1	Grain scale processes recorded by oxygen isotopes in olivine-		
2	hosted melt inclusions from two MORB samples		
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15			
16	Abstract		

17 Although olivine-hosted melt inclusions from mid ocean ridge basalts (MORB) are 18 commonly used as a proxy for mantle composition, these melt inclusions generally show 19 larger elemental and isotopic compositional variation than their host lavas and the origin of 20 these heterogeneities remains disputed. Here we present oxygen isotope data from melt 21 inclusions hosted in olivine from two samples from the Mid-Atlantic ridge. Melt inclusions 22 from different crystals within the same sample show more than 2.5‰ δ^{18} O variation within 23 each sample, which is nearly eight times the analytical error of 0.3% (2 standard deviations) and five times the δ^{18} O range in unaltered MORB. Measured δ^{18} O in melt inclusions do not 24 correlate with common magmatic tracers, and $\delta^{18}O$ measured in the host olivines suggest a 25 maximum of 1‰ δ^{18} O source heterogeneity. Less than half of the melt inclusions from each 26 27 sample are in equilibrium with their host crystals; the remaining melt inclusions have either 28 lower or higher olivine-melt oxygen isotope partition coefficients compared to the theoretical 29 equilibrium values. Here we discuss several potential processes that could contribute to these 30 observations, but none satisfactorily explain the olivine-melt inclusion oxygen disequilibrium that we observe in these samples. Nevertheless, it seems clear that the variability of δ^{18} O in 31 32 melt inclusion from two MORB samples do not record only common magmatic process(es), but rather a localized grain scale process. Any δ^{18} O variation in melt inclusions should thus be 33 34 interpreted with caution.

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36 Keywords

Olivine-hosted melt inclusions; MORB; Mid-Atlantic Ridge; oxygen isotopes; grain scale
processes; heterogeneity; SIMS

39

40 Introduction

Olivine-hosted melt inclusions are often used to study the origin and evolution of mantlederived magmas (for a review, see Kent, 2008; Schiano, 2003 and references therein). Melt inclusions are physically isolated from their host magma, which undergoes different processes during magmatic evolution after the entrapment of melt inclusions (for example, magma mixing, assimilation, degassing). An important observation is that larger elemental and 46 isotopic compositional variations are generally observed in melt inclusions when compared to 47 whole-rock or glass compositions. A widespread interpretation is that variations measured in 48 melt inclusions reflect magmatic diversity that is not recorded in whole rock compositions 49 due to obscuration during later processes (e.g., Kent, 2008). In some cases, variations could 50 also be related to small-scale reactions in the melt in which the olivine is growing 51 (Danyushevsky et al., 2004).

52 In mid-oceanic ridge basalts (MORB), most melt inclusions have trace-element compositions 53 in the range of MORB lavas, but there are also some clear exceptions to this general 54 observation (see Schiano, 2003 for a review). These 'anomalous' trace-element compositions 55 have usually been interpreted to reflect near-fractional melting of a depleted mantle and/or 56 spinel lherzolite, or of hydrothermally altered peridotite (e.g., Gurenko and Chaussidon, 1995; 57 Nielsen et al., 2000; Sobolev and Shimizu, 1993). The anomalous trace-element compositions 58 are not found in MORB rocks, as MORBs represent aggregates of different melt fractions 59 (e.g., Jin et al., 1994; McKenzie, 1984; Sobolev and Shimizu, 1993). Melt reaction within the 60 crust, i.e. within a crystal mush, may also explain the larger compositional variations 61 observed in some MORB melt inclusions compared to lavas (e.g., Danyushevsky et al., 2004; 62 Kamenetsky and Gurenko, 2007).

63 Major and trace element variability in melt inclusions from MORB samples are also 64 sometimes observed within a single phenocryst (Shimizu, 1998), which indicates that melt 65 inclusions are not necessarily in equilibrium with their hosts. Such small-scale diversity could 66 be generated by trace element diffusion within the melt during melt inclusion formation by host dissolution-reprecipitation, or by boundary layer entrapment during fast, diffusion-67 controlled growth or melt transport (e.g., Baker, 2008; Cottrell et al., 2002; Faure and 68 69 Schiano, 2005; Manzini et al., 2017; Van Orman et al., 2002). Other processes, such as diffusion of some elements (e.g., H, Fe) toward the host olivine or the host melt are 70

responsible, in some cases, for elemental variation in melt inclusions (e.g., Danyushevsky et
al., 2000a; Gaetani et al., 2012).

73 To date, no stable isotope data from melt inclusions within olivine from typical MORB samples have been published. Although $\delta^{11}B$ and $\delta^{18}O$ were reported in some olivine-hosted 74 75 melt inclusions from Iceland, the petrogenetic history of these samples is fundamentally 76 different from typical MORB settings (Gurenko and Chaussidon, 1997, 2002; Hartley et al., 2013). In this study, we focus on δ^{18} O values from melt inclusions within MORB samples, as 77 78 the systematics of this stable isotope system are well constrained. Based on a compilation of 79 δ^{18} O data in Neogene volcanic rocks, Harmond and Hoefs (1995) concluded that bulk MORB rocks have δ^{18} O compositions of 5.7% \pm 0.2%. Because the variation reported is similar to 80 the typical analytical uncertainty reported in the literature for individual δ^{18} O measurements 81 82 of silicate rocks at that time, the authors concluded that MORB have a uniform δ^{18} O 83 signature. More recently, fresh MORB glasses from the Mid-Atlantic Ridge, Indian Ocean 84 and east Pacific Ridge were analyzed by laser fluorination and show a narrow range of 85 oxygen isotope ratios from 5.37 ± 0.01 to 5.81 ± 0.04 ‰ (1 σ), with an average of +5.50 ‰ (Eiler et al., 2000). In addition, δ^{18} O is thought to not fractionate during melting (e.g., Eiler, 86 87 2001). As such, if melt inclusions are representative of the MORB source, they should show 88 similar values within a relatively restricted range; however, as melt inclusions generally show larger chemical variations than those recorded by the bulk rock, δ^{18} O in melt inclusions from 89 90 MORB samples might also display larger isotopic variabilities than MORB glasses. 91 Constraining the variability of oxygen isotopes in melt inclusions from MORB samples is important, as δ^{18} O variations within these melt inclusions serve to assess the importance of 92 93 fluid interaction, magma mixing and/or assimilation of oceanic crust (Gurenko and 94 Chaussidon, 2002; Hartley et al., 2013). In this study we report δ^{18} O data from melt inclusions within two different MORB samples from the northern Mid-Atlantic Ridge. We observe a 95

96 never before reported large range of δ^{18} O compositions within these melt inclusions, up to 97 2.5‰, and discuss the reported data in term of magmatic processes, olivine dissolution-98 precipitation and diffusion.

99

100 Sample description and geological context

101 The studied olivine-hosted melt inclusions are from two dredged basalts from the North Atlantic ridge. Sample ARP73-10-03 is a picrite from the FAMOUS zone (36°8372'N, 102 103 33°2482'W) that contains a little more than 20% olivine phenocrysts (Fo_{87,2-91,3}), rare 104 clinopyroxene, and plagioclase xenocrysts. The crystals are surrounded by a hyalocrystalline 105 groundmass composed of glass, plagioclase (An₇₅₋₈₀) and olivine microlites (Fo₈₆) (Laubier et 106 al., 2007). Laubier et al. (2007) suggest that the variable major and trace element 107 compositions of ARP73-10-03 melt inclusions reflect polybaric partial melting of a 108 homogeneous mantle, followed by mixing in various proportions at either different degrees of 109 melting or in different parts of the melting system.

110 CH77-DR6-203 is a dredged basalt from the 14° MAR triple junction (14.12°N, 45°W). This 111 fresh sample is highly enriched in volatiles, with CO₂ in vesicles and water in the glass, 112 (popping rock; Javoy and Pineau, 1991) and shows a compositional anomaly with enriched 113 (Nb/Zr)_N, (Ta/Hf)_N and (La/Sm)_N compared to adjacent segments which may result from 114 mantle heterogeneity (Bougault et al., 1988). The mantle heterogeneity hypothesis is also 115 supported by Sr and Nd isotopes in MORB rocks from different segments of the triple 116 junction (Dosso and Bougault, 1986). The melt inclusions studied here are hosted in 1-5 mm, 117 polyhedral olivines containing one or several melt inclusions. They are glassy, sub-spherical, 118 and have a radius of 40-350 µm. They contain no daughter minerals. Rare shrinkage bubbles 119 are present, but represent only a small fraction of the total volume.

121 Methods

122

123 Sample preparation

124 Polished olivines containing exposed melt inclusions were pressed into 1-inch indium mounts, 125 along with the glass standards (BHVO-2G, BCR-2G and BIR-1G) and San Carlos olivine. 126 Care was taken to mount them within 8 mm of the center of the mount. A separate mount was 127 made for each sample. SEM images were acquired at the University of Lausanne using a 128 Tescan Mira LMU field-emission scanning electron microscope operated at 20 kV and 0.2-0.4 129 nA probe current to ensure that the melt inclusions were totally glassy. Melt inclusions with 130 cracks or crystals were discarded. Topography of each mount was measured using a Brucker 131 GTA-K white light interferometer.

132

133 Electron microprobe

134 Major element compositions and X-ray distribution maps of melt inclusions and olivines were 135 acquired using a JEOL 8200 Superprobe electron microprobe analyzer. Analytical conditions 136 for the quantitative measurement of olivine were 15kV at 20nA and a 5 µm beam diameter. 137 Counting times were 30 s for Si, Mg, Mn and Fe and 40 s for Al, Ca, Ni and Cr. Analytical 138 conditions for analyses of melt inclusions were 15kV, 10 nA and a 10 µm beam diameter. 139 Counting times were 30 s for all elements except for K (20 s) and Na (16 s). A glass standard 140 (KL2-G) was used for SiO₂ and Al₂O₃. All other elements were calibrated on minerals. A 141 grain of ML3B-G was used as an internal standard to check the calibration.

142

143 Secondary ion mass spectrometer

Oxygen isotope ratios, volatiles (H₂O, CO₂, Cl, F) and trace-element (Ti, V, Sr, Y, Nb, Ba,
La, Ce, Nd, Sm, Eu, Yb) concentrations were determined using the CAMECA ims1280-HR at
the SwissSIMS laboratory.

147 Oxygen isotopes

148 Analytical conditions were similar to those described by Seitz et al. (2017): Throughout 149 ¹⁸O/¹⁶O measurements of melt inclusions and olivines, the sample was sputtered with a Cs⁺ 150 primary beam of 2 nA, which was raster over 15 µm. An electron gun was utilized in order to 151 compensate for sample charging. Samples were pre-sputtered for 30 s, followed by 80 s of analysis (20 cycles of 4 s). Automatic centering of secondary ions on field and contrast 152 apertures were performed before each analysis. Measurements of ¹⁶O and ¹⁸O were performed 153 simultaneously via two Faraday cups, using 10^{10} and 10^{11} resistors, respectively. The intensity 154 of ¹⁶O was typically ~ 2.2×10^9 counts per second for glasses and ~ 1.7×10^9 counts per second 155 156 for olivine. Internal error (2 standard error), reflecting the counting statistics of one analysis, 157 was usually better than 0.3% for melt inclusions and olivines.

For melt inclusions, instrumental mass fractionation (IMF) was corrected using a set of 9 158 159 international glass standards with compositions ranging from basaltic to rhyolitic (NKT-1G, 160 GOR132-2G, BIR-2G, BHVO-2G, ML3B-2G, BCR-2G, StHs6/80G, UoE-Lipari, ATHO-G; 161 Hartley et al., 2012; Jochum et al., 2006). The matrix effect was corrected using a linear 162 regression of IMF with SiO₂ (Figure DR1-A). Two to three standards (BHVO, BIR and BCR), 163 with compositions closely bracketing the melt inclusion compositions, were added on each 164 mount to monitor and correct for potential instrumental drift. For olivines, IMF was corrected 165 using an in-house San Carlos olivine standard (Fo 90.1), as well as three in-house olivine 166 reference materials (Fo 99.6, Fo 91.9 and Fo 37.5; Fig DR1-B).

167 At the outset of each session, we performed at least eight measurements of a running standard 168 (i.e., measured repeatedly during the session; BHVO for melt inclusions sessions and San 169 Carlos for olivine sessions), in order to determine standard reproducibility (noted hereafter in 170 2 standard deviation, 2SD). The 2 SDs were better than 0.3% for melt inclusions sessions, 171 and 0.4‰ for olivines sessions. To monitor the instrument stability, running standards were 172 measured every 8 to 12 unknowns and gave a reproducibility of 0.3‰ 2SD, similar to the 173 eight measurements at the outset of each session. Accuracy was checked using one of the two 174 to three standards inserted in each mounts and was similar to reproducibility (0.3‰ 2SD and 175 0.4‰ 2SD for melt inclusions and olivines, respectively).

176

177 Trace elements

178 Phosphorous concentrations within melt inclusions were measured using the same analytical 179 conditions described by Manzini et al. (2017): Trace-element compositions of melt inclusions 180 and olivine were analyzed using a 5-6 nA O⁻ primary ion beam, resulting in an analysis spot of about 20 µm. A mass resolution of 5000 was used. Samples were pre-sputtered for 90 s, 181 using a 25 µm raster. Trace elements were quantified by measuring the signal from the 182 following isotopes: ⁴⁸Ti, ⁵¹V, ⁸⁸Sr, ⁸⁹Y, ⁹³Nb, ¹³⁸Ba, ¹³⁹La, ¹⁴⁰Ce, ¹⁴²Nd, ¹⁵²Sm, ¹⁵³Eu, ¹⁷²Yb. 183 These measurements were performed in mono-collection mode by peak switching throughout 184 one analytical run on an electron multiplier (EM) collector. ⁴⁰Ca was measured on a FC 185 186 collector in the same analytical run as the other trace elements and was used as reference element. Using ⁴⁰Ca instead of ³⁰Si (or ²⁸Si) during melt inclusion analyses allowed us to 187 check for possible contamination by olivine throughout the analyses. A waiting time was used 188 189 to obtain a steady state sputtering condition, and each analysis consisted of six cycles. 190 Waiting and counting time, typical standard reproducibility and typical internal error are 191 given in Table DR1. BHVO-2G was used as standard for melt inclusions, using the USGS192 certified values for the trace elements.

193

194 Volatile elements

195 A Cs⁺ primary ion beam with a current of 1.5 nA was used to measure volatile elements 196 (H₂O, CO₂, Cl, F, S); throughout these analyses an electron gun was used to compensate the 197 charges and to avoid charging of the sample surface. The secondary beam was accelerated at 198 10 kV, and the sample was pre-sputtered applying a 15 μ m raster for 210 s. Elements were 199 measured in mono-collection mode, with a 10 µm raster during 12 cycles. Total analysis time, 200 including pre-sputtering, was approximately 14 minutes. The field aperture was set to 3000 201 µm, entrance slit to 60 µm and exit slit to 240 µm. This configuration allowed a mass 202 resolution of 5000, sufficient to fully discriminate the interferences. BHVO-2G, BCR-2G, 203 BIR-1G, ALL, NS and ALV (Helo et al., 2011) were used as standards. BHVO-2G was then 204 used as a running standard to check instrument stability. Background was determined by 205 analysis of a synthetic anhydrous olivine. Additional information is given in Table DR1.

206

207 **Results**

A total of 34 melt inclusions within 30 olivines were measured from sample CH77-DR6-203, as were 17 melt inclusions within 13 olivines from sample ARP73-10-03. The results of these analyses are summarized in Table DR2. Oxygen isotope ratios in olivines are reported in Table DR3.

Analyzed olivines are characterized by a high forsterite content (Fo = Mg/Fe+Mg) ranging from 88.8 to 91.4 for ARP73-10-03 and from 84.0 to 90.9 for CH77-DR6-203. Major-element compositions of melt inclusions were corrected for the effect of post-entrapment 215 crystallization (PEC) of olivine at the wall of melt inclusion, using the Petrolog software 216 package (Danyushevsky and Plechov, 2011). Parameters used for Petrolog calculations are 217 summarized in the footnote of Table DR2. Our calculated PEC indicates that between 5.9 and 218 18.1% of olivine crystallized after melt inclusion trapping. Major elements are less variable after PEC correction (e.g., SiO₂ variation of 2.8 wt. % before and 1.6 wt. % after correction; 219 220 Table DR2), and all major-element compositions mentioned hereafter are corrected for PEC. 221 Melt inclusions from ARP73-10-03 have a narrow range of major-element compositions, with 222 SiO₂ ranging from 50.0-50.4 wt. %, MgO from 11.2-12.8 wt. % and K₂O from 0.10-0.15 wt. 223 %. CH77-DR6-203 shows a slightly larger compositional range in major elements, with SiO₂ 224 content from 49.2-50.8 wt. %, MgO from 7.8-10.0 wt. %, and K₂O from 0.20-0.55 wt % 225 (Figure 1A). Major-, volatile- (Cl, F, S) and trace-element concentrations measured in 226 ARP73-10-03 fall within the same range as those reported by Laubier et al. (2012) for melt 227 inclusions within the same sample and are consistent with glass N-MORB compositions 228 (Jambon et al., 1995; Naumov et al., 2014; Wallace and Carmichael, 1992). Water and CO₂ 229 concentrations are also in the typical range of MORB glass values (Danyushevsky et al., 230 2000b; Magenheim et al., 1995; Wallace and Carmichael, 1992). The melt inclusions from 231 CH77-DR6-203 are enriched in volatiles (to a lower extent for CO₂) when compared with 232 MORB glass values, and have trace-element compositions similar to typical E-MORB glasses 233 (e.g., Figure 1B modified from Gale et al., 2013).



Figure 1: Oxygen isotope values for MI plotted against the K₂O concentration (A) and the La/Sm chondrite-normalized ratio (McDonough and Sun, 1995) (B). Line separating D-, N- and E-MORB are from Gale et al., 2013a. Fresh MORB glasses studied represented by white triangles in the figure, are plotted for comparison (Eiler et al., 2000).

Oxygen isotopes from melt inclusions show > 2.3% variation within each sample ($\delta^{18}O =$ 240 $+5.0 \pm 0.17$ to $+7.2 \pm 0.16$ %, with an average of +6.2% for CH77-DR6-203; and $\delta^{18}O = +4.5$ 241 \pm 0.3 to +7.1 \pm 0.6‰, average +5.7‰ for ARP73-10-03). These variations are about eight 242 times larger than the analytical uncertainty (0.3‰, 2SD) and about five times larger than the 243 variations observed for bulk δ^{18} O of unaltered MORB glasses (<0.5%; Figure 2). Before IMF 244 correction, data show similar δ^{18} O variations (2.2‰). This is explained by the relatively 245 246 narrow range of SiO₂ of the melt inclusions within a sample, which indicates that the applied 247 IMFs are relatively similar ($\pm 0.13\%$) for all melt inclusions from the same sample. ARP73-248 10-03 is the closest to MORB glass composition in term of major, trace elements and oxygen isotopes. Nevertheless, 8 melt inclusions out of 21 have 'anomalous' δ^{18} O composition (i.e., 249 statistically – 2 SD – outside the range of MORB glass bulk values). Within a single olivine, 250 251 δ^{18} O from different melt inclusions are statistically different in the ARP73-10-03 sample (3) 252 pairs, variation up to 2.2‰), whereas in the CH77-DR6-203 sample, melt inclusions hosted in a single olivine have values within analytical uncertainty δ^{18} O (3 pairs, variation less than 0.4‰; Figure 3). In both samples, olivines show smaller variability than their hosted melt inclusions: the ARP73-10-03 olivine have δ^{18} O values of +4.9 to +5.7‰ and olivines from CH77-DR6-203 have δ^{18} O ranging from +5.0 to +6.5‰ (Table DR3, Figure 2).





Figure 2: Comparison of oxygen isotope variations in MORB glasses from Mid-Atlantic Ridge, Indian ocean and east-Pacific Ridge (n = 28) (Eiler et al., 2000) and melt inclusions (MIs) from the two studied MORB samples (ARP73-10-03 and CH77-DR6-203). For a comparison, range of δ^{18} O composition measured in the olivines hosting the melt inclusions are also plotted (dashed line contoured fields). Melt inclusions show larger variability compare to MORB glasses and to their host olivines.



Figure 3: Oxygen isotope ratio of melt inclusions, compared to the δ^{18} O of their host olivine crystals. Melt-olivine equilibrium fractionation (Δ^{18} O; black line) is from Matthews et al. (1998). Only 42 and 46% of melt inclusions from ARP_73-10-03 and CH77-203-DR6 are in equilibrium with their host, within analytical error.

269

270 Discussion

271 To date, no oxygen isotope data have been reported for olivine-hosted melt inclusions from 272 MORB samples. Only two studies (Gurenko and Chaussidon, 2002; Hartley et al., 2013) 273 reported δ^{18} O in olivine-hosted melt inclusions, and these data are from 7 samples taken from Iceland. These studies found δ^{18} O variation reaching a maximum of 1.2 ‰ (+4.0 to +5.2‰, 274 275 sample 14161G, n = 3 melt inclusions; Gurenko and Chaussidon, 2002) and 0.95 ‰ (+3.05 to 276 +4.0‰, sample N18, n = 9 melt inclusions; Hartley et al., 2013), respectively. These 277 variations were interpreted to reflect either mixing of the ascending magma with melt lenses that had previously interacted with the low- δ^{18} O Icelandic crust (Hartley et al., 2013), or 278 279 heterogeneity in the Icelandic mantle or mixing of two end-members (Gurenko and 280 Chaussidon, 2002). Melt inclusions from the two MORB samples measured in our study both 281 show more than 2.3‰ variation in δ^{18} O, a larger variation than that reported for olivine-282 hosted melt inclusions from Iceland. Hereafter, the δ^{18} O variability is discussed in term of 283 analytical versus magmatic processes.

284

285 Analytical artefacts

286 Oxygen isotope analyses via SIMS are very sensitive to the sample geometry, i.e. the placement of the analyzed grains in the mount and the topography of the analyzed material 287 288 (e.g., Kita et al., 2009). All grains and reference material have been placed within 8 mm of the 289 center of the mount, in order to avoid X-Y effects due to deformation of the electrostatic field 290 close to the edge of the sample holder (Peres et al., 2013) (see methods section for more 291 details). Reproducibility of the measurements of a homogeneous reference material for 292 oxygen isotope is typically <0.3% (2 SD) when the surface relief is 1 µm, whereas it increases up to 3‰ (2SD) when topography is on the order of 10-40 µm (Kita et al., 2009). 293 294 Our mounts show $< 4 \mu m$ topography both between the indium and grains, and between 295 olivine and melt inclusions. Two different cases are presented in Figure 4: First, Figure 4-A shows olivine CH77-DR6-203 37 hosting 2 melt inclusions with similar δ^{18} O (within the 296 297 0.3% 2SD calculated for a session, $+6.37\pm0.21\%$ and $+5.97\pm0.16\%$) and a topography of 1µm. Olivine ARP73-10-03 16, shown in Figure 4-B, has δ^{18} O in melt inclusions which vary 298 299 from $+5.15 \pm 0.21\%$ to $+6.42 \pm 0.26\%$. This olivine has a topographical variation of up to 4 300 µm, with a difference in topography of 1.5 µm between melt inclusions A, C and D 301 (maximum 1.3% difference in δ^{18} O), and 2.5 µm between melt inclusions A and B (similar 302 δ^{18} O). Melt inclusions are less than 1 µm lower compared to the olivine, which should not 303 affect the analyses: 17 analyses made following 2 profiles along perpendicular axes from wall

to wall in melt inclusion A, and 7 in melt inclusion B yield a reproducibility better than 0.3% 304 305 (2SD) (Table DR2). Also, multiple measurements (n = 75) located at the edge or in the center of the olivine grain ARP73-10-03 16, which shows the largest topographical variation, yield 306 data clustering within 0.45‰ (2SD; Table DR3), comparable to the reproducibility obtained 307 308 during the same session for San Carlos reference material that has negligible topography (0.40‰, 2SD). Moreover, δ^{18} O from melt inclusions within ARP73-10-03 16 A and B were 309 310 measured twice, in two different sessions and similar values were obtained. As the mount was placed in the sample holder with a different orientation, this confirms that measured $\delta^{18}O$ 311 312 variation cannot be due to sample topography or orientation. Based on these observation, we find it highly unlikely that topography of the grains is responsible for the large variation of 313 314 δ^{18} O in the melt inclusions analyzed in this study. As the calibration of the SIMS analyses 315 reported here were made using several reference materials covering the range of melt inclusions major elements (Fig. DR2). Hence we conclude that the observed δ^{18} O variations 316 317 are real in melt inclusions and do not represent analytical artefacts.



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Figure 4: Example of profilometer images of the studied samples. A – ARP73-10-03_16 olivine; B –
CH77-DR6-203_37 olivine. The number of points is specified when more than one analyses was
obtained in the same inclusion, and the error (2SD) represents the reproducibility on the repeated
measurements.

324 Effects of different magmatic processes

Partial melting causes large variations in the major- and trace- element concentrations of MORBs (e.g., Frey et al., 1993; Schilling et al., 1983; Zindler et al., 1979), but is not expected to largely fractionate oxygen isotopes. Indeed, Bindeman et al. (2012) estimate that extraction of 15% of the melt, assuming equilibrium is maintained throughout the melting, will change δ^{18} O by only 0.05‰. Even high degrees of partial melting (up to 42%) of a mantle source will

create less than 0.2 $\% \delta^{18}$ O variations (e.g. Eiler, 2001). Variation in the degree of partial 330 331 melting will thus not be resolvable by SIMS, given the typical precision of ~0.3‰ 2SD. Hence, the lack of correlation between δ^{18} O and the abundance of incompatible elements such 332 333 as [La/Sm]_N for example (Figure 1), which are sensitive to degree of melting, indicates that variable extents of partial melting cannot explain the large δ^{18} O scatter measured in this study. 334 335 Interestingly, ARP73-10-03 has a composition similar to that of typical MORB glass (N-336 MORB) in terms of [La/Sm]_N, K₂O and volatile contents, and average δ^{18} O composition (e.g., 337 Eiler et al., 2000; Gale et al., 2013), whereas CH77-DR6-203 has higher (La/Sm)_N, volatiles, K₂O contents, and δ^{18} O than that of N-MORB glass (Figure 1). Differences in (La/Sm)_N, 338 339 volatiles and K₂O contents between the two samples could reflect different degrees of melting, but the different average δ^{18} O rather suggests melting of an enriched mantle source 340 341 for CH77-DR6-203. Despite the melting of different mantle sources for the two samples, resulting in different δ^{18} O for their respective melt inclusions, a similar variation of δ^{18} O 342 343 $(\sim 2.5\%)$ is observed in melt inclusions for both samples.

344 The lack of correlation between δ^{18} O of the melt inclusions and the forsterite content of their 345 host olivines do not support large fractionation of oxygen isotopes during magmatic evolution 346 of the studied samples (Figure 5). This observation is consistent with an experimental study 347 examining crystallization of a MORB melt dominated by olivine and plagioclase fractionation 348 (Tormey et al., 1987), which shows that the oxygen isotopic composition of the melt should not vary by more than 0.1 ‰ for a MgO content between 3 to 8 wt% (Eiler, 2001). Recently, 349 350 Bucholz et al. (2017) modelled change in δ^{18} O in tholeiite during fractional crystallization and also found minor change in δ^{18} O (up to 0.35% for composition comparable to ARP73-10-03 351 352 and CH77-203-DR3 melt inclusions).



Figure 5: Plot of oxygen isotopes of olivine and of melt inclusions plotted against the Fo content ofthe host olivines. Full symbols are for melt inclusions, empty symbols represent olivines.

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357 Considering the low volatile concentrations in MORB lavas, the effect of magma degassing 358 on oxygen isotopes should be negligible. In fact, the total devolatilization of MORB magmas 359 will change the δ^{18} O by less than 0.1 ‰ (Eiler, 2001). Similarly, oxygen fugacity changes in 360 the magmatic system should not create any oxygen isotope ratio variations as the fractionation 361 of oxygen isotopes in the melt is not controlled by fO_2 (Lester et al., 2013).

Assimilation of altered oceanic (AOC) crust with variable δ^{18} O (1-6‰ for lower AOC and 7-362 363 15‰ for upper AOC; e.g. Alt, 2003; Alt and Bach, 2006) may also affect the δ^{18} O variability 364 observed in melts (e.g., Genske et al., 2013). The assimilation process can be modeled by 365 simple mixing, because MORB melts and AOC have very similar oxygen concentrations. If we consider a typical mantle composition of $\delta^{18}O=5.5$ % (Eiler, 2001; Mattey et al., 1994) 366 367 and mix it with an altered upper oceanic crust with $\delta^{18}O=15$ ‰ (e.g., Alt, 2003), a minimum of 17% assimilation of upper AOC is required to explain the highest δ^{18} O measured in melt 368 inclusions (7.1‰) from ARP73-10-03. This amount of upper AOC assimilation (>17%) 369 370 would, for example, result in Ba content of >62 ppm, assuming an average Ba content of the

371 upper AOC of 300 ppm (e.g., Kelley et al., 2003) and 13.9 ppm for uncontaminated MORB 372 (Hofmann, 1988). As melt inclusions from ARP73-10-03 (N-MORB sample), have Ba 373 contents typical of N-MORB (around 20 ppm; maximum 34 ppm Ba), we find that 374 assimilation of upper crustal material is not likely to be responsible for the high δ^{18} O values 375 measured in these melt inclusions. Similarly, a minimum of 22% assimilation of the upper 376 AOC is required to obtain the lowest δ^{18} O (4.5%) of the melt inclusions, assuming an extreme lower AOC composition of $\delta^{18}O=1\%$ (as in hydrothermal amphibolite veins; e.g., 377 Alt and Bach, 2006). Assimilation of serpentinites that have δ^{18} O values of 2.0 to 6.7%, with 378 379 most of the values <5% (Barnes et al., 2014; Boschi et al., 2008) would require a larger 380 amount of assimilation compared to AOC, and could not account for the δ^{18} O up to 7.1 % or 381 down to 4.5 %.

The absence of correlation between elements that are similarly incompatible in the dissolution reactions of pyroxene, plagioclase and spinel (e.g., S and Ce; Danyushevsky et al., 2004) suggests that local grain-scale dissolution-reaction mixing processes (Danyushevsky et al., 2004) is an unlikely process to explain the δ^{18} O variations in melt inclusions. In fact, open system local dissolution of minerals formed in equilibrium with a basaltic melt will not change the δ^{18} O composition of the ascending basaltic melt.

388 The δ^{18} O variability of 1.1 and 1.4 ‰ in the olivine crystals (δ^{18} O_(Ol)) from ARP73-10-03 and 389 CH77-DR6-203 (Table DR3), respectively, is an indicator that these olivine crystals record 390 crystallization in a heterogeneous environment, or record a heterogeneous crystallization 391 history. Considering the typical reproducibility of olivine measurements of 0.4‰, these 392 variations are not likely to reflect analytical artefacts and should therefore represent real 393 heterogeneities. In fact, individual crystals are homogeneous within measurement uncertainty 394 of ca. 0.4 ‰, while different crystals from the same sample are more variable. In terms of 395 major elements, each crystal is homogeneous (with the exception of P heterogeneities in some

olivines; Manzini et al., 2017), but all olivines have different Fo contents (Table DR3). Within each sample, no clear trend is observed between Fo content and average $\delta^{18}O_{(OI)}$ (Fig. 5). At a given Fo content, $\delta^{18}O_{(OI)}$ varies up to 0.9‰. This observation suggests that these olivine crystals from both samples grew in slightly heterogeneous environments (~0.9-1‰ variation).

401

402 Olivine-melt inclusion isotopic disequilibrium

Equilibrium fractionation between olivine and melt ($\Delta^{18}O_{ol-melt}$) at 1250°C is -0.6 %, based on 403 Matthews et al. (1998), which is equivalent to twice the uncertainty of the measured 404 compositions. As mentioned above, Δ^{18} O variability in the studied melt inclusions is larger 405 406 than the heterogeneity of their host olivines. Only 42 and 46% of melt inclusions from 407 ARP 73-10-03 and CH77-203-DR6 are in equilibrium with their host (Figure 3). The remaining melt inclusions have either lower or higher Δ^{18} O_{ol-melt} compared to the theoretical 408 409 equilibrium value at 1250°C. While the proportion of melt inclusions in equilibrium with their 410 host olivine is similar for both samples, the proportion of melt inclusions with higher versus lower $\Delta^{18}O_{ol-melt}$ is different for each sample. In detail, within an olivine containing several 411 412 exposed melt inclusions, $\Delta^{18}O_{ol-melt}$ can vary significantly. No correlation is observed between 413 olivine-melt isotopic disequilibrium and Fo content of the host olivine, which indicates that 414 disequilibrium conditions could have prevailed at any time during the samples' magmatic 415 history. An important observation is that melt inclusions in equilibrium with their host olivines do not show significant δ^{18} O variation (5.7±0.3‰ for ARP73-10-03 and 6.3±0.5‰ 416 417 for CH77-203-DR6). Assuming equilibrium between olivine and melt, variations in temperature throughout crystal growth could affect $\Delta^{18}O_{ol-melt}$. However, the large variation of 418 Δ^{18} O_{ol-melt} would require a large and unrealistic temperature range, from 700°C for the lowest 419 Δ^{18} O_{ol-melt} to >2500 °C for the positive Δ^{18} O_{ol-melt}. 420

We propose that the measured $\Delta^{18}O_{ol-melt}$ variations instead indicate disequilibrium. Lower or 421 higher $\Delta^{18}O_{ol-melt}$ compared to theoretical equilibrium require either a process able to shift the 422 423 equilibrium toward both higher and lower values, or a combination of different processes. 424 Oxygen diffusion along the moving interface between melt and crystal during crystal growth 425 or after melt inclusion entrapment is a process capable of generating isotopic disequilibrium 426 between melt inclusions and their host olivine. Rapid crystal growth can lead to a local 427 disequilibrium in the growth media, creating boundary layers in which elements with the 428 lowest diffusivity in the melt and/or lowest partition coefficients (e.g., S, Cl, P, Al) will 429 concentrate. It has been demonstrated that this process can create melt inclusion compositions 430 with different major elements (Faure and Schiano, 2005; Watson and Müller, 2009), volatiles 431 (S, Cl) and/or P content (Baker, 2008) compared to the bulk melt composition. Oxygen is also a slow diffusing element in relatively dry melts, with ¹⁶O diffusing faster than ¹⁸O (e.g., 432 433 Lesher, 2010); therefore, such a boundary layer should have a higher δ^{18} O composition than 434 the melt in which the crystal in growing. Melt inclusions from ARP-73-10-03 have major, 435 trace and volatile elements compositions similar to MORB, with no enrichment in Al, S, Cl or 436 Ca; however, some melt inclusions within this sample do have higher P contents due to 437 olivine dissolution near P-rich (fast growing) domains (Manzini et al., 2017). No significant difference in $\delta^{18}O_{(OI)}$ between P-rich and P-poor domains within each olivine are observed 438 439 (Figure DR2). Moreover, $\delta^{18}O_{(OI)}$ compositions of the recrystallized olivine (P-depleted halo) 440 are similar to those of the primary olivine (Figure 6A). Figure 6B shows the δ^{18} O of melt 441 inclusions from both samples, as a function of their location (in primary olivine, or in 442 recrystallized domains), based on phosphorus maps of the host olivines. No systematic shift in δ^{18} O as a function of the olivine domain in which the melt inclusion is trapped can be 443 444 resolved. Also, isotopic disequilibrium can be found in melt inclusions trapped in olivine with or without P zoning (Table DR2), suggesting that the MI-Olivine δ^{18} O disequilibrium and 445

446 large δ^{18} O variations in melt inclusions are not related to initial rapid crystal growth creating 447 P-rich domains, nor to the dissolution of P-rich olivine domains.



449 **Figure 6**: A: Comparison of δ^{18} O measured in the ARP73-10-03 16 olivine (P-enriched and P-normal 450 zones), in the re-precipitated part (P-depleted zone around melt inclusion) and in the 2 melt inclusions 451 situated in the P-zoned olivine part. No differences in oxygen isotope can be seen between the 452 different parts of the olivine. B: Comparison of δ^{18} O measured in melt inclusions located in 453 recrystallized olivine (i.e. close to a P-zoned region of the olivine) and in unzoned olivine parts, 454 probably in initial olivine, that did not suffer from dissolution and re-precipitation. No differences can 455 be observed between the two populations, but in general CH77-DR6-203 has higher δ^{18} O values than 456 ARP73-10-03.

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As argued above, olivine-hosted melt inclusions within the samples studied here suffered from PEC of olivine. Manzini et al. (2017) have shown that this thin (< 1 μ m) layer crystallizes a few hours before eruption. This fast crystallization results in a strong P enrichment of the late-crystallized olivine. Post-entrapment crystallization could thus generate a thin boundary layer with a distinct (high) $\delta^{18}O_{(OI)}$ composition and lower the initial $\delta^{18}O$ of the melt inclusions. Given the small volume of the PEC (1 μ m layer compared to melt inclusion diameters of > 50 μ m), it should not largely influence the melt inclusion 465 composition. This assumption is verified by the absence of a correlation between amount of 466 PEC or P with δ^{18} O of the melt inclusions or Δ^{18} O _{ol-melt}.

467 Diffusion of O in and out of the melt inclusions could in theory produce higher or lower $\Delta^{18}O_{ol-melt}$. Oxygen can diffuse as mono or multi-species. Indeed, O is present in CO₂ and 468 469 H₂O molecules. It is known that water can diffuse in or out of melt inclusions (e.g., Chen et 470 al., 2013; Gaetani et al., 2012; Portnyagin et al., 2008), but mostly in the form of H⁺. Given 471 the low water content involved in MORB systems, this should not produce large δ^{18} O 472 variations in melt inclusions (Zhang and Ni, 2010). It is possible that CO₂ could largely degas 473 in melt inclusion shrinkage bubbles (e.g., Moore et al., 2015); migration of CO₂ in the bubble may fractionate δ^{18} O, but as ¹⁶O is diffusing faster this could only generate higher δ^{18} O. Melt 474 475 inclusions with the lowest $\Delta^{18}O_{ol-melt}$ do not necessarily contain shrinkage bubbles, which 476 indicates that fractionation of O isotopes is not related to shrinkage bubble formation.

477 Diffusion of O after melt inclusion entrapment would imply that melt inclusions were initially 478 not in equilibrium with their host at the time of entrapment and thus cannot explain the origin of the large variation of δ^{18} O in the studied melt inclusions. If O isotopic diffusion occurred 479 480 after melt inclusion entrapment, it should be limited based on the following observations: 481 First, based on P profiles made from melt inclusions in olivines, melt inclusions were trapped 482 for a few hours to a few days before quenching (Manzini et al., 2017). Using the diffusion data of Dohmen et al. (2002) (this dataset yields a diffusion coefficient of $\sim 9.3 \times 10^{12}$ in olivine 483 484 at 1250°C), this amount of time would result in a small amount of O diffusing from melt 485 inclusion to olivine over a profile no longer than 10 microns, which is not resolvable using the techniques employed in this study. This is confirmed by δ^{18} O profiles obtained from three 486 487 different olivines, in which melt inclusions are either in disequilibrium (Figures 7A and DR2) 488 or equilibrium (Figure 7B), starting at ~15-30µm from the inclusion wall, do not show any significant increase or decrease of δ^{18} O values toward the MI. Second, δ^{18} O profiles measured 489

490 within a single melt inclusion, from olivine wall to MI center, give similar values (Figure 8). 491 Lastly, there is no correlation between the melt inclusion size and the δ^{18} O value of the melt 492 inclusion.



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Figure 7: Oxygen isotopes profiles in olivines, starting from melt inclusion. A: oxygen isotopes profile in olivine ARP73-10-03_12, for which melt inclusion "a" is not in equilibrium. Profile starts from the melt inclusion. B: different δ^{18} O profiles done around the melt inclusion (in equilibrium) entrapped in olivine ARP-10-03_7. The 3 profiles start from 3 different locations around the melt inclusion and all are made in different directions. No clear diffusion profile from the melt inclusion toward the olivine can be observed. Black lines represent the average composition of olivine and dashed lines the 2 SD variations of the data along the profiles.

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Figure 8: Oxygen isotopes profiles in melt inclusion APR-73-10-03_16A, from wall to wall. Two
profiles were performed in this melt inclusion, perpendicular to one other.

506 In theory, oxygen diffusion could occur in response to a thermal gradient (Soret diffusion). 507 Indeed, Kyser (1998) and Bindeman et al. (2013) reported significant isotopic fractionation of 508 oxygen isotopes (up to 5.0‰) in anhydrous basaltic melt due to thermal diffusion, with migration of ¹⁸O towards the colder regions. However, such a process has never convincingly 509 510 been demonstrated in natural samples. Kyser (1998) pointed out that, even if thermally induced diffusion can lead to large δ^{18} O fractionation, the rapid decay of thermal gradient (< 511 512 10 days for 100° gradient at 1 cm from contact) makes it an unlikely process to happen in 513 natural samples. Given the short timescale associated with the olivine residence time and melt 514 inclusion formation in our samples (Manzini et al., 2017), a locally large thermal gradient (> 515 250°C) between migrating magma and conduit walls might be sustained before melt inclusion trapping. Assuming such a scenario, melt inclusions with δ^{18} O lower might form in olivine 516

517 located in the center of a conduit, whereas melt inclusions with higher δ^{18} O would form close 518 to the walls of the conduit; however, olivine crystallizing around the melt inclusions should 519 also have variable δ^{18} O, which is not observed. This observation indicates that O diffusion in 520 response to a thermal gradient is not responsible for the δ^{18} O variation in melt inclusions.

521

522 **Concluding remarks**

523 The large variations (up to 2.5‰) observed in oxygen isotope ratios within melt inclusions from 2 different Atlantic MORB samples (N-MORB and E-MORB) are not correlated with 524 525 major, trace or volatile elements. Oxygen isotope compositions of host olivines suggest a 526 small (<1‰) source heterogeneity for each sample. Less than half of the melt inclusions from 527 each sample are in isotopic equilibrium with their host olivines. The remaining melt 528 inclusions have either lower or higher isotopic fractionation compare to theoretical equilibrium values. Boundary layer entrapment will only shift δ^{18} O toward higher values, and 529 530 the present dataset does not support this hypothesis. Fractionation diffusion of O during CO₂ migration into the shrinkage bubble will shift δ^{18} O toward lower values, but no evidence of 531 this process has been found. Another process capable of generating δ^{18} O fractionation both 532 higher and lower than equilibrium values is O diffusion in due to the presence of a thermal 533 534 gradient before melt inclusion entrapment (i.e. Soret diffusion). Although this process could explain the large δ^{18} O variation measured, it would imply that a large, unrealistic, thermal 535 536 gradient (> 250°C) is sustained during the formation of melt inclusions. In addition, the olivine directly surrounding the melt inclusion should also have a different $\delta^{18}O$ in this 537 scenario, which we do not observe. 538

539 Despite the marked disequilibrium in a majority of melt inclusions, our data do not show any540 significant O diffusion in or out the melt inclusions. This is due to the short residence time at

high temperature after melt inclusion entrapment and the relatively large spatial resolution of the SIMS (10-15 microns). The exact process responsible for the melt inclusion-host $\delta^{18}O$ disequilibrium is thus not yet fully understood. Nonetheless, we argue that this dataset clearly shows that melt inclusions do not necessarily record the $\delta^{18}O$ composition of the magma in which their host grew. Caution should thus be taken before interpreting $\delta^{18}O$ variation as source process, especially when not related to any other tracer of common magmatic processes.

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560 **References**

- Alt, J.C., 2003. Stable isotopic composition of upper oceanic crust formed at a fast spreading ridge,
- 562 ODP Site 801. Geochemistry, Geophys. Geosystems 4, 1–11.
- 563 https://doi.org/10.1029/2002GC000400
- Alt, J.C., Bach, W., 2006. Oxygen isotope composition of a section of lower oceanic crust, ODP Hole
 735B. Geochemistry, Geophys. Geosystems 7. https://doi.org/10.1029/2006GC001385
- 566 Baker, D.R., 2008. The fidelity of melt inclusions as records of melt composition. Contrib. to Mineral.

567 Petrol. 156, 377–395. https://doi.org/10.1007/s00410-008-0291-3

- 568 Barnes, J.D., Beltrando, M., Lee, C.T.A., Cisneros, M., Loewy, S., Chin, E., 2014. Geochemistry of
- Alpine serpentinites from rifting to subduction: A view across paleogeographic domains and
 metamorphic grade. Chem. Geol. 389, 29–47. https://doi.org/10.1016/j.chemgeo.2014.09.012
- 571 Bindeman, I.N., Kamenetsky, V.S., Palandri, J., Vennemann, T., 2012. Hydrogen and oxygen isotope
- behaviors during variable degrees of upper mantle melting: Example from the basaltic glasses
- from Macquarie Island. Chem. Geol. 310–311, 126–136.
- 574 https://doi.org/10.1016/j.chemgeo.2012.03.031
- Bindeman, I.N., Lundstrom, C.C., Bopp, C., Huang, F., 2013. Stable isotope fractionation by thermal
 diffusion through partially molten wet and dry silicate rocks. Earth Planet. Sci. Lett. 365, 51–62.
 https://doi.org/10.1016/j.epsl.2012.12.037
- 578 Boschi, C., Dini, A., Früh-Green, G.L., Kelley, D.S., 2008. Isotopic and element exchange during
- 579 serpentinization and metasomatism at the Atlantis Massif (MAR 30°N): Insights from B and Sr
- isotope data. Geochim. Cosmochim. Acta 72, 1801–1823.
- 581 https://doi.org/10.1016/j.gca.2008.01.013
- 582 Bougault, H., Dmitriev, L., Schilling, J.G., Sobolev, A., Joron, J.L., Needham, H.D., 1988. Mantle
- 583 heterogeneity from trace elements: MAR triple junction near 14°N. Earth Planet. Sci. Lett. 88,

- 584 27–36. https://doi.org/10.1016/0012-821X(88)90043-X
- 585 Bucholz, C.E., Jagoutz, O., VanTongeren, J.A., Setera, J., Wang, Z., 2017. Oxygen isotope trajectories
- 586 of crystallizing melts: Insights from modeling and the plutonic record. Geochim. Cosmochim.
- 587 Acta 207, 154–184. https://doi.org/10.1016/j.gca.2017.03.027
- 588 Chen, Y., Provost, A., Schiano, P., Cluzel, N., 2013. Magma ascent rate and initial water
- 589 concentration inferred from diffusive water loss from olivine-hosted melt inclusions. Contrib. to
- 590 Mineral. Petrol. 165, 525–541. https://doi.org/10.1007/s00410-012-0821-x
- 591 Cottrell, E., Spiegelman, M., Langmuir, C.H., 2002. Consequences of diffusive reequilibration for the
 592 interpretation of melt inclusions. New York 3.
- 593 Danyushevsky, L. V., Della-Pasqua, F.N., Sokolov, S., 2000a. Re-equilibration of melt inclusions
- trapped by magnesian olivine phenocrysts from subduction-related magmas: Petrological

implications. Contrib. to Mineral. Petrol. 138, 68–83. https://doi.org/10.1007/PL00007664

- 596 Danyushevsky, L. V., Eggins, S.M., Falloon, T.J., Christie, D.M., 2000b. H₂O abundance in depleted
- 597 to moderately enriched mid-ocean ridge magmas; Part I: Incompatible behaviour, implications
- for mantle storage, and origin of regional variations. J. Petrol. 41, 1329–1364.
- 599 https://doi.org/10.1093/petrology/41.8.1329
- 600 Danyushevsky, L. V., Leslie, R.A.J., Crawford, A.J., Durance, P., 2004. Melt inclusions in primitive
- 601 olivine phenocrysts: The role of localized reaction processes in the origin of anomalous

602 compositions. J. Petrol. https://doi.org/10.1093/petrology/egh080

- Danyushevsky, L. V., Plechov, P., 2011. Petrolog3: Integrated software for modeling crystallization
- processes. Geochemistry, Geophys. Geosystems 12. https://doi.org/10.1029/2011GC003516
- Dohmen, R., Chakraborty, S., Becker, H.-W., 2002. Si and O diffusion in olivine and implications for
- 606 characterizing plastic flow in the mantle. Geophys. Res. Lett. 29, 2030.
- 607 https://doi.org/10.1029/2002GL015480
- Dosso, L., Bougault, H., 1986. A hot spot at 14 N on the Mid-Atlantic Ridge: Isotopic (Sr, Nd) and
- trace element data. Eos Trans. AGU 67, 410.

- 610 Eiler, J.M., 2001. Oxygen Isotope Variations of Basaltic Lavas and Upper Mantle Rocks. Rev.
- 611 Mineral. Geochemistry 43, 319–364. https://doi.org/10.2138/gsrmg.43.1.319
- 612 Eiler, J.M., Schiano, P., Kitchen, N., Stolper, E.M., 2000. Oxygen-isotope evidence for recycled crust
- 613 in the sources of mid-ocean-ridge basalts. Nature 403, 530–534.
- Faure, F., Schiano, P., 2005. Experimental investigation of equilibration conditions during forsterite
- growth and melt inclusion formation. Earth Planet. Sci. Lett. 236, 882–898.
- 616 https://doi.org/10.1016/j.epsl.2005.04.050
- 617 Frey, F.A., Walker, N., Stakes, D., Hart, S.R., Nielsen, R., 1993. Geochemical characteristics of
- basaltic glasses from theamar and famous axial valleys, Mid-Atlantic Ridge (36°–37°N):
- 619 Petrogenetic implications. Earth Planet. Sci. Lett. 115, 117–136. https://doi.org/10.1016/0012-
- 620 821X(93)90217-W
- 621 Gaetani, G. a., O'Leary, J. a., Shimizu, N., Bucholz, C.E., Newville, M., 2012. Rapid reequilibration
- of H₂O and oxygen fugacity in olivine-hosted melt inclusions. Geology 40, 915–918.
 https://doi.org/10.1130/G32992.1
- 624 Gale, A., Dalton, C.A., Langmuir, C.H., Su, Y., Schilling, J.-G., Dalton, C.A., Langmuir, C.H., Su, Y.,
- 625 Schilling, J., 2013. The mean composition of ocean ridge basalts. Geochem. Geophys. Geosyst
- 626 14, 489–518. https://doi.org/10.1029/2012GC004334
- 627 Garcia, M.O., Ito, E.M.I., Eiler, J.M., Pietruszka, A.J., 1998. Crustal Contamination of Kilauea
- 628 Volcano Magmas Revealed by Oxygen Isotope Analyses of Glass and Olivine from Puu Oo
 629 Eruption Lavas 39, 803–817.
- 630 Genske, F.S., Beier, C., Haase, K.M., Turner, S.P., Krumm, S., Brandl, P.A., 2013. Oxygen isotopes
- 631 in the Azores islands: Crustal assimilation recorded in olivine. Geology 41, 491–494.
- 632 https://doi.org/10.1130/G33911.1
- 633 Gurenko, A.A., Bindeman, I.N., Chaussidon, M., 2011. Oxygen isotope heterogeneity of the mantle
- beneath the Canary Islands: insights from olivine phenocrysts. Contrib. to Mineral. Petrol. 162,
- 635 349–363. https://doi.org/10.1007/s00410-010-0600-5

- 636 Gurenko, A.A., Chaussidon, M., 2002. Oxygen isotope variations in primitive tholeiites of Iceland :
- 637 evidence from a SIMS study of glass inclusions, olivine phenocrysts and pillow rim glasses. 638 Earth Planet. Sci. Lett. 205, 63-79.
- 639 Gurenko, A.A., Chaussidon, M., 1997. Boron concentrations and isotopic composition of the Icelandic 640 mantle: evidence from glass inclusions in olivine. Chem. Geol. 135, 21-34.
- 641 Gurenko, A.A., Chaussidon, M., 1995. Enriched and depleted primitive melts included in olivine from

642 Icelandic tholeiites: origin by continuous melting of a single mantle column. Geochim.

643 Cosmochim. Acta 59, 2905–2917. https://doi.org/10.1016/0016-7037(95)00184-0

- 644 Harmon, R.S., Hoefs, J., 1995. Oxygen isotope heterogeneity of the mantle deduced from global 180
- 645 systematics of basalts from different geotectonic settings. Contrib. to Mineral. Petrol. 95-114.
- 646 Hartley, M.E., Thordarson, T., Fitton, J.G., 2013a. Oxygen isotopes in melt inclusions and glasses

647 from the Askja volcanic system, North Iceland. Geochim. Cosmochim. Acta 123, 55–73. 648

- https://doi.org/10.1016/j.gca.2013.09.008
- 649 Hartley, M.E., Thordarson, T., Fitton, J.G., 2013b. Oxygen isotopes in melt inclusions and glasses
- 650 from the Askja volcanic system, North Iceland. Geochim. Cosmochim. Acta 123, 55-73.
- 651 https://doi.org/10.1016/j.gca.2013.09.008
- 652 Hartley, M.E., Thordarson, T., Taylor, C., Fitton, J.G., EIMF, 2012. Evaluation of the effects of
- 653 composition on instrumental mass fractionation during SIMS oxygen isotope analyses of glasses. 654 Chem. Geol. 334, 312–323. https://doi.org/10.1016/j.chemgeo.2012.10.027
- 655 Helo, C., Longpré, M.A., Shimizu, N., Clague, D.A., Stix, J., 2011. Explosive eruptions at mid-ocean
- 656 ridges driven by CO 2-rich magmas. Nat. Geosci. 4, 260–263. https://doi.org/10.1038/ngeo1104
- 657 Hofmann, A.W., 1988. Chemical differentiation of the Earth : the relationship between mantle,
- 658 continental crust, and oceanic crust 90, 297-314.
- 659 Jambon, A., Déruelle, B., Dreibus, G., Pineau, F., 1995. Chlorine and bromine abundance in MORB:
- 660 the contrasting behaviour of the Mid-Atlantic Ridge and East Pacific Rise and implications for
- 661 chlorine geodynamic cycle. Chem. Geol. 126, 101-117. https://doi.org/10.1016/0009-

662 2541(95)00112-4

Javoy, M., Pineau, F., 1991. The volatiles record of a "popping" rock from the Mid-Atlantic Ridge at
14°N: chemical and isotopic composition of gas trapped in the vesicles. Earth Planet. Sci. Lett.

665 107, 598–611. https://doi.org/10.1016/0012-821X(91)90104-P

- Jin, Z.-M., Green, H.W., Zhou, Y., 1994. Melt topology in partially molten mantle peridotite during
 ductile deformation. Nature 372, 164.
- 668 Jochum, K.P., Stoll, B., Herwig, K., Willbold, M., Hofmann, A.W., Amini, M., Aarburg, S.,
- Abouchami, W., Hellebrand, E., Mocek, B., Raczek, I., Stracke, A., Alard, O., Bouman, C.,
- 670 Becker, S., Dücking, M., Brätz, H., Klemd, R., de Bruin, D., Canil, D., Cornell, D., de Hoog, C.-
- J., Dalpé, C., Danyushevsky, L., Eisenhauer, A., Gao, Y., Snow, J.E., Groschopf, N., Günther,
- D., Latkoczy, C., Guillong, M., Hauri, E.H., Höfer, H.E., Lahaye, Y., Horz, K., Jacob, D.E.,
- 673 Kasemann, S.A., Kent, A.J.R., Ludwig, T., Zack, T., Mason, P.R.D., Meixner, A., Rosner, M.,
- 674 Misawa, K., Nash, B.P., Pfänder, J., Premo, W.R., Sun, W.D., Tiepolo, M., Vannucci, R.,
- 675 Vennemann, T., Wayne, D., Woodhead, J.D., 2006. MPI-DING reference glasses for in situ
- 676 microanalysis: New reference values for element concentrations and isotope ratios.
- 677 Geochemistry, Geophys. Geosystems 7, n/a-n/a. https://doi.org/10.1029/2005GC001060
- 678 Kamenetsky, V.S., Gurenko, A.A., 2007. Cryptic crustal contamination of MORB primitive melts
- 679 recorded in olivine-hosted glass and mineral inclusions. Contrib. to Mineral. Petrol. 153, 465–

680 481. https://doi.org/10.1007/s00410-006-0160-x

- Kelley, K. a., Plank, T., Ludden, J., Staudigel, H., 2003. Composition of altered oceanic crust at ODP
 Sites 801 and 1149. Geochemistry, Geophys. Geosystems 4, n/a-n/a.
- 683 https://doi.org/10.1029/2002GC000435
- Kempton, P.D., Hawkesworth, C.J., Fowler, M., 1991. Geochemistry and isotopic composition of
- gabbros from layer 3 of the Indian Ocean crust, Hole 735B. Proc., Sci. results, ODP, Leg 118,
- 686 Fract. Zo. Drill. Southwest Indian Ridge 127–143.
- 687 https://doi.org/10.2973/odp.proc.sr.118.118.1991

- 688 Kent, A.J.R., 2008. Melt Inclusions in Basaltic and Related Volcanic Rocks. Rev. Mineral.
- 689 Geochemistry 69, 273-331. https://doi.org/10.2138/rmg.2008.69.8
- 690 Kita, N.T., Ushikubo, T., Fu, B., Valley, J.W., 2009. High precision SIMS oxygen isotope analysis
- 691 and the effect of sample topography. Chem. Geol. 264, 43–57.
- 692 https://doi.org/10.1016/j.chemgeo.2009.02.012
- 693 Kyser, T.K., Lesher, C.E., Walker, D., 1998. The effects of liquid immiscibility and thermal diffusion
- 694 on oxygen isotopes in silicate liquids. Contrib. to Mineral. Petrol. 133, 373-381.
- 695 https://doi.org/10.1007/s004100050459
- 696 Laubier, M., Gale, A., Langmuir, C.H., 2012. Melting and crustal processes at the FAMOUS segment
- 697 (mid-atlantic ridge): New insights from olivine-hosted melt inclusions from multiple samples. J.
- 698 Petrol. 53, 665–698. https://doi.org/10.1093/petrology/egr075
- 699 Laubier, M., Schiano, P., Doucelance, R., Ottolini, L., Laporte, D., 2007. Olivine-hosted melt
- 700 inclusions and melting processes beneath the FAMOUS zone (Mid-Atlantic Ridge). Chem. Geol.
- 701 240, 129–150. https://doi.org/10.1016/j.chemgeo.2007.02.002
- 702 Lecuyer, C., Reynard, B., 1996. High-temperature alteration of oceanic gabbros by seawater (Hess
- 703 Deep, Ocean Drilling Program Leg 147): Evidence from oxygen isotopes and elemental fluxes. J. 704
- Geophys. Res. 101, 15883-15897.
- 705 Lesher, C.E., 2010. Self-diffusion in Silicate Melts: Theory, Observations and Applications to
- 706 Magmatic Systems. Rev. Mineral. Geochemistry 72, 269-309.
- 707 https://doi.org/10.2138/rmg.2010.72.7
- 708 Lester, G.W., Kyser, T.K., Clark, A.H., 2013. Oxygen isotope partitioning between immiscible silicate
- 709 melts with H₂O, P and S. Geochim. Cosmochim. Acta 109, 306-311.
- 710 https://doi.org/10.1016/j.gca.2013.01.037
- 711 Magenheim, A.J., Spivack, A.J., Michael, P.J., Gieskes, J.M., 1995. Chlorine stable isotope
- 712 composition of the oceanic crust: Implications for Earth's distribution of chlorine. Earth Planet.
- 713 Sci. Lett. 131, 427-432. https://doi.org/10.1016/0012-821X(95)00017-7

- 714 Manzini, M., Bouvier, A.S., Baumgartner, L.P., Müntener, O., Rose-Koga, E.F., Schiano, P., Escrig,
- S., Meibom, A., Shimizu, N., 2017. Weekly to monthly time scale of melt inclusion entrapment
- prior to eruption recorded by phosphorus distribution in olivine from mid-ocean ridges. Geology
- 717 45, 1059–1062. https://doi.org/10.1130/G39463.1
- Mattey, D., Lowry, D., Macpherson, C., 1994. Oxygen isotope compositions of mantle peridotite.
 Earth Planet. Sci. Lett. 128, 231–241.
- Matthews, A., Stolper, E.M., Eiler, Epstein, S., 1998. Oxygen isotope fractionation among melts,
 minerals and rocks. Goldschmidt Conf. Toulouse 971–972.
- 722 McDonough, W.F., Sun, S. -s., 1995. The composition of the Earth. Chem. Geol. 120, 223–253.

723 https://doi.org/10.1016/0009-2541(94)00140-4

- McKenzie, D.P., 1984. The generation and compaction of partial melts. J. Petrol. 25, 713–765.
 https://doi.org/https://doi.org/10.1093/petrology/25.3.713
- 726 Moore, L.R., Gazel, E., Tuohy, R., Lloyd, A.S., Esposito, R., Steele-MacInnis, M., Hauri, E.H.,
- Wallace, P.J., Plank, T., Bodnar, R.J., 2015. Bubbles matter: An assessment of the contribution
- of vapor bubbles to melt inclusion volatile budgets. Am. Mineral. 100, 806–823.
- 729 Naumov, V.B., Dorofeeva, V.A., Girnis, A. V., Yarmolyuk, V. V., 2014. Comparison of major,
- volatile, and trace element contents in the melts of mid-ocean ridges on the basis of data on
- inclusions in minerals and quenched glasses of rocks. Geochemistry Int. 52, 347–364.
- 732 https://doi.org/10.1134/S0016702914050073
- Nielsen, R.L., Sours-Page, R.E., Harpp, K.S., 2000. Role of a Cl-bearing flux in the origin of depleted
 ocean floor magmas. Geochemistry, Geophys. Geosystems 1.
- 735 https://doi.org/10.1029/1999GC000017
- 736 Peres, P., Kita, N.T., Valley, J.W., Fernandes, F., Schuhmacher, M., 2013. New sample holder
- 737 geometry for high precision isotope analyses. Surf. Interface Anal. 45, 553–556.
- 738 https://doi.org/10.1002/sia.5061
- 739 Portnyagin, M., Almeev, R., Matveev, S., Holtz, F., 2008. Experimental evidence for rapid water

- exchange between melt inclusions in olivine and host magma. Earth Planet. Sci. Lett. 272, 541–
- 741 552. https://doi.org/10.1016/j.epsl.2008.05.020
- 742 Schiano, P., 2003. Primitive mantle magmas recorded as silicate melt inclusions in igneous minerals.

743 Earth-Science Rev. 63, 121–144. https://doi.org/10.1016/S0012-8252(03)00034-5

- 744 Schilling, J.G., Zajac, M., Evans, R., Johnston, T., White, W., Devine, J.D., Kingsley, R., 1983.
- Petrologic and geochemical variations along the Mid-Atlantic Ridge from 29 degrees N to 73

746 degrees N. Am. J. Sci. 283, 510–586. https://doi.org/10.2475/ajs.283.6.510

- 747 Seitz, S., Baumgartner, L.P., Bouvier, A.S., Putlitz, B., Vennemann, T., 2017. Quartz Reference
- 748 Materials for Oxygen Isotope Analysis by SIMS. Geostand. Geoanalytical Res. 41, 69–75.
- 749 https://doi.org/10.1111/ggr.12133
- 750 Shimizu, N., 1998. The geochemistry of olivine-hosted melt inclusions in a FAMOUS basalt ALV
- 751 519-4-1. Phys. Earth Planet. Inter. 107, 183–201. https://doi.org/10.1016/S0031-9201(97)00133752 7
- Sobolev, A. V., Shimizu, N., 1993. Ultra-depleted primary melt in an olivine from the Mid-Atlantic
 Ridge. Nature 363, 151–154.
- 755 Stakes, D.S., 1991. Oxygen and hydrogen isotope compositions of oceanic plutonic rocks: High-
- temperature deformation and metamorphism of oceanic layer 3. Stable Isot. Geochemistry A
 Tribut. to Samuel Epstein 77–90.
- Tormey, D.R., Grove, T.L., Bryan, W.B., 1987. Experimental petrology of N-MORB near the Kane
 Fracture Zone: 22-25°N, mid-Atlantic ridge. Contrib. Miner. Pet. 96, 121–139.
- 760 Van Orman, J.A., Grove, T.L., Shimizu, N., 2002. Diffusive fractionation of trace elements during
- production and transport of melt in Earth's upper mantle. Earth Planet. Sci. Lett. 198, 93–112.
 https://doi.org/10.1016/S0012-821X(02)00492-2
- 763 Wallace, P., Carmichael, I.S.E., 1992. Sulfur in basaltic magmas. Geochim. Cosmochim. Acta 56,
- 764 1863–1874. https://doi.org/10.1016/0016-7037(92)90316-B

765	Watson, E.B., Müller, T., 2009. Non-equilibrium isotopic and elemental fractionation during	
766	diffusion-controlled crystal growth under static and dynamic conditions. Chem. Geol. 267, 111-	
767	124. https://doi.org/10.1016/j.chemgeo.2008.10.036	
768	Zhang, Y., Ni, H., 2010. Diffusion of H, C, and O Components in Silicate Melts, in: Reviews in	
769	Mineralogy and Geochemistry. Mineralogical Society of America, pp. 171-225.	
770	https://doi.org/10.2138/rmg.2010.72.5	
771	Zindler, A., Hart, S.R., Frey, F.A., Jakobsson, S.P., 1979. Nd and Sr isotope ratios and rare earth	
772	element abundances in Reykjanes Peninsula basalts evidence for mantle heterogeneity beneath	

773 Iceland. Earth Planet. Sci. Lett. 45, 249–262. https://doi.org/10.1016/0012-821X(79)90127-4

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777 Supplementary material:

- **Figure DR1**: Calibration line for IMF correction for oxygen isotope analyses in glasses and
- olivine.
- **Figure DR2**: P-elemental X-ray map of the ARP73-10-03_16 olivine
- 781 Table DR1: SIMS conditions
- **Table DR2:** Major, trace, volatile, elements and δ^{18} O composition of melt inclusions
- 783 **Table DR3:** Oxygen isotopes in olivines

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