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New isotopic constraints on age and magma genesis of an embryonic oceanic crust: The Chenaillet Ophiolite in the Western Alps

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Abstract

The Chenaillet Ophiolite is one of the best-preserved remnants of the Piemont-Liguria oceanic basin, a branch of the Central Atlantic that opened during the separation of Adria/Africa from Europe. Despite numerous studies of structure, petrology, geochemistry and isotope geochronology, the timing and genesis of various magmatic rocks within the Chenaillet Ophiolite are still controversial. We provide in this study integrated in situ analyses of zircon U–Pb age and O–Hf isotopes for the troctolite and albitite within the Chenaillet Ophiolite. Our new results indicate that the troctolite and albitite crystallized synchronously at ~165 Ma. Zircons from the troctolite have homogeneous Hf and O isotopic compositions, with εHf(T) = + 13.5 ± 1.0 (2SD) and δ18O = 5.4 ± 0.4‰ (2SD), indicating crystallization from magmas that were derived from a depleted, MORB-like mantle. The albitite zircons give consistent εHf(T) values (+ 13.0 to + 13.5) within errors with those of troctolite zircons, but variable δ18O values. The altered zircon domains have relatively low δ18O values of 4.7 ± 0.6‰ (2SD) due to subsolidus hydrothermal alteration, whilst the least-altered zircon domains give δ18Ozir values of 5.1 ± 0.4‰ (2SD), indistinguishable within errors with the troctolite zircons and the igneous zircons from the Mid-Atlantic and Southwest Indian Ridges gabbros, norites, and plagiogranites of modern oceanic crust. In situ zircon O–Hf isotopic data suggest that the troctolites and albitites are most likely cogenetic, with the albitites being formed by extreme fractional crystallization from the basaltic magma.

Based on our new age results and compilation of the reliable literature U–Pb age data, the ophiolitic gabbros from Eastern, Central and, Western Alps, Liguria and Corsica crystallized nearly synchronously at 158–166 Ma, suggesting a short life span of < 11 m.y. for the formation of the Piemont-Liguria oceanic domain, rather than a ~ 30 m.y. life span as previously thought. The Chenaillet ophiolite is likely a remnant of embryonic oceanic crust, rather than a piece of “mature” oceanic crust. Provided the spreading velocities of < 3 cm/yr full rate for magma-poor MOR sequences, the maximum width to the Piemont-Liguria oceanic floor would have been in the order of 300 km.
Keywords: Zircon U–Pb age; Zircon O–Hf isotopes; Plagiogranites; Chenaillet ophiolite; Alpine Tethys; Piemont-Liguria oceanic basin

1. Introduction

The ophiolites exposed in the Western Alps, the Ligurides, the Northern Apennines and Corsica (Fig. 1A) represent remnants of the Piemont-Liguria basin that opened as a branch of the Central Atlantic during the separation of Adria/Africa from Europe (Lemoine et al., 1987). The timing of the opening, nature of the crust and development and life-span of this embryonic oceanic domain are crucial to the understanding of the Mesozoic evolution of the Alpine Tethyan realms and, in a more general way, to the understanding of how continents break apart and new oceans form.

Fig. 1. : (A) Distribution of Ophiolite massifs in the Tertiary orogens from the Western Alps, the Ligurides, the Northern Apennines, and Corsica; (B) simplified geological map of the Chenaillet Ophiolite. Figures are modified after Manatschal and Müntener (2009) and Manatschal et al. (2011).

One of the best-preserved ophiolites in the Western Alps is the Chenaillet Ophiolite exposed along the French–Italian border south of Montgenèvre (Fig. 1B). The fact that this ophiolite escaped high-pressure metamorphism made it a preferred target to study the structure, petrology, geochemistry and isotope geochronology of a piece of ancient Jurassic oceanic crust (Bertrand et al., 1987, Caby, 1995 and Costa and Caby, 2001; Chalot-Prat et al., 2003; Chalot-Prat, 2005, Lewis and Snewing, 1980, Manatschal and Müntener, 2009, Manatschal et al., 2011 and Mével et al., 1978). In particular the present isotopic age data for the Chenaillet Ophiolite that cover a time span of ~ 30–50 m.y., which is by far the largest among the dated Jurassic ophiolites in the Western Alps, casting some problems for the interpretation of this ophiolite as part of a normal spreading oceanic crust. In order to test the validity of these data,
we carried out integrated *in situ* analyses of zircon U–Pb age and O–Hf isotopes for the troctolite and albitite from the Chenaillet Ophiolite. The aim of the study was to (1) provide robust age constraints on the timing of the opening of the oceanic domain preserved at the Chenaillet; (2) investigate the genetic relationships between the mafic and felsic magmatic rocks within the Chenaillet Ophiolite using integrated *in situ* zircon O–Hf isotopes. Our work indicates that the troctolites and albitites within the Chenaillet Ophiolite were not only contemporaneous, as they both crystallized at ~ 165 Ma, but also cogenetic. These new isotopic data have significant implications for the timing of formation and the magmatic evolution of the Piemont-Liguria ocean basin.

### 2. Geological background

The Chenaillet Ophiolite exposed south of Montgenèvre along the Franco–Italian border represents a klippe emplaced over the strongly deformed blueschist facies bearing Replatte-Lago Nero unit. The Chenaillet Ophiolite preserves primary oceanic contacts between serpentinized mantle and mafic intrusive rocks, sediments and MOR basalts (MORB) exposed over an area of about 25 km² (Fig. 1B). In contrast to most ophiolites from the Western Alps that experienced a high-pressure/low-temperature metamorphism and polyphase tectonic overprint during their emplacement within the Alpine nappe stack, the Chenaillet Ophiolite never exceed prehnite-pumpellyite facies and the Alpine tectonic overprint is minor (Manatschal et al., 2011, and references therein). Major lithologies forming the Chenaillet Ophiolite comprise, from bottom to top, serpentinized mantle rocks, intrusive mafic and minor felsic rocks, volcanic sequences and sediments. The mantle rocks consist of strongly serpentinized lherzolites and harzburgites, as well as minor pyroxenites, dunes and wehrlites (Bertrand et al., 1987).

Intrusive mafic rocks consist of troctolites, gabbros, diorites and dolerites. A thin sequence of layered troctolites and olivine-bearing gabbros occur within highly serpentinized mantle rocks, forming discrete bodies of less than 1 km in diameter and showing evidence of syn-magmatic deformation (Mével et al., 1978). The troctolites comprise plagioclase and olivine with minor clinopyroxene, whilst the gabbros consist of plagioclase and clinopyroxene, with minor olivine and opaque minerals. Costa and Caby (2001) interpreted the gabbros as remnants of an oceanic lower crust preserving a paleo-Moho with the mantle rocks. More recently, Manatschal et al. (2011) demonstrated that the mafic rocks form discrete bodies (< 1 km in diameter) intrusive into serpentinized mantle. In contrast to the work of Chalot-Prat (2005), we are not able to demonstrate that the gabbros are sill-like intrusions. They seem more to correspond to small magmatic bodies like those described from other ophiolites in the Alps (e.g. Desmurs et al., 2002). Dioritic rocks occur as veins, dykes with either diffuse or sharp boundaries with the gabbros (Costa and Caby, 2001 and Manatschal et al., 2011). Rock-forming minerals of the diorites include plagioclase, clinopyroxene and pargasitic amphibole. Dolerites and basaltic dykes outcrop within the sheared gabbros, particularly in the vicinity of high-angle faults, throughout the Chenaillet Ophiolite (Manatschal et al., 2011).

Albitites form dykes and sills that were emplaced within the serpentinized peridotites and the gabbros (Caby, 1995 and Chalot-Prat, 2005), and possibly within the pillow basalts as well (e.g. description of Chapelle, 1990). Albitites are the only felsic intrusive rocks identified within the Chenaillet Ophiolite. They are fine-grained, consisting predominantly of albite (> 90%), magmatic amphibole pseudomorphosed into tremolite–actinolite, and accessory minerals such as allanite, apatite and zircon (Costa and Caby, 2001).
A volcanic sequence overlies the mantle and intrusive mafic rocks, and locally also sediments (Manatschal et al., 2011). In contrast to Chalot-Prat (2005), we interpret the volcanic sequence that is locally up to 400 m thick to be formed by several magmatic events. This sequence consists of lava flows, pillow basalts, pillow breccias and hyaloclastites affinitive geochemically to the MORB (Bertrand et al., 1987 and Chalot-Prat, 2005). Sediments occur either between exhumed serpentinized mantle and intrusive mafic rocks and the volcanic sequence, or within the volcanic sequence (Manatschal et al., 2011).

A troctolite sample ALP3-6.3 (44°53′47″N, 06°44′10″E) and an albitite sample ALP3-6.1 (44°53′40″N, 06°43′53″E) were collected from the southwestern Chenaillet Ophiolite complex (Fig. 1B) for in situ zircon U–Pb age and O–Hf isotope study.

3. Analytical methods

3.1. Zircon U–Pb dating

Zircon concentrates were separated from a ca. 5-kg of troctolite and a ca. 2-kg of albitite using standard density and magnetic separation techniques. Zircon grains, together with standard zircons TEMORA 2, 91500, Penglai and an in-house standard zircon Qinghu, were mounted in an epoxy mount that was then ground and polished to section the crystals in half for analysis. Zircons were documented with transmitted and reflected light photomicrographs as well as cathodoluminescence (CL) images to reveal their internal structures, and the mount was vacuum-coated with high-purity gold prior to SIMS analyses.

Measurements of U, Th and Pb isotopes were conducted using a Cameca IMS 1280 SIMS at the Institute of Geology and Geophysics, Chinese Academy of Sciences in Beijing. Analytical procedures are similar to those reported by Li et al. (2009). The primary O$_2^-$ ion beam spot is about 20 × 30 μm in size. Positive secondary ions were extracted with a 10 kV potential. In the secondary ion beam optics, a 60 eV energy window was used, together with a mass resolution of ca. 5400 (at 10% peak height), to separate Pb$^+$ peaks from isobaric interferences. A single electron multiplier was used in ion-counting mode to measure secondary ion beam intensities by peak jumping mode. Analyses of the standard zircon TEMORA 2 were interspersed with unknown grains. Each measurement consists of 7 cycles. Pb/U calibration was performed relative to zircon standard TEMORA 2 ($^{206}$Pb/$^{238}$U age = 417 Ma, Black et al., 2004); U and Th concentrations were calibrated against zircon standard 91500 (Th = 29 ppm, and U = 81 ppm, Wiedenbeck et al., 1995). Measured compositions were corrected for common Pb using non-radiogenic $^{204}$Pb, and an average of crustal composition (Stacey and Kramers, 1975) is used for the common Pb. Data reduction was carried out using the Isoplot/Ex v. 2.49 program (Ludwig, 2001). In order to monitor the external uncertainties of SIMS U–Pb zircon measurements, an in-house standard zircon Qinghu was alternately analysed as an unknown together with other unknowns. Twenty measurements on Qinghu zircon (Appendix Table 1) yield a Concordia age of 160 ± 1 Ma (2SE, MSWD of concordance = 0.92), which is in good agreement within errors with the recommended U–Pb age of 159.5 ± 0.2 Ma (Li et al., 2009). Zircon U–Pb age data are presented in Appendix Table 1.

3.2. Zircon oxygen isotopes

Zircon oxygen isotopes were measured using the same Cameca IMS 1280 SIMS, with analytical procedures similar to those reported by Li et al. (2010a). Oxygen isotopic analyses
were obtained on the same spots of zircon grains that were previously analyzed for U–Pb age determinations. After U–Pb dating, the sample mount was re-ground and re-polished to ensure that any oxygen implanted in the zircon surface from the O$_2^-$ beam used for U–Pb determination was removed. The Cs$^+$ primary ion beam was accelerated at 10 kV, with an intensity of ~ 2.5 nA corresponding to a beam size of ~ 10 μm in diameter. The normal incidence electron flood gun was used to compensate for sample charging. Negative secondary ions were extracted with a ~ 10 kV potential. Oxygen isotopes were measured in multi-collection mode using two off-axis Faraday cups. The mass resolution used to measure oxygen isotopes was ca. 2500 (at 10% peak height). Measured $^{18}\text{O}/^{16}\text{O}$ ratios were normalized by using Vienna Standard Mean Ocean Water compositions (VSMOW, $^{18}\text{O}/^{16}\text{O} = 0.0020052$), and then corrected for the instrumental mass fractionation factor (IMF) using zircon 91500 as a reference standard with δ$^{18}\text{O}$ value of 9.9‰ (Wiedenbeck et al., 2004). Fifteen measurements of Penglai zircon standard during the course of this study yield a weighted mean of δ$^{18}\text{O} = 5.28 \pm 0.34$‰ (2SD), which is consistent within errors with the reported value of 5.31 ± 0.10‰ (Li et al., 2010b). Zircon oxygen isotopic data are listed in Appendix Table 2.

3.3. Zircon Lu–Hf isotopes

*In situ* zircon Lu–Hf isotopic analysis was carried out on a Neptune multi-collector ICP-MS equipped with a Geolas-193 laser-ablation system at IGGCAS, and the analytical procedures were similar to those described by Wu et al. (2006). Lu–Hf isotopic analyses were obtained on the same zircon grains that were previously analyzed for U–Pb and O isotopes, with ablation pits of 63 μm in diameter, ablation time of 26 seconds, repetition rate of 10 Hz, and laser beam energy density of 10 J/cm$^2$. During laser ablation analyses, the isobaric interference of $^{186}$Lu on $^{176}$Lu Hf is negligible due to extremely low $^{176}$Lu Lu/$^{177}$Lu Hf in zircons (mostly < 0.01). Isobaric interference of $^{176}$Lu Yb on $^{176}$Lu Hf is corrected using independent mass bias factors for Hf and Yb for correction, since the contribution of $^{176}$Lu Yb to $^{176}$Lu Hf could affect the accuracy of the measured $^{176}$Lu Hf/$^{177}$Lu Hf ratio. The mean 173 Lu Yb/$^{172}$Lu Yb ratio of the individual spots was used to calculate the fractionation coefficient ($\beta_{Yb}$) and the contribution of $^{176}$Lu Yb to $^{176}$Lu Hf by applying ratios of $^{176}$Lu Yb/$^{172}$Lu Yb = 0.5887 and $^{176}$Lu Yb/$^{172}$Lu Yb = 0.73925 (Wu et al., 2006). Measured $^{176}$Lu Hf/$^{177}$Lu Hf ratios were normalized to $^{176}$Hf/$^{177}$Hf = 0.7325. The determined $^{176}$Lu Hf/$^{177}$Lu Hf value of 0.282497 ± 0.000018 (2SD, n = 20) for zircon standard Mud Tank during the course of this work is in agreement within errors with the recommended value of 0.282507 ± 0.000006 by solution MC-ICP-MS measurements after chemical separation (Woodhead and Hergt, 2005). The reported $^{176}$Lu Hf/$^{177}$Lu Hf value is adjusted relative to the Mud Tank standard value of 0.282507, and the results are listed in Appendix Table 2.

Because zircons from the albitite sample ALP3-6.1 have exceptionally high $^{176}$Lu Yb/$^{177}$Lu Hf (> 0.3) that would result in inaccurate isobaric interference correction for the measured $^{176}$Lu Hf/$^{177}$Lu Hf ratio, we carry out solution MC-ICP-MS measurements for two zircon fractions from this sample. After decomposition of zircon grains, the sample solution was split into two parts, ca. 1/3 of the solution was used for determination of Yb, Lu and Hf concentrations, and the remaining 2/3 solution for MC-ICP-MS measurements of $^{176}$Lu Hf/$^{177}$Lu Hf after chemical purification. The procedures for the separation of Hf from Zr and REE were similar to those reported by Goolaeers et al. (2004). The determined $^{176}$Lu Hf/$^{177}$Lu Hf value is 0.282172 ± 0.000010 (2SE) for JMC-475 standard during the course of this work. The reported $^{176}$Lu Hf/$^{177}$Lu Hf values were adjusted relative to a JMC-475 $^{176}$Lu Hf/$^{177}$Lu Hf ratio
of 0.282160. The Lu–Hf isotopic data obtained by solution MC-ICP-MS measurements is presented in Appendix Table 2.

4. Analytical results

4.1. Zircon U–Pb age

4.1.1. Troctolite sample ALP3-6.3

Zircons from this sample are transparent, 50–100 μm in length, and have aspect ratios of ~ 1:1 to 2:1. In CL images they are quite homogeneous, with weak, broad zonings (Fig. 2). Fifteen analyses were conducted on 15 zircons during a single analytical session (Appendix Table 1). These zircons have moderate U (70–509 ppm) and Th (47–421 ppm) contents, and Th/U ratios between 0.51 and 1.38. Common Pb is low; value for $f_{206}$ (the proportion of common $^{206}$Pb in total measured $^{206}$Pb) is mostly lower than 1.0%, apart from two analyses (spots 5 and 6) that have slightly higher $f_{206}$ value of 1.2–1.5%. All the measured Pb/U ratios are concordant within analytical errors, yielding a Concordia age of 165 ± 1 Ma (2σ, MSWD of concordance = 0.01) (Fig. 2). This age is interpreted as the best estimate of crystallization age of the troctolite.

![Fig. 2. U–Pb Concordia diagram for zircons from troctolite sample ALP3-6.3 and CL images for representative zircons. The ellipses in the CL images represent the spots of SIMS U–Pb and O isotope analyses. The numbers on the left and right are the U–Pb dates (Ma) and $\delta^{18}$O values, respectively. The white bars are 50 μm in length for scale.](image)

4.1.2. Albitite sample ALP3-6.1

Zircons are semi-transparent, stubby, 80–150 μm in length, and have aspect ratios of ~ 1:1 to 1.5:1. Microfractures are visible in most zircons. In CL images most zircons are characterized by magmatic concentric zonings, associated with, more or less, chaotic/botryoidal domains (Fig. 3A) that are most likely attributed to alteration (Corfu et al., 2003 and Grimes et al.,...
Twenty-one analyses were conducted on 21 zircons during a single analytical session (Appendix Table 1). They have moderate U (66–399 ppm) and Th (19–253 ppm) contents, with Th/U ratios between 0.20 and 0.94. Common Pb is highly variable, with value of $f_{206}$ ranging from 0.16% to 25.6%. The common Pb-uncorrected $^{206}_{238}$Pb/$^{238}$U and $^{207}_{206}$Pb/$^{206}$Pb ratios define a discordia line on the inverse (Tera-Wasserburg) U–Pb concordia plot (Fig. 3A), with a lower-intercept age of 165 ± 2 Ma (2σ, MSWD = 1.3). It is noteworthy that nine of twenty-one analyses are conducted on the altered zircon domains, providing imprecise $^{206}_{238}$Pb/$^{238}$U dates ranging from 163 Ma to 184 Ma, with uncertainties up to ± 16 Ma (1σ) (Appendix Table 1), due to high proportion (3.45–25.6%) of common Pb. The remaining twelve analyses are made on the least-altered domains (Fig. 3B) that are relatively low in common Pb ($f_{206}$ value = 0.16–2.49%). Their Pb/U ratios are concordant within analytical errors, yielding a U–Pb Concordia age of 165 ± 2 Ma (2σ, MSWD of concordance = 0.53) (Fig. 3B). We interpret the age of 165 ± 2 Ma as the best estimate of the timing of crystallization of albitite sample ALP3-6.1.
Fig. 3. U–Pb Concordia diagram for zircons from albitite sample ALP3-6.1 and CL images for representative zircons. (A) All twenty-one analyses; (B) twelve analyses for the least-altered zircons that are shown are low in common Pb. The ellipses in the CL images represent the spots of SIMS U–Pb and O isotope analyses. The numbers on the left and right are the U–Pb dates (Ma) and $\delta^{18}$O values, respectively. The white bars are 50 μm in length for scale.

4.2. *In situ* zircon O–Hf isotopes

*In situ* O–Hf isotopic measurements are conducted on fifteen zircons from the troctolite sample ALP3-6.3. Values of the measured $\delta^{18}$O$_{zr}$ are between 5.1‰ to 5.8‰ (Appendix Table 2), forming a Gaussian distribution. Thus, this troctolite is homogeneous in oxygen
isotopes, with a mean $\delta^{18}O$ value of $5.4 \pm 0.4\%$ (2SD) (Fig. 4A). Fifteen analyses yield $^{176}\text{Yb}/^{177}\text{Hf}$ ratios between 0.023 and 0.117 (Appendix Table 2), which are reasonably low for accurate isobaric interference correction. The analyses give fairly homogeneous $^{176}\text{Hf}/^{177}\text{Hf}$ ratios between 0.283015 and 0.283079 (Appendix Table 2), with an average of $0.283058 \pm 0.000030$ (2SD), corresponding to $\varepsilon\text{Hf}(t) = +13.5 \pm 1.0$ (2SD). In order to confirm the LA-MC-ICPMS analyses, we conduct on solution MC-ICP-MS measurement for one zircon fraction after decomposition of zircon grains and chemical purification (Appendix Table 2). The solution analysis yields $^{176}\text{Yb}/^{177}\text{Hf} = 0.0614$ and $^{176}\text{Hf}/^{177}\text{Hf} = 0.283061 \pm 0.000005$ (2SE). The calculated $\varepsilon\text{Hf}(t)$ is +13.7, which is in good agreement within errors with the average ($+13.5 \pm 1.0$) of LA-MC-ICPMS analyses.

![Histograms of zircon $\delta^{18}O$ values for the troctolite (ALP3-6.3) and albitite (ALP3-6.1) samples.](image)

Fig. 4: Histograms of zircon $\delta^{18}O$ values for the troctolite (ALP3-6.3) and albitite (ALP3-6.1) samples.
Twenty-one zircons from the albitite sample ALP3-6.1 were analyzed for oxygen isotopes. The measured δ18O values range from 4.2‰ to 5.4‰ (Fig. 4B). It is noted, despite being indistinguishable within analytical errors, the altered zircon domains have slightly lower δ18O values between 4.2‰ to 5.0‰ (averaging at 4.7 ± 0.6‰, 2SD) than the least-altered domains that give δ18O values between 4.8‰ to 5.4‰ (averaging at 5.1 ± 0.4‰, 2SD). Overall, the measured δ18O values for the troctolite zircons and the least-altered zircon domains from the albitite are consistent within analytical uncertainty with those of igneous zircons from the Mid-Atlantic and Southwest Indian Ridges gabbros, norites, and plagiogranites of modern oceanic crust (5.3 ± 0.8‰, Cavosie et al., 2009; 5.2 ± 0.5‰, Grimes et al., 2011) and with igneous zircons in high-temperature equilibrium with primitive magma compositions of the mantle (5.3 ± 0.6%, Valley et al., 1998, Valley et al., 2005 and Valley, 2003). These albitite zircons have exceptionally high 176Yb/177Hf ratios between 0.320 and 0.861 (Appendix Table 2) and highly variable 176Hf/177Hf ratios between 0.283139 and 0.283547. The measured 176Hf/177Hf ratios, corresponding to εHf(t) = + 15.6 to + 28.9, are not only much higher than those of troctolite sample ALP3-6.3, but also correlated positively with the measured 176Yb/177Hf ratios (not shown). Such anomalously high 176Hf/177Hf ratios are most likely attributed to inaccurate isobaric interference correction of 176Yb on 176Hf. To clarify this, we conducted solution MC-ICP-MS measurements for two zircon fractions after decomposition of zircon grains and chemical purification. These two zircon fractions yield the measured 176Yb/177Hf = 0.1080 and 0.3216 and 176Hf/177Hf = 0.283060 ± 0.000009 and 0.283064 ± 0.000007 (2SE), respectively (Appendix Table 2). The calculated εHf(t) values are between + 13.5 and + 13.0, consistent within uncertainty with the average of the troctolite zircons (+ 13.5 ± 1.0).

5. Discussion

5.1. Formation age for the Chenaillet ocean-floor sequence

Costa and Caby (2001) and Chalot-Prat et al. (2003) carried out systematic isotopic age determinations for various rock types of the Chenaillet Ophiolite. They reported Sm–Nd whole-rock isochron ages of 198 ± 22 Ma (Costa and Caby, 2001) and 142 ± 23 Ma (Chalot-Prat et al., 2003) for the gabbros, and an ID-TIMS U–Pb zircon age of 156 ± 3 Ma and 148 ± 2 Ma for a leucodioritic vein and an albitite lens (Costa and Caby, 2001), respectively. These Sm–Nd and U–Pb zircon ages were interpreted as the timing of crystallization of the dated rocks. These results implied a multi-stage magmatic process that occurred over ~ 30–50 m.y., which is unusual for true oceanic sequences.

However, Manatschal et al. (2011) questioned the validity of the Sm–Nd isochron age of 198 ± 22 Ma for the gabbros, because (1) this Sm–Nd age is clearly older than the U–Pb age of 156 ± 3 Ma for the leucodiorites that are thought to be genetically related with the gabbros (Desmurs et al., 2002), and (2) the Sm–Nd age of 198 ± 22 Ma for the Chenaillet gabbros would invoke their emplacement during continental rifting, prior to first mantle exhumation at 165–160 Ma in the Piemont-Liguria domain (Manatschal and Müntener, 2009). In addition, we noted that five zircon fractions from a leucodiorite sample (MG3511) reported by Costa and Caby (2001) are variably discordant in U–Pb isotopes. While four of the five analyzed fractions yield an average U–Pb age of 156 ± 3 Ma, the fifth fraction gives significant older U–Pb age of ~ 170 Ma. This older zircon fraction was interpreted by the authors as inheritance. However it is quite unusual that such a significant portion (~ 20%) of inherited
zircons was present during the formation of the diorite. While the U–Pb age of 148 ± 2 Ma for the albitite (Costa and Caby, 2001) is consistent within errors with the Sm–Nd isochron date of 142 ± 23 Ma (Chalot-Prat et al., 2003), it is ~30 to 50 m.y. and ~3 to 13 m.y. younger than the U–Pb ages for the gabbro and leucodiorite (Costa and Caby, 2001), respectively. Costa and Caby (2001) and Manatschal et al., (2011) interpreted that the albitites might correspond to a late, possibly off-axis magmatic event associated with the very last stage of oceanic deformation, whereas others considered that the albitites might represent highly evolved liquids differentiated from a gabbroic or dioritic magma (Bertrand et al., 1987, Caby, 1995 and Desmurs et al., 2002).

Our new SIMS U–Pb zircon dating results demonstrate that the troctolite and albitite within the Chenaillet Ophiolite crystallized synchronously at ~165 Ma. Zircon morphology and internal structure of the troctolite (Fig. 2) are well consistent with those of zircons crystallized from mafic magmas (Corfu et al., 2003). Furthermore, troctolite zircons give typical mantle-like δ18O (5.4 ± 0.4‰) and highly positive zircon εHf(t) (+13.5 ± 1.0), suggesting crystallization from magmas that were produced by partial melting of a depleted, MORB-like mantle.

Zircons from the albitite sample ALP3-6.1 have more complex internal textures than those from the troctolite; they mostly show microfracture and alteration to varying degrees (Fig. 3). Despite careful selection of the targeting domains for SIMS dating at a scale of 20 × 30 μm, nine of the twenty-one analyses still have variably high proportion (3.45–25.6%) of common Pb as a result of alteration and/or metamictization. To our knowledge, such altered zircons are difficult to be precisely dated by using conventional ID-TIMS method that requires a few milligrams of zircons for analysis. Similar to our imprecise dating results for the altered zircon domains by SIMS, the ID-TIMS zircon U–Pb analyses of the albitite sample MG542 also give high proportion (18–27%) of common Pb (Costa and Caby, 2001), making the reported U–Pb age of 148 ± 2 Ma problematic. Contrarily, taking the advantage of SIMS microanalytical technique, we are able to precisely date the albitites by targeting zircon domains that show weak and/or negligible alteration at a small scale of 20 × 30 μm. Thus, our new SIMS U–Pb zircon dating result of 165 ± 2 Ma constrained by 12 analyses of the least-altered zircon domains is reliable, providing the best estimate of the crystallization age of the albitite.

Troctolites and albitites bracket the intrusive magmatism of the Chenaillet Ophiolite. Our new SIMS zircon U–Pb dating results indicate that these intrusive rocks are coeval, placing a robust constraint on the formation age of ~165 Ma for the Chenaillet Ophiolite.

5.2. Constraints on the albitite genesis

Plagiograrnites, referred collectively to the suite of felsic plutonic rocks (including diorite, quartz diorite, tonalite, trondjhemite and albitite/anorthosite) found in modern ocean crust and ophiolites (e.g. Coleman and Donato, 1979), usually formed as the most differentiated and latest-stage magmas in ocean crust. Despite volumetrically minor, the genesis of oceanic plagiograrnites is controversial, with different processes being involved, including (1) extreme fractional crystallization of MORB, (2) liquid immiscibility with Fe-enriched tholeiitic melts, and (3) hydrous partial melting of oceanic gabbros (see review of Koepke et al., 2007). In contrast to the Platta ophiolite in the Central Alps where the albitites are in close association with ferrogabbros (Desmurs et al., 2001 and Schaltegger et al., 2002), the Chenaillet albitites do not show direct relationships with the ferrogabbros in the field. Process 1 and process 3
have been proposed for diorites and albitites from the Chenaillet Ophiolite (e.g., Bertrand et al., 1987, Caby, 1995 and Costa and Caby, 2001). It should be noted that element geochemistry and radiogenic isotopic systems such as Sm–Nd and Lu–Hf in which the parental isotopes have half-lives exceeding billions of years are not able to clearly distinguish the process 1 from the process 3.

Alternatively, zircon oxygen isotopes of the plagiogranites could provide an effective approach to this problem. This is because (1) zircons, which are usually retentive of the magmatic oxygen isotopes, in equilibrium with mantle-derived magmas have very consistent $\delta^{18}O_{Zir}$ value of 5.3 ± 0.6‰ (2σ) (Valley, 2003, Valley et al., 1998 and Valley et al., 2005). This mantle zircon value is insensitive to magmatic differentiation, because the increase in bulk rock $\delta^{18}O$ value is compensated by an increase in zircon/liquid $\delta^{18}O$ fractionation, (2) zircons are highly resistant to alteration except for extreme cases (e.g., hydrothermal alteration and metamictization), and commonly preserve primary magmatic $\delta^{18}O$ values (Grimes et al., 2011, Page et al., 2007, Valley, 2003 and Valley et al., 2005). Identification of altered and/or metamict zircons are readily feasible by combined determination of zircon internal textures in CL images and correlation of age concordance, Pb-loss and/or common-Pb proportion to $\delta^{18}O_{Zir}$ value (e.g., Booth et al., 2005 and Corfu et al., 2003), and (3) hydrothermal alteration of oceanic gabbros by infiltration of seawater leads to $\delta^{18}O$ value higher than 6‰ at temperature below 200–250 °C, but lower than 5‰ above 200–250 °C (e.g., Alt and Bach, 2006, Bindeman et al., 2005 and Muehlenbachs, 1986). Thus, plagiogranites formed by the process 1 and 3 would have significantly different $\delta^{18}O_{Zir}$ values, i.e., plagiogranites formed by process 1 should have mantle-like $\delta^{18}O_{Zir}$ values, whereas, those formed by process 3 would have $\delta^{18}O_{Zir}$ values either lower or higher than the mantle values depending on temperature of hydrothermal alteration of the gabbro protoliths.

Zircons from the albitite sample ALP3-6.1 show features of variable alteration, suggesting that the albitite is overprinted by subsidious hydrothermal alteration. This is consistent with the pseudomorphosed amphibole into secondary tremolite/actinolite. Nine altered zircons show $\delta^{18}O_{Zir}$ values of 4.2‰ to 5.0‰ (averaging at 4.7 ± 0.6‰, 2SD) due to high-temperature alteration. Similar phenomenon has been reported for the altered ocean crust zircons that give $\delta^{18}O_{Zir}$ values from 0.0 to 5.0‰ (Fu et al., 2012; Grimes et al., 2011). On the contrary, the least-altered zircon domains from the albitite give a mean $\delta^{18}O_{Zir}$ value of 5.1 ± 0.4‰ (2SD) that is well consistent with the average $\delta^{18}O_{Zir}$ value of 5.2 ± 0.5‰ determined for 107 normal magmatic zircons from plagiogranites (Grimes et al., 2011), and also indistinguishable within uncertainties from the average value (5.4 ± 0.4‰) for the troctolite sample ALP3-6.3 and the range of 5.3 ± 0.6‰ observed for zircons in high-temperature equilibrium with primitive mantle (Valley, 2003, Valley et al., 1998 and Valley et al., 2005). Therefore, the least-altered albitite zircons do not show any detectable seawater signature, indicating that the albitite should have formed prior to significant interactions between the ocean crust and seawater. In other words, the albitite is most likely generated by the aforementioned process 1. If the albitites were formed by hydrous partial melting of oceanic gabbros (i.e. the process 3), their zircon oxygen isotopic features request that the gabbros should preserve MORB-like $\delta^{18}O$ value at the time of melting. This would invoke very low degrees of hydrothermal alteration of the protolith gabbros prior to melting and zircon crystallization. In consideration of the synchronous crystallization at ~ 165 Ma and consistent zircon O–Hf isotopes of the troctolite and albitite, we suggest that the troctolites and albitites are most likely co-genetic, and the albitites were formed by extreme fractional crystallization from a gabbroic magma (Bertrand et al., 1987). This interpretation is consistent with the identical zircon $\varepsilon$Hf(T) values for the
troctolite (+ 13.5 ± 1.0) and albite (+ 13.2 ± 0.2) and is also similar to the processes described from other Alpine ophiolites (Desmurs et al., 2002 and Schaltegger et al., 2002).

While the process 2 (liquid immiscibility with Fe-enriched tholeiitic melts) is not favoured for the Chenaillet albitites in terms of the field relationship between the albitites and ferrogabbros, our isotopic study alone could not entirely rule out this possibility. Examination of this possibility awaits further petrological and geochemical investigations.

5.3. Implications for age and nature of the crust and width of the Piemont–Liguria domain

The discovery of symmetric seafloor spreading recorded by magnetic anomalies in oceanic domains enabled to propose a simple relationship between a location in the ocean, its distance from the mid ocean ridge and its spreading rate. Applying this concept to ophiolites enables to determine a location relative to the edge of the former ocean assuming that age of accretion, i.e. age of the formation of the mafic crust, and continental breakup are known and off-axis magmatic activity can be excluded. Based on this concept, ages obtained from ophiolites can be used to restore ancient oceanic domains.

In the case of the Alpine Tethys, numerous radiometric data have been reported since 1980's, providing a ~ 30 m.y. life span for the formation of the Piemont-Liguria oceanic basin (e.g., Costa and Caby, 2001). A compilation of zircon U–Pb age data from the ophiolitic gabbros and plagiogranites from the Central and Western Alps, Liguria and Corsica is presented in Fig. 5. Sm–Nd isochron and Ar/Ar or K–Ar age data are not included, because they are either imprecise or susceptible to Alpine overprinting, and do therefore not necessarily date the age of accretion of a given piece of ophiolite.
Fig. 5. Compilation of U–Pb age determinations for mafic (solid circles) and plagiogranitic (open circles) rocks from the Piemont-Liguria ocean. Data sources: Bill et al. (1997), Borsi et al. (1996), Costa and Caby (2001), Kaczmarek et al. (2008), Liati et al. (2005), Ohnenstetter et al. (1981), Rossi et al. (2002), Rubatto and Scambelluri (2003), Rubatto et al. (1998), Rubatto et al. (2008), Schaltegger et al. (2002), Sticki et al. (2003); this study.

The U–Pb zircon dates for the plagiogranites are highly variable ranging from 142 to 169 Ma (Borsi et al., 1996, Costa and Caby, 2001, Ohnenstetter et al., 1981, Rossi et al., 2002, Rubatto and Scambelluri, 2003, Rubatto et al., 2008, Schaltegger et al., 2002 and Sticki et al., 2003). While the plagiogranites from the Platta, Lanzo and Chenaillet Ophiolites are dated at 161–165 Ma, coeval with the co-exiting MOR gabbros (Rubatto et al., 2008 and Schaltegger et al., 2002; this study), others are either poorly dated or differ in age significantly from the co-existing gabbros. Because most plagiogranite zircons are subject to oceanic alteration and/or high-pressure metamorphism, their crystallization ages await further investigations.

In contrast to the highly variable age data for the plagiogranite, all U–Pb zircon ages for the ophiolitic gabbros are between 166 and 155 Ma (Bill et al., 1997, Kaczmarek et al., 2008, Liati et al., 2005, Rubatto and Scambelluri, 2003 and Rubatto et al., 1998; Schaltegger et al., 2002; this study). It is noted that two metagabbros from Antrona gave younger U–Pb ages of 155–156 Ma (Liati et al., 2005) than others from the Western Alps. Zircons from these two metagabbros are strongly affected by the Alpine overprint, and the “co-magmatic domains” have SHRIMP $^{206}{_{\text{Pb}}}/^{238}{_{\text{U}}}$ dates vary from 129 to 163 Ma (Liati et al., 2005). Average of such highly variable dates could not give a statistically meaningful age. We consider that the maximum $^{206}{_{\text{Pb}}}/^{238}{_{\text{U}}}$ dates of 160 ± 3 Ma and 163 ± 2 Ma (1σ), which might have been subjected to the least radiogenic Pb loss, possibly provide the lower age limits for the two metagabbros, consistent with the U–Pb zircon ages for the other gabbros from the Western Alps. Apart from the two Antrona metagabbros, all other ophiolitic gabbros have crystallization ages clustering around 158–166 Ma. These consistent U–Pb age data thus suggest a life span of ≤ 8 m.y. for the formation of the Piemont-Liguria oceanic basin, rather than a ~ 30 m.y. life span as previously thought. Taking into account an external age uncertainty of about ± 2–3 Ma for these middle Jurassic rocks by SIMS and/or LA-ICPMS U–Pb zircon dating technique, the maximum life span documented by the ophiolitic gabbros is about 11 m.y. This short time range also corresponds to the ages of the Radiolarian cherts that are the oldest sediments overlying the ophiolites (Bill et al., 2001). Assuming spreading velocities that are < 3 cm/yr full rate, commonly considered as maximum values for magma-poor MOR sequences corresponding to those described from the Alps (Cannat et al., 2009), the maximum width to the paleo-ocean floor would have been in the order of 300 km (Fig. 6). This width also corresponds to the length of the slab imaged beneath the Alps and commonly interpreted as the subducted remnant of the former Piemont-Liguria ocean (Kissling et al., 2006). The available age data also do not show significant geographic difference for gabbros across the Piemont-Liguria oceanic basin. For instance, the ~ 165 Ma Chenaillet Ophiolite, is nearly synchronous with gabbros from the ~ 166 Ma Gets and ~ 161 Ma Platta nappes that intruded into subcontinental mantle and are associated with continent derived extensional allochthons, characteristic for ocean-continent transition zone (Bill et al., 1997, Kaczmarek et al., 2008 and Manatschal and Müntener, 2009; Schaltegger et al., 2002). Therefore, the Chenaillet Ophiolite may not represent a piece of “mature” oceanic crust, but rather represent a remnant of embryonic oceanic crust surrounded by a hyper-extended domain comparable to the present northern Red Sea (Manatschal and Müntener, 2009). However, such embryonic oceanic domains are not necessarily symmetric and the processes controlling their formation...
are yet little understood. The study of the Chenaillet Ophiolite shows that this domain was characterized by the scarcity of mafic plutonic rocks and the occurrence of exhumed mantle, the absence of a sheeted dike complex linking the intrusives with the volcanic pillow lavas, and the occurrence of top-basement detachment faults associated with tectono-sedimentary breccias, also referred to as oceanic core complexes (e.g. Manatschal et al., 2011).

Fig. 6. Conceptual 3D model showing the temporal and spatial evolution of the Alpine domain from an embryonic oceanic stage (Early Cretaceous (~100 Ma)) to the onset of collision (Middle Eocene (~45 Ma)). The age of the dated rocks from the Piemont-Liguria domain range between 166 and 158 Ma, which gives, including error bars, a maximal duration of seafloor accretion of 11 m.y. Assuming maximum velocity rates of 3 cm/yr, corresponding to the upper bound of accretion rates associated with mantle exhumation in modern spreading systems, would result in a maximum width of about 300 km for the Piemont-Liguria basin. 300 km corresponds also to the length of the slab observed today beneath the Alps (3D block is modified after Beltrando et al., 2010).

6. Conclusions

In conclusion, our new SIMS U–Pb zircon dating results demonstrate that the troctolites and albitites within the Chenaillet Ophiolite crystallized synchronously at ~165 Ma. In situ analyses of zircon O–Hf isotopes indicate that the troctolites and albitites are cogenetic, the albitites were formed by extreme fractional crystallization from a MOR basaltic magma. Our
new dating results and compilation of the reliable literature U–Pb age data suggest a short life span of less than 11 m.y. for the formation of the Piemont-Liguria oceanic domain with a maximum width in the order of 300 km. The Chenaillet ophiolite may represent a remnant of embryonic oceanic crust.

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