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Unraveling the exhumation history of high-pressure ophiolites using magnetite (U-Th-Sm)/He thermochronometry



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ABSTRACT

Magnetite is a ubiquitous oxide in ultramafic and mafic rocks, which is present in a large range of geological and tectonic settings. In the case of high-pressure ultramafic rocks, exhumation timing is commonly constrained by geochronometers in nearby lithologies. The development of the magnetite (U-Th-Sm)/He method (MgHe) has opened new perspectives to refine the exhumation history of such rocks. However no thermal history has been deduced using MgHe data so far. Here we applied MgHe dating to magnetite from an HP-LT alpine ophiolite body (Rocher Blanc) from the Schistes lustrés units (Western Alps, France) where part the P-T-t history is constrained to further refine the thermal history and demonstrate validity of the method. Textural and geochemical characterizations of the magnetite grains reveal two crystallization environments leading to distinct grain morphologies, euhedral and pseudo-euhedral, with different amounts of mineral inclusions (titanite, chlorite and augite-aegirine). Both magnetite types crystallized during the same retrograde path at $T > 250\,^{\circ}\text{C}$, under hydrothermal conditions shown by Si and V oscillatory zoning and low Ti content. MgHe ages derived from euhedral magnetite crystals range from 14.8 ± 2.1 to 20.9 ± 3.0 Ma, between zircon and apatite fission track (ZFT) and (AFT) ages deduced from the literature. Pseudo-euhedral grains display older ages interpreted as resulting from the implantation of He from U-rich neighboring minerals such as titanite. For the first time, MgHe data are inverted along with AFT and ZFT data to refine the thermal history of an ophiolite related to their exhumation. The MgHe data suggest that the Rocher Blanc sample has undergone an exhumation in ductile conditions related to subduction dynamics and underplating processes, followed by an exhumation reflecting denudation by erosion under brittle conditions. This exhumation regime change is found to take place at \sim 20 Ma, as opposed to a less constrained \sim 26 Ma cooling as would be inferred from the ZFT and AFT ages alone. Considering published thermochronometric data along the Western Alps, we propose that the exhumation is controlled by the upward indentation of the Apulian mantle producing regional tilting and associated erosion. With the present Schistes lustrés ophiolite example, we demonstrate that the MgHe method can be used as a low-temperature thermochronometer (closure temperature in the 200-250 °C range) that aids in unraveling the cooling history associated with exhumation of high-pressure ultramafic and mafic rocks at the brittle to ductile transition.

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1. Introduction

Ultramafic and mafic rocks are widely distributed on the Earth's surface and are associated with a large range of geological and geodynamic settings, from oceanic accretion to orogenic zones (e.g., Guillot et al., 2015). In contrast to their abundance, mafic and ultramafic rocks only yield a few minerals suitable for geo-

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thermochronometry such as U-Th-bearing accessory minerals (zircon, titanite or apatite). However, recent advances in the analysis of radiogenic isotopes have extended the application of (U-Th)/He geo-thermochronology to minerals incorporating U and Th at the ppb level such as magnetite (Blackburn et al., 2007; Cooperdock and Stockli, 2016) or spinel (Cooperdock and Stockli, 2018). Magnetite (U-Th)/He geo-thermochronometry, abbreviated MgHe below, opens new perspectives with respect to the use of basic and ultrabasic rocks for dating geodynamic processes.

Indeed, magnetite $(Fe^{2+}Fe_2^{3+}O_4)$ is ubiquitous in these rock types where it can either crystallize from the parent magma or precipitate later in the rock history from hydrothermal fluids at lower temperature. The MgHe method is particularly attractive for dating geodynamic processes involving serpentinite formation, metamorphism or exhumation. Magnetite is produced during serpentinization of the lithospheric mantle at slow-spreading ridges for T > 200 °C (Klein et al., 2013, 2014) and controls thereby hydrogen production (e.g., see Brunet, 2019 for a review). Magnetite in subducted serpentinite can survive temperature conditions up to the antigorite breakdown (Debret et al., 2014; Merkulova et al., 2017). Therefore, in addition to constraining the timing of mafic and ultramafic rocks exhumation, dating of magnetite by (U-Th-Sm)/He geochronometry can also put time constraints on the functioning of fossil hydrothermal systems and on serpentinite metamorphism in subduction-collision zones.

The MgHe method is based on radiogenic ⁴He production and accumulation in the crystal lattice during alpha decay of U, Th, and Sm isotopes (238 U, 235 U, 232 Th and 147 Sm) that lead to stable isotopes (206Pb, 207Pb, 208Pb and 143Nd respectively). The alpha particle is ejected with a high kinetic energy of several MeV and travels a mean distance of \sim 15 μm in the magnetite lattice (Ketcham et al., 2011). A correction for He loss due to alpha ejection (called F_T) needs to be applied which can range from few percent to 20-30% for smaller grains with sizes of \sim 100 μ m (Farley et al., 1996). Otherwise, abrasion of the first \sim 15 µm of the crystals or selection of the inner parts of a large crystal can be done to avoid ejection or implantation from neighboring minerals (Blackburn et al., 2007; Cooperdock and Stockli, 2016). The remaining He atoms in the crystal can diffuse outside of the crystal with a rate of diffusion that is a function of the temperature and crystal volume. Based on experimental He diffusion coefficients obtained on magmatic magnetite by Blackburn et al. (2007), the closure temperature Tc (Dodson, 1973) is as 250 ± 40 °C for a crystal size of 500 μm and a cooling rate of 10 °C/Ma. The Tc for He in magnetite falls in the same range as for the zircon fission track (ZFT) system, which is broadly estimated to be \sim 240 \pm 50 °C (Gallagher et al., 1998). Ketcham (2019) derived a Tc value of 281 °C based on fitting experimental data sets from relatively young zircons (Murakami et al., 2006), and Rahn et al. (2004) showed that radiation damage may lower annealing resistance.

The number of studies dealing with the MgHe method remains limited. MgHe has been applied to volcanic rocks (Blackburn et al., 2007), ore deposits (Fanale and Kulp, 1962) as well as serpentinites and chloritoschists (Cooperdock and Stockli, 2016). These studies have clearly demonstrated the relevance of the MgHe thermochronometer to obtain cooling or crystallization ages. However, MgHe data have never previously been used to infer the thermal history of rocks using inverse modeling, which is the main purpose of collecting thermochronometric data.

Here, the MgHe method is applied to an ophiolite from the Schistes lustrés units (Western Alps, France), for which the cooling history is relatively well constrained, as the mineralogy and P-T history of the area have been characterized (Pognante and Kienast, 1986; Tricart and Schwartz, 2006) and ZFT and AFT data are available (Schwartz et al., 2007). The obtained MgHe data are crit-

ically evaluated in the framework of the ophiolite thermal history constrained by the ZFT and AFT literature data.

2. Geological setting

The studied samples were collected in the internal domain of the western Alps (Fig. 1A) that underlies the boundary between the European and African plates (e.g., Lemoine et al., 1986). This domain (Fig. 1B) derives from a part of the Tethyan ocean (Piedmont zone) and its northwestern European margin (Briançonnais zone). In the study area, the Piedmont zone is composed of the Schistes lustrés units and the Monviso ophiolite, which rest on the internal crystalline massif of Dora Maira (Fig. 1C). These units were juxtaposed during subduction and collision processes in Late Cretaceous to Late Tertiary times. The Piedmont zone represents different levels of a paleo-subduction zone, preserving the low thermal gradient (5-8°C/km) associated with the subduction dynamics (Schwartz et al., 2001; Agard et al., 2002). In this context, the Schistes lustrés units correspond to a fossiliferous sedimentary accretionary wedge metamorphosed under high pressure and low temperature (HP-LT) conditions during the late Cretaceousearly Eocene subduction of the Tethyan ocean (Lafay et al., 2013). This domain is derived from Mesozoic oceanic sediments that were originally composed of marls, clays, and limestones associated with fragments of oceanic lithosphere. These sediments were strongly deformed and metamorphosed during subduction, and they crop out today as foliated and polydeformed calcschists enclosing boudinaged meter- to kilometer-sized Jurassic ophiolites. The P-T conditions increase toward the east, from low-temperature blueschist (LT-BS, Fig. 1C) to high-temperature blueschist facies conditions (HT-BS, Fig. 1C) (Tricart and Schwartz, 2006). The blueschist domain is structurally located above the Monviso eclogitic ophiolite (Fig. 1C). In this massif, the metasedimentary component is small (<20 vol.%) relative to the voluminous oceanic lithosphere (Schwartz et al., 2001).

The P-T evolutions of the Schistes lustrés units are well documented and show clockwise loop paths, keeping a record of the early prograde evolution, and the retrograde metamorphism stages have remained in the field typical of greenschist paragenesis (Fig. 1D), with no temperature increase (Tricart and Schwartz, 2006). For the studied Rocher Blanc ophiolite located in the central part of the Schistes lustrés units (Fig. 1C), the metamorphic peak conditions are estimated at 10 ± 2 kbar and $>350\,^{\circ}\text{C}$ (Tricart and Schwartz, 2006; Schwartz et al., 2013), with laser probe Ar-Ar ages ranging from 62 to 55 Ma (Agard et al., 2002). The final exhumation is constrained by apatite and zircon fission track (AFT-ZFT) analysis (Schwartz et al., 2007) (Fig. 1C), which reveals a diachronous exhumation of the HP-LT metamorphic rocks from east to west during Oligocene to Miocene time. Taking into account the cooling rates from Schwartz et al. (2007), the Rocher Blanc ophiolite is considered to have passed through the ~240-280 °C isotherm at 24.8 \pm 1.6 Ma and the \sim 110 °C isotherm at 12.3 \pm 2.0 Ma (Fig. 1C).

3. Samples and methods

3.1. Sampling and petrologic characterization

Sample RB1 was collected from the Rocher Blanc massif (RB; WGS84 UTM31N, 6°57′08.01″E, 44°39′53.25″N; elevation: 2810 m a.s.l.) that corresponds to a km-scale ophiolitic body composed of mafic and ultramafic rocks embedded in the Schistes lustrés units (Fig. 1C) that record middle-temperature blueschist facies conditions (MT-BS) (Pognante and Kienast, 1986; Tricart and Schwartz, 2006). The high-pressure blueschist paragenesis in mafic rocks

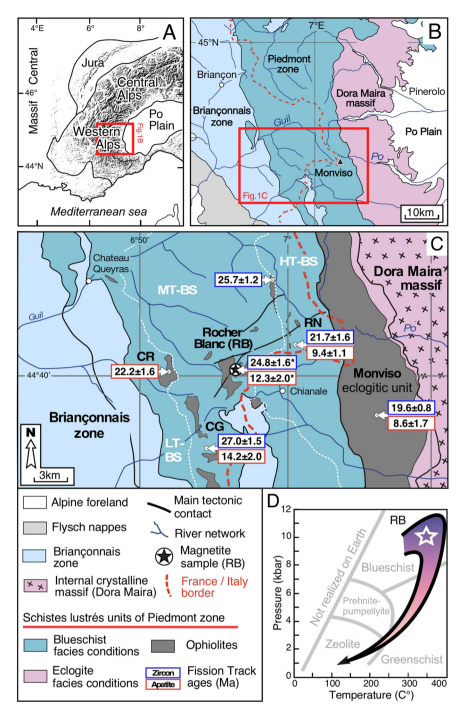


Fig. 1. (A) Geographical setting of the Alps showing the location of Fig. 1B. (B) Simplified structural and geological map of the Western Alps between France and Italy, with location of Fig. 1C. (C) Structural sketch-map of the Schistes Lustrés units of Piedmont zone, with published apatite and fission tracks ages (modified from Schwartz et al., 2007) and location of the Rocher Blanc ophiolite massif (RB), with extrapolated AFT and ZFT ages (in *). The location of CG (Cirque de la Gavie) and RN (Rocca Nera) are also indicated. (D) Pressure-Temperature path from the Rocher Blanc ophiolite (modified from Schwartz et al., 2009).

(glaucophane, zoisite) has been partially to totally retrogressed at greenschist facies conditions (chlorite, titanite and augite-aegirine). In ultramafic rocks, high pressure is characterized by the presence of antigorite (Schwartz et al., 2013). The studied sample (RB1) corresponds to a magnetite-bearing chloritized Fe-Ti meta-gabbro (chloritoschist), some tens of centimeters thick, located at the interface between meter-size lenses of Fe-Ti gabbro and serpentinite (Fig. 2A). The sample contains visible mm-scale euhedral magnetite crystals as shown in Fig. 2B, which are superimposed over the main schistosity defined by the chlorite – titanite assemblage (Fig. 2C).

3.2. Mineralogical and geochemical analysis

The chemistry of both euhedral and pseudo-euhedral magnetite crystals of the RB1 sample were characterized at the microscale. Semi-quantitative analyses and imaging of five magnetite and their inclusions were performed with quantitative analyses and mapping of minor and trace elements in magnetite (Si, Ti, Cr, V, Mn and Mg). Several SEM images are shown in Figs. 2D, E, F, and concentrations of Si, Ti, Sr, V, Mn, and Mg obtained on magnetite crystals are reported in Table 1. Fig. 3 presents the Si and V contents of one euhedral and one pseudo-euhedral magnetite crystal identi-

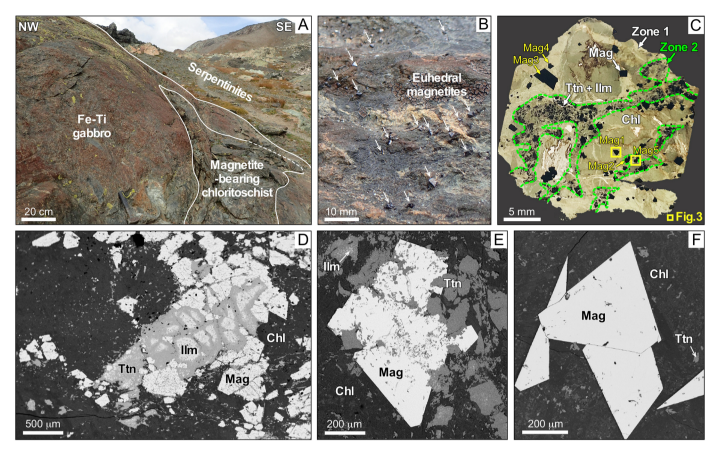


Fig. 2. Photographic views of the Rocher Blanc ophiolite massif, polished thin section of RB1 sample and SEM backscattered electron images. (A) Fe-Ti gabbro embedded in serpentinite and developing magnetite-bearing chloritoschist at its interface. (B) Millimeter scale euhedral magnetite crystals visible on outcrop indicated by white arrows. (C) Thin-section of the RB1 magnetite-bearing chloritoschist sample. The magnetite crystal analyzed is presented in Fig. 3. (D), (E) and (F) are SEM-BSE images of the sample RB1. The sample presents two mineralogical zones (2 and 1), rich in titanite and ilmenite (D and E) and chlorite (F), respectively. Mag, Ttn, Ilm and Chl mean magnetite, titanite, ilmenite and chlorite.

Table 1Elemental concentration in ppm obtained on representative magnetite crystals from zone 1 and zone 2 of the RB1 sample.

**			-			
Element	Si	Ti	Cr	V	Mn	Mg
Detection Limit (ppm)	24	30	30	30	35	25
Euhedral magnetite						
RB1-Mag1 (n = 76) Size 500 μm	680 ± 474	138 ± 102	55 ± 11	2119 ± 244	432 ± 35	B.d.l.
Profile B – Fig. 3 RB1-Mag3 (n = 69) Size 650 μm	1019 ± 638	72 ± 48	48 ± 10	1669 ± 267	420 ± 24	B.d.l.
RB1-Mag4 (n = 61) Size 600 μm	744 ± 349	60 ± 38	45 ± 10	1802 ± 230	429 ± 19	B.d.l.
Pseudo-euhedral magnetite	2					
RB1-Mag2 (n = 53) Size 550 μm	993 ± 650	59 ± 31	43 ± 11	1653 ± 329	418 ± 34	B.d.l.
RB1-Mag5 (n = 42) 480 μm Profile F - Fig. 3	608 ± 456	n.a.	n.a.	1863 ± 257	n.a.	n.a.
RB1 (n = 301)	$\textbf{809} \pm \textbf{513}$	$\textbf{82} \pm \textbf{55}$	$\textbf{48} \pm \textbf{10}$	$\textbf{1821} \pm \textbf{265}$	$\textbf{425} \pm \textbf{28}$	B.d.l.

B.d.l.: Below detection limits.

n.a. not analyzed.

fied also in Fig. 2C. Additional information can be found in the supplementary section.

In-situ uranium and thorium concentrations in chlorites, titanites and magnetites from sample RB1 were quantified by laser ablation inductively coupled plasma mass spectrometry (LA-ICPMS). Results are presented in Table 2 and on Fig. 4. Nevertheless, 2σ internal errors are reported in Table 2 and additional details can be found in the supplementary section.

3.3. Crystal selection using CT scanning

Magnetite grains scans were performed following guidelines by Cooperdock and Stockli (2016, 2018), and addition details can be found in the supplementary section. The blob3D software (Ketcham, 2005a; Cooperdock et al., 2019) was used to determine precisely the volume, surface/volume ratio, sphere-equivalent radius and alpha-ejection factor F_T (Ketcham et al., 2011) for

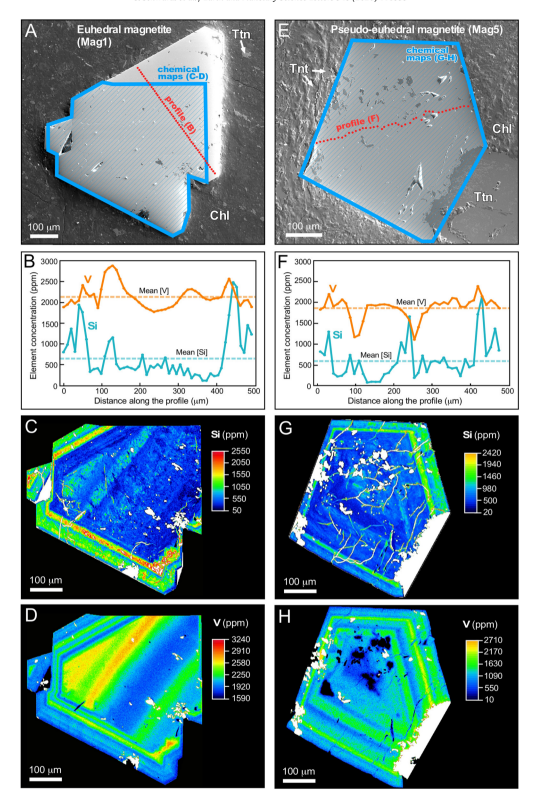


Fig. 3. Geochemical characterization of magnetite crystal from the chlorite-rich Zone 1 and titanite-rich Zone 2 shown in Fig. 2C (RB1-Mag1 and RB1-Mag5, Table 1). **(A,E)** EMP-SE image of the magnetite crystal with EMP elemental X-ray maps (1 μm beam size, 1.5 μm step size and 1 s dwell time) and profile. **(B,F)** Si and V concentration in ppm along the profile. **(C,G)** and **(D,H)** map of the Si and V concentration in ppm. Inclusions of silicate minerals appear in white on the chemical maps.

each grain, and the presence of large inclusions ($>5~\mu m$). 3D images of the magnetite crystals were generated to aid in selecting crystals for (U-Th-Sm)/He dating and documenting their euhedral and pseudo-euhedral shapes. 3D grain shape images of MgHe-analyzed magnetite crystals are reported in Figs. 5A and B.

3.4. (U-Th-Sm)/He on magnetite

Non-abraded single mm-size euhedral and pseudo-euhedral magnetite crystals were degassed under-vacuum in order to determine the He content using the protocol described in Allard et al. (2018). He content is corrected from the blank level, which is

Table 2 In-situ U and Th concentrations obtained by LA-ICP-MS in sample RB1 (thick section).

Magnetite	M1	M2	M3	M4c ^{\$}	M4r ^{\$}	М5с	М6с	M6r
	Zone 1 -	euhedral mag.				Zone 2 - ps	eudo euhedr	al mag.
$U \pm 2\sigma \text{ (ppb)}$ Th $\pm 2\sigma \text{ (ppb)}$	$15\pm2\\3\pm3$	5 ± 0.4 bdl	17 ± 4 6 ± 3	$\begin{array}{c} 8\pm2 \\ 1\pm0.7 \end{array}$	$\begin{array}{c} 21\pm2 \\ 6\pm9 \end{array}$	7 ± 1 3 ± 0.6	$\begin{array}{c} 7\pm 3 \\ 4\pm 1 \end{array}$	$\begin{array}{c} 24\pm16 \\ \text{bdl} \end{array}$
Titanite	T1	T2	T3	T7	T8	T4*	T5*	T6*
	Zone 2							
$\begin{array}{c} U \pm 2\sigma \; (ppb) \\ Th \pm 2\sigma \; (ppb) \end{array}$	94 ± 8 28 ± 22	$1569 \pm 141 \\ 628 \pm 546$	704 ± 56 109 ± 9	217 ± 72 163 ± 11		9 27 ± 5 4 ± 1	194 ± 16 36 ± 3	38 ± 6 6 ± 1
Chlorite		C4	C5		C1	C2		C3
		Zone 1			Zone 2			
$U \pm 2\sigma \text{ (ppb)}$ Th $\pm 2\sigma \text{ (ppb)}$		16 ± 8 5 ± 1	15 ± 7 13 ± 5		61 ± 6 7 ± 1	42 ± 13 ±		68 ± 10 17 ± 12

 $c^{\$}$ and $r^{\$}$ mean magnetite core and rim.

bdl = below detection limit.

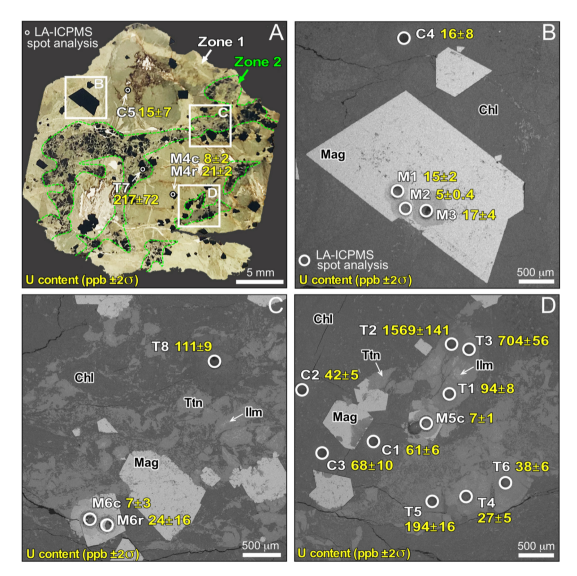


Fig. 4. In-situ U concentration of magnetite, chlorite and titanite from Zone 1 and Zone 2. (A) Thin-section of the RB1 magnetite-bearing chloritoschist sample, with location of the U analysis zones (B), (C) and (D). Zoom of the (B), (C) and (D) areas and U content in ppb.

from 40 to 160 times lower than the He signal. Helium content error is estimated to 2% accordingly. The wet chemistry digestion protocol was adapted from Blackburn et al. (2007) allowing the

determination of the 238 U, 232 Th, 147 Sm content. Additional details are placed in the supplementary section. Analytical uncertainties in the U, Th and Sm content translate into a 15% error on alpha-

^{*} Mn-rich titanite.

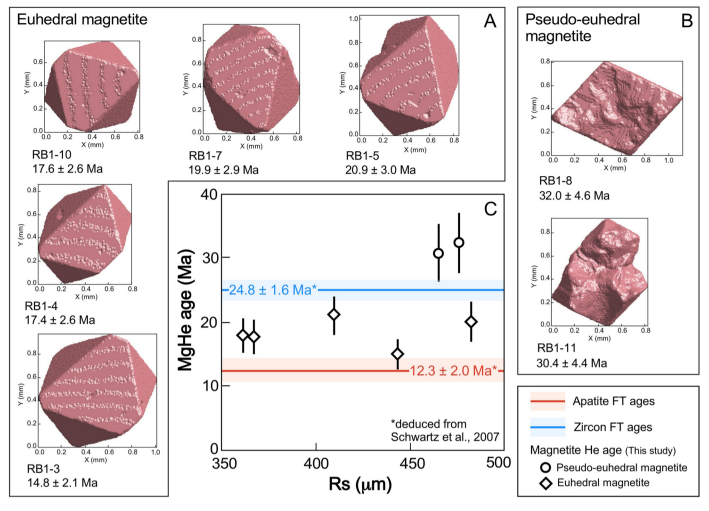


Fig. 5. Magnetite (U-Th-Sm)/He ages from Rocher Blanc ophiolite. (A) and (B) are 3D reconstructions from CT-scan images of each euhedral and pseudo-euhedral crystals. (C) MgHe age as a function of the sphere equivalent radius (Rs) estimated from CT-scan analysis (Ketcham, 2005a; Cooperdock et al., 2019). The AFT and ZFT age ranges from Schwartz et al. (2007) and shown in Fig. 1C are shown.

ejection-corrected MgHe ages. Sample mass, volume, He, U, Th, Sm content, raw and alpha-ejection-corrected (U-Th-Sm)/He ages are reported in Table 3.

4. Results

4.1. Magnetite phase relationships and chemical composition

At the thin section scale, magnetite crystallizes in two zones of distinct mineralogy, which are likely inherited from the initial distribution of magmatic minerals in the Fe-Ti gabbro (Fig. 2C). The mineralogy of these two zones is either dominated by a chlorite + magnetite \pm relict magmatic pyroxene (Zone 1) or titanite + magnetite \pm chlorite \pm relict magmatic ilmenite (Zone 2) assemblages. In Zone 1, the augite-aegirine pyroxene may stably coexist with titanite and chlorite contains a 10-20% jadeitic component. In Zone 2, titanite, magnetite and chlorite appear texturally as destabilization products of magmatic ilmenite and generates local reaction coronas (Fig. 2D). Plagioclase has not been identified in either of the two zones. Magnetite crystal habits differ between the two zones. In Zone 2, magnetite is pseudo-euhedral to anhedral, possibly polycrystalline, and contains numerous inclusions (Fig. 2D-E) of titanite and chlorite. In Zone 1, magnetite is euhedral, and can form larger grains exceeding 500 µm with rare inclusions (Fig. 2F) of augite-aegirine, chlorite and titanite.

Chemical characterization of euhedral magnetite crystals from Zone 1 and pseudo-euhedral magnetite crystals from Zone 2 are reported in Table 1. Significant Mn, Cr and Ti contents are present but remain below 500 ppm (Table 1). Mg is below detection limit (i.e., $<\!25$ ppm), while Si and V are the two most concentrated cations after Fe, with mean concentrations of 809 ± 513 and 1821 ± 265 ppm, respectively (Table 1). The Si and V distribution patterns are the same in euhedral and in pseudo-euhedral magnetite grains indicating similar growth process and history. It is characterized by oscillatory zoning with a relatively good correlation between Si and V-rich growth zones (Fig. 3), although vanadium zoning is sharper. The nature of the inclusions and their distribution seem independent of the chemical variations recorded by these two elements.

4.2. Magnetite, chlorite and titanite in-situ U content

The U content of magnetite derived from laser ablation ranges from 5 ± 0.4 to 24 ± 16 ppb and is \sim five times higher in the rim compared to the core (Table 2, Fig. 4). In addition, the U content is similar to the U value obtained by wet chemistry for the dated crystals with value of 8 ± 1 to 29 ± 2 ppb (Table 3). The U content is also similar between magnetite from Zone 1 (euhedral) and Zone 2 (pseudo-euhedral) (Table 2 and 3). The U content in chlorite from Zone 1 of \sim 16 \pm 8 ppb is lower than the content in chlorite from Zone 2 which ranges from 42 ± 5 to 68 ± 10 ppb (Table 2, Fig. 4).

CT-scan and (U-Th-Sm)/He results of the magnetite-bearing chloritoschist RB1 sample

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Aliquot	Volume	Surface area	Rs	F	Mass	⁴ He	^{238}U	²³² Th	¹⁴⁷ Sm	⁴ He ± s	238U	²³² Th	¹⁴⁷ Sm	eU	Th/U	Raw date	Corrected date \pm s	
	(mm ³)	(mm ²)	(mm)		(mg)	(lomd)	(gu)	(gu)	(ng)	(blomd)	(qdd)	(qdd)	(qdd)	(qdd)		(Ma)	(Ma)	
Euhedral	Euhedral magnetite																	
RB1-3	0.29	2.72	4 4 4	0.98	1.54	2.33×10^{-3}	0.025	0.020	0.018	1.49 ± 0.03	16.2	13.0	11.7	19	8.0	14.4 ± 2.2	14.8 ± 2.1	
RB1-4	0.16	1.82	368	0.97	0.84	1.85×10^{-3}	0.019	0.007	0.021	2.18 ± 0.04	22.7	8.4	25.1	25	0.4	16.9 ± 2.5	17.4 ± 2.6	
RB1-5	0.21	2.28	410	0.97	1.12	1.16×10^{-3}	0.009	0.007	0.015	0.88 ± 0.02	7.8	6.5	13.3	6	8.0	20.4 ± 3.1	20.9 ± 3.1	
RB1-7	0.25	2.46	483	0.98	1.32	4.71×10^{-3}	0.038	0.027	0.075	1.85 ± 0.04	28.7	20.2	56.9	34	0.7	19.5 ± 2.9	19.9 ± 2.9	
RB1-10	0.16	1.82	362	0.97	0.85	2.03×10^{-3}	0.020	0.009	0.015	2.31 ± 0.05	23.2	11.2	17.5	26	0.5	17.1 ± 2.6	17.6 ± 2.6	
Pseudo-e	Pseudo-euhedral magnetite	ıetite																3
RB1-8	0.18	2.12	477	0.98	0.94	3.23×10^{-3}	0.016	0.012	0.020	3.26 ± 0.07	17.3	12.3	21.9	20	0.7	31.3 ± 4.7	32.0 ± 4.8	. <i>S</i> t
RB1-11	0.13	1.80	466	0.98	0.71	1.60×10^{-3}	0.008	0.007	0.004	2.39 ± 0.05	11.7	9.7	6.2	14	8.0	29.7 ± 4.5	30.4 ± 4.6	ILVV
																		ш

Rs and F_T are the sphere equivalent radius and alpha ejection factor. (U-Th-Sm)/He date has been corrected from alpha ejection factor (F_T) (Ketcham et al., 2011).

Although variable, the U content of titanite is by far the highest with values ranging from 27 ± 5 to 1569 ± 141 ppb. In the RB1 sample titanite is the U-bearing phase.

4.3. Magnetite (U-Th-Sm)/He

Five euhedral magnetite crystals (Fig. 5A) and two pseudo-euhedral grains (Fig. 5B) were selected for MgHe dating, utilizing the CT scan data to select grains with the fewest (for pseudo-euhedral magnetite) or no inclusions. The selected crystals have equivalent-sphere radii (Rs) ranging from 362 to 483 μm (Table 3). Helium content of these seven magnetite grains ranges from 0.9 to 3.3 pmol/g and effective uranium (U + 0.234xTh) ranges from 9 to 34 ppb (Table 3). Ejection-corrected MgHe ages for euhedral and pseudo-euhedral grains range from 14.8 \pm 2.1 to 20.9 \pm 3.0 Ma and 30.4 \pm 4.4 to 32.0 \pm 4.6 Ma, respectively (Fig. 5C).

5. Discussion

5.1. Conditions of magnetite formation

Magnetite is a common mineral in ultramafic and mafic rocks and appears over a large range of P-T and $\rm fO_2$ conditions (Nadoll et al., 2014). In the case of magnetite created during the oceanic serpentinization process, it is classically assumed to form at temperatures above $\sim\!200\,^{\circ}\text{C}$ (e.g., Alt et al., 2013; Klein et al., 2013, 2014), even if magnetite could potentially crystallize at temperatures even below $120\,^{\circ}\text{C}$ (Alt et al., 2013). In the case of subducted ophiolites (serpentinite and gabbro lithologies), magnetite is observed from eclogite paragenesis (Li et al., 2004; Debret et al., 2014) up to greenschist facies conditions related to the retrograde stage for temperatures ranging from 550 to 250 $^{\circ}\text{C}$ (Schwartz et al., 2013).

The petrological and geochemical studies of the RB1 sample indicate that the euhedral and pseudo-euhedral magnetite grains crystallized during the retrograde path under greenschist facies conditions assisted by metamorphic hydrothermal fluids and are not derived from magmatic origin. Indeed, the Ti content of magnetite crystals is under 100 ppm (Table 1), too low to be of magmatic origin (Nadoll et al., 2014), and the host inclusions (titanite, chlorite, augite-aegirine) that are representative of HP-LT paragenesis demonstrate that magnetite grains crystallized during the greenschist retrograde path. In this context, the oscillatory V and Si chemical zoning, with a Si-richer rim observed for euhedral and pseudo-euhedral magnetite grains (Fig. 3) record the changing conditions and compositions of the metamorphic fluid related to magnetite growth (Dare et al., 2014). The Si and V are the main minor elements at the \sim 1000 to 2000 ppm level, compared to Ti, Cr, Mn, and Mg (Table 1); these Si and V contents reveal high solubility indicating crystallization from a high temperature fluid (Dare et al., 2014). Such high Si content has already been encountered in hydrothermal systems (skarns, metamorphic veins; e.g., Shimazaki, 1998) and Si oscillatory zoning has been described in hydrothermal magnetite (Deditius et al., 2018). Hydrothermal conditions (250-350 °C, Fig. 6A) in the fluid-circulation zone marked by the magnetite-bearing chloritoschists may have favored Si and V incorporation, the latter element being supplied locally by the Fe-Ti gabbro itself.

Based on previous thermobarometric studies of the host metamorphic Fe-Ti gabbro (e.g., Pognante and Kienast, 1986), and petrological investigation in this study, the magnetite begins to crystallize most probably near the blueschist – greenschist transition, as residual augite-aegirine is found in inclusions, and magnetite crystallization continues during the entire greenschist retrograde path (Fig. 6B).

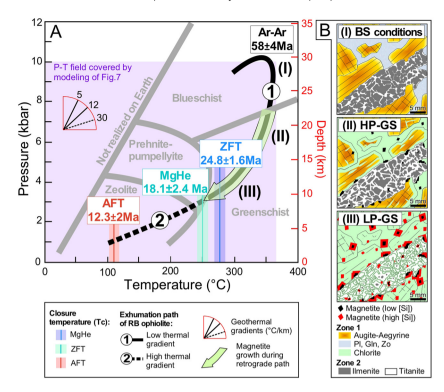


Fig. 6. Pressure-temperature-time pathway record by the RB ophiolite and magnetite growth condition. (A) Pressure-Temperature-time path of the HP-LT Rocher Blanc ophiolite modified from Schwartz et al. (2009, 2013). The AFT and ZFT estimated ages from RB (Fig. 1C), MgHe ages from euhedral magnetites are reported for their associated closure temperature condition (Blackburn et al., 2007; Gallagher et al., 1998; Ketcham, 2019). The age of HP conditions are taken from Agard et al. (2002). (B) Sketch of magnetite crystals growth during the retrograde pathway in three main steps, from (I) blueschist conditions (BS), (II) High-Pressure greenschist (HP-GS) and (III) Low-Pressure greenschist (LP-GS) conditions. Zones 1 and 2 refer to the initial magnatic sites with augite-plagioclase and Fe-Ti oxide (ilmenite) respectively. (I) The HP blueschist conditions are marked by the partial mineralogical transformation of augite to aegirine, and the development of glaucophane and zoisite in the plagioclase site (Pognante and Kienast, 1986). In Zone 2, ilmenite is progressively transformed to titanite. (II) The HP greenschist conditions are marked by the progressive destabilization of augite-aegirine, glaucophane, and zoisite to chlorite and magnetite in Zone 1, whereas, ilmenite continues to be destabilized in titanite and ± magnetite and chlorite. The magnetite grains show low Si content whatever the considered zones. (III) Finally, the LP greenschist facies conditions, show the pervasive development of the chlorite and magnetite in the two zones. In the zone 2, only relics of ilmenite remain locally in the core of titanite crystals. The magnetites present higher Si content.

5.2. Magnetite He age distribution

The MgHe ages for all crystals are younger than the highpressure ages recorded for the Schistes lustrés units (62-54 Ma; Agard et al., 2002), and potentially represent exhumation ages, i.e., related to the retrograde path. Interestingly, the euhedral grains are quite homogeneous with a mean age of 18.1 \pm 2.4 Ma that falls between the AFT and ZFT ages as deduced from Schwartz et al. (2007) for the Rocher Blanc area (Fig. 5C), and are thus compatible with cooling ages associated with the exhumation history of the area. Indeed, the MgHe ages are younger than ZFT ages, and older than AFT ages, in agreement with the published thermal sensitivity of MgHe (250 \pm 50 °C; Blackburn et al., 2007), ZFT $(240 \pm 50 \,^{\circ}\text{C}, \text{ Gallagher et al., } 1998; \sim 281 \,^{\circ}\text{C}, \text{ Ketcham, } 2019)$ and AFT (110 \pm 10 °C, Gallagher et al., 1998). Conversely, the ages obtained on pseudo-euhedral crystals are significantly older than the euhedral crystal and ZFT ages (Fig. 5C). This is not consistent with petrological data which show that euhedral and pseudo-euhedral magnetite all crystallized simultaneously under greenschist conditions at T > 250 °C. Thus, we interpret these ages as related to the presence of excess He compared to euhedral crystals with similar size and eU content (Table 3); they cannot be considered as cooling ages. Excess of He may be due to (i) the impact of alpha implantation from neighboring minerals or circulating rich U-Th fluids, and/or (ii) the impact of inclusions with higher U-Th content on the global He budget of magnetite.

Alpha implantation has been frequently observed or modeled for the (U-Th)/He method, especially for apatite (e.g., Spiegel et al., 2009; Gautheron et al., 2012) and magnetite (Blackburn et al., 2007; Cooperdock and Stockli, 2016). When minerals contain lower

or equivalent U and Th content compared to neighboring phases, the alpha implantation effect can be significant. This is particularly the case for magnetite, which typically contains low U-Th content (<30 ppb) compared to minerals with high U-Th content of >100 ppb like titanite in the present case (Table 2). In Zone 1, magnetite is embedded in chlorite which is U-poor with contents \sim 15 ppb. It must be noted that the higher U content in Zone 2 results in a higher U content, from 40 to 60 ppb, in chlorite as well (Table 2, Fig. 4). In the RB1 sample, for the chlorite area (Zone 1) alpha implantation in magnetite should be negligible whereas in the titanite area (Zone 2) alpha implantation in magnetite could be significant. As the RB1 grains are large (Rs >400 μ m) with small alpha ejection corrections ($F_T > 0.97$; Ketcham et al., 2011), a 20-40 ppb matrix would simply cancel out the F_T correction, and only thousands of ppb U-Th will have an age effect approaching 10% (Gautheron et al., 2012). For euhedral magnetite, MgHe ages younger than ZFT imply no significant alpha implantation and reinforce the hypothesis that euhedral magnetites crystallized in Zone 1 (chlorite-rich environment). Similarly, MgHe ages older than ZFT observed for the pseudo-euhedral magnetite may reflect implantation from U-Th-rich neighboring minerals such as titanite, consistent with the pseudo-euhedral magnetites crystallizing in Zone 2, in agreement with petrological observations.

Inclusions richer in U-Th than the host minerals can also impact (U-Th)/He ages by adding He to the crystal lattice, as already reported for apatite (Farley, 2002) and magnetite (Blackburn et al., 2007; Cooperdock and Stockli, 2016). An excess of He can occur if the total U-Th content of host mineral and inclusions is incompletely retrieved by the chemical digestion protocol that is usually designed to target the host mineral. For magnetite, the digestion

protocol is not adapted to dissolve completely silicates (e.g., chlorite, titanite, zircon) or other oxide minerals (e.g., titanite). A small fraction of the U-Th budget coming from inclusions may be retrieved, but the total U-Th budget is incomplete, leading to spuriously older MgHe ages with no geological significance. For the two pseudo-euhedral magnetites from RB1, the MgHe ages older than the euhedral magnetites with similar U-Th content (Table 2) would be consistent with the absence of U-Th-rich inclusions in the euhedral crystals. However, because the CT scan analysis indicated minimal detectable inclusions, and because the primary U-Th-bearing phase, titanite, has eU compositions only ~100x higher than the magnetite (Table 2, 3), we do not believe inclusions can explain the magnitude of excess He. We thus interpret that MgHe ages older than ZFT reflect implantation across large areas of grain boundaries shared with titanite (Fig. 4D), and for the rest of the discussion we do not use or further discuss the two pseudo-euhedral crystals, as their measured ages are not geologically meaningful. However, we can conclude that the MgHe ages from the euhedral crystals of the RB1 sample are not significantly impacted by implantation and reflect cooling due to exhumation along the retrograde path.

Finally, the results of this study give some methodological insight into MgHe dating. First, CT imaging is essential for detecting the presence of mineral inclusions larger than \sim 5 µm, as already stated by Cooperdock and Stockli (2016). Comparison of U-Th content for crystals with older MgHe ages can provide indirect evidence of the impact of rich U-Th inclusions below the CT-scan detection limit ($<1 \mu m$). Second, the removal of the 20 μm outer margin of the magnetite crystals, as recommended by Blackburn et al. (2007), is not always necessary. In fact, the result of this study shows that implantation is not always significant and depends on the petrological environment of magnetite (i.e. chlorite versus titanite zones), as well as the size of the crystal. Furthermore, alpha implantation along long grain boundaries shared with titanite, as documented in our zone 2 non-euhedral magnetites, if combined with diffusional migration of helium into the magnetite grain (Gautheron et al., 2012) may render abrasion an insufficient remedy, depending on cooling history. Thorough petrological characterization of mineral textures and compositions thus appears to be a necessary step of the MgHe method.

5.3. Origin of MgHe age dispersion

Significant MgHe age dispersion is encountered for euhedral RB1 magnetites (MgHe age $=18.1\pm2.4$ Ma, 13% at 1σ). MgHe age dispersion has also been reported in previous work in the HP-LT subduction complex on the island of Syros, Greece (Cooperdock and Stockli, 2016). This age dispersion is higher than typical analytical error (i.e. $\sim\!5$ -10%, for apatite and zircon) but similar or lower than frequently found in practice for apatite or zircon (U-Th)/He methods (e.g., Ault et al., 2019; Ketcham et al., 2018). Using investigations already published on the origin of apatite or zircon (U-Th)/He age dispersion, some factors affecting MgHe ages related to the impact of (i) U-Th zonation, (ii) slight alpha-implantation, or (iii) crystal-dependent He diffusion coefficient can be anticipated.

First, a slight impact of alpha-implantation could affect the MgHe age, as the U-Th content of the neighboring minerals could be slightly different For example, U and Th content for chlorite of $\sim\!15$ ppb has been determined for Zone 1 (Table 2), but slightly higher U-Th values are possible (Table 2). However, the large crystal size of the magnetite from this study reduces the impact of implantation. Nevertheless, other parameters can account for dispersion. For instance, Cooperdock and Stockli (2016) found similar MgHe age dispersion on abraded crystals (i.e. mechanical removal of the 20 μ m border of the crystal that can have been affected by alpha implantation).

Second, due to the low U-Th content, analytical determination of zoning is difficult, and up to a \sim 4x higher U concentration in the rim has been measured for three magnetite crystals (Table 2). Because the ablation spot is large (150 µm), the exact size of the U-Th-enriched rim is not known but can be similar to the one observed by Si-V analysis, i.e. \sim 100 μ m (Fig. 3). In this case, the presence of a ~100 µm enriched U-Th rim will not have much impact on the ejection factor as it is much larger than the mean stopping distance (15 µm). If finer zoning exists, modeling can be performed (e.g., Farley et al., 1996; Gautheron et al., 2012) assuming that U-Th zoning follows Si and V zoning (Fig. 3C and D). Using the Monte Carlo code associated with Gautheron et al. (2012), we modeled a 400 µm radius sphere with a 50 µm rim depleted or enriched in U-Th by 50%, and as a worst-case scenario a rim containing all of the crystal's U-Th (i.e., no U-Th in the core). We obtained F_T ejection factors of 0.99, 0.96, and 0.76, respectively, implying a 25% MgHe age difference for the worst case. However, in comparing our bulk and in situ core and rim analyses we find no evidence of such severe zoning in U-Th, and we can estimate a maximum MgHe age dispersion component of only up to a few percent. Given that implantation can be from minerals with much more than 50% higher eU, it is likely to be a source of equal or greater dispersion.

Third, a defect- and/or composition-dependent He diffusion coefficient could also be invoked to explain age dispersion, as has been proposed and modeled for other minerals such as apatite and zircon (Gautheron et al., 2009; Flowers et al., 2009; Guenthner et al., 2013). Indeed, damage due to radioactivity, and also crystal defects could trap or facilitate He diffusion within the crystal structure. A damage content of some 5% would be required to change the He diffusion coefficient significantly, as modeled for apatite (e.g., Gerin et al., 2017), which is unlikely given the low eU content of magnetite (\ll 1%). Alternatively, substitution by minor or trace elements could deform the crystal structure and thus modify He diffusion, as shown in apatite at the atomic scale (Djimbi et al., 2015). The magnetite crystals from RB1 have \sim 0.5 wt% of minor elements, and if we assume similar effect as for apatite (Djimbi et al., 2015), no significant change in the closure temperature compared to pure magnetite is expected. This is highly speculative, as the composition of the volcanic magnetite used by Blackburn et al. (2007) is not known. The effect of minor elements in magnetite on He volume diffusion still needs to be investigated. It is also possible that fluid inclusions can act as traps for He, as has been proposed for apatite and zircon (Zeitler et al., 2017; Danišík et al., 2017); this possibility could be further pursued with higherresolution imaging and/or ramped heating experiments (McDannell et al., 2018).

Finally, we stress that there is currently only one experimental study documenting He diffusion kinetics in magnetite (Blackburn et al., 2007), and even that data set shows signs of complex release behavior. Further data are certainly required to build a more robust database of kinetic parameters and explore possible anomalous behaviors.

5.4. MgHe data inversion: resolving exhumation rate from the Rocher Blanc ophiolite

The petrological and geochemical arguments deduced from the RB1 sample analysis indicate that magnetite crystalized at $T>250\,^{\circ}\text{C}$, as illustrated in Fig. 6, at temperature higher than the MgHe closure temperature (Tc), and thus that MgHe ages obtained from euhedral magnetite of ca. 18.1 ± 2.4 Ma correspond to cooling ages of the Rocher Blanc ophiolite (Fig. 6). Specifically, for the dated RB1 magnetites, which all have similar sizes (Table 3), Tc ranges from 242 to 248 °C for sphere-equivalent radii of 362 to 483 μ m and a cooling rate of $10\,^{\circ}\text{C/Ma}$. MgHe ages derived from

Table 4 Thermal history inverse modeling parameters.

1. Thermochronologic data	a
Samples and data used	AFT and ZFT data from Schwartz et al. (2007)
	MgHe data from this paper: Table 2, samples RB1-3, 4, 5, 7, and 10
Data treatment: Dates, und	certainties, and other relevant constraints
AFT data	Kinetics: Apatite assumed to have $Cl = 0$ apfu
	Uncertainty model: Asymmetric 95% bounds
ZFT data	Initial mean track length: N/A; no track length data available
	Uncertainty model: Asymmetric 95% bounds
MgHe data	Uncertainty model: Uniform 15% (1SE) uncertainty
2. Additional geologic info	rmation and constraint placement
Present-day temperature	20°C
Initial constraint	Assumed peak HP-LT conditions: $350 \pm 20^{\circ}\text{C}$ at $58 \pm 4\text{Ma}$
T-t path complexity	HeFTy code E3/50: Paths episodic, halved 3 times, with maximum cooling
	rate 50°C/m.y.
3. System- and model-spec	ific parameters
AFT annealing model: Ketc	ham et al. (1999)
ZFT annealing model: Ketc	ham (2019)
MgHe diffusion parameters	s: Blackburn et al. (2007)
Modeling code: HeFTy v1.9	0.3
Statistical fitting criteria: H	IeFTy default
Ending condition: Models	run until 50 good fits found

are sandwiched between ZFT and AFT ages as expected by their thermal sensitivity, and in fact shed further light on the thermal sensitivity of the ZFT method, supporting a ZFT Tc >260 °C. Fig. 6 summarizes the P-T path for RB1 sample, including the magnetite growth field during the retrograde path and the time constraint obtained from AFT, ZFT and MgHe ages.

5.4.1. MgHe data inversion

The thermal history of the studied area was retrieved using the HeFTy software (Ketcham, 2005b) by inverse modeling of MgHe, AFT and ZFT low temperature thermochronometric data. HeFTy uses a constrained Monte Carlo scheme that poses a large number of candidate time-temperature (t-T) paths, and retains those that fit the thermochronometric data to a "good" (closer than statistical expectation for a random sample) or "acceptable" (does not fail a 95% confidence test) extent. For the inversions the fission-track annealing models by Ketcham et al. (1999) and Ketcham (2019) were used for AFT and ZFT data, respectively. For MgHe ages, the sphere-equivalent radii (Rs) of each crystal and He diffusion parameters from Blackburn et al. (2007) were used. In detail, the HeFTy software models (i) the fission-track production and annealing in apatite and zircon, and (ii) helium production, taking into account ejection and diffusion, in magnetite. In this way, the data inversion identifies a family of potential cooling histories by using 17 t-T nodes randomly placed to define a cooling-only path with a maximum allowed cooling rate of 50 °C/m.y. over any segment. In addition to the MgHe, AFT and ZFT data, the peak HP-LT metamorphic conditions for the Rocher Blanc ophiolite massif as reported in Fig. 6, estimated to be 350 \pm 20 °C at 58 \pm 4 Ma (Agard et al., 2002) is used at a starting point. Further model inputs are listed in Table 4.

Thermal history modeling was performed with AFT-ZFT alone, with MgHe data alone and with all three combined (AFT-ZFT and MgHe). First, modeling was done with interpolated AFT and ZFT ages for RB1 sample assuming that ages vary linearly with the sample position on a W-E transect (Schwartz et al., 2007) (Fig. 1), and with and without MgHe data. Second, inversions were also performed with AFT and ZFT data from RN and CG samples, that represent the available data geographically framing the RB ophiolite (Fig. 1C), and with and without MgHe data. Finally, for comparison, the AFT-ZFT data and the MgHe data separately for RB1 sample were inverted. For modeling of MgHe data, all euhedral grains were individually included. The computed thermal histories

are reported on Fig. 7. In the on-line supplement, Fig. S1 presents the modeling inversion results obtained for AFT and ZFT data of CG and RN samples.

5.4.2. Thermal history modeling

The inversion of the MgHe data alone or with the AFT and ZFT literature data allows us to refine the final t-T exhumation history of the Rocher Blanc ophiolite massif, while also verifying good consistency with AFT and ZFT data. First, model outputs showing the set of good and acceptable fits, and their weighted mean, demonstrate that the MgHe data provide time constraints at ca. 230-240 °C (Fig. 7A) whereas ZFT and AFT ages put constraints on the Rocher Blanc ophiolite cooling history at ~280 and ~100 °C (Fig. 7B) respectively. In particular, the MgHe data refine the thermal history by requiring that the rapid phase of cooling did not commence until after 20 Ma (Fig. 7C). The three systems are thus complementary with respect to the temperature range that they cover. The apparent consistency between MgHe, AFT and ZFT cooling ages suggests that the He diffusion data obtained at 1 bar by diffusion experiments in vacuum at temperatures between 350 and 620°C over several days by Blackburn et al. (2007) can be extended to geological conditions and timescales. As the AFT and ZFT data for RB1 sample are extrapolated from a different locality, we also ran models with AFT and ZFT ages from samples CG and RN (Fig. S1). The resulting cooling histories only slightly differ from those obtained with the extrapolated Rocher Blanc ophiolite ages (Fig. S1) supporting the relevance of this thermal modeling approach.

5.5. Final exhumation of the Rocher Blanc ophiolite: geodynamic implications

The thermal history obtained using the combination of thermochronometric methods (AFT-ZFT and MgHe data) brings strong constraint on the final exhumation of the Rocher Blanc ophiolite and also on the Schistes lustrés units of the Piedmont zone (Western Alps). Our thermal modeling reveals that the RB1 sample has undergone nonlinear cooling and records a major change in the thermal pathway at 20 Ma (Fig. 7C). During the first exhumation stage, from 55 to 20 Ma, the sample recorded slow cooling from 350 to 300 °C associated with an extensive exhumation from 30 to 10 km (Fig. 6). The exhumation of ∼20 km of rocks from the Alpine orogenic wedge cannot be accounted for by sur-

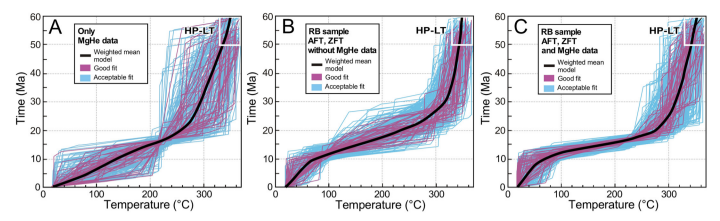


Fig. 7. Thermochronological data inversion modeling of the HP-LT Rocher Blanc RB ophiolite using HeFTy software (Ketcham, 2005). (A) T-t pathways obtained for MgHe data inversion only. (B) T-t pathways obtained for AFT and ZFT data inversion only. (C) T-t paths obtained for joint AFT, ZFT and MgHe data inversion.

face denudation (i.e. erosion) only. Large exhumation with limited cooling rather corresponds to exhumation in ductile conditions related to subduction dynamics and underplating processes (Agard et al., 2001). During ductile exhumation the sample passes from blueschist to greenschist conditions with a mean exhumation rate lower than 1 mm/yr, calculated using the depth and time constraints (Fig. 6A and 7). Over the second exhumation stage, from 20 Ma to 10 Ma, the RB1 sample records rapid cooling from 300 to 100°C (Fig. 7) associated with an exhumation from 10 to 3 km (Fig. 6A). This exhumation and strong change of cooling rate reflect denudation in the brittle conditions. A mean exhumation rate of 0.7 mm/yr can be deduced (Fig. 6A and 7). The last information of the thermal history is given by the final exhumation stage recorded by AFT data and an exhumation of 3 km over 10 Ma leading to a mean lower exhumation rate of 0.3 mm/yr (Fig. 7). Interestingly, the sample records similar exhumation rate in ductile and brittle domains around <1 mm/yr, but significant cooling rate variations. The change at \sim 20 Ma corresponds to the passage of the Rocher Blanc ophiolite below the transition from ductile to brittle conditions in the lower grade of the greenschist facies conditions (\sim 250 °C) during the retrograde path.

The final exhumation recorded by the thermal history reflects denudation at brittle conditions (Fig. 7), where denudation is related to erosion linked to relief creation. The relief creation is associated to a westward tilting of the Schistes lustrés units (Tricart et al., 2004; Schwartz et al., 2009), as a response to the upward indentation of the Dora Maira massif. Indeed, the crustal scale geometry of the internal zone is characterized by the presence of Apulian mantle in a superficial position (Fig. 8), at 12 kilometers below the Dora Maira massif (Lardeaux et al., 2006; Zhao et al., 2015). This portion of cold and rigid mantle acts as an indenter that allowed the deformation of the thinned Briançonnais margin by crustal underthrust process constituting a continental subduction wedge that forms the current basement of the Piedmont zone (oceanic subduction wedge in Fig. 8). This continental subduction wedge was overthrust on the European crust by the activation of the crustal Penninic Frontal Thrust (PFT in Fig. 8) at \sim 34 Ma (Dumont et al., 2012).

Taking in consideration the MgHe data of this study, the AFT-ZFT data of Piedmont zone (Schwartz et al., 2007) and AFT data of the internal crystalline Massif of Dora Maira (Beucher et al., 2012), we can illustrate the evolution of the area with their respective positions though time in the orogenic wedge (Fig. 8). At present-day, all samples crop out at similar elevations (>2000 m a.s.l.) for the Piedmont zone and at lower elevation (<1500 m a.s.l.) for Dora Maira. At present-day the global shortening affects the front of the European foreland (Schwartz et al., 2017), and allows the uplift and

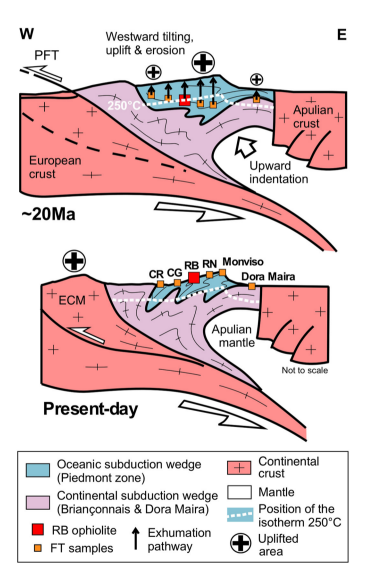


Fig. 8. Schematic evolution of the Western Alps between France and Italy ${\sim}20$ Ma ago and at present-day. At ${\sim}20$ Ma, the Penninic Frontal Thrust (PFT) is already active since 10 Myr associated with the mantellic upward indentation, and the AFT-ZFT and MgHe dated samples have started to be exhumed in the brittle condition (above 250 °C). The samples record differential exhumation related with the westward tilting and uplift. One representative sample from Dora Maira massif is represented, where AFT ages of 20-25 Ma have recorded by Beucher et al. (2012).

exhumation of the external crystalline massifs (Pelvoux, Beucher et al., 2012) (Fig. 8). Using the AFT-ZFT and MgHe ages and thermal sensitivity of the thermochronometric systems for all the samples, we can reconstruct their position at depth \sim 20 Ma ago. Firstly, the samples CR from Schistes lustrés and from Dora Maira present AFT age of 20-25 Ma indicating a shallow position around 3 km (assuming an AFT closure temperature of 100 °C and a thermal gradient of 25 °C). The position at depth for the other samples from the Piedmont zone need to be progressively deeper from east to west (Fig. 8). RB1 sample was passing the $250\,^{\circ}\text{C}$ isotherm $\sim\!20$ Ma based on the thermal history obtained in this study (Fig. 7). From \sim 20 Ma to present-day, the Piedmont zone has recorded a differential exhumation with a westward titling and uplift that is recorded by the youngest ages to the east in the Monviso (Fig. 8). This exhumation is controlled by the upward indentation of the Apulian mantle producing the global tilting of the Piedmont zone (Fig. 8).

6. Conclusions

This study contributes to the understanding of the formation conditions of magnetite in HP-LT ophiolite and to the use of magnetite (U-Th-Sm)/He as a thermochronometer to quantify the exhumation history. We demonstrate that a comprehensive characterization of magnetite crystals and crystallization environment is needed in conjunction with MgHe dating in order to interpret the age data.

We specifically focused on one sample (RB1) from the Rocher Blanc massif that is an ophiolitic body embedded in the Schistes lustrés units (Western Alps, France), which contains abundant magnetite. This sample recorded middle-temperature blueschist facies conditions and was overprinted by greenschist conditions during exhumation. We show that for the RB1 sample, magnetite crystallized in two zones of distinct chemistry, that originated from magmatic relictual zones, with euhedral and pseudoeuhedral grains with a dominant chlorite or titanite mineralogy, respectively. In addition, the petrological and magnetite geochemical analyses (e.g. V and Si) indicate that magnetite crystallized during the retrograde metamorphic path under greenschist conditions (T > 250 °C). The Si and V zoning observed in euhedral and pseudo-euhedral magnetite grains reinforce the fact that both type of magnetite crystallized under same physical conditions. The pseudo-euhedral magnetite, crystallizing in dominant titanite mineralogy, present older MgHe ages compared to euhedral magnetite. The old MgHe ages are interpreted by the impact of alphaimplantation from U-Th-rich neighbor minerals (e.g., titanite). On the contrary, alpha-implantation in euhedral magnetite crystallizing in chlorite dominant zone is insignificant, as chlorite contain similar U-Th content than magnetite. The MgHe ages derived from euhedral magnetite range from 14.8 ± 2.1 to 20.9 ± 3.0 Ma, sandwiched between ZFT and AFT ages as expected. The MgHe data along with AFT and ZFT data were inverted for the first time to refine the local thermal history. The MgHe data suggest that fast cooling was delayed until \sim 20 Ma, as opposed to the less constrained ~26 Ma that would be inferred from the ZFT and AFT ages alone. The MgHe data suggest that the RB1 sample has undergone an exhumation in ductile conditions related to subduction dynamics and underplating processes, followed by an exhumation reflecting denudation by erosion at brittle conditions. Considering all thermochronometric data along the Western Alps, we propose that the exhumation is controlled by the upward indentation of the Apulian mantle producing the global titling of area and associated erosion.

Additional work on U-Th zoning in magnetite and He diffusion in magnetite will be necessary in the future, as the thermal inversion interpretation is contingent on the Blackburn et al. (2007)

kinetic parameters being correct. Nevertheless, the present study further demonstrates that MgHe can be used as a low-temperature thermochronometer to date HP ultramafic final exhumation and the ductile-brittle transition opening new avenues of research. This method effectively complements the available thermochronological tools, making it a powerful tool for emerging applications.

Declaration of competing interest

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

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Appendix A. Supplementary material

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.epsl.2020.116359.

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