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Titanium and water-rich metamorphic olivine in high-pressure serpentinites from the Voltri Massif (Ligurian Alps, Italy): evidence for deep subduction of high-field strength and fluid-mobile elements

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Abstract

Titanium and water-rich metamorphic olivine (Fo 86-88) is reported from partially dehydrated serpentinites from the Voltri complex, Ligurian Alps. The rocks are composed of mostly antigorite and olivine in addition to magnetite, chlorite, clinopyroxene and Ticlinohumite. In-situ SIMS data shows that metamorphic olivine has very high and strongly correlated H_2O (up to 0.7 wt.%) and TiO_2 contents (up to 0.85 wt.%). Ti-rich olivine shows colourless to yellow pleochroism. Olivine associated with Ti-clinohumite contains low Ti, suggesting that Ti-rich olivine is not the breakdown product of Ti-clinohumite. FTIR absorption spectra show peaks of serpentine, Ti-clinohumite and OH-related Si vacancies. Combining FTIR and SIMS data we suggest the presence of clustered planar defects or nanoscale exsolutions of Ti-clinohumite in olivine. These defects or exsolutions contain more H_2O ($x\sim0.1$ in the formula $4Mg_2SiO_4\cdot(1-x)Mg(OH,F)_2\cdot xTiO_2$) than Ti-clinohumite in the sample matrix (x=0.34-0.46).

In addition to TiO_2 and H_2O , secondary olivine contains significant Li (2-60 ppm), B (10-20 ppm), F (10-130 ppm) and Zr (0.9-2.1 ppm). It is enriched in ^{11}B ($\delta^{11}B = +17$ to +23%). Our data indicates that secondary olivine may play a significant role in transporting water, high-field strength and fluid-mobile elements into the deeper mantle as well as introduce significant B isotope anomalies. Release of hydrogen from H_2O -rich olivine subducted into the deep mantle may result in strongly reduced mantle domains.

Keywords: metamorphic olivine, clinohumite, serpentinite, high-pressure metamorphism, subduction recycling of elements, Voltri Massif

1. Introduction

Serpentinites play an important role in the cycling of water and fluid-mobile elements in subduction zones (Scambelluri et al. 2004a; John et al. 2011; e.g., Kodolanyi et al. 2012). Hydrated oceanic crust will dehydrate upon subduction and release fluids and fluid-mobile elements to the overlying mantle wedge (Ulmer and Trommsdorff 1995). This fluid causes hydration of the overlying forearc-mantle peridotites, producing more serpentinites at the base of the mantle wedge (Hattori and Guillot, 2003). These forearc-mantle serpentinites have high H₂O contents (up to 13 wt%) and are enriched in fluid-mobile metals, such as As, Sb and Pb (e.g., Hattori and Guillot 2007; Savov et al. 2007). Their downward movement by mantle

flow eventually causes the serpentinites to dehydrate at a depth of ~100 km, releasing water and fluid-mobile elements into the interior of the mantle wedge, where they induce partial melting related to arc magmatism (Ulmer and Trommsdorff 1995). Mantle-wedge serpentinites thus act as a sponge transferring fluid-mobile elements from the fore-arc to subarc mantle (Hattori and Guillot 2003).

Dehydration is accompanied by the generation of secondary, metamorphic olivine. It is generally thought that most fluid-mobile elements will be lost from the dehydrating serpentinite to the mantle wedge during their dehydration. However, nominally anhydrous minerals (NAMs) such as olivine may contain considerable amounts of water in crystal defects in their structure (e.g., Kitamura et al. 1987; Khisina and Wirth 2002; Peslier 2010) and deep subduction of B and Li by olivine and clinohumite was suggested Scambelluri et al. (2004b). Hence, secondary olivine may potentially introduce large amounts of water and fluid-mobile elements to deeper depths, which has important implications for rheology of mantle peridotites, compositions of mantle fluids and partial melts, and deep recycling of elements into the mantle. To evaluate the retention of elements during serpentinite dehydration in the mantle, we examined the compositions of secondary olivine grains in the Erro-Tobbio peridotite from the Voltri Massif, which formed during serpentinite dehydration at depths of about 75 km (Scambelluri et al. 1995).

1.1. Sample description and petrography

Samples were collected from the Erro-Tobbio metaperidotite in the Voltri Massif (Ligurian Alps, Italy) where the rocks are well exposed along the Gorzente River (Piccardo et al. 1988; Scambelluri et al. 1991; 1995). The metaperidotite is a slice of abyssal peridotite that was serpentinised on the pre-Jurassic ocean floor and underwent partial dehydration during its subduction up to pressures of 25 kbar in Cretaceous time (Cimmino et al. 1979; Scambelluri et al. 1991; 1995). This dehydration resulted in the formation of secondary olivine and minor humite minerals, which form discontinuous veins of several centimetres width in serpentinites (Scambelluri et al. 1995).

The sample which is the focus of this paper (*VT8-3*, 44°33.677' N, 8°48.907' E), is a chlorite-bearing serpentinite that shows a prominent foliation defined by the alignment of antigorite and chlorite and bands of secondary olivine and clinohumite (**Fig. 1A-F**). It contains magnetite and acicular diopside as well as rare porphyroblastic augitic clinopyroxene. Olivine (Ol) occurs as large but heavily fractured grains as well as fine-grained domains. They contain abundant inclusions of low-Ti magnetite (Mag), confirming their secondary

origin. Fine-grained needle-like diopside is associated with olivine and clinohumite.

Magnetite is often finely disseminated and associated with clinohumite-rich domains.

Several generations of secondary olivine are recognized based on microtexture and mineral chemistry. The early metamorphic olivine is Ti-poor and forms porphyroblastic discrete grains in an antigorite matrix and less commonly in Ol-Chu-Mag veins. The second generation also forms porphyroblastic discrete grains in the antigorite matrix, but it is Ti-rich (up to 0.8 wt.%). Locally it forms overgrowths around earlier Ti-poor olivine. The third generation olivine is fine-grained Ti-poor olivine forming the bulk of the Ol-Chu-Mag veins, probably the result of dynamic recrystallisation during late-stage deformation. This also resulted in recrystallisation of fine-grained humite group minerals (**Fig. 1C**). In addition, large cm-sized masses of heavily fractured olivine occur, which appear to predate all other generations.

Three different sections were prepared from a single hand specimen: a thick (\sim 100 µm) section suitable for LA-ICP-MS analysis (VT8-3A), a regular polished thin section (VT8-3B) and a double-polished thick (\sim 150 µm) section suitable for FTIR analysis (VT8-3C).

2. Analytical methods

2.1. Electron probe micro analysis (EPMA)

Major and minor elements were measured by electron probe Cameca SX-100 at EMMAC, University of Edinburgh using a dedicated high-precision setup developed for low concentrations of minor elements in olivine. Accelerating voltage was 20 kV with a defocused 5 μm beam. Mg, Si and Fe were measured at 10 nA for 60s, whereas minor elements were measured at 80 nA for 240s (Na, Al, Ca, Cr) or 120s (Mn, Ni). Pure forsterite and fayalite were used to calibrate Si, Mg and Fe, respectively, in addition to wollastonite, Mg spinel, jadeite and pure metals for remaining elements. Accuracy and precision were monitored by repeated analysis of San Carlos olivine from USGS (USNM# 111312/44; Jarosewich et al. 1980), in-house olivine standard DC0212 (see De Hoog et al. 2010, for preparation and details) and an in-house Cr-diopside standard, the latter to monitor elements which are low in olivine (Na, Al, Ca, Cr). Additional analyses were performed with the same setup but with shorter counting times on other minerals (humite group, serpentine, oxides, some olivines).

Additional mineral analyses were done at Carleton University, Ottawa using a CAMEBAX MBX electron probe in the wavelength dispersive mode. Counting times were 15–20 s per element, except for Ni (40 s). A 15 kV accelerating voltage and a 20 nA beam current were

applied. The calibration used orthoclase (Si), wollastonite (Ca), synthetic spinel (Al), synthetic Cr_2O_3 (Cr), forsterite (Mg), synthetic $MnTiO_3$ (Mn and Ti), albite (Na), fayalite (Fe in silicates), metallic Ni (Ni), and synthetic Fe_2O_3 (Fe in oxides). Results of both EPMA laboratories were in excellent agreement.

2.2. Secondary ion mass spectrometry (SIMS)

Water contents and selected trace elements were measured by SIMS Cameca 4f at the Edinburgh Ion Microprobe Facility (EIMF), University of Edinburgh. Polished samples were gold coated and kept under high vacuum ($<5\times10^{-8}$ mbar) for at least 48 hrs prior to analyses. We used a 5 nA primary beam of negative ¹⁶O ions accelerated to 14.5 kV. Every spot was pre-sputtered for 3 minutes with 30 µm rastered beam, then spots were analysed with a 20µm beam using a small field aperture to avoid stray H ions from the beam periphery entering the mass spectrometer. Depth of the analysis pits was <2 µm. This setup allowed us to reach a background of ~50 ppm H₂O in San Carlos olivine, which value was subtracted from all subsequent water analyses. Collected isotopes are ¹H, ⁷Li, ¹¹B, ¹⁹F, ²³Na, ²⁶Mg, ²⁷Al, ³⁰Si, ³⁹K, ⁴⁰Ca, ⁴¹K, ⁴³Ca, ⁴⁴Ca, ⁴⁵Sc, ⁴⁶Ti, ⁴⁷Ti, ⁴⁸Ti, ⁴⁹Ti, ⁵¹V, ⁵²Cr, ⁵³Cr, ⁵⁵Mn, ⁸⁸Sr, ⁸⁹Y, ⁹⁰Zr using an energy window of 50±25 V. Raw count rates were corrected for molecular interferences such as MgO⁺ (mass 40-42), MgOH⁺ (mass 41-43), SiO⁺ (mass 44-46), SiOH⁺ (mass 45-47), MgMg⁺ (mass 48-52) and MgSi⁺ (52-56). Concentrations of trace elements were calculated against GSE1-G (Jochum and Nohl 2008) using Si as an internal standard. Water concentrations of olivine and humite-group minerals were calibrated using a set of in-house basaltic glass standards ranging in concentration from 0.32-4.63 wt.% H₂O (Hall 1998; Shishkina et al. 2010). The use of basalt standards may introduce a bias in measured olivine and humite-group mineral water contents, but the measured values for the latter are very close to those calculated from stoichiometry. This suggests that matrix-induced fraction for olivine is likely small. The presence of water-bearing inclusions was monitored using the ¹H/³⁰Si signal, but no such inclusions were detected.

Boron isotope compositions were measured using a Cameca 1270 Secondary Ion Mass Spectrometry (SIMS) at EIMF, University of Edinburgh. A primary beam of $^{16}O_2$ with a current of 5 nA was used with a spot size of ca. 15×20 μ m. Secondary ions were extracted at a constant 10 kV voltage and collected by a single electron multiplier measuring ^{10}B and ^{11}B alternately for 4 and 2s, respectively, for a total of 900s per analysis. The area for the analysis was pre-sputtered for about 3 min, which included ion beam centring time. No energy filtering of secondary ions was applied. A small image field (50 μ m) and field aperture (20

μm) ensured maximum intensity whilst minimising the effect of secondary ion beam aberrations. Mass resolution ($\Delta m/m$) of ca. 1200 was used to avoid interferences from 10 B 1 H on 11 B and 9 Be 1 H on 10 B peaks.

Instrumental mass fractionation was corrected using GSD1-G basalt ($^{11}B/^{10}B$ ratio of 4.0849 based on a $\delta^{11}B$ value of +10.2 ± 0.5% reported by Jochum et al., 2008), GOR128-G komatiite (+13.6 ± 0.2%; (Rosner and Meixner 2004) and StHs6/80-G dacite (-4.4 ± 0.3%; (Rosner et al. 2008). As small matrix-dependent fractionations were observed between these glasses (-2.6% for GOR128-G komatiite and -1.6% for StHs6/80-G relative to GSD1-G), we used komatiite GOR128-G as the primary standard, as it is closest in composition to olivine. Boron isotope compositions are presented as permil notation relative to SRM951 using the following equation: $\delta^{11}B_{sample} = 1000 \times (R_{sample}/R_{STD} \times R_{STD}/R_{LSVEC}^* - 1)$ where R is the measured $^{11}B/^{10}B$ ratio and R* is the true $^{11}B/^{10}B$ ratio. The SRM951 $^{11}B/^{10}B$ ratio of 4.0437 was taken from Catanzaro et al. (1970).

2.3. Fourier transform infrared spectroscopy (FTIR)

FTIR was used to determine the structural position of hydroxyl in olivine on a doubly-polished section of 150 μ m thickness (VT8-3C). The section was kept in an oven at 120 °C for more than 1 day before the measurement. A Nicolet 6700 instrument with a Continuum IR Microscope was used at the Tectonophysics Laboratory at SEES in Seoul National University. The water content within a single crystal was measured using an IR beam size of 30×30 μ m. Nitrogen gas was flushed during the FTIR measurement to avoid measuring the moisture in the atmosphere. The measurements were carried out using an unpolarised light source, a KBr beam-splitter and an MCT detector. A series of 128 scans was averaged for each spectrum with a resolution of 4 cm⁻¹.

2.4. Laser ablation ICP-MS

Selected minerals were measured in thick section (VT8-3A) for their trace-element contents by laser ablation ICP-MS in the Department of Earth Sciences at the University of Oxford using a custom-built New Wave 193nm ArF Excimer laser system attached to a Thermo-Finnegan Element2 magnetic sector ICP-MS in low resolution mode. Ablation took place in He atmosphere and spot size was 90-180 µm. NIST SRM-612 was used for calibration (Pearce et al. 1997). Further details including identification and correction of molecular interferences for olivine can be found in De Hoog et al. (2010).

2.5. Electron backscatter diffraction (EBSD)

EBSD analysis was performed using the HKL Channel 5 electron backscatter diffraction (EBSD) system on a Philips XL30CP SEM at EMMAC, University of Edinburgh, operating at 20 kV accelerating voltage with a beam current of ca. 1 nA and a 20 mm working distance with a tilt angle of 70°. For each point of the map the diffraction pattern was collected and solved by the HKL software for the orientation. Prior to EBSD analysis the sample (VT8-3A) was polished with colloidal silica in order to remove damage to the surface caused by prior mechanical polishing.

3. Mineral compositions

3.1. Olivine

3.1.1. Major and trace elements (EPMA)

Two types of secondary olivine are identified based on their chemistry (**Table 1**): low Ti olivine (2-250 ppm Ti) and Ti-rich olivine (500-2600 ppm Ti). The latter shows pleochroism from colourless to light yellow in plane-polarised transmitted light and usually occurs as small porphyroclasts within the antigorite matrix (**Figs. 1D,F**). Both types show identical EBSD patterns to olivine. Occasionally high-Ti olivine forms rims around low-Ti grains (e.g., VT8-3C grain C, Table 2). Low-Ti olivine occurs as small porphyroclasts in the antigorite matrix, as large porphyroclastic olivine and as recrystallised fine-grained olivine which constitutes most of the Ol-Chu-Mag veins but also mantles Ti-poor and Ti-rich olivine porphyroclasts.

Apart from their variable TiO₂ content all olivines have very similar major element compositions spanning a narrow range of Fo contents (86.5 to 87.6%), which is slightly lower than typical mantle olivine. NiO contents (0.23-0.39 wt.%) are comparable to typical mantle olivine but NiO and Fo show no correlation. MnO contents are relatively high (0.26-0.43 wt.%), which is in agreement with olivine being of secondary (metamorphic) origin. This is also supported by abundant inclusions of magnetite, which is a product of earlier serpentinisation. CaO contents are low but variable (0.001-0.029). Cr₂O₃ contents range from <0.001 to 0.070 w.t% but are on average higher in Ti-rich olivine than typical mantle olivine (0.017 and 0.005, respectively; De Hoog et al. 2010).

Ti-rich olivine shows a distinct Si deficiency in its calculated formula, as #Si is 0.995-1.005 in Ti-poor olivine but down to 0.970 in Ti-rich olivine (based on 4O, **Table 1**). This

may be explained by the presence of other separate phases as exsolution or inclusions, as indicated by FTIR spectra (Ti-clinohumite defects or exsolutions; Section 3.1.4).

3.1.2. Trace elements (SIMS and LA-ICP-MS)

Selected olivines were analysed for trace elements by SIMS (**Table 2**). Many trace elements are enriched in Ti-rich olivines compared to Ti-poor olivines, e.g., F (15-51 vs. 5-35 ppm), Al (22-51vs. 2-3 ppm), Sc (67-117 vs. 18-40 ppm), V (9-11 vs. 2-3 ppm), Cr (0.038-0.044 vs. 0.004-0.007 wt.%), Y (0.7-0.9 vs 0.2 ppm) and Zr (0.9-2.1 vs 0.2-0.3 ppm). In contrast, Li and B contents show little difference (Li: 31-44 ppm in Ti-rich olivine and 6-17 ppm in regular olivine; B: 14-16 vs. 9-13 ppm).

Results from LA-ICP-MS (**Table 3**) are very similar to those measured by SIMS except Sc, V and Cr, which show lower values by 20-50%. The discrepancies of these elements may be caused by complex interference corrections for these elements during SIMS analysis (see Section 2.2). Nevertheless, differences between V, Cr and Sc of Ti-poor and Ti-rich olivine are similar to that measured by SIMS. The contents of Co, Cu, Zn, As, Sb and Sn are similar in both olivine types.

3.1.3 Boron isotopes

Selected olivines (including the two porphyroclasts and one recrystallised olivine in **Fig. 1F**) were analysed in-situ for boron isotopic composition by SIMS (**Table 5**). δ^{11} B values range from +17 to +23‰, and are on average the same for yellow Ti-rich olivine and colourless Ti-poor olivine. Variations up to 5‰ were observed within individual grains despite having constant major-element compositions, which likely reflects variable fluid composition during the growth of secondary olivine.

3.1.4. Water content and structural position

Water contents of olivine measured with SIMS show a large variation from 100 ppm in Tipoor olivine to 6400 ppm in Tiprich olivine. The values are high compared to typical mantle-derived olivine (<100 ppm; Peslier 2010). The high contents of water are not due to inclusions of hydrous minerals, such as serpentine and clinohumite, as these inclusions were avoided by using a small beam (<20 μ m) and the shallow depth of penetration (<2 μ m) during SIMS analysis. The H₂O contents show a strong positive correlation with TiO₂ contents (**Fig. 2**).

The structural position of OH was characterised using FTIR spectroscopy (**Fig. 3**). Five FTIR spectra obtained from optically clear areas of two grains (grains A and B) in a thick (150 µm) section show sharp absorption peaks at 3687, 3612, 3564, 3522 and 3473 cm⁻¹ and a

broad peak around 3400-3415 cm⁻¹ (**Fig. 3**). Although the beam was focused to 30 μm, the FTIR collects data through the entire thickness of sample, hence it was unavoidable to measure minor inclusions. The peak at 3687 cm⁻¹ is related to serpentine inclusions (e.g., Kitamura et al. 1987; Matsyuk and Langer 2004; Khisina et al. 2008; Jung, 2009), whereas the peak at 3612 cm⁻¹ is most likely related to OH incorporated as Mg₂[]OH₄, where [] represents an Si vacancy (Berry et al. 2005). The peaks at 3564 and 3522 cm⁻¹ may be associated with Ti-clinohumite-type point defects composition of MgTi[]O₂(OH)₂ (Berry et al. 2005), although a peak position of at 3572 cm⁻¹ was reported by the authors. Alternatively, the FTIR spectra indicate the presence of Ti-clinohumite exsolutions or clustered planar defects (Matsyuk and Langer 2004), consistent with the broad peak around 3400-3415 cm⁻¹ as well as the spectrum of pure Ti-clinohumite (Berry et al. 2005). However, Hermann et al. (2007) reported a peak at 3564 cm⁻¹ (not 3572 cm⁻¹) for olivine that showed high densities of clustered planar Ti-clinohumite defects, which were imaged by TEM. It appears therefore difficult distinguish between defects and exsolutions based on FTIR data without additional TEM imaging. As Voltri olivines appeared homogeneous during EBSD and back-scatter electron imaging at high magnifications, any exsolution of phases, if present, must be nanometre-scale.

The 3473 cm⁻¹ peak could not be related to any known OH stretching modes in olivine or any known hydrous minerals or inclusions. A peak at 3476-80 cm⁻¹ has been attributed to Si vacancy substitution of hydrogen into olivine in previous studies (Lemaire et al. 2004; Bali et al. 2008; Kovacs et al. 2010). However, the peak reported by previous researchers is slightly offset from the one reported here and consistently very small relative to other peaks in contrast to the intense 3473 cm⁻¹ peak observed in our samples.

3.2. Ti-clinohumite and related minerals

The humite mineral group consists of four members with the general formula $n\text{Mg}_2\text{SiO}_4\cdot\text{Mg}(F,\text{OH})_2$ where n=1 for norbergite (Nrb), 2 for chondrodite (Chn), 3 for humite (Hu) and 4 for clinohumite (Chu) (Jones et al. 1969). Most natural humite-group minerals contain substantial amounts of TiO₂ (Evans and Trommsdorff 1983). Incorporation of TiO₂ up to x=0.5 is accommodated by the exchange reaction TiO₂=Mg(OH)₂, leading to the general formula $n\text{Mg}_2\text{SiO}_4\cdot(1-x)\text{Mg}(\text{OH},F)_2\cdot x\text{TiO}_2$. Note that the IMA reserves the names clinohumite for the F endmember and hydroxylclinohumite for the OH endmember (http://pubsites.uws.edu.au/ ima-cnmnc/); for simplicity the name clinohumite will be used throughout this article.

Humite-group minerals with their reddish brown colour were easily distinguishable under a microscope and were identified by their M/Si atomic ratio where M is the sum of all cations including Ti. The M/Si ratios are 2.25 for Chu, 2.33 for Hu and 2.5 for Chn. Ti-clinohumite is the most common humite-group mineral in sample VT8-3 (Figs. 1A, 1B, 1C, 1E). It occurs in contact with magnetite, antigorite, and Ti-poor olivine. Small grains of chondrodite and humite were observed during the composition analysis with EPMA (Table 1, Fig. 2). They occur together with other humite group minerals and serpentine; electron back-scattered images suggest most individual grains are homogeneous in compositions. Rare lamellar intergrowths of clinohumite and chondrodite are found in contact with Ti-poor olivine (Fig. 1E). There is no textural evidence suggesting the replacement of olivine by humite group minerals and vice versa. None of humite-group minerals is in contact with Ti-rich olivine. The presence of humite together with other humite group minerals is rarely reported. Apparent humite compositions can be obtained from fine-grained intergrowths of clinohumite and chondrodite (Wirth et al. 2001), but whereas lamellar intergrowths were occasionally observed in other grains (Fig. 1E), the humite grains were optically homogeneous.

All humite-group minerals in our sample have high TiO_2 contents (Chu: 3.6-4.9 wt.%; Hu: 5.4-6.0 wt.%; Chn: 7.2-7.8 wt%), which equals to a range of x =0.35-0.46 in the general formula $nMg_2SiO_4\cdot(1-x)Mg(OH,F)_2\cdot xTiO_2$. They contain appreciable MnO (0.6-0.8 wt.%), small amounts of Cr_2O_3 (0.01-0.18 wt.%), and little Al_2O_3 (<0.07 wt.%) and CaO (<0.01 wt.%). Different humite group minerals have similar major element compositions, including Mg# (85-87 for all). This is common for co-existing humite-group minerals, as shown in the Buell Park kimberlite, Arizona (Aoki et al. 1976), in serpentinised dunites from Isua, Greenland (Dymek et al. 1988) and ultrahigh-pressure garnet pyroxenites from Dabie Shan, China (Hermann et al. 2007). The values of Mg# are similar to that of co-existing olivine. This is also observed in serpentinites elsewhere (Dymek et al. 1988, and references therein). Water contents based on stoichiometry (cf. Jones et al. 1969) are 1.5-1.8 wt.% in clinohumite, 2.1-2.2 wt.% in humite and 3.0 wt% in chondrodite. Water contents measured by SIMS give similar values (1.5-1.8 wt.%, 2.0 wt.% and 2.6-3.0 wt.%, respectively). Fluorine contents are low (0.03-0.11 wt.%).

Trace element analysis revealed that humite-group minerals are enriched in volatiles and high field strength elements (HFSE) (**Table 4**). Lithium contents (1-17 ppm) are lower, whereas B contents (21-43 ppm) are higher than co-existing olivine (4-48 and 9-22 ppm, respectively). Zirconium contents are rather high (12-126 ppm), whereas V (2-24 ppm), Sr (1-4 ppm) and Y (0.3-2 ppm) contents are similar to those observed in olivine. Apart from

slightly higher F contents of chondrodite, trace elements show little difference between various humite minerals.

3.3. Serpentine

Serpentine (antigorite) has a narrow range of Mg# (94.5-94.8) and contains 1.3-2.6 wt.% Al_2O_3 and 0.1-0.2 wt.% Cr_2O_3 . Some MnO (0.01-0.05 wt.%) and NiO (0.07-0.17) were also detected. Compared to co-existing olivine it has low contents of Ti (8-13 ppm), Li (<0.15 ppm) and B (6-9 ppm) whereas F contents (15-47 ppm) fall within the same range.

3.4. Clinopyroxene

Two types of clinopyroxene are present. The first type is coarse-grained deformed augite, which exhibits well-developed cleavages. Its Mg# falls between 90.6-91.8 and it contains 4.9-7.2 wt.% Al₂O₃, 1.0-1.2 wt.% Cr₂O₃, 0.3-0.5 wt.% TiO₂, and 0.3-0.4 wt.% Na₂O. The high concentrations of Ti, Al and Cr suggest this type of clinopyroxene to be of relic mantle origin and survived serpentinisation.

In contrast, the second type has an acicular habit and is often associated with clinohumiterich domains. It is nearly pure diopside, with high Mg # (94.3-96.4) and low TiO_2 (<0.01-0.4 wt.%), Al_2O_3 (0.03-0.66 wt.%), Cr_2O_3 (0.02-0.3 wt.%) and Na_2O (0.1-0.3 wt.%) and is therefore considered to be of metamorphic origin.

3.5. Oxides

Oxides (var. magnetite) are closely associated with clinohumite-rich veins, but also occur within large patches of secondary olivine. They contain significant Cr_2O_3 (3.5-6.1 wt.%) but low TiO_2 (<0.5 wt.%) and MnO (<0.1 wt.%). In addition they contain ca. 0.07 wt.% V_2O_3 , 0.10 wt.% NiO and <0.01 wt% ZnO. No chromite was observed in the sample.

4. Interpretation and discussion

4.1. Origin of high Ti and H₂O in olivine

The good positive correlation between H_2O and TiO_2 contents of secondary metamorphic olivine from Voltri (**Fig. 2**) suggests that these elements are incorporated by a common mechanism. FTIR spectra of olivine indeed indicated the presence of Ti-clinohumite planar defects or exsolutions (Section 3.1.4). However, Voltri olivines have excess water compared to pure Ti-humite defects as well as Ti-clinohumite grains in the sample matrix (**Fig. 2**), which suggests that the planar defects or exsolutions must have a composition that differs

from the ideal formula of MgTi[]O₂(OH)₂. FTIR spectra do show an abundance of serpentine micro-inclusions in addition to clinohumite defects or exsolutions, which could explain the increased H₂O signal, but such inclusions were effectively avoided during SIMS analyses. We therefore infer that olivine contains clinohumite defects or exsolutions with considerably less Ti than the ideal formula MgTi[]O₂(OH)₂. Based on the relative amounts of TiO₂ and H₂O in olivine, we derive a value of ca. 0.1 for x in the formula (1-x)Mg(OH)₂·xTiO₂. This value is significantly less than that of humite-group minerals in the sample matrix, which have a value of x ranging from 0.34 to 0.46.

Very high solubility of H₂O in olivine under water-saturated conditions was indicated by experimental data at high pressures (e.g., Kohlstedt et al. 1996 report up to 0.5 wt% H₂O at 13 GPa). On the other hand, water contents of mantle olivines are typically <100 ppm, with an average of 43 ppm calculated based on water contents of mantle-derived melts (Peslier 2010). The maximum solubility of TiO₂ in olivine is ca. 1.3 wt.% in anhydrous systems at temperatures in excess of 1200°C (Hermann et al. 2005) but are strongly controlled by coexisting Ti-bearing phases. Solubility is ca. 0.2 wt.% in natural water-bearing systems containing ilmenite at 1200°C (Ulmer et al. 1998). Olivines from mantle xenoliths have maximum TiO₂ contents of ca. 0.06 wt.% (De Hoog et al. 2010). Our results indicate that the presence of clinohumite defects or nano-exsolutions can accommodate H₂O and TiO₂ in olivine well above those of typical mantle values. Hermann et al. (2007) pointed out that the humite-group minerals and olivine form a polysomatic series with decreasing hydrous interlayers relative to olivine layers going from norbergite (n=1) to clinohumite (n=4) to olivine $(n=\infty)$. If TiO₂-rich olivines from Voltri are expressed using the general formula for humite group minerals, $nMg_2SiO_4\cdot(1-x)Mg(OH)_2\cdot xTiO_2$, where n=4 for clinohumite and ∞ for pure olivine, the value for *n* reaches a minimum value of 16 for Voltri olivine.

High TiO₂ contents of olivine have been used to infer an ultradeep origin of rocks, as typical mantle olivine has Ti concentrations less than 350 ppm or ca. 0.06 wt% (Hermann et al. 2005; De Hoog et al. 2010). For example, Ti contents of over 0.6 wt.% and rutile/ilmenite exsolutions in olivines from garnet peridotites from Alpe Arami were used to infer an origin in excess of 300 km (Dobrzhinetskaya et al. 1996), although the pressure dependence of Ti solubility was later questioned by experiments by Hermann et al. (2005). In addition, olivine with 0.19 wt.% TiO₂ was reported by Hermann et al. (2007) as a breakdown product of Ti-chondrodite under high pressure in garnet-pyroxenites from Dabie Shan UHP belt. The Ti contents of olivine in our sample (up to 0.9 wt.%) indicate that olivine containing even higher TiO₂ contents can be formed at relatively moderate pressures and temperatures (600°C and 25

kbar; Scambelluri et al. 1995). Thus, even such high TiO₂ contents of olivine cannot be used to demonstrate deep mantle conditions.

4.2. Formation conditions of Ti-rich hydrous olivine and humite-group minerals

The peak P-T conditions, 600° C and 25 kbar, of the metaperidotite were estimated from boudins of metagabbros (Scambelluri et al. 1995). This agrees with the assemblage Atg + Fo + Ti-Chu + Di of the metaperidotite. The stability field is bracketed by the antigorite-out reaction (Atg + Di = Fo + Tr + H₂O) and the brucite-out reaction (Atg + Brc = Fo + H₂O). Since antigorite breaks-down to olivine and orthopyroxene at high pressures, the upper stability limit is defined by Atg = Fo + En + H₂O. Some minor secondary Opx was reported in these rocks by Scambelluri et al. (1995).

The upper stability limit of Ti-clinohumite is provided by the breakdown reaction Ti-Chu = Fo + Gk (MgTiO₃) + H₂O, which is close to the antigorite stability limit in F-free assemblages (Engi and Lindsley 1980; Trommsdorff and Evans 1980; Weiss 1997). The absence in our samples of geikilite (MgTiO₃), a breakdown product of clinohumite commonly reported from other ultramafic localities (Trommsdorff and Evans 1980; Okay 1994; Sanchez-Vizcaino et al. 2005; e.g., Hermann et al. 2007) limits the temperature of the formation to below ca. 675°C (Engi and Lindsley 1980).

A distinctive feature of our sample is the occurrence of abundant Ti-chondrodite (ca. 25% of all humite-group minerals). Unfortunately, the relative stability of different humite-group minerals is not well constrained. Chondrodite may form preferentially in Mg-rich rocks (Mg#>95; Engi and Lindsley 1980; Dymek et al. 1988), but our sample has relatively low Mg# of 86-87. Engi and Lindsley (1980) suggested that chondrodite is stable at higher pressures than clinohumite. Thus, re-equilibration during exhumation would form clinohumite at the expense of chondrodite, but microtextures in our sample show no clear evidence of humite-group minerals replacing each other. Weiss (1977) showed that temperature may play an important role, as clinohumite breaks down to more Ti-rich norbergite and olivine (+ fluid) with increasing temperature over a narrow 20°C interval. Taking into account the proximity of Voltri P-T conditions to the start of the F-free clinohumite breakdown reaction we infer that clinohumite started to break down to chondrodite and olivine according the reaction Ti-OH-Chu = Ti-OH-Chn + 2 Ol, but that the temperature (ca. 600°C; Scambelluri et al. 1995) was not high enough for further breakdown to ilmenite and fluid.

The conditions and timing of the formation of Ti-rich olivine relative to formation of Ticlinohumite are not clear. Previously reported occurrences of high-TiO₂ olivine are in chlorite harzburgites from the Almirez Massif in Spain (Trommsdorff et al. 1998; Ruiz Cruz et al. 1999; Puga et al. 1999; Garrido et al. 2005; Sanchez-Vizcaino et al. 2005) and in garnet pyroxenites from Dabie Shan UHP belt, China (Hermann et al. 2007). In the Almirez Massif, brown olivine is intergrown with Ti-clinohumite and contains numerous Ti-rich solid inclusions. It is interpreted to be the breakdown product of clinohumite, which still partially survived as lamellae and patchy inclusions in olivine. An origin of Ti-rich olivine as a breakdown product of Ti-clinohumite and Ti-chondrodite was also proposed for garnetpyroxenites from Dabie Shan (Hermann et al. 2007). In contrast, Ti-rich yellow olivine documented here from Voltri appears to have a different origin, as it occurs away from Ticlinohumite-rich domains, contains no lamellae or patchy inclusions of clinohumite and lacks inclusions of Mg-rich ilmenite (a breakdown product of Ti-clinohumite). In addition, olivine closely associated with humite-group minerals is Ti-poor. Ti-rich olivine forms overgrowths on Ti-poor olivine, but humite-group minerals do not occur in contact with Ti-rich olivine. In Voltri, clinohumite typically occurs in veins hosted by massive antigorite, which formed during prograde Alpine subduction and deformation (Scambelluri et al. 1995; Scambelluri and Rampone 1999). Hence, clinohumite formed together with metamorphic olivine during dehydration near peak pressures of 18-25 kbar. Since peridotites originally contain low TiO₂, variable influx of fluids from external rocks during prograde metamorphism may have resulted in formation of Ti-poor olivine, Ti-rich olivine and Ti-clinohumite in the same sample.

4.3. Implications for transfer of water, HFSE and fluid-mobile elements beyond arc depths

Our results indicate that secondary olivine which formed during subduction dehydration of serpentinite may contain considerable amounts of H₂O and TiO₂ and may therefore play an important role in the redistribution of these elements in subduction zones. Fluorine in olivine behaves similar to H₂O as it shows a similar positive correlation with TiO₂ (**Fig. 4**). As F contents fall in-between typical mantle olivine values (0.5-14 ppm; Guggino et al. 2007) and humite-group minerals, we suspect that F enters humite defects in olivine. Similarly, the contents of Zr and Nb positively correlate with TiO₂ contents, which probably reflects HFSE partitioning into Ti-clinohumite defects in olivine. As both F and HFSE are considerably enriched in secondary olivine, anomalously high contents of these elements can be introduced into the mantle. Likewise, Garrido et al. (2005) reported chloritised harzburgite from the Almirez Massif acting as a sink for HFSE during dehydration of serpentinite. The main

carriers of HSFE in their study were Cr-magnetite and ilmenite exsolutions in brown olivine containing Ti-clinohumite intergrowths (Ruiz Cruz et al. 1999; Garrido et al. 2005). Although metamorphic conditions in Voltri were not high enough to reach Ti-clinohumite breakdown (Scambelluri et al. 1995), our observations are similar to those of Garrido et al. (2005), hence HFSE enrichment in the secondary olivine may be important in understanding the distribution of these elements, as they have been shown to be mobile in subducting slabs at depths below volcanic arcs (Gao et al. 2007). Upon deeper subduction, increasing temperatures lead to Ti-clinohumite defect breakdown and formation of Mg-rich ilmenite (Puga et al. 1999; Risold et al. 2001; Hermann et al. 2007), which will retain HFSE in the rock. Also, inclusions and humite-type defects likely remain stable in olivine under high pressures (Kitamura et al. 1987; Wirth et al. 2001; Matsyuk and Langer 2004).

In contrast to HFSE and F, Li shows no correlation with TiO₂ content in olivine. It is enriched in both Ti-poor and Ti-rich olivine compared to antigorite and clinohumite, as well as typical upper mantle olivine (**Fig. 4**). As antigorite has low Li contents and its breakdown can liberate only little Li, this requires an alternative source of Li in olivine during serpentinite dehydration. Possible sources are the breakdown of relic mantle Cpx and external fluids. Relic clinopyroxene in VT8-3 contains only 5-9 ppm Li, considerable lower than Ti-rich olivine, and therefore an unlikely source of Li found in olivine. External fluids may be derived from gabbros and mafic volcanic rocks during eclogitisation (Scambelluri and Rampone 1999; Scambelluri et al. 2004a).

The behaviour of B during serpentine dehydration is of particular interest as its behaviour in arc settings is thought to be dominated by serpentinite (Scambelluri and Tonarini 2012). Similar to Li, B is enriched in Ti-poor as well as Ti-rich olivine compared to antigorite. Its abundance in olivine is therefore not related to the presence of Ti-clinohumite exsolutions but to a distinct solution mechanism such as B(OH) = SiO (Sykes et al. 1994). The B contents are considerably higher than typical mantle values (**Fig. 4**), in agreement with Scambelluri et al. (2004b) who observed high residual B contents in dehydrated serpentinites of ca. 10 ppm. Boron isotope compositions show high δ^{11} B (+17 to +23‰, **Table 5**) compared to typical mantle values of ca. -3‰ (Hart et al. 1999). The observed values for the metamorphic olivine are virtually identical to the bulk-rock data for the high-pressure veins from which our sample is derived (+20 to +24‰, Scambelluri and Tonarini, 2012). Scambelluri and Tonarini (2012) consider the high δ^{11} B values due to a supra-subduction zone setting of the serpentinites, although similar values (+5 to +40‰) have been reported for sea-floor altered serpentinites

(Vils et al. 2009). Scambelluri and Tonarini (2012) reported little fractionation of B isotopes with increasing metamorphic grade of serpentinites and little B loss during transition from lizardite to antigorite. Our olivine data confirms retention of B and limited B isotopic fractionation during the metamorphic transformation of serpentine to olivine, so that even after complete dehydration the residue will retain its B isotope signature. Therefore subduction of secondary olivine may introduce significant B isotope anomalies into the deep mantle.

Finally, the deep subduction of serpentinites and their dehydration contribute to redox changes in the Earth's mantle. Serpentinisation, the hydration of peridotites, is accompanied by the oxidation of Fe to form magnetite and the release of reduced gases, such as H₂ and CH₄. (Evans 2008). In reverse dehydration reaction, H₂O-rich olivine is formed and may be subducted to even deeper mantle. When such olivine comes in contact with hot dry peridotites it may readily release H₂, as diffusion rate for hydrogen is much higher than those of metals in olivine (Ingrin and Skogby 2000; Hwang et al. 2008). In addition, at the reducing lower mantle conditions, H₂ is immiscible with hydrous fluids (Bali et al. 2013). Thus fluid released from subducted hydrous olivine may lead to reduced mantle domains, and, together with carbon brought down by sinking slabs, it may induce favourable conditions for the formation of diamond.

4.4. Summary

Secondary (metamorphic) olivine in dehydrating serpentinite from the high-pressure Voltri Complex is enriched in H_2O (up to 0.7 wt.%) and TiO_2 (up to 0.86 wt.%); the two show a positive correlation in their concentrations. Based on FTIR data we suggest that these elements are hosted by clustered planar Ti-clinohumite defects or nano-scale exsolutions in olivine.

Secondary metamorphic olivine is significantly enriched in Li (2-60 ppm), B (10-20 ppm), F (10-130 ppm) and Zr (0.9-2.1 ppm) compared to mantle olivine. It shows high $\delta^{11}B$ values ranging from +17 to +23‰, which are similar to those of bulk serpentinites. The dehydration of serpentine to form olivine did not release much of B and F. Our data indicates that secondary olivine may play a significant role in transporting HFSE, water, fluid-mobile elements and ^{11}B to the deep mantle. Release of hydrogen from subducted hydrous olivine in the deep mantle may result in strongly reducing mantle domains.

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FIGURE CAPTIONS

- Fig. 1. Petrography of Voltri serpentinite (section VT8-3B). Scale bars in bottom-left corners of each photograph are 100 µm wide except for A (1mm). Rectangles indicate locations of detailed photographs. Note that B, E and F are 90° rotated. The area numbers indicated in the detailed photographs match those of the grain numbers in the compositional tables (Tables 1-5). Mineral abbreviations from Whitney and Evans (2010). A. Transmitted light scanned image of thin section. B. Clinohumite in contact with Ti-poor olivine porphyroclast in antigorite matrix. Some of the orange grains are chondrodite and humite (plane-polarised light). C. Coarse-grained Ti-poor olivine and clinohumite recrystallised to fine-grained crystal aggregates. The small orange grains are dominantly clinohumite with about one third chondrodite. Recrystallised olivine has low TiO₂ contents (plane-polarised light). D. Titanium-rich olivine in antigorite matrix exhibiting yellow (left) to colourless (right) pleochroism (plane-polarised light). Also note the presence of exsolved and deformed clinopyroxene interpreted to be relic mantle pyroxene. E. Chondrodite lamellae (dark brown) in clinohumite (light brown) in plane-polarised light (left) and under crossed polarisers exhibiting anomalous interference colours (right). F. Porphyroblastic Ti-rich olivine (yellowish) associated with colourless Ti-poor porphyroblastic olivine (plane-polarised light).
- **Fig. 2.** TiO_2 vs H_2O for hydrous minerals from Voltri serpentinite. Olivine data measured by SIMS, serpentine and humite-group minerals by EPMA with water contents calculated from stoichiometry. Humite-group minerals (orange symbols) are subdivided into clinohumite, humite and chondrodite based on their chemistry. Dashed lines indicate predicted compositional variation of humite minerals following $TiO_2 = Mg(OH)_2$ exchange up to the maximum of X_{Ti} =0.5. Inset shows detailed view of olivine compositions. Note the strong correlation between TiO_2 and H_2O for olivines, which follows a trend different than predicted for pure humite defects $MgTi[]O_2(OH)_2$.
- **Fig. 3.** FTIR spectra of two secondary olivine grains (A and B) from sample VT8-3. Drawn for comparison are spectra for Ti-clinohumite and olivine equilibrated with rutile (Ti-Chu vacancy) and periclase (Si vacancy) after Berry et al. (2005).

- **Fig. 4.** TiO₂ vs fluid-mobile elements F, H₂O, Li and B for olivine (Ol), antigorite (Atg) and humite-group minerals (Chu). Fields for mantle olivine values compiled from Kent and Rossman (2002), Guggino et al. (2007), De Hoog et al. (2010) and Peslier (2010).
- **Fig. 5.** Boron isotopes vs. B concentrations for Voltri secondary olivine (this study) and Voltri HP serpentinites (Scambelluri and Tonarini 2012). Fields for subducted oceanic crust and hydrated mantle wedge after Scambelluri and Tonarini (2012), serpentinised oceanic mantle (which extend up to +40‰) and depleted mantle after Vils et al. (2009) and references therein.

TABLES

Table 1. Major element data of selected minerals by EPMA

Mineral	Ol1	Ol2	Chu	Hu	Chn	Atg	Mag	Cpx1	Cpx2
Section	VT8-3B	VT8-3B	VT8-3B	VT8-3B	VT8-3B	VT8-3B	VT8-3B	VT8-3B	VT8-3C
Grain	C-2	D-5	C-1	A-2	C-4	A-7	C-1	D-1	F-3
Lab	UoE	UoE	UoE	UoE	UoE	OU	UoE	OU	OU
SiO ₂	40.03	39.26	36.53	34.92	32.69	41.54	0.17	51.26	54.46
TiO_2	<d.1.< td=""><td>0.861</td><td>4.72</td><td>5.39</td><td>7.76</td><td>0.01</td><td>0.41</td><td>0.40</td><td>0.01</td></d.1.<>	0.861	4.72	5.39	7.76	0.01	0.41	0.40	0.01
Al_2O_3	<d.1.< td=""><td><d.l.< td=""><td><d.l.< td=""><td><d.1.< td=""><td>0.01</td><td>2.56</td><td>0.05</td><td>5.46</td><td>0.05</td></d.1.<></td></d.l.<></td></d.l.<></td></d.1.<>	<d.l.< td=""><td><d.l.< td=""><td><d.1.< td=""><td>0.01</td><td>2.56</td><td>0.05</td><td>5.46</td><td>0.05</td></d.1.<></td></d.l.<></td></d.l.<>	<d.l.< td=""><td><d.1.< td=""><td>0.01</td><td>2.56</td><td>0.05</td><td>5.46</td><td>0.05</td></d.1.<></td></d.l.<>	<d.1.< td=""><td>0.01</td><td>2.56</td><td>0.05</td><td>5.46</td><td>0.05</td></d.1.<>	0.01	2.56	0.05	5.46	0.05
Cr_2O_3	<d.1.< td=""><td>0.045</td><td>0.08</td><td>0.04</td><td>0.12</td><td>0.11</td><td>3.48</td><td>1.12</td><td>0.02</td></d.1.<>	0.045	0.08	0.04	0.12	0.11	3.48	1.12	0.02
Fe_2O_3							63.93		
FeO	12.60	12.13	13.02	12.17	12.71	3.66	30.63	2.72	1.91
MnO	0.347	0.294	0.44	0.39	0.44	0.05	0.09	0.05	0.10
NiO	0.315	0.296	0.20	0.14	0.14	0.07	0.24	0.02	0.00
MgO	46.82	46.71	44.13	44.78	42.64	37.00	0.43	15.26	17.57
CaO	0.007	0.005	0.00	0.01	0.01	0.02	<d.1.< td=""><td>22.91</td><td>25.58</td></d.1.<>	22.91	25.58
Na ₂ O	<d.1.< td=""><td><d.l.< td=""><td>n.a.</td><td>n.a.</td><td>n.a.</td><td><d.l.< td=""><td>n.a.</td><td>0.31</td><td><d.1.< td=""></d.1.<></td></d.l.<></td></d.l.<></td></d.1.<>	<d.l.< td=""><td>n.a.</td><td>n.a.</td><td>n.a.</td><td><d.l.< td=""><td>n.a.</td><td>0.31</td><td><d.1.< td=""></d.1.<></td></d.l.<></td></d.l.<>	n.a.	n.a.	n.a.	<d.l.< td=""><td>n.a.</td><td>0.31</td><td><d.1.< td=""></d.1.<></td></d.l.<>	n.a.	0.31	<d.1.< td=""></d.1.<>
H ₂ O ^b	0.036	0.680	1.49	2.27	2.97	12.52	n.a.	n.a.	n.a.
total	100.19	100.47	100.62	100.11	99.52	97.54	99.31	99.51	99.70
#(O,OH,F)	4	4	18	14	10	9	4	6	6
#Si	0.994	0.967	4.047	3.000	2.019	1.991	0.006	1.875	1.989
#Ti	0.001	0.016	0.393	0.349	0.361	0.000	0.012	0.011	0.000
#Al	0.000	0.000	0.000	0.000	0.001	0.144	0.002	0.235	0.002
#Fe ³⁺							1.855		
#Cr	0.000	0.001	0.007	0.003	0.006	0.004	0.106	0.032	0.001
#Fe ²⁺	0.262	0.250	1.206	0.874	0.657	0.147	0.988	0.083	0.058
#Mn	0.007	0.006	0.041	0.028	0.023	0.002	0.003	0.002	0.003
#Ni	0.006	0.006	0.018	0.010	0.007	0.003	0.008	0.001	0.000
#Mg	1.733	1.715	7.288	5.736	3.926	2.643	0.025	0.832	0.956
#Ca	0.000	0.000	0.000	0.001	0.001	0.001	0.000	0.898	1.000
#Na	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.022	0.000
#cations	3.003	2.961	13.000	10.000	7.000	4.935	3.000	3.991	4.010
#OH	0.006	0.112	1.099	1.301	1.222	4.000			
3.6.11	0.010	0.0=4	0.070	0.0.10	0.0==	0.01-	0.05/	0.005	0.044
Mg#	0.869	0.873	0.858	0.868	0.857	0.947	0.024	0.909	0.943
M/Si	2.02	2.06	2.21	2.33	2.47				
$x(TiO_2)$		h xx o	0.46	0.35	0.38			10 F P 2+x	3.6/01

 $x(11O_2)$ 0.40 0.55 0.50 $x(11O_2)$ by stoichiometry, except olivine by SIMS; Mg# = Mg/(Mg+Fe²⁺); M/Si = (#cations -#Si)/#Si; $x(TiO_2)$ = value of x in general formula $nMg_2SiO_4.(1-x)Mg(OH)_2.xTiO_2$; <d.l. = less than detection limit; n.a. = not analysed. Atg = antigorite, Ol = olivine, Chu = clinohumite, Hu = humite, Chn = chondrodite, Mag = magnetite, Cpx = clinopyroxene, lab codes: UoE = University of Edinburgh, OU = Ottawa University

Table 2. Trace element and water contents of olivine by SIMS

Section	Grain	H_2O	Li	В	\mathbf{F}	Al	Ca	Sc	Ti	\mathbf{V}	Cr	MnO	Sr	Y	Zr
VT8-3A	A-1	1785	17	14	29	317	75	n.a.	1086	n.a.	n.a.	0.36	1.1	0.53	3.2
VT8-3A	A-2	2090	22	16	34	463	98	n.a.	771	n.a.	n.a.	0.38	1.5	0.81	0.7
VT8-3A	B-2	338	47	10	9	55	27	n.a.	211	n.a.	n.a.	0.33	0.5	0.22	0.9
VT8-3A	C-1	684	5	17	16	2	39	n.a.	223	n.a.	n.a.	0.35	3.9	0.38	2.5
VT8-3A	E-1	817	5	11	16	1	26	n.a.	17	n.a.	n.a.	0.28	0.5	0.08	0.5
VT8-3A	F-1	590	34	11	18	32	24	n.a.	491	n.a.	n.a.	0.36	0.3	0.24	0.3
VT8-3B	A-1	655	12	15	12	1	23	34	124	1	33	0.31	2.4	0.55	0.7
VT8-3B	B-1	460	16	10	15	3	<1	40	114	3	43	0.34	0.9	0.24	0.3
VT8-3B	B-2	223	6	11	7	2	5	18	2	2	51	0.33	0.7	0.16	0.3
VT8-3B	C-2	363	17	9	5	2	7	40	80	3	34	0.35	0.7	0.17	0.2
VT8-3B	D-1	5777	31	16	47	51	4	67	2434	10	271	0.36	0.9	0.45	2.1
VT8-3B	D-2	4784	39	16	31	31	8	82	2036	11	265	0.36	0.9	0.36	1.8
VT8-3B	D-3	5460	35	14	51	27	2	72	2558	10	259	0.36	0.7	0.38	1.8
VT8-3B	D-4	1689	44	14	15	22	3	117	535	9	301	0.35	0.8	0.37	0.9
VT8-3B	D-5	6802	17	18	137	10	<1	42	4545	11	199	0.30	0.5	0.43	2.4
VT8-3C	A-1	1124	5	20	11	1	9	15	559	1	47	0.32	0.5	0.36	1.2
VT8-3C	A-2	1017	6	20	24	5	11	15	611	2	51	0.34	0.4	0.28	1.4
VT8-3C	A-3	2279	4	23	22	2	5	14	1417	3	42	0.30	0.6	0.48	2.0
VT8-3C	B1-1c	537	33	12	27	1	9	28	211	4	125	0.30	0.5	0.19	0.3
VT8-3C	B1-2c	891	22	12	27	1	4	27	364	4	110	0.29	0.6	0.19	0.4
VT8-3C	B2-1c	140	13	15	10	1	3	94	14	8	184	0.29	0.5	0.21	0.4
VT8-3C	B5r	5431	7	20	111	2	8	20	4184	8	106	0.32	1.0	0.34	2.0
VT8-3C	B3r	4535	12	14	64	2	3	36	2263	8	123	0.29	0.5	0.22	0.7
VT8-3C	C-1	267	27	12	7	0	6	26	100	4	136	0.31	0.6	0.18	0.3
VT8-3C	I-1	151	27	12	6	1	7	34	9	3	67	0.28	0.6	0.19	0.3

All concentrations in ppm, except MnO in wt.%

Table 3. Trace element contents of olivine and serpentine by LA-ICP-MS

Section	Grain	Li	Al	Ca	Sc	Ti	\mathbf{v}	Cr	Mn	Co	Cu	Zn	As	Sr	Y	Zr	Nb	Sn	Sb
VT8-3A	A-1	18	55	78	29	867	3.8	92	2678	164	0.16	84	< 0.05	2.0	0.51	1.12	0.049	0.52	0.004
VT8-3A	A-2	22	69	62	43	1169	5.7	125	2568	161	0.16	85	< 0.05	1.6	0.42	1.23	0.052	0.54	0.004
VT8-3A	B-1	40	35	47	30	289	4.0	106	2474	158	0.15	84	0.09	0.7	0.30	0.60	0.016	0.58	< 0.003
VT8-3A	C-1	7	6	90	19	189	2.6	21	2594	156	0.26	83	0.21	3.4	0.41	0.66	0.000	0.58	0.011
VT8-3A	E-1	9	<5	166	20	20	1.4	39	2459	152	0.42	83	0.05	1.9	0.30	0.27	0.003	0.51	0.004
VT8-3A	G-1	6	<5	57	22	59	1.5	15	2773	158	0.14	84	0.20	1.5	0.29	0.33	0.002	0.53	0.003
VT8-3A	G-2	3	<5	28	11	10	0.9	8	2502	160	0.89	80	0.30	0.7	0.27	0.19	0.003	0.61	< 0.003
VT8-3A	G-3	3	<5	21	14	30	1.1	9	2378	157	0.11	78	0.21	0.3	0.15	0.10	0.000	0.60	< 0.003
VT8-3A	G-4	12	<5	95	51	7	3.1	37	2376	158	0.18	77	0.15	0.5	0.24	0.08	0.001	0.59	< 0.003
VT8-3A	H-1	7	<5	118	21	80	0.9	14	2613	164	0.22	79	0.14	3.6	0.65	0.81	0.012	0.45	0.004
VT8-3A	H-2	7	<5	46	20	62	0.8	28	2736	158	0.14	83	0.10	1.1	0.43	0.71	0.005	0.56	0.013
VT8-3A	I-1	14	<5	70	22	108	2.7	65	2442	153	0.14	82	< 0.05	1.3	0.26	0.19	0.005	0.54	0.007
VT8-3A	I-2	21	<5	52	35	75	3.9	100	2446	153	0.16	82	< 0.05	1.0	0.26	0.17	0.001	0.58	0.004
VT8-3A	J-1	9	<5	74	17	44	2.1	38	2540	158	0.24	85	0.09	2.3	0.30	0.35	0.001	0.50	< 0.003
VT8-3A	J-2	6	<5	41	18	53	2.1	34	2500	159	0.18	83	0.08	0.4	0.17	0.17	0.003	0.54	0.013

All concentrations in ppm.

Table 4. Trace element and H_2O contents of humite-group minerals and serpentine by SIMS

Section	Grain	H ₂ O wt%	TiO ₂ wt%	Li ppm	B ppm	F ppm	V ppm	Cr ppm	Mn ppm	Sr ppm	Y ppm	Zr ppm
clinohumit	e											
VT8-3B	C-1	1.7	4.2	9	29	472	9	581	5364	1.3	0.4	16
VT8-3B	C-2	1.8	3.9	6	40	335	16	759	6020	1.3	0.6	25
VT8-3B	C-3	2.1	3.6	1	32	416	2	56	5690	0.9	0.5	26
VT8-3B	A-1	1.8	3.6	2	43	751	11	387	3366	1.2	1.2	51
VT8-3B	H-1	1.8	4.2	17	32	725	21	599	4412	0.9	0.5	12
VT8-3C	J-1	1.5	3.8	1	36	675	6	59	4323	0.7	0.3	12
humite												
VT8-3B	A-2	2.0	5.5	5	22	616	19	391	3729	1.5	1.8	13
VT8-3C	I-1	2.0	5.9	14	33	805	12	811	3658	1.4	1.0	25
chondrodi	te											
VT8-3B	C-4	2.9	7.0	7	21	596	7	851	6088	1.3	0.5	20
VT8-3B	C-5	3.0	6.3	12	28	669	5	1237	4751	4.4	0.7	39
VT8-3B	H-2	2.6	7.4	6	29	1129	24	536	4653	0.9	1.0	126
VT8-3C	K-1	3.0	7.0	6	36	1116	12	527	4375	2.2	0.5	30
antigorite												
VT8-3A	E-1	11.7	0.002	0.1	6	17			305	0.3	0.6	0.8
VT8-3B	D-1	18.5	0.001	0.1	10	85	103	2298	301	0.6	0.5	0.7
VT8-3B	A-1	13.3	0.002	0.1	10	335	73	2103	257	0.4	0.5	0.5

Table 5. Boron isotope compositions of olivine by SIMS

Section	Grain	B (ppm)	$\delta^{11}B$ ‰	δ^{11} B
			SRM951	1s ‰
VT8-3B	F-1-1 (Ti-rich porphyroclast)	10.1	+18.7	±1.3
VT8-3B	F-1-2	12.0	+18.9	±1.1
VT8-3B	F-1-3	14.0	+17.4	±1.0
VT8-3B	F-2-1 (Ti-poor recrystallised)	17.3	+22.7	± 0.8
VT8-3B	F-3-1 (Ti-poor porphyroclast)	13.6	+22.0	±1.1
VT8-3B	F-3-2	15.2	+17.3	±1.1
VT8-3B	I-1 (Ti-poor matrix)	10.5	+16.7	±1.3
Average		13.3	+19.1	±2.4

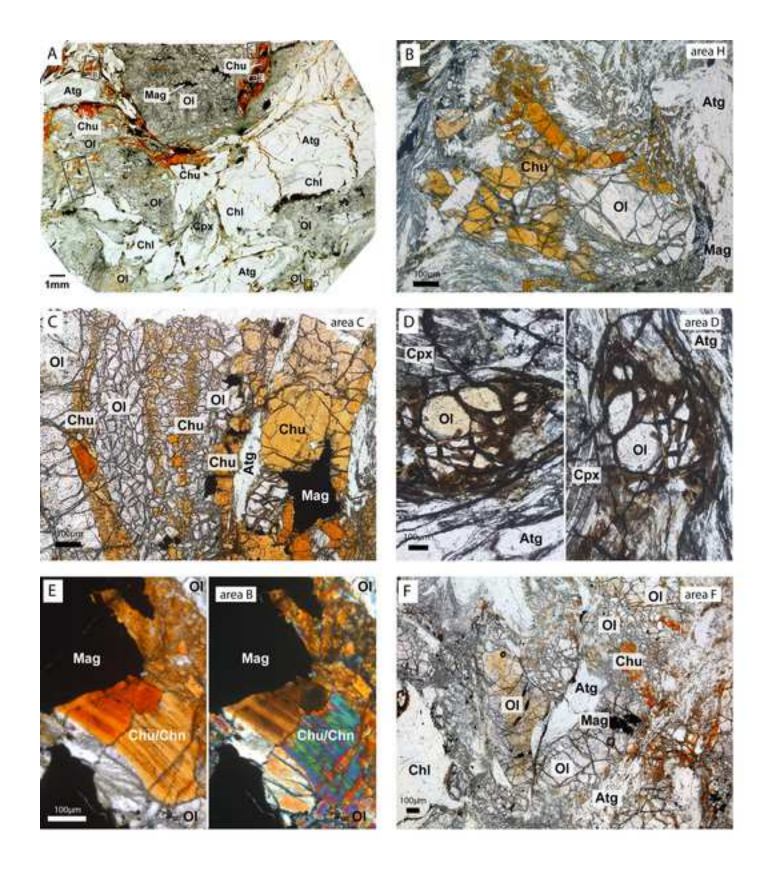


Figure 1. De Hoog et al. Voltri secondary olivine

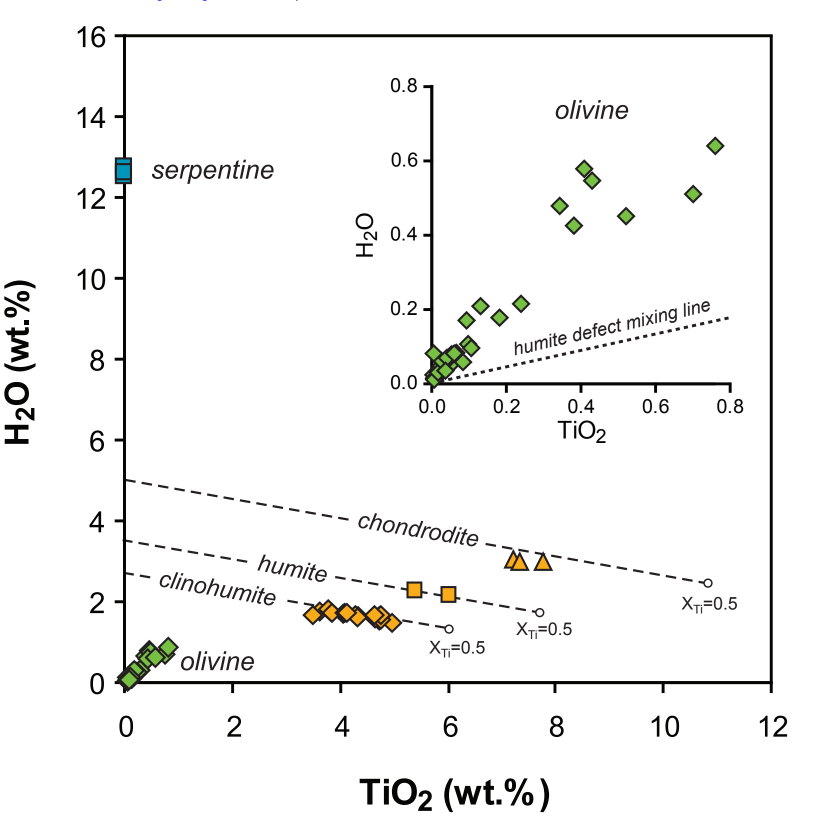


Fig. 2 De Hoog et al. Voltri secondary olivine

Figure 3
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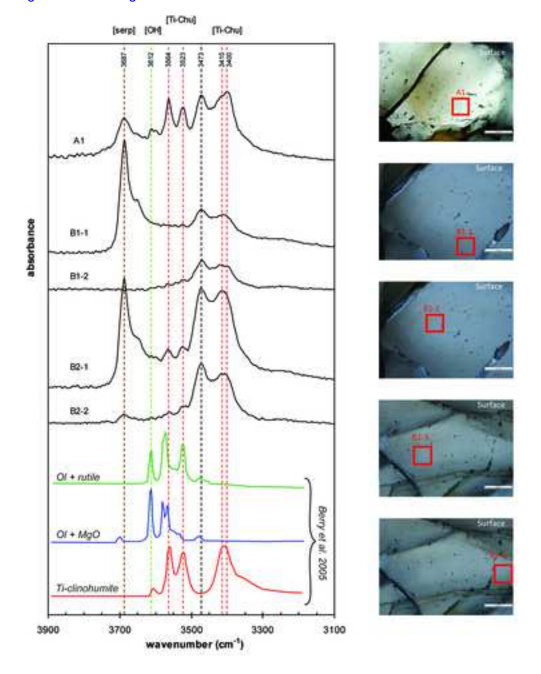


Figure 3 De Hoog et al. Voltri secondary olivine

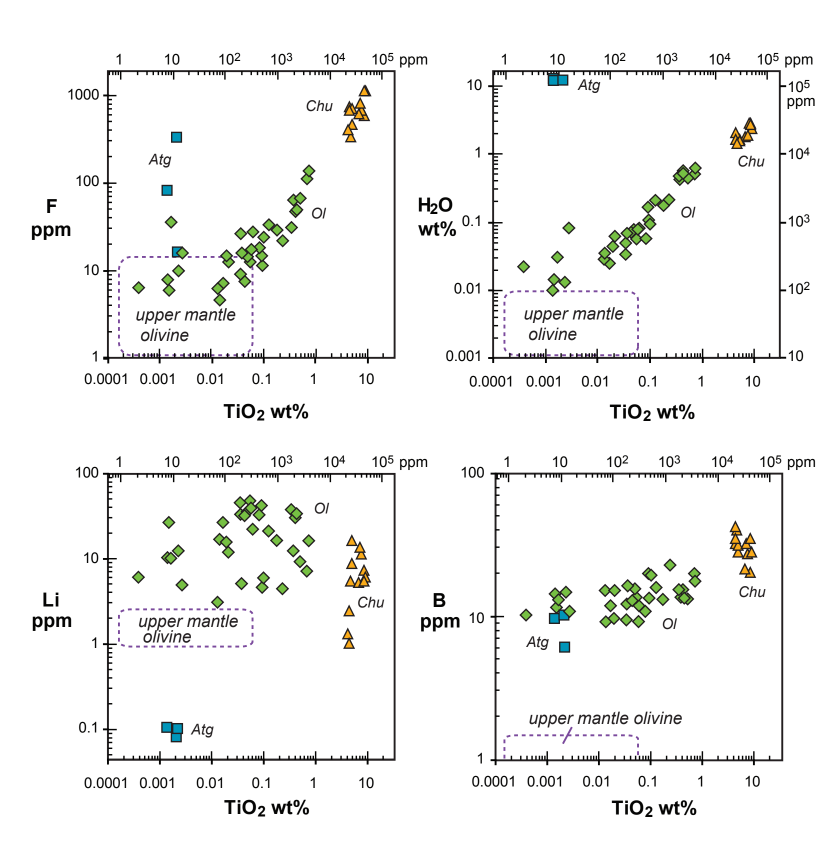


Fig. 4 De Hoog et al. Voltri secondary olivine

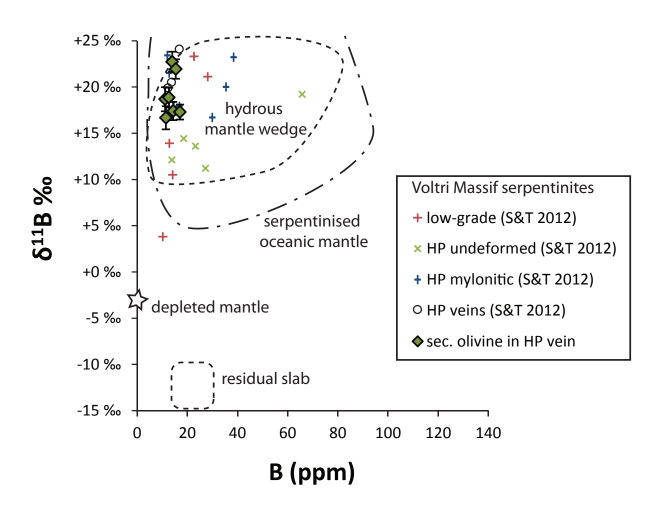


Fig. 5 De Hoog et al. Voltri secondary olivine