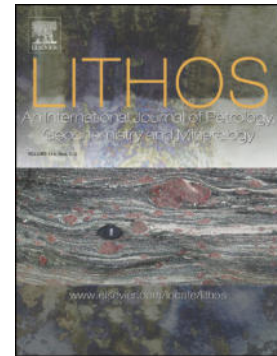


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Native copper formation associated with serpentinization in the Cheshmeh-Bid ophiolite massif (Southern Iran)

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Abstract

In the Cheshmeh-Bid district of the Khajeh-Jamali ophiolitic massifs (Southern Iran), mantle peridotites are intruded by abundant pyroxenite dykes. A few of these dykes are remarkable for the occurrence of native copper associated with the development of a metasomatic reaction zone. The dykes are progressively reacted, from their margins towards the center, with an amphibole + antigorite selvage, followed by a centimeter-thick clinopyroxene + antigorite assemblage and, finally, by the native copper-bearing zone consisting of clinopyroxene + chlorite + antigorite. Native Cu occurs along cleavages and partially healed fractures in clinopyroxene, and as massive grains intergrown with antigorite. Copper isotope signatures and thermodynamic calculations show that the main driver for reaction zone formation is Ca-metasomatism. Native copper forms at the expense of chalcocite in the reaction zone. Such a reaction can only occur in reducing conditions, in agreement with the analysis of fluid inclusions composition displaying H_2 and CH_4 . Such fluids presumably originated from the hydration of mantle rocks. The observed reaction zone and native copper mineralization are thus interpreted as the result of Ca-metasomatism during hydrothermal alteration of the oceanic lithosphere. This is consistent with U/Pb dating of titanite, suggesting formation during the Albian when the dykes were exposed on the seafloor in a supra-subduction setting. The source for copper mineralization, as revealed by Cu isotopes, is probably mantle-like.

Keywords: Native copper; serpentinization; Ca-metasomatism; pyroxenite; Cheshmeh-Bid ophiolitic massif; Iran

1. Introduction

Native copper has been documented in ultramafic and mafic rocks from ophiolitic and orogenic massifs. It was observed in partially serpentinized peridotites of Costa Rica (Schwarzenbach et al., 2014), pillow basalts from la Désirade, Lesser Antilles island arc (Nagle et al., 1973), rodingitized gabbro in serpentinites of the Braszowice-Brzeznicza

Massif, Poland (Gunia, 1986), basal cumulate sequence of the massif from the Zambales ophiolite, Philippines (Abrajano and Pasteris, 1989), plagioclase lherzolites from the Horoman peridotite complex, Hokkaido, northern Japan (Ikehata and Hirata, 2012) and websterite from the Totalp ultramafic massif, Swiss Alps (Van Acken et al., 2007). Three hypotheses have been put forward to explain the genesis of native copper in peridotites and basalts: ((i) crystallization in equilibrium with a mafic magma (e.g. Cabral and Beaudoin, 2007; Ikehata and Hirata, 2012); (ii) low-temperature in situ alteration of magmatic or hydrothermal Cu-sulfides at highly reducing conditions (e.g. Lorand and Grégoire, 2006; Schwarzenbach et al., 2014); (iii) precipitation from low-temperature hydrothermal fluids (e.g. Dekov et al., 2013; Ikehata et al., 2016).

Thermodynamic calculations show that hypothesis (ii) requires reducing conditions for native copper to precipitate in hydrothermal environments (Schwarzenbach et al., 2014). The observation of Fe-Ni alloys and native metals in serpentinized peridotites (e.g. Klein and Bach, 2009; Schwarzenbach et al., 2014) shows that highly reducing conditions prevail during serpentinization. This is consistent with vent fluid compositions measured in ultramafic-hosted hydrothermal fields (Charlou et al., 2002) and gas seep compositions analyzed in ophiolites (e.g. Abrajano et al., 1988). During serpentinization, hydrogen production relates to water reduction coupled with iron oxidation during replacement of primary olivine and pyroxene by secondary magnetite and serpentine (McCullom and Bach, 2009; Malvoisin et al., 2012). Water-rock interaction is also intimately associated with Ca-metasomatism of mafic rocks in which the removal of silica (SiO_2) and addition of Ca are the main mass transfers (Coleman 1967).

In this contribution, we report a new occurrence of native copper mineralization within Ca-metasomatically altered pyroxenites from the Cheshmeh-Bid ophiolitic massif in the Khajeh-Jamali area, Southern Iran. Based on field and petrographic observations combined with new

geochronological and geochemical data we aim to: (1) define the paragenetic sequence of the ore and silicate minerals; (2) produce a thermodynamic model for the origin of Ca-metasomatism and native copper; (3) determine the sources of Cu in the mineralization system; (4) constrain the timing of the copper mineralization and, finally, (5) develop a conceptual model of native copper genesis within metasomatised pyroxenite veins.

2. Geological setting

The Iranian Ophiolites (Fig. 1a) represent the remnants of the Tethyan oceanic lithosphere in the Anatolian segment of the Alpine–Himalayan Orogen. Mesozoic ophiolites of Iran have been divided into five ophiolitic belts (Shafaii Moghadam and Stern, 2015): (1) Late Cretaceous Zagros Outer Belt (ZOB) ophiolites along the Main Zagros Thrust including Kermanshah-Kurdistan, Maku-Khoy-Salmas, Moyrizic/Khajeh-Jamali and Haji Abad ophiolites; (2) Late Cretaceous Zagros Inner Belt (ZIB) ophiolites including Nain, Dehshir, Shahr-e-Babak and Balvard-Baft ophiolites along the main boundaries of the Central Iranian Micro-continental block; (3) Late Cretaceous–Early Paleocene Sabzevar-Torbate-Heydariieh ophiolites in the northeast of Iran; (4) Early to Late Cretaceous ophiolites including Nehbandan, Birjand and Teheh-e-Kureh ophiolites between the Afghan and Lut blocks; and (5) Late Jurassic–Cretaceous Makran ophiolites in the southeast of Iran. The Khajeh-Jamali ophiolites consist of four ophiolitic massifs belonging to ZOB in the Zagros Mountains (Southwestern Iran), and outcrop about 20 km from the Main Zagros Thrust (Fig. 1a). These massifs are dominantly composed of mantle harzburgite-dunite tectonites and a Moho Transition Zone (MTZ) (Fig. 1b; e.g. Rajabzadeh, 1998; Rajabzadeh and Nazari Dehkordi, 2013; Eslami et al., 2015). Several concordant or subconcordant orthopyroxenites and clinopyroxenites intrude both mantle tectonites and the MTZ. Boundaries between the pyroxenite intrusions and peridotite host rocks are generally sharp. The pyroxenite dykes/veins are usually coarse-grained and range from a few cm to 35 cm in thickness.

A typical cumulate sequence and a sheeted dyke complex are missing in the Khajeh-Jamali area. Peridotites are overlain by a very thin nappe of basaltic pillow lavas, reddish-purple radiolarian cherts and Late Cretaceous pelagic limestones (Rajabzadeh, 1998; Eslami et al., 2015). The ophiolitic assemblage is thrust onto the Cenomanian/Turonian shallow water deposits of the Sarvak Formation in the western portion of the Khajeh-Jamali area (Alavi, 1994). The Cheshmeh-Bid ophiolitic massif covers rugged mountainous area of $\sim 25 \text{ km}^2$ in the northwest of the Khajeh-Jamali area. This massif hosts large economical concentrations of chromitites showing massive and high-grade disseminated texture (Rajabzadeh, 1998; Eslami et al., 2015). In the Cheshmeh-Bid chromitite mine, a few pyroxenite dykes and veins crosscutting massive chromitite and serpentinized dunite are partially (Fig. 2a, b) or completely (Fig. 2c) metasomatised with remarkable native copper mineralization.

3. Analytical methods

Analytical work has been focused on five representative samples of metasomatically altered pyroxenite veins (KJPX04, KJPX05, KJPX06, KJPX11 and KJPX12) and two fresh pyroxenites (KJPX01 and KJPX20). Polished thin sections of both fresh and altered pyroxenites were examined under transmitted light using a Leica optical microscope at University of Milan. Samples KJPX04, KJPX05 and KJPX06 were selected for petrography observation because they show typical metasomatic zoning on the micrometer scale.

3.1 Major and trace element analyses in minerals and whole-rocks

Quantitative chemical analyses of silicate minerals (except chlorite) were obtained using a JEOL JXA-8230 electron probe micro-analyzer (EPMA) equipped with five wavelength-dispersive spectrometers (WDS) at the Institut des Sciences de la Terre, Université Grenoble Alpes, France. Analytical conditions were: 15 kV acceleration voltage, 12 nA beam current, $3 \mu\text{m}$ beam size. Natural minerals, pure metal and synthetic oxides were used as standards and ZAF correction was applied. Spectral interference ($\text{V K}\alpha$ vs $\text{Ti K}\beta$) was corrected using the

software-calculated correction factor. Elemental distribution maps were done at an accelerating voltage of 15kV, a beam current of 100 nA and a dwell time of 200 ms. Major element composition of chlorites, sulfides and spinels were analyzed using a JEOL JXA 8200 Superprobe equipped with five wavelength-dispersive (WDS) spectrometers, an energy dispersive (EDS) spectrometer, and a cathodoluminescence detector (accelerating potential 15 kV, beam current 15 nA), at the Dipartimento di Scienze della Terra “Ardito Desio”, University of Milano (Italy).

Major elements and Cr and Ni in whole-rock samples were analysed by X-ray Fluorescence (XRF) with a BRUKER S4-Pioneer instrument at the Instituto Andaluz de Ciencias de la Tierra (IACT, Granada, Spain), using standard sample preparation and analytical procedures. The accuracy of analyses was assessed by repeated analyses of international reference material JP-1 (peridotite) handled as unknown, which show good agreement with accepted values for this standard (Govindaraju, 1997). Whole-rock trace elements (REE, Ba, Th, Pb, Nb, Ta, Sr, Y and Sc) were analyzed by an Agilent 8800 QQQ ICP–MS (Inductively Coupled Plasma–Mass Spectrometer) at the IACT. Sample digestion was performed following the HF/HClO₄ dissolution procedure described in detail by Ionov et al. (1992), and element concentrations were determined by external calibration using aqueous solutions. Accuracy of the ICP–MS analyses has been assessed analyzing the BIR-1 basalt standard as an unknown, which show good agreement with reference concentrations of Jochum et al. (2016).

3.2 Copper isotope analysis

Grains of native copper were separated through crushing and hand picking. Samples were processed similar to the techniques in previous studies, namely Bornhorst and Mathur (2017), Mathur et al. (2009a) and Wall et al. (2011). The native copper samples were dissolved in Teflon beakers with 8 mL of ultrapure aquaregia at 140°C for 24 hours or until complete dissolution occurred. Samples were diluted to 100 ng/g for Cu isotope analysis with a

Neptune MC-ICP-MS at Pennsylvania State University. Copper isotope compositions are reported in the traditional per mil format compared to the NIST 976 international standard. Mass bias was corrected for using standard-sample-standard bracketing. All samples matched the intensities of the standards within 25 %. An internal copper metal standard (USA 1838 Cent, reported in Mathur et al. 2009b) was measured 4 times during the session and produced a $\delta^{65}\text{Cu} = 0.04 \pm 0.05$ per mil value. This value is identical to accepted $\delta^{65}\text{Cu}$ of the standard (-0.02 per mil ± 0.08).

3.3 U-Pb LA-ICP-MS analysis of titanite

Titanite grains were ablated in thin section using a 213 nm laser beam with diameter of 25 μm at 5 Hz and 40 % power (NWA ablation system). ICP-MS analyses were carried out using a Plasmaquad instrument. Data were collected on ^{206}Pb , ^{207}Pb , and ^{238}U . Immediately prior to each analysis, the spot was briefly pre-ablated over a larger area than the beam diameter to clean the surface. Following a 10 s period of baseline accumulation the laser sampling beam was turned on and data were collected for 25 s followed by a 50 s washout period. About 150 measurement cycles per sample were produced and ablation pits are about 15 μm deep. Data were edited and reduced using custom VBA software (UTILLAZ program) written by the author. $^{206}\text{Pb}/^{238}\text{U}$ show only slight fractionation caused by hole depth through the run and most of the $^{207}\text{Pb}/^{206}\text{Pb}$ and $^{206}\text{Pb}/^{238}\text{U}$ data can be averaged. Pb-204 was not measured since this peak is relatively small and would require a large amount of measurement time as well as being subject to interference from ^{204}Hg in the Ar. Ages are determined by projecting data along a common Pb mixing line to the concordia curve (see Results below). Titanite from a quartz diorite sample in northwest Ontario, DD81-29, previously dated at 2700 ± 2 Ma (Davis and Edwards, 1986) was used as a standard.

3.4 Raman Spectroscopy

Raman spectroscopic analyses were carried out at the Research and Industrial Relations Center of the Faculty of Science at the Eötvös University, Budapest. Analyses were carried out with a Horiba LabRAM HR800 spectrometer, using a 532 nm (green) laser with a maximum laser energy of 130 mW (~25 mW on the sample surface). A 100 μm confocal pinhole and an optical grating with 1800 grooves/mm were used. The spectral resolution of the measurements was 0.8 cm^{-1} at 1707.9 cm^{-1} (defined as the measured full width at half maximum values of neon atomic emission lines). During the measurements an objective with 100 \times magnification was used (NA=0.9). Analyses ran with 2 to 5 repetitions for an accumulated 200-500 seconds.

3.5 Thermodynamic modelling

We used thermodynamic modelling to determine (1) the conditions prevailing during native copper formation, and (2) the factors controlling metasomatism. The calculations were performed differently for these two objectives. We therefore constructed phase diagrams in the Fe-Ni-Cu-O-S system with SUPCRT92 (Johnson et al., 1992) for (1). We used the thermodynamic database provided in Klein and Bach (2009) in which the SUPCRT92 database is extended by including thermodynamic data for Fe-Ni sulfides. The diagrams are computed as a function of the activities of $\text{H}_{2,\text{aq}}$ ($a_{\text{H}_{2,\text{aq}}}$) and $\text{H}_{2\text{S},\text{aq}}$ ($a_{\text{H}_{2\text{S},\text{aq}}}$) at $a_{\text{H}_2\text{O}} = 1$ and 50 MPa. This latter pressure is used to compute equilibrium constants in Klein and Bach (2009), and is relevant for serpentinization on the seafloor.

We used Gibbs energy minimization with Perple_X (Connolly, 2005) for (2) because it provides solid solutions for the phases observed here. This ensures accurate modelling of the distribution of major elements during metasomatism. The equilibrium assemblages were determined for the composition of the orthopyroxenite (sample IRKJPX20; Supplementary Table 10) as a function of the chemical potential of CaO and temperature at 50 MPa. We did not include components with a concentration below 0.5 wt.% in the calculation. The chemical

potential of CaO was also computed as a function of temperature for three metasomatised samples (IRKJPX04, IRKJPX11 and IRKJPX12). We used the thermodynamic database of Holland and Powell (1998a) and the solid solution models of Holland and Powell (1996) for orthopyroxene and clinopyroxene, Holland and Powell (1998a) for olivine, spinel and staurolite, Holland and Powell (1998b) for chlorite, Diener et al. (2007) for amphibole, Padrón-Navarta et al. (2013) for antigorite and White et al. (2000) for garnet. The solid solutions for talc and brucite were considered as ideal. H₂O was added in excess and modelled with the CORK equation of state (Holland and Powell, 1998). This explains why the orthopyroxenite is not composed of orthopyroxene in the calculations but rather of its hydration products (talc and antigorite). We did not consider here the kinetic effects that led to orthopyroxene preservation.

Copper, and iron-nickel sulfides are not all available in Perple_X. We therefore constructed phase diagrams in the Fe-Ni-Cu-O-S system with SUPCRT92 (Johnson et al., 1992) for (2). We used the thermodynamic database provided in Klein and Bach (2009) at 50 MPa in which the SUPCRT92 database is extended by including thermodynamic data for Fe-Ni sulfides. The diagrams are computed as a function of the activities in H_{2, aq} ($a_{H_2, aq}$) and H_{2S, aq} ($a_{H_2S, aq}$) at $a_{H_2O} = 1$ and at the pressure at which the thermodynamic data are available (50 MPa).

4 Results

4.1 Petrography of the host peridotites

Petrography of the Cheshmeh-Bid host peridotites and chromitites has been discussed in detail by previous authors (e.g. Rajabzadeh, 1998; Eslami, 2015). The scarcity of bastite suggests that the host peridotites of the studied pyroxenites was originally a dunite. The host dunites consist of >95 % mesh-textured serpentine, cut by serpentine veins, plus 1–3 % spinel minerals. The Cheshmeh-Bid harzburgites are composed of 70-85 modal% olivine and 10-20 modal% orthopyroxene. The minor phases are fine-grained anhedral clinopyroxene (3–

5 modal%) and subhedral spinel (0.5–3 modal%). Pervasive alteration in harzburgite includes the formation of serpentine minerals after olivine and orthopyroxene. In the Cheshmeh-Bid peridotites, Fe, Cu and Ni commonly form fine-grained opaque minerals including oxides (magnetite), sulfides (pentlandite, heazlwoodite) and alloys (awaruite) that occur together with (accessory) platinum-group minerals (PGM). Base metal sulfide inclusions inside alloys show desulfurization effects in the matrix of the Cheshmeh-Bid chromitite and host dunite and harzburgite (Eslami, 2015).

4.2 Petrography and mineral chemistry of pyroxenite and reaction zones in contact with host peridotites

Along the interface with the peridotite host rock, orthopyroxenite veins developed three mineralogically and chemically distinct alteration zones. These are described below. The transition between least altered and altered zones can either be sharp or progressive.

4.2.1. Orthopyroxenite. The least altered mineral assemblage of the pyroxenite vein comprises orthopyroxene and chromian spinel. It occurs in a zone with an average thickness of 1.5 cm (hereafter Opx-rich zone, Fig. 3a). Clinopyroxene, olivine and antigorite are secondary minerals in this zone. This zone displays a porphyroclastic texture with 1 to 3 mm large pyroxene grains surrounded by antigorite. The composition of orthopyroxene in this zone is mostly enstatitic (supplementary Fig.1), with Mg# [$\text{Mg}/(\text{Mg}+\text{Fe}_{\text{tot}})$] of 0.90–0.93. Orthopyroxene has Cr_2O_3 , Al_2O_3 and TiO_2 contents of 0.12–0.46 wt.%, 0.40–1.38 wt.% and <0.05 wt.%, respectively (Supplementary Table S1). The orthopyroxene crystals contain numerous ovoid and lamellar exolutions of clinopyroxene (Fig. 3b). Clinopyroxene exolutions in the Opx-rich zone have high CaO contents ranging between 21.02– 23.62 wt.% (with average of 22.94 wt.%), low Al_2O_3 contents of 0.50–2.20 wt.% (with average of 1.33 wt.%) and Cr_2O_3 contents of 0.26 to 0.94 wt.% with average of 0.63 wt.% (Supplementary Table S2). The Mg# of clinopyroxene exolutions, observed in large orthopyroxenes, range

from 0.91 to 0.99. Clinopyroxene from pyroxenites are mostly diopside (Supplementary Fig. 1).

In Opx-rich zone of Cu-rich pyroxenites, secondary olivine occurred as vein-like texture in the middle of antigorite around the orthopyroxene porphyroclasts (Fig. 3c). Representative analyses of olivine with vein-like texture from the Cu-bearing pyroxenites are given in Supplementary Table S3. Olivine has low forsterite contents comprised between 75 and 83 mol.%. Compared to the vein-like textured olivine in Cu-bearing pyroxenites, primary olivines found in dunite and harzburgite of the studied area have considerably higher Fo content (91-95 mol %) (Fig. 4; Rajabzadeh and Nazari-Fenizordi, 2013). MnO contents for vein-like textured olivines vary from 0.34 to 0.82 wt % whereas they are lower than 0.21 wt.% for olivine in associated harzburgite and dunite (Fig. 4a). NiO contents for vein-like textured olivines are homogeneous in all samples and lower (0.08 to 0.16 wt.%) compared to those in associated dunite and harzburgite (0.27-0.66 wt.%) (Fig. 4b). MnO contents and Mg# are negatively correlated for vein-like textured olivines whereas there is a broad positive correlation between NiO contents and Mg# for olivines from associated harzburgite and dunites.

Euhedral or subhedral chromian spinel (50 μm to 2.5 mm in diameter) is interstitial or included in orthopyroxene grains (Fig. 3d). Chromian spinel grains show alteration to ferrian chromite across their cracks and fractures. Rarely, spongy reaction rims developed around homogeneous chromian spinel grains. Sieve-textured rims of chromian spinel results from the occurrence of several euhedral to anhedral inclusions of chlorite and clinopyroxenes (Fig. 3e). Inclusion sizes range from 5 to 100 μm . Fresh cores of spinel show Cr# $[\text{Cr}/(\text{Cr} + \text{Al})]$ ranging from 0.65 to 0.73 (average of 0.70) and Mg# $[\text{Mg}/(\text{Mg} + \text{Fe}^{2+})]$ from 0.49 to 0.59 (average of 0.55) (Supplementary Table S4; Fig. 5). These high-Cr# spinels are characterized

by low TiO₂ content < 0.18 wt.%, MnO contents of 0.04-0.24 wt.%, FeO contents of 18.29-25.21 wt.%, Al₂O₃ contents of 13.38–20.29 wt.% (Fig. 5).

The products of three successive reactions overprinted the Opx-rich zone. We distinguished three reaction zones from inner to outer portions of the orthopyroxenite dykes:

4.2.2. *Zone I* is characterized by a narrow selvage of amphibole with subordinate antigorite between Opx-rich zone and Zone II. The thickness of this zone is 50 µm to 100 µm (Fig. 3a). The chemical composition of amphibole from the Cheshmeh-Bid pyroxenites is given in Supplementary Table S5. Based on the nomenclature of Leake et al. (1997), amphiboles show a wide range of compositions from tremolite to magnesian-hornblende and edenite. They are characterized by Mg# between 0.92 and 0.98, TiO₂ < 0.7 wt.%, Cr₂O₃ contents are between 0.12 and 2.50 wt.% and Al₂O₃ contents between 0.26-1.30 wt.%.

4.2.3. *Zone II* is composed of clinopyroxene megacrysts with subordinate antigorite and very rare orthopyroxene. The thickness of this zone is from 1 cm to 1.5 cm (Fig. 3a). Clinopyroxenes in Zone II show a wide range of Mg# (0.75-0.98) and are characterized by CaO contents of 21.95– 26.1 wt.% with average of 23.25 wt.%, Al₂O₃ and Cr₂O₃ contents lower than 2.23 wt.% and 1.50 wt.%, respectively (Supplementary Table S2; Supplementary Fig. 1). Light grey bands of zoned clinopyroxenes in Zone II have lower Mg# values (0.76-0.80) and higher Cr₂O₃ contents compared to dark grey zones (Fig. 3f). Sporadic inclusions of amphiboles are discernible along the cleavage of clinopyroxenes (Fig. 3g). Amphibole inclusions in Zone II are magnesian-hornblende to edenite. They are characterized by Mg# between 0.90 and 0.97 and TiO₂ contents < 0.23 wt.%, Cr₂O₃ contents of 0.80-2.53 wt.% and Al₂O₃ contents of 4.14-9.71 wt.% (Supplementary Table S5). Orthopyroxene inclusions in diopside megacrysts of Zone II show lower Mg# (~ 0.87). These inclusions have average Al₂O₃ = 1.5 wt.% and Cr₂O₃ = 0.51 wt.%. Clinopyroxene is locally replaced by calcite with perfect rhombohedral cleavage (Fig. 3h, 3i).

4.2.4. *Zone III* is the mineralized zone consisting dominantly of clinopyroxene, chlorite, antigorite and native copper. Zone III range from 0.5 cm to 1.5 cm in width and is typically massive in texture (Fig. 3a). Antigorite forms well-shaped tablets or laths that penetrate deeply into the clinopyroxenes. Chlorite forms patches or more commonly occurs interstitially in between the antigorite plates (Fig. 3j). It shows chemical variations (dominantly clinochlore) in each set of samples (Supplementary Table S6). Generally, chlorites in Zone III from the studied pyroxenites display low FeO (<5.08 wt.%), high MgO (28.46-33.26 wt.%) and variable Cr₂O₃ contents (0.17-3.84 wt.%). Chlorites are mainly clinochlore with subordinate pennine (Supplementary Fig. 2). In backscattered electron images, the Fe-rich chlorites and Fe poor have the same texture (Fig. 3i). Rarely, apatite is interstitial with respect to chlorite and antigorite.

Native Cu with pure composition occurs as flakes intergrown with antigorite (Figs. 6a and 6b; Table S7), along cleavage planes and along partially healed fractures cutting clinopyroxene (Fig. 6c). Occasionally, native copper occurs in fractures of chromian spinel as well as along chromian spinel grain boundaries (Fig. 6d). Occasionally, individual titanite grains are closely associated with native copper and chlorite (Fig. 6e) in zone III. Rare Cu-Au alloy is found in Zone III (Fig. 6e). Small inclusions of chalcocite (Cu₂S) are fully embedded within native copper (Fig. 6f). Chalcocites hosted by native copper as vein and inclusion shows a relatively narrow range of Cu (76.69-80.92 wt.%) and S (20.05-21.71 wt.%) (Table S7). Occasionally, cadmium sulfides (Greenockite) occur as inclusions in native copper and have Cu (3.87-6.78 wt.%) and Cd contents (76.5-78.1 wt.%) (Table S7).

4.3. Chemical changes across the reaction zone

X-ray element mapping shows that the contact between Opx-rich and metasomatic zones (I and II) of Cu-rich pyroxenite samples is sharp (Fig. 7). The formation of amphiboles in Zone

I is associated with increase in Ca, Na and Al. Sporadic high concentrations of these elements also occur in Zone II where they correspond to amphibole inclusions. Aluminium transfer may have also a significant role in the formation of chlorite in zone III. The high volumetric proportion of clinopyroxene in zone II is responsible for an increase in Ca.

4.4 Fluid inclusions in Zone II

A study of fluid inclusions (FIs) in clinopyroxene from zone II was carried out for the sample PX05 (Fig. 8a). FIs appear in cloudy clinopyroxenes either oriented randomly or along cleavage planes and healed cracks. The FIs mainly range in size from 3 to 10 μm and display various shapes (Fig. 8a). The inclusions are often partially or fully decrepitated and whiskers may occur (Fig. 8a). At room temperature two phases (liquid + vapor) can be observed (Fig. 8a). The FIs were discriminated based on their petrographic position following Roedder (1984) and Van den Kerkhof and Hein (2001). The FIs are considered to have a secondary origin, since they mostly occur along healed fractures and cleavage planes, indicating that they were captured after the formation of the host clinopyroxenes.

We determined the nature of the phases in the FIs using Raman microspectroscopy. The molar proportion of the fluid components (shown in Supplementary Table S8) in the fluid (vapor + liquid) at room temperature was calculated based on 1) the integrated band area of their characteristic Raman bands and 2) Raman cross-sections following the method of Dubessy et al. (1989). In all studied FIs we identified CH_4 and H_2 based on their highest intensity bands (at ~ 2918 ; ~ 588 and $\sim 4156 \text{ cm}^{-1}$, respectively) (Fig. 8b). The fraction of H_2 widely varies in the FIs from 20 to 98 vol.%.

4.5 U-Pb LA-ICP-MS dating of titanite

Laser ablation ICP-MS U-Pb analyses including U and Pb contents, isotopic ratios and ages of seven titanite grains from the Cheshmeh-Bid Cu-rich pyroxenite are given in Supplementary Table S9. All titanite grains contain some U ranging from 5 to 38 $\mu\text{g/g}$ as well

as some initial Pb. Concordia plots of titanite U–Pb isotopic results are shown in Figure 9. All titanite grains contain significant initial common Pb and therefore lie on a mixing line between the $^{207}\text{Pb}/^{206}\text{Pb}$ ratio of the common Pb component and the age of the radiogenic component as defined by the lower concordia intercept (Fig 9). Regressing data from the titanite grains using Isoplot (Ludwig, 2003) gives a line that projects to an age of 101 ± 22 Ma. The MSWD of 3.8 indicates scatter outside of measurement error, probably because of excess variations in the Pb/U ratio. To some extent the 95% confidence error takes account of this since it scales as the square root of the MSWD (Ludwig, 2003).

4.6 Copper isotope composition of native copper

Copper isotopic results of eight native copper grains from the Cheshmeh-Bid Cu-rich pyroxenite are given in Supplementary Table S10. The eight native copper grains have Cu isotope compositions ranging from $\delta^{65}\text{Cu} = -0.20\text{‰}$ to $+0.28\text{‰}$ (Fig. 10). Errors for all the analyses are $\pm 0.08\text{‰}$. An average $\delta^{65}\text{Cu}$ of Bulk Silicate Earth (BSE) value of $+0.07 \pm 0.10\text{‰}$ (2SD) (Savage et al., 2015) was used to examine and characterize the Cu isotope composition of Cheshmeh-Bid native copper. The copper isotope values for the studied native copper lie within the previously reported copper isotope range for bulk mantle rocks (Fig. 10; Ben Othman et al., 2006; Ikehata and Hirata, 2012; Liu et al., 2015; Zou et al., 2019). Native copper grains from the Cheshmeh-Bid pyroxenites display significant isotope fractionation compared to primary native copper in Horoman peridotite complex but there is significant isotope fractionation for Cheshmeh-Bid native copper.

4.7 Bulk-rock chemistry

Whole rock geochemical data of the Cheshmeh-Bid pyroxenite samples are shown in Supplementary Table S11. Compared to fresh pyroxenite samples (PX01 and PX20), matasomatized pyroxenites show slightly higher Mg# values (0.91-0.96), lower SiO_2 (46.37-51.76 wt.%), higher Al_2O_3 (2.60-8.84 wt.%), CaO (5.18-18.96 wt.%) and TiO_2 contents

(0.07-0.17 wt.%) (Supplementary Table S11). The Cheshmeh-Bid metasomatised pyroxenites have high REE concentrations (3.79-15.64 $\mu\text{g/g}$).

We use in the following an isocon diagram of Grant (2005) to estimate element mobility associated with metasomatism in the Cheshmeh-Bid pyroxenites (Fig. 11). We used average composition of the Cheshmeh-Bid fresh orthopyroxenites (samples IRKJPX01 and IRKJPX20) and metasomatic pyroxenite samples (samples IRKJPX04, IRKJPX05, IRKJPX06, IRKJPX11 and IRKJPX12). The isocon diagrams allow to quantify the loss and gain of elements during the alteration of fresh samples by using TiO_2 and Sc to define the isocon (Fig. 11). These elements are commonly considered as relatively immobile during alteration (Van Baalen, 1993; Grant, 2005; Beinlich et al., 2010). The elements plotting above the isocon were gained during metasomatism whereas those plotting below were lost. Concentration of Al_2O_3 , CaO and REE increased during the metasomatism of the Cheshmeh-Bid orthopyroxenites, whereas MgO, FeO, SiO_2 , MnO, K_2O and Ba decreased (Fig. 11). The slope of the constant composition line in the isocon diagram is 1.9. This indicates that the total mass and the total volume decreased of 47 % and 61 % during the alteration, respectively. This is consistent with the observation that most major elements were lost during metasomatism.

4.8 Thermodynamic modelling of Ca-metasomatism

Thermodynamic modelling of phase equilibria indicates that the talc + antigorite + amphibole assemblage is stable in the presence of fluid at temperatures below $\sim 420^\circ\text{C}$ and at the lowest chemical potential of CaO investigated here (μCaO ; Fig. 12). This assemblage is replaced at higher μCaO by amphibole + antigorite (I), then by clinopyroxene + antigorite (II) and finally by clinopyroxene + antigorite + chlorite (III). The samples having experienced Ca metasomatism are predicted to be composed of one of these three mineralogical assemblages (assemblage I for IRKJPX04 and assemblage III for IRKJPX11 and IRKJPX12). At the

highest μCaO investigated in the simulation, phases such as brucite, wollastonite and vesuvianite are stable. These latter phases are not observed in the natural samples. The olivine-antigorite transition occurs at a higher temperature at low μCaO (450°C at -730 kJ/mol) than at high μCaO (250°C at -675 kJ/mol).

4.9 Stability of native copper

The Cu-Fe-Ni-O-S phase relations are displayed in Figure 13 as a function of the activities in $\text{H}_{2,\text{aq}}$ and $\text{H}_{2\text{S},\text{aq}}$ at 50 MPa and 200 °C, 300 °C and 400 °C. In the Fe-Ni-O-S system, the stability fields are identical to the ones calculated in Klein and Bach (2009). Native copper is stable at $a_{\text{H}_{2,\text{aq}}} > 10^{-3}$ and $a_{\text{H}_{2\text{S},\text{aq}}} < 10^{-3}$ together with awaruite or heazlewoodite in the Cu-Fe-Ni-O-S system. Its stability field is bound by chalcocite (Cu_2S) at low $a_{\text{H}_{2\text{S},\text{aq}}}$ and bornite (Cu_5FeS_4) at higher $a_{\text{H}_{2\text{S},\text{aq}}}$. The stability field of bornite is bound by chalcopyrite (CuFeS_2) at $a_{\text{H}_{2\text{S},\text{aq}}} > 10^{-3}$ at 200°C and at $a_{\text{H}_{2\text{S},\text{aq}}} > 10^{-2}$ at 400°C. In the Cu-O-S system, the stability field of native copper extends towards higher $a_{\text{H}_{2\text{S},\text{aq}}}$ and is entirely bound by chalcocite.

5. Discussion

5.1. Ca-metasomatism: the main driver for reaction zone formation

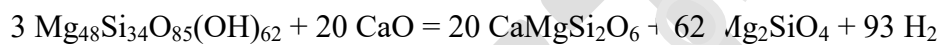
Thermodynamic calculations predict the progressive replacement of orthopyroxene by amphibole, clinopyroxene and chlorite as μCaO increases (Figure 12). This mineralogical sequence is exactly the one observed in the reaction zone where amphibole, clinopyroxene and clinopyroxene + chlorite are the main components of zones I, II and III, respectively (Fig. 3a). This suggests that calcium input is the main driver for metasomatism. Clinopyroxene crystals in zone II show a wide range of Mg# (0.76-0.98) due to almost unity Mg-Fe partition coefficient between clinopyroxene and associated antigorite and chlorites. Clinopyroxenes cogenetic with chlorites, antigorite and/or tremolite have metasomatic origin. Aluminum transfer may also play a role for the formation of zone III in which significant amounts of chlorite are observed. The chromian spinel of the orthopyroxenite appears to react during

metasomatism and may provide the Al needed to form chlorite. Chromian spinel alteration may also explain the high Cr content measured in chlorite (up to 3.41 wt.%). The presence of chlorite inclusions and ferrian chromite in porous textured chromian spinel can be attributed to metamorphism or hydrothermal events (e.g. Beinlich et al 2020). Gervilla et al. (2012) suggested that during the early stage of chromite alteration, pristine chromite may react with olivine in the presence of reducing fluids to form chlorite and secondary high Cr-chromite with porous texture. Textural evidence (Fig. 3d) suggests the possibility that chromite grains from the Cheshmeh-Bid pyroxenites also reacted with percolating Ca^{2+} - H_2 -rich fluids during metasomatism. Possible support for this scenario arises from the abundant large inclusions of chlorite and metasomatic clinopyroxene in porous chromian spinel rims.

Ca-metasomatism is thought to occur by interaction between the pristine orthopyroxenite and Ca-OH fluids derived from serpentinization of ultramafic rocks. The observation of secondary/metamorphic olivine may also strengthen the interpretation of reaction zone formation by Ca-metasomatism. The mode of occurrence and mineral chemistry of olivine ($\text{Mg}\# = 0.75\text{-}0.83$) in Opx-rich zone of Cu-rich pyroxenites is indeed in agreement with a secondary origin. It is interesting to note that the $\text{Mg}\#$ of olivine is also controlled by the phases in equilibrium (i.e. antigorite) and low- $\text{Mg}\#$ olivine is quite ordinary in low-T metaperidotites (Arai and Uyama, 1981).

Several hypotheses have been proposed to explain the dehydration of serpentine producing olivine: (i) injection of magma into meta-peridotites or -serpentinites (e.g. Vance and Dungan, 1977); (ii) Barrovian-type metamorphism (e.g. Evans, 1977). (iii) de-serpentinization in deep portion of subducting slab that has been linked to intermediate-depth intraslab earthquake (e.g. Hacker et al., 2003;; Plümper et al., 2017); (iv) in-situ dehydration of serpentinized mantle within oceanic lithosphere prior to subduction and/or obduction processes (Iyer et al., 2010). Figure 12 shows that serpentine breakdown occurs at

temperatures above 400°C at 50 MPa at low μCaO (-730 kJ/mol). This is in agreement with experimental constraints in the Mg-Si-O-H system (Ulmer and Trommsdorff, 1999; Padrón-Navarta et al., 2010). Furthermore, the addition of CaO significantly shifts the antigorite-olivine equilibrium towards lower temperature (250°C at $\mu\text{CaO} = -675$ kJ/mol; Figure 12). The evolution of the modes of the rock-forming minerals (Supplementary Fig. 2) indicates that the reaction of olivine formation at high μCaO can be written in the Mg-Si-Ca-O-H system as:



There is no evidence for high-pressure metamorphism in the studied pyroxenites. This suggests that serpentine breakdown and olivine formation are the result of Ca-metasomatism rather than heating at high-pressure in the Cheshmeh-Bid pyroxenites.

Fluids originated during serpentinization usually referred as “serpentinization-buffered fluids”, are Si-undersaturated and Ca-rich (e.g. Bach and Klein, 2009). The source of Ca is still a matter of debate and two ultimate sources have been suggested: (i) breakdown of pyroxenes during serpentinization, which may release Ca^{2+} into aqueous fluids (e.g. Coleman, 1967; Allen and Seyfried, 2003). Bach and Klein (2009) proposed that addition of Ca^{2+} from serpentinizing peridotite is likely driven by diffusive mass transfer in the form of hydroxo species (CaOH^+); (ii) external hydrothermal solutions (e.g. Hatzipanagiotou and Tsikouras, 2001). The isocon diagram (Fig. 11) confirms mobility of major elements and REE (La-Lu) during metasomatism. Significant increase of CaO and decrease of SiO_2 reflect the Ca-metasomatism reaction during alteration of the Cheshmeh-Bid orthopyroxenites. Although REEs are generally regarded as immobile elements during fluid-rock reaction, they can be mobilized during metasomatic process and hydrothermal alteration (e.g., Salvioli-Mariani et al., 2020). Composition of fluid(s), pH conditions of the fluid phase, availability of

ligands in the aqueous fluids and water/rock ratio are key factors controlling the pronounced REE mobilization during hydrothermal alteration and/or metasomatic processes. The formation of calcite and apatite during Ca-metasomatism of the Cheshmeh-Bid pyroxenites is likely related to high activities in CO_3^{2-} and PO_4^{3-} , respectively. Complexation of LREE may be triggered by carbonate ions (e.g. Gimeno-Serrano et al., 2000). High concentration of LREE with respect to the HREE can be explained by preferential complexation of carbonates with LREE. High pH and relatively basic conditions are favourable for the REE-carbonate complexation (e.g. Haas et al., 1995), which is consistent with alkaline nature of fluids during serpentinization of peridotites (e.g. Barnes and O'Neil, 1969).

The Ca-metasomatism observed in the case of Cheshmeh-Bid Cu-rich pyroxenites leads to the formation of a reaction zone. The dykes are surrounded by harzburgites and dunites having experienced extensive serpentinization, suggesting that clinopyroxene breakdown in the ultramafic rocks during hydration may play a key role for the Ca-metasomatism described here. In the following, we further investigate the links between Ca-metasomatism and serpentinization by looking at evidence for reducing conditions during fluid/rock interaction.

5. 2. Reducing conditions during metasomatism

Serpentinization leads to reducing conditions due to the coupled reduction of water to form H_2 and the oxidation of the ferrous iron initially contained in the primary minerals (mostly olivine) to form ferric minerals (e.g. magnetite) (McCullom and Bach, 2009; Malvoisin et al., 2012).

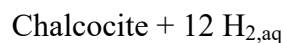
H_2 (e.g. Barnes et al., 1967) is commonly observed in fluids associated with serpentinized peridotites and Ni-Fe alloys (e.g. Ramdohr, 1950) have been described in serpentinized peridotites. Methane is also found in serpentinizing environments; it is generated through reduction of CO_2 or CO by H_2 (e.g. Charlou et al., 2002; McCullom, 2016). The observation of H_2 - CH_4 -bearing fluid inclusions in metasomatic diopside from the Cheshmeh-Bid

pyroxenites indicates that Ca-metasomatism occurred under reducing conditions, most likely associated with the serpentinization of the surrounding rocks.

The source of copper may be either desulfurization of primary Cu-sulfides of the pyroxenite during the hydrothermal overprint or transportation of Cu in a S-poor fluid from the surrounding harzburgites-dunites into the orthopyroxenite dike during hydrothermal alteration. Thermodynamic modelling indicates that native copper can be stable at high H_2 activity and/or at a low H_2S activity (Fig. 13). The fluid inclusions data reveal the presence of H_2 during metasomatism (Figs. 8a and 8b), suggesting that a high activity in H_2 prevailed during native copper formation in the Cheshmeh-Bid pyroxenites. H_2S activity during alteration is probably relatively low (Fig. 13) and thus difficult to constrain. Some mineral equilibria allow H_2S activity to be retrieved from H_2 activity. Seyfried et al. (2004) proposed that a_{H_2} and a_{H_2S} may be buffered by the equilibrium magnetite + bornite + chalcocite in serpentinized peridotites. Klein and Barr (2009) used mineralogical observations in serpentinized peridotites collected on the seafloor and measurements of H_2S and H_2 concentrations in fluids venting at peridotite-hosted hydrothermal fields (Rainbow and Logatchev, Charlou et al., 2002) to determine that the heazlewoodite/pentlandite equilibrium buffers a_{H_2S} where serpentinization occurs (HP buffer). The sulfide mineralogy in the serpentinized peridotites surrounding the orthopyroxenites described here is dominated by pentlandite and heazlewoodite suggesting that the HP buffer may fix a_{H_2} and a_{H_2S} during native Cu formation. This would imply fast H_2 and H_2S transport between the serpentinized peridotites and the orthopyroxenites. H_2 is known to diffuse rapidly in water (Kallikragas et al., 2014). Figure 13 shows that native copper is not stable at a_{H_2} and a_{H_2S} fixed by the HP buffer in the Cu-Fe-Ni-O-S system. Bornite (Cu_5FeS_4) is indeed the stable Cu-bearing mineral along the HP buffer under reducing conditions. Bornite and other iron-bearing sulfides have not been observed in the samples from the Cheshmeh-Bid Ophiolite. This may

be interpreted as evidence for i) $a_{\text{H}_2\text{S}}$ below the one fixed by the HP buffer or ii) a low iron availability preventing Fe-bearing sulfide formation in zone III during metasomatism (Fig. 13). It is difficult to select one of these two options since the mobility of H_2S and the availability of iron are poorly constrained in the studied system. The isocon diagram (Fig. 11) indicates that Fe is lost during metasomatism but the concentration in Fe is still sufficient to form bornite instead of native copper in zone III. The thermodynamic calculations performed here in the Cu-Fe-Ni-O-S system assume the presence of Fe in excess and thus do not allow for Fe distribution to be investigated. Thermodynamic calculation of iron distribution in a Cu-bearing system is theoretically possible, for example, with Gibbs energy minimization. However, it cannot be currently performed due to the lack of a thermodynamic database containing Cu-sulfides and realistic silicate solid solutions.

The scarcity of chalcocite in the studied sample and its occurrence as inclusions in native copper (Fig. 6f) suggest that native copper is formed at the expense of chalcocite during alteration. If $a_{\text{H}_2\text{S}}$ is below the HP buffer (i), native copper formation can be explained by desulfurization induced by an increase in H_2 activity associated with serpentinization in the surrounding peridotites. If a low iron availability is assumed and bornite formation is excluded from thermodynamic calculations (ii), native Cu can be stable at the activities in $\text{H}_{2,\text{aq}}$ and $\text{H}_{2\text{S},\text{aq}}$ fixed by the HP buffer (Fig. 13), allowing the following equation to be written:



This equilibrium can be used to estimate the minimum a_{H_2} achieved during native copper formation (see Fig. 13-14). Klein and Bach (2009) estimated that H_2 gas could be produced after saturation in the liquid during serpentinization providing an estimate for a_{H_2} variation

with temperature (Fig. 14). If a_{H_2} is fixed on the gas saturation line, native Cu is stable at temperature below ~ 325 °C, whereas chalcocite is stable at higher temperature. Native Cu may thus be formed through chalcocite desulfurization associated with a decrease in temperature.

5.3. Model of native copper formation: Ca-metasomatism at the seafloor in a supra-subduction setting

5.3.1. Cu isotope constraints on the Cu provenance

The copper isotope values for the studied native copper are similar to those reported for bulk mantle rocks (Fig. 10; Ben Othman et al., 2006; Ikehata and Hirata, 2012; Liu et al., 2015; Zou et al., 2019). As discussed above, native Cu formation in the Cheshmeh-Bid pyroxenites can be related to addition of external Cu or desulfurization of pre-existing Cu sulfides. These two hypotheses can be assessed by copper isotopes of other mafic-ultramafic lithologies. Copper sourced from the surrounding mafic ophiolitic rocks provides the most straightforward interpretation of the data. The study by Dekov et al. (2013) augments this interpretation. In their study of native copper associated with modern oceanic crust, they proposed that Cu was mobilized within the basalt with no significant copper fractionation, whereas highly fractionated copper derived from seawater was found in the sedimentary sections of the drilled cores. Ikehata and Hirata (2012) also showed copper isotope value of primary native copper in peridotite was same as that of the host rock and demonstrated that there was no significant copper isotope fractionation during high-temperature magmatic processes.

In-situ alteration of the copper sulfides by metamorphic fluid certainly occurred and could have impacted the copper isotope values in the native copper. How metamorphic fluids could alter the copper isotope composition in this tectonic environment are not well studied, however, Höhn et al. (2017) demonstrated that the metamorphic processes associated with

sulfide deposition generated a relatively tight range of copper isotope value from -0.3 to +0.4 ‰. Equally interesting and related to the formation of native copper in association with basalts, the Michigan native copper associated with secondary mobilization of copper also display a relatively tight range of copper isotope values that overlap the values reported here (Bornhorst and Mathur, 2017; Larson et al., 2003; Mathur et al., 2014). Therefore, significant copper isotope fractionation could not have occurred during secondary alteration by metamorphic fluids. This interpretation is in agreement with the textural and thermodynamic evidence discussed.

Most certainly, the copper isotope values seen in the native copper samples are not related to secondary supergene processes or high temperature magmatic processes. Baggio et al. (2017) demonstrated that copper sourced from local basalts was later redistributed through supergene process that causes the 2 ‰ variation measured. Mathur et al. (2005) and Mathur et al. (2009a) clearly show that highly fractionated copper isotope values in supergene minerals resulted from low temperature oxidation. Ikehata et al. (2011) found that copper isotope values of supergene native copper (from +1.4 to +1.7 ‰) were significantly higher than those of primary chalcopyrite (from -0.3 to -0.1 ‰) from the same deposit.

Due to textural equilibrium of the Cheshmeh-Bid native copper with antigorite and chlorite, high temperature magmatic processes suggested by Ikehata and Hirata (2012) for the primary native copper in the peridotite are unlikely for its formation. Zou et al. (2019) suggested that remarkable copper isotope heterogeneity for Balmuccia pyroxenites is due to variable extent of sulfide segregation as well as melt-peridotite reaction. The absence of any main sulfide phases (e.g. pentlandite, chalcopyrite and pyrrhotite) in the Cheshmeh-Bid pyroxenites precludes the possibility of magmatic sulfide segregation and/or melt-peridotite reaction. Unfortunately, we could not obtain the copper isotope values of the magmatic chalcocite grains because they are very small (Fig. 6f). However, the chalcocites probably has the same

$\delta^{65}\text{Cu}$ values to bulk mantle rocks. Previous studies of copper isotope values of sulfide minerals from igneous-hosted ore deposits are tightly clustered around 0 ‰ (–0.6 to 0.4 ‰) (e.g. Larson et al., 2003).

Our petrographic and geochemical results show that metasomatic fluids were involved for the formation of the Cheshmeh-Bid native copper. Copper isotope variations of native copper grains from the Cheshmeh-Bid pyroxenites are slightly large in comparison with those of non-metasomatised peridotite (Fig. 10; Ben Othman et al., 2006; Ikehata and Hirata, 2012; Liu et al., 2015), probably due to Cu isotope heterogeneity of the magmatic chalcocite grains and isotope fractionations during desulfurization of the sulfides. Our study also indicates that Cu isotopes can be a powerful tool to trace the source of copper in ophiolitic rocks.

5.3. 2. Timing of copper formation

Based on $^{40}\text{Ar}/^{39}\text{Ar}$ dating on hornblendes from plagiogranites and diabase, the genesis of Neyriz ophiolite in the southeast of Neyriz/Khajeh-Jamali ophiolite belt was constrained to Late Cretaceous (range between 83.6 ± 8.4 Ma and 93.19 ± 2.48 Ma) (Lanphere and Pamić, 1983; Babaie et al., 2006). U-Pb zircon dating of plagiogranite and gabbro intrusions provided formation ages of 100.1 ± 2.3 to 93.4 ± 1.3 Ma for the Neyriz ophiolite (Monsef et al., 2018). The nature and formation age of Khajeh-Jamali ophiolitic massifs in the northwest of Neyriz/Khajeh-Jamali ophiolite belt remains currently debated. The uncertainty of geochronological data for the Khajeh-Jamali ophiolitic massifs made it difficult to understand the formation age of these massifs. Uranium–lead dating carried out in this study on titanite grains in association with native copper yield an age of 101 ± 22 Ma (Fig. 9) which may be thus considered as the age of copper mineralization. If the previously reported age interval (83.6 ± 8.4 Ma– 100.1 ± 2.3 Ma) could be attributed to the Khajeh-Jamali ophiolitic massifs, our U-Pb geochronological data on titanite indicates seafloor serpentinization, Ca-

metasomatism and native copper precipitation during formation of an intra-oceanic forearc setting.

5.3.3 A model for copper formation

Considering all available data, a genetic model is proposed for the formation of native Cu occurrence during Ca-metasomatism affecting orthopyroxenites (Fig. 15). High Cr-spinels of the studied pyroxenites bear remarkable chemical resemblance to the peridotites and mantle-hosted chromitites from the studied area (Fig. 5). High Cr#, low TiO₂ content of the Cheshmeh-Bid pyroxenites are consistent with crystallization from arc-related magmas such as high-Mg andesite or boninite. At the first stage, orthopyroxenite dykes formed by magmatic segregation from Mg-andesite or boninite melt within a supra-subduction mantle at oceanic-arc system.

At the second stage, the associated mantle peridotites underwent sub-sea floor hydrothermal alteration. Mantle exhumation was possibly enhanced by the trench-slab roll-back system (e.g. Barth et al., 2008). In such a scenario, Ca²⁺ and H₂ released from the serpentinizing peridotite metasomatised ultramafic lithologies (here pyroxenite dykes). Circulation of hydrothermal fluids formed reaction zones I (amphibole + antigorite), II (clinopyroxene + antigorite) and III (clinopyroxene + chlorite + antigorite), respectively. During late stage of sea-floor serpentinization, native copper was likely formed by low-temperature alteration of Cu-sulfides in the pyroxenites at highly reducing conditions.

6. Concluding remarks

Alteration of pyroxenites from the Cheshmeh-Bid massif led to the formation of a reaction zone composed of three distinct metasomatic mineral assemblages (i) amphibole + antigorite (ii) clinopyroxene + antigorite and (iii) clinopyroxene + chlorite + antigorite (III). Metasomatism was driven by calcium transport and low aSiO₂ fluids which could be related to interaction with Ca-rich fluids formed during clinopyroxene breakdown in the surrounding

serpentinized peridotites. The fluid inclusions data reveal the presence of H₂ inclusions along the cleavage planes and healed cracks of clinopyroxenes in Zone II, revealing that a high activity in H₂ prevailed during native copper formation in the Cheshmeh-Bid pyroxenites. These observations are consistent with thermodynamic calculations showing that native copper can be stable at high H₂ activity. The reducing conditions associated with serpentinization allowed for native copper formation under decreasing T conditions. Tightly clustered copper isotope values of native copper also indicate secondary alteration by metamorphic fluids. According to the Cu isotope composition a mantle origin can be inferred for copper. Serpentinization and Ca-metasomatism likely occurred in an intra-oceanic forearc setting during Albian.

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Figure captions

Fig. 1. a. Map of Mesozoic ophiolites of Iran, showing the Khajeh-Jamali ophiolitic massifs (within red square) in the Zagros Suture Zone; b. simplified geological map of Khajeh-Jamali area (modified after Rajabzadeh, 1998).

Fig 2. Hand samples of: a-b: partly metasomatised pyroxenite with native copper mineralization; c: highly metasomatised and deformed pyroxenite containing native copper mineralization. Scale of clips is 2 cm in length. Opx-rich zone is composed of orthopyroxene and minor clinopyroxene.

Fig. 3. Back-scatter-electron (BSE) images of the Cheshmeh-Bid Cu-rich pyroxenites. (a) Whole profile from Opx-rich portion and metasomatic zones (I) amphibole (Amp) + antigorite (Atg), (II) clinopyroxene (Cpx)+ antigorite and (III) clinopyroxene + chlorite (Chl) + antigorite. Zone I (antigorite+amphibole) is characterized by small narrow selvage of amphibole and antigorite; (b) Ovoid and lamellae inclusions of clinopyroxene in orthopyroxene; (c) Spinifex-like textured olivine (Ol) in the middle of antigorite; (d) Sieve-textured chromite (Chr) included in orthopyroxene (Opx); (e) Euhedral inclusions of chlorite and clinopyroxene in chromite; (f) Zoned clinopyroxene in Zone II; (g) Inclusions of amphibole and orthopyroxene in metasomatic clinopyroxene; (h) Pseudomorph of clinopyroxene replaced by antigorite, chlorite and calcite; (i) Replacement of orthopyroxene by calcite (Cal); (j) Fe-poor (dark grey) and Fe-rich (light grey) chlorite.

Fig. 4. Chemical compositions of olivines from the Cheshmeh-Bid peridotites. a Mg# vs. MnO (wt.%); b. Mg# vs. NiO (wt.%). The olivine compositions from the Cheshmeh-Bid dunites and harzburgites (Rajabzadeh and Nazari-Dehkordi, 2013) are included for comparison.

Fig. 5. Composition of Cr-spinels from the Cheshmeh-Bid orthopyroxenites. (a) Compositional plot of Cr-spinels on the Cr-Al-Fe³⁺ ternary diagram (b) Cr/(Cr+Al) vs.

Mg/(Mg+Fe²⁺) (c) Cr/(Cr+Al) vs. TiO₂ (wt.%). Compositional fields of Cr-spinel from abyssal peridotites, forearc peridotites and podiform chromitites (Miura and Arai, 2014) are shown for comparison.

Fig. 6. Photomicrograph and back-scatter-electron (BSE) images of copper assemblage (zone III): (a-b) Reflected light (a) and cross polarized (b) images of native copper (Cu) intergrown with antigorite (Atg); (c) Native Cu along cleavage planes and along partially healed fractures cutting clinopyroxene (Cpx); (d) native copper filling fracture of chromian spinel (Chr); (e) association of Cu, Cu-Au alloy and titanite (Ttn) in metasomatic clinopyroxene. Cu and Au maps are shown as small insets; (f) chalcocite (Cct) inclusions in native copper.

Fig. 7. Back-scattered electron (BSE) imaging and multi-element wavelength-dispersive spectroscopy (WDS) mapping of sharp contact between orthopyroxene-rich and metasomatised portion of Cu-rich pyroxenite sample.

Fig. 8. (a) Fluid inclusions in the cloudy clinopyroxene of sample X05. In the zoomed area, the fluid inclusions appear in healed fractures and along the cleavage planes of the clinopyroxene. The white arrow shows a whisker, suggesting decrepitation of the FI. Abbreviations: L - liquid phase; V - vapor phase. (b) Main Raman bands of the components (CH₄ and H₂) of the FIs. The stars note the Raman bands of the host clinopyroxene.

Fig. 9. Concordia plots of titanite U–Pb isotopic results

Fig. 10. Variations of $\delta^{65}\text{Cu}$ values in the Cheshmeh-Bid Cu-rich pyroxenites. Data of peridotites from previous studies are also shown for comparison. The grey rectangle represents the estimated $\delta^{65}\text{Cu}$ value of the Bulk Silicate Earth (BSE: $0.07\pm 0.10\%$, Savage et al., 2015).

Fig. 11. Isocon diagram for major (a), and trace and rare earth elements (b) using the method of Grant (2005) comparing average composition of Cheshmeh-Bid fresh orthopyroxenites and average composition of five metasomatised samples. The error bars correspond to +/- one standard deviation. Scaling factors have been used to display a better dispersion of the elements. Scaling factors are shown on isocon diagrams. The lines for constant total mass (black dashed line), constant total volume (black dotted line) and the best fit of the composition in immobile elements (isocon assuming Sc and TiO₂ as immobile; black plain line) are displayed. No change in composition is assumed along the isocon. The grey dashed lines correspond to compositions calculated for different mass gain(+)/loss(-). The densities used for constant volume and volume change calculations are 2896 kg/m³ for the fresh orthopyroxenites and 3870 kg/m³ for the Cu-bearing metasomatised samples. They are retrieved with image analysis of the modal composition in the fresh orthopyroxenite and in zone III. *: estimated based on image analysis.

Fig. 12. Temperature- μ CaO pseudosection for the composition of sample IRKJPX20 (orthopyroxenite) at 50 MPa. μ CaO calculated as a function of temperature is displayed for the orthopyroxenite and three samples affected by Ca-metasomatism (IRKJPX04, IRKJPX11 and IRKJPX12). Tlc: talc; Arg: antigorite; Amp: amphibole; Cpx: clinopyroxene; Chl: chlorite; Ol: olivine; Bro: brocrite; Wo: wollastonite; Mont: monticellite; Ves: vesuvianite; Rnk: rankinite.

Fig. 13. Sulfide and oxide stabilities as a function of $a_{\text{H}_2,\text{aq}}$ and $a_{\text{H}_2\text{S},\text{aq}}$ in the Cu-Fe-Ni-O-S system. The phase diagrams are computed at 50 MPa, $a_{\text{H}_2\text{O}} = 1$ and 200°C (a), 300°C (b) and 400°C (c). The boundaries in the Fe-O-S, Cu-O-S, Fe-Ni-O-S, and Cu-Fe-O-S systems are displayed with grey dashed lines, green plain lines, black plain lines and red plain lines, respectively. The blue spot corresponds to the intersection of the pentlandite/heazlewoodite and the chalcocite/Cu boundaries. This intersection is used to determine the stability of native

copper in a system where $a_{\text{H}_2,\text{aq}}$ and $a_{\text{H}_2\text{S},\text{aq}}$ are buffered by the pentlandite/heazlewoodite equilibrium.

Fig. 14. a_{H_2} as a function of temperature for the $\text{H}_{2,\text{aq}} = \text{H}_{2,\text{g}}$ equilibrium (black line) and the 2 Pentlandite + 12 H_2O + 20 Native Copper = 3 Heazlewoodite + 3 Magnetite + 10 Chalcocite + 12 $\text{H}_{2,\text{aq}}$ equilibrium (red line). This latter equilibrium allows to determine the activity in hydrogen at which the transition from chalcocite to native copper occurs in the rock. Hydrogen partial pressure and hydrostatic pressure are assumed to be equal (50 MPa).

Fig. 15. Schematic model for the genesis of Cu-rich pyroxenite from the Cheshmeh-Bid massif. Figures (a) and (b) are presented as meso-scale.

Supplementary Figure Captions

Supplementary Fig. 1. Pyroxene ternary diagram showing clino- and orthopyroxene compositions from the Cheshmeh-Bid Cu-rich pyroxenites.

Supplementary Fig. 2. Plot of the analysed chlorites on their classification diagram (after Hey, 1954).

Supplementary Fig. 3. Modes (mol.%) of the phases as a function of temperature and chemical potential in CaO for the pseudosection calculated with the composition of sample IRKJPX20 at 50 MPa. (a) Olivine. (b) Antigorite. (c) Clinopyroxene. (d) Amphibole.

Supplementary Table Captions

Table S1. Representative analyses of orthopyroxenes from the Cheshmeh-Bid Cu-bearing pyroxenites

Table S2. Representative analyses of olivine with vein-like texture from the Cu-bearing pyroxenites

Table S3. Representative analyses of chromian spinels in the Cheshmeh-Bid Cu-bearing pyroxenite

Table S4. Representative analyses of amphiboles from the Cheshmeh-Bid Cu-bearing pyroxenites

Table S5. Representative analyses of clinopyroxenes from the Cheshmeh-Bid Cu-bearing pyroxenites

Table S6. Representative analyses of chlorites in Zone III from the Cu-bearing pyroxenites

Table S7. Chemical compositions of sulfides in Zone III

Table S8. Composition of the fluid phase (liquid+vapor) in studied clinopyroxene-hosted fluid inclusions from the PX05 sample.

Table S9. Laser ablation ICP-MS U-Pb analyses including U and Pb contents, isotopic ratios and ages of seven titanite grains from the Cheshmeh-Bid Cu-bearing pyroxenites

Table S10. Copper isotopic results of eight native copper grains from the Cheshmeh-Bid Cu-bearing pyroxenite

Table S11. Whole-rock analyses of the Cheshmeh-Bid pyroxenites.

Declaration of interests

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

- Native copper associated with a metasomatic reaction found in the Cheshmeh-Bid pyroxenite
- Metasomatic reaction zones are interpreted as the result of a Ca-metasomatism on the seafloor during formation of an intra-oceanic forearc setting.

- The source for copper mineralization, as revealed by Cu isotopes, is probably mantle-like.
- Native copper was formed by desulfurization of Cu-sulfides in the pyroxenites at highly reducing conditions.

DRAFT

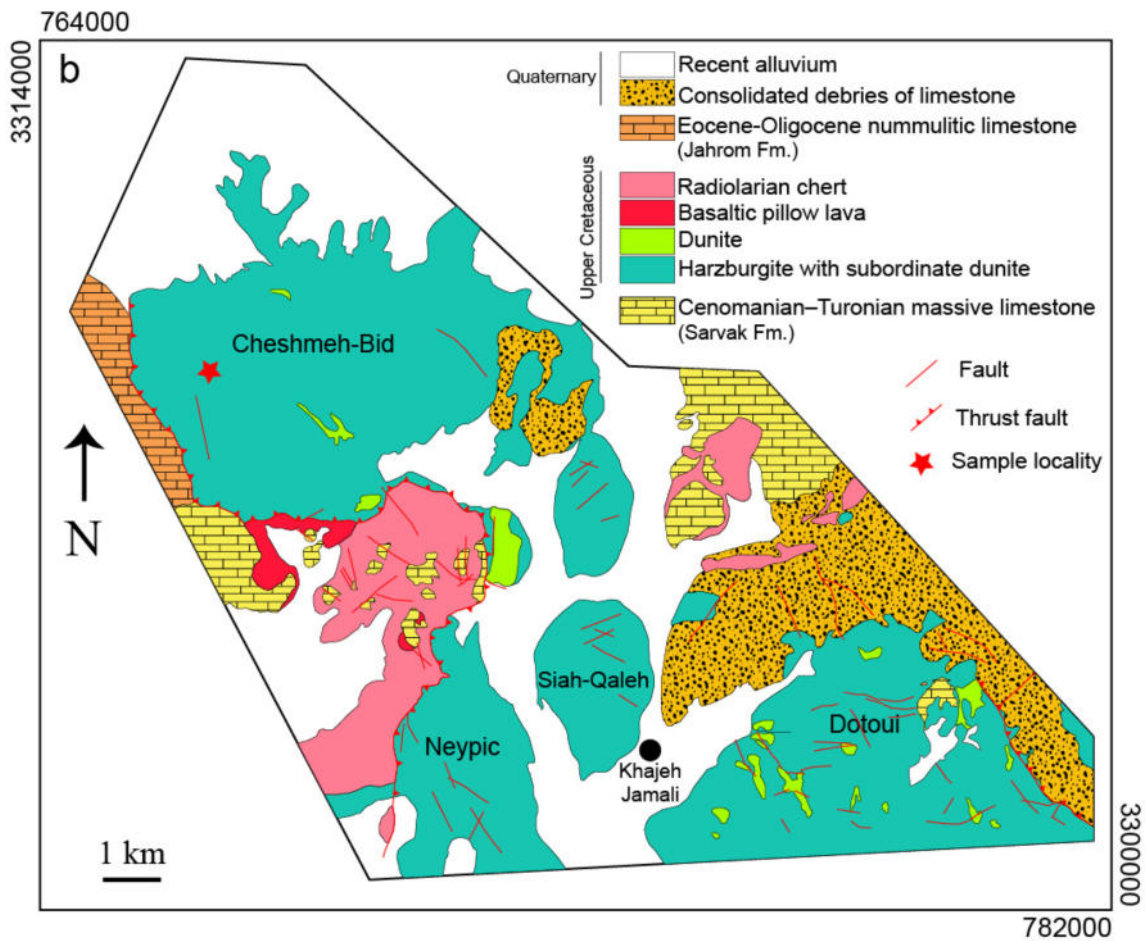
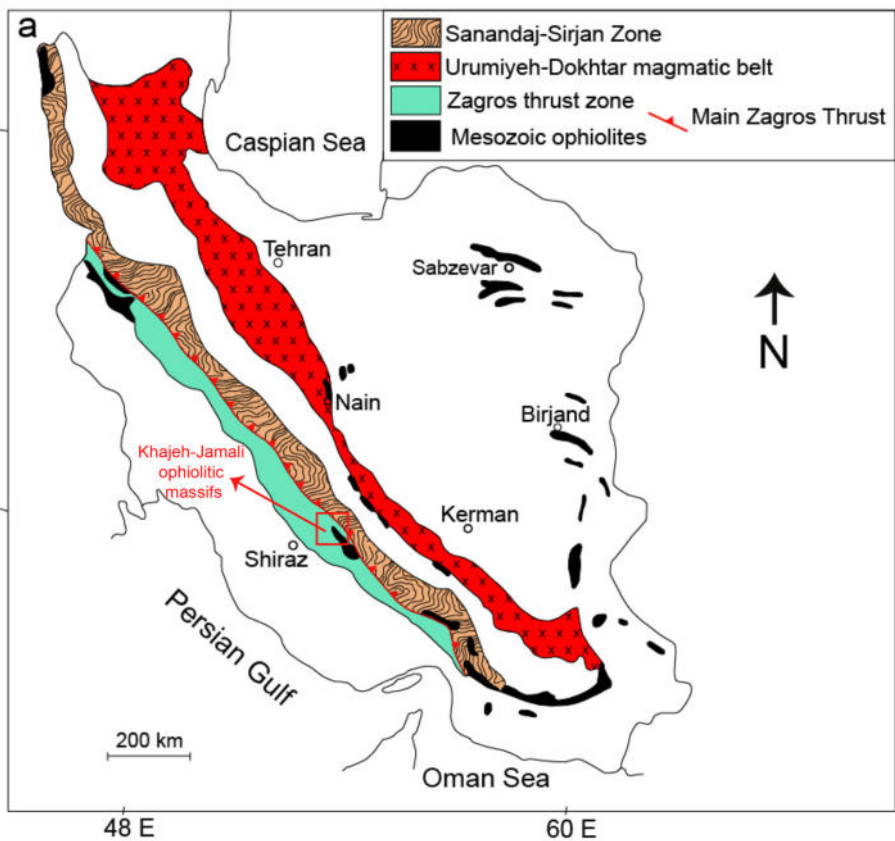


Figure 1

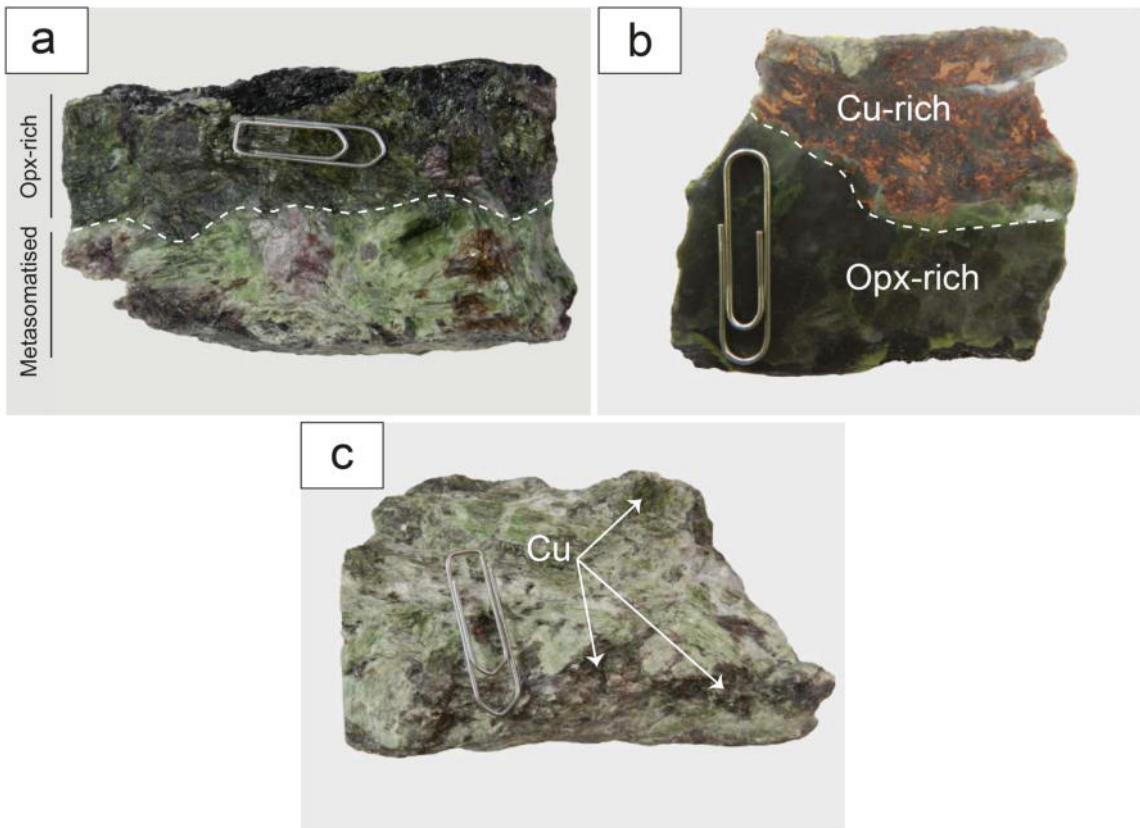


Figure 2

Zone III

Zone II

Opx-rich

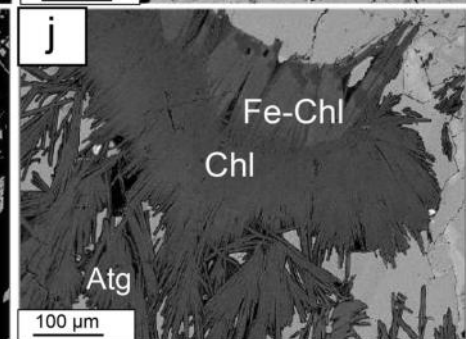
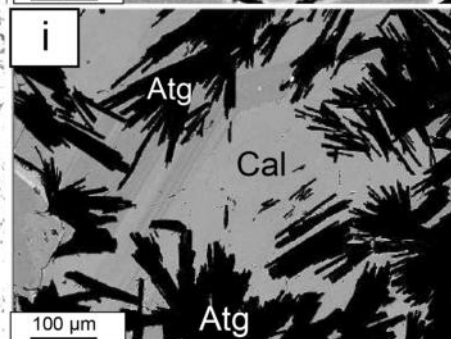
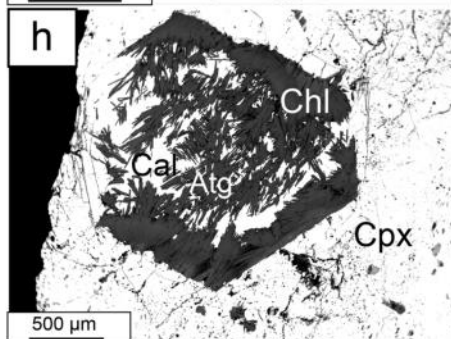
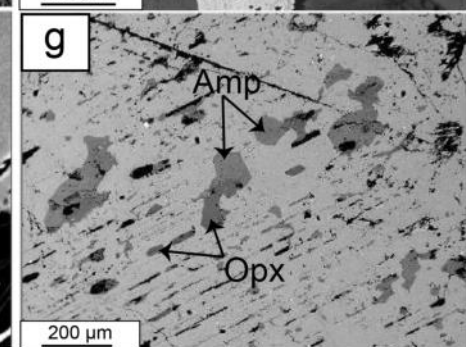
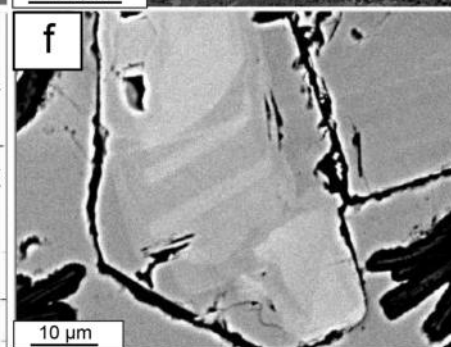
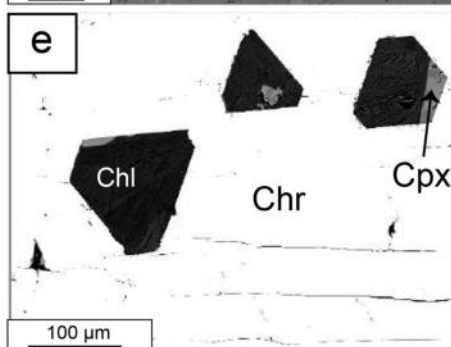
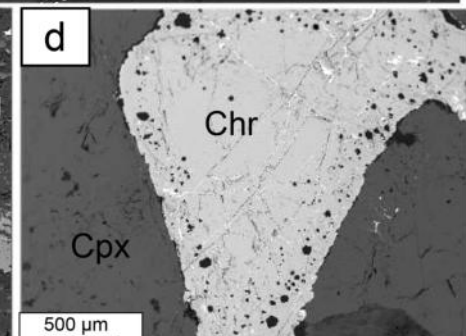
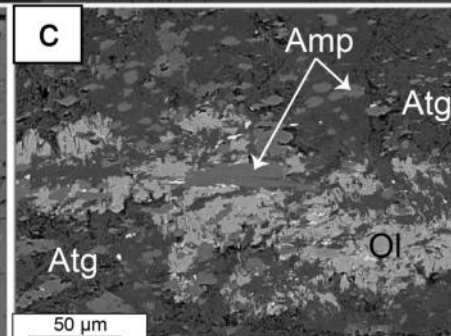
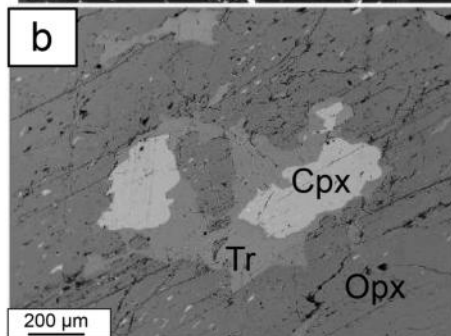
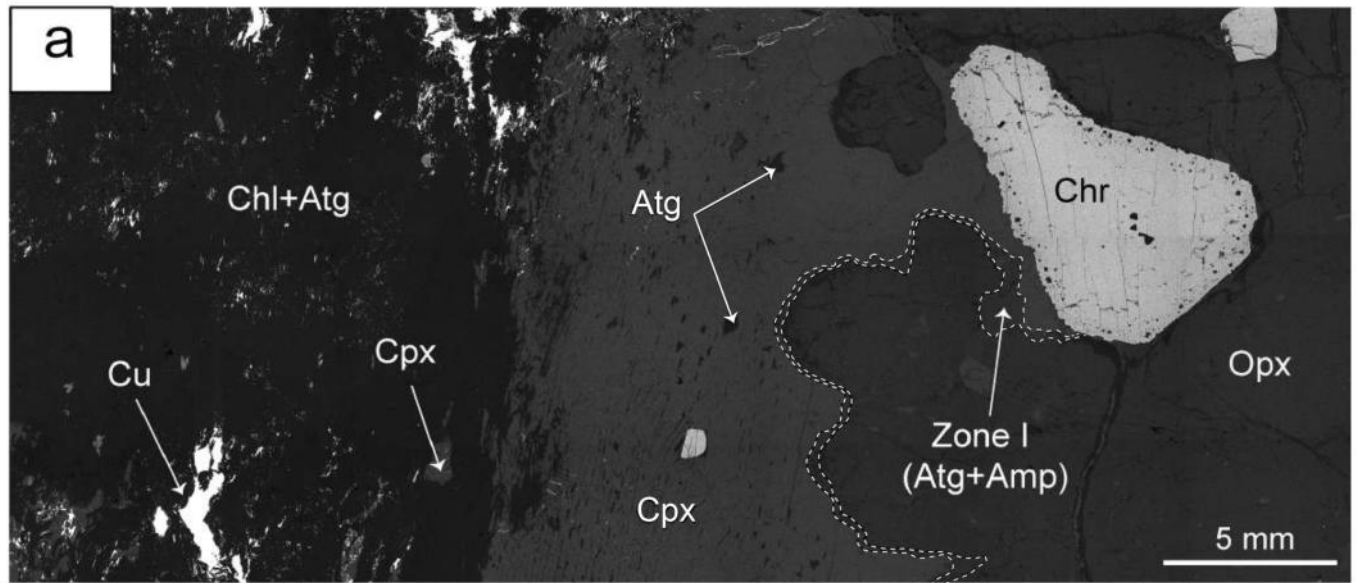


Figure 3

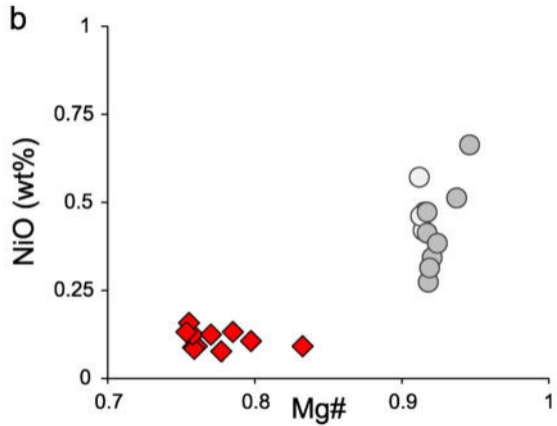
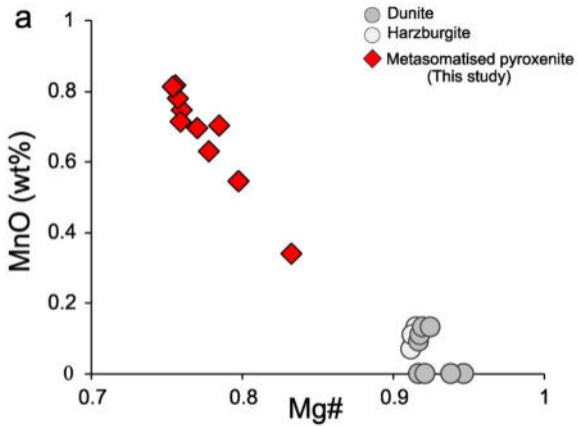


Figure 4

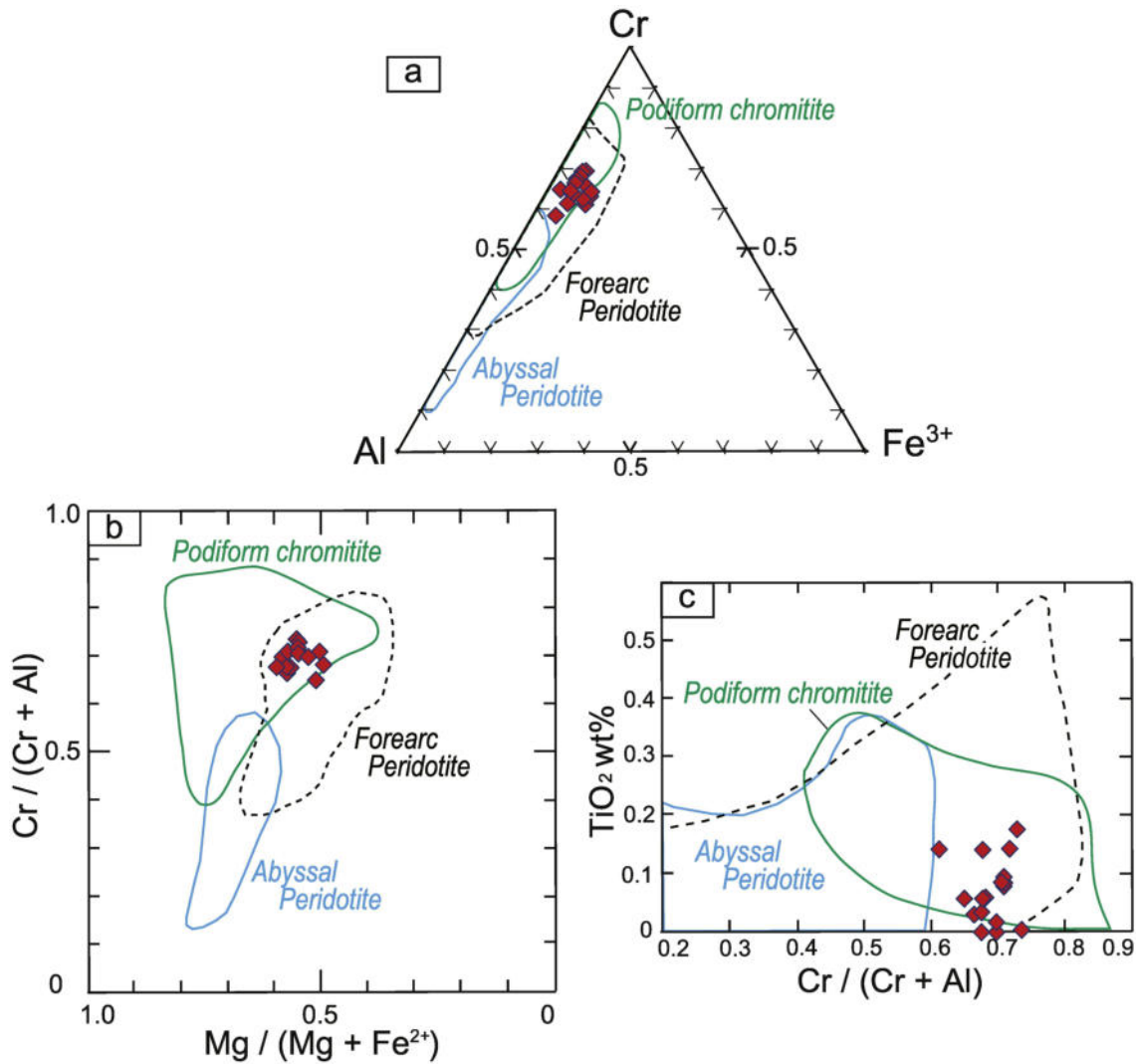


Figure 5

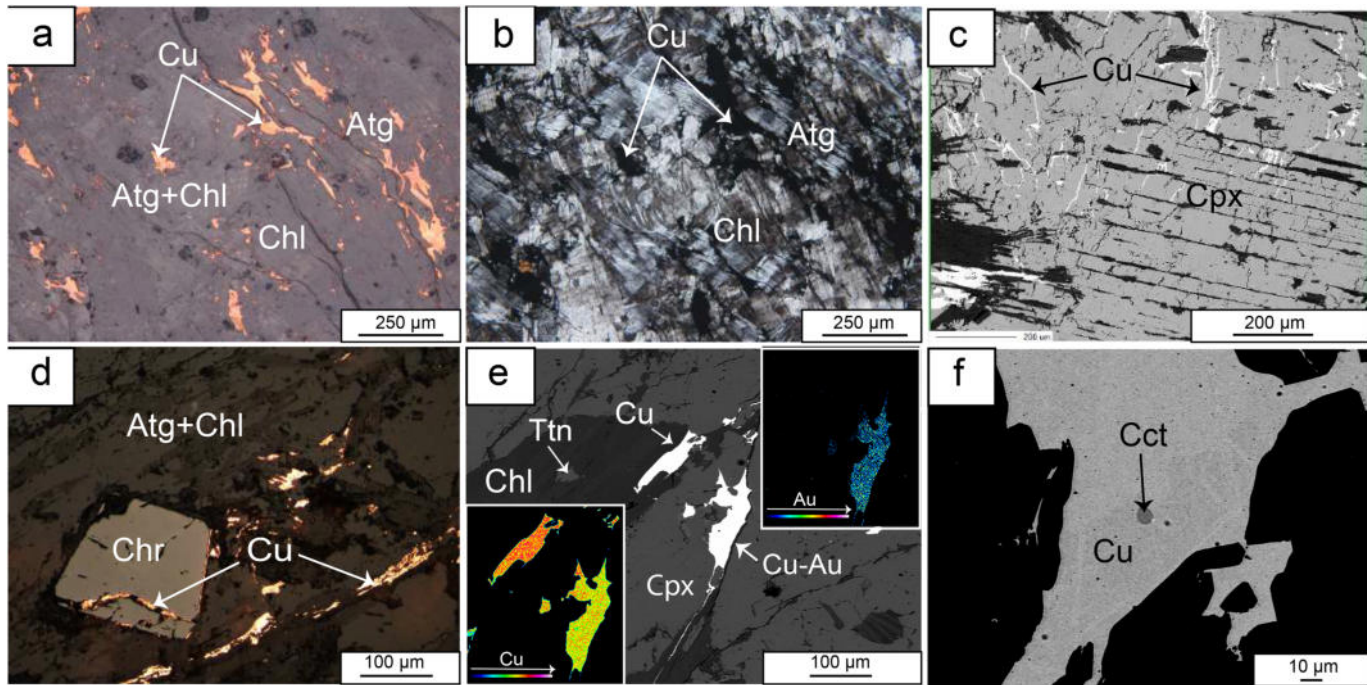


Figure 6

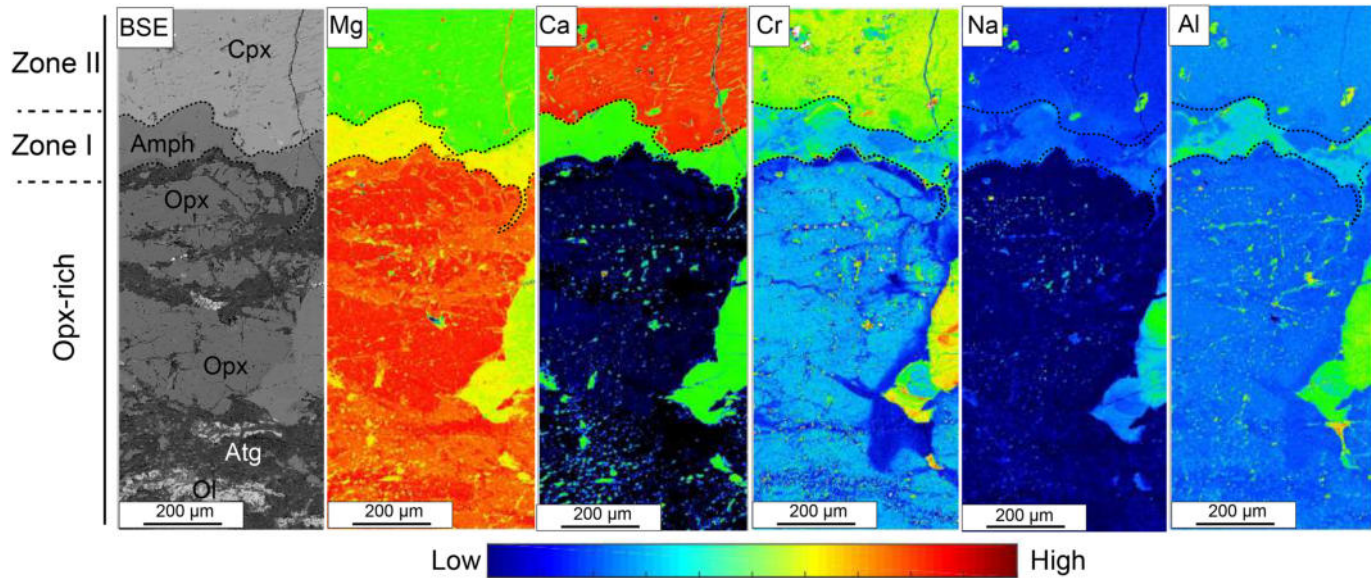


Figure 7

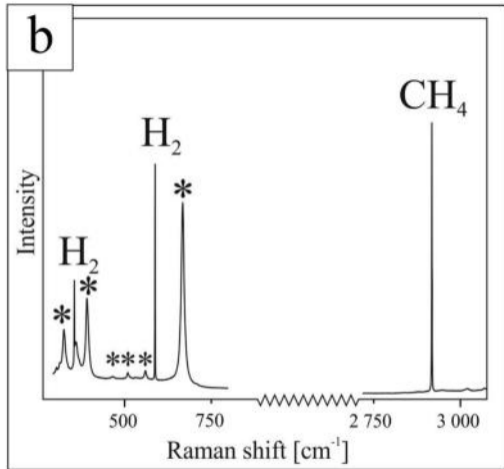
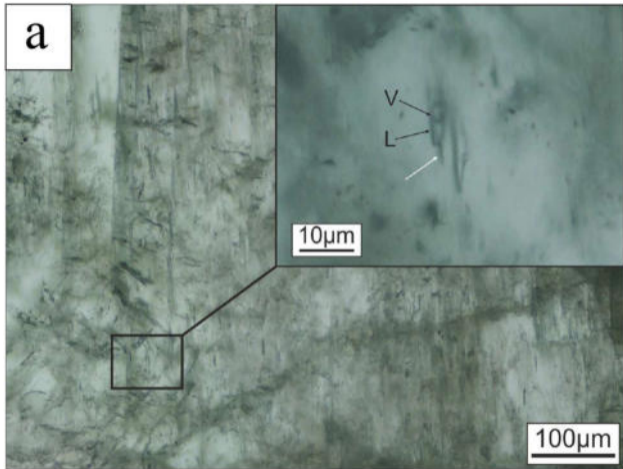


Figure 8

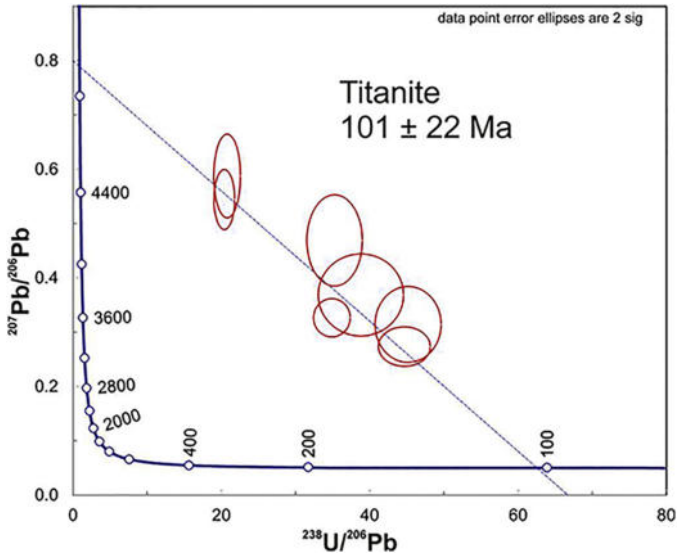


Figure 9

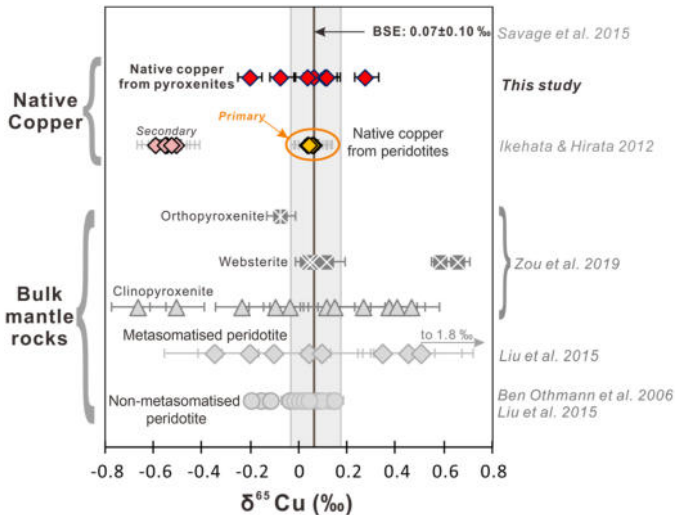
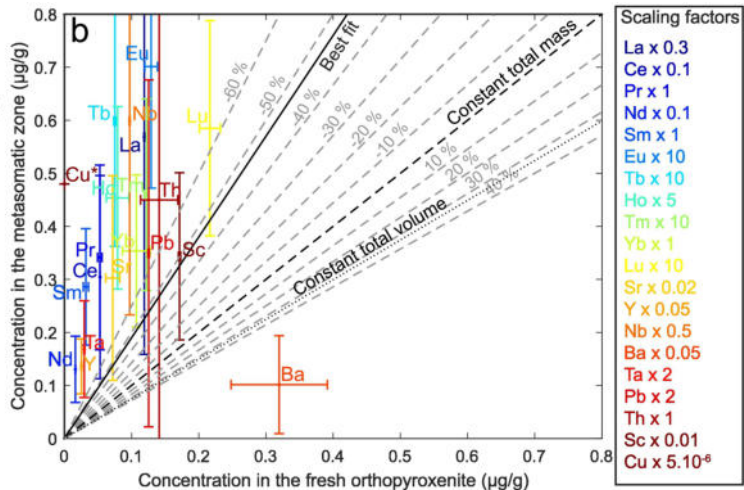
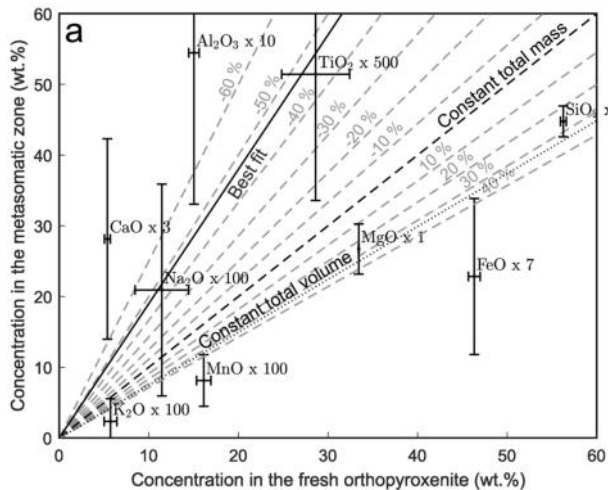


Figure 10



Scaling factors

- La x 0.3
- Ce x 0.1
- Pr x 1
- Nd x 0.1
- Sm x 1
- Eu x 10
- Tb x 10
- Ho x 5
- Tm x 10
- Yb x 1
- Lu x 10
- Sr x 0.02
- Y x 0.05
- Nb x 0.5
- Ba x 0.05
- Ta x 2
- Pb x 2
- Th x 1
- Sc x 0.01
- Cu x 5.10⁻⁶

Figure 11

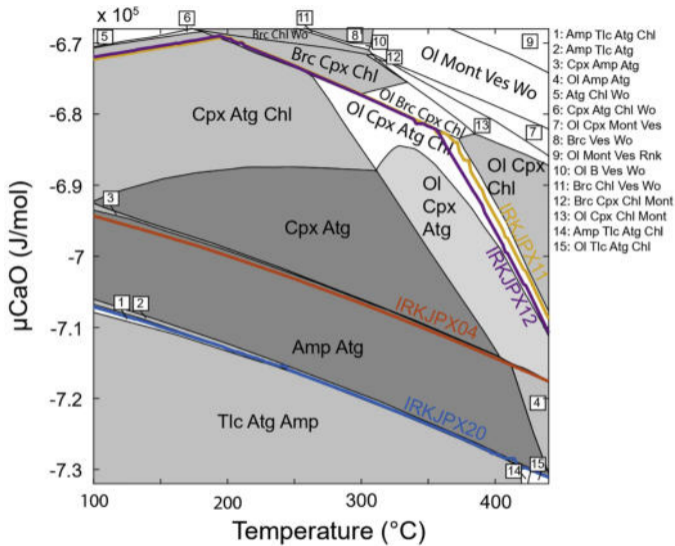


Figure 12

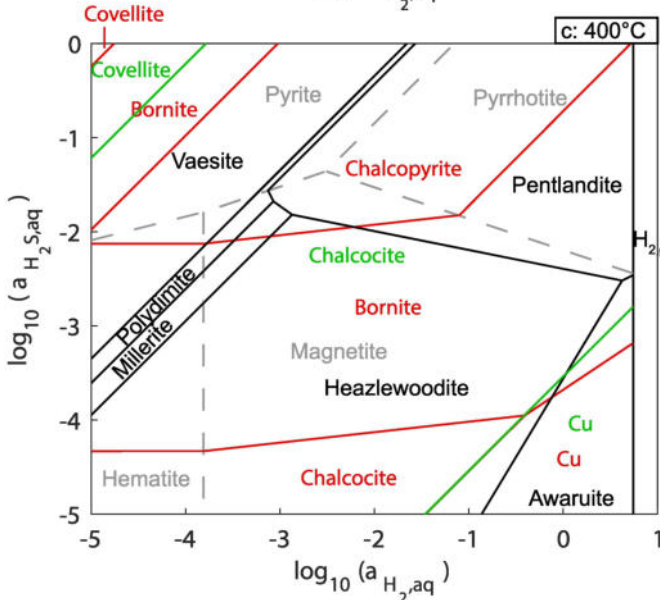
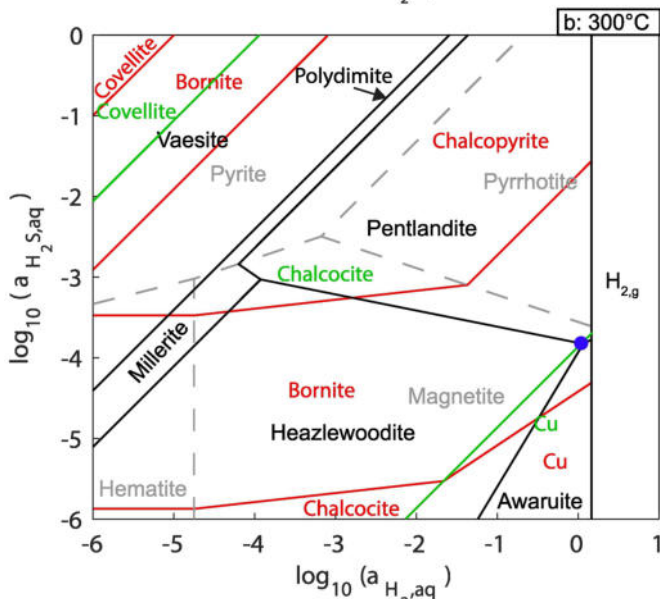
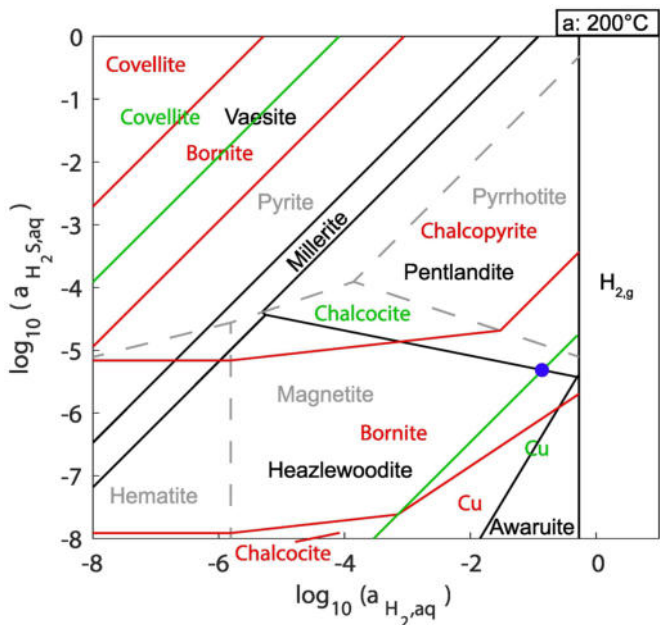


Figure 13

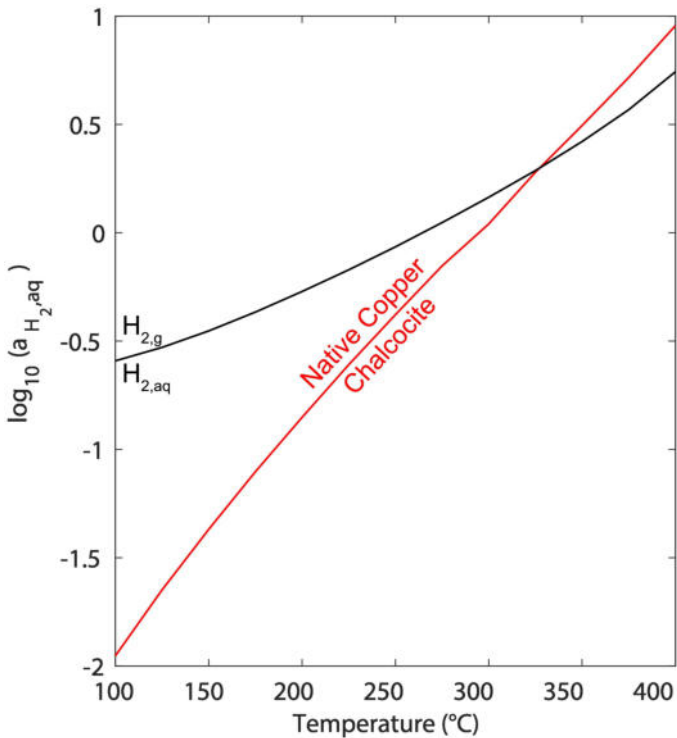


Figure 14

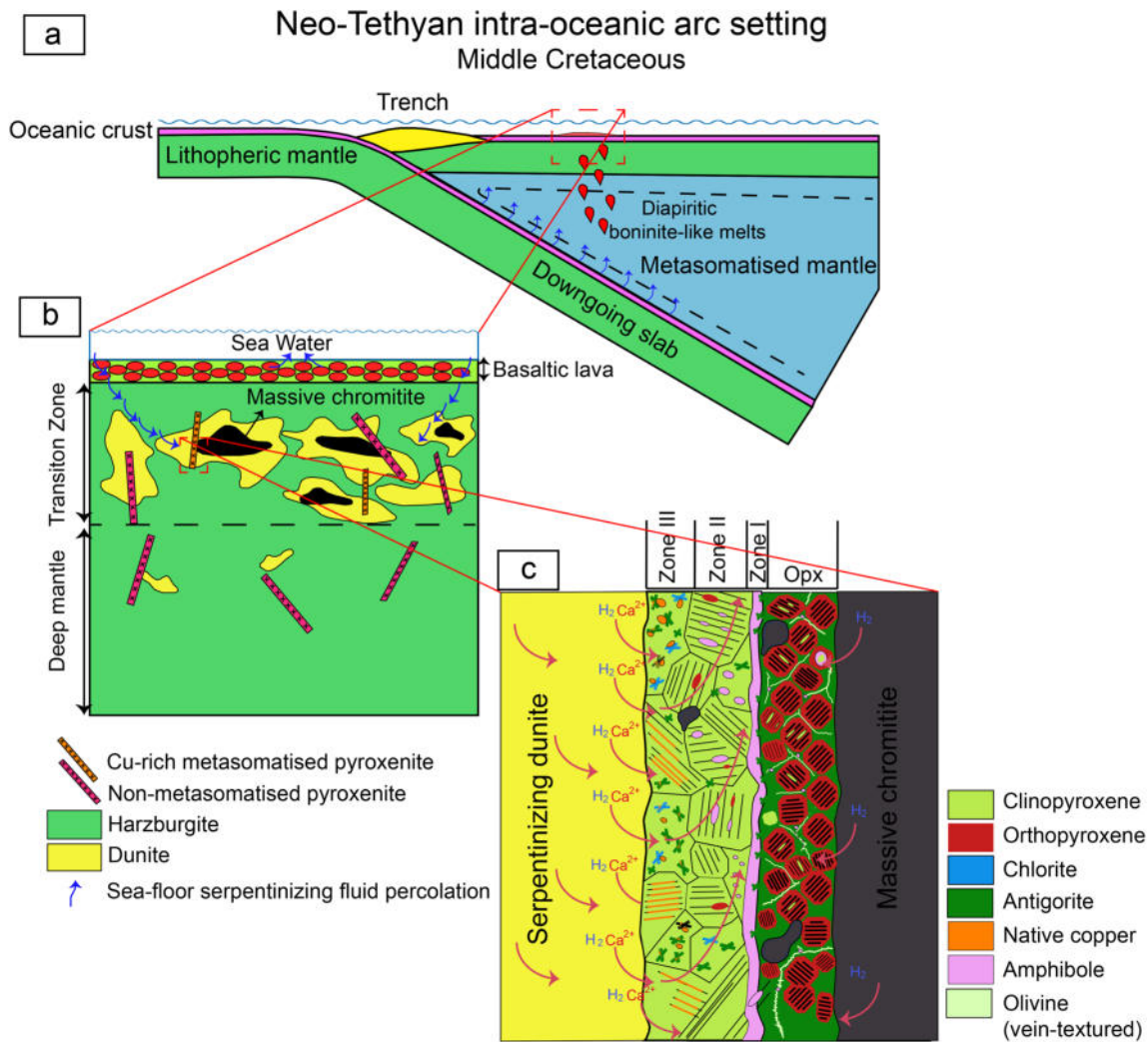


Figure 15