized by high V concentrations (Table S5), and (ii) Fe–Ti oxides for which V is a compatible element (Best, 2003). The concentrations of Ni and Cr are overall low (< 30 ppm and 30–100 ppm, respectively).

The chondrite-normalized REE patterns (Fig. 11) of fine-grained amphibolites and medium-grained amphibolite PN4 from Pietre Nere are characterized by LREE enrichment over HREEs (LaN / YbN = 4.2–10.5 for YbN = 5.7–8.4). The REE pattern of these samples also displays a positive Eu anomaly (Eu / Eu* = 1.5–1.8). Fine-grained samples are slightly LREE enriched and HREE depleted relative to sample PN4. The REE pattern of medium-grained amphibolite PN3 differs from those of other Pietre Nere rocks for showing slightly higher MREE and HREE contents (by a factor of ~2), a slight depletion of LREEs relative to MREEs and negative Eu anomaly (Eu / Eu* = 0.6), which may be related to a relatively high modal amphibole/plagioclase ratio (Table 1). The chondrite-normalized REE pattern of the
garnet amphibolite shows a steady decrease from LREEs to HREEs ($\La_{N}/\Yb_{N} = 8.1$ for $\Yb_{N} = 15$) and a weak positive Eu anomaly ($\Eu/\Eu^{*} = 1.1$). The quartz amphibolite displays LREE depletion relative to MREEs, which are in turn enriched over HREEs ($\Gd_{N}/\Yb_{N} = 1.7$ for $\Yb_{N} = 26$), and a negative Eu anomaly ($\Eu/\Eu^{*} = 0.5$). It is noteworthy that REE patterns of the quartz and garnet amphibolites are approximately parallel to those of medium-grained sample PN3 and fine-grained rocks from Pietre Nere, respectively, but at slightly higher concentration levels.

In the chondrite-normalized multi-element diagram, Pietre Nere amphibolites display the enrichment of Ba, Nb and Sr relative to Rb, Th, Zr and Y (Fig. 11). Fine-grained amphibolite PN1 differs from other Pietre Nere samples in showing higher contents of Rb, Ba and Th and Zr enrichment over Y. Amphibolites interlayered with the paragneiss are characterized by a significant enrichment of Rb, Ba, Th and Nb with respect to Sr, Zr and Y.

6 Discussion

6.1 The protolith of Palmi amphibolites

In the Palmi area (southwestern Calabria, Italy), a Variscan lower–intermediate crust is exposed and mainly consists of amphibole-bearing tonalitic gneisses and migmatitic paragneiss. Amphibolites occur in close spatial relationship with migmatitic paragneiss (Fig. 1). In particular, amphibolites are present as foliated, decimeter-thick layers within migmatites and as a decametric main body (i.e., Pietre Nere body) adjacent to the paragneiss (Fig. 2). The Pietre Nere body is mainly fine-grained and variably foliated, even though unfoliated, medium-grained portions rarely occur. Amphibolites from the Pietre Nere body and interlayered with paragneiss share a similar mineralogical composition mainly composed of anorthite-rich plagioclase ($\An_{80−91}$), amphibole consisting of cummingtonite rimmed by hornblende and biotite. Amphibolite layers within paragneiss contain minor quartz, accessory allanite and locally garnet. Accessory allanite also locally occurs in foliated, fine-grained Pietre Nere amphibolites.

Based on their $\SiO_{2}$ contents (47 wt%–49 wt%, calculated water free), all amphibolites have a basic composition and fall in the basaltic (gabbroic) field of the total alkali vs. silica diagram (TAS, Best, 2003; Fig. S1 in the Supplement). All samples are characterized by high amount of normative anorthite and hypersthene (Table S6). Furthermore, one sample from the Pietre Nere body and one interlayered with paragneiss are slightly olivine normative, thereby indicating that amphibolites likely derived from a mafic igneous protolith. Relic of texture that may be reconciled with a magmatic origin is represented by the poikilitic morphology of amphibole in unfoliated, medium-grained rocks from Pietre Nere (Fig. 4a and b). The above analogy in terms of modal, normative and whole-rock major-element compositions and the fact that amphibolites interlayered with paragneiss have approximately parallel REE patterns to amphibolites from Pietre Nere (Fig. 11) suggest a genetic relationship among these rocks. Furthermore, whole-rock compositions suggest that the protolith of amphibolites had a cumulus origin, in particular the low $\SiO_{2}/\Al_{2}O_{3}$ values (Table S6), and the frequent occurrence of positive Eu anomaly in REE patterns.
(Fig. 11) provide evidence for a plagioclase-bearing cumulate nature for these rocks (see Kempton and Harmon, 1992). The relatively low Mg# values (49 to 59) and low concentrations of Ni and Cr (Table S6) suggest a formation by melts that had previously undergone fractional crystallization.

The late-Variscan mafic intrusion of Rovale (Silà Massif, northern Calabria) includes olivine-bearing cumulates ranging from norites to amphibole–gabbros, emplaced at mid-crustal level ($P \sim 4$ kbar, pressure, Caggianelli et al., 1994). They formed by early precipitation of plagioclase and olivine and late crystallization of orthopyroxene, amphibole and biotite. Olivine has a low forsterite component (63 mol %–68 mol %), and orthopyroxene is characterized by Mg# ranging from 71 to 75. Amphibole, essentially Ti-pargasite, occurs as poikilitic grains enclosing plagioclase, which overall has 78 mol %–90 mol % of anorthite component. The Rovale gabbroic rocks have enriched Nd and Sr isotopic signatures indicating a significant crustal contribution in their origin ($\varepsilon_{Nd} = -6.1$ and $^{87}Sr / ^{86}Sr = 0.7091 - 0.7095$). Amphibolites from Palmi have comparable Mg# and Al$_2$O$_3$ with Rovale gabbros (Fig. 10). Palmi amphibolites and Rovale gabbros also consist of similar anorthite-rich plagioclase. In addition, Palmi amphibolites have similar chondrite-normalized REE patterns to those of gabbros from Rovale controlled by plagioclase accumulation and high amphibole modal amount (Fig. 11). The original mineralogical and geochemical composition of the protolith of Palmi amphibolites could approximately resemble that of cumulate gabbroic rocks from the Rovale mafic intrusion, thereby suggesting that the late-Variscan evolution of the Calabria terrane was characterized by mid–lower-crust intrusion of chemically comparable basic melts.

6.3 Origin of amphibole by reactive melt migration

Amphibole is ubiquitous in all amphibolites from the Palmi area and mostly consists of cummingtonite cores rimmed by hornblende (Fig. 4). Cummingtonite is often described in rocks formed under conditions ranging from upper greenschist to the beginning of granulite facies conditions (e.g., Mottana et al., 1994; Schumacher, 2007). The entire stability range for cummingtonite was thermodynamically constrained at 400–800°C and < 1–15 kbar (Evans and Ghiorsò, 1995). Cummingtonite was also observed as a magmatic phase in dacites and rhyolites (e.g., Geschwind and Rutherford, 1992; Deering et al., 2008). Fractional crystallization experiments conducted at 0.7 GPa for hydrous, calc–alkaline to arc-tholeiitic magmas showed that cummingtonite crystallizes at temperatures below 780°C, coexisting with a rhyolitic liquid and a mineralogical assemblage characterized by a plagioclase with $\sim 40 \text{ mol} %$–65 mol % of anorthite component (Nandedkar et al., 2014, 2016). Furthermore, the thermodynamic model by Evans and Ghiorsò (1995) showed that cummingtonite, when present, should be associated with relatively Na-rich plagioclase. In the Palmi amphibolites, the primary crystallization of cummingtonite in association with the anorthite-rich plagioclase is thus incompatible with experiments and natural occurrence of this amphibole in magmatic rocks of silicic compositions. The involvement of an exotic agent after the formation of early plagioclase is required to explain the crystallization of cummingtonite in the Palmi amphibolites.

In gabbros, the origin of cummingtonite is sometimes attributed to interaction between orthopyroxene and a late mafic magmatic aqueous fluid phase (e.g., Roberts et al., 2000) or SiO$_2$-rich hydrous melt (e.g., Elburg, 1996). In metabasic rocks from the Serre and Sila massifs (central and northern Calabria) the origin of cummingtonite was related to hydration of granulite facies orthopyroxene, according to the following reaction (mineral abbreviations after Whitney

$$\text{en} + \text{pl} = \text{cumm} + \text{hbl} + \text{cl}$$

where en = enstatite, pl = plagioclase, cumm = cummingtonite, hbl = hornblende, and cl = clinopyroxene.
The following reaction for the replacement of orthopyroxene by cummingtonite may be envisaged as follows:

\[
\text{Opx} + \text{SiO}_2 - \text{rich hydrous melt} = \text{Cum}. \tag{R1}
\]

The proposed cummingtonite-forming reaction implies an open system reactive melt flow, characterized by a coupled dissolution–precipitation process. A similar scenario was proposed for the formation, in the lower and mid crust, of hornblende-rich, hornblende or biotite-rich rocks by modification of precursor, largely anhydrous gabbroic material (Meek et al., 2019; Piazolo et al., 2020). Cummingtonite formed by replacing precursor orthopyroxene that became unstable, reacted and dissolved in the melt. Coupling of dissolution and precipitation may result in new mineral replacing the volume occupied by the dissolving phase, thereby explaining the local poikilitic habit of amphibole in Pietre Nere amphibolites, which most likely preserves the igneous texture of precursor orthopyroxene. It was indicated that, during high-temperature annealing, the process of dissolution–precipitation may erase microstructures such as interstitial grains with low dihedral angles, commonly related to the presence of migrating melt in a rock (Piazolo et al., 2006, 2020). In Palmi amphibolites, the common occurrence of triple junctions among plagioclase and locally amphibole indicates that annealing could have been responsible for erasing microstructures commonly referred to the presence of a migrating melt.

In Palmi amphibolites, cummingtonite is generally associated with biotite and locally also with allanite (Fig. 4f). The markedly LREE-depleted pattern locally shown by cummingtonite confirms the presence of allanite during cummingtonite formation, because allanite is a REE-rich mineral showing a strong preference of LREEs over HREEs (Fig. 9). Notably, in the Pietre Nere body, the REE pattern of plagioclase from the sample containing allanite is not LREE depleted as the associated cummingtonite but is similar to those of plagioclase from other allanite-free amphibolites (Fig. 9). It suggests that cummingtonite was not at equilibrium with plagioclase, most likely because the duration of heating could have been too short for reaction to go to completion, thereby hindering element diffusion (Vernon, 2004; see also Sect. 6.4). In amphibolites interlayered with paragneiss, both plagioclase and cummingtonite show...
LREE-depleted patterns, which indicates that plagioclase crystallized in association with allanite and cummingtonite. Since minor quartz occurs in association with cummingtonite and biotite in amphibole interlayered with paragneiss, the cummingtonite-forming reaction for Palmi amphibolites may be expressed as follows:

\[
\text{Opx} \pm \text{PI(1)} + \text{SiO}_2 \rightarrow \text{rich hydrous melt} = \text{Cum} + \text{Bt} \pm \text{Pl(2)} \pm \text{Qtz} \pm \text{Aln.}
\] (R2)

This formulation implies that plagioclase could locally recrystallize simultaneously with the melt–rock reaction. Development of plagioclase after Reaction (R2) may explain the slightly higher anorthite content shown by plagioclase from amphibolites interlayered with paragneiss relative to that from Pietre Nere (Table 2). The incorporation of Ca in Pl (2) could have been also responsible for the minor precipitation of hornblende within amphibole interlayered with paragneiss (Table 1).

In Palmi amphibolites, cummingtonite is generally rimmed by hornblende (Fig. 4e and g), frequently exhibiting optical continuity, which indicates they form part of a single crystal. Similar hornblende–cummingtonite associations were documented in granulites and amphibolites derived from mafic and ultramafic protoliths (Angus and Kanaris-Sotiriou, 1982; Mongkoltip and Ashworth, 1986; Schenk, 1984). Hornblende (+ quartz) was interpreted to derive by reaction between anorthite–rich plagioclase and cummingtonite, which in turn had formed from precursor orthopyroxene (see also Evans and Ghiorso, 1995). In metabasic rocks from lower-crustal, granulite facies units of the northern Serre from southern Calabria, the derivation of hornblende after cummingtonite was inferred to be coupled with a decrease in the anorthite component in plagioclase (i.e., from An90 to An70, Schenk, 1984). Quartz is absent in Pietre Nere amphibolites, which are characterized by the highest modal amounts of hornblende (Table 1), and only occurs in amphibolites interlayered with paragneiss. It is also noteworthy that plagioclase has a relatively homogeneous composition and does not exhibit a decrease in the anorthite content from core to rim, which would be expected in response to the hornblende-producing reaction. I therefore argue that hornblende crystallization after cummingtonite in Palmi amphibolites did not involve other minerals.

The envisioned melt–rock reaction initiated with the development of cummingtonite. Following the cummingtonite formation, the reacting melt could enrich in elements having low affinity with respect to cummingtonite, e.g., Ca, Na and Al, which, with decreasing temperature, could promote crystallization of hornblende on cummingtonite. This is consistent with the optically continuity frequently shown by cummingtonite cores to hornblende rims. This hypothesis is also consistent with the increase in REE concentrations from cummingtonite to hornblende because experimental studies for a liquid line of descent of a calc–alkaline basalt at 0.7 GPa determined that amphibole–melt partition coefficients increase systematically with decreasing temperature (Nandekar et al., 2016); in particular, REE partition coefficients increase up to 1 order of magnitude. Note that the ratios of REEs in hornblende to cummingtonite slightly decrease with an increasing atomic number (Fig. 8), in agreement with a higher compatibility of HREEs relative to LREEs and MREEs in cummingtonite (Tiepolo et al., 2007). Hornblende formation may be thus related to cooling and late-stage crystallization of the migrating melt, incorporating elements that have an extremely low affinity for cummingtonite. The complete reaction for the process leading to the formation of Palmi amphiboles may be schematized as follows:

\[
\text{Opx} \pm \text{Pl(1)} + \text{SiO}_2 \rightarrow \text{rich hydrous melt} = \text{Cum} + \text{Bt} + \text{Hbl} \pm \text{Pl(2)} \pm \text{Qtz} \pm \text{Aln} \pm \text{residual melt.} \] (R3)

Melt has been included among the reaction products, it could correspond to any melt fraction possibly able to migrate out of the system. The development of amphibole in Palmi amphibolites may be thus attributed to a reaction between precursor orthopyroxene (± plagioclase) and a SiO2-rich hydrous melt, by a coupled dissolution–precipitation process. This reaction was also responsible for the precipitation of biotite ± allanite ± quartz ± plagioclase.

In the Variscan sectors of the Calabria terrane, rocks from the lower crust, after peak metamorphism under granulite facies conditions at 300 ± 10 Ma (U–Pb on zircon and monazite), 790 ± 30 °C and 7.5 ± 0.5 kbar, experienced a nearly isothermal decompression up to 290–280 Ma (Schenk, 1989, 1990; Fornelli et al., 2011). The end of the granulate facies metamorphism was induced by a tectonic event during which the lower crust was uplifted into mid–lower-crust level (~5.6 kbar), where it cooled isobarically up to 25 Ma (Schenk, 1990). The temperatures estimated on the basis of hornblende composition for Palmi amphibolites indicate that hornblende crystallization occurred at ~800 °C. The two amphiboles show nearly parallel REE patterns with hornblende having higher concentrations levels of about 1 order of magnitude than cummingtonite (Fig. 8). The origin of mineral grains (i.e., clinopyroxene and amphibole) with a similar trace-element variability in gabbros and basalts is commonly attributed to interplay between crystal-growth phenomena and kinetic effects during relatively rapid cooling (e.g., Renna et al., 2011, 2016). It was experimentally shown that at high cooling rates, the crystal growth rate exceeds diffusion rate of components, thereby resulting in the development of a boundary layer at the crystal–melt interface enriched in rejected component (Loftgren et al., 2006; Schwandt and McKay, 2006). I speculate that conditions of relatively rapid crystal growth responsible for hornblende crystallization on cummingtonite rim could be related to the onset of the exhumation and cooling of granulite facies rocks at mid-crustal levels.
6.4 Origin of the garnet amphibolite

Garnet poikiloblast in the amphibolite interlayered with paragneiss contains inclusions of plagioclase and cummingtonite showing variable grain size, rounded to elliptical shape and frequent embayed morphology, thereby indicating that they were partially digested by garnet growth. The idioblastic crystal shape with sharp edges and the inclusion of cummingtonite and anorthite-rich plagioclase (Table 2) that are not present in the neighboring paragneiss (Rochira, 2014) indicate that garnet is not a xenocryst derived from the immediately adjacent migmatisites, but it formed in the amphibolite layer.

The embayed morphology of garnet-hosted plagioclase and cummingtonite, in tandem with the absence of plagioclase and cummingtonite in the corona around garnet, suggests that garnet was produced by a reaction involving cummingtonite and plagioclase. The reaction may be referred to that proposed by Hollocher (1991) for Fe-rich metabasic rock in the transitional amphibolite to granulite facies terrane from central Massachusetts (USA): \[ \text{Cum} + \text{Pl} + \text{Bt} + \text{Ilm} = \text{Hbl} + \text{Grt} + \text{Qtz} + \text{H}_2\text{O}. \]

On the compatibility triangular ACF diagram for metamorphic mafic rocks (Best, 2003) created using the compositions of minerals from the garnet amphibolite, this reaction is confirmed by the intersection between the hornblende–garnet and the anorthite–cummingtonite–biotite tie lines (Fig. S2). Agreement with the occurrence of irregularly shaped biotite inclusions in garnet. Hence, quartz included and along the rim of garnet and the rare garnet-hosted hornblende likely formed as a product of a cummingtonite breakdown reaction. The Ca enrichment shown by garnet rim may be attributed to plagioclase breakdown, in agreement with its absence in the corona around the poikiloblast (Fig. 3). Notably, the garnet-forming reaction proposed by Hollocher (1991) implies fluids release and, consequently, their possible interaction with minerals from the matrix. The migration of fluids in the garnet amphibolite is indicated by zircons, which have internal structure characterized by curved patchy pattern and convolute zonating (Fig. 5), commonly related to a reaction with a metamorphic fluid through a coupled dissolution–reprecipitation process (Tomaschek et al., 2003; Geisler et al., 2007; Rubatto et al., 2008). Zircons mostly preserved their euhedral/subhedral habit because the process of dissolution–reprecipitation is inferred to be responsible for the complete replacement of one zircon by another without losing the external shape or crystal morphology of the parent zircon. The numerous inclusions of xenotime in zircons represent another typical sign of their recrystallization because they are generally interpreted to form from the expulsion of non-essential trace elements (i.e., Y and REEs) from the recrystallizing zircons (see also Spandler et al., 2004). It is therefore likely that garnet formed by reaction of cummingtonite + plagioclase + biotite; the garnet-forming reaction was responsible of liberating fluids whose migration was cause for zircon recrystallization.

Garnet poikiloblast shows Mg- and HREE-depleted rim with respect to the core. The lower-Mg zones in garnet rim are most likely the result of diffusive Fe–Mg exchange with the adjacent biotite from the surrounding corona. The HREE-depleted signature of rim to core is consistent with a pro-grade garnet growth because a progressive decrease in the mass fraction of compatible elements (i.e., Y and HREEs) from core to rim is generally attributed to a Rayleigh-type fractionation process (Otamendi et al., 2002; Rubatto et al., 2020). Subsolidus Fe–Mg and Mg–Mn exchange occurred between garnet and included mafic minerals, as indicated by the higher Mg# and lower MnO shown by cummingtonite within garnet with respect to that in the matrix and the lower \( \text{Fe}^{2+}/(\text{Fe}^{2+} + \text{Mg}) \) shown by biotite included relative to that in the matrix (Fig. 6; Table 5). Cummingtonite inclusion was also affected by HREE re-equilibration with the host garnet, most likely owing to high-temperature conditions, as indicated by its HREE-depleted pattern with respect to that of cummingtonite in the matrix.

Garnet is absent in the Pietre Nere body and occurs preferentially along the contact between amphibolite layers and migmatisites. Smith et al. (2015) showed that metamorphic rocks with heterogeneous texture and mineral abundance may be produced by different extents of reaction and annealing in high-grade gneiss adjacent to intruding plagioclase-rich felsic dikes. The extent of reaction and annealing is greatest in areas adjacent to dike–host boundaries, owing to a long duration of relatively high temperature, whereas further away the reaction is incomplete. This results in an increase in grain size and abundance of reaction products close to the felsic dike. It can be proposed that garnet could develop in amphibolite layers by the cummingtonite–plagioclase–biotite reaction, in close proximity to migmatitic paragneiss, because these could be areas of long durations of high temperatures generated by migmatization and migration of anatectic melts. Exposure to high temperature for a relatively long period may explain the absence in garnet amphibolite of hornblende rim around cummingtonite because its development has been related to conditions of relatively rapid cooling causing rapid crystal growth (see Sect. 6.3). A garnet-forming reaction most likely proceeded in tandem with a high extent of annealing, thereby resulting in the nucleation and growth of coarse-grained garnet. Notably, conditions of a higher extent of annealing in amphibolite interlayered with paragneiss relative to the Pietre Nere body are consistent with evidence for plagioclase recrystallization simultaneously with allanite, amphibole and biotite, namely during the melt–rock reaction process. The Pietre Nere body, further away from the contact with the migmatitic paragneiss, could experience high temperature for a relatively shorter time lapse, which hindered the garnet-forming reaction and was associated with a relatively short time annealing, not

enough for plagioclase recrystallization during the melt–rock reaction.

6.5 Chemical and mineralogical variation in Palmi amphibolites

The amphibolites from Palmi show systematic variations in terms of modal and chemical composition. The highest and lowest amounts of amphibole were found in medium-grained Pietre Nere amphibolites and in amphibolites interlayered within paragneiss, respectively; intermediate amphibole amounts characterize fine-grained Pietre Nere amphibolites (Table 1). Amphibolites interlayered with paragneiss have the highest amounts of biotite, which is minor in fine-grained amphibolites and accessory in medium-grained amphibolites from the Pietre Nere body. Quartz is ~ 7 vol % in amphibolites interlayered within the paragneiss but absent in Pietre Nere amphibolites. The whole rock, cummingtonite and hornblende share a similar variation of Mg#, which increases from the quartz amphibolite to fine- and medium-grained amphibolites of Pietre Nere (Figs. 6, 7 and 10). Furthermore, the Fe$_{tot}^{2+}$/($Fe_{tot}^{2+} + Mg$) of biotite decreases from amphibolites interlayered with paragneiss to fine- and medium-grained Pietre Nere amphibolites (Table 5). Quartz and garnet amphibolites have higher bulk-rock content of K$_2$O than Pietre Nere amphibolites. Notably hornblende and biotite from amphibolite layers within paragneiss are also enriched in K$_2$O relative to their counterparts from Pietre Nere, thereby indicating that the K$_2$O enrichment observed for bulk-rock composition is not related to a low-temperature alteration. A gradual decrease in K$_2$O-Al$_2$O$_3$ and increase in SiO$_2$ with increasing Mg# is remarkably shown by hornblende compositions (Fig. 7). Moreover, from the amphibolite interlayered with paragneiss to those from Pietre Nere, the contents of MnO in cummingtonite decrease. Major-element variations may be overall explained by a variation in the composition of the reactive migrating melt in response to dissolution of pre-existing minerals and fractionation of new phases. Dissolution of orthopyroxene and crystallization of cummingtonite and biotite could be responsible for the increase in Mg and decrease in Fe in the reactive migrating melt. Biotite fractionation also resulted in a decrease in K$_2$O and Al$_2$O$_3$ of the reacting melt. A decrease in SiO$_2$ and MnO could reflect the crystallization of quartz and cummingtonite, respectively. The process of reaction, dissolution and precipitation also resulted in an increase in Ca and Na in the migrating melt because these elements are incorporated in negligible amounts in cummingtonite and biotite. Ca and Na enrichment was most likely responsible for a progressive increase in hornblende crystallization as the melt migrated, reacted and crystallized, thereby explaining the increase in hornblende modal amount from the quartz amphibolite to the fine- and medium-grained Pietre Nere amphibolites (Table 1). Hence, the migrating melt could ultimately lead to a predominance of hornblende with respect to cummingtonite in the mineral reaction products.

Mg# of cummingtonite equilibrium melts may be computed assuming amphibole–basalt Fe–Mg partition coefficients of 0.399 as resulting from experiments carried out at 780°C and 0.7 GPa by Nandedkar et al. (2014). Calculations showed that cummingtonite from the quartz amphibolite, matrix of the garnet amphibolite, fine-grained Pietre Nere amphibolite and medium-grained Pietre Nere amphibolite were in equilibrium with a melt having 29.8, 33.0, 32.7 and 37.2 Mg#, respectively. It was shown that chemical variation of a melt reactively migrating within gabbroic material may be exemplified by a process of assimilation and fractional crystallization (AFC) involving assimilation by the reactively migrating melt of various proportions of pre-existing minerals (e.g., Lissenberg et al., 2013). To simulate whether a melt–rock reaction can reproduce the Mg# of melts at equilibrium with cummingtonite and to discuss the role of the fractionating assemblage I computed an AFC process involving the dissolution of orthopyroxene during the crystallization of the reactively migrating melt. I used the following parameters. (i) The initial melt composition is that of melt in equilibrium with the cummingtonite showing the lowest Mg# (quartz amphibolite). (ii) The model assumes assimilation of orthopyroxene (Ma = 100 % Opx) and crystallization of cummingtonite and biotite. (iii) The composition of assimilated material is that of orthopyroxene from norites of the Variscan Rovale sequence (Sila Massif, Caggianelli et al., 1994). (iv) The partition coefficients used for the AFC are those of Nandedkar et al. (2014). The Mg# of the melt in equilibrium with cummingtonite from fine-grained Pietre Nere amphibolites was reproduced by a fractionating assemblage of 0.35 cummingtonite + 0.65 biotite and an assimilated mass/crystallized mass ratio (Ma/Mc) of 0.85. The Mg# of the melt in equilibrium with cummingtonite from medium-grained Pietre Nere amphibolites was reproduced by a fractionating assemblage of 0.60 cummingtonite + 0.40 biotite and a Ma/Mc ratio of 0.52. It can be thus concluded that modal and major-element variations among Palmi amphibolites reflect an evolution of the reactively migrating melt characterized by variable fractionating assemblage and assimilation to crystallization rates.

Amphibolites are characterized by variable contents of Allanite, which is relatively frequent in amphibolites interlayered with paragneiss, but it is rare in the Pietre Nere body where it was only found in foliated fine-grained amphibolite. It is noteworthy that amphibolites interlayered with paragneiss are enriched in most incompatible elements (i.e., Rb, Ba, Th, U) and Zr relative to Pietre Nere samples (Fig. 11). Rb and Ba enrichments are paralleled by cummingtonite and hornblende compositions (Fig. 8). Furthermore, cummingtonite, hornblende and plagioclase from amphibolites interlayered with paragneiss have chondrite-normalized MREE and HREE contents higher by about an order of magnitude than amphibole and plagioclase from Pietre Nere amphibol-
lites. These high concentrations could be related to a high contribution from the reactive migrating melt, which could be also responsible for the high K concentrations shown by the whole rock, hornblende and biotite of amphibolites interlayered with paragneiss. In the Pietre Nere body, fine-grained amphibolites have (i) more fractionated LREE–HREE patterns than medium-grained portions (Fig. 11) and (ii) intermediate Rb/N/Y ratios between medium-grained portions and amphibolites interlayered with paragneiss (~1.0, ~0.4 and 1.5–6.1, respectively, Table S6). It is also noteworthy that foliated fine-grained Pietre Nere amphibolite has both bulk rock and amphibole enriched in Rb and Ba relative to other samples from the Pietre Nere body and Zr bulk-rock contents similar to amphibolites interlayered with paragneiss. These chemical signatures could be related to a higher contribute from the migrating melt in fine-grained relative to medium-grained amphibolites. Hence, allanite abundance and trace-element variations among Palmi amphibolites indicate a decrease in REEs and most incompatible-trace-element contents from amphibolite layers within paragneiss to fine- and medium-grained amphibolites of Pietre Nere. This may reflect a progressive depletion of the migrating melt as it reacts and promotes crystallization of minerals (e.g., amphibole and allanite) able to incorporate a relatively highREE amount and most incompatible elements.

In summary, the reactive melt migration imparted a chemical and mineralogical signature reflecting a variable modification of the protolith initial compositions. In particular, the melt–rock reaction determined an enrichment in SiO$_2$, Al$_2$O$_3$, K$_2$O and incompatible elements in amphibolites interlayered with paragneiss relative to those from the Pietre Nere body, most likely promoting the crystallization of minor biotite, hornblende from fine-grained Pietre Nere amphibolite PN2. The amphibole–melt partition coefficients determined experimentally for a melt having SiO$_2$ = 63.9 wt % were used (Nandedkar et al., 2016). $\text{Amp/La}\mid \text{D}$ values are $\text{La} = 0.329$, $\text{Ce} = 0.714$, $\text{Nd} = 2.334$, $\text{Sm} = 4.19$, $\text{Eu} = 5.052$, $\text{Gd} = 5.43$, $\text{Tb} = 6.062$, $\text{Dy} = 6.041$, $\text{Ho} = 6.46$, $\text{Y} = 6.113$ and $\text{Er} = 5.97$. Whole-rock composition of late-Variscan peraluminous granites (Rottura et al., 1993; Fiannacca et al., 2019) are reported for comparative purposes.

Figure 12. Calculated chondrite-normalized (Anders and Ebihara, 1982) RREE and Y compositions of melt in equilibrium with hornblende from fine-grained amphibolite PN2. The amphibole–melt partition coefficients determined experimentally for a melt having SiO$_2$ = 63.9 wt % were used (Nandedkar et al., 2016). $\text{Amp/La}\mid \text{D}$ values for $\text{La} = 0.329$, $\text{Ce} = 0.714$, $\text{Nd} = 2.334$, $\text{Sm} = 4.19$, $\text{Eu} = 5.052$, $\text{Gd} = 5.43$, $\text{Tb} = 6.062$, $\text{Dy} = 6.041$, $\text{Ho} = 6.46$, $\text{Y} = 6.113$ and $\text{Er} = 5.97$. Whole-rock composition of late-Variscan peraluminous granites (Rottura et al., 1993; Fiannacca et al., 2019) are reported for comparative purposes.

The development of biotite, quartz and allanite, in tandem with amphibole, suggests that the reactive migrating melt had a felsic composition. Furthermore, the high MREE and HREE contents and marked negative Sr and Eu anomalies shown by amphibole from amphibolite interlayered with paragneiss indicate that the reacting melt had an evolved trace-element signature (Fig. 8). Notably, the chondrite-normalized patterns of Rb–Ba–Th–Nb–Sr–Zr–Y of quartz and garnet amphibolites nearly coincide with those of late-Variscan, peraluminous granites from central and southern Calabria (Fig. 11). RREE compositions of melts in equilibrium with amphiboles from Palmi amphibolites could give the basis of the olivine–basalt partition coefficient reported by Médard and Grove (2008; $K_{d\text{Ol}} / \text{basalt}^{Fe/Mg} = 0.32$) and orthopyroxene–basalt partition coefficient reported by Mün tener et al. (2001; $K_{d\text{Opx}} / \text{basalt}^{Fe/Mg} = 0.28$), I calculated Mg# of the melt for the least evolved Rovale gabbros, in order to reduce possible contributes from contamination and/or fractional crystallization. Computed Mg# of the melts in equilibrium with the most primitive olivine and orthopyroxene are 41 and 42, respectively. These values are slightly higher than the Mg# calculated at equilibrium with cummingtonite from medium-grained Pietre Nere amphibolites (~37), thereby suggesting that reactive migration of the SiO$_2$-rich hydrous melt imparted minor chemical variation in these rocks. It can be concluded that fine-grained, foliated rocks represented a high-porosity domain enhancing reactive migration of low viscous hydrous melt and that medium-grained domains could represent a low-porosity barrier possibly preventing further melt migration.

6.6 Inferences on the composition of the migrating melt

The development of biotite, quartz and allanite, in tandem with amphibole, suggests that the reactive migrating melt had a felsic composition. Furthermore, the high MREE and HREE contents and marked negative Sr and Eu anomalies shown by amphibole from amphibolite interlayered with paragneiss indicate that the reacting melt had an evolved trace-element signature (Fig. 8). Notably, the chondrite-normalized patterns of Rb–Ba–Th–Nb–Sr–Zr–Y of quartz and garnet amphibolites nearly coincide with those of late-Variscan, peraluminous granites from central and southern Calabria (Fig. 11). RREE compositions of melts in equilibrium with amphiboles from Palmi amphibolites could give...
information about the compositions of the reactively migrating melt. Experimental studies support that melt composition, namely SiO\(_2\) content, is the main parameter controlling the absolute values of REE amphibole–melt partition coefficients (Tiepolo et al., 2007; Nandedkar et al., 2016). The SiO\(_2\) content of the parent melt can be inferred based on the major-element mineral chemistry. In particular, the compilation of amphibole compositions by Putirka (2016) allows for calculation of the SiO\(_2\) content of the hornblende equilibrium melt, thereby permitting the choice of suitable amphibole–melt partition coefficients for the system of interest. Calculations indicate that the melts at equilibrium with hornblende had high SiO\(_2\) contents of \(\sim 60\) wt\% in the quartz amphibolite, \(\sim 63\) wt\% in fine-grained Pietre Nere amphibolites and \(\sim 68\) wt\% in medium-grained Pietre Nere amphibolites. Amphiboles whose LREE and MREE compositions were affected by the simultaneous crystallization of allanite (i.e., fine-grained amphibolite PN1 and quartz amphibolite) were not considered for the calculation of equilibrium melts. Hence, REE and Y equilibrium melt compositions were computed using a set of hornblende–melt partition coefficients obtained experimentally for melts having SiO\(_2\) = 63.9 wt\% (Nandedkar et al., 2016), which is the value obtained for the melt in equilibrium with hornblende from fine-grained amphibolite PN2. Computed REE compositions resemble those of late-Variscan peraluminous granites (Fig. 12). The intrusion in the upper crust of these peraluminous granites was related to a protracted episode of partial melting of lower-crust metasedimentary rocks, which started at \(\pm 300\) Ma and ended at \(\sim 280\) Ma (Fiannacca et al., 2008; Micheletti et al., 2008). A whole-rock chemistry and U–Pb zircon dating study recently proposed that the dominant components of the late-Variscan peraluminous granite’s magma source was a deep-seated metasediment compositionally similar to migmatitic paragneiss widespread along the southern sector of the Calabria–Peloritani terrane (Fiannacca et al., 2019). A hybrid origin for these granites was also proposed because some of their geochemical and isotopic features may reflect a contribution from mantle-derived components (Rottura et al., 1993). I conclude that the hydrous, SiO\(_2\)-rich melt reactivity of migmatites is a viable mechanism by which major and trace elements are transferred from the anatectic source to adjacent mafic rocks. The melt–rock reaction promoted amphibole and accessory mineral (i.e., allanite) crystallization, which played an important role in the distribution of most incompatible trace elements and REEs during the migration of migmatite-related melt in mafic crust out of the anatectic source. The reactive melt migration could derive by partial melting of metasediments akin to neighboring migmatitic paragneiss.

Late-Variscan orogenic collapse of western European Variscides recorded the widespread intrusions of chemically similar basic melts with depleted to slightly enriched isotopic compositions, in different sectors of the original Europe–Adria lithosphere from the crust–mantle boundary (Malenco, Hermann et al., 2001) to the lower crust (e.g., Ivrea Zone, Quick et al., 1994; External Liguride units, Renna and Tribuzio, 2009), mid crust (e.g., Sondalo, Tribuzio et al., 1999; Mont Collon, Monjoie et al., 2005) and shallow crust (Corsica batholith, Renna et al., 2007; Tribuzio et al., 2009). These intrusions were dated from \(\sim 306\) to \(\sim 280\) Ma and were associated with lithosphere thinning/extension and uprise of asthenospheric mantle. The emplacement of gabbroic sequences of western European Variscides was nearly coeval with the proposed late-Variscan intrusions of mantle-derived magmas in several sectors of the Calabria terrane (e.g., Caggianelli et al., 1994; Renna et al., 2020). In a geodynamic scenario characterized by lithospheric extension and attenuation subsequent to the Variscan collision, high heat input from the rising mantle could have contributed to producing partial melting of lower-crust metasedimentary rocks.

7 Summary and concluding remarks

This work deals with a portion of the Variscan lower to intermediate crust exposed in the Palmi area (southwestern Calabria, Italy). Amphibolites occur in close spatial relationship with migmatitic paragneiss, namely as decimeter-thick layers within the migmatites and as a decametric main body (i.e., Pietre Nere body) adjacent to the paragneiss. The decametric body is mainly fine-grained and weakly to moderately foliated; unfoliated medium-grained portions also rarely occur. Amphibolites derived from mafic rocks of cumulus origin. Amphibole development likely resulted from a coupled dissolution–precipitation process as a consequence of the reaction between a migrating SiO\(_2\)-rich hydrous melt and precursor orthopyroxene. This reaction was also responsible for the crystallization of biotite ± allanite ± quartz ± plagioclase and most likely occurred in conjunction with a subsolidus re-equilibration under granulite facies conditions. The SiO\(_2\)-rich hydrous melt had a similar composition to that of late-Variscan peraluminous granites and could derive by partial melting of metasediments akin to neighboring migmatitic paragneiss. Garnet formed by reaction between cummingtonite, plagioclase and biotite in amphibolites interlayered with paragneiss, most likely due to relatively long period of high temperatures generated by anatexis.

This study provides evidence that reactive migmatite-related melt migration is a viable mechanism by which major and trace elements are transferred from the anatectic source to adjacent mafic rocks. The melt–rock reaction promoted amphibole and accessory mineral (i.e., allanite) crystallization, which played an important role in the distribution of most incompatible trace elements and REEs during the migration of migmatite-related melt in mafic crust out of the anatectic source. The reactive melt migration was enhanced in fine-grained, foliated rocks which could represent a high-porosity domain enabling better interaction with precursor minerals and nucleation of new mineral phases. Medium-grained domains preserved compositions more similar to pristine mafic rocks and
could represent a low-porosity barrier possibly preventing further melt migration.

Data availability. All data derived from this research are presented in the enclosed figures and tables.

Supplement. The supplement related to this article is available online at: https://doi.org/10.5194/ejm-35-1-2023-supplement.

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