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| 2 | Phlogopite-Peridotite): Insights into the source of the Triassic-Jurassic magmatism at the |
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27 Abstract

A gabbroic dyke swarm containing magmatic sapphirine occurs in the Finero phlogopite-peridotite 28 (FPP), one of the major mantle massifs in the Ivrea–Verbano Zone (IVZ; western Southern Alps). 29 Sapphirine is part of a particular mineral assemblage, including plagioclase, titanian pargasite, 30 titanian phlogopite, and Cl-rich apatite; the latter mineral hosts calcite inclusions. The dykes cut the 31 mantle foliation at a high angle, are bounded by orthopyroxenite layers, and show symmetric 32 33 internal banding, represented by two outer hornblendite selvages and an inner leucogabbro band. The sapphirine occurs in up to 3 cm-thick irregular patches in both hornblendite salvages, along 34 with Al-rich amphibole and green spinel. 35

We present major and trace elements of minerals and bulk rock, as well as mineral O, Sr, and Nd isotopic compositions of dykes and the host peridotite from two different outcrops in the FPP area.

Our data show that early melt migration developed through porous flow within cm-thick channels 38 39 and was characterised by orthopyroxene dissolution. Following progressive percolation and reaction, the melt became silica saturated with segregation of orthopyroxenite in the centres of the 40 41 channels. The banded internal structure of the dykes was caused by three different evolutionary stages, involving opening and enlargement of the conduits. The sapphirine and green spinel 42 segregation took place at T > 1,000 °C, in the presence of melt with transient composition, which 43 44 interstitially migrated and reacted with the cumulus minerals to form the hornblendite layers. The mineral chemistry of the newly-formed amphiboles indicates that the sapphirine parental melt was 45 Al-rich, depleted to strongly depleted in Hf, Zr, Nb, Ta, Ti, Sc, V, and middle and heavy rare earth 46 elements, and characterised by a positive Eu anomaly and $(Zr/Hf)_N < 1$. These data suggest a 47 parental melt with a significant amount of normative plagioclase. However, the studied veins do not 48 show evidence of plagioclase assimilation, and we argue that this process could have occurred in 49 magmatic bodies that are not outcropping today to the surface or in the melt source. 50

51 The δ^{18} O values of vein amphiboles and plagioclases vary from 6.9 to 8.6‰ SMOW, which is well 52 above the mantle range, even when considering fractionation upon cooling. Given that orthopyroxene from the wall has "normal" mantle δ^{18} O values (5.8‰), reaction with the host metasomatised peridotite cannot be responsible for the heavy δ^{18} O signature, and the latter must have been imparted by crustal components deeper in the mantle.

56 Our petrographic and geochemical evidence demonstrates that the northern IVZ records an 57 extremely prolonged release, from the Variscan orogenic cycle to the Mesozoic exhumation, of K-58 H₂O-rich mantle-derived melts, mixed with subduction-related components. This finding provides 59 valuable insights into the Triassic–Jurassic magmatism and the geodynamic environment at the 60 Europe–Africa boundary.

61

62 **1. Introduction**

It is well known that the subcontinental lithospheric mantle (SCLM) may record significant 63 heterogeneities in terms of both lithology and geochemical composition, resulting from the 64 65 development of different tectono-magmatic events over a long time span (Mukasa and Shervais, 1999; Rivalenti et al., 2007a, b; Mazzucchelli et al., 2009; 2010; 2016; Borghini et al., 2017; 66 67 Princivalle et al., 2014; Ponce et al., 2015; Rocco et al., 2017; Consuma et al., 2019). The petrochemical records of the lithospheric mantle and uprising melts can be very different based on 68 the geochemical affinity of the melt, the composition of the peridotite mantle column, the 69 modalities of melt migration, and the P, T, fO₂, and fH₂ conditions of the system. An exceptional 70 locality where this can be studied in great detail is the phlogopite-peridotite mantle unit of Finero 71 (Cawthorn, 1975; Siena & Coltorti, 1989). The Finero phlogopite-peridotite (FPP) is one of the 72 most studied mantle massifs on Earth. It crops out in the northernmost part of the Ivrea-Verbano 73 Zone (IVZ, western Southern Alps, Italy; Fig. 1) and consists of an association of phlogopite-74 bearing amphibole harzburgites and dunites, both locally associated with phlogopite-bearing 75 76 amphibole pyroxenites. This is the result of an important episode of pervasive to channelled porous flow migration of melts containing a large volume of crustal components that induced virtually 77 complete metasomatic recrystallisation of an older sequence of harzburgite and pyroxenite 78

associations (Lensch, 1968; Cawthorn, 1975; Griffin & O'Reilly, 1986; Siena & Coltorti, 1989; 79 Shervais and Musaka, 1991; Hartmann & Wedepohl, 1993; Zanetti et al., 1999, 2016; Grieco et al., 80 2001, 2004; Morishita et al., 2003, 2008; Raffone et al., 2006; Selverstone & Sharp, 2011; 81 Mazzucchelli et al., 2014; Tommasi et al., 2017). The crustal affinity of these metasomatic melts is 82 implied by high δ^{18} O values and by the Nd, Sr, Pb, H, S, Cl, and noble gas isotopic compositions of 83 the FPP rocks (Hunziker and Zingg, 1982; Voshage et al., 1987, 1988; Cumming et al., 1987; 84 Hartmann and Wedepohl, 1993; Obermiller, 1994; Seitz and Woodland, 2000; Downes, 2001; 85 Matsumoto et al., 2005; Selverstone and Sharp, 2011). The melt migration processes have been 86 mainly attributed to supra-subduction environments (e.g. Zanetti et al., 1999; Grieco et al., 2001, 87 88 2004; Morishita et al., 2003, 2008; Matsumoto et al., 2005), but also to extensional settings (Garuti et al., 2001; Zaccarini et al., 2004). 89

The available geochronological data indicate that this main event of melt migration took place in the Palaeozoic (Zanetti et al., 2016; Malitch et al., 2017), but many field, petrochemical, and geochronological data point to the development of several other tectono-magmatic events in the Mesozoic as well (Stähle et al., 1990, 2001; Grieco et al., 2001; Matsumoto et al., 2005; Morishita et al., 2003, 2008; Zanetti et al., 1999; 2016; Malitch et al., 2017).

95 Giovanardi et al. (2013) studied the occurrence of a late sapphirine-bearing gabbroic dykelet swarm 96 previously reported by Siena & Coltorti (1989) and discovered by M. Mazzucchelli (Giovanardi et 97 al., 2013). Based on a detailed petrographic and geochemical survey, they provided evidence for a 98 multi-stage dyke formation, which involved fractional crystallisation associated with different 99 patterns of melt–rock interactions.

With the aim of better constraining the nature of the primitive melt and petrological processes and inducing the segregation of sapphirine in the dykes, in this paper, new and detailed petrochemical data are reported for dykes and host peridotites close to the area studied by Giovanardi et al. (2013) and on a second finding in an adjacent area. Particular care has been dedicated to characterising the compositional variability in terms of major and trace element mineral chemistry, as well as the 105 mineral O, Nd, and Sr isotopic composition, to document the geochemical fractionation of the 106 flowing melt following reaction with the host peridotite and early cumulates. The genetic and 107 temporal relationships between the dyke emplacement and other events of melt migration recorded 108 by the FPP unit, as well as the related geodynamic scenarios, are also addressed.

109

110 **2. Geological setting**

111 In the Finero area (Fig. 1), the IVZ crops out in a pseudo-antiform structure with the FPP mantle unit at the core, flanked by what has been called the Finero Mafic Complex (Cawthorn, 1975; Siena 112 & Coltorti, 1989; Lu et al., 1997a, b; Zanetti et al., 2013, 2014; Giovanardi et al., 2014; 113 Mazzucchelli et al., 2014; Langone et al., 2017). The Mafic Complex is divided into three different 114 units: the stratigraphically lower Layered Internal Zone unit (LIZ), in contact with the FPP; the 115 Amphibole-Peridotite (Amphibole-Pd); and the External Gabbro (EG). The EG is placed in tectonic 116 117 contact with the Amphibole-Peridotite by a Mesozoic high-T shear-zone (Langone et al., 2018). A tectonic contact also characterises the transition toward the amphibolite-facies metasediments and 118 119 metabasites of the Kinzigite Formation (KF), which represent the metamorphic basement of the 120 Adria Plate. Septa of KF rocks are embedded in the EG.

The Finero mafic-ultramafic sequence presents several differences with respect to the southern and 121 central sectors of the IVZ (i.e. the Baldissero and Balmuccia peridotites and the Val Sesia Complex; 122 Quick et at., 1995; Correia et al., 2012; Mazzucchelli et al., 2014 and references therein), providing 123 evidence of a different geological evolution (Zanetti et al., 2013, 2014, 2016; Langone et al., 2017, 124 2018). In particular, the FPP has been completely recrystallised by several events of migrating melts 125 with crustal affinity (Zanetti et al., 1999; 2016; Mazzucchelli et al., 2014 and references therein), 126 and the relationships with the parental melts of the surrounding Mafic Complex are far from being 127 understood (Giovanardi et al., 2014). Conversely, the peridotites cropping out in the central and 128 southern part of IVZ did not undergo similar melt-induced recrystallisation and were emplaced in 129

the Kinzigite Formation as tectonic slivers before the intrusion of the central and southern Mafic
Complex (Quick et at., 1995; Mazzucchelli et al., 2014 and references therein).

The FPP is mainly represented by phlogopite-bearing amphibole harzburgites and associated 132 phlogopite-bearing pyroxenites (Cawthorn, 1975; Siena & Coltorti, 1989; Hartmann & Wedephol, 133 1993; Zanetti et al., 1999; Raffone et al., 2006; Selverstone & Sharp, 2011; Mazzucchelli et al., 134 2014; Giovanardi et al., 2018). These lithologies derived from pervasive metasomatism of a 135 136 depleted peridotite, which formed secondary orthopyroxene, amphibole, and phlogopite (Zanetti et al., 1999; Tommasi et al., 2017). Channelled migration events formed dunite bodies containing 137 stratiform to podiform chromitites and, rarely, pyroxenite and hornblendite layers (Grieco et al., 138 2001, 2004; Zanetti et al., 2016). 139

Late stages of porous-flow melt migrations crystallised apatite- and dolomite-bearing wehrlites and apatite-rich, orthopyroxene-bearing peridotites, which sometimes contain carbonate-bearing domains showing marked modal and geochemical gradients with the host rocks (Zanetti et al., 1999; Morishita et al., 2003, 2008; Matsumoto et al., 2005; Raffone et al., 2006). U–Pb analyses on apatite and isotopic noble gas data have yielded Triassic ages (Morishita et al., 2008).

Apatite and calcite also occur in dykes of nepheline-bearing syenites of the Triassic age, associated
with hornblendites (Stähle et al., 1990, 2001). An Early Jurassic U–Pb zircon age was determined
for an alkali pegmatite (Grieco et al., 2001).

The rocks of the FPP related to the main event and the dunite bodies show similar geochemical 148 features and absence of geochemical gradients. The harzburgite-pyroxenite association and the 149 chromitite and pyroxenite layers in dunite bodies are depleted in Nb and high field strength 150 elements and significantly enriched in K, Rb, Ba, Sr, and light rare earth elements (LREEs) 151 (Hartmann & Wedephol, 1993; Zanetti et al., 1999, 2016; Mazzucchelli et al., 2014). The 152 mineralogical and compositional features have been considered by several authors as evidence of 153 the role of a slab-derived crustal component in the percolating melts (Hartmann & Wedephol, 1993; 154 Mazzucchelli et al., 1995, 2016; Rivalenti et al., 1995, 2007a; Zanetti et al., 1999, 2016; Grieco et 155

al., 2001, 2004; Morishita et al., 2003, 2008; Ponce et al., 2015; and others). Thus far, a variety of 156 different geochemical components has been identified based on isotopic data. In particular, the Hf 157 (in zircon) and O (in zircon and pyroxene) isotopic compositions of chromitites suggest the 158 presence of large volumes of continental crust in the migrating melts (Zanetti et al., 2016; Malitch 159 et al., 2017), whereas the hydrogen, oxygen, and chlorine isotopic compositions of amphibole and 160 phlogopite from the harzburgite-pyroxenite association show variability (δD from -29 to -86‰, 161 δ^{18} O from 4.9 to 6.1‰, and δ^{37} Cl from -2.0 to +2.1‰) consistent with mixtures of magmatic fluids 162 contaminated by seawater (Hartmann & Wedepohl, 1993; Selverstone & Sharp, 2011). 163

Giovanardi et al. (2013) reported the occurrence in the FPP of late sapphirine-bearing gabbroic dykes. These dykes crosscut at high angles the pervasive mantle foliation and other lithologies, showing different mineralogical and major element mineral chemistry features with respect to other FPP rocks.

168

169 **3. Samples and petrography**

Two different dykes and their host peridotites from the FPP unit were investigated. The first dike (sample FI09C06, Fig. 2) was collected along the road that connects National Road 631 to a peridotite quarry located on the right flank of the Rio Creves valley (less than 100 m from the outcrops studied by Zanetti et al., 1999). The second dike (sample FI9664) is from a boulder along the Rio Creves, about 30 m upstream of its intersection with the Rio Cannobino. The host peridotite was collected 8 cm from the FI09C06 dyke, and close to the contact with the dyke represented by sample FI9664.

The gabbroic dykes are centimetric in thickness (mostly 2–5 cm; Fig. 2). They show variable
strikes, usually crosscutting the harzburgite-pyroxenite association at high angles (Fig. 2).

A Mesozoic age for these sapphirine-bearing dykes is constrained by the observation that they also crosscut the foliation of protomylonites in the external domains of Mesozoic shear zones (Matysiak & Trepmann, 2015 and references therein), but are themselves deformed in a few cm-wide mylonitic to ultramylonitic bands parallel to the protomylonite foliation (Tommasi et al., 2017). Such shear zones were active at different crustal levels over a very long, Triassic–Jurassic time interval (235–180 Ma; Langone et al., 2018 and references therein). This finding is also consistent with the observation that all the late intrusive bodies, which are discordantly intruding the pervasive mantle foliation by hydraulic fracturing, do not show ages older than Triassic (Stähle et al., 1990, 2001; Grieco et al., 2001; Matsumoto et al., 2005).

The gabbroic dykes contain sapphirine (Giovanardi et al., 2013) and show a banded symmetric structure consisting of melanocratic zones at the peridotite contacts and a leucocratic zone representing the dyke core (Fig. 2). The melanocratic zones, in turn, can be divided in three different bands.

192 From the host peridotite to the dyke core, the following layers can be recognised:

An orthopyroxenite zone (hereafter, the Orthopyroxene Zone), established within the ambient
 peridotite. Orthopyroxene locally shows recrystallised rims with the growth of fine neoblasts of
 orthopyroxene and rarely olivine. The orthopyroxene sometimes presents exsolution lamellae.
 Locally, black spinel and phlogopite occur as accessory phases. Phlogopite is concentrated in
 interstitial positions but rarely fills fractures within orthopyroxene crystals (Fig. 2). Giovanardi et
 al. (2013) reported the occurrence of a sapphirine- and amphibole-bearing recrystallisation front in
 the Orthopyroxene Zone.

200 2) A first melanocratic zone (hereafter, the Early Amphibole Zone) inside veins formed by dark-201 brown amphibole (up to 1 cm long, named 'early amphibole') and associated small plagioclase 202 grains, spinel, phlogopite, and apatite; in these zones, the magmatic texture is preserved, as 203 evidenced by amphibole twinning. Apatite and spinel mainly occur as rounded inclusions in 204 amphibole, whereas phlogopite is in interstitial position.

3) A second melanocratic zone (hereafter the Late Amphibole Zone), consisting of light-brown to
green amphibole (named 'late amphibole'), green spinel, sapphirine, and phlogopite. Sapphirine
occurs in three textural positions: inclusions within late amphibole, coronas rimming spinel, and

isolated/aggregated crystals in interstitial positions (Fig. 2). Phlogopite is an accessory phase in 208 interstitial positions. Late amphibole is smaller than early amphibole and is euhedral to anhedral in 209 shape. Intermediate green-brown amphibole, often associated with spinel and sapphirine, is 210 211 recognised near and through the Early Amphibole Zone-Late Amphibole Zone contact. Recrystallisation zones with a fine-grained texture occur. The Late Amphibole Zone is not 212 continuous through the dykes. It forms patches, which can also occur only on one side of the 213 leucocratic core of the dyke, or that can extend up to the Orthopyroxene Zone. The Late Amphibole 214 215 Zone is more developed in sample FI9664 (up to about 1.5 cm in thickness) than in sample FI09C06 (up to 1 cm in thickness). In addition, sapphirine crystals in sample FI9664 can reach up to 216 217 1.5 mm in size, whereas in sample FI09C06, they are commonly < 0.2 mm.

4) A leucocratic zone (Leucocratic Zone, hereafter) formed by plagioclase and subordinate 218 amphibole (both magmatic and relict from melanocratic zones). Apatite occurs as an accessory 219 phase, sometimes included in plagioclase or in relicts of brown early amphibole. The apatite rarely 220 contains calcite inclusions. Plagioclase shows twinning (mainly pericline and subordinately crossed 221 222 albite and pericline), which is often partially or totally erased by recrystallisation induced by late 223 deformation. Magmatic amphibole has greenish pleochroism, whereas relict amphibole, ripped from the melanocratic zones, is brown. Amphibole often forms single-crystal alignments parallel to the 224 dyke strike, like in a flow texture. Phlogopite is rare and is associated with amphibole. 225 Recrystallisation zones show fine-grained equigranular texture. Rarely, Fe-oxides, Fe-Ni sulphides, 226 and pyrite occur. 227

The host peridotite away from the contact is a hornblende-harzburgite in modal composition with a porphyroclastic texture (Giovanardi et al., 2018). It is characterised by the presence of olivine and orthopyroxene porphyroclasts, with a secondary, undeformed mineral assemblage dominated by amphibole (15% by volume), in association with orthopyroxene, spinel, phlogopite, and clinopyroxene, strictly similar to the dominant peridotite type described by Zanetti et al. (1999) and Tommasi et al. (2017). A detailed petrographic inspection highlights the occurrence of a reacted peridotite zone approaching the orthopyroxene zone. It is characterised by the presence of a secondary mineral assemblage, modally dominated by long (up to 5 mm) phlogopite lamellae, with associated subordinate amounts of undeformed orthopyroxene, spinel, amphibole, and clinopyroxene. In the reacted zone, olivine was stable; conversely, the modal orthopyroxene content is slightly lower than that in the peridotite far from the vein. Primary (e.g. olivine) and secondary minerals in the reacted zones display elongation sub-parallel to the present-day vein strike.

240

241 4. Analytical methods

Sample FI9664, representing the gabbroic dyke and the contact host harzburgite, was analysed for whole-rock major and trace elements. Whole-rock major elements and Sc, V, Cr, Co, Ni, Cu, and Zn were analysed by X-ray fluorescence spectrometry (XRF), whereas Li, Rb, Sr, Y, Zr, Nb, Ba, rare earth element (REE), Hf, Ta, and Pb contents were analysed by inductively coupled plasmamass spectrometry (ICP-MS). Analyses were performed following the methods described by Mazzucchelli et al. (2010) (data reported in Supplementary Material A).

Samples FI9664 and FI09C06 were analysed for mineral major elements with the electron
microprobe JEOL 8200 Super Probe housed at the University of Milano. Analyses were performed
following methods described by Ponce et al. (2015). Data are reported in Supplementary Material
A.

Mineral trace elemental contents were determined (as in Rivalenti et al., 2007b) with a LA-ICP-MS 252 housed at Istituto di Geoscienze e Georisorse, Consiglio Nazionale delle Ricerche, Sede Secondaria 253 di Pavia (data are reported in Supplementary Material A) using a Perkin Elmer SCIEX DCR-e 254 coupled with a solid-state laser source (Q-switched Nd:YAG, Quantel Brilliant). Data reduction was 255 performed using the GLITTER software (Griffin et al., 2008). NIST SRM 610 was used as an 256 257 external standard. Ca was used as an internal standard for clinopyroxene, amphibole, and plagioclase, Si for olivine, orthopyroxene, and phlogopite, and Mg for spinel. Precision and 258 accuracy were assessed by repeated analysis of the standard BCR-2g and were found to be better 259

than 10% at the ppm concentration level. Further information is reported by Giovanardi et al.(2017).

Sample FI09C06 was selected for O, Nd, and Sr isotopic analyses in mineral separates, after the combination of preliminary electron microprobe and LA-ICP-MS analyses suggested a more primitive nature for its parental melt.

O isotopes of pure mineral separates from sample FI09C06 were analysed at the I.G.G.-C.N.R., Pisa, by conventional laser fluorination with a Finnigan Delta Plus mass spectrometer. For each phase, 1–1.5 mg aliquots were necessary to measure the oxygen isotope composition. Analyses were performed following the methods described by Perinelli et al. (2011). Data are reported in Table 1.

Sr and Nd isotopic ratios of amphibole and plagioclase separates from sample FI09C06 and the host 270 harzburgite were analysed at the Marine Environmental Sciences Laboratory (LEMAR) of the 271 272 Institut Universitaire Européen de la Mer (IUEM), Université de Bretagne Occidentale. Analyses were carried out after dissolution and chromatographic separation using a TRITON Thermo-273 274 Ionisation Mass Spectrometer (TIMS), following the procedure described by Janin et al. (2012). Analyses were corrected for the reference material NBS987 (87 Sr/ 86 Sr = 0.710241 ± 0.000019, n = 275 6) for Sr and the La Jolla standard (143 Nd/ 144 Nd = 0.511847 ± 0.000009, n = 3) for Nd. Data are 276 277 reported in Table 2.

278

279 **5. Bulk rock chemistry**

The gabbroic dyke FI9664 presents higher TiO₂, Al₂O₃, CaO, Na₂O, and K₂O contents and lower
FeO and MgO contents than the host harzburgite collected at the contact.

The gabbroic dyke has trace element abundances that are one order of magnitude higher than those of the host harzburgite; only Rb and Li contents are similar in both rocks (Fig. 3; Supplementary Material A). The two rock types show some similarities in element fractionation: marked enrichments in LREEs with respect to middle-heavy rare earth elements (M-HREE; (La/Yb)_N values are 5.5 and 16–30 for the dyke and host harzburgite, respectively; primitive mantle data from McDonough and Sun, 1995); (Th/U)_N and (Zr/Hf)_N always < 1; and positive Ba and Pb anomalies. Conversely, the (Nb/Ta)_N values are 1.6 for the gabbroic dyke and 0.7 for the host harzburgite. Linearly fractionated LREE-enriched patterns were also shown by the nepheline-bearing alkaline

dykes described by Stähle et al. (2001). However, the latter had significantly higher trace element
contents than those of gabbroic dyke FI9664. The dykes reported by Stähle et al. (2001) also
displayed significant peculiarities in terms of fractionation of highly incompatible trace elements,
such as U, Th, Nb, Ta, Ba, and Pb, with respect to REEs.

The spidergram of the peridotite close to the contact with dike FI9664 shows some relevant differences with respect to those reported by Hartmann & Wedephol (1993): large positive Pb and Hf anomalies, greater Rb, U, Ba, Ta, and Nb contents, and slightly lower LREE contents (Fig. 3).

297

298 6. Major element mineral chemistry

299 6.1. Host harzburgites

The mineral composition of the host harzburgite far from the veins is similar to that of the harzburgite-pyroxenite association reported by Zanetti et al. (1999). The amphibole is pargasitic in composition, with only one analysis of Mg-hornblende composition. The mica is phlogopite, and the clinopyroxene is diopside.

Minor differences include a slightly Fe-enriched composition shown by olivine (Fo = $100 \times Mg/(Mg + Fe^{2+}_{tot})$, molar ratio of 90.4–91.1), orthopyroxene, spinel, and amphibole (Fig. 4). Orthopyroxene and spinel (hercynite to spinel in composition) are also richer in Al. Because of their low Cr#, the FI09C06 oxides straddle the hercynite–spinel boundary, whereas oxides in the FPP are chromite. Lower Cr content is shown by amphibole, which is also characterised by lower Na and higher K contents.

At the contact with the vein, the Mg# does not change significantly in the minerals. Aluminium decreases in orthopyroxene, spinel, amphibole, and phlogopite, whereas Ti content is higher in both

amphibole and phlogopite. In amphibole, Na decreases, balanced by higher K content (Fig. 5). 312 Conversely, the Na/K ratio is very variable in phlogopite. 313

Orthopyroxene from the Orthopyroxene Zone has lower Mg# and CaO content and higher Al₂O₃ 314 content (Fig. 4). Similarly, its composition is different from orthopyroxene from the sapphirine-315 bearing rock found in the LIZ northern unit of the Finero Mafic Complex by Sills et al. (1983). The 316 latter has lower Mg# and higher Al₂O₃ content (Fig. 4). 317

318

6.2. Gabbroic dykes 319

Notwithstanding the similar internal banding, the two studied dykes show marked differences in 320 terms of major elemental mineral chemistry. Significant compositional changes are also shown for 321 the different types of amphibole (i.e. the Early and Late Amphibole Zones and the Leucocratic 322 Zone; Fig. 6). The unique common feature of the amphiboles of both gabbroic dykes is the higher 323 324 Al content exhibited by the late amphibole (Giovanardi et al., 2013). Amphibole from gabbroic dykes is mostly pargasitic in composition (unit formula calculated according to Ridolfi et al., 2018), 325 326 but sometimes Al substitution for Si in late amphibole is higher than 2 a.p.f.u., thereby entering the sadaganaite compositional field. 327

In particular, amphibole from sample FI09C06 has distinctly lower Mg# (0.73-0.82) than sample 328 FI9664 (0.85-0.87). In sample FI09C06, Mg# is lower in early amphibole than in late and 329 leucocratic amphibole, whereas it is exactly the reverse in sample FI9664 (Fig. 6). 330

FI09C06 amphibole also displays the lowest CaO and Na₂O and the highest TiO₂ and K₂O contents 331

(TiO₂ 0.27–2.19 wt.% and K₂O 0.31–1.28 wt.% respectively) (Fig. 6). 332

In sample FI09C06, TiO₂ linearly increases with the decrease in Mg#. Instead, TiO₂ is higher in the 333 FI9664 early amphibole than in the leucocratic amphibole, but distinctly higher TiO₂ contents are 334 shown by late amphibole. Moreover, in sample FI9664 early amphibole, the increase in Mg# is 335 336

As a whole, amphibole from the gabbroic dykes has higher Al_2O_3 content and lower Mg# and Cr_2O_3 content with respect to others FPP lithologies, resulting in similarity to the amphibole from the sapphirine-bearing rock in the LIZ (Sills et al., 1983) (Fig. 6). Amphibole from the two gabbroicdykes samples shows lower CaO and K₂O and higher Na₂O contents than the respective host amphibole (Fig. 6).

Phlogopite from sample FI9664 presents a narrow range of major elemental contents with respect to
sample FI09C06. In sample FI9664, phlogopite has higher Mg# values than phlogopite from sample
FI09C06 (0.92–0.94 and 0.70–0.90, respectively) and commonly lower TiO₂ contents (Fig. 5).
Phlogopite from the gabbroic dykes shows higher Al₂O₃ content and lower Mg# than the phlogopite
from the host peridotite and the harzburgite-pyroxenite association (Fig. 5).

Spinel is mainly found in the Late Amphibole Zone. Unlike the spinel from the host harzburgite, the harzburgite-pyroxenite association reported by Zanetti et al. (1999), and the chromitite layers in dunite bodies (Grieco et al., 2001; 2004; Zanetti et al., 2016), the spinel from the gabbroic dykes does not contain Cr_2O_3 and can be classified as spinel (Mg# 0.46–0.75).

Plagioclase from sample FI9664 is commonly more anorthitic than the plagioclase from sample FI09C06 (An contents of 82–93 and 36–87, respectively). In both samples, some reversely zoned plagioclases are recognised.

The sapphirine composition falls near the 7:9:3 composition on the SiO_2 -(FeO + MgO)-(Al₂O₃ + Cr₂O₃ + Fe₂O₃) diagram (Fig. 7). Sapphirine from sample FI9664 is higher in Al₂O₃ content with respect to sample FI09C06, whereas it is lower in SiO₂ content.

357 Scanning electron microscope (SEM) investigations at the Centro Interdipartimentale Grandi 358 Strumenti laboratories of the Università di Modena e Reggio Emilia, performed with an 359 Environmental SEM Quanta-200 (Fei Company-Oxford Instruments), suggest that apatites from 360 different zones of the dykes are Cl-apatites.

361

362 7. Trace elemental compositions

363 *7.1. Host harzburgite*

Amphibole and clinopyroxene from harzburgite FI09C06, 8 cm away from the contact with the gabbroic dyke, are characterised by LREE-enriched linearly fractionated patterns (Figs. 8, 9 and 10). Their compositions are similar to those of the FPP harzburgite-pyroxenite association (Zanetti et al., 1999). Proceeding toward the contact (i.e. sample FI9664), amphibole and clinopyroxene show upward-convex REE patterns characterised by high variability in absolute contents (Figs. 8, 9 and 10). These patterns are similar to those from apatite-rich domains reported by Zanetti et al. (1999) (Figs. 9 and 10).

371

372 *7.2. Gabbroic dykes*

The two gabbroic dykes show different trace element compositions. In sample FI09C06, 373 amphiboles from the Early Amphibole Zone and Leucocratic Zone have L-MREE-enriched 374 375 upward-convex patterns, similar to those of harzburgite near the contact (Fig. 8). In the Late Amphibole Zone, some crystals show more fractionated patterns characterised by lower M-HREE 376 377 contents and positive Eu anomalies (Fig. 8). These variations are associated with marked depletion in Ta, Zr, Hf, Y, Sc and V (Fig. 10, V is not shown). As a whole, amphiboles from different zones 378 of sample FI9664 show more fractionated REE patterns characterised by enrichment in LREEs and 379 depletion in HREEs (Fig. 8) with a nearly flat pattern in the LREE region. Amphibole from the 380 Early Amphibole Zone of sample FI9664 presents a small positive Eu anomaly ((Eu/Eu*)_N within 381 1.04–1.65), which becomes more evident in the amphibole from the Late Amphibole Zone 382 ((Eu/Eu*)_N within 1.29–1.62). Amphibole from sample FI9664 is also more enriched in Th, U, and 383 Pb (and to lesser extent in Sr) compared with amphibole from sample FI09C06 (Fig. 10). 384

Similar to the amphibole, REE patterns of plagioclase from sample FI9664 are more fractionated than those of plagioclase from sample FI09C06, (Fig. 8). In both samples, the trace elemental abundances of the plagioclase cores are greater than those of the plagioclase rims. Apatite displays the typical LREE enrichment and high Th, U and Pb contents (12.31–17.35 ppm, 4.35–5.92 ppm, and 2.18–4.19 ppm, respectively). Nb, Ta, Zr, Hf, Ti, and Sc show negative anomalies with values that are often below the detection limit.

391 No systematic trace elemental variations were found in phlogopite.

392

393 8. O isotopes

The δ^{18} O in silicates of sample FI09C06 shows a steady increase from the contact (Orthopyroxene Zone) to the leucocratic gabbro in the vein core, through the Early and Late Amphibole Zones (Fig. 8, 10, and 11 and Table 1). In particular, it varies from 5.81‰ (standard deviation (std. dev.) 0.11) in orthopyroxene from the Orthopyroxene Zone to 6.9‰ (std. dev. 0.05) in amphibole from the Early and Late Amphibole Zones, and to 8.60‰ (std. dev. 0.01) in the plagioclase of the gabbroic core.

The δ^{18} O values of orthopyroxene from the Orthopyroxene Zone lie within the mantle range. They are higher than the δ^{18} O value reported by Hartmann & Wedephol (1993) for clinopyroxene from the phlogopite-bearing amphibole harzburgite (Fig. 11), and for orthopyroxene from olivinechromitites of the FPP, but significantly lower than the orthopyroxene from orthopyroxenechromitites (Zanetti et al., 2016).

Vein amphibole and plagioclase have δ^{18} O values significantly higher than the mantle range (Fig. 11). They are close to the highest values found in amphibole, phlogopite, orthopyroxene, clinopyroxene, and zircon from the phlogopite-bearing amphibole harzburgite-pyroxenites (Hartmann & Wedephol, 1993; Selverstone and Sharp, 2011) and chromitite layers of the FPP (Zanetti et al., 2016).

410 The δ^{18} O values obtained for the green spinel associated with sapphirine are markedly lower than 411 those of the associated late amphibole (δ^{18} O = 4.38‰, std. dev. 0.10), as expected based on crystal-412 chemical constraints (see Bindeman, 2008 and references therein).

413

414 9. Sr and Nd isotopes

The trace elemental concentrations of the mineral separates of early and late amphibole match the 415 differences highlighted by LA-ICP-MS analysis of thin sections. The Sr and Nd isotopic 416 compositions of early amphibole and late amphibole from the FI09C06 dyke are coincident within 417 analytical uncertainty (Table 2). Plagioclase from the FI09C06 Leucocratic Zone shows the same Sr 418 isotopic composition of the amphiboles, and only a slightly more radiogenic ¹⁴³Nd/¹⁴⁴Nd ratio. 419 Similar Sr isotopic compositions were documented in amphibole and apatite of discordant veins 420 from the FPP, which were characterised by higher ¹⁴³Nd/¹⁴⁴Nd ratios (Morishita et al., 2008). The 421 isotopic composition of the dyke minerals is enriched with respect to the depleted mantle reference 422 value and the mid-oceanic ridge basalt (MORB) field; these minerals fall within the ocean island 423 basalt (OIB) field (Fig. 12). In particular, they lie between the isotopic compositions of FPP 424 hornblende-syenite dykes (bulk rock from Stähle et al., 1990, 2001) and peridotites (amphibole and 425 426 clinopyroxene from Obermiller, 1994). Amphiboles separated from the host harzburgite (collected far from the contact) show the most radiogenic Sr and unradiogenic Nd values, but have never 427 428 before been documented in the literature for FPP rocks (Fig. 12).

429

430 **10. Discussion**

431

432 10.1. Petrogenesis of the sapphirine-bearing gabbroic dykes

The two veins studied here share exactly the same internal banding as the sample described by Giovanardi et al. (2013). This suggests that all three veins were part of an interconnected swarm, which allowed them to record the same sequence of petrologic events. The symmetric structure of the dykes suggests that the intrusion occurred through several events of melt percolation. In the following section, we discuss, in detail, the different stages and processes that produced the gabbroic dyke swarm.

440 *10.1.1. Melt percolation though the peridotite*

According to the process governing the emplacement of other dyke swarms in mantle sectors (e.g. Mazzucchelli et al., 2010 and references therein), the melts flowing in the metasomatic haloes and in the conduit were originally similar, undergoing severe fractionation owing to assimilation and fractional crystallisation.

The characterisation of the wall peridotite has revealed some peculiarities. It is a common 445 observation that metasomatic haloes of cm-to-m-scale surround late dykes and veins formed by 446 melt segregation under mantle conditions. In most cases, the metasomatic haloes show marked 447 geochemical and mineralogical gradients as a function of the distance from the dike contact (Zanetti 448 et al., 1999; Ionov et al., 2002, Mazzucchelli et al., 2010; Borghini et al., 2016, 2017), which are 449 interpreted as the result of porous flow percolation of melt escaping from the open conduits. 450 Conversely, no progressive mineralogical or compositional variation is documented in the wall 451 452 peridotite of sample FI9664 at a variable distance from the vein (see Supplementary Material B). Newly-formed clinopyroxene actually shows trace element heterogeneities, but these are randomly 453 454 distributed. Conversely, the trace element composition of amphibole is very homogeneous, indicating late crystallisation from a unique melt. 455

The particular L/MREE-enriched convex-upward patterns shown by clinopyroxene and amphibole from the wall peridotite of sample FI9664 also demonstrate that their parental melt could not have been the same as that from which the adjacent early amphibole of dyke FI9664 crystallised (which show linearly-fractionated REE patterns). However, based on the amphibole trace element composition, the melt recorded by the wall peridotite of sample FI9664 must have been quite similar to the melt that produced the Early Amphibole Zone of dyke FI09C06.

The textural evidence that primary and secondary minerals from the metasomatic haloes display elongation sub-parallel to the present-day vein strike confirms that the development of melt migration channels was associated with some local deformation. These observations can be reconciled by assuming that the metasomatic haloes were not produced by melt escaping normally from the conduit. They were likely established during an early stage of melt migration via focused porous flow along channels developed in correspondence with structural weakness, with a direction roughly parallel to the strike of the present-day veins.

The absence of resorption textures in olivine of the reacted wall peridotite indicates that this mineral was stable. Conversely, the modal content of orthopyroxene is slightly lower than in the peridotite far from the vein. This evidence leads us to consider that the orthopyroxene saturation of the melts, indicated by the segregation of the reactive Orthopyroxene Zone and the presence of few orthopyroxenes in the veins (Giovanardi et al., 2013), was not a clear, primary characteristic, but possibly a consequence of an early stage of reactive melt migration characterised by olivine precipitation and orthopyroxene dissolution (see Piccardo et al., 2017).

The mineralogical mismatch between the nearly monomineralic, hydrous-mineral-free 476 477 Orthopyroxene Zone and the Early Amphibole Zone suggests that they were segregated by compositionally different melts. This hypothesis is confirmed by large differences in terms of O 478 479 isotopic compositions recorded by the orthopyroxene (5.8‰ standard mean ocean water (SMOW)) 480 from the Orthopyroxene Zone and the amphibole from the Early Amphibole Zone (6.9‰ SMOW) within the vein. The lighter O isotopic composition of the Orthopyroxene Zone would support a 481 hybrid composition of its parental melt caused by reactive porous flow through the FPP, with 482 buffering of O isotopic composition at mantle values. 483

This conclusion led us to consider the possibility that the Orthopyroxene Zone formed before the segregation of the Amphibole Zones, possibly at the centre of the migration channels, as observed elsewhere in the FPP. It was successively split into two parts by the opening of the fracture.

487

488 10.1.2. Melt segregation in the open conduit

489 The Early Amphibole Zone was segregated when melts started flowing in open fractures. It is likely 490 that amphibole crystallisation was triggered by the presence of an ultramafic wall, suggesting that 491 plagioclase was unstable in contact with the orthopyroxenite layers. The development of 492 amphibole-rich selvages where hydrous gabbroic rocks come in contact with ultramafic layers (e.g. 493 in the LIZ of the Finero Complex; Mazzucchelli et al., 2014) has previously been documented in the 494 literature.

The two gabbroic dykes of this study show significant differences in terms of major and trace 495 element mineral compositions. In particular, amphibole from the FI9664 vein shows an unusual 496 497 relative enrichment in highly incompatible trace elements (U, Th, LREEs, and Na) and compatible elements (Mg, Ca, and Cr) with respect to those from the FI09C06 vein. Consistently, phlogopite is 498 more enriched in Mg and Na, and plagioclase is more anorthitic. Assuming that the dyke swarm 499 was produced by the injection of a unique melt, this relationship cannot be reconciled based on a 500 melt evolution governed only by fractional crystallisation. Mazzucchelli et al. (2010) documented 501 502 the same correlation between compatible and incompatible elements in minerals from cm-thick diorite dykes, demonstrating that such geochemical features can be modelled considering an 503 assimilation of host minerals concomitant with fractional crystallisation (i.e. assimilation and 504 505 fractional crystallisation (AFC) processes). Accordingly, we argue that the parental melt of vein minerals in sample FI9664 can be considered more evolved through AFC processes than that of 506 dyke FI09C06. The concomitant enrichment of Na and Ca in the vein minerals of sample FI9664 507 508 evidences the role of amphibole in the assimilated component. This is also confirmed by the trace element patterns of FI9664 vein amphiboles, in which highly incompatible elements (such as 509 LREEs, Th, and U) increase and moderately incompatible element (such as HREEs, Ti, and Y) 510 decrease, moving toward the fractionation shown by those of host harzburgite away from the 511 contact. The decrease of moderately incompatible elements with the progression of the melt 512 evolution may be indicative of Amphibole/LD higher than 1 (Tiepolo et al., 2007). It is noteworthy that 513 514 the Al content in sample FI9664 early amphibole is comparable with to slightly lower than in those from the FI09C06 vein, suggesting that the assimilation of host minerals was not effective in 515 boosting the Al concentration of the evolving melt. 516

518 10.1.3. Formation of sapphirine-bearing patches

The mm-to-cm-thick sapphirine-bearing patches and stripes are randomly distributed from the 519 internal end of the hornblendite seam to the Orthopyroxene Zone, although they are generally 520 concentrated towards the centres of the veins. They are apparently the results of interstitial 521 migration of a melt in chemical disequilibrium with the early cumulus minerals, as indicated by the 522 recrystallisation fronts inside large early amphiboles. Both interstitial and recrystallised patches 523 show that the injected melt was saturated in late amphibole, sapphirine, and spinel, whereas the 524 saturation in phlogopite is uncertain, because it occurs only interstitially and the textural 525 relationships are equivocal. 526

The following series of petrographic and geochemical features suggest that the sapphirine-bearing mineral assemblages were not simply related to interaction with the parental melt of the Leucocratic Zone, but were segregated because of the injection in the dyke swarm of an additional melt component coming from outside the system:

531 1) Early amphibole at the contact with or embedded by the plagioclase of the Leucocratic Zone532 does not show any evidence of comparable reaction.

2) Notwithstanding the significantly different major elemental composition of early amphibole 533 documented in the two veins of this study, the major elemental chemistry of sapphirine-534 associated late amphibole converges toward specific Al and Mg# values. Phlogopite shows 535 consistent variation, even though its attribution to early or late amphibole assemblages is 536 doubtful because of its interstitial position. The data reported by Giovanardi et al. (2013) 537 confirm this trend. This finding indicates that a unique component with a specific 538 composition was responsible for the sapphirine segregation in the three sectors of the dyke 539 540 swarm.

3) Normalised patterns of late amphibole from sapphirine-bearing areas show particular
features: stronger LREE/HREE fractionation, depletion in M-HREEs, Zr, Hf, Nb, Ta, Ti, Sc,

and V, and the appearance of a positive Eu anomaly. The results also show inversion of the (Zr/Hf)_N values, which are > 1 in the early amphibole, but < 1 in the late amphibole. The overall fractionation mimics that normally exhibited by plagioclase.

546 The latter evidence, consistent with the apparent increase of the Al content in the system, suggests 547 that the injected melt was characterised by a high normative plagioclase content.

548 The O, Nd, and Sr isotopic compositions of early and late Amphibole are very similar, suggesting 549 that the plagioclase component belonged to the same magmatic cycle that produced the dyke 550 swarm.

However, petrographic surveying has not provided evidence of plagioclase assimilation and/or replacement by amphibole or another mineral, excluding plagioclase components derived by local assimilation. Thus, we suggest that the plagioclase signature was acquired by the injected melt through assimilation of cumulus plagioclase in hidden magmatic bodies and/or addition in the melt source, where it was likely present as a high-P metamorphic equivalent of pristine plagioclase.

556

557 10.1.4. Segregation of the Leucocratic Zone

The petrographic evidence confirms that the formation of the Leucocratic Zone was a high energy event, which determined the partial disaggregation of the Early Amphibole Zone, with evident breaking of large early amphibole crystals. It can be locally recognised that two parts of a formerly unique crystal apparently lie on different sides of a vein. Giovanardi et al. (2013) suggested that this stage was accompanied by enlargement of the conduits.

The Nd and Sr isotopic compositions of the plagioclase indicate that the parental melt had a cognate origin with those of the hornblendite selvages. This inference is also confirmed by the trace element composition of the leucocratic amphibole, which is strictly similar to that of the associated early amphibole.

567 Nevertheless, the observation that the major elemental chemistry of the leucocratic amphibole is 568 intermediate between that of early and late amphibole leads us to conclude that the parental melt of 569 the sapphirine-bearing assemblages could be still present in diluted proportions.

570

571 *10.2. Nature of the parental melts*

Evaluating the nature of the parental melt of magmatic segregates belonging to narrow, cm-thick veins/dykes intruding mantle peridotites at high-T-P mantle conditions must be done with particular caution, with an effort first to characterise the modifications imparted by fractional crystallisation and the reaction/assimilation processes with the host rock (e.g. Mazzucchelli et al., 2010 and references therein). As discussed in the previous section, sample FI09C06 shows that its parental melt underwent the lowest degree of interaction with the host peridotite among the known dykes, and thus is the best candidate to obtain constraints on the nature of the parental melts.

An important contribution in defining the geochemical components in the melts is provided by the marked zoning in terms of O isotopic composition documented in sample FI09C06 between the Orthopyroxene Zone, the hornblendite selvage, and the Leucocratic Zone.

The δ^{18} O values of early and late amphibole (6.9‰ SMOW) well exceed the mantle range 582 (Bindeman, 2008; Polat et al., 2018), suggesting a large contribution by crustal components in the 583 parental melt of sample FI09C06 hornblendite. The proportion of the crustal component is even 584 larger in the leucocratic band, in which a δ^{18} O value of 7.8% SMOW can be calculated for the 585 amphibole in hypothetical equilibrium with the plagioclase composition (8.6% SMOW). This is 586 done by conservatively considering a low closure T for the FPP of ~850 °C (provided by pyroxene-587 solvus geothermometry), an average value of An70 for plagioclase, and the $\delta^{18}O$ mineral 588 fractionation values reported by Bindeman (2008). 589

590 Published data on FPP minerals show a pronounced heterogeneity of the O isotopic composition, 591 with δ^{18} O values from below to well above the mantle range (Hartmann and Wedephol, 1993; Selverstone and Sharp, 2011; Zanetti et al., 2016). This heterogeneity may be due to multiple melt-migration events.

594 Values of δ^{18} O approaching those of early and late amphibole have sometimes been documented in 595 harzburgites and chromitites (Hartmann & Wedephol, 1993; Zanetti et al., 2016), but the values of 596 the leucocratic layers are markedly out of range.

In addition, the δ^{18} O value shown by orthopyroxene (5.81‰ SMOW) from the Orthopyroxene Zone provides a very important constraint, indicating how migrating melts, after prolonged interaction with the FPP, may have oxygen isotopic compositions buffered to mantle values. The decrease of δ^{18} O values in the minerals from the hornblendite selvages with respect to those from the dyke core (i.e. the Leucocratic Zone) confirms such a buffering effect.

We conclude that the high δ^{18} O shown by dyke minerals cannot be the result of interaction between melts and the FPP but is a primary feature of the melts, which indicates the occurrence of large amounts of crustal components, either added to the melt in the source regions or at greater mantle depths.

A subduction-related component was identified based on noble gas isotopic composition in Late Triassic apatite-rich layers (Matsumoto et al., 2005, Morishita et al., 2003, 2008), overprinting the crustal affinity of the host FPP. Amphiboles from such apatite layers show the same radiogenic Sr isotopic composition as the minerals of sample FI09C06, coupled to a slighter more radiogenic Nd isotopic composition.

The isotopic compositions of early amphibole, late amphibole, and plagioclase from the FI09C06 dyke are significantly richer in radiogenic Sr and unradiogenic Nd isotopes compared with those reported from nepheline-bearing Triassic intrusions cutting the FPP (Stähle et al., 1990, 2001), which were interpreted as derived from alkaline melts of OIB affinity.

Combining data from nepheline-bearing alkaline dykes, sapphirine-bearing dykes, and the FPP, we
can define a trend at low radiogenic Nd and high radiogenic Sr isotopic ratios, suggesting mixing
between asthenospheric components (OIB, according to Stähle et al., 1990; 2001 and Schaltegger et

al., 2015) with materials derived from the continental crust (see compositional fields of Casetta et
al., 2018a). We speculate that the crustal components were located at lithospheric deep levels after
the Variscan orogenic cycle (e.g. Bonadiman et al., 1994) and mobilised by asthenospheric
magmatism. This scenario is supported by the evidence of continental crustal metasomatism
exhibited by mantle bodies involved in the Variscan collisional orogeny, such as the FPP, the Ulten
Zone (Sapienza et al., 2009), and some of the Bohemian Massif (Becker et al., 1999; Schulmann et
al., 2014).

In this scenario, it can be noted that the Nd and Sr isotopic compositions of sample FI09C06 lie on the mixing line between the composition of the Triassic alkaline dykes and the host harzburgite away from the contact. According to AFC simulation, assimilation of significant degrees of the metasomatised peridotite material (15%) into OIB must be considered to account for the composition of the parental melt of the FI09C06 veins.

630 Comparison of the trace elemental patterns of the amphibole from the vein apparently less 631 contaminated by the host peridotite (FI09C06) with those of amphiboles segregated by primary 632 hydrous alkaline basalts (e.g. Demeny et al., 2005) shows similarity in REE fractionation, but the 633 early amphibole has an absolute content that is nearly one order of magnitude higher. In addition, a 634 correlation with primary alkaline mantle melts is not directly supported by the negative Nb, Ta, and 635 Ti anomalies shown by the early amphibole.

Equilibrium liquids calculated from the trace elemental composition of early amphibole from the 636 FI09C06 dyke and amphibole-melt partition coefficients experimentally determined at 1,015 °C in 637 the presence of moderately polymerised melts (dataset T2 1015; Tiepolo et al., 2007) match the 638 REE contents and fractionation exhibited by shoshonitic rocks of the Triassic magmatism of the 639 640 Dolomites area (Casetta et al., 2018a; 2018b; see Supplementary Material C). The calculated melts also show consistent negative Ti anomalies, (Nb/La)_N values close to 1, and slight positive U, Th, 641 and Pb anomalies; the latter are more pronounced in the shoshonitic melts. However, a best match 642 in terms of U, Th, and Pb concentrations is shown by equilibrium liquids calculated in equilibrium 643

with amphiboles from 9,664 dykes. These observations suggest that the sapphirine-bearing gabbroic dykes may record the deep mantle input of Triassic K-rich calc-alkaline to shoshonitic melts erupted into the eastern Southern Alps, also documenting the fractionation trends responsible for the enrichment in Th, U, and Pb. This hypothesis is also supported by the Sr and Nd isotopic compositions of Triassic K-rich calc-alkaline to shoshonitic intrusive rocks and lavas of the Eastern Alps (Casetta et al., 2018a; Figure 12), which approach the isotopic trend defined by our model.

650

651 *10.3. Constraints on the geodynamic evolution of the Europe–Africa boundary*

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653 10.3.1. P–T constraints on the FPP

Geothermobarometric estimates constrain the intrusion of sapphirine-bearing dykes at very high-P 654 and T conditions. Ab-initio calculations indicate that the primary crystallisation field of sapphirine 655 in the MAS diagram (SiO₂-MgO-Al₂O₃) becomes considerably larger over 1.0 GPa and shrinks at 656 2.0 GPa (Belmonte et al., 2014). This evidence confirms the experimental results of Liu & Presnall 657 (1990, 2000) and Milholland & Presnall (1998), indicating that magmatic sapphirine in FPP veins 658 likely crystallised at P \geq 1 GPa. Equilibrium T estimates based on the sapphirine-spinel Mg-Fe²⁺ 659 exchange thermometer of Sato et al. (2006) and the amphibole-plagioclase thermometer of Holland 660 & Blundy (1994) are mostly higher than 1,000 °C (up to 1,085 °C), confirming the T estimates of 661 Giovanardi et al. (2013). The melt T was thus significantly higher than that in the host harzburgite, 662 which shows solidus T typically corresponding to the water-oversaturated peridotite (965 $^{\circ}$ C at P = 663 1.1 GPa; Giovanardi et al., 2013). The absence of partial melting evidence in the host FPP confirms 664 that the source of the uprising melts was at greater mantle depths. The high T, in combination with 665 the high water and volatile contents, may have allowed the melt to migrate via porous flow along 666 667 the direction of structural weakness (see Tommasi et al., 2017) before the opening of the conduits. The high-P emplacement conditions are consistent with the scenario where the FPP, after 668

of the Anzola-Val Grande high-T shear zone (namely, from north to south, Premosello, Balmuccia,
and Baldissero; Quick et al., 1995), until its exhumation at shallower levels at ~180 Ma (Zanetti et
al., 2013, 2016; Langone et al., 2017, 2018; Decarlis et al., 2017, Malitch et al., 2017; Petri et al.,
2019).

674

675 10.3.2. Constraints on the Mesozoic mantle sources at the Africa–Europe Boundary

The outcomes of this study indicate that in the Mesozoic, melts extremely rich in volatiles (H₂O, P, CO₂, and Cl), K, Na, and highly incompatible elements rose from mantle depths toward the surface. The large crustal component present in the melts, as indicated by the O isotopic composition, leads us to consider that the high Al content, enrichment in large-ion lithophile elements and LREEs, and enriched Nd and Sr isotopic compositions are a primary feature inherited from the source.

This finding confirms the extreme complexity of the tectono-magmatic scenario recorded by the 681 682 FPP. In particular, the northern IVZ records an extremely prolonged release of K-H₂O-rich mantlederived melts mixed with subduction-related components, which lasted from the Variscan orogenic 683 cycle to the Mesozoic exhumation of lithospheric mantle at shallower levels. This explains why the 684 FPP records many generations of phlogopite-bearing mineral assemblages, showing variable field 685 relationships, geochemical signatures, and ages (Hartmann & Wedephol, 1993; Zanetti et al., 1999; 686 2013; 2016; Stähle et al., 1990; 2001; Greco et al., 2001; 2004; Morishita et al., 2003, 2008; 687 Malitch et al., 2017). It also provides a new interpretative frame for previous data indicating the 688 emplacement of melts with subduction-related components in the Triassic (Mastumoto et al., 2005; 689 Morishita et al., 2008, Malitch et al., 2017). 690

691 This magmatism-bearing subduction component appears to have roughly overlapped with the ascent

of silica-undersaturated alkaline melts of OIB affinity (Stähle et al., 1990, 2001, Schaltegger et al.,
2015), which likely also have some counterparts in the central IVZ (Fiorentini et al., 2018; Galli et
al., 2019).

The FPP developed in a concomitant extensional-transtensional tectonic regime, the origin and 695 geodynamic scenario of which are still strongly debated (Cassinis et al., 2008; Zanetti et al., 2013; 696 Casetta et al., 2018a, b). Further investigations are needed to address whether the crustal 697 components are remnants of old subduction events (Bonadiman et al., 1994), possibly located at 698 lithospheric levels, reactivated by asthenospheric magmatism with OIB or depleted mantle affinity; 699 represent crustal material recycled into asthenospheric mantle sources (Locmelis et al., 2016); or 700 were added by active Mesozoic subduction (Cassini et al., 2008; Schmid et al., 2008; Morishita et 701 702 al., 2008; Selverstone and Sharp, 2011; Zanetti et al., 2013).

It is a matter of fact that the major elements (in particular the enrichment in K and Al), trace elements, and Sr and Nd isotopic compositions of the sapphirine-bearing gabbroic rocks approach the geochemical affinity shown by Triassic K-rich calc-alkaline to shoshonitic magmatism of the Dolomites area (Eastern Alps) (Casetta et al., 2018a, b). Thus, the study of magmatic events at the roots of the continental crust of the Adria Plate can provide a unique opportunity to constrain the tectono-magmatic evolution of the Europe–Africa boundary.

709

710 **11. Concluding remarks**

A detailed survey of Mesozoic sapphirine-bearing gabbroic dykes within the FPP unit led us to recognise different stages of melt migration (from porous-flow migration in peridotite channels to flow in open conduits) and constrain the presence in the parental melts of a large continental crustal component acquired in the source region or at deeper lithospheric mantle levels.

The reaction between dyke melts and the strongly metasomatised FPP enhanced the crustalsignature.

The high Al content of these melts allowed the segregation of magmatic sapphirine, which is amarker of high-P-T intrusive conditions.

This dyke swarm possibly represents a record of the input to the mantle of the K-rich calk-alkaline to shoshonitic melts that characterised the magmatism widespread during the Triassic in the Southern Alps area.

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- Figure captions

1022 Figure 1: Geological map of the Finero area, modified after Mazzucchelli et al. (2014).

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Figure 2: A) Sample FI09C06 crosscutting the host harzburgite foliation. The centre of the dyke is formed by the Leucocratic Zone, whereas the melanocratic zones (i.e. the Orthopyroxene Zone, Early Amphibole Zone, and Late Amphibole Zone) are indistinguishable; B) phlogopite vein cutting the Orthopyroxene Zone; C–F) occurrences of sapphirine in sample FI09C06 (C and D) and sample FI9664 (E and F). The figures show the increase of sapphirine size, from μ m (C) to mm (D and E) and cm scale (F).

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Figure 3: Primitive mantle (PM)-normalised bulk rock trace elemental patterns of a gabbroic dyke and host harzburgite. PM values are from McDonough and Sun (1995). Literature values from the harzburgite-pyroxenite association of (1) Hartmann and Wedephol (1993) and nepheline-bearing hornblende syenitic dykes of (2) Stähle et al. (2001) are reported for comparison.

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Figure 4: Orthopyroxene major elemental contents from host harzburgites, dykes, and literature
data. Plotted literature data are the harzburgite-pyroxenite association orthopyroxene from (1)
Zanetti et al. (1999) and orthopyroxene from sapphirine-bearing rock from the Mafic Complex from
(2) Sills et al. (1983).

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Figure 5: Phlogopite major elemental contents from host harzburgites, dykes, and literature data.
Plotted literature data are: harzburgite-pyroxenite association phlogopite from (1) Zanetti et al.
(1999).

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Figure 6: Amphibole major elemental contents from host harzburgites, dykes, and literature data.
Plotted literature data are: harzburgite-pyroxenite association amphibole compositions from (1)
Zanetti et al. (1999) and (2) Morishita et al. (2008), amphibole from apatite-rich veins in the FPP

from (1) Zanetti et al. (1999) and (2) Morishita et al. (2008), and amphibole from sapphirinebearing rock from the Mafic Complex from (3) Sills et al. (1983).

1050

Figure 7: Sapphirine compositions plotted in the (MgO + FeO)–(Cr₂O₃ + Fe₂O₃ + Al₂O₃)–SiO₂
diagram (mol. %). Literature data are from sapphirine of the Finero Mafic Complex (Sills et al.,
1983) and sapphirine 1 of Higgins et al. (1979).

1054

Figure 8: REE patterns of clinopyroxene and amphibole from the gabbroic dykes and the respective host rocks divided for samples and position. Values are normalised to Chrondrite-I (CI, values from Lyubetskaya and Korenaga (2007); δ^{18} O values (normalised to SMOW) from phases of sample FI09C06 are reported near the REE patterns according to their positions in the sample.

1059

Figure 9: PM-normalised trace elemental patterns of clinopyroxene from host harzburgite. PM values are from McDonough and Sun (1995). Plotted literature data are: clinopyroxene and amphibole average compositions from the harzburgite-pyroxenite association from (1) Zanetti et al. (1999) and (2) Morishita et al. (2008), clinopyroxene and amphibole average from apatite-rich veins in the FPP from (1) Zanetti et al. (1999) and (2) Morishita et al. (2008), reported as apatite veins.

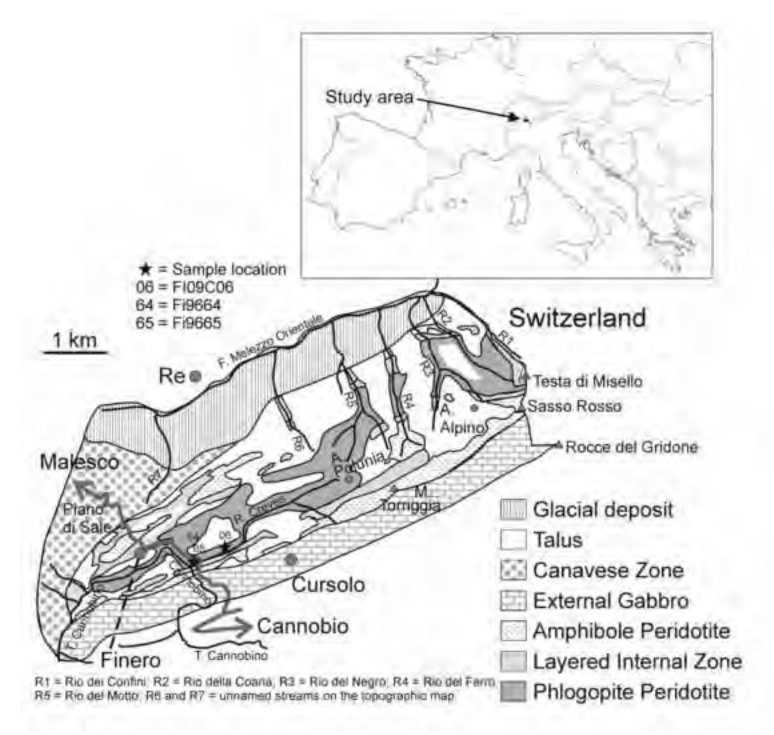
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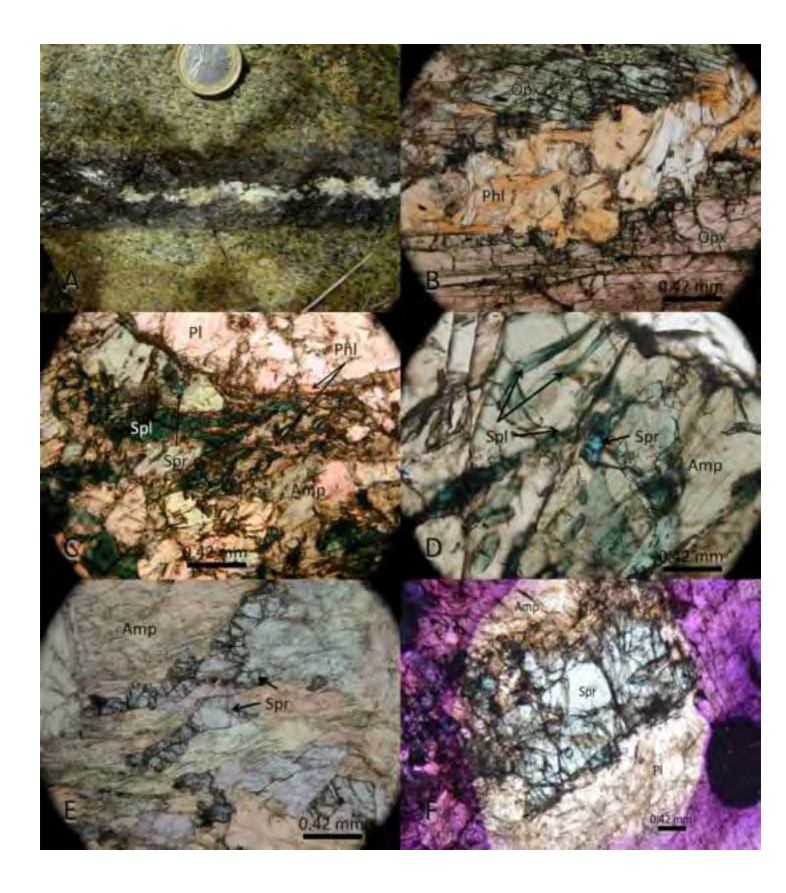
Figure 10: PM-normalised trace elemental patterns of amphiboles from gabbroic dyke and hostharzburgite. PM values are from McDonough and Sun (1995).

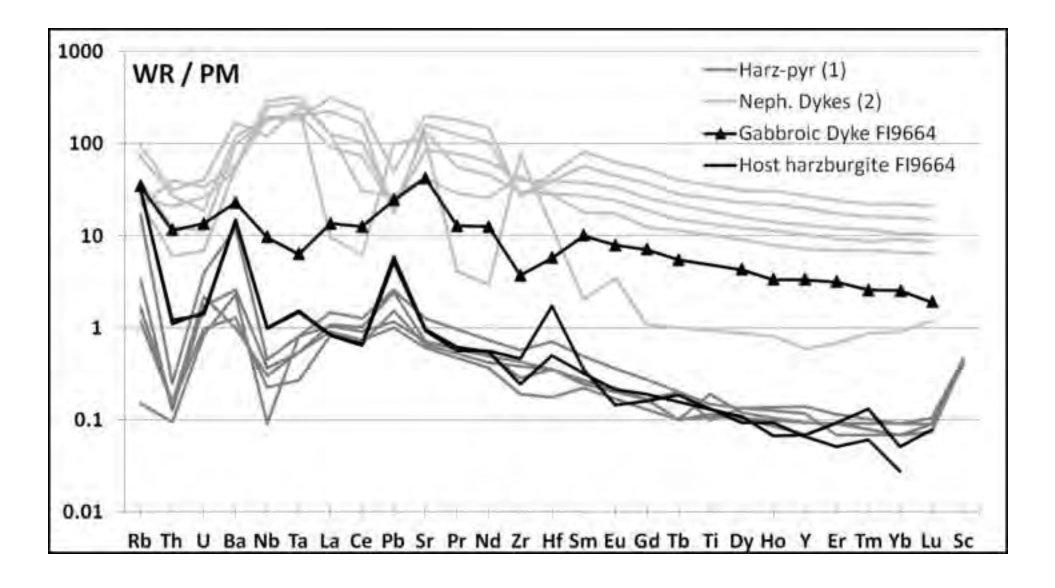
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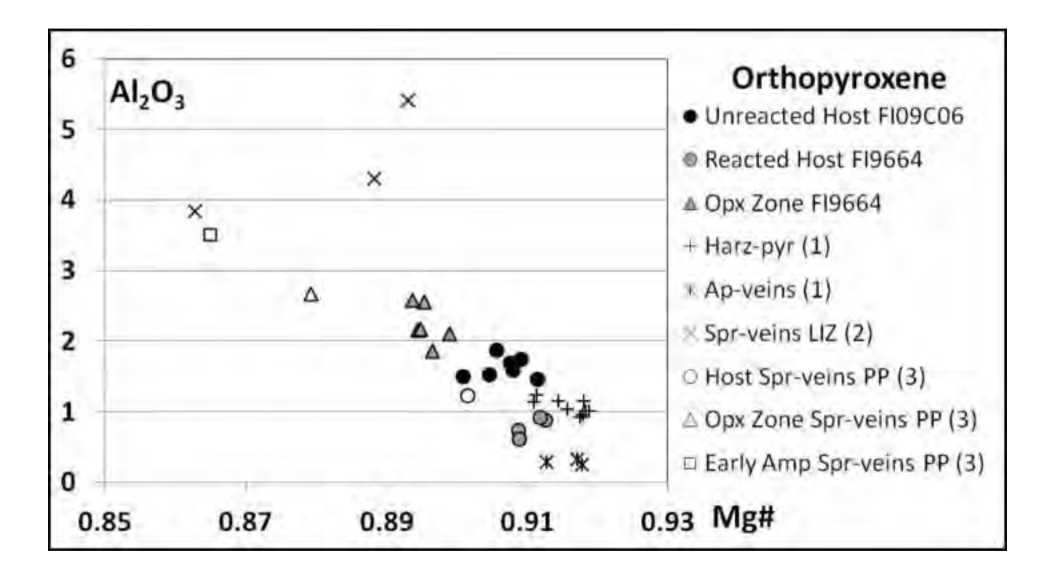
Figure 11: δ^{18} O normalised to SMOW. Values from literature are the field of mantle and variation of MORB from Rollinson (1993), and mantle ultramafics and mantle-derived melts (*) from Bindeman (2008). Finero literature data are reported from the harzburgite-pyroxenite association of Hartmann and Wedephol (1993) and Selverstone and Sharp (2011) and from chromitite in dunite bodies from Zanetti et al. (2016). 1074

Figure 12: ¹⁴³Nd/¹⁴⁴Nd vs ⁸⁷Sr/⁸⁶Sr recalculated at 225 Ma for amphibole and plagioclase from the 1075 various zones of sample FI09C06 (host and dyke): (1) amphibole data of the FPP from Obermiller 1076 (1994); (2) bulk rock data of the Finero Mafic Complex from Lu et al. (1997b); (3) data from an 1077 alkaline dyke in the FPP from Stähle et al. (1990); (4) data from an alkaline dyke in the FPP from 1078 Stähle et al. (2001); (5) amphibole and apatite data from discordant apatite-bearing veins in the FPP 1079 from Morishita et al. (2008); (6) Mesozoic shoshonitic magmatism (SS: silica saturated; US: 1080 1081 undersaturated in silica) in the Predazzo area from Casetta et al. (2018a, b); Depleted MORB Mantel (DMM) from Workman and Hart (2005). The mixing model was calculated between the two 1082 end-members: the alkaline dyke S9 of Stähle et al. (1990) and the hosting peridotite FI09C06. Data 1083 for the melt in equilibrium with the alkaline dyke S9 are: ${}^{143}Nd/{}^{144}Nd = 0.512607$, ${}^{87}Sr/{}^{86}Sr =$ 1084 0.703720 (from Stähle et al., 1990; recalculated at 225 Ma), Nd = 8.333 ppm, and Sr = 830 ppm 1085 1086 (calculated from LA-ICP-MS plagioclase analysis of albitite dykes similar to the dyke of Stähle et al., 1990, using the K_d of Dohmen and Blundy, 2014; average Nd = 0.25 ppm, Sr = 5810 ppm). 1087 Data for the melt in equilibrium with the host FPP peridotite are ${}^{143}Nd/{}^{144}Nd = 0.512130$, ${}^{87}Sr/{}^{86}Sr =$ 1088 0.708501 (recalculated at 225 Ma), Nd = 67.5 ppm, and Sr = 909 ppm (calculated using the K_d of 1089 Ionov et al., 2002). 1090











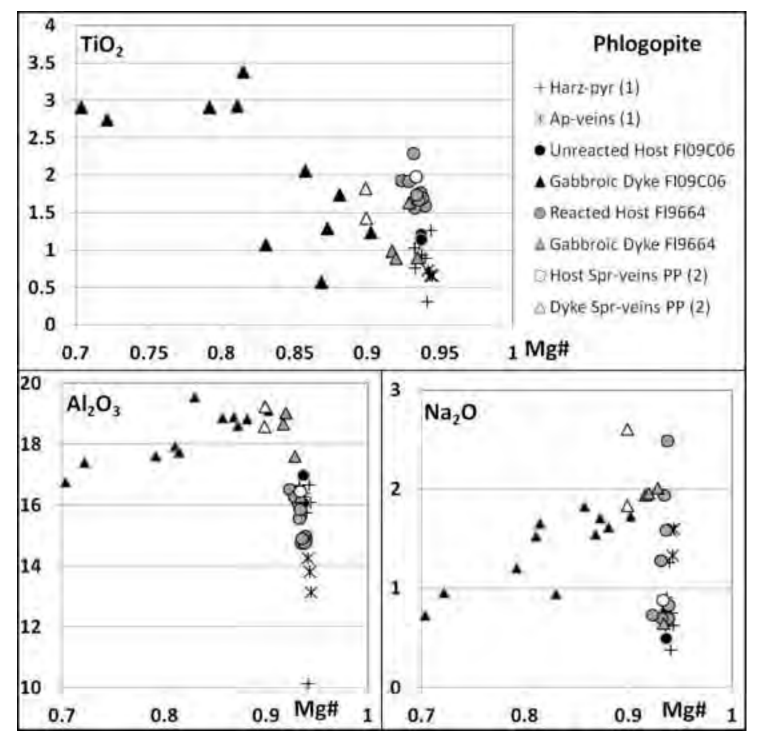
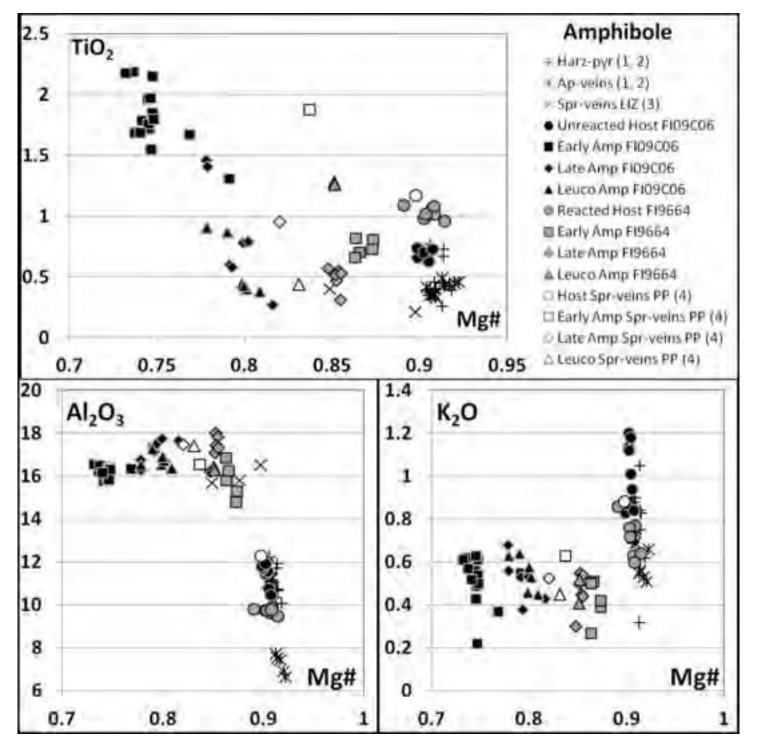
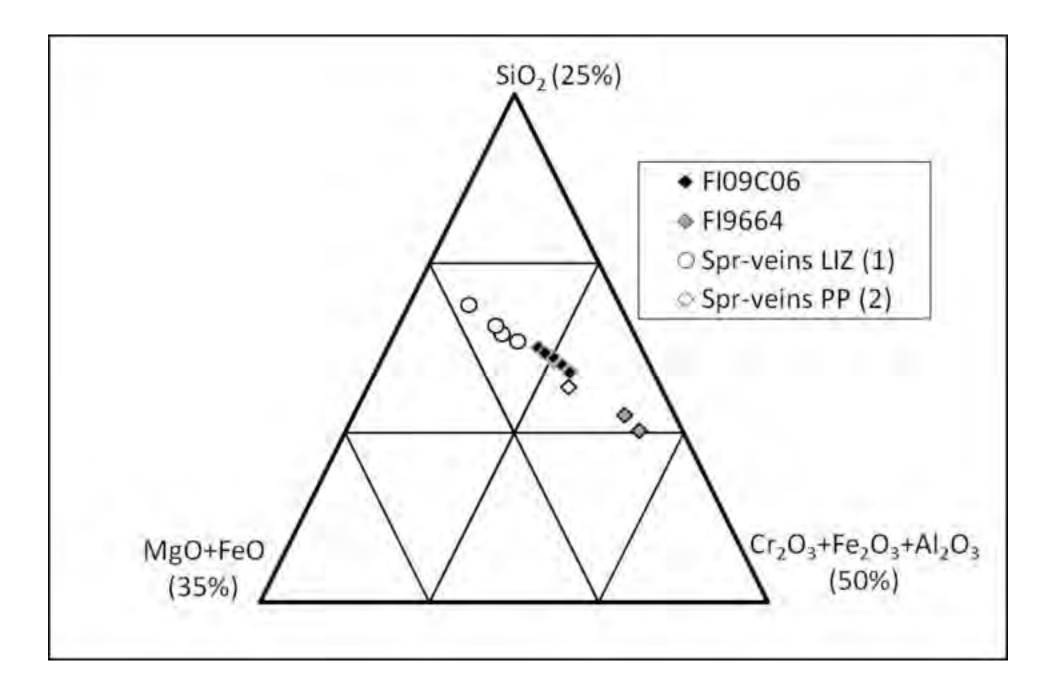
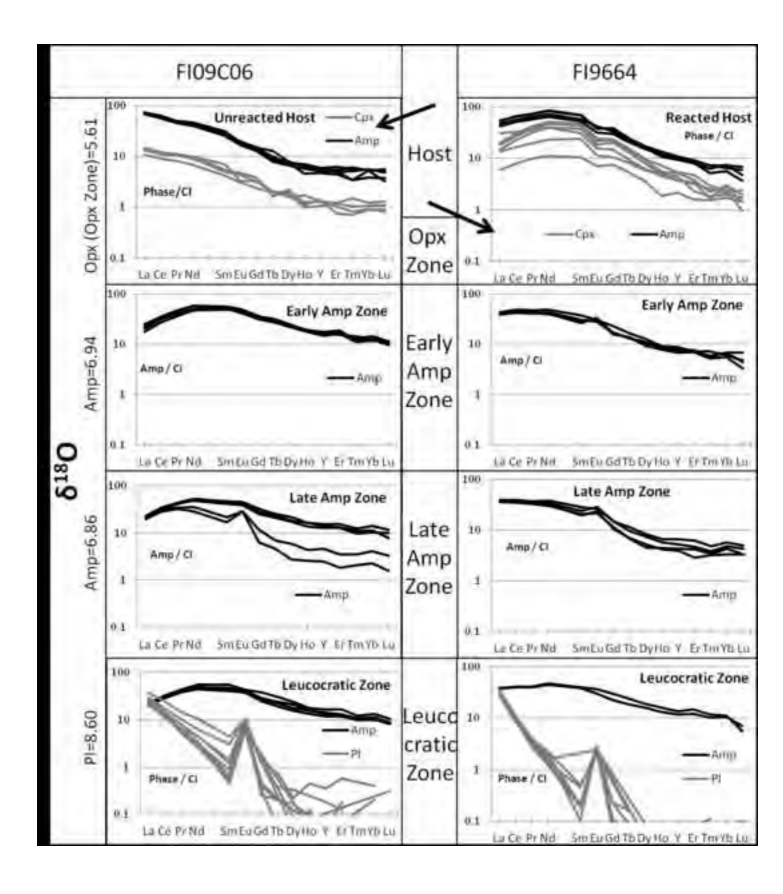


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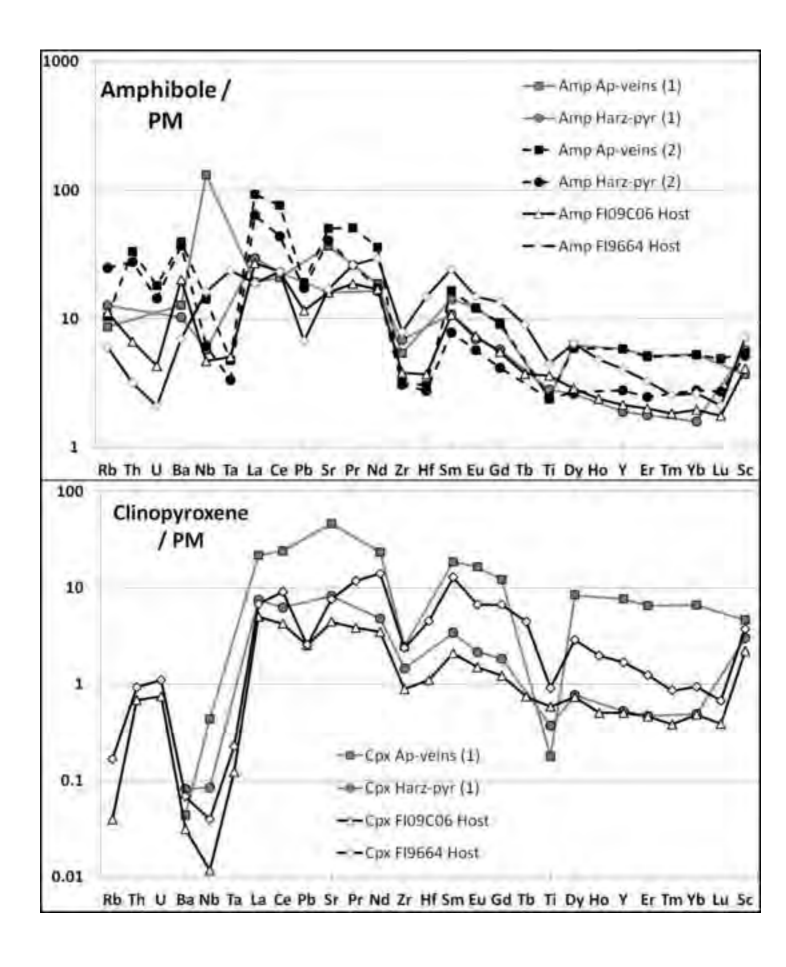
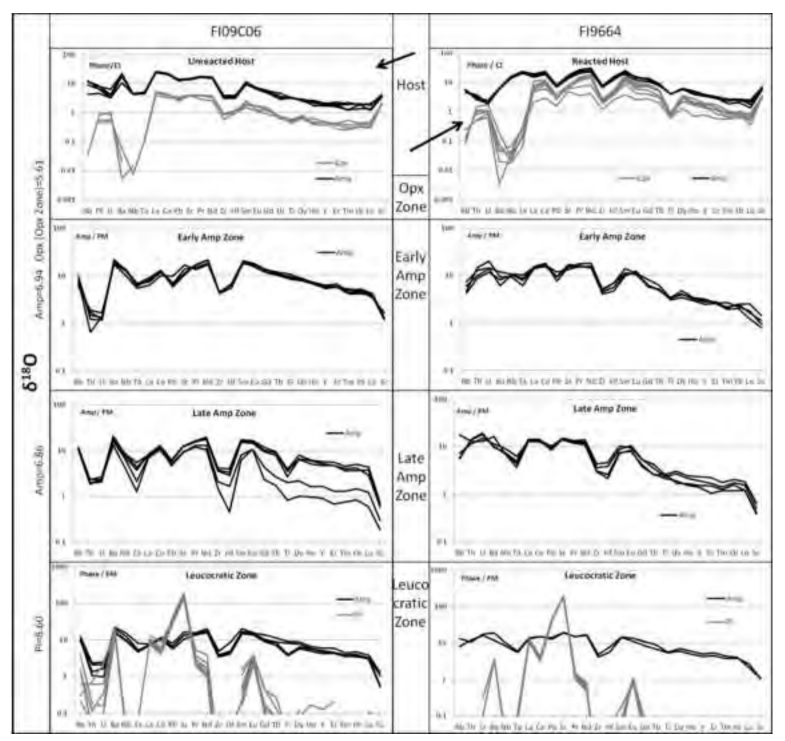
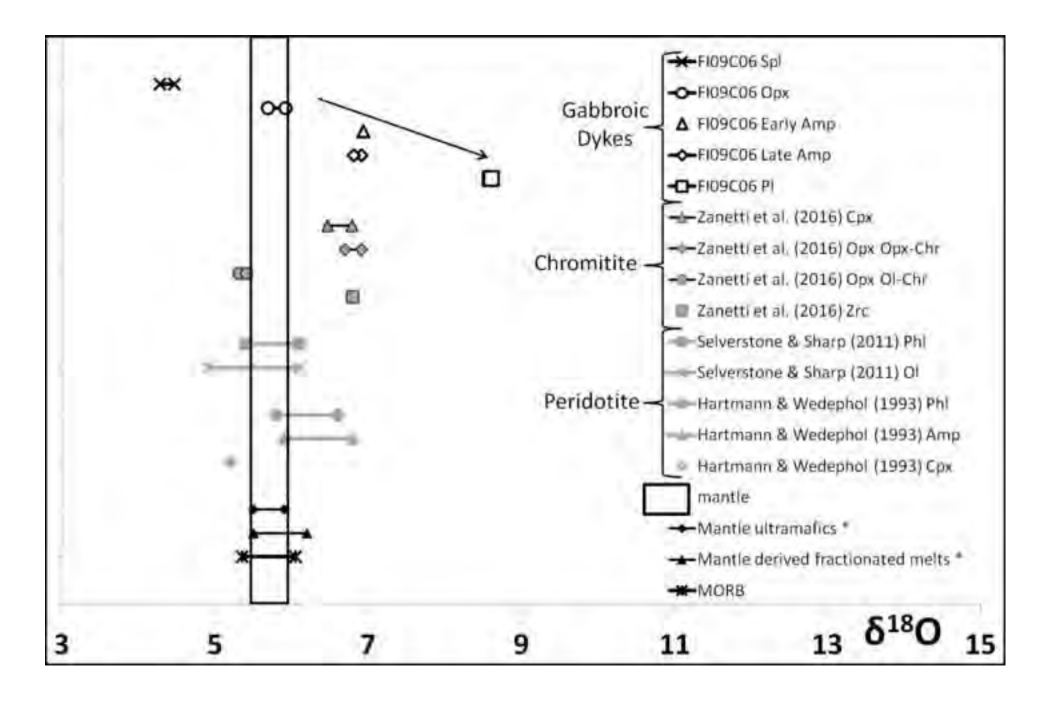


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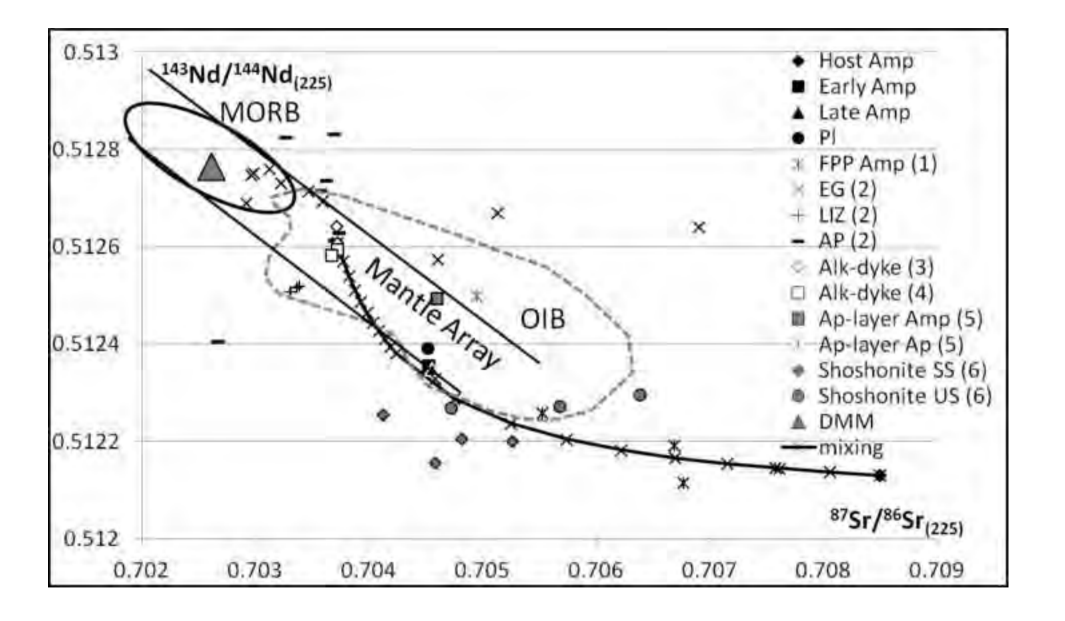


Table 1: O isotopic composition of minerals from the Spr-bearing gabbroic dykes.

| Sample | Phase | δ ¹⁸ Ο | std. dev. | | |
|---------|-----------|-------------------|-----------|--|--|
| | | | | | |
| FI09C06 | Early Amp | 6.94 | 0.00 | | |
| FI09C06 | Late Amp | 6.86 | 0.05 | | |
| FI09C06 | P1 | 8.60 | 0.01 | | |
| FI09C06 | Opx | 5.81 | 0.11 | | |

| Rock | Phase | Rb | Sr | ⁸⁷ Sr/ ⁸⁶ Sr | 2SE | ⁸⁷ Rb/ ⁸⁶ Sr | Nd | Sm | ¹⁴³ Nd/ ¹⁴⁴ Nd | 2SE | 147 Sm/ 144 Nd |
|------|-----------|-----|--------|------------------------------------|----------|------------------------------------|------|-----|--------------------------------------|----------|--------------------------|
| Host | Amp | 6.5 | 281.2 | 0.708713 | 0.000008 | 0.066372 | 19.7 | 4.2 | 0.512317 | 0.000008 | 0.126796 |
| Dyke | Early Amp | 6.8 | 324.4 | 0.704722 | 0.000008 | 0.060834 | 24.8 | 8.1 | 0.512646 | 0.000008 | 0.197559 |
| Dyke | Late Amp | 8.6 | 189.5 | 0.704971 | 0.000008 | 0.131167 | 16.8 | 4.0 | 0.512559 | 0.000006 | 0.142593 |
| Dyke | Pl | 0.6 | 3109.7 | 0.704519 | 0.000008 | 0.000568 | 2.2 | 0.2 | 0.512482 | 0.000006 | 0.061238 |

Table 2: Sr and Nd isotopic composition of minerals from sample FI09C06 from the Spr-bearing gabbroic dykes and the host peridotite.