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#### Mantle metasomatic influence on water contents in continental lithosphere:

### new constraints from garnet pyroxenite xenoliths (France & Cameroon

volcanic provinces)

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#### Abstract

Quantifying water contents in the lith sp...ric mantle is key to our understanding of global geodynamics, mantle composition,  $\infty$  d related physical properties. Most mantle lithologies (peridotite) contain little wate: (~50 ppm), but petrological heterogeneities such as pyroxenites are more hydrous (~300 ppm) relative to the mantle rocks. Pyroxenites also melt at lower temperatures than period tites and are thus important to magma genesis. Thus, quantifying pyroxenite water contents provides new information on the distribution of water in the mantle. Here, we present phase-specific FTIR measurements of the water contents in pyroxenite mantle xenoliths from two continental lithospheric domains that experienced intense metasomatism: the French Massif Central (FMC, France) and the Adamawa Volcanic Plateau (AVP, Cameroon). The AVP garnet pyroxenites are more hydrated ([H<sub>2</sub>O]<sub>Clinopyroxene</sub> = 124–155 ppm; [H<sub>2</sub>O]<sub>Garnet</sub> < 0.5 ppm) than FMC ones ([H<sub>2</sub>O]<sub>Clinopyroxene</sub> = 112–465 ppm; [H<sub>2</sub>O]<sub>Orthopyroxene</sub> = 61–104 ppm; [H<sub>2</sub>O]<sub>Garnet</sub> < 0.5 ppm). These water concentrations are homogenous at the grain and correlate with equilibrated major element concentrations, indicating that they are representative of lithospheric water, although

the FMC pyroxenites were dehydrated during metasomatism by a carbonatitic fluid (based on the correlation between  $La_N/Sm_N$  and Ti/Eu ratios); the water contents of AVP pyroxenites were likely not affected by metasomatism. FMC pyroxenites show peculiar FTIR spectra that may reflect the preferential dehydration of specific sites in the pyroxene structure. In both regions, metasomatism modified the light rare Earth element contents (e.g., Ce) of the pyroxenites, resulting in highly variable H<sub>2</sub>O/Ce ratios. Therefore, we conclude that the utility of the H<sub>2</sub>O/Ce ratio to identify the involvement of pyroxenites in magmas genesis is limited. **Keywords:** nominally anhydrous minerals; garnet pyroxenite<sup>-</sup>, n<sub>2</sub> droxyl; mantle

metasomatism; Adamawa Volcanic Plateau; French Massif Cer tral

#### 1. Introduction

Water in the silicate Earth is mainly concentrated in the mantle, which contains a mass of water equivalent to 4–10 times that in the oceans (Marty et al., 2016; Peslier et al., 2017). Water in the mantle can be hosted an aydrous minerals such as amphiboles or micas (e.g., up to 2 wt% H<sub>2</sub>O in amphibole; Bizlimis and Peslier, 2015; Green, 2015) or as hydrogen point defects in nominally anhydrous unnerals (NAMs) due to the replacement of cations in their mineral structures by hydrogen atoms (Demouchy and Bolfan-Casanova, 2016). Although NAMs do not company free molecular H<sub>2</sub>O, their hydrogen concentrations are generally expressed as ppin H<sub>2</sub>O equivalent. In the upper mantle rocks (peridotites and pyroxenites), different NAMs have various storage capacities: garnet is generally anhydrous and contains little water (Rossman and Smyth, 1990; Bell and Rossman, 1992; Matsyuk et al., 1998; Beran and Libowitzky, 2006), as does olivine (up to 40 ppm H<sub>2</sub>O; Yu et al., 2011; Doucet et al., 2014; Hui et al., 2015), whereas clinopyroxene and orthopyroxene can contain 100–400 and 50–200 ppm H<sub>2</sub>O, respectively (Fig. 1, Table 1; Peslier et al. (2002, 2012), Peslier (2010), Xia et al. (2010, 2013), Yu et al. (2011), Hao et al. (2012, 2014, 2016a, 2016b, 2018), Doucet et al. (2014), Warren and Hauri (2014), Bizimis and Peslier (2015), Demouchy

et al. (2015), Li et al. (2015, 2018), and Peslier and Bizimis (2015), Kilgore et al. (2018, 2020), Schaffer et al. (2019), Ashley et al. (2020)).

Despite its low concentrations in the mantle, water has many effects on mantle properties: it decreases mineral electrical resistivities (Selway, 2014), olivine aggregate viscosities (Hirth and Kohlstedt, 1996), and peridotite melting temperatures (Asimow et al., 2004; Green et al., 2010; Kovács et al., 2012). Water also acts as a lubricant for plate tectonics by weakening olivine (Karato, 1990; Hirth and Kohlstedt, 1996<sup>.</sup> Fei et al., 2013; Peslier and Bizimis, 2015). Therefore, it is necessary to quantify the distribution and concentrations of water in the mantle to better understand geodynamics at the gle bal scale.

Water is heterogeneously distributed in man<sup>1</sup> e d, mains. The mantle source of midocean ridge basalts (MORBs), i.e., the depleted MCRB mantle (DMM), contains a relatively uniform 70–160 ppm H<sub>2</sub>O (Salters and Stracke, 2004; Workman and Hart, 2005), although these results should be considered with caulon because MORBs likely represent homogenized final products. The manue source of ocean island basalts (OIBs), i.e., the FOcal ZOne (FOZO), high-µ (HIMU), \* rimi ive mantle (PM), and enriched mantle (EM) reservoirs, is rather heterogeneous and more hydrated (96–750 ppm  $H_2O$ , depending on the specific reservoir; McDonough a id S in, 1995; Dixon and Clague, 2001; Dixon et al., 2002; Wallace, 2002; Workman et al., 2014, 2006; Cabral et al., 2014; Kendrick et al., 2014, 2015). The mantle transition zone is also likely water-rich (e.g., Bolfan Casanova et al., 2012). Therefore, water is likely present throughout the upper mantle and at different concentrations in the individual reservoirs. Variations also exist within each reservoir; for example, EM reservoirs are thought to contain between 90 and 400 ppm H<sub>2</sub>O (Bizimis and Peslier, 2015). Because hydrogen is both volatile and incompatible, its concentration in minerals is modified by processes including partial melting, crystallization, and metasomatism. Furthermore, given the large variation of hydrogen concentrations between olivine and pyroxene, different mantle

lithologies (e.g., dunite, harzburgite, pyroxenite, etc.) likely have diverse bulk water contents (0–120 and 0–350 ppm H<sub>2</sub>O in peridotites and pyroxenites, respectively; Fig. 1, Table 1).

The lithospheric upper mantle may contain 5% pyroxenites (Kornprobst, 1969; Pearson et al., 1993), comprising eclogites (recycled oceanic crust; Allègre and Turcotte, 1986; Viljoen et al., 2005; Gonzaga et al., 2010; Montanini et al., 2012) and high-pressure cumulates fractionated from basaltic magmas percolating through the lithospheric mantle (O'Hara and Yoder, 1967; Viljoen et al., 2005; Downes, 2007; Gonzaga et al., 2010; Perinelli et al., 2011; France et al., 2015), which have distinct geochemical signatures (e.g., O isotopic data, trace elements) from one another (e.g., Gonzaga et al, 2010; Pezzali et al., 2015). Despite the low abundance of pyroxenites in lithosphaic mantle domains, their relatively high-water contents make them important lithologie: for constraining the global water budget of the upper mantle. Indeed, because of their d fferent chemical and modal compositions (<40% olivine, although some exceptions vist at high pressure; Green et al., 2010; Kovács et al., 2012; Lambart et al., 2016), the malting temperatures of most pyroxenites are lower than those of peridotites (Lambart et a'., 2010). This implies that pyroxenites can contribute up to 40% of the total melt formed during basalt magma genesis (Mallick et al., 2015). In contrast, if the surrounding mantle is varially molten, the consumption of pyroxenite-derived melts is limited to less than 20% the to liquid reactivity (Lambart et al., 2012). Evidence of such heterogeneities is visible in percolated mantle domains (e.g., Borghini et al., 2016; Sanfilippo et al., 2017) and in magmas erupted at the surface in various geodynamic settings (rift related, Rooney et al., 2014; MORBs and OIBs, Sobolev et al., 2007; Shorttle et al., 2014). Furthermore, in some cases, melts derived from eclogite-bearing peridotite sources can react with the surrounding mantle to form secondary pyroxenites (Borghini et al., 2016). Hence, pyroxenites are at least as involved in magma genesis as peridotites (Hirschmann and Stolper, 1996; Lambart et al., 2009). Because the mantle sources of magmas are commonly at depths

where garnet pyroxenites are stable (deeper than 45 km), those lithologies are essential to our understanding of magma genesis. Nevertheless, few data are available on the water contents of garnet pyroxenites (Bizimis and Peslier, 2015; Hao et al., 2018; Li et al., 2018).

In this study, we present new data on the water contents of NAMs in garnet pyroxenites from two continental domains: the French Massif Central (FMC, France) and the Adamawa Volcanic Plateau (AVP, Cameroon). We use our results to constrain the contribution of garnet pyroxenites to the water distribution in the subcontinental lithospheric mantle. By comparing these water contents to trace element data, we disc ass the effects of mantle metasomatism on the distribution and concentration of water in the subcontinental lithospheric mantle. Our results reveal varying degree a content dehydration of NAMs of FMC pyroxenites and thus highly variable H<sub>2</sub>O/C a rations, limiting the reliability of H<sub>2</sub>O/Ce ratios for tracking the influence of garnet pyrc remes during magma genesis.

#### 2. Geological settings and sample (le.c. iptions

The studied garnet pyrox mite xenoliths record some of the mineralogical and petrological heterogeneities of u e continental lithospheric mantle beneath the FMC and the AVP. Petrological and genetic mical characteristics (textures, bulk-rock and phase-specific major and trace element concentrations, and O-Sr-Nd isotopes), together with geodynamic interpretations of the genesis and pressure-temperature-time evolution of these samples are available in France et al. (2015). These garnet pyroxenites are mantle cumulates that crystallized from percolating melts at 1–2 GPa during post-collisional magmatism (possibly of Hercynian age in the FMC and Pan-African age in the AVP; France et al., 2015). Their textures and chemical compositions were subsequently fully re-equilibrated at various pressure-temperature conditions (France et al., 2015). Here, we briefly summarize the geological histories of these two regions.

#### 2.1. French Massif Central

The lithospheric mantle beneath the FMC is composed of two domains north and south of ~45°30'N (Lenoir et al., 2000; Fig. 2a) inherited from the Hercynian orogeny and characterized by different petrological and geochemical properties. The northern domain is refractory, enriched in light rare-earth elements (LREEs), and similar to the cratonic subcontinental lithospheric mantle; it is thus considered to be a ore-Variscan lithospheric block. The southern domain is more juvenile and likely accreted ouring the Hercynian orogeny (Lenoir et al., 2000). Both domains have been enriched by metasomatic fluids. Based on REE data, Wittig et al. (2007) interpreted that the northern and southern domains were metasomatized by carbonatitic fluids and silicate metas, respectively. However, Gu et al. (2016, 2018) recently showed that the southern. domain was also metasomatized by either fluid with carbonatitic affinities or melt/flui.<sup>1</sup> interactions related to subducted materials.

Garnet pyroxenite xenolith: n a nly comprise clinopyroxene and garnet and are from two volcanic areas in the souther. domain: the Devès (Marais de Limagne and Saint-Didier d'Allier basaltic maars) and the languedoc (Le Pouget basaltic maar) volcanic fields (France et al., 2015; Fig. 2a). In the corresponding pyroclastic deposits, most xenoliths are peridotitic, pyroxenites are less common, and garnet pyroxenites are extremely rare: only five samples have been reported in the FMC, one in Touron (2005) and four in France et al. (2015), corresponding to those studied herein. All samples from the Devès volcanic field (LN78, LP27, and SD53) show evidence of metasomatism (e.g., amphiboles in LN78), whereas that from Le Pouget has not been affected by any metasomatic process (France et al., 2015). Amphiboles in these samples are not part of the primary cumulate assemblage, but result from metasomatic interactions between the primary magmatic mineral assemblage (garnet, clinopyroxenes, orthopyroxenes, and spinel) and a metasomatic fluid (France et al., 2015).

#### 2.2. Adamawa Volcanic Plateau

The AVP basement formed during the Eburnean orogeny 2.1 Ga and was reworked during the Pan-African orogeny and by subsequent granite intrusions ~520 Ma (Castaing et al., 1994; Fig. 2b). During the break-up of Pangea, the lithospheric mantle domain was metasomatized by a hydrous silicate melt (Lee et al., 1996; Temdjim et al., 2004; France et al., 2015; Njombie et al., 2018). The alkaline AVP is in the norther part of the Cameroon Volcanic Line, a large plutonic and volcanic province emplaced from 65 Ma to present (see chronological summary by Njome and de Wit, 2014).

Pyroxenite xenoliths (YK01, YK03, YK05, YK12, YK13, and YK16) were sampled at the Youkou basaltic maar. In the corresponding ryroclastic deposits, most xenoliths are peridotitic, and pyroxenites are less common. Pyroxenites mainly comprise clinopyroxene and garnet (Table 2). All xenoliths were metasonalized, and all samples except YK01 contain rare secondary amphiboles that result from an etasomatic interactions at depth between the primary magmatic mineral assemblage (garnet, clinopyroxenes, orthopyroxenes, and spinel) and a metasomatic agent (France et al. 2015). Two groups of xenoliths have been defined based on their petrographic characteristics; the first (YK01, YK05 and YK16) contains cm-sized megacrysts of clinopyror one with exsolved garnet crystals, whereas the second (YK03, YK12 and YK13) is composed of mm-sized granular assemblages (France et al., 2015).

#### 3. Petrographic characterization and metasomatic interactions

#### 3.1. Pyroxenites as metacumulates

Here, we summarize the main petrographic characteristics of the pyroxenite xenoliths to highlight that the studied samples are mantle metacumulates that suffered mantle

metasomatism. A detailed description and interpretation of the origin and evolution of these samples in their geodynamic context is available in France et al. (2015).

The FMC pyroxenites show recrystallized granoblastic textures. Samples LN78, LP27, and SD53 have average grain sizes of 1 mm, whereas the Le Pouget sample is finer grained with pyroxenes averaging 0.25 mm and spinel and garnet 0.75 mm . Based on the corresponding bulk-rock heavy REE (HREE) enrichments, garnet was likely part of the liquidus assemblage that formed the cumulates (see section 3.2 and details in France et al., 2015). Nevertheless, the garnets currently present in the samples are likely metamorphic; garnets often enclose green spinel and contain numerous spinel plagioclase, orthopyroxene, and clinopyroxene inclusions surrounded by radial fractores (Fig. 3a). Those inclusions record the transition from the garnet-free Seiland pyroxenit, mandle facies to the garnet-bearing Ariegite pyroxenite facies under changing pre-sure-temperature (P-T) conditions (likely decreasing P; France et al., 2015). Severar ' rge clinopyroxene grains display orthopyroxene exsolutions (e.g., LN78, Fig. 3b), indicating an early decrease in temperature. Local assemblages of finer grained plagloclase, brown spinel, and orthopyroxene are in contact with both clinopyroxene and garnet, subgesting a late re-equilibration in the Seiland subfacies (France et al., 2015).

The AVP pyroxer ites of the first petrological group (France et al., 2015) contain clinopyroxene megacrysts (0.5–3 cm) with garnet exsolutions (Fig. 3d), and additional garnet is present around spinel grains. Sample YK05 records crystal-plastic deformation (Fig. 3d) and shows additional orthopyroxene exsolutions in clinopyroxene. The second group of AVP pyroxenites comprise mm-sized granular assemblages (~3 mm) with clinopyroxenes containing exsolutions of garnet, amphibole, and spinel (Fig. 3c; France et al., 2015). Garnet is both exsolved from and bordering clinopyroxene grains. Very thin (~1–2  $\mu$ m) spinel exsolutions are also present in clinopyroxenes. Exsolutions of garnet and orthopyroxenes in

clinopyroxenes are likely related to a decrease in temperature following crystallization, suggesting chemical and mineralogical re-equilibration at different P-T conditions (Ariegite facies). As in the FMC, these petrographic characteristics indicate that the mantle cumulates record the transition from a garnet-free to a garnet-bearing facies, likely with decreasing T (France et al., 2015).

The above petrographic features and corresponding thermobarometric estimates highlight that the FMC and AVP pyroxenites record the protracted evolution of the lithospheric mantle domains, with re-equilibrations at various r - r conditions (France et al., 2015). Thus, these cumulates are now metacumulates who is mineral compositions have been modified by metamorphism and no longer represent the initial magmatic composition.

#### 3.2. Pyroxenite trace element ( on positions

The chondrite-normalized REE patterns of the three FMC pyroxenites from the Devès massif are enriched in HREEs inc LREEs, but depleted in middle REEs (MREEs) (see Fig. 6a, b in France et al., 2015 . Thus, the chondrite-normalised REE shaped pattern decreases from the HREE to the MREE and increases from the MREE to the LREE (spoonshaped REE pattern). The Le Pouget sample displays similar M-HREE contents, but is strongly depleted in LREEs (by up to a factor of 100 relative to M-HREEs). These enrichments in HREEs compared to MREEs attest to the initial presence of magmatic garnet. Additionally, the Devès samples are enriched in U and Th and depleted in Nb, Zr, and Hf, which, along with LREE enrichment, have been attributed to metasomatism in peridotite xenoliths of the FMC (Lenoir et al., 2000; Féménias et al., 2003; Dautria et al., 2010). Thus, the presence of these anomalies in the studied pyroxenites attest to a similar metasomatic stage in which a LREE-depleted protolith was enriched by a LREE-rich percolating agent. Moreover, sample LN78 contains amphibole-rich and amphibole-poor areas, indicating the

localized percolation of the metasomatic agent through the sample. Furthermore, clinopyroxene grains near amphiboles have REE patterns similar to those of amphiboles, i.e., highly enriched in LREEs and less depleted in HREEs than other clinopyroxenes, suggesting equilibration between the two species. Thus, FMC pyroxenites were metasomatized; amphiboles are not primary and LREE compositions do not reflect the initial magmatic composition.

The chondrite-normalized REE patterns of AVP pyroxenites display flat M-HREE patterns indicating a lack of magmatic garnet during their initial crystallization. These samples (especially YK13 and YK16) are enriched in LREEs relative to MREEs and enriched in Ba, Sr, U, and Th, and depleted in Zr and Hf relative to other trace elements with similar incompatibilities during mantle melting (see Fig. 6c, d in France et al., 2015). Similar signatures in spinel lherzolites from this regio: (e.g., Temdjim et al., 2004) have been attributed to metasomatic interactions, and the accordingly interpret these geochemical signatures as evidence of metasomaticm (France et al., 2015).

#### 4. Fourier-transform infrarea :pectroscopy

We determined the v ater contents of NAMs by Fourier-transform infrared spectroscopy (FTIR), a con-destructive method with a low detection limit (<1 ppm H<sub>2</sub>O). Measurements were performed on doubly polished sections 150–300 µm thick that were cut at the Laboratoire Magmas et Volcans (Clermont-Ferrand, France) and the Université Jean Monnet (Saint Etienne, France); residue of glue used during section preparation was carefully removed in successive acetone and alcohol baths. Unpolarized and polarized spectra were acquired at the Laboratoire de Spectroscopy pour les Interactions, la Réactivité et l'Environnement (Université de Lille, France) using a Bruker Vertex 70 infrared spectrometer coupled to a HYPERION 3000 microscope and a Cassegrain 15× lens. Absorption spectra

were collected over 256 scans at 4 cm<sup>-1</sup> spectral resolution between 600 and 4000 cm<sup>-1</sup> in transmission mode with aperture sizes ranging between 40 × 40  $\mu$ m and 100 × 100  $\mu$ m depending on the size and quality (e.g., inclusions, alteration, etc.) of each mineral. The majority of analyses were performed using unpolarized infrared light, which are as reliable as polarized analyses if at least ten different grains with distinct random orientations are measured and the maximum linear unpolarized absorbance is <0.15 (Kovacs et al., 2008). To maximize the reliability of our analyses, between 16–34 grains of both clinopyroxene and orthopyroxene were analyzed in every sample (Table 3). As gamen is isotropic and anhydrous (there were no discernal absorption peaks to analyze, see socion 5.3; Figs. 4, 5), only 5 analyses were performed per sample.

Because sample YK01 comprises large, cm-2iz, d clinopyroxenes, it was not possible to analyze >10 grains; therefore, we conducted pointized analyses. Two perpendicular sections of each clinopyroxene grain weight of and several polarized analyses were performed at complementary orientations (e.g., 96<sup>-/1</sup>80°, 105°/195°). Because the summed absorbances of the two sections of each grain were constant regardless of the complementary orientations used, we report results using the 90°/180° for each grain to quantify the water contents of YK01 clinopyroxenes.

All spectra we normalized to section thickness, and mean clinopyroxene and orthopyroxene spectra for each sample were calculated from the respective individual spectra. Baseline corrections for the mean spectra was performed following the procedure of Hao et al. (2014). To subtract the baseline, several methods are possible. The one used here is manual. For each considered spectrum three baselines, one optimum and two extremes are generated in order to define the uncertainty values. At least three anchor points are defined on either side of the spectrum peaks (Supplementary Fig. S2). These points are connected by a polynomial function. This method is user-dependent; however, it ensures the reliability of the choice of

the best baseline. Thus, the measurement uncertainty is defined according to the maximum and minimum areas defined under the spectra. Then, the errors due to the Bell et al. (1995) and Libowitzky and Rossman (1997) absorption coefficients are propagated until the final uncertainty obtention. The area under the OH<sup>-</sup> peak was integrated between 3000 and 3800  $cm^{-1}$  for clinopyroxenes and orthopyroxenes. Water concentrations were calculated from the thickness-normalized absorbances using a modified Beer-Lambert law and two calibrations: the mineral-dependent calibration of Bell et al. (1995; using integral specific absorption coefficients  $l'_{Cpx} = 7.09 \pm 0.32$  and  $l'_{Opx} = 14.84 \pm 0.59$  for cline and orthopyroxene, respectively) and the wavenumber-dependent calibration of Libowitzky and Rossman (1997; using mean wavenumbers  $v_{Cpx} = 3550 \text{ cm}^{-1}$  and  $v_{Opx} = 350^{\circ} \text{ cm}^{-1}$ ). For diopside, the wavenumber-dependent calibration is considered to he more reliable than the mineraldependent calibration (Weis et al., 2018). How zv r 'ne mineral-dependent calibration was developed using natural minerals similar to those analyzed herein and is thus particularly suitable for our purposes. Furthermore, the mineral-dependent calibration has been more widely used than the wavenumber-deperdent calibration, better allowing for comparisons with literature data. Thus, althe 'gh we used both calibrations, we use only the water concentrations calculated using the Bell et al. (1995) calibration for comparison with previous data. The total  $2\sigma$  uncervatures on water concentrations in clinopyroxenes and orthopyroxenes are between 15 and 25% (Table 2, Supplementary Fig. S2, Fig. S4).

#### 5. Results

Clinopyroxene and orthopyroxene water contents are presented in Table 3, and related spectra are displayed in Figs. 4 and 5. We measured core-rim profiles to demonstrate the homogeneity of water contents in clinopyroxene and orthopyroxene grains from all the studied pyroxenites (Fig. 6).

#### 5.1. Clinopyroxene water contents

All clinopyroxenes from the AVP and Le Pouget (FMC) show similar mean spectra with absorption bands centered around ~3630 cm<sup>-1</sup>, ~3530 cm<sup>-1</sup>, and ~3470 cm<sup>-1</sup> (Fig. 4). These bands are common in the literature (e.g., Bell et al., 1995; Bizimis and Peslier, 2015; Patkó et al., 2019), and the largest peak occurs at the highest wavenumber (~3630 cm<sup>-1</sup>). Here, we refer to such spectra as 'Type 1'. In contrast, the other FMC samples (LN78, LP27, and SD53) show uncommon spectra with bands centered around ~3610 cm<sup>-1</sup>, ~3518 cm<sup>-1</sup>, and ~3445 cm<sup>-1</sup> (Fig. 4), and the largest peak occurring at . lov/er wavenumber (~3518 cm<sup>-1</sup>) compared to Type-1 spectra. Furthermore, the contribution of the first band, with the largest peak occurring at ~3610 cm<sup>-1</sup>, to the spectra increases progressively from LN78 to Le Pouget clinopyroxenes (Fig. 4). These spectra are here, fter referred to as 'Type 2'.

Water concentrations in AVP clinopy oxenes vary between xenoliths from  $386^{+78}_{-85}$  to  $685^{+92}_{-154}$  ppm H<sub>2</sub>O using the Bell et al ( . 995) absorption coefficient (Fig. 7a); from  $296^{+28}_{-44}$  to  $525^{+16}_{-81}$  ppm H<sub>2</sub>O using the Libowitzky and Rossman (1997) (Supplementary Fig. S3) absorption coefficient. Clinopyroxene water concentrations in samples YK01, YK03, YK05, YK12, and YK13 are '101000 neous, with average values of  $455 \pm 24$  (1 $\sigma$ ) ppm H<sub>2</sub>O using the Bell et al. (1995) absorption coefficient and ( $349 \pm 19$  (1 $\sigma$ ) ppm H<sub>2</sub>O using the Libowitzky and Rossman (1997) dusorption coefficient. Only clinopyroxenes in YK16 have higher water contents of  $685^{+92}_{-154}$  ppm H<sub>2</sub>O using Bell et al. (1995) absorption coefficient. Unlike the AVP clinopyroxenes, water concentrations in FMC clinopyroxenes differ more markedly between xenoliths, varying from  $112^{+25}_{-15}$  ppm H<sub>2</sub>O in LN78 to  $465^{+185}_{-100}$  ppm H<sub>2</sub>O in the Le Pouget sample using Bell et al. (1995) absorption coefficient (Fig. 7a); from  $85^{+9}_{-9}$  ppm H<sub>2</sub>O in LN78

to  $357^{+96}_{-52}$  ppm H<sub>2</sub>O in the Le Pouget sample using the Libowitzky and Rossman (1997) (Supplementary Fig. S3) absorption coefficient.

#### 5.2. Orthopyroxene water contents

Orthopyroxenes from the AVP and Le Pouget (FMC) show similar mean spectra with bands centered around ~3590 cm<sup>-1</sup>, ~3518 cm<sup>-1</sup>, and 3417 cm<sup>-1</sup> (Fig. 4). These spectra are referred to hereafter as 'Type 1', and are the most common in the literature. Orthopyroxenes from the other FMC samples (LN78 and LP27) have spectra with only two bands centered around ~3570 cm<sup>-1</sup> and ~3517 cm<sup>-1</sup> (Fig. 4), which we refered as 'Type 2' hereafter. In all orthopyroxenes, the largest peak, with the highest line or a psorption, corresponds to the highest wavenumber (3590 cm<sup>-1</sup> and 3570 cm<sup>-1</sup> for Type-1 and Type-2 spectra, respectively).

Water concentrations in AVP or ' iop /roxenes (YK03, YK05, YK12, and YK13) vary between xenoliths from  $124_{-15}^{+15}$  to  $15.1_{-35}^{+35}$  ppm H<sub>2</sub>O using Bell et al. (1995) absorption coefficient (Fig. 7b); from  $160_{-8}^{+6}$  to  $2.01_{-30}^{+26}$  ppm H<sub>2</sub>O using the Libowitzky and Rossman (1997) (Supplementary Fig. S2, absorption coefficient, but are homogeneous with an average value of  $143 \pm 13$  (1 $\sigma$ ) ppm  $1^{+}\sigma$ C using Bell et al. (1995) absorption coefficient and  $184 \pm 17$ (1 $\sigma$ ) ppm H<sub>2</sub>O using the Libowitzky and Rossman (1997) absorption coefficient. Water concentrations in FMC orthopyroxenes (LN78, LP27, and Le Pouget) vary from  $61_{-13}^{+22}$  ppm H<sub>2</sub>O in LN78 to  $104_{-16}^{+14}$  ppm H<sub>2</sub>O in Le Pouget using Bell et al. (1995) absorption coefficient (Fig. 7b); from  $78_{-12}^{+19}$  ppm H<sub>2</sub>O in LN78 to  $135_{-14}^{+6}$  ppm H<sub>2</sub>O in Le Pouget using the Libowitzky and Rossman (1997) (Supplementary Fig. S3) absorption coefficient. Although SD53 (FMC) contains orthopyroxenes, they were not analyzed because they show secondary fluid inclusions and some alteration features.

#### 5.3. Garnet water contents

None of the garnet spectra show OH bands between 3000 and 3800 cm<sup>-1</sup> (Fig. 4). Water concentrations in these garnets are therefore below the FTIR detection limit for garnet and this range of sample thickness (<0.5 ppm H<sub>2</sub>O; Bizimis and Peslier, 2015). Therefore, we hereafter consider the studied garnets to be anhydrous. In the Le Pouget sample, rare areas near the edges of some garnets or in fracture zones present a small peak between 3000 and  $3800 \text{ cm}^{-1}$  that may result from a late episode of fluid diffusion through fractures (Suzuki and Nakashima, 2004). Despite this event, the cores of those grain a remain anhydrous.

#### 6. Discussion

To better understand the distribution of writer in the lithospheric mantle and its implications, we divide the discussion of r unrest its into in three parts. We first discuss whether the quantified water contents represent lithospheric mantle signatures or were modified during xenolith transport transport to the surface (section 6.1). We then explore the potential influence of metasomatism on the lithospheric water contents (section 6.2), and finally discuss the use of H<sub>2</sub>O/Ce to track the in fluence of pyroxenite during magma genesis (section 6.3).

6.1. Water contents: Mantle signature preservation and whole-rock water concentrations

A key concern when studying water systematics in NAMs is whether the xenoliths retained their primary water contents; here, that is whether the pyroxenites reliably preserve mantle water from before their ascent to the surface in the host magma, or if the mantle water was modified during ascent. Indeed, the solubility of water varies with pressure and temperature, and its diffusivity in the melt and minerals may enable quick re-equilibration

with the host magma at magmatic temperatures, even at the timescale of an individual eruption (e.g., Zhao et al., 2004; Ingrin and Blanchard, 2006; Demouchy and Bolfan-Casanova, 2016). Moreover, degassing can potentially disturb xenolith water contents. Hence, mineral water contents, especially olivine, could be disturbed during xenolith ascent and emplacement (e.g., Demouchy et al., 2006, 2015; Peslier and Bizimis, 2015; Hao et al., 2016a). Here, we consider three criteria to determine if the pyroxenite water contents effectively document water contents in the lithospheric mantle.

First, our profile analyses highlight the homogeneity of Cinopyroxene, orthopyroxene, and garnet grains in all studied pyroxenites (Fig. 6). Indeed, the water contents of pyroxene cores and rims are identical within uncertainties, whereas the diffusion of H into or out of pyroxene would result in increase 1 or decreased rim water contents, respectively. Consequently, there is no eviden to that hydrogen diffused into or out of the FMC and AVP pyroxenes at the timescale of magina ascent and emplacement.

Second, in the FMC pyroxe  $n_{1}$  s, orthopyroxene and clinopyroxene water contents correlate with their major element concentrations, such as Mg# ( $R^2 = 0.91$ ) and Ca ( $R^2 = 0.99$ ) in clinopyroxene (Fig. 8a, b) and Mg# ( $R^2 = 0.99$ ) in orthopyroxene (Fig. 8c). The diffusivities of these major elements prec. de their re-equilibration at the grain scale over periods characteristic of mantle x nolith ascent times (a few hours to days; e.g., Demouchy et al., 2006). Therefore, the water and major element concentrations in FMC pyroxenites are interdependent and consistent with water contents at equilibrium conditions prior sampling by the magma. In contrast, there are no such correlations in AVP pyroxenites (Fig. 8), indicating that their water was likely not acquired during the stage of major element equilibration or at least much less sensitive to these changes.

Lastly, clinopyroxenes in FMC and AVP pyroxenites contain 1.5–3.5 times more water than orthopyroxenes from the same sample (Fig. 9), consistent with the range of water

partition coefficients between clinopyroxene and orthopyroxene  $(D_{H_2O}^{Cpx-Opx})$  determined in natural samples: 1.5–3.5 (Warren and Hauri, 2014; Bizimis and Peslier, 2015; Demouchy and Bolfan-Casanova, 2016; Demouchy et al., 2017; Peslier et al., 2017; Xia et al., 2017; Hao et al., 2018; Li et al., 2018). Hence, clinopyroxene and orthopyroxene water contents are considered to be in equilibrium for all samples studied herein.

Based on these three arguments, the FMC and AVP pyroxenites likely preserved their lithospheric water contents during xenolith ascent. Based on this assumption, bulk-rock water concentrations can be estimated by considering the modal percent. ges and water contents of each mineral. This calculation shows that bulk-rock AVP pyrox enite water concentrations are homogeneous with an average value of  $309 \pm 95 (1\sigma)$  pp. H<sub>2</sub>O (ranging from  $231_{-51}^{+47}$  to  $480_{-108}^{+64}$  ppm H<sub>2</sub>O using the Bell et al. (1995) ab so ption coefficient. Bulk-rock water concentrations of FMC pyroxenites vary from  $52_{-9}^{+15}$  ppm H<sub>2</sub>O for LN78 to  $287_{-61}^{+113}$  ppm H<sub>2</sub>O for the Le Pouget sample (Table 3, Fig. 7c).

6.2. Influence of metacomatism on water contents: unaffected AVP pyroxenites vs dehydraica r'MC pyroxenites

Because hydrogen is both volatile and incompatible, both modal and cryptic metasomatism likely can affect the water contents of upper mantle minerals. Nevertheless, Denis et al. (2015) showed that neither modal (e.g., presence of hydrated minerals such as amphibole) nor cryptic (e.g., LREE and MREE enrichments) metasomatic interactions have had a clear effect on the water contents of FMC peridotite xenoliths. They suggested that only high-melt/rock-ratio interactions have the potential to dehydrate xenoliths (Denis et al., 2018). In contrast, other studies have attributed high water concentrations in pyroxenites and peridotites to metasomatic agents (e.g., Peslier and Bizimis, 2015; Li et al., 2018; Patkó et al.,

2019). In this perspective, we explore the influence of metasomatism on AVP and FMC pyroxenite water contents to determine if they represent those acquired during metasomatism or rather document the subsequent equilibration conditions of the mantle metacumulates. Several lines of evidence indicate that some degree of metasomatism occurred after igneous crystallization, such as bulk-rock and mineral LREE enrichments associated with specific elemental enrichments or depletions (e.g., U enrichments and depleted high field strength element concentrations in the FMC; U and Th enrichments in the AVP); these features are similar to geochemical signatures of the host peridotitic lithosphere, domains that have been attributed to metasomatism (e.g., Lenoir et al., 2000; Fémé na. et al., 2003; Temdjim et al., 2004; Dautria et al., 2010).

Geochemical tracers can be used to identify and characterize metasomatic agents. The Cameroon lithospheric mantle, especially beneath the AVP, experienced metasomatism during the early break-up of Pangea aroun.<sup>1</sup> 200 Ma, which accounts for the incompatible element enrichments in related perido<sup>1</sup>te. (Lee et al., 1996; Temdjim et al., 2004; Njombie et al., 2018). The studied AVP pyror en<sup>1</sup>tes are similarly enriched in LREEs, U, and Th relative to MREEs and other incompa<sup>+</sup>ib<sup>+</sup>, trace elements, and therefore also experienced metasomatism (France et al., 2015). Two types of mantle-melt interactions are recorded in lherzolites from the AVP. (i) cryptic metasomatism and trace element enrichments are attributed to a regional mantle plume and the percolation of hydrous silicate melts, whereas (ii) enrichments in CaO relative to Al<sub>2</sub>O<sub>3</sub> during refertilization are attributed to the percolation of carbonatite or carbonated silicate melts (Njombie et al., 2018).

The La<sub>N</sub>/Sm<sub>N</sub> ratio (the subscript 'N' denotes chondrite-normalized values; Anders and Grevesse, 1989) is widely used to identify mantle metasomatism, whereas the Ti/Eu ratio is a good proxy for carbonatitic metasomatism (Ashley et al., 2020), which is characterized by Ti, Zr, and Hf depletions relative to REEs. Indeed, as Ti, Zr and Hf are less incompatible than

REEs in a carbonatic melt, the mineral, which interacts with this carbonatitic melt would record its signature (Ashley et al., 2020). This correlation relies on the corresponding partition coefficients that imply that REE are strongly enriched in carbonatite melts when HFSE (Ti, Nb-Ta, Zr-Hf) are strongly depleted (e.g., Coltorti et al., 1999; Martin et al., 2013). The studied AVP pyroxenites likely record a metasomatic stage related to hydrous silicate melts because Ti/Eu and La<sub>N</sub>/Sm<sub>N</sub> are not correlated (Fig. 10). Thus, to track the metasomatic influence on the pyroxenite water contents, we plotted clinopyroxene and whole-rock water concentrations against La<sub>N</sub>/Sm<sub>N</sub> (Fig. 11a, b). Clinopyroxene water contents are not correlated with  $La_N/Sm_N$  whereas whole-rock contents show a rough  $los^{\dagger}$  ive correlation ( $R^2 = 0.57$ , Fig. 11b). The whole-rock enrichments in LREEs relative to MPEEs in these samples might be related to the widespread metasomatic stage that affer te., the Cameroon lithospheric domain (Lee et al., 1996; Temdjim et al., 2004; France et 4, 2015; Njombie et al., 2018). Nevertheless, because clinopyroxene waar concentrations do not correlate with tracers of metasomatism, we propose that the rough whole-rock correlation reflects the modal composition of the pyroxenites and net r metasomatic event. Indeed, the most water-rich AVP pyroxenite YK16 contains the vost clinopyroxene (the most hydrous mineral phase present). Furthermore, samples YK01, VV.05, and YK16 contain megacrysts, decreasing the reliability of the results. Therefore, we consider that all AVP pyroxenites have roughly similar water contents that were not affected by metasomatism by the hydrous silicate melts, a result similar to what Denis et al. (2018) observed in the FMC. The relative homogeneity of water contents of AVP pyroxenites likely documents a previous equilibration stage in the lithosphere during their PT evolution that is not directly related to metasomatism.

The lithospheric mantle beneath the FMC also experienced metasomatic interactions: the northern domain was metasomatized by carbonatitic melts (Lenoir et al., 2000; Wittig et al., 2007), whereas the southern domain was metasomatized by both hydrous silicate melts

and carbonatitic melts (Wittig et al., 2007; Gu et al., 2018). The FMC samples studied herein show a robust correlation between  $La_N/Sm_N$  and Ti/Eu ( $R^2 = 0.88$ , Fig. 10), consistent with recent evidence of carbonatitic metasomatism in the FMC (e.g. Gu et al., 2018). The negative correlations between clinopyroxene and whole-rock  $La_N/Sm_N$  ratios and water concentrations (Fig. 11a, b) can be interpreted as indicative of a common process for LREE enrichment and water depletion. Indeed, according to Keppler (2003) and Sokol et al. (2013), carbonatitic melts can be considered as "dry" metasomatic agents with low water activity in the melt, and have the potential to extract water from even water-poor rocks cuch as peridotite or pyroxenite.

The presence of amphiboles could also explain the dehydration of FMC pyroxenes. Indeed, the metasomatism would have stabilized am<sub>i</sub>, hibole, which by being OH-rich, resulted in dehydration of these pyroxenes. He wever, Denis et al (2015) showed that the presence of amphibole does not affect the tater content of NAMs. There is also a continuous variation in water content between the four samples, from the OH-rich sample (Le Pouget) to the OH-poor sample (LN78). Moreover, LP27 also contains little water while no amphibole is present. Finally, all samples from Cameroon (but YK01) contain small amounts of amphibole, and all of them are OH-rion. Thus, the dehydration of FMC pyroxenites is preferentially due to the carbonatitic metaso natism.

Furthermore, the strength of the metasomatic imprint (i.e., the degrees of LREE enrichment and Ti/Eu depletion) correlates with the type of FTIR spectra observed (type 1 vs. type 2): the more pronounced the carbonatitic metasomatic imprint, the weaker the high-wavenumber water band (type 2, see the most metasomatized sample LN78 in Fig. 4). The spectral signature of water in minerals is characterized by various bands, each representing precise vibrational interactions, and the several hydrogen storage defects present in the pyroxene lattice account for the various bands observed between 3000 and 4000 cm<sup>-1</sup> (Ingrin

and Skogby, 2000). Several processes can explain the incorporation of water in NAM structures, including cation vacancies (e.g.,  $2H^+$  vs.  $Mg^{2+}$  or  $4H^+$  vs.  $Si^{4+}$ ), charge balance (e.g.,  $Al^{3+} + H^+$  vs.  $Si^{4+}$ ), and redox reactions (Fe<sup>3+</sup> – O<sup>2+</sup> + 1/2H<sub>2</sub> = Fe<sup>2+</sup> + OH<sup>-</sup>). However, according to several authors (e.g., Skogby, 1994; Yang et al., 2019) there is no clear correlation between band positions and clinopyroxene chemical composition. It is therefore difficult to attribute specific band (between 3000 and 4000 cm<sup>-1</sup>) to specific OH locations in the crystalline structure. We suggest here that the carbonatitic metasomatism of the FMC pyroxenites preferentially dehydrated a specific OH site in the restructure, resulting in the uncommon type-2 spectra observed herein (Ingrin et al , 2019).

Type-2 spectra have also been reported scare *iy* in the literature and were associated with low structural hydroxyl contents (<360 ppm: Packó e. al., 2019), in agreement with our findings (Denis et al., 2018; Gu et al., 2018; P tko et al., 2019). However, contrary to the observations of Patkó et al. (2019), there is no correlation of type-2 spectra (and related low water content) with anomalously high hydroxyl partition coefficients between clinopyroxene and orthopyroxene: the ratio of cline pyroxene to orthopyroxene water contents in LN78 and LP27 are within 1.5-3.5, matching literature data (Fig. 9; Peslier et al., 2002, 2012; Peslier, 2010; Xia et al., 2010, 2010, Yu et al., 2011; Hao et al., 2012, 2014, 2016a, 2016b; Doucet et al., 2014; Warren and Ha, ri, 2014; Bizimis and Peslier, 2015; Demouchy et al., 2015; Hui et al., 2015; Li et al., 2015; Peslier and Bizimis, 2015; Denis et al., 2018; Hao et al., 2018). However, because Patkó et al. (2019) focused on peridotites whereas we focus on pyroxenites, differences in mineral partitioning or exchanges may explain the different observed partition coefficients. Several other parameters could also influence structural hydroxyl contents, and thus the observed spectra, such as oxidation caused by infiltration of the host magma (which in some cases, lowers the solubility of water in minerals, resulting in dehydration during peridotite partial melting; Peslier et al., 2002; Tollan and Hermann, 2019) or decreased water

activity due to the infiltration of a "dry" agent (Patkó et al., 2019). Because the whole-rock water contents of FMC and AVP pyroxenites are not correlated with redox state (France et al., 2015; Supplementary Fig. S1;  $R^2 = 0.01$  and  $R^2 = 0.13$  respectively), the infiltration of an oxidized melt seems unlikely in the case of the FMC pyroxenites. Finally, because the water contents and major element concentrations (e.g., Al or Mg#) of FMC pyroxenites are correlated (see section 5.1), the water contents are negatively correlated with the major element equilibration temperatures determined by France et al. (2015): the least hydrous and most metasomatized samples have the highest equilibrium temperatures (Table 2, Fig. 12), consistent with their partial dehydration during the metasomatized sam of the FMC lithospheric mantle by a hot carbonatitic melt or fluid.

In conclusion, FMC pyroxenites were metalomalized by a carbonatitic agent (Wittig et al., 2007; Gu et al., 2018) and AVP pyroxenites by a hydrous silicate one (Njombie et al., 2018). These interactions modified their number or and trace element compositions and the carbonatitic melt likely dehydrated the Funct pyroxenites, seemingly from a specific structural site, resulting in their unusual specific signatures.

#### 6.3. Pyroxeni<sup> $\cdot$ 2</sup>s, m<sup> $\epsilon$ </sup> tasomatism, H<sub>2</sub>O/Ce, and basalt sources

Pyroxenites are generally more water-rich than peridotites (Fig. 1) and are a key lithology for quantifying the water distribution in the upper mantle. Despite their relatively low abundance in mantle domains, their lower solidus temperatures relative to peridotites result in their disproportionate implication in magma genesis (e.g., Hirschmann and Stolper, 1996; Bizimis and Peslier, 2015). Nonetheless, quantifying their importance in magma genesis relative to peridotites has remained elusive because the major element compositions of the melts produced by melting each lithology are relatively similar (e.g., Le Roux et al., 2010; Lambart et al., 2013). Although radiogenic isotopes (e.g., Sr-Nd-Pb) are powerful

tracers of the involvement of mantle reservoirs evolving separately over billions of years (e.g., the DMM, HIMU, or EM reservoirs), only a few minor elements have proven reliable for tracking mantle compositional heterogeneities such as the presence of pyroxenites in basalt mantle sources. For example, the implication of pyroxenites in mantle melting is reflected by higher Ni/Mg and lower Mn/Fe or Ca/Fe ratios in olivine phenocrysts or by higher Zn/Fe and Ge/Si ratios in the formed basalts (Sobolev et al., 2007; Le Roux et al., 2010; Yang et al., 2020).

Alternatively, H<sub>2</sub>O/Ce ratios may document the chemical composition of basalt mantle sources (Michael, 1995). Indeed, because H<sub>2</sub>O and Ce 1 ave similar partition coefficients during peridotite mantle melting, the H<sub>2</sub>O/Ce ratio is expected to remain unchanged during this process, allowing mantle water contents to be estimated from basalts (Dixon et al., 1988; Michael, 1988, 1995). This was confirmed by Hauri et al. (2006) who showed that, despite the different H<sub>2</sub>O and C e partition coefficients for olivine, clinopyroxene, orthopyroxene, and game, the overall melting process buffers the H<sub>2</sub>O/Ce ratio, which remains almost constant during melting. Bizimis and Peslier (2015) also suggested that the implication on 'ow H<sub>2</sub>O/Ce ratio gamet pyroxenites (H<sub>2</sub>O/Ce < 200; [H<sub>2</sub>O] > 350 ppm) in mantle melting could account for the low H<sub>2</sub>O/Ce ratios of some fertile magmas (Hawaii, Samoa, Society). Relying on those important results and based on the observation that pyroxenites from other settings are also characterized by low H<sub>2</sub>O/Ce ratios (e.g., Hao et al., 2018; Li et al., 2018; Fig. 13), it seems plausible that the H<sub>2</sub>O/Ce ratio of basalts might track the involvement of pyroxenite in their genesis. We here explore this possibility in light of our new results.

Because the partition coefficients of  $H_2O$  and Ce vary among minerals, the evolution of the  $H_2O/Ce$  ratio during the melting of peridotite or pyroxenite also depends on mineralogical and modal compositions. Indeed, although Hauri et al. (2006) showed that the

H<sub>2</sub>O/Ce ratio is buffered and remains almost constant during peridotite melting, this may not be the case for pyroxenite. The use of basalt H<sub>2</sub>O/Ce ratios to quantify the water content of the mantle source may thus be inappropriate when pyroxenitic lithologies are involved. To test this, we calculated the H<sub>2</sub>O and Ce partition coefficients for bulk-rock pyroxenites based on literature constraints and mineral partition coefficients.  $D_{H_2O}^{Cpx-melt}$  varies between 0.015 and 0.019 among different samples and  $D_{\rm H_2O}^{\rm Opx-melt}$  between 0.009 and 0.015 (compositionally dependent values from O'Leary et al., 2010; Table 3).  $D_{H_2O}^{Garnet-nelt} = 0.0032$  and does not vary with pressure (Novella et al., 2014), representing the value at upper mantle depths. The partition coefficients of Ce between the liquid and const<sup>i</sup> uent minerals are  $D_{Ce}^{Cpx-melt} =$ 0.0858 (Hart and Dunn, 1993),  $D_{Ce}^{Opx-melt} = 0.0017$  (Alpha and Green, 2006), and  $D_{Ce}^{Garnet-melt} = 0.0034$  (Klemme et al., 2002). Hence,  $D_{H_2O}^{Pyroxenite-melt}$  varies between 0.011 and 0.014 and  $D_{Ce}^{Pyroxenite-melt}$  between 0.338 and 0.068 depending on modal composition. Therefore, we reach the same conclusion .s Bizimis and Peslier (2015): Ce is 4-5 times more compatible than H<sub>2</sub>O in pyroxeni'es, and the H<sub>2</sub>O/Ce ratio is likely modified during the melting process (e.g., increasing Juring partial melting).

To verify the efficiency of this tool as a source tracer, we ran a non-modal, incremental batch-melting model using modal mineralogies representative of our samples and assuming similar melting proportions of the main phases (1/3 clinopyroxene, 1/3 orthopyroxene, and 1/3 garnet; Fig. 14). At low degrees of melting, H<sub>2</sub>O/Ce can be up to three times the starting values of the studied pyroxenites, especially for the Le Pouget sample. With increasing degrees of partial melting, the ratios reach values similar to those of the melting pyroxenite at ~10% melting (Fig. 14). Thus, at least 10% partial melting of the pyroxenite assemblage is required to obtain a H<sub>2</sub>O/Ce similar to that of the magma sources. In the case of peridotites, such differences between the H<sub>2</sub>O and Ce partition coefficients, and thus between

the H<sub>2</sub>O/Ce ratios of the partial melting products and the source, are only apparent at <3%partial melting (Hauri et al., 2006), and the H<sub>2</sub>O/Ce ratios of basalts can thus be used to estimate the water content of the mantle source at higher degrees of melting. Because pyroxenites are fertile lithologies, e.g., up to 20% pyroxenite partial melting can be attained before the onset of lherzolite melting (Hirschmann and Stolper, 1996), it is likely that pyroxenite signatures can be detected in basalts. It follows that, if pyroxenites display different H<sub>2</sub>O/Ce ratios than peridotites, the H<sub>2</sub>O/Ce ratio can be used as a tracer of pyroxenites in the mantle source, and thus to attempt to quantify the involvement of pyroxenites during magmas genesis. However, both the AVr and FMC pyroxenites have H<sub>2</sub>O/Ce ratios spanning a wider range of values (from  $20^{+8}_{-5}$  to  $2,388^{+945}_{-512}$ ) than in previous studies (Bizimis and Peslier, 2015; Hao et al., 2018; Li e. al., 2018), and H<sub>2</sub>O contents varying from  $58^{+15}_{-9}$  to  $480^{+64}_{-108}$  ppm (Fig. 13). Thus, r. os. c. the variability of H<sub>2</sub>O/Ce ratios herein is governed by variations in Ce content (and the symptotic symptot symptotic symptot symp ratio is observed for the Ce-poor Le Poue et sample, the least metasomatized pyroxenite from the FMC. The range of H<sub>2</sub>O/Ce ratios in the studied pyroxenites is larger than that reported for the peridotitic mantle, reducing the utility of H<sub>2</sub>O/Ce for tracking pyroxenites in the magma source.

The question that remains is: why are such extreme H<sub>2</sub>O/Ce ratios (e.g.,  $2,388_{-512}^{+945}$ ) not observed in lavas at the surface when pyroxenites are generally more involved than peridotites in magma genesis. Although H<sub>2</sub>O/Ce systematics have not been studied in the FMC, magmas erupted at the surface likely result from the aggregation of heterogeneous magma batches formed by the partial melting of different mantle domains (e.g., Grove et al., 1992; Morgan et al., 1992; Saal et al., 1998; Koornneef et al., 2019). Lava compositions therefore likely represent the product of magma mixing, with little chance of preserving extreme values, especially in the case of incompatible elements such as Ce (e.g., Stracke et

al., 2019). For example, Stracke et al. (2019) identified depleted magmatic melt inclusions in basaltic lavas from the Azores, although this heterogeneity was not detected in bulk analyses. Under similar conditions, the melting of pyroxenites similar to the Le Pouget sample might not be detected in erupting magmas because such depleted melts do not directly rise to the surface, but rather mix with various magma batches formed by the melting of surrounding peridotites. Finally, given the H<sub>2</sub>O/Ce variations recorded in the studied pyroxenite samples (from  $30^{+8}_{-512}$ ) and the striking effect of metasomatism on their signature, the reliability of the H<sub>2</sub>O/Ce systematic to track the presence of pyroxe. ites in magma sources worldwide is limited; it is most applicable in areas where t<sup>4</sup> ie 1<sup>4</sup><sub>2</sub>O/Ce ratios of pyroxenites differ markedly from those of peridotites (e.g., Bizimis and Peslier, 2015).

#### 7. Conclusions

This study reports new FTIR data claracterizing water contents in the continental lithospheric mantle beneath the FMC (Trance) and AVP (Cameroon). We quantified bulk-rock and NAM water contents in pyrchemics that represent mantle heterogeneities in the lithospheric domains of each region. NAMS are homogeneous at the grain and sample scales (no evidence of diffusior) and in equilibrium according to available partitioning coefficients, and water contents correlate with major element concentrations or equilibrium temperature for FMC pyroxenites. Therefore, we interpret that the water contents of FMC and AVP pyroxenites were not noticeably modified during xenolith ascent and emplacement and are thus representative of lithospheric conditions.

Correlations between water concentration and proxies of metasomatism suggest that the FMC pyroxenite water contents were modified by metasomatic agents in the lithospheric mantle. Based on previous geochemical constraints on the metasomatic agents affecting FMC pyroxenites, we conclude that a carbonatitic metasomatic agent caused the dehydration of the

pyroxenite minerals. Based on their peculiar FTIR absorption spectra, we suggest that this dehydration preferentially affected specific OH storage sites in pyroxenes. The water contents of AVP pyroxenites have not been noticeably affected by metasomatism.

Finally, we investigated the validity of the  $H_2O/Ce$  systematic in our samples, which has been proposed as a tool for tracking the involvement of pyroxenite during magma genesis. Based on our new analyses we conclude that the increased range of  $H_2O/Ce$  ratios for bulkrock pyroxenites limits the utility of this parameter as a magma source tracer.

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#### **Research Data**

Research Data associated with this article can be accessed at doi:10.17632/jc46vyk76v.1 on Mendeley Data.

#### Supplementary data

Supplementary Figure S1

Supplementary Figure S2

Supplementary Figure S3

Supplementary Figure S4

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Fig. 1. The distribution of orthopyroxene, clinopyroxene, garnet\_a.d whole-rock water contents (ppm) in peridotite (left) and pyroxenite (right). Data are from Peslier et al. (2002, 2012), Peslier (2010), Xia et al. (2010, 2013), Yu et al. (2011), Hao et al. (2012, 2014, 2016a, 2016b, 2018), Doucet et al. (2014), Warren and Hauri (2014), Bizimis and Peslier (2015), Demouchy et al. (2015), Li et al. (2015, 2018) a. d Feslier and Bizimis (2015), Kilgore et al. (2018, 2020), Schaffer et al. (2019), Ashlav (t al. (2020). Data of this study are in green. On average, pyroxenites are more water-rich than peridotites.

Fig. 2. Simplified geologic ma<sub>r</sub>'s and sampling locations (red stars) in (a) the French Massif Central and (b) the Adan awa Volcanic Plateau (modified from France et al., 2015), black areas represent volcanic p ovinces. The stars indicate the samples location.

Fig. 3. (a, b) Photomicrographs and (c, d) X-ray elemental maps of the studied samples. (a) Sample LN78 showing garnets that often enclose green spinel (g-Spl) and contain numerous spinel (also brown spinel, b-Spl), plagioclase, orthopyroxene, and clinopyroxene inclusions surrounded by radial fractures. (b) A large grain of pyroxene with clinopyroxene and orthopyroxene exsolutions in sample LN78. (c) Millimetric granular assemblages in sample

YK03; thin garnet exsolutions are visible in clinopyroxenes. (d) Deformed clinopyroxene megacryst containing exsolutions of garnet and orthopyroxene in sample YK05.

Fig. 4. Sample-averaged unpolarized FTIR spectra of clinopyroxene, orthopyroxene, and garnet in samples from the FMC (LN78, LP27, SD53, Le Pouget) and AVP (YK03, YK05, YK12, YK13, YK16). Each average spectrum is normalized to 1-cm sample thickness and the reported water concentrations were calculated using the minera -epecific calibration of Bell et al. (1995) for clinopyroxenes and orthopyroxenes. The water concentrations in garnets are below the detection limit (<0.5 ppm). Clinopyroxene and o.the pyroxene in FMC samples (LN78, LP27, SD53) are atypical ('type 2'), with the nan water band occurring at lower wavenumbers (between 3515 and 3520 cm<sup>-1</sup>) the arise the other samples ('type 1'). The dashed lines at 3630 cm<sup>-1</sup> and 3515 cm<sup>-1</sup> highlight the main water bands in type-1 and type-2 spectra, respectively.

Fig. 5. Polarized FTIR spectra obtained of (a, b) clinopyroxene and (c) the average spectrum of garnets in sample YK01. Garnets are anhydrous (below the detection limit, <0.5 ppm). All spectra were normalized to 1-cm sample thickness. (a) Spectra over 3000–4000 cm<sup>-1</sup>, with peaks corresponding to OH bands. (b) Spectra over 1200–2400 cm<sup>-1</sup>, with peaks corresponding to lattice overtones specific to each orientation.  $Cpx_1^{90^\circ}/Cpx_1^{180^\circ}$  and  $Cpx_2^{90^\circ}/Cpx_2^{180^\circ}$  correspond to the orientations of the first and second thin sections, respectively.

Fig. 6. Comparison of the integrated absorptions of mineral cores and rims in (a) clinopyroxenes and (b) orthopyroxenes. Each point indicates the ratio of the integrated

absorption values measured at the rim and core of the same crystal. Most samples plot along on the 1:1 correlation line, highlighting the lack of core-rim zonation. This result indicates that the minerals did not experience diffusive re-equilibration, suggesting that there was no loss or gain of water during xenolith transport. The integrated absorption at the core and rim of each mineral are from profile measurements along the crystal; because the spectra are similar along the entire profiles, only the core and rim values are shown for clarity.

Fig. 7. (a) Clinopyroxene, (b) orthopyroxene, and (c) whole-rock (WR) water contents in FMC and AVP pyroxenites calculated using the calibration of '3ei' et al. (1995). The range shaded in blue indicates water contents of FMC peridotite cencliths reported by Gu et al. (2018) and Denis et al. (2015). The low water concentrations of whole-rock peridotites are related to their low modal proportion of pyroxenes and high modal proportion of olivine, which contains relatively little water.

Fig. 8. Major element contents versu s  $v_{4}$  ter concentrations in clinopyroxene and orthopyroxene in FMC and AVP pyroxenites. Mg# (= 100 × Mg/(Mg + Fe)) versus water content in (a) clinopyroxene and (c) orthopyroxene. (b) CaO versus water content in clinopyroxene. (d) A! O<sub>3</sub> ver us water content in orthopyroxene. Dashed purple lines highlight linear relation. Inps between major element contents and water concentrations in FMC pyroxenite minerals; any correlation is not obvious for the AVP samples. These results indicate that the water and major element contents of pyroxenites were acquired during a common equilibration (or re-equilibration) episode in the FMC, but during separate events in the AVP. Major element compositions are from France et al. (2015).

Fig. 9. Clinopyroxene vs. orthopyroxene water concentrations in peridotites from the

literature (Peslier et al., 2002, 2012; Peslier, 2010; Xia et al., 2010, 2013; Yu et al., 2011; Hao et al., 2012, 2014, 2016a, 2016b; Doucet et al., 2014; Warren and Hauri, 2014; Demouchy et al., 2015; Hui et al., 2015; Li et al., 2015; Peslier and Bizimis, 2015; Denis et al., 2018), pyroxenites from the literature (Warren and Hauri, 2014; Bizimis and Peslier, 2015, Hao et al., 2016a, 2018), and pyroxenites in this study (colored symbols). Red lines define the 1.5:1 and 3.5:1 partitioning of water between clinopyroxene and orthopyroxene; this equilibrium range encompasses the values of the pyroxenites studied herein, suggesting that the pyroxenes in these samples are in equilibrium with each other. Lines of  $D_{H-0}^{Cpx-G_Fx} = 1.5$  and 3.5 are added for reference.

Fig. 10. Whole-rock Ti/Eu ratios as a function of whole-rock (La/Sm)<sub>N</sub> ratios (normalized to chondrites; Anders and Grevesse, 1989). The (La/Sm)<sub>N</sub> ratio is used here as a proxy of metasomatic overprinting. The Ti/Eu ratios of FMC pyroxenites are negatively correlated with (La/Sm)<sub>N</sub> ratios, suggesting a carbot at the metasomatic overprint. See text for further details.

Fig. 11. (a) clinopyroxene (calculated using the calibration of Bell et al., 1995) and (b) whole-rock (WR) water concentrations (calculated from mineral-specific water contents and sample modal mineralogy) as a function of  $La/Sm_N$ . Both the clinopyroxene and whole-rock water contents are negatively correlated with  $(La/Sm)_N$  for the FMC pyroxenites.

Fig. 12. (a) Clinopyroxene, (b) orthopyroxene, and (c) whole-rock (WR) water concentrations (ppm H<sub>2</sub>O; calculated using the calibration of Bell et al., 1995) as a function of temperature (TNG09, °C; France et al., 2015). Correlations with  $R^2 = 0.95$  and 0.99 are observed for FMC minerals and whole-rocks, whereas only the whole-rock water contents show good correlation

with temperature estimates in the AVP with  $R^2 = 0.98$ .

Fig. 13. Whole-rock water concentrations as a function of  $H_2O/Ce$  ratio for pyroxenites from the literature and the present study, with mantle reservoirs shown for comparison. Ce and water concentration values for the reservoirs (DMM, FOZO, HIMU, EM, and PM) are from Bizimis and Peslier (2015) and are presented as a reference frame for magmas worldwide. Types I and II pyroxenites are from Li et al. (2018), Hawaiian  $_{\rm I}$  v<sup>-</sup>oxenites are from Bizimis and Peslier (2015), and Cr-, Al-, and Grt-pyroxenites are from Hav et al (2018). The most metasomatized samples from the literature (Hao et al., 2012: sy mbols outlined in black) are characterized by low H<sub>2</sub>O/Ce ratios.

Fig. 14. Non-modal, incremental batch-r elting models of representative FMC and AVP garnet pyroxenites presented on a  $[H_2O_1^1 (ppm)$  versus  $H_2O/Ce$  diagram.  $[H_2O]$  and  $H_2O/Ce$  strongly vary at low degrees of melting, and reach  $H_2O/Ce$  values similar to those of the starting pyroxenites after >10% partial melting.

Table 1: The distribution of o thopyroxene, clinopyroxene, garnet, and whole-rock water contents (ppm) in peridotice and pyroxenite as a function of geological origin.

Source	Minerals	Mean water content in peri-	Standard deviation	Mean water content in py-	Standard de- viation
		dotite (ppm)		roxenite (ppm)	
Craton <sup>[b][c]</sup>	Орх	116	94	56	49
[d] [e] [f] [g] [h] [l]	Срх	240	185	146	94
[j] [k] [l] [p] [q] [t]	Grt	17	25	9	17
	WR	58	48	216	224
Off-craton	Орх	146	69	143	171
[a] [c] [m] [s] [u]	Срх	347	143	363	180
	Grt	36	92	-	-
	WR	81	36	268	72
Oceanic	Орх	125	97	201	49
lithosphere	Срх	290	111	425	78

[c] [n] [o] [r] [v]	Grt	-	-	23	5
	WR	38	36	333	65

<sup>[a]</sup> Peslier et al.  $(2002)^{[b]}$  Peslier et al.  $(2012)^{[c]}$  Peslier  $(2010)^{[d]}$  Xia et al.  $(2010)^{[e]}$  Xia et al.  $(2013)^{[f]}$  Yu et al.  $(2011)^{[g]}$  Hao et al.  $(2012)^{[h]}$  Hao et al.  $(2014)^{[i]}$  Hao et al.  $(2016a)^{[j]}$  Hao et al.  $(2016b)^{[k]}$  Hao et al.  $(2018)^{[l]}$  Doucet et al.  $(2014)^{[m]}$  Warren and Hauri  $(2014)^{[n]}$  Bizimis and Peslier  $(2015)^{[o]}$  Demouchy et al.  $(2015)^{[p]}$  Li et al.  $(2015)^{[q]}$  Li et al.  $(2018)^{[r]}$  Peslier and Bizimis  $(2015)^{[s]}$  Kilgore et al.  $(2018)^{[t]}$  Kilgore et al.  $(2018)^{[t]}$  Kilgore et al.  $(2019)^{[v]}$  Schaffer et al.  $(2019)^{[v]}$  Ashley et aal. (2020)

Location	French	French Massif Central			Adoma va Volcanic Plateau					
Sample	LN78	LP27	SD53	Le Pouget	Y.701	YK03	YK05	YK12	YK13	YK16
Thickness of sections (µm)	233- 316	260- 306	154- 274	179- 207	500- 560	239- 295	229- 298	167- 328	231- 293	169- 273
Modal composition										
Clinopyroxene (%)	40	70	55	61	60	45	60	45	65	70
Orthopyroxene (%)	22	5	10	3	0	15	17	40	17	0
Garnet (%)	30	15	20	33	30	26	23	10	8	28
Spinel (%)	8	10	5	3	10	14	0	5	10	2
Equilibrium temperature from major elements <sup>b</sup>	S									
T (°C)	1017	930	866	716		872	916	888	977	
(La/Sm) <sub>N Cpx</sub> <sup>b</sup>	1.018	1.454	0.227	0.034	0.244	0.774	0.111	0.176	1.875	1.380
$(La/Sm)_{NWR}^{b}$	1.771	1.880	0.808	0.037	0.248	0.728	0.327	0.181	1.771	1.536

Table 2. Modal compositions, equilibrium temperatures, and  $(r_{2})^{\prime}$ m)<sub>N</sub> ratios.

<sup>a</sup> Although SD53 contains orthopyroxene, they are too altered and contain too many fluid

inclusions for reliable analyses.

<sup>b</sup> Data from France et al. (2015).

Location	French	Massif C	Central		Adamay	wa Volca	nic Plate	eau		
Sample Integral absorption	LN78	LP27	SD53	Le Pouget	YK01	YK03	YK05	YK12	YK13	YK16
Cpx (cm <sup>-</sup> <sup>2</sup> )	264	554	849	1098	911	1106	1065	1118	1179	1619
$\frac{\text{Error (cm}^{-2})}{2}$	(+29;- 16)	(+46;- 86)	(+66;- 102)	(+299; -158)	(+87;- 136)	(+77;- 130)	(+69;- 55)	(+101; -124)	(+112; -142)	(+51;- 250)
Number of cpx analyses per sample	29	31	30	24		29	23	30	28	34
$Opx (cm^{-2})$	301	383		515		765	711	614	731	
$\frac{\text{Error (cm}^{-2})}{2}$	(+74;- 47)	(+39;- 36)		(+22;- 53)	J.C	(+97;- 116)	(+20;- 49)	(+21;- 33)	(+92;- 34)	
Number of opx analyses per sample	30	30	2.	16		30	27	28	23	
Water concentrat ion (Bell et al., 1995)		5	5							
Cpx [H <sub>2</sub> O] (ppm)	112	234	359	465	386	468	451	473	499	685
Error (ppm)	(+25 ; -15)	(+45 ; -53)	(+66 ; -69)	(+185 ; -100)	(+78 ; -85)	(+82 ; -89)	(+77; -59)	(+94 ; -87)	(+102 ; -96)	(+92 ; -154)
Opx [H <sub>2</sub> O] (ppm)	61	77		104		155	144	124	148	
Error (ppm)	(+22 ; -13)	(+15 ; -12)		(+14 ; -18)		(+35; -33)	(+17; -20)	(+15 ; -15)	(+33 ; -17)	
Water										

# Table 3. Water concentrations, $H_2O/Ce$ ratios, and water partition coefficients.

concentrat ion (Libowitz ky and Rossman, 1997)										
Cpx [H <sub>2</sub> O] (ppm)	85	179	275	357	296	360	344	362	382	525
Error (ppm)	(+9 ; - 5)	(+15 ; -28)	(+21 ; -33)	(+96 ; -52)	(+28 ; -44)	(+25 ; -43)	(+23 ; -17)	(+33; -40)	(+36 ; -46)	(+16 ; -81)
Opx [H <sub>2</sub> O] (ppm)	78	100		135		201	185	160	191	
Error (ppm)	(+19 ; -12)	(+10 ; -10)		(+6 ; - 14)		(+26 ; -30)	(~5;- 13)	(+6 ; - 8)	(+24 ; -9)	
Other data						$\mathbb{N}_{-}$				
[H <sub>2</sub> O] Whole rock (ppm) <sup>a</sup>	58	168	198	287	231	234	295	263	349	480
Error (ppm)	(+15 ; -9)	(+32; -38)	(+36 ; -38)	(+113 ; -61,	(+47 ; -51)	(+42 ; -45)	(+49 ; -39)	(+48 ; -45)	(+ 72 ; -66)	(+64 ; -108)
$[H_2O]$ Whole rock (ppm) <sup>b</sup>	51	130	151	2 22	177	192	238	227	281	368
Error (ppm)	(+8 ; - 5)	(+11 · -?0)	(† 12 ; -18)	(+59 ; -32)	(+17 ; -26)	(+15 ; -24)	(+14 ; -13)	(+17; -22)	(+28 ; -31)	(+12 ; -57)
H <sub>2</sub> O/Ce Whole rock <sup>c</sup>	30	44	166	2388	257	202	508	370	79	201
Error	(+8 ; - 5)	(+8 ; - 10)	(+31 ; -32)	(+945 ; -512)	(+52 ; -56)	(+36 ; -39)	(+84 ; -66)	(+68 ; -64)	(+16 ; -15)	(+27 ; -45)
D <sup>Cpx/Melt</sup> <sub>d</sub> H2	0.019	0.019	0.018	0.016	0.017	0.015	0.015	0.015	0.017	0.017
Error	(+0.01 0;- 0.006)	(+0.01 0;- 0.007)	(+0.01 0;- 0.006)	(+0.00 9;- 0.006)	(+0.00 9;- 0.006)	(+0.00 8;- 0.005)	(+0.00 7;- 0.005)	(+0.00 8;- 0.005)	(+0.00 9;- 0.006)	(+0.00 9;- 0.006)
D <sup>Opx/Melt</sup> <sub>d</sub> H2	0.015	0.015		0.009		0.010	0.010	0.010	0.010	
Error	(+0.00	(+0.00		(+0.00		(+0.00	(+0.00	(+0.00	(+0.00	

6;-	6;-	3;-	3;-	3;-	3;-	3;-	
0.004)	0.004)	0.002)	0.002)	0.002)	0.002)	0.003)	

<sup>a</sup> Water concentrations calculated using the method of Bell et al. (1995) and modal

composition.

<sup>b</sup> Water concentrations calculated using the method of Libowitzky and Rossman (1997) and modal composition.

<sup>c</sup> Data from France et al. (2015).

<sup>d</sup> Calculated following O'Leary et al. (2010) as

 $\ln D_{\rm H_20}^{\rm (Cpx-Melt)} = -4.2(\pm 0.2) + 6.5(\pm 0.5) X_{\rm Al^{\rm IV}}^{\rm Cpx} - 1.0(\pm 0.2) X_{\rm Ca}^{\rm Cpx-Melt)} = -4.2(\pm 0.2) + 6.5(\pm 0.5) X_{\rm Al^{\rm IV}}^{\rm Cpx} - 1.0(\pm 0.2) X_{\rm Ca}^{\rm Cpx-Melt)} = -4.2(\pm 0.2) + 6.5(\pm 0.5) X_{\rm Al^{\rm IV}}^{\rm Cpx} - 1.0(\pm 0.2) X_{\rm Ca}^{\rm Cpx}$  and  $\ln D_{\rm H_20}^{\rm (Opx-Melt)} = -4.2(\pm 0.2) + 6.5(\pm 0.5) X_{\rm Al^{\rm IV}}^{\rm Cpx} - 1.0(\pm 0.2) X_{\rm Ca}^{\rm Cpx}$ 

 $-5.66(\pm 0.11) + 8.4(\pm 1.1)X_{Al^{VI}}^{Opx} + 10(\pm 2)X_{Ca}^{Opx}.$ 

#### **Declaration of interests**

The authors declare that they have no know... competing financial interests or personal relationships that could have appeared to inf<sup>2</sup> aence the work reported in this paper. The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:







b.





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Figure 4



Figure 5



Figure 6



#### Figure 7





Figure 9



Figure 10



Figure 11





Figure 13



Figure 14