

Evolution of Fe redox state in serpentine during subduction



Baptiste Debret^{a,b,c,*}, Muriel Andreani^d, Manuel Muñoz^e, Nathalie Bolfan-Casanova^{a,b,c}, Julie Carlut^f, Christian Nicollet^{a,b,c}, Stéphane Schwartz^e, Nicolas Trcera^g

^a Clermont Université, Université Blaise Pascal, Laboratoire Magmas et Volcans, Clermont-Ferrand, France

^b CNRS, UMR6524, LMV, Clermont-Ferrand, France

^c IRD, R163, LMV, Clermont-Ferrand, France

^d Laboratoire de Géologie de Lyon, UMR5276, ENS – Université Lyon 1, Villeurbanne, France

^e Institut des Sciences de la Terre, Université Grenoble I, Grenoble, France

^f Institut de Physique du Globe de Paris, France

^g Synchrotron SOLEIL, Paris, France

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ABSTRACT

Serpentinities are an important component of the oceanic lithosphere that formed at slow to ultra-slow spreading centers. Serpentine could thus be one of the most abundant hydrous minerals recycled into the mantle in subduction zones. Prograde metamorphism in subducted serpentinites is characterized by the destabilization of lizardite into antigorite, and then into secondary olivine. The nature of the fluid released during these phase transitions is controlled by redox reactions and can be inferred from oxidation state of Fe in serpentine minerals. We used bulk rock analyses, magnetic measurements, SEM observations and μ XANES spectroscopy to establish the evolution of $\text{Fe}_2\text{O}_3^{\text{Tot}}_{(\text{BR})}$ and magnetite content in serpentinite and Fe oxidation state in serpentine minerals from ridge to subduction settings.

At mid-ocean ridges, during the alteration of peridotite into serpentinite, iron is mainly redistributed between magnetite and oceanic serpentine (usually lizardite). The $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratio in lizardite and the modal percentage of magnetite progressively increase with the degree of local serpentinization to maxima of about 0.8 and 7 wt%, respectively, in fully serpentinized peridotites. During subduction, the $\text{Fe}_2\text{O}_3^{\text{Tot}}_{(\text{BR})}$ of serpentinite remains constant (~ 7 – 10 wt%, depending on the initial Fe content of the peridotite) while the modal percentage of magnetite decreases to less than 2% in eclogite facies rocks. The $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratio in serpentine also decreases down to 0.2–0.4 in antigorite at eclogite facies.

Our results show that, in the first 70 km of subduction, the transition from lizardite to antigorite is accompanied by a reduction of Fe in bulk rock samples and in serpentine minerals. This redox reaction might be coupled with the oxidation of reduced oceanic phases such as sulfides, and the formation of oxidized fluids (e.g. SO_x , H_2O , CO_x). At greater depths, the beginning of antigorite dehydration leads to an increase of $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ in relict antigorite, in agreement with the preferential partitioning of ferric iron into serpentine rather than into olivine.

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1. Introduction

Fluids released during subduction of the oceanic lithosphere are the primary cause of melting in the mantle wedge and arc magmatism. Those magmas commonly display higher $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratios relative to those of mid-ocean ridge and ocean-island basalts (MORB and OIB, respectively) suggesting that the fluids released from the slab upon subduction are oxidized, i.e. dominated by H_2O ,

CO_2 and SO_x species (Arculus, 1994; Evans and Tomkins, 2011; Kelley and Cottrell, 2009; Stolper and Newman, 1994). Those conclusions are consistent with geochemical studies of mantle wedge xenoliths (Andersen and Neumann, 2001; Parkinson and Arculus, 1999). However, recent results based on V/Sc ratios (Lee et al., 2010), fluid inclusions (Song et al., 2009) and thermochemical modeling (Malvoisin et al., 2011) have suggested that reduced fluids are present in subduction zone. These discrepancies emphasize the difficulty in assessing the nature of slab-derived fluids and their respective oxidation states.

The oxidation state of the rocks forming the lithosphere controls the speciation of C-S-O-H-bearing fluids (Frost and McCammon, 2008) that play a fundamental role in metasomatic processes

* Corresponding author. Present address: Department of Earth Sciences, Durham University, Durham, DH1 3LE, UK.

E-mail address: baptiste.debret@durham.ac.uk (B. Debret).

in subduction zones. As hydrous minerals are intimately linked to subduction processes, the nature of the released fluids is controlled by redox reactions during hydrous mineral phase change or dehydration. Serpentinites form through the hydration of oceanic lithosphere at slow spreading centers (Mével, 2003) and are an important component of subduction zones (Hattori and Guillot, 2007; Reynard, 2013). Furthermore, since they contain up to ~12 wt% of water, they constitute a large reservoir of water compared to other hydrous rocks forming the oceanic lithosphere and may therefore play a major role in the transfer of fluids in subduction zones.

At slow or ultra-slow spreading ridges, the upper 3–6 km of oceanic lithosphere are highly serpentinized (Cannat et al., 2000, 1995, 2010). The serpentinization of ultra-mafic rocks is associated with magnetite formation (Bach et al., 2006; Oufi et al., 2002), while the fluids discharged from serpentinites can be H₂ and CH₄ rich (Charlou et al., 2002; Marcaillou et al., 2011). These observations indicate that olivine-hosted Fe becomes oxidized as water is reduced during serpentine crystallization (Berndt et al., 1996; Klein and Bach, 2009; Klein et al., 2009; McCollom and Bach, 2009; Seyfried et al., 2007). Recent studies have shown that the increase of the serpentinization degree of the peridotite is correlated with an increase of the magnetite mode and Fe³⁺/Fe_{Total} ratio of the serpentine (Andreani et al., 2013; Marcaillou et al., 2011). These observations imply that the serpentinites that constitute the upper oceanic lithosphere and which are ultimately subducted are highly oxidized relative to mantle peridotites.

During subduction, serpentinites are situated in the upper 3–6 km of the slab (Debret et al., 2013a; Reynard et al., 2010). With prograde metamorphism, the progressive serpentine phase changes (lizardite → antigorite → olivine/pyroxene) drive fluid release to the mantle wedge (Hattori and Guillot, 2007) until depths of about 150–180 km, where dehydration of serpentine to form olivine, enstatite and chlorite assemblages (Trommsdorff et al., 1998; Fumagalli and Poli, 2005) should be complete (Ulmer and Trommsdorff, 1995; Wunder and Schreyer, 1997). Geochemical observations based on fluid mobile element (B, Li, As, Sb, Ba, Rb and Cs), halogens (F, Cl) and volatiles (S) behavior during prograde metamorphism in subduction zones reveal a direct link between the composition of the fluid released during serpentine phase transitions and those of arc magmas (Debret et al., 2013b, 2014; Savov et al., 2005, 2007; Scambelluri and Tonarini, 2012; Vils et al., 2011). Nonetheless, there is no overall consensus on the evolution of the redox state of serpentinites during subduction.

Exhumed high-pressure serpentinites in ophiolites from localities such as the Western Alps provide an alternative means of constraining the redox state of slab-derived fluids. Indeed, these ophiolites are considered to have been highly hydrated and serpentinized during an oceanic stage (distal continental margin or mid-ocean ridge environment) prior to partial dehydration during prograde metamorphism (Debret et al., 2013a, 2013b; Hattori and Guillot, 2007; Lafay et al., 2013; Li et al., 2004; Schwartz et al., 2013; Vils et al., 2011). Here we propose to use the redox state of iron in serpentinites, an abundant element in this type of rock, in order to assess the possible nature of the released fluids during serpentine phase changes in subduction zones. We provide the first in-situ (μXANES spectroscopy) and bulk rock chemistry measurements of Fe redox state from alpine serpentinites, which record different *P–T* conditions representative of a cross section through a subducting slab. In contrast to arc magmas, which may have undergone processes such as crystal fractionation or assimilation, these samples provide a relatively direct means of constraining the redox state of subducted lithosphere. Furthermore, the study of high-pressure metamorphic rocks allows changes in Fe redox state as a function of serpentine metamorphism during subduction to be directly constrained.

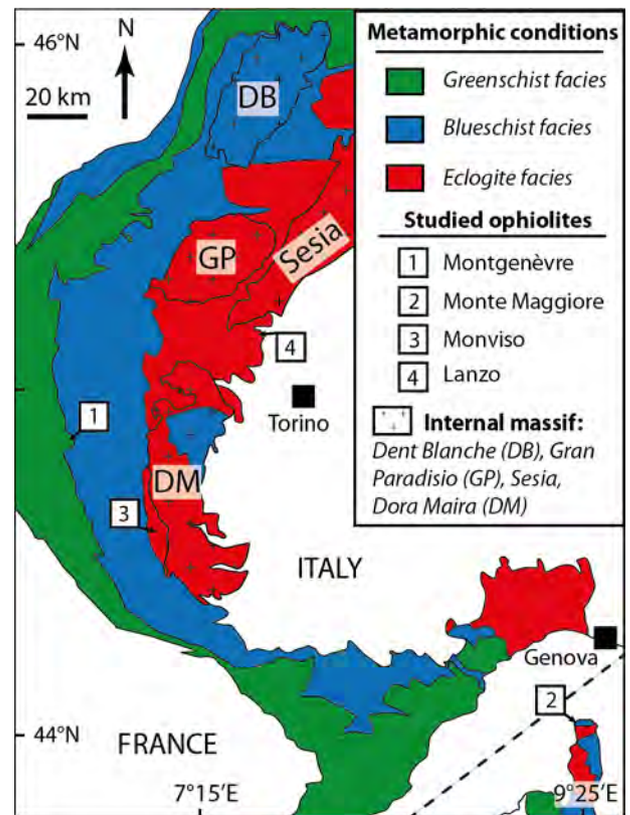


Fig. 1. Geological map of the Western Alps showing the metamorphic facies and the spatial distribution of studied ophiolites. The numbers indicate the studied ophiolites: 1. The Mont Genève ophiolites (Chenaillet and Punta Rascia massifs); 2. Monte Maggiore ophiolite; 3. Mon Viso ophiolite and 4/Lanzo ophiolite.

2. Geological setting and petrographic observations

The Western Alps formed as the result of the subduction of the Tethyan oceanic lithosphere beneath Apulia between late Jurassic and early Tertiary (Lombardo et al., 2002). The Tethyan oceanic lithosphere is an equivalent to the modern Atlantic Ocean lithosphere (Lagabrielle and Cannat, 1990) and is composed of intrusive gabbroic pods surrounded by serpentinites and sometimes topped by basalts and/or sediments (Cannat et al., 1995). In order to study the redox state of Fe along prograde metamorphism, we sampled various alpine meta-ophiolites recording different metamorphic conditions (Fig. 1) representative of a subduction gradient. Two main phase transitions are observed in the serpentinites of this metamorphic suite: the transition from lizardite to antigorite that occurs from greenschist to blueschist facies at ~300–400 °C (Evans, 2004; Schwartz et al., 2013), and the dehydration of antigorite into secondary olivine at *T* > 600 °C in the eclogitic facies (Bromiley and Pawley, 2003).

Oceanic serpentinites have been widely described and are known to be mostly composed of lizardite, the low-pressure/low-temperature (LP/LT) variety of serpentine (Evans, 2004), and of chrysotile (Andreani et al., 2007; Mével, 2003). They form pseudomorphic mesh and bastite textures, replacing olivine and orthopyroxene respectively (Mével, 2003). The mesh texture appears grey under cross polarized light and forms homogenous areas of lizardite with finely disseminated magnetite. The mesh cell is delimited by fibrous lizardite rims associated with stringy magnetite aggregates (Fig. 2a). The magnetite strings consist of euhedral magnetite grains in equilibrium with the surrounding lizardite (Fig. 2b). Bastite textures consist of white serpentine grains elongated parallel to the original orthopyroxene cleavages (Fig. 2a) while clinopyroxene is typically resistant to low temper-

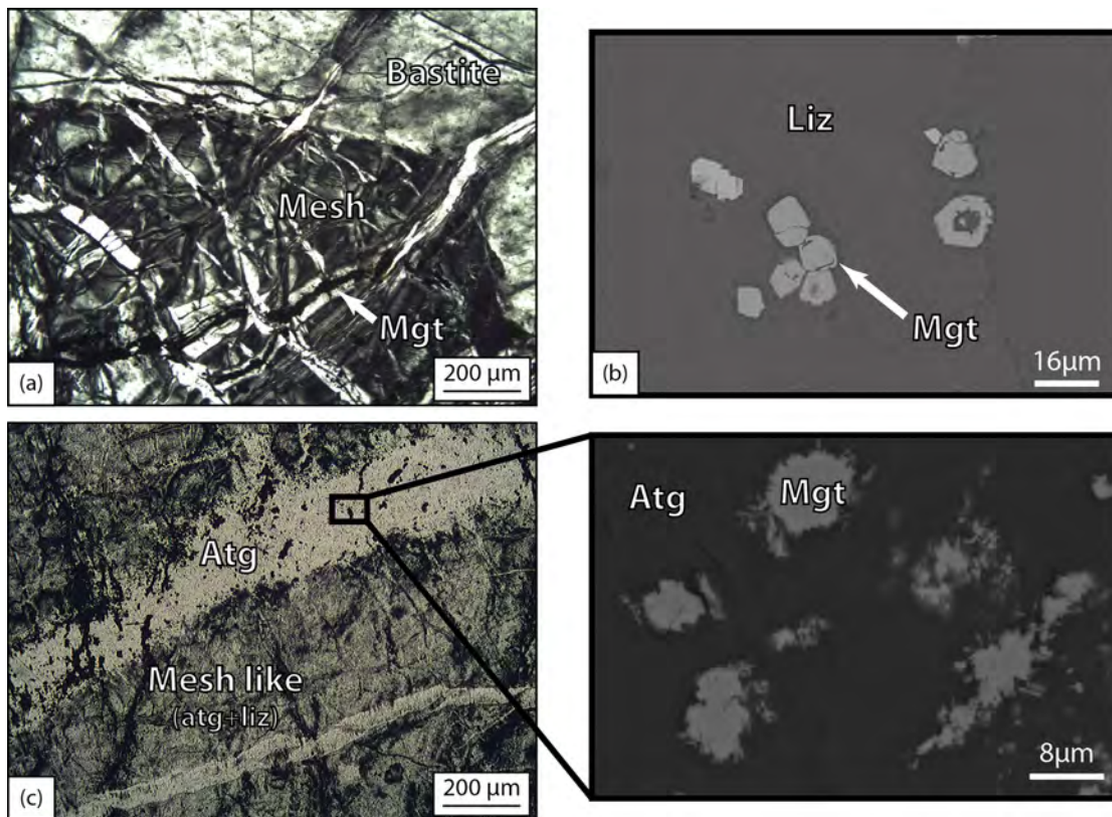


Fig. 2. (a) Photomicrograph in crossed polarized light showing typical mesh and bastite textures. The mesh rims are associated with a fine string of magnetite. (b) SEM photomicrograph in back scattered electron (BSE) of magnetite grains associated with mesh texture. (c) Photomicrograph in plane polarized light (left photo) and BSE (right photo) of a serpentinite from the Montgenèvre ophiolite. The mesh like texture is composed of lizardite (Liz) and antigorite (Atg) intergrown on a micrometer scale. A late antigorite vein crosses the mesh texture. The magnetite grains associated with both textures display indented boundaries (right photo).

ature serpentinization (Andreani et al., 2007; Klein et al., 2013; Mével, 2003).

2.1. The Montgenèvre ophiolite

The Montgenèvre ophiolite is located in the external Piemonte zone, 6 km west of Briançon (Fig. 1). It is a thin klippe resting onto the Lago Nero Unit (Caby, 1995). The massif is composed of metagabbro pods occasionally topped by basalts that display greenschist parageneses (Mével et al., 1978), and are surrounded by massive serpentinites (serpentinization >80%). The massif is commonly interpreted as an oceanic portion of the upper part of the Tethyan oceanic lithosphere (Chalot-Prat, 2005; Manatschal et al., 2011). Most of serpentinites are composed of lizardite and chrysotile assemblages displaying mesh and bastite textures (Liz-serpentinites) crystallized in an oceanic setting (Schwartz et al., 2013; Lafay et al., 2013). The local crystallization of antigorite, high-pressure and high-temperature serpentine variety (HP/HT) (Wunder et al., 2001), at the expense of lizardite (Atg/Liz-serpentinites) can be interpreted in terms of an increase in P – T conditions during subduction. It is marked in thin section by the partial recrystallization of mesh and bastite textures into antigorite, which display intermediate Raman spectra between lizardite and antigorite (mesh-like and bastite-like textures; see Debret et al., 2013a for further details), and by the occurrence of pure antigorite veins crossing oceanic textures (Fig. 2c). The magnetite grains observed in antigorite veins and within mesh-like rims display indented boundaries interpreted as dissolution textures (Fig. 2c). In contact with pure antigorite, the primary spinel is zoned: it has an Al–Cr–Fe core surrounded by a thin double corona (<30 μ m) of chromite and magnetite.

2.2. The Monte Maggiore ophiolite

The Monte Maggiore ophiolite is located in the northern end of the Cap Corse, Northern Corsica (Fig. 1). It is an ultramafic body of ~ 4 km² surrounded by eclogitized continental units. The massif is a fragment of oceanic lithosphere generated in an ocean–continent transition (OCT) context (Jackson and Ohnenstetter, 1981; Piccardo and Guarnieri, 2010). It is composed of slightly serpentinized peridotites (SSP; <20% serpentinization) intruded by mafic pods and dykes displaying blueschist facies paragenesis (Nicollet et al., 2001; Vitale Brovarone et al., 2013). The SSP are plagioclase- and/or spinel-harzburgites and lherzolites. They are affected by two main serpentinization episodes (Debret et al., 2014): the first one corresponds to the crystallization of lizardite veins from primary minerals during massif oceanization and the second event corresponds to antigorite veins formed from lizardite and primary minerals during subduction prograde HP metamorphism. In the southern part, the massif is bordered by a serpentinite envelope of <1 km-wide. This envelope is affected by an East–West foliation that surrounds boudins of massive serpentinite. The foliated serpentinites are mostly composed of antigorite (Atg-serpentinites) while the massive boudins are composed of mesh- and bastite-like textures and pure antigorite (Atg/Liz-serpentinites). At mesh-like rims, the lizardite and magnetite are replaced by thin lamellae of antigorite with only rare relicts of magnetite displaying indented boundaries (Fig. 3a). In these rocks, the primary spinel is coronitic: at the contact of antigorite, it is recrystallized into a chromite–magnetite (Mgt2) assemblage (Fig. 3b). The clinopyroxene has a coronitic texture composed of antigorite lamellae associated with magnetite grains (Fig. 3c). The magnetite grains are sub-euhedral attesting that they are equilibrated with the surrounding antigorite (Fig. 3c): they constitute a new generation of magnetite (Mgt2).

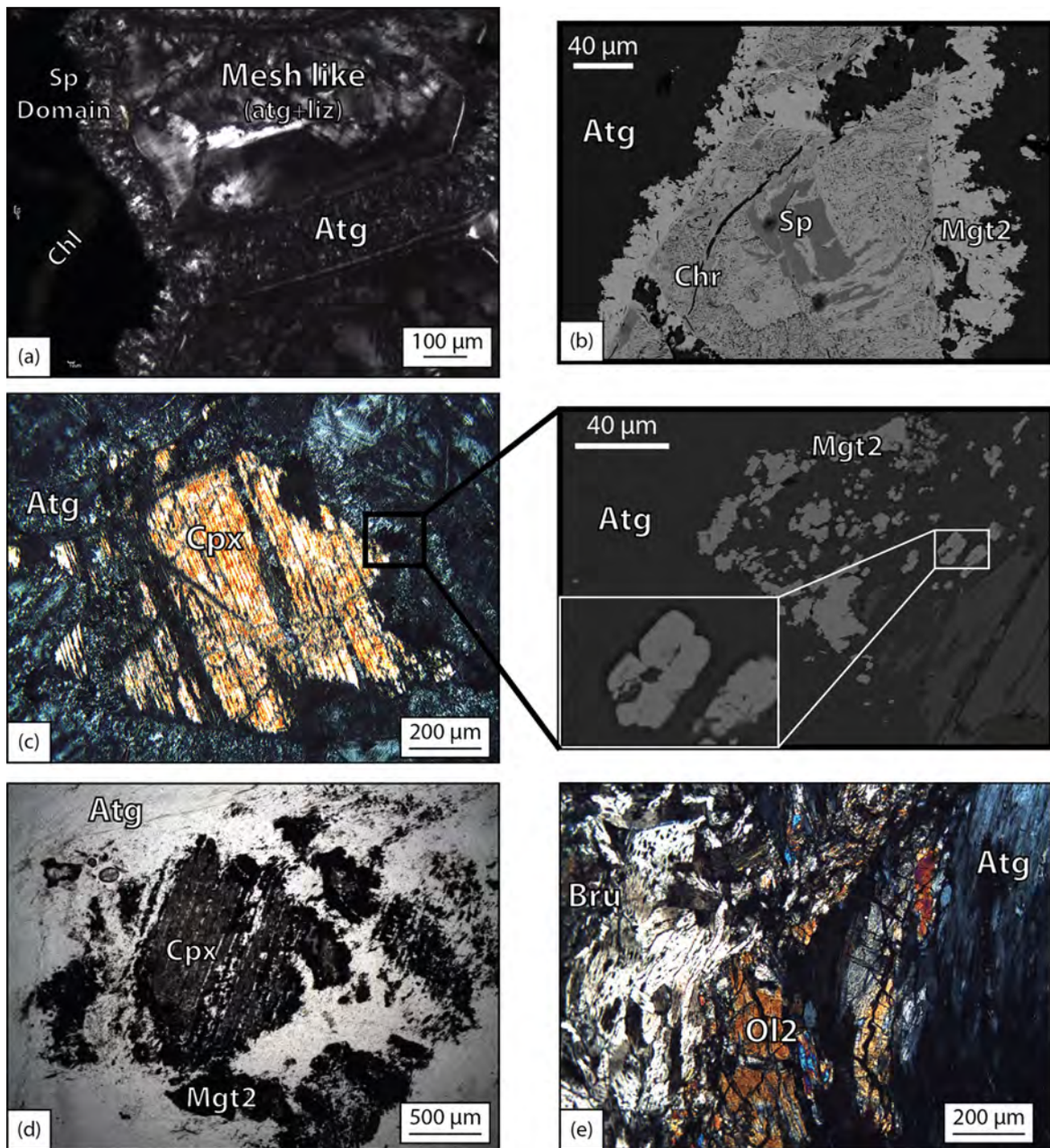


Fig. 3. (a) Photomicrograph in crossed polarized light of serpentinite from Monte Maggiore. Mesh rims are recrystallized into thin lamellae of antigorite. The mesh core is composed of lizardite and antigorite associated at nano-scale (Mesh-like texture). (b) SEM microphotography in BSE of a spinel domain from Monte Maggiore serpentinite. In contact with antigorite, the primary spinel (Sp) displays a double corona composed of chromite (Chr) and magnetite (Mgt2). (c) Photomicrograph in crossed polarized light (left photo) and in BSE (right photo) of a serpentinite from Monte Maggiore. The clinopyroxene relict is surrounded by an antigorite and magnetite corona. The magnetite associated with the antigorite corona is sub-euhedral (Mgt2, right photos). (d) Photomicrograph in plane polarized light of a clinopyroxene relict partly recrystallized into antigorite and secondary magnetite. In the left corners of the photo, the olivine domain is composed of antigorite without magnetite. (e) Photomicrograph in crossed polarized light of secondary olivine (Ol2) formed at the expense of brucite (Bru) and antigorite from a Monviso serpentinite.

2.3. The Monviso massif

The Monviso massif is buried between the Dora Maira massif and the Queyras accretionary prism (Fig. 1). It represents oceanic lithosphere fragments blended during massif exhumation (Schwartz et al., 2001). It is composed of metagabbroic pods metamorphosed under the eclogite facies and massive serpentinites boudins surrounded by foliated serpentinites (Schwartz et al., 2001). Massive serpentinites are mostly composed of antigorite (Atg-serpentinites). Few relicts of mesh-like textures are observed in these rocks and can be interpreted as relicts of oceanic serpen-

tinization. The foliated serpentinites are greatly recrystallized into antigorite lamellae (Fig. 3d): the olivine domain is replaced by thin antigorite lamellae of $<5 \mu\text{m}$ and the orthopyroxene domain by antigorite lamellae of $\sim 20\text{--}50 \mu\text{m}$ long. The spinel domain is composed of chromite and magnetite assemblages of $\sim 100 \mu\text{m}$ width. Clinopyroxene has recrystallized into oriented antigorite lamellae of $\sim 20\text{--}50 \mu\text{m}$ long associated with secondary magnetite aggregates; the primary clinopyroxene is sometimes preserved in the center of these aggregates (Fig. 3d). Locally, prograde antigorite is partly dehydrated into secondary olivine at eclogite facies conditions (Schwartz et al., 2013).

2.4. The Lanzo massif

The Lanzo massif is an eclogitized ultramafic body of 150 km² bounded by the sediments of the Po plain to the East and South, by meta-ophiolites and the Schistes Lustrés unit in the West, and by the HP metamorphic continental unit of Sesia in the North (Pelletier and Müntener, 2006). The massif is a portion of oceanic lithosphere preserving its oceanic structure (Debret et al., 2013a; Lagabrielle et al., 1989). It is composed of a central part of SSP surrounded by a serpentinite envelope of 3–5 km width. The ultramafic rocks record different steps of serpentinitization (Debret et al., 2013a): lizardite crystallization during an oceanic serpentinitization step; prograde and retrograde antigorite crystallization during subduction and massif exhumation; locally secondary olivine granoblasts of ~20 µm formed at the expense of prograde antigorite and magnetite at metamorphism peak condition of the massif. As observed in the Monviso serpentinites, in the serpentinite envelope, the olivine domain is replaced by thin antigorite lamellae thinner than 5 µm, the spinel domain by chromite and magnetite assemblages of ~100 µm widths, and the pyroxene domain by antigorite lamellae of ~20–50 µm sometimes associated with magnetite masses of ~20 µm widths.

A set of 13 serpentinites and 6 SSP representatives of serpentine phase changes during prograde metamorphism were selected for this study (Table 1). Most of these samples have already been characterized in previous petrological (Debret et al., 2013a; Schwartz et al., 2013) and geochemical (Debret et al., 2013b; Lafay et al., 2013) studies. The metamorphic conditions and petrographic characteristics of the studied samples are summarized in Table 1. In the Liz/Atg- and Atg-serpentinites no retrograde phases (e.g. talc, chrysotile amphiboles; Padrón-Navarta et al., 2011; Müntener et al., 2000) are observed suggesting that they were little affected by retrograde metamorphism. In the Liz/Atg-serpentinites, antigorite is observed to have formed at the expense of mesh and bastite textures (Figs. 2c and 3a). We therefore assume that antigorite crystallization is relative to prograde metamorphism stage.

3. Methods

3.1. Bulk rock analyses

Bulk rock powders for major elements measurements were prepared with the addition of Li-metaborate. The melts obtained by fusion of this mixture in a magnetic induction oven were subsequently quenched and diluted by a factor of 2000 for analysis. Major element concentrations were determined by ICP-AES at the Laboratoire Magmas et Volcans of Clermont-Ferrand (France). Concentrations were calibrated against the DRN basaltic glass. Precision on samples were determined by repeated analyses of the reference basalt BHVO. This typically resulted in a better than 1% precision for all measured elements. Additional Fe²⁺ analyses of bulk rock (BR) samples were performed at the SARM-CRPG (Nancy, France). These analyses were done by automatic titration at the equivalent point with potassium dichromate after dissolution of the sample in a HF/H₂SO₄ mixture, in the presence of H₃BO₃ and H₃PO₄. Subsequently Fe³⁺/Fe_{Total(BR)} ratios were calculated from the measuring Fe_{2O₃Tot(BR)} and Fe²⁺_{Tot(BR)} values (e.g. Andreani et al., 2013).

3.2. Hysteresis measurements

The amount of magnetite in serpentinite samples can be estimated from the saturation magnetization derived from hysteresis magnetic measurements. Hysteresis cycles at room temperature up to maximum field of 500 mT to 700 mT were measured using

a Princeton Vibrating Sample Magnetometer (VSM) at IGP Paris. At least two fragments of ~1 cm³ for each serpentinite samples were used and the saturation magnetization was then determined by taking the average value. The saturation magnetization of the serpentinite depends linearly on concentration of magnetite; this calculation is made using a proportionality factor of 92 Am²/kg between J_s and the nominal weight of magnetite (Malvoisin et al., 2012; O'Reilly, 1984).

3.3. In-situ analyses

In situ major element analyses of minerals of ultramafic rocks were performed with a microprobe CAMECA SX 100 at the Laboratoire Magmas et Volcans in Clermont Ferrand (France). Microprobe analysis mean values and FeO maximal and minimal values of repeated analysis of serpentine and secondary olivine forming serpentinites and SSP are given in Appendix A, MMC 1.

Iron speciation was measured by X-ray absorption spectroscopy at the iron K-edge at the LUCIA beamline of SOLEIL synchrotron (Source Optimisée de Lumière d'Énergie Intermédiaire du LURE, France). Measurements were operated with a current and energy of 400 mA and 2.75 GeV, respectively. The X-ray absorption near edge structure (XANES) spectra were collected using a Si(311) double crystal monochromator. The energy calibration was performed using an iron foil. XANES spectra were measured in fluorescence mode using a four-element silicon drift diode (SDD) detector with a total active area of 40 mm². The beam spot size was set to 4 × 4 µm² by using two dynamically bendable mirrors in Kirkpatrick–Baez configuration.

XANES spectra were acquired from 7050 eV to 7300 eV with a sampling step of 0.1 eV in the pre-edge region. In order to minimize potential photo-oxidation effects (Appendix A, MMC 2), the dwell time was adjusted to 0.5 s per point, resulting in a total acquisition time of 12 min. To improve signal-to-noise ratio, each XANES spectrum is the average of 4 spectra acquired on 4 distinct locations, separated by 30 µm, in a same serpentine texture. X-ray fluorescence (XRF) chemical maps were used to identify minerals in the thin sections and to selected spots for XANES measurements.

XANES spectra were normalized using the Athena© software (Ravel and Newville, 2005). Following previous XANES studies (e.g. Berry et al., 2003, 2010; Bolfan et al., 2012; Farges et al., 2001; Muñoz et al., 2013; Wilke et al., 2001), the Fe³⁺/Fe_{Total} ratio of serpentines were derived by fitting the pre-edge region (shown in Fig. 4) using PeakFit© software. The background of the pre-edge region was modeled using the tail of a Gaussian function and the pre-edge peak was deconvoluted using two pseudo-Voigt functions. This treatment results in absolute uncertainties for the determination of the pre-edge centroid of ±0.05 eV in energy, and ±0.025 in integrated area (e.g. Muñoz et al., 2013; Wilke et al., 2001). The Fe³⁺/Fe_{Total} was derived directly from the variogram of Wilke et al. (2001). This variogram was reproduced by measuring three powdered standard compounds namely olivine (^{VI}Fe²⁺), andradite (^{VI}Fe³⁺) and sanidine (^{IV}Fe³⁺), previously characterized in other studies (Andreani et al., 2013; Marcaillou et al., 2011; Muñoz et al., 2013).

Previous studies reported important changes for XANES spectra and pre-edge peaks depending on crystal orientation relative to the polarized X-ray beam (Bolfan et al., 2012; Evans et al., 2014; Muñoz et al., 2013). Nevertheless, a recent comparison between the XANES spectra of antigorite crystals and powders revealed that maximal orientation effects of antigorites were reduced relative to other phyllosilicates (Muñoz et al., 2013). Due to the unknown orientation of antigorite crystals, Fe³⁺/Fe_{Total} might be underestimated by a maximum of 0.1 or overestimated by 0.05.

Table 1

Bulk rock analysis of serpentinites and SSP. The error on major element content is smaller than 0.5 wt%.

Rock	Serpentinite								Blueschist				Eclogite				SSP	
Metamorphic facies	Greenschist				Blueschist				Eclogite				Blueschist		Eclogite			
Locality	Mont Genève				Monte Maggiore				Lanzo				Monte Maggiore		Lanzo			
Sample name	ICH02	Bch10	Bch6	Bch9	MM8	MM15	MM19	MM2	Vis1	LZ14b	RO1	LZ27a1	Vis5b	MAG33	MAG26	LZ17a	LZ19	
Rock type	Liz-serpentinite		Atg/Liz-serpentinite		Atg/Liz-serpentinite		Atg-serp		Atg/Liz-serp.		Atg-serpentinite		Ol ₂ /Atg-serpentinite					
Serpentine minerals	Liz	Liz	Liz ± Atg	Liz, Atg	Liz, Atg	Liz, Atg	Atg ± Liz	Atg	Liz, Atg	Atg	Atg ± Ol2	Atg, Ol2	Atg, Ol2	Liz ± Atg	Atg ± Liz	Liz, Atg	Atg, Ol2	
SiO ₂	39.6	38.8	39.2	40.2	38.6	39.5	39.1	40.1	40.0	39.0	39.7	40.0	38.3	41.1	42.1	42.6	41.9	
Al ₂ O ₃	1.0	2.0	2.6	3.0	1.9	2.1	2.0	1.8	2.0	2.4	2.8	2.3	1.3	2.3	2.2	2.6	3.0	
Fe ₂ O ₃	7.8	8.1	7.9	7.9	9.5	7.9	8.9	8.0	8.1	8.4	8.1	8.5	9.2	8.4	8.3	8.9	8.9	
MgO	37.9	37.8	36.5	36.0	37.4	36.3	38.3	37.8	36.2	38.1	36.0	38.5	40.1	39.5	40.1	39.5	38.6	
CaO	0.1	1.0	1.3	1.8	0.9	1.9	0.4	0.7	2.6	0.0	2.1	1.5	0.9	2.2	2.4	2.6	3.1	
Na ₂ O	b.d.l.	0.0	0.0	0.0	0.0	0.0	0.0	0.0	b.d.l.	0.3	0.2	0.3	b.d.l.	0.0	0.0	0.1	0.1	
K ₂ O	b.d.l.	0.0	0.0	b.d.l.	b.d.l.	0.0	0.0	0.0	b.d.l.	0.3	0.2	0.2	0.0	0.1	0.1	b.d.l.	b.d.l.	
TiO ₂	0.0	0.1	0.1	0.1	0.1	0.1	0.1	0.0	0.0	0.1	0.1	0.1	0.0	0.1	0.1	0.1	0.1	
MnO	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.1	0.0	0.1	0.1	0.1	0.1	0.1	0.1	0.1	
P ₂ O ₅	b.d.l.	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0	
LOI	13.2	12.1	11.8	11.0	12.1	11.3	10.9	11.6	10.4	11.3	10.5	7.9	9.9	6.0	4.2	3.1	3.4	
Tot	99.6	100.1	99.6	100.3	100.6	99.3	99.7	100.2	99.3	100.1	99.9	99.5	99.9	99.8	99.6	99.7	99.3	
FeO (Fe ²⁺ titration)	2.1	2.7	2.8	3.3	2.8	3.7	4.8	3.0	3.5	3.6	3.6	6.3	4.2	5.5	7.1	6.5	7.0	
Fe ³⁺ /Fe ^{Tot}	0.7	0.6	0.6	0.6	0.7	0.5	0.4	0.6	0.5	0.5	0.5	0.2	0.5	0.3	0.1	0.2	0.2	
%Mag (stdr deviation)	6(0.4)	6.1(1.5)	3.4(0.9)	3.2(1.0)	4.9(1.3)	2.2(1)	1.2(0.2)	1.4(0.5)	3.4(0.8)	1.8(0.7)	1(0.1)	<1	1.2(0.9)	<1	<1	<1	<1	
Analyzed fragments	3	4	4	2	2	2	2	2	2	4	2	1	2	1	1	1	1	

b.d.l.: below detection limit.

Table 2
Pre-edge characteristics of XANES standards and serpentine minerals. The XANES standards have been measured with exactly the same conditions as the sample data set and during the same run.

Sample	Mineral	Texture	Pre-edge centroid		Fe ³⁺ /Fe _{Total}	Error Fe ³⁺ /Fe _{Total} (±)
			Area	Position		
Standards	Olivine		0.079	7112.96	0	
	Sanidine		0.457	7114.19	1	
	Andradite		0.158	7114.19	1	
Serpentinities						
ICh02	Liz	Mesh	0.184	7114.14	0.93	0.06
BCh6	Liz ± Atg	Mesh	0.165	7114.07	0.85	0.06
	Liz, Atg	Vein	0.156	7114.17	0.97	0.06
	Liz, Atg	Vein	0.171	7114.08	0.85	0.06
	Liz, Atg	Vein	0.207	7114.08	0.84	0.06
BCh9	Liz, Atg	Mesh	0.188	7114.02	0.76	0.07
	Liz, Atg	Mesh	0.176	7114.07	0.83	0.06
	Liz, Atg	Mesh	0.199	7114.05	0.80	0.08
	Liz, Atg	Mesh	0.151	7113.97	0.74	0.06
	Atg	Vein	0.153	7113.94	0.68	0.06
	Atg	Vein	0.139	7113.92	0.69	0.05
	Atg	Vein	0.142	7113.77	0.50	0.06
MM8	Liz, Atg	Mesh	0.134	7114.14	0.93	0.06
	Liz, Atg	Mesh	0.133	7114.02	0.80	0.06
	Liz, Atg	Mesh	0.129	7114.06	0.85	0.06
	Liz, Atg	Mesh	0.126	7114.04	0.83	0.06
	Liz, Atg	Mesh	0.124	7114.02	0.80	0.06
	Liz, Atg	Mesh	0.139	7114.09	0.87	0.05
	Atg	Mesh rim	0.133	7114.03	0.81	0.05
	Atg	Mesh rim	0.125	7114.07	0.86	0.05
	Atg	Cpx rim	0.177	7114.04	0.80	0.06
	Atg	Cpx rim	0.162	7114.07	0.84	0.06
	Atg	Cpx rim	0.138	7114.08	0.87	0.05
	Atg	Sp rim	0.141	7114.15	0.95	0.06
	Atg	Sp rim	0.144	7114.06	0.84	0.06
	LZ14b	Atg	–	0.120	7113.74	0.52
Vis5b	Atg	Associated with Ol2	0.114	7113.84	0.63	0.05
LZ27a1	Atg	Associated with Ol2	0.131	7113.69	0.44	0.06
SSP						
LZ17	Liz	Vein	0.123	7113.94	0.72	0.06
	Atg	Vein	0.145	7113.69	0.39	0.08
Mag32	Liz	Vein	0.134	7113.86	0.62	0.05
	Liz	Vein	0.122	7113.90	0.67	0.05
LZ35c	Atg	Vein	0.122	7113.64	0.40	0.05
	Atg	Vein	0.131	7113.50	0.13	0.10
LZ19	Atg	Vein associated with Ol2	0.123	7113.72	0.49	0.05
LZ34b	Atg	Vein associated with Ol2	0.128	7113.85	0.62	0.06
	Atg	Vein associated with Ol2	0.134	7113.78	0.53	0.05

Moreover, the same authors mention that areas of the pre-edge do not change significantly with crystal orientation.

The centroid energy and integrated area of the pre-edge peaks are displayed in Figs. 5a and 5b and are reported in Table 2; including uncertainty estimates. The entire dataset of serpentine plots in the ^{VI}Fe²⁺–^{VI}Fe³⁺–^{IV}Fe³⁺ region (Fig. 5). This suggests the absence of Fe²⁺ in serpentine tetrahedra and the occurrence of Fe³⁺ in serpentine octahedra and tetrahedra as proposed in previous crystallographic and chemical studies (Andreani et al., 2013; Fuchs et al., 1998; Marcaillou et al., 2011; O’Hanley and Dyar, 1993).

4. Results

4.1. Bulk rock analysis

Among the selected samples, there is no obvious change in Fe₂O₃^{Tot}_(BR) (BR: Bulk Rock) with prograde metamorphism, indicating that Fe is immobile in serpentinities during subduction (Table 1). The Fe₂O₃^{Tot}_(BR) content ranges from 7.79 to 9.53 wt%

(Table 1) and these slight variations most probably reflect variations in the modal abundances of the protolith minerals; it has previously been shown that the major element compositions of serpentinities are mainly controlled by primary mineralogy (Godard et al., 2008; Mével, 2003; Paulick et al., 2006). Furthermore, according to previous geochemical studies on these samples (Debret et al., 2013b; Lafay et al., 2013), the selected serpentinities have primary mineral modes similar to plagioclase/spinel lherzolites. The different Fe₂O₃^{Tot}_(BR) contents among the selected samples are thus attributed to slight variations in clinopyroxene, spinel and plagioclase contents. The Fe₂O₃^{Tot}_(BR)-poor serpentinities are formed from spinel- and clinopyroxene-poor lherzolites while the Fe₂O₃^{Tot}_(BR)-rich serpentinities are formed from spinel- and clinopyroxene-rich lherzolites.

In the serpentinities, the Fe³⁺/Fe_{Total}(BR) decreases with the metamorphic grade (Fig. 6a). At the Montgenèvre ophiolite, the Liz-serpentinities have a Fe³⁺/Fe_{Total}(BR) ratio ranging from 0.65 to 0.7 while the Atg/Liz-serpentinities display lower Fe³⁺/Fe_{Total}(BR) ratio ranging from 0.56 to 0.62. At the Monte Maggiore, the Atg/Liz-serpentinities have a Fe³⁺/Fe_{Total}(BR) ratio varying from 0.51 to 0.68

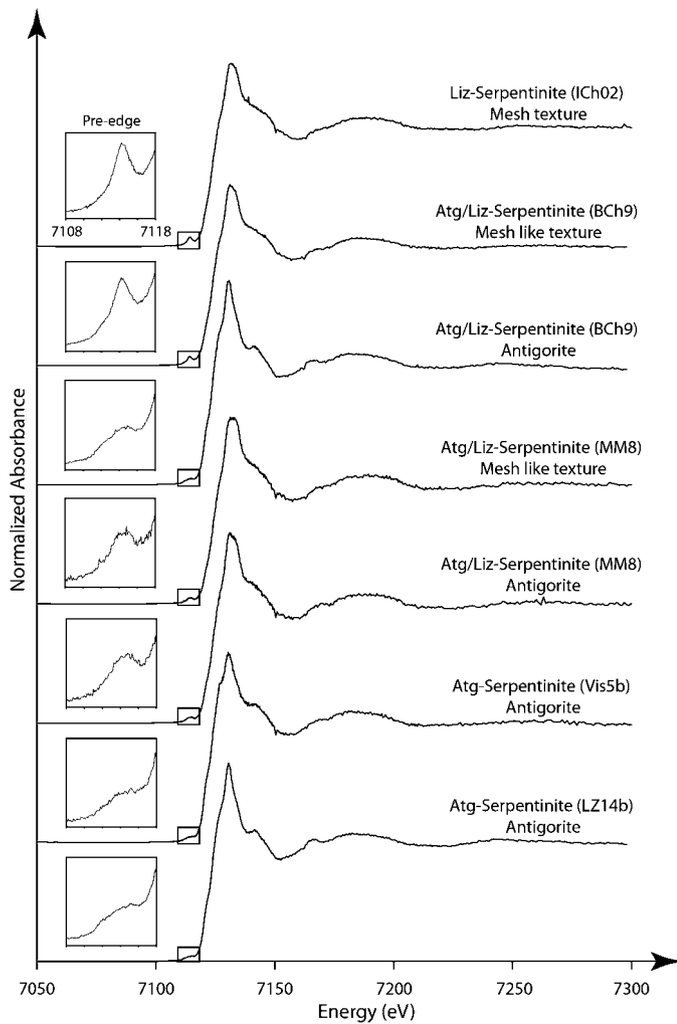


Fig. 4. Spectral signature at the K-edge for the different serpentinite textures in the Montgenèvre (ICh02, BCh9), Monte Maggiore (MM8), Monviso (Vis5b) and Lanzo (LZ14b) massifs. The box shows the pre-edge region of each XANES spectra.

and the one of Atg-serpentinites varies from 0.43 to 0.6. Under eclogite facies (Monviso and Lanzo massif), the $Fe^{3+}/Fe_{Total} (BR)$ ratio in the Atg/Liz-serpentinites and Atg-serpentinites ranges from 0.53 to 0.54. The occurrence of secondary olivine in Atg/Ol2-serpentinites is marked by a decrease of this ratio to 0.24–0.52.

The measured magnetite content of selected serpentinites and SSP is reported in Table 1. The magnetite mode progressively decreases with metamorphic grade (Fig. 6b). At the Montgenèvre ophiolite, the Liz-serpentinites have a mean magnetite mode of ~6% whereas the mean magnetite mode of the Atg/Liz-serpentinite is about 3.3% (Table 1). At the Monte Maggiore, Atg/Liz-serpentinites display a mean magnetite mode varying from 2.2 to 4.9%, while Atg-serpentinites mean magnetite mode is about 1.3% (Table 1). At eclogite facies (Monviso and Lanzo massif), the mean magnetite mode is 3.4% for Atg/Liz-serpentinite and between 1 and 1.8% for Atg-serpentinite and Atg/Ol2-serpentinites (Table 1).

The selected SSP are plagioclase/spinel lherzolites. They display $Fe_2O_3^{Tot} (BR)$ contents varying from 8.31 to 8.87 wt% and a $Fe^{3+}/Fe_{Total} (BR)$ ratio varying from 0.13 to 0.33 (Table 1). Their magnetite mode is less than 1%. According to Andreani et al. (2013), the $Fe^{3+}/Fe_{Total} (BR)$ ratio and magnetite mode of those rocks highly dependent on the local serpentinization degree of the rocks. Thus the observed variations in $Fe^{3+}/Fe_{Total} (BR)$ ratios and magnetite modes in these samples are most likely linked to

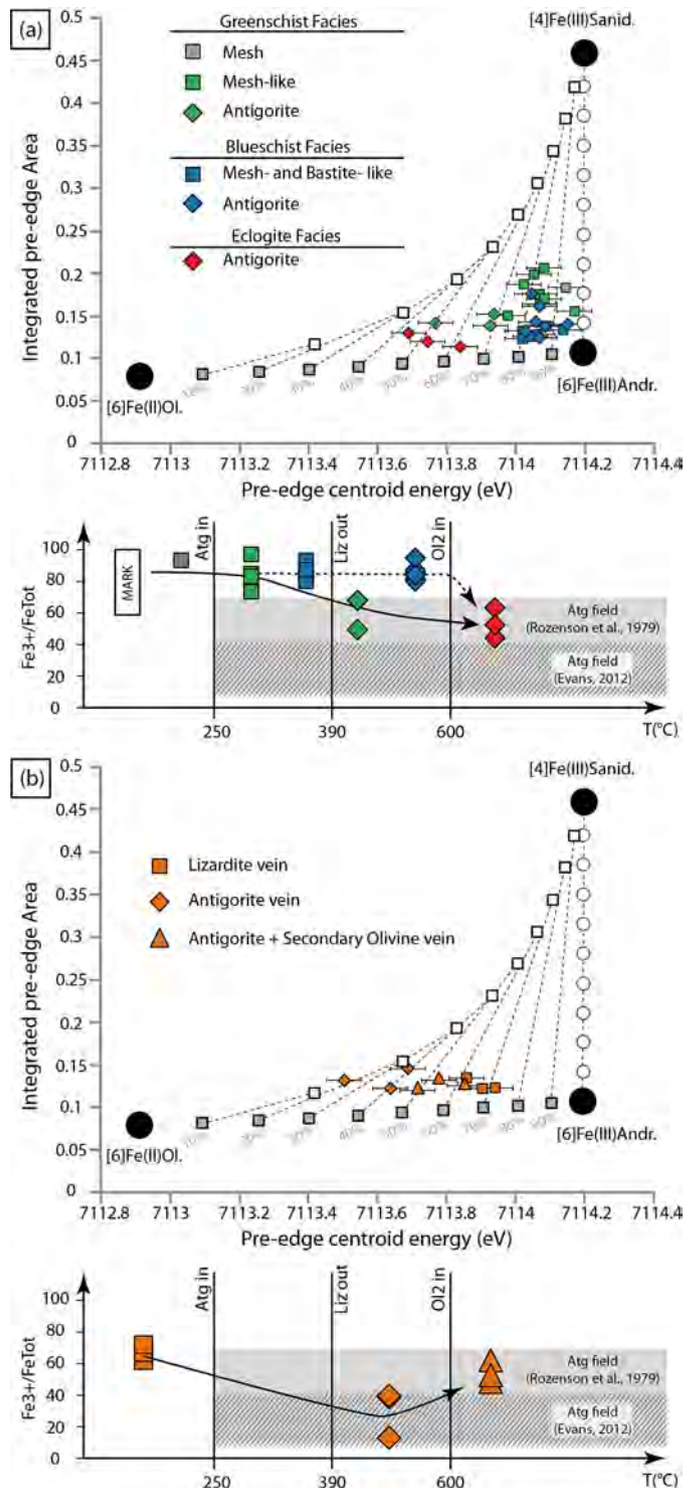


Fig. 5. Calibration grids modified after Wilke et al. (2001) and quantification of the Fe^{3+}/Fe_{Total} ratio in serpentinite. The pre-edge centroid energy error is 0.05 eV (Wilke et al., 2001). (a) Evolution of serpentinite pre-edge parameters (Integrated pre-edge Area Vs Pre-edge centroid energy) and corresponding evolution of the Fe^{3+}/Fe_{Total} ratio in serpentinites with prograde metamorphism. Two evolutions are displayed: the black arrow correspond to low Fe samples and the dashed arrow, to high Fe samples. MARK values are from Andreani et al. (2013). (b) Evolution of serpentinite pre-edge parameters and corresponding evolution of the Fe^{3+}/Fe_{Total} ratio in serpentinite veins from SSP. (See also Rozenson et al. (1979), Evans et al. (2012).)

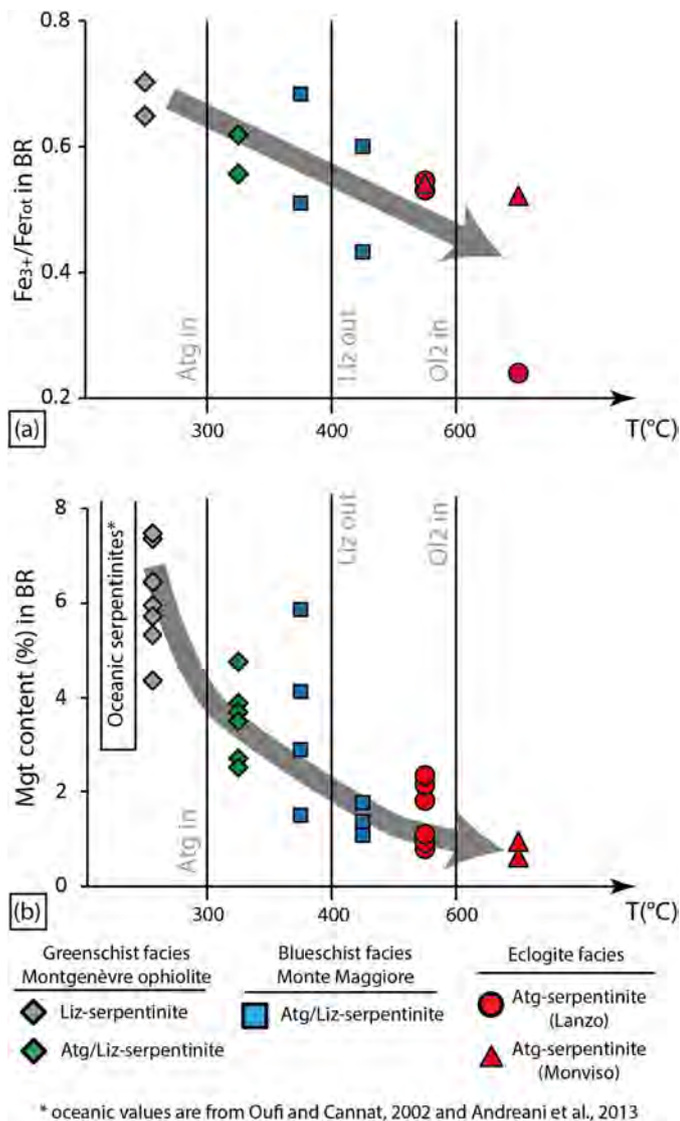


Fig. 6. Diagram presenting the evolution of (a) $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}(\text{BR})$ and (b) modal magnetite versus temperature. Sample temperature calibration is based on serpentine phase transitions drawn from thermodynamical data (Evans, 2004; and references therein), field observations (Schwartz et al., 2013) and experimental data (Bromiley and Pawley, 2003). The progressive increase of temperature is first marked by the growth of antigorite at 300 °C and then by the disappearance of lizardite at 400 °C (Evans, 2004; Schwartz et al., 2013) and finally by the occurrence of secondary olivine at 600 °C (Bromiley and Pawley, 2003).

the variable serpentinization degrees of the rocks, rather than the phase change of serpentine along prograde metamorphism.

4.2. In situ electron microprobe analysis

In situ measurements have been performed on 10 of the 13 serpentinites and on 5 of the 6 SSP samples. The microprobe analyses of serpentine minerals and secondary olivines are reported in Appendix A.

At the Montgenèvre ophiolite, the oceanic mesh and bastite textures of Liz-serpentinite display FeO contents of 2.4 to 3.5 wt%. In the Atg/Liz-serpentinites, the FeO contents of mesh- and bastite-like textures (Fig. 2c) range from 2.7 to 4.4 wt%, and the FeO contents of antigorite veins (Fig. 2c) vary from 3.6 to 4.4 wt%. In the Monte Maggiore ophiolite, samples MM8 and MM15 display heterogeneity in FeO: FeO contents vary respectively from 1.7 to 2.5 wt% and 2.8 to 4.4 wt%. In the Atg/Liz-serpentinites of Monviso ophiolite, the FeO content of mesh-like relicts ranges from 1.2 to

3 wt% and antigorite FeO contents vary from 0.8 to 3.5 wt%. In the Atg-serpentinites of Monviso and Lanzo ophiolites, the FeO content of antigorite (Fig. 3d) ranges from 3–5 wt%. In the same ophiolites, the FeO content of antigorite from Atg/Ol2-serpentinites (Fig. 3e) ranges from 1.8–4.8 wt%, while the composition of the secondary olivine is highly variable: in the Lanzo ophiolite, it varies from 12.4 to 15.4 wt% and in the Monviso ophiolite, the secondary olivine rims (FeO = 5.9–7.1 wt%) are depleted in FeO relative to the core (FeO = 10.6–11.4 wt%).

In the SSP, the FeO composition of serpentine veins is highly variable and depends on vein width or mineralogical site (olivine or pyroxene). The lizardite FeO content veins vary from 1.8 to 8.25 wt%. The FeO contents of antigorite veins ranges from 3.1 to 5.1 wt%. The secondary olivine FeO contents vary from 9.7 wt% to 11.5 wt%.

4.3. In situ XANES and pre-edge micro-analysis

The phase change from lizardite to antigorite is accompanied by a shift of the pre-edge peak centroid energy to lower values and a decrease of its integrated area (Fig. 4 and Fig. 5a) which corresponds to a decrease in serpentine $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ from 0.93 ± 0.06 at oceanic facies to 0.44 ± 0.06 at eclogite facies.

At the Montgenèvre ophiolite, in Atg/Liz-serpentinites (BCh6 and BCh9 samples), similar $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratios (0.97 ± 0.06 and 0.74 ± 0.06) are observed in mesh-like textures relative to oceanic textures of Liz-serpentinites from the same massif (ICh02, $\text{Fe}^{3+}/\text{Fe}_{\text{Total}} = 0.93 \pm 0.06$) and from modern mid-oceanic ridge ($\text{Fe}^{3+}/\text{Fe}_{\text{Total}} = 1-0.6$; Andreani et al., 2013). Antigorite veins in these samples have lower $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratios ranging from 0.69 ± 0.05 to 0.50 ± 0.06 relative to mesh textures. At the Monte Maggiore (MM8 sample), the mesh-like textures and antigorites display similar $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratios ranging respectively from 0.93 ± 0.06 to 0.80 ± 0.06 and from 0.95 ± 0.06 to 0.80 ± 0.06 . These values are close to oceanic mesh textures. At eclogite facies (Monviso and Lanzo ophiolites), the antigorite $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratios vary from 0.63 ± 0.05 to 0.44 ± 0.06 .

In the SSP, the lizardite veins display lower $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratios (0.72 ± 0.06 to 0.62 ± 0.05) relative to oceanic mesh textures from serpentinites (Fig. 5b). As observed previously, antigorite crystallization at the expense of lizardite is marked by a decrease in $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ (0.40 ± 0.05 to 0.13 ± 0.10). The crystallization of secondary olivine on prograde antigorite is marked by a new increase of $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ (0.62 ± 0.05 to 0.49 ± 0.05).

5. Discussion

5.1. The transition lizardite to antigorite in serpentinites

Numerous studies have shown that most of the major elements, in particular iron, are immobile during the serpentinization processes at the oceanic ridge (Andreani et al., 2013; Godard et al., 2008; Mével, 2003; Paulick et al., 2006) and during prograde metamorphism in subduction zones (Debret et al., 2013a, 2013b; Garrido et al., 2005; López Sánchez-Vizcaíno et al., 2005). In agreement with these previous results, no significant change of Fe content is observed in our serpentinites samples during prograde metamorphism (Table 1). This suggests that the fluid released from serpentinites during prograde metamorphism does not carry significant amount of Fe. Thus, a direct transfer of Fe from the slab to the mantle wedge cannot be considered as a mechanism for increasing the $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ of the mantle wedge peridotites and those of arc magmas. Nevertheless, modification of the $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ of the mantle wedge peridotites can result from the circulation of oxidized (H_2O , CO_2 , SO_2 , ...) or reduced (H_2 , CH_4 , H_2S , ...) fluids

and the production of such fluids will be a function of redox reactions occurring in the slab during prograde metamorphism, which should be reflected in the $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ of the prograde mineral assemblages.

Serpentinites from the studied ophiolites were formed close to the oceanic floor (Angiboust et al., 2011; Caby, 1995; Debret et al., 2013a). In such context, serpentinites are expected to be formed in a fluid dominated system and to have high magnetite and Fe^{3+} contents (Andreani et al., 2013; Frost et al., 2013; Oufi et al., 2002). The Montgenèvre ophiolite preserves poorly metamorphosed Liz-serpentinites which have similar magnetite content and $\text{Fe}^{3+}/\text{Fe}_{\text{Total(BR)}}$ ratio relative to oceanic serpentinites from modern settings (Fig. 6). Furthermore, in this ophiolite the mesh and mesh-like textures display similar $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratio to oceanic serpentine from modern settings (Fig. 5). This suggests that the studied serpentinites started with high $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratios prior to subduction that were close to those displayed by modern abyssal serpentinites.

During the first stage of subduction, from greenschist to eclogite facies, the transition from lizardite to antigorite is first marked by the formation of Atg/Liz-serpentine and then of Atg-serpentine. During this phase transition, the $\text{Fe}^{3+}/\text{Fe}_{\text{Total(BR)}}$ ratio decreases from 0.65–0.7 in Liz-serpentinites to 0.53–0.54 in Atg-serpentinites (Fig. 6a). The selected samples correspond to peridotites that have been largely serpentinized during the oceanic stage (distal continental margin or mid-ocean ridge environment; Debret et al., 2013a, 2013b; Hattori and Guillot, 2007; Lafay et al., 2013; Schwartz et al., 2013) and then evolved without any interaction with external fluids during subduction (Debret et al., 2013b), in contrast to serpentinites from the Queyras accretionary prism, which evolved in close proximity to metasediments (Schwartz et al., 2013) and experienced substantial fluid contamination as evidenced from As, Sb or Cs enrichments in antigorite (Lafay et al., 2013). Hence the variations in $\text{Fe}^{3+}/\text{Fe}_{\text{Total(BR)}}$ observed in our samples are unlikely to reflect either contamination processes operating during subduction or variable serpentinization degrees of the initial oceanic peridotite. However, these results show that the reaction lizardite to antigorite is a redox reaction accompanied by a reduction of about 25% of the $\text{Fe}_2\text{O}_3^{\text{Tot(BR)}}$ (Fig. 6a). Among the minerals involved during this phase transition, magnetite and serpentine minerals are the main iron carriers and they are both affected by reduction processes. Indeed, during subduction, the growth of antigorite at the expense of lizardite is associated with a decrease of modal magnetite (Fig. 6b) due to a progressive dissolution of the magnetite grains at mesh rims (Fig. 2c). This decrease is more rapid in Cpx- and Sp-poor samples (i.e. Fe-poor samples), than in Cpx- and Sp-rich samples (i.e. Fe-rich samples, Table 1) because Cpx and Sp are respectively transformed into Mgt2 + Atg and Mgt2 + Cr-Sp + Atg assemblages (Fig. 3a–d). Nevertheless, in pure Atg-serpentinites, irrespective of the primary mineral mode, the oceanic lizardite, mantle clinopyroxene or spinel are absent, and the magnetite modal abundance drops down to 1%. Hence, magnetite has been progressively consumed and may even act as a reactant during the prograde redox reaction lizardite to antigorite. This observation also suggests that magnetite is metastable in Atg-serpentine at eclogite facies.

The decrease in modal magnetite content is inversely correlated to the Fe content of serpentine (Fig. 7), which reflects the fact that the Fe lost during magnetite dissolution is transferred to serpentine. Thus, there is no room for Fe addition by fluids to the rock during the transition from lizardite to antigorite in our samples (Frost, 1991; O'Hanley and Dyar, 1993). This is in good agreement with our previous geochemical study attesting that the alpine ophiolites have evolved in relative closed system during subduction (Debret et al., 2013b). It also confirms the immobile character of Fe during serpentinization at the metric scale.

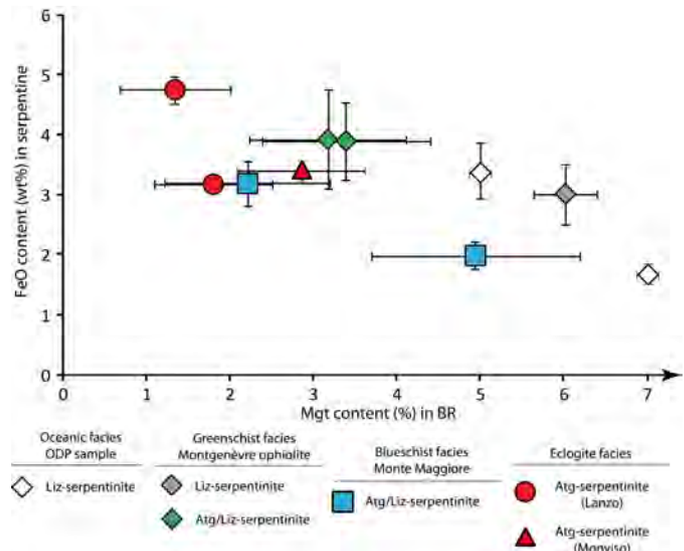
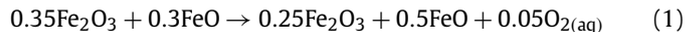


Fig. 7. Diagram of the mean values of FeO content in serpentine versus modal magnetite in bulk rock (BR). The error bars correspond to the standard deviation of the mean values. The data indicate these parameters are inversely correlated.

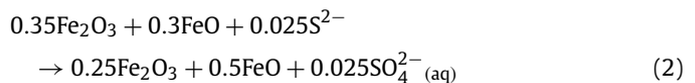
Fig. 5a shows that $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratio in serpentine minerals decreases non-linearly with prograde metamorphism and this tendency varies with $\text{Fe}_2\text{O}_3^{\text{Tot(BR)}}$ content of the rock during first reaction stages. At the Montgenèvre ophiolite, in Fe-poor Atg/Liz-serpentinities, the $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ ratio decreases from 0.74–0.97 in the mesh-like texture to 0.50–0.69 in the antigorite. However, at the Monte Maggiore locality, in Fe-rich Atg/Liz-serpentinities, this ratio is constant during the recrystallization of mesh-like texture (0.80–0.93) into antigorite (0.80–0.95), and remains close to oceanic serpentinites. At eclogite facies, in Atg- and Atg/Ol2-serpentinities, the antigorite $\text{Fe}^{3+}/\text{Fe}_{\text{Total}}$ (0.44–0.63) is lower irrespective of the $\text{Fe}_2\text{O}_3^{\text{Tot(BR)}}$ content of the bulk rock (Table 1). Thus, the redox reaction associated with the whole lizardite \rightarrow antigorite transition finally represents a reduction of $\sim 50\%$ of the serpentine Fe.

The reduction of Fe in the bulk rock is the result of a decrease of the amount of magnetite and a reduction of the Fe in serpentine. However, this decrease does not constitute the full redox reaction taking place during the lizardite/antigorite transition. Indeed, considering a decrease of $\text{Fe}^{3+}/\text{Fe}_{\text{Total(BR)}}$ ratio from 0.7 to 0.5 during the lizardite/antigorite transition (Table 1), the iron reduction can be expressed as:



According to this reaction, the reduction of iron liberates oxygen that could oxidize reduced oceanic species.

In intra-oceanic serpentinites, sulfides occur as disseminated micrometric grains in mesh or bastite textures (Alt and Shanks, 2003; Delacour et al., 2008a). During subduction, S is removed from serpentinites during the prograde transition lizardite to antigorite suggesting that sulfides are dissolved and then released in a fluid phase (Debret et al., 2014). Considering the reduction of iron expressed in Eq. (1), a more complete reaction involving the oxidation of sulfides could be written as:



Based on the iron content of serpentinites, Eq. (2) should lead to a decrease of about 4000 ppm in sulfide concentration in the bulk rock during the lizardite to antigorite transition. Such a mobilization is higher than the sulfur decrease of about 500 ppm measured

by Debret et al. (2014). Potentially, other reduced species may also be oxidized which would offset this effect, such as reduced oceanic carbon (Delacour et al., 2008b; Pasini et al., 2013), residual CH₄ or H₂ (e.g. Evans et al., 2012, 2013), and/or the release of oxygen as mentioned in Eq. (1). Furthermore, reaction (1) and (2) are a subset of a continuum of reactions occurring into the slab and other fluid/rock interactions need to be considered in order to better constrain the entire redox budget of the slab.

5.2. The antigorite dehydration into secondary olivine.

In alpine serpentinites, few variations of serpentine Fe redox state have been observed with antigorite breakdown (Tables 1 and 2). The secondary olivine assemblages represent less than ~10% of the rock. Their formation is not sufficient to modify the redox state of the remaining antigorite. On the other hand, in the SSP, the serpentine veins correspond to relatively closed systems where the local dehydration degree can reach ~40%.

In the SSP, the oceanic lizardite veins display lower Fe³⁺/Fe_{Total} ratio than the ones observed in Liz-serpentinites. This is inherited from the lower serpentinization degree of the SSP on the seafloor since experimental and natural results have shown that the magnetite mode and Fe³⁺/Fe_{Total} ratio of the serpentine progressively increase with the serpentinization degree (Andreani et al., 2013; Marcaillou et al., 2011).

In SSP, antigorite veins have a lower Fe³⁺/Fe_{Total} ratio than lizardite veins. This corresponds to a reduction of about 50% of the Fe contained in serpentine. The occurrence of secondary olivine is accompanied with a new increase of the Fe³⁺/Fe_{Total} ratio in antigorite that increases from 0.1–0.4 in prograde antigorite to 0.5–0.6 (Fig. 5b) in the remaining antigorite associated with secondary olivine.

Previous studies realized on the meta-peridotites of Almirez massif have proposed that antigorite dehydration results in the growth of enstatite, olivine and chlorite and the release of H₂O (Trommsdorff et al., 1998). The prograde antigorite breakdown occurs through a series of continuous reactions forming antigorite associated with olivine and chlorite (Chollet et al., 2011; Padrón-Narvarta et al., 2008, 2011, 2013). Even if enstatite and olivine mainly incorporate Fe²⁺, the full redox equilibria of dehydration reactions are poorly constrained because the changes in Fe oxidation state of antigorite and chlorite as well as the mode of magnetite remain unknown. In alpine ophiolites, only the first stage of serpentine dehydration has been observed in the Monviso and Lanzo ophiolites. This episode is accompanied by the quasi-total disappearance of magnetite (Fig. 6b) and by the formation of a Fe³⁺ bearing-antigorite and secondary olivine assemblage (Debret et al., 2013a; Schwartz et al., 2013). Our XANES measurements indicate a redistribution of Fe during the first stages of antigorite breakdown. Indeed, the solubility of Fe³⁺ in olivine is extremely low, and thus it is preferentially partitioned into antigorite during the dehydration reaction. The incorporation of Fe³⁺, as well as Al³⁺, into the antigorite structure could stabilize it to higher temperature and pressure (Bromiley and Pawley, 2003; Egger and Ehmman, 2010; Padrón-Narvarta et al. 2008, 2011, 2013) allowing its crystallization in equilibrium with secondary olivine.

6. Conclusions

During subduction, with the increase of *P*–*T* conditions from greenschist to blueschist facies, the oceanic lizardite and chrysotile assemblages are progressively recrystallized into antigorite. This episode is accompanied by a decrease of Fe³⁺/Fe_{Total(BR)} without changes of Fe₂O₃^{Tot}(BR) attesting for a reduction of Fe in the serpentine. Among the minerals involved during this phase change, magnetite and serpentine are the main iron carriers and

they are both affected by these reduction processes. Magnetite is progressively dissolved during lizardite destabilization, and in Atg-serpentinites, the antigorite has a lower Fe³⁺/Fe_{Total} than its precursor. The reduction of iron in the bulk rock is accompanied by the release of oxygen, which therefore potentially drives to the oxidation and mobilization of reduced phases such as sulfide or carbon into a fluid phase (e.g. H₂O, SO_x or CO₂). These fluids could have a great oxidizing potential, e.g., during mantle wedge metasomatism by SO₂ fluids, 1 mole of S (S²⁺) can oxidize 6 moles of Fe²⁺ during the SO₂ reduction to sulfide (S²⁻). It also suggests that the redox change during the transition lizardite to antigorite controls the formation of a fluid phase and thus the mobility of volatile and/or trace elements.

At greater depth, the nature of the released fluids remains unclear. The first stage of antigorite breakdown is accompanied by a redistribution of iron forming a Fe³⁺-rich antigorite associated with Fe²⁺-olivine while previous studies (e.g. Padrón-Narvarta et al., 2011) have shown that during the full dehydration of the serpentine, the antigorite is replaced by enstatite, olivine and chlorite assemblages. Even if the redox state of Fe in chlorite and the magnetite mode evolution remain unknown, it seems that this episode is accompanied by a reduction of Fe and a massive release of H₂O (Padrón-Narvarta et al., 2011).

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Appendix A. Supplementary material

Supplementary materials (MMC 1 and MMC 2) related to this article can be found online at <http://dx.doi.org/10.1016/j.epsl.2014.05.038>.

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