Formation of Amphibole-Bearing Peridotite and Amphibole-Bearing Pyroxenite through Hydrous Melt-Peridotite Reaction and In Situ Crystallization: An Experimental Study

Chunguang Wang¹, Yan Liang², and Wenliang Xu^1

¹Jilin University ²Brown University

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Abstract

Amphibole is a common hydrous mineral in mantle rocks. To better understand the processes leading to the formation of amphibole-bearing peridotites and pyroxenites in mantle rocks, we have undertaken an experimental study reacting lherzolite with hydrous basaltic melts in Au-Pd capsules using the reaction couple method. Two melts were examined, a basaltic andesite and a basalt, each containing 4 wt% of water. The experiments were run at 1200°C and 1 GPa for 3 or 12 h, and then cooled to 880°C and 0.8 GPa over 49 h. The reaction at 1200°C and 1 GPa produced a melt-bearing orthopyroxenite-dunite sequence. The cooling stimulates crystallization of orthopyroxene, clinopyroxene, amphibole, and plagioclase, leading to the formation of an amphibole-bearing gabbronorite–orthopyroxenite–peridotite sequence. Compositional variations of minerals in the experiments are controlled by temperature, pressure, and reacting melt composition. Texture, mineralogy, and mineral compositional variation trends obtained from the experiments are similar to those from mantle xenoliths and peridotite massif from the field including amphibole-bearing peridotites and amphibole-bearing pyroxenite and amphibolite that are spatially associated with peridotites, underscoring the importance of hydrous melt-peridotite reaction in the formation of these amphibolebearing rocks in the upper mantle. Amphiboles in some field samples have distinct textual and mineralogical features and their compositional variation trends are different from that defined by the melt-peridotite reaction experiments. These amphiboles are either crystallized from the host magma that entrained the xenoliths or product of hydrothermal alterations at shallow depths.

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4	Chunguang Wang ^{1, 2, *} , Yan Liang ² , and Wenliang Xu ¹
5	¹ College of Earth Sciences, Jilin University, Changchun 130061, China
6 7	² Department of Earth, Environmental and Planetary Sciences, Brown University, Providence, RI 02912, USA
8	Corresponding author: Chunguang Wang (c_wang@jlu.edu.cn)
9	Key Points:
10 11	• Consequence of hydrous melt-peridotite reaction followed by in-situ crystallization was investigated experimentally.
12 13	• Hydrous melt-lherzolite reactive crystallization forms amphibole-bearing peridotite and amphibole-bearing pyroxenite.
14 15	• Chemical composition of amphibole can be used to identify hydrous melt and peridotite interaction in the upper mantle.

16 Abstract

Amphibole is a common hydrous mineral in mantle rocks. To better understand the 17 processes leading to the formation of amphibole-bearing peridotites and pyroxenites in mantle 18 rocks, we have undertaken an experimental study reacting lherzolite with hydrous basaltic melts 19 in Au-Pd capsules using the reaction couple method. Two melts were examined, a basaltic 20 21 andesite and a basalt, each containing 4 wt% of water. The experiments were run at 1200°C and 1 GPa for 3 or 12 h, and then cooled to 880°C and 0.8 GPa over 49 h. The reaction at 1200°C 22 and 1 GPa produced a melt-bearing orthopyroxenite-dunite sequence. The cooling stimulates 23 crystallization of orthopyroxene, clinopyroxene, amphibole, and plagioclase, leading to the 24 formation of an amphibole-bearing gabbronorite-orthopyroxenite-peridotite sequence. 25 Compositional variations of minerals in the experiments are controlled by temperature, pressure, 26 27 and reacting melt composition. Texture, mineralogy, and mineral compositional variation trends obtained from the experiments are similar to those from mantle xenoliths and peridotite massif 28 from the field including amphibole-bearing peridotites and amphibole-bearing pyroxenite and 29 amphibolite that are spatially associated with peridotites, underscoring the importance of hydrous 30 melt-peridotite reaction in the formation of these amphibole-bearing rocks in the upper mantle. 31 Amphiboles in some field samples have distinct textual and mineralogical features and their 32 compositional variation trends are different from that defined by the melt-peridotite reaction 33 34 experiments. These amphiboles are either crystallized from the host magma that entrained the

35 xenoliths or product of hydrothermal alterations at shallow depths.

36 **1. Introduction**

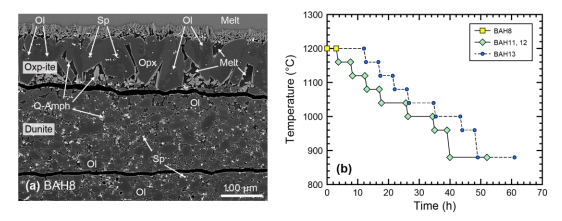
Amphibole is commonly observed in mantle xenoliths (e.g., Wilshier and Schwarzman, 37 1971; Dautria et al., 1987; Moine et al., 2001; Xu et al., 2010) and massif peridotites (e.g., 38 Agrinier et al., 1993; Zanetti et al., 1996; Ohara and Ishii, 1998; Li et al., 2018) from a range of 39 geological settings, such as arcs (e.g., Ohara and Ishii, 1998; Ishimaru et al., 2007; Bénard and 40 Ionov, 2013), ocean islands (e.g., Clague and Bohrson, 1991; Moine et al., 2001), continental 41 margins (Francis, 1976a, 1976b; Agrinier et al., 1993), and intracontinental regions (e.g., 42 Wilshier and Schwarzman, 1971; Dautria et al., 1987; O'Reilly et al., 1991; Xu et al., 2010). 43 Typical lithologies include amphibole-bearing peridotite, amphibole-bearing pyroxenite, and 44 amphibolite. In general, amphibole occurs either interstitial to anhydrous minerals in peridotites 45 or in amphibole-rich veins or bodies in peridotites. 46

Based on field studies, a number of mechanisms have been proposed for the formation of 47 amphibole in mantle xenoliths and massif peridotites. Frey and Prinz (1978) reported a set of 48 amphibole-bearing peridotite and pyroxenite xenoliths in basanites from Grand Canyon and San 49 Carlos, Arizona, USA. They suggested that the poikilitic amphiboles enclosing olivines and 50 spinels were accumulations of basaltic magma. Agrinier et al. (1993) described different types of 51 amphibole in a peridotite massif from Zabargad Island, Red Sea. Amphiboles that exist as 52 individual grains interstitial between olivines and spinels were inferred to be products of reaction 53 between the peridotites and hydrous fluids, and amphiboles that occur in shear zones and 54 mylonitic bands to be results of seawater alteration after emplacement of the massif. Coltorti et 55 al. (2004) and Bonadiman et al. (2014) inferred that the disseminated amphiboles and vein 56 57 amphiboles in peridotite entrained by basalts from Victoria Land, Antarctica were formed by reaction between clinopyroxene and a batch of metasomatic melt. The different occurrences are 58 due to different melt-to-rock ratios. The chief objective of the present study is to develop a set of 59

criteria that can be used to distinguish amphiboles formed by hydrous melt and peridotite
 reaction from those produced by fluid-peridotite interaction or hydrothermal alteration.

The processes and consequences of melt-peridotite interaction have been extensively 62 studied through laboratory dissolution experiments for a range of starting compositions and P-T 63 conditions in anhydrous systems (e.g., Daines and Kohlstedt, 1994; Yaxley and Green, 1998; 64 Takahashi and Nakajima, 2002; Morgan and Liang, 2003, 2005; Beck et al., 2006; Lambart et 65 al., 2009, 2012; Van den Bleeken et al., 2010, 2011; Wang et al., 2013). However, comparable 66 studies in hydrous systems are limited (Sekine and Wyllie, 1983; Carroll and Wyllie, 1989; 67 Johnston and Wyllie, 1990; Kelemen et al., 1990; Sen and Dunn, 1994; Rapp et al., 1999; Mallik 68 et al., 2015; Mandler and Grove, 2016; Wang et al., 2016). Sen and Dunn (1994) conducted 69 reaction experiments by packing a powdered spinel lherzolite layer against a powdered 70 71 amphibolite (1.5 wt% water) layer in graphite-lined Pt capsules at 1.5-2.0 GPa and 950-1025°C. They observed amphiboles along the boundaries of olivine, pyroxenes, and spinel. The 72 amphiboles have compositions similar to those in amphibole-bearing peridotite from the field. 73 They inferred that reaction of olivine, primary orthopyroxene, clinopyroxene, and spinel with 74 hydrous melt forms amphibole and Fe-rich orthopyroxene and that a further reaction would form 75 amphibole-bearing harzburgite (free of clinopyroxene and spinel) and finally amphibole-bearing 76 orthopyroxenite (free of clinopyroxene, spinel, and olivine). In a recent melt-peridotite reaction 77 78 study (Wang et al., 2016), we reacted a layer of lherzolite with a layer of water-added (4 wt%) basaltic andesite in a Au-Pd lined graphite capsule at 1200°C and 1 GPa (run BAH8 in Wang et 79 al., 2016). Figure 1a is a back-scattered electron (BSE) image showing the orthopyroxenite-80 dunite sequence formed in the peridotite half of the reaction couple and quenched amphibole 81 crystals in the interstitial melts around the orthopyroxenite-dunite boundary. The presence of 82 quenched amphiboles in this isothermal hydrous melt-peridotite reaction experiment is 83 intriguing. The reacting melt around the orthopyroxenite must be near amphibole saturation at 84

85 1200°C and 1 GPa.



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Figure 1. (a) Back-scattered electron (BSE) image that recalls the result (interface region) of hydrous basaltic
 andesite and peridotite reaction experiments conducted at 1200 °C and 1 GPa (BAH8, Wang et al., 2016) using the

same capsule setup as those in the present study. Ol = olivine; Opx = orthopyroxene; Sp = spinel; Q-Amph = quench

amphibole; Opx-ite = orthopyroxenite. (b) Temperature histories of the melt-rock reaction experiments. Zero time

marks the beginning of the experiment, when the target temperature of 1200°C was reached. Experiments BAH11

and MBH12 were run using the same temperature-pressure path. BAH13 was hold at 1200°C for a longer duration

93 (12 h) than BAH11 and MBH12 (3 h). Temperature and duration used in the isothermal experiment BAH8 (1200°C,

3 h, without step cooling, Wang et al., 2016) is also shown for comparison.

95 In this study, we further expand the isothermal hydrous melt-peridotite reaction

96 experiments of Wang et al. (2016) by considering two hydrous melts (basalt and basaltic

andesite) and by allowing in situ crystallization of reacting melts through step cooling of

experimental charges (to 880°C and 0.8 GPa) at the end of isothermal reaction at 1200°C and 1
 GPa (Fig. 1b). The quenched experimental charge enables us to examine the texture, mineralogy,

and mineral compositional variation formed after the hydrous melt and peridotite reaction.

101 Results from these experiments allow us to better understand the processes leading to the

102 formation of amphibole-bearing peridotites and pyroxenites.

103 The remaining part of this paper is organized as follows. In the next section, we describe 104 the starting materials and experimental methods. We present the textural and chemical features 105 of the run products, compare the results with those observed in previous experimental studies, 106 and discuss the processes leading to the formation of amphibole-bearing peridotites and 107 pyroxenites. We then compare texture and mineral compositions observed in the experiments 108 with those from amphibole-bearing peridotites and pyroxenites in mantle xenoliths and massif 109 peridotites. We show that water and melt-peridotite reaction are important in determining the

110 lithology and mineral compositional variation in the mantle rocks.

Oxide (wt%)	Basaltic andesite	MORB	Spinel lherzolite			
	MONT147	D44A	WFY-2 ^b			
			Olivine	Opx	Срх	Spinel
SiO ₂	54.86	50.13	40.25	54.39	51.45	0.08
TiO ₂	0.77	1.25	0.01	0.09	0.39	0.15
Al_2O_3	18.78	16.31	0.04	4.44	5.49	54.26
Cr_2O_3			0.03	0.39	0.81	12.83
FeO ^a	8.96	8.85	9.82	6.25	2.93	10.87
MnO	0.21	0.17	0.12	0.12	0.11	0.15
MgO	3.66	8.72	48.41	32.46	16.27	20.50
CaO	8.80	12.50	0.09	0.88	20.55	0.01
Na ₂ O	3.08	2.63	0.00	0.12	0.98	0.02
K ₂ O	0.78	0.02	0.00	0.01	0.00	0.01
NiO			0.38	0.11	0.04	0.35
P_2O_5	0.14	0.08				
Total	100.04	100.66	99.15	99.26	99.02	99.23
Mg#	42.13	63.72	89.78	90.25	90.82	77.07
Cr#						13.69

 Table 1 Starting compositions

 $Mg\# = 100 \times Mg/(Mg + Fe)$, atomic ratio; $Cr\# = 100 \times Cr/(Cr + Al)$, atomic ratio; Opx = orthopyroxene; Cpx = clinopyroxene.

^a Total Fe as FeO

^b WFY-2 consists of 60 % olivine, 23 % opx, 12 % cpx, and 5 % spinel

111 **2. Methods**

112 Hydrous melt-peridotite reaction and in situ crystallization experiments were conducted 113 using the reaction couple method and a 19.1-mm piston-cylinder apparatus at Brown University. 114 Reaction couples were made by juxtaposing a powdered starting basalt layer against a powdered

spinel lherzolite layer at a length ratio of $\sim 5:3$ in Au-Pd lined graphite capsules. Each experiment

116 consists of four stages: pressurization to 1 GPa at room temperature, a dwell at 1200°C and 1

GPa, a subsequent step cooling to 880°C and 0.8 GPa, and a final dwell at 880°C and 0.8 GPa. The starting melt and mineral compositions are listed in Table 1. Run conditions for each

The starting melt and mineral compositions are listed in Table 1. Run conditions for each experiment are listed in Table 2, and the temperature-time paths are shown in Fig. 1b.

- 120 2.1. Starting compositions

The starting lherzolite (sample WFY-2) was obtained by mixing optically clean olivine 121 (60%), orthopyroxene (23%), clinopyroxene (12%), and spinel (5%) from a disaggregated fertile 122 spinel lherzolite xenolith entrained by alkali basalt from Huinan, China. The starting materials 123 for the reacting melts are a basaltic andesite (sample Mont147) from Montserrat and a mid-ocean 124 ridge basalt (MORB, sample D44A) from Southern East Pacific Rise to which we added 4 wt% 125 126 water during sample loading. The starting lherzolite and basaltic andesite were used in isothermal hydrous melt-rock reaction experiments of Wang et al. (2016), and the MORB was 127 used in anhydrous reaction and crystallization experiments of Tursack and Liang (2012) and 128 129 Saper and Liang (2014). Results from these earlier laboratory studies serve as references or benchmarks for the hydrous reaction and crystallization experiments reported in this study. The 130 starting samples were ground separately in ethanol using an agate mortar and pestle for 2-4 h and 131

132 stored at 110°C before use.

133 2.2. Experimental procedures

The furnace assembly consists of a Au-Pd lined graphite capsule in a MgO sleeve 134 sandwiched between two crushable MgO spacers in Pyrex tubing, NaCl sleeve, and straight-135 walled graphite heater. The cylindrical graphite capsule (6.5 mm OD, 7 mm long) was lined with 136 a Au₇₅-Pd₂₅ inner capsule (3 mm OD, 2.8 mm ID, 5 mm long). To make a reaction couple, we 137 138 first packed the peridotite powder in the lower part of the Au-Pd capsule and filled the remaining part of the capsule with the basaltic andesite or MORB powder. We then added deionized water 139 using a micro syringe and sealed the Au-Pd capsule using a micro arc welder immediately after 140 water loading. We weighed the capsule before and after each loading to ensure that the water-to-141 basalt weight ratio is around 4:96 (i.e., 4 wt% water). The arc welder has a thin tungsten 142 electrode (0.6 mm in diameter) and operates under a stream of protective argon gas. It allows 143 pinpoint welding with a very small heat affected area. The weight loss due to the welding is less 144 than 0.5% (evaporation of water and Au-Pd alloy). Finally, we loaded the sealed Au-Pd capsule 145 in the graphite capsule and capped the graphite capsule with a 0.5 mm thick graphite lid. The 146 furnace assembly and capsule used in this study are the same as those used in experiments BAH8 147 and BADH15 in Wang et al. (2016). Oxygen fugacity of the experiments is maintained at -8 to -9 148 logarithmic unit, estimated using the method of Barr and Grove (2010). 149

To conduct a reaction experiment, the charge was cold pressurized to 1 GPa, followed by heating to 1200°C at 75°C/min while maintaining pressure. The run was held at 1200°C and 1 GPa for 3 h for runs BAH11 and MBH12 and 12 h for BAH13. The run was then cooled to 880°C over 7 steps in 49 h (Fig. 1b). Each step cooling was performed at a rate of 1°C/min, followed by a 4 or 8 h dwell at the specific temperature. The pressure, which was not adjusted during cooling, gradually decreased to 0.8 GPa at the final dwell (880°C). At the end of experiment, the charge was quenched by cutting the power supply while maintaining the

- 157 pressure. The heating and step cooling were operated through a Eurotherm controller, and
- temperature was measured using a $W_{97}Re_3$ - $W_{75}Re_{25}$ thermocouple with an uncertainty of 10°C
- 159 (Morgan and Liang, 2005). No friction correction was applied to the pressure. The retrieved
- 160 experimental charge was checked under a binocular microscope to ensure no crack in the Au-Pd
- capsule. The capsule was then mounted in epoxy and polished gradually using a 600-mesh SiC
 polishing paper in water until the cylindrical axis was exposed. During polish, the Au-Pd capsule
- 162 pointing paper in water until the cymunical axis was exposed. During pointin, the Au-1 d capsul 163 was checked using a microscope under reflecting light to ensure no leak exist; the exposed
- 164 surface was soaked in epoxy and stored in a vacuum chamber for several times, so that cracks
- formed during quench were puttied. The exposed surface was finally polished to 1 μ m finish for
- 166 electron microprobe analysis.
- 167 2.3. Microprobe analysis

Collection of back-scattered electron images and X-ray concentration maps and
 measurements of mineral compositions were carried out using a Cameca SX-100 electron
 microprobe at Brown University. An accelerating voltage of 20 kV and a beam current of 25 nA
 were used to collect X-ray concentration maps. A 10 µm diameter beam was used for glass

analyses and a focused beam for mineral analyses. Accelerating voltage and beam current were

- 173 15 kV and 10 nA for glass analyses, 20 kV and 15 nA for plagioclase analyses, and 20 kV and 25
- nA for olivine, pyroxene, amphibole, and spinel analyses, respectively. Sodium was analyzed
- 175 first with counting times of 10 s for peak and 5 s for background to minimize volatilization.

176 Counting times for other elements were 20-30 s for peak, and 10-15 s for background on each

side of the peak. Natural standards were used for calibration.

Run#	BAH8 ^a	BAH11	BAH13	MBH12
Starting melt Basaltic andesite Bas		Basaltic andesite	Basaltic andesite	MORB
Pressure (GPa)	1	1 (0.8) ^b	1 (0.8) ^a	1 (0.8) ^a
Initial temp (°C)	1200 (3 h) ^c	1200 (3 h) ^c	1200 (12 h) ^c	1200 (3 h) ^c
Final temp (°C)		880 (12 h) ^c	880 (12 h) ^c	880 (12 h) ^c
Cooling process	Quench	Step cooling ^d	Step cooling ^d	Step cooling ^d
Total duration (h)	3	52	61	52
RBL lithologies	Glass	Amph gabbronorite	Amph gabbronorite	Amph gabbronorite
	Opx-ite	Amph-bearing opx-ite	Amph-bearing opx-ite	Amph-bearing opx-ite
	Dunite	Amph-bearing harz	Amph-bearing harz	Amph-bearing lherz

 Table 2 Summary of experimental conditions

RBL = reactive boundary layer; Amph = amphibole; Opx-ite = orthopyroxenite; Harz = harzburgite; Lherz = lherzolite.

^a Isobaric and isothermal experiment from Wang et al. (2016)

^b Number in parenthesis is the pressure at final temperature

^c Number in parenthesis is the duration at the initial or final temperature

^d Step cooling is at a rate of 1°C/min to specific temperatures maintained for 4 or 8 h

178 **3. Results and discussion**

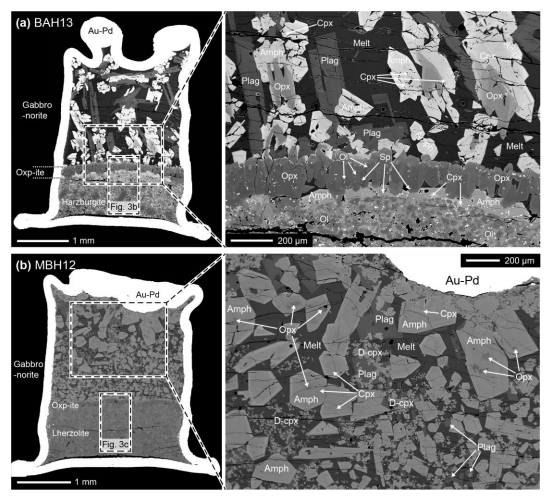
179 Three hydrous melt-peridotite reaction and crystallization experiments were conducted

using the two starting melts. They were built on melt-peridotite reaction experiments using

similar setups and starting compositions that are either hydrous but isothermal (Wang et al.,

182 2016) or step-cooling but anhydrous (Tursack and Liang, 2012; Saper and Liang, 2014).

- 183 Experiments BAH11 and BAH13 use the 4 wt% water-added basaltic andesite as the starting
- melt, and experiment MBH12 uses the 4 wt% water-added MORB as the starting melt. We did
- not measure water contents in the experimental glasses because of the crystalline nature of the
- experimental charges. Judging from the presence of amphibole and vesicles in the experiments and integrity of the Au-Pd capsules (Fig. 2 and supporting Figure S1), we conclude that these
- capsules retained a significant fraction of water in the system and our reacting melts were indeed
- hydrous at run conditions. Mineralogical and textural features of the experiments are
- summarized in Table 2. Representative BSE images of the run products are displayed in Figs. 2
- and 3. Mineral compositions that were obtained from spot microprobe analyses are presented in
- Figs. 4-10, and 12. Additional BSE images and mineral composition profiles (Figures S1-S8) can
- be found in the Supporting Information, and electron microprobe data are archived at Mendeley
- 194 Data (http://dx.doi.org/10.17632/895f6y8chd.1).

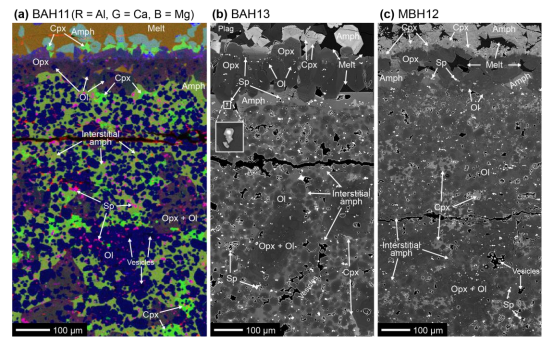




- 196 Figure 2. Back-scattered electron (BSE) images showing the results of the reactive crystallization experiments. (a)
- 197 BAH13 (hydrous basaltic andesite vs. lherzolite, 12 h reaction). (b) MBH12 (hydrous MORB vs. lherzolite, 3 h
- reaction). The left panels are the entire experimental charges, and the right panels are the views of the gabbronorite
- regions. Ol = olivine; Opx = orthopyroxene; Cpx = clinopyroxene; D-cpx = disseminated clinopyroxene; Sp = 200 spinel: Amph = amphibole: Plag = plagioglase: Opx ite = orthopyroxenite
- 200 spinel; Amph = amphibole; Plag = plagioclase; Opx-ite = orthopyroxenite.

201 3.1. Lithologies and textures

After reaction at 1200°C and 1 GPa and step cooling to 880°C and 0.8 GPa, each 202 experimental charge consists of three distinct lithological zones: an amphibole gabbronorite, an 203 amphibole-bearing peridotite, and an amphibole-bearing orthopyroxenite (100-300 µm thick) 204 that separates the former two lithological zones (Fig. 2 and Figure S1). The amphibole 205 gabbronorite, which is absent in the isothermal experiments of Tursack and Liang (2012) and 206 Wang et al. (2016), consists of orthopyroxene, clinopyroxene, amphibole, plagioclase, and 207 quenched melt (Fig. 2). Orthopyroxenes are subhedral and often surrounded by large euhedral 208 amphiboles or present as inclusions around the central part of amphiboles (Fig. 2 and Figure S1). 209 Clinopyroxene grains are anhedral and small. They are generally enclosed around rim parts of 210 the large amphiboles, and some clinopyroxenes in gabbronorite from MBH12 are dispersed in 211 melt or enclosed in plagioclases (Fig. 3b). Plagioclases are tabular-shaped with modal abundance 212 decrease from Runs BAH11 and BAH13 to Run MBH12. Ulvöspinel is observed in the 213 gabbronorite section in experiment BAH11 (Figure S1), a byproduct of reactive crystallization at 214 the reducing condition. 215



216

Figure 3. False-colored composite X-ray concentration map (a) and back-scattered electron (BSE) images (b and c) showing the amphibole-bearing orthopyroxenite and peridotite regions of experimental charges. (a) BAH11 (hydrous basaltic andesite vs. lherzolite, 3 h reaction). (b) BAH13 (hydrous basaltic andesite vs. lherzolite, 12 h reaction). (c) MBH12 (hydrous MORB vs. lherzolite, 3 h reaction). Ol = olivine; Opx = orthopyroxene; Cpx = clinopyroxene; Sp = spinel, Amph = amphibole; Plag = plagioclase.

The amphibole-bearing peridotite consists of olivine, orthopyroxene, clinopyroxene, spinel, and amphibole (Fig. 3). Olivines are round in shape and small in size ($<60 \mu$ m in diameter). Orthopyroxenes are large (up to 350 µm in length) and core-to-rim zoned in the X-ray concentration map and BSE images (Fig. 3). The orthopyroxenes enclose small rounded olivines and spinels forming a poikilitic texture. Clinopyroxenes are embayed in shape and low in modal abundance ($<7 \nu$ ol%). Clinopyroxene is more abundant in the peridotite when the reacting melt

is the hydrous MORB (run MBH12) than those when the reacting melt is the hydrous basaltic

andesite (BAH11 and BAH13, cf. Fig. 3c to Figs. 3a and 3b). Some spinels exhibit euhedral

darker rims in the BSE images (insert in Fig. 3b). Amphibole is crystallized interstitial to other

- minerals throughout the peridotite. Empty vesicles are present where small olivine grains are
- abundant, but absent in the poikilitic orthopyroxenes. Plagioclase and interstitial melt are not
- observed in the peridotite region.

234 The amphibole-bearing orthopyroxenite layer between the amphibole gabbronorite and peridotite is mainly composed of large euhedral orthopyroxenes (up to 200 µm in length) (Fig. 235 3). These orthopyroxene grains contain small olivine and spinel inclusions (Fig. 3), similar to 236 those formed in the isothermal hydrous reaction experiment BAH8 from Wang et al. (2016). 237 Clinopyroxene and amphibole are observed around the gabbronorite-orthopyroxenite boundary, 238 whereas interstitial melt is observed around the orthopyroxenite-peridotite interface. 239 Interestingly, on the gabbronorite side of the orthopyroxenite in MBH12, orthopyroxenes are 240 partially dissolved and replaced by a layer of clinopyroxene (Fig. 3c), suggesting a reaction 241 relationship between the two minerals. 242

243 3.2. Variations in mineral composition

Compositional variations in minerals from the hydrous melt-peridotite reaction 244 experiments are observed both within grain and across lithological regions. Grain-scale variation 245 exists in orthopyroxene, amphibole, plagioclase, and spinel. Cores of large orthopyroxene and 246 amphibole grains have higher Mg# [Mg# = $100 \times Mg/(Mg + Fe)$, in molar] than the rims 247 (Figures S2 and S3). Plagioclase displays oscillatory zoning with an overall decreasing in An# 248 $[An # = 100 \times Ca/(Ca + Na), in molar]$ and Al_2O_3 content from core to rim (Figure S4). Spinel 249 grains exhibit core-to-rim zoning in the BSE images (insert in Fig. 2b). The darker rims likely 250 have lower Cr# [Cr# = $100 \times Cr/(Cr + Al)$, in molar] and higher Mg# than the lighter cores. 251 These core-to-rim variations are consistent with crystallization trends. 252

Figures 4 and 5 display compositional variations of orthopyroxene and amphibole across 253 the gabbronorite, orthopyroxenite, and peridotite sequence. (Compositional profiles for other 254 minerals can be found in Figures S5-S8.) Zero distance marks the approximate position of the 255 gabbronorite-orthopyroxenite interface at the end of the run. This interface is also the boundary 256 separating the crystalline region (melt-bearing orthopyroxenite and dunite) and the crystal-free 257 reacting melt during the reaction at 1200°C (Fig. 1a). Mineral compositions are distinct across 258 the interface for both orthopyroxene and amphibole, although the trends are somewhat obscured 259 by core-to-rim variations. The differences are more pronounced in run BAH11 (basaltic andesite 260 as reacting melt, 3 h reaction at 1200°C, cyan symbols) than the other two runs (blue and red 261 symbols). For longer reaction time (BAH13) and more primitive reacting melt composition 262 (MBH12), compositions of amphiboles and orthopyroxenes in gabbronorite become more similar 263 to those in the peridotite. SiO₂ content in amphibole and Mg# in orthopyroxene and amphibole 264 from the gabbronorite layer are lower than those from the peridotite and orthopyroxenite layers, 265 whereas TiO₂ and MnO contents are higher in the gabbronorite. Na₂O abundance decreases in 266 orthopyroxenes (Fig. 4g) and increases in amphibole (Fig. 5h) from peridotite to gabbronorite. 267 Orthopyroxene and amphibole compositions from each experiment exhibit large variabilities at 268 the same distance in the compositional profiles, consistent with the core-to-rim compositional 269 270 variations (Figs. 4 and 5).

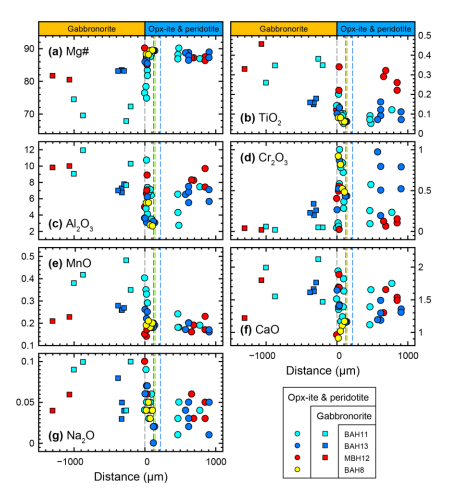


Figure 4. Plots of Mg# and oxide abundances (in wt%) in orthopyroxene as a function of distance from the gabbronorite-orthopyroxenite interface. Circles represent orthopyroxene in the amphibole-bearing orthopyroxenite

273 gabbionome-orthopyroxenic interface. Circles represent orthopyroxene in the amphibole-bearing orthopyroxen274 (opx-ite) and peridotite regions, and squares represent those in the amphibole gabbronorite region of each

experiment. Also shown are compositions of orthopyroxene in orthopyroxenite from the isothermal reaction

experiment BAH8 from Wang et al. (2016). The dashed gray lines mark the position of gabbronorite-

277 orthopyroxenite interface, and the dashed cyan, blue, red, and yellow lines mark the positions of orthopyroxenite-

278 peridotite interfaces in BAH11, BAH13, BAH12, and BAH8, respectively.

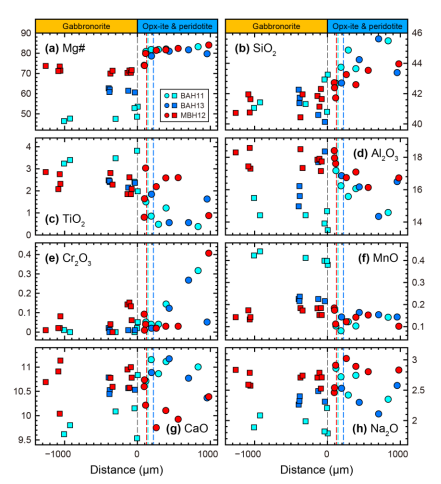
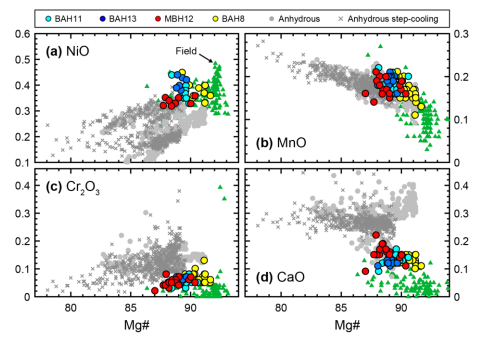


Figure 5. Plots of Mg# and oxide abundances (in wt%) in amphibole as a function of distance from the
 gabbronorite-orthopyroxenite interface. Circles represent amphibole in the amphibole-bearing orthopyroxenite (opx ite) and peridotite regions, and squares represent those in the amphibole gabbronorite region of each experiment.
 The dashed gray lines mark the position of gabbronorite-orthopyroxenite interface, and the dashed cyan, blue, and
 red lines mark the positions of orthopyroxenite-peridotite interfaces in BAH11, BAH13, and BAH12, respectively.

285 3.3. Comparison with previous melt-rock reaction experiments

The lithological sequence and mineral compositions of the hydrous melt-peridotite 286 reaction experiments are significantly different from those of the anhydrous cases that have 287 nearly identical starting melt and peridotite compositions. The most remarkable difference 288 between the hydrous reactive crystallization experiments presented here and the anhydrous ones 289 290 reported by Tursack and Liang (2012) and Saper and Liang (2014) is the formation of amphibole-bearing orthopyroxenite and amphibole-bearing peridotite in the hydrous experiments 291 (Figs. 2 and 3). The two anhydrous reactive crystallization studies (reaction at 1 GPa and 1300-292 293 1320°C, followed by a step cooling to 1200°C or 1050°C) used the same starting basalt (sample D44A) as the hydrous run MBH12 but without addition of water. These experiments produced a 294 295 gabbronorite-wehrlite or a gabbronorite-wehrlite-peridotite sequence. Plagioclase is observed in 296 the wehrlite and peridotite in experiments where the starting lherzolite was mixed with 12% 297 basalt (Saper and Liang, 2014). Plagioclase is not observed in the peridotite layer in the present 298 study.



300 Figure 6. Variations of oxide abundances (in wt%) vs. Mg# in olivine from melt-peridotite reaction experiments and 301 those from field observations of amphibole-bearing mantle rocks (green triangles). Color circles are olivines from the hydrous melt-peridotite reaction experiments BAH11, BAH13, and BAH12 from this study and BAH8 from 302 303 Wang et al. (2016). Gray circles are olivines from anhydrous isothermal melt-peridotite reaction experiments 304 (Harzdis24 from Morgan and Liang, 2003; Lherzdis4 and Lherzdis5 from Morgan and Liang, 2005; PDET1 from 305 Tursack and Liang, 2012), and gray crosses are those from anhydrous reactive crystallization experiments (PDET1b, 306 PDET1c, and PDET2b from Turack and Liang, 2012; PDLS6 and PDSL11 from Saper and Liang, 2014), all 307 conducted in the same laboratory using similar experimental design and starting compositions. The Mantle xenoliths 308 are from the North China Craton (Xu and Bodinier, 2004; Xu et al., 2010), western US (Wilshier and Schwarzman, 309 1971; Francis, 1976a, 1976b; Wilshier et al., 1980), Hawaii (Clagu and Bohrson, 1991), Antarctica (Coltorti et al., 310 2004; Bonadiman et al., 2014), southeastern Australia (O'Reilly et al., 1991; Powell et al., 2004), Kerguelen Islands 311 (Moine et al., 2001), Rhenish Uplands (Witt and Seck, 1989; Witt-Eickschen et al. 2003), Avacha (Bénard and 312 Ionov, 2013), and Ahaggar (Dautria et al., 1987). The massif peridotites are from Lherz (Zanetti et al., 1996), 313 Mariana Trench (Ohara and Ishii, 1998), Zabargad Island (Agrinier, 1993), and Sulu Orogen (Li et al., 2018).

Figures 6-10 compare mineral compositions obtained from the hydrous melt-peridotite 314 reaction experiments with those from the anhydrous ones (for olivine, pyroxenes, spinel, and 315 plagioclase) and phase equilibrium experiments (for amphibole and plagioclase). In general, NiO 316 is positively and MnO is negatively correlated with Mg# in olivine from the melt-rock reaction 317 experiments (Figs. 6a and 6b). Olivine composition from the hydrous melt-rock reaction 318 experiments covers a considerably narrower range than both the isothermal and the step-cooling 319 experiments conducted under anhydrous conditions (Fig. 6). They cluster at the high-Mg# and 320 high-NiO end in the NiO vs. Mg# diagram (Fig. 6a). Olivines from the hydrous experiments 321 have lower CaO and Cr₂O₃ contents and slightly higher MnO content than those from the 322 anhydrous ones (Figs. 6b-6d). Among the hydrous reaction experiments, olivine from the step-323 cooling ones (cvan, blue, and red circles in Fig. 6) have slightly lower Mg# than the hydrous 324 isothermal one BAH8 (yellow circles in Fig. 6). 325

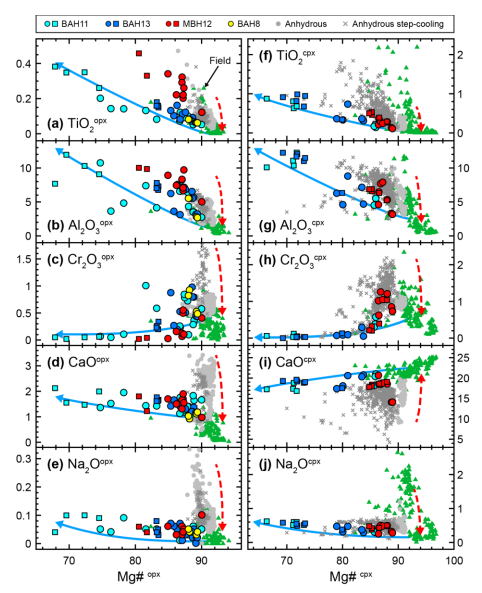


Figure 7. Variations of oxide abundances (in wt%) vs. Mg# in orthopyroxene (a-e) and clinopyroxene (f-j) from melt-peridotite reaction experiments and those from field observations of amphibole-bearing mantle rocks (green triangles). Circles are orthopyroxene in the amphibole-bearing orthopyroxenite and peridotite regions, and squares are those in the amphibole gabbronorite regions in experiments from this study. Source of the experimental and field data are the same as in Fig. 6. The dashed red arrows mark the melting and re-equilibration trend, and the solid blue arrows mark the melt-rock reaction-crystallization trend in orthopyroxene and clinopyroxene compositional variations.

Two types of variation trends are observed in orthopyroxene and clinopyroxene 334 compositions from the melt-rock reaction experiments: the melting and reequilibration trend and 335 the reaction-crystallization trend (Wang et al., 2013). The former is characterized by decreases of 336 incompatible elements (e.g., TiO₂, Al₂O₃, and Na₂O) and a slight increase of Mg# in pyroxenes 337 (dashed red arrows in Fig. 7), and the latter by gradual enrichments of the incompatible elements 338 with a significant decrease of Mg# (blue arrows in Fig. 7). Compositions of orthopyroxene from 339 the isothermal hydrous melt-rock reaction experiment BAH8 plot at the high-Mg# ends on the 340 oxide-Mg# variation diagrams. Compositions of pyroxenes from the hydrous reactive 341 crystallization experiments further expand the melt-rock reaction trend that are mainly 342

- established by pyroxenes from the anhydrous reactive crystallization experiments of Tursack and
 Liang (2012) and Saper and Liang (2014) with Mg# down to lower than 70.
- Figure 8 plots Cr# against Mg# and TiO₂ content in spinel from the melt-rock reaction experiments. Spinel from the hydrous reactive crystallization experiments is scattered in Cr# (6-55), consistent with the observed core-to-rim variations in spinels from the hydrous reactive
- crystallization experiments (inset in Fig. 3b). Spinels from the isothermal run BAH8 are plotted
- at the high-Cr# end of data from the hydrous experiments. Mg# (46-61) and TiO₂ content (0.19-0.40 wt%) in spinel from the hydrous experiments are relatively uniform, both lower than those
- in the anhydrous experiments.

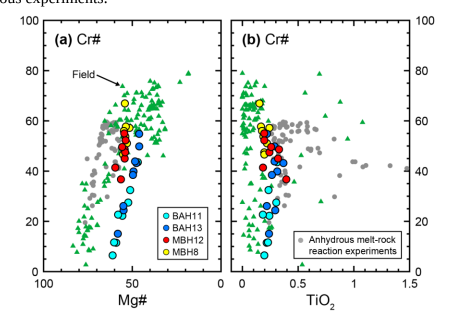


Figure 8. Variations of Cr# as a function of Mg# or TiO₂ abundance (in wt%) in spinel from melt-peridotite reaction experiments and those from field observations of amphibole-bearing mantle rocks (green triangles). Color circles are spinels in experiments from this study. Gray circles are spinels from anhydrous reaction experiments (PDET1, PDET1b, and PDET1c) of Turack and Liang (2012). Source of the field data are the same as in Fig. 6.

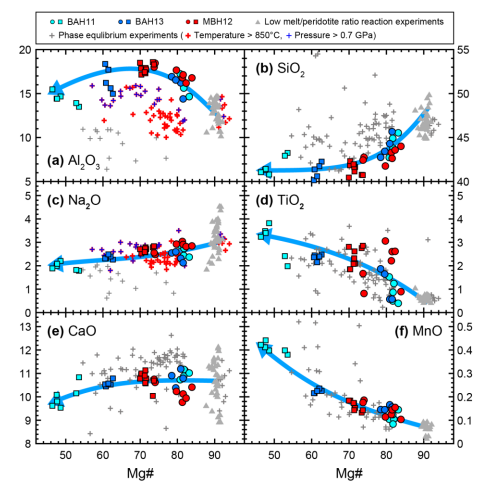
Figure 9 compares compositions of amphibole from this study with those from the meltperidotite reaction experiments of Sen and Dunn (1994), Mallik et al. (2015), and Mandler and Grove (2016) that have low starting melt-to-peridotite ratios (gray triangles). Also shown in Fig.

360 9 are amphiboles from phase equilibrium experiments for hydrous basaltic and andesitic

361 compositions (plus symbols). Compositions of amphibole from the melt-rock reaction

- 362 experiments broadly vary along continuous trends (arrows), different from the phase equilibrium
- experiments which have relatively scattered amphibole compositions. Amphiboles from the low
- melt-to-peridotite ratio reaction experiments are plotted at the high-Mg# end (88-92) of the oxide
- vs. Mg# variation diagrams. Amphiboles in the peridotite and orthopyroxenite regions from our hydrous reaction experiments have lower Mg#, and those from the gabbronorite region have the
- lowest Mg#. Amphiboles from phase equilibrium studies have lower Al_2O_3 and higher SiO₂ than
- those from the melt-rock reaction experiments at a given Mg# (Figs. 9a and 9b), whereas they
- 369 broadly overlap the melt-rock reaction trends in the CaO, TiO₂, Na₂O, and MnO vs. Mg#

diagrams (Figs. 9c-9f).



372 Figure 9. Variations of oxide abundances (in wt%) as a function of Mg# in amphibole from melt-peridotite reaction 373 experiments (circles, squares, and triangles) and phase equilibrium experiments (plus symbols, Nicholls and Harris, 1980; Green and Pearson, 1985; Latourrette et al., 1995; Feig et al., 2006; Mercer and Johnston, 2008; Parat et al., 374 375 2008; Pietranik et al., 2009; Nandedkar et al., 2016). Circles are amphiboles in the amphibole-bearing orthopyroxenite and peridotite regions, and squares are those in the amphibole gabbronorite region in reaction 376 377 experiments from this study (identical to those in Fig. 5). Triangles are amphiboles in the low melt-to-rock ratio 378 reaction experiments (Sen and Dun 1994; Mallik et al. 2015; Mandler and Grove 2016). Blue arrows mark the 379 amphibole compositional variation trends established by the melt-peridotite reaction experiments. Data from the 380 phase equilibrium studies are discriminated by experimental temperature and pressure in (a) and (c).

Figure 10 compares plagioclase compositions from melt-rock reaction experiments with those from phase equilibrium experiments. Plagioclase compositions from the reaction experiments are generally low in An#, FeO and MgO contents than the phase equilibrium experiments except some experiments that were run at subsolidus conditions (Borghini et al., 2010; Fumagalli et al., 2017). Plagioclase from hydrous melt-rock reaction experiments has similar An# and Al₂O₃ and FeO contents to, but significantly lower MgO content than, those from anhydrous ones in Saper and Liang (2014) (Fig. 10c).

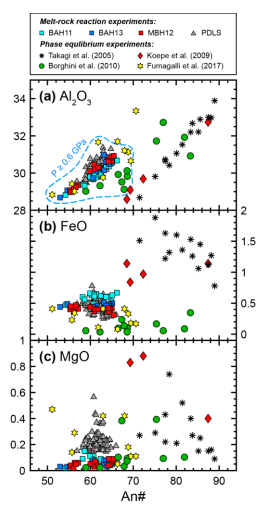
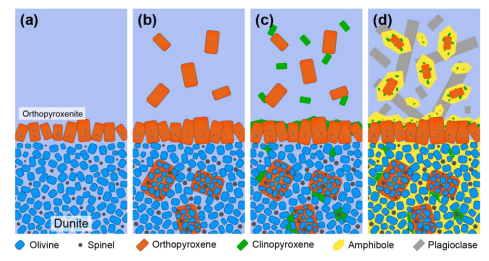


Figure 10. Variations of Al₂O₃, FeO, and MgO abundances (in wt%) vs. An# in plagioclase from melt-peridotite reaction experiments and phase equilibrium experiments. The reaction experiments include those from this study and the anhydrous ones (PDLS) from Saper and Liang (2014). The phase equilibrium experiments are from Takagi et al. (2005), Koepke et al. (2009), Borghini et al. (2010), and Fumagalli et al. (2017). The dashed field in (a) shows the range of data from experiments that were run at pressures greater than 0.6 GPa.

394 3.4. Origin of texture and mineral compositional variations

395 Texture of the melt-rock reaction experiments depends on major element composition and water content of the reacting melt, and P-T-t history. Reaction between the hydrous basaltic 396 andesite and lherzolite at 1200°C and 1 GPa produced a high-porosity orthopyroxenite-dunite 397 sequence (run BAH8 in Wang et al., 2016). The dunite is formed by hydrous melting of 398 lherzolite, and the orthopyroxenite is formed by reaction between olivine and the interface melt 399 with orthopyroxene as the liquidus phase (Fig. 11a). Oversaturation of orthopyroxene in the 400 401 hydrous reacting melt leads to the crystallization of orthopyroxene upon subsequent cooling, forming the orthopyroxene oikocrysts in the peridotite (Figs. 3 and 11b). Clinopyroxene and 402 amphibole are near-liquidus phases of interstitial melts in the peridotite and orthopyroxenite. 403 Stability of clinopyroxene on the liquidus increases from hydrous basaltic andesite to hydrous 404 basalt (e.g., Weaver et al., 2011). This explains the more abundant clinopyroxene in run MBH12 405 than in runs BAH11 and BAH13, and the replacement of orthopyroxene by clinopyroxene 406 around the gabbronorite-orthopyroxenite interface in MBH12 (Figs. 3 and 11c). Orthopyroxene 407

- and clinopyroxene are surrounded by amphiboles (Figs. 2, 3, and 11d), suggesting the delayed
- 409 crystallization of the latter. In the anhydrous reactive crystallization experiments, clinopyroxene
- 410 is the main phase crystallized during cooling (Tursack and Liang, 2012) and plagioclase is
- 411 precipitated when melt infiltration is extensive (Saper and Liang, 2014). The presence of water in 412 reacting melt in the hydrous reactive crystallization experiments stabilizes amphibole which
- 412 reacting ment in the hydrous reactive crystallization experiments stabilizes amphibole which413 crystallizes throughout the peridotite region (Fig. 3). Crystallization of amphibole is at the
- expenses of clinopyroxene and hydrous melt, resulting in the embayed clinopyroxenes in the
- 415 peridotite and the small clinopyroxene inclusions near the rims of large amphiboles in the
- 416 gabbronorite (Figs. 3 and 11d). The empty vesicles formed during quench represent water
- 417 exsolved from interstitial melt upon cooling. Surface tension and capillary suction drive water to
- the porous olivine matrix. The euhedral shape of plagioclase grains in the gabbronorite region
- suggests that plagioclase is the last phase of crystallization.



420

421 Figure 11. Schematic diagrams illustrating processes leading to the formation of amphibole-bearing gabbronorite –

422 orthopyroxenite - peridotite sequence in the hydrous melt and lherzolite reactive crystallization experiments. (a)

Formation of orthopyroxenite-dunite sequence by reaction between hydrous melt and peridotite at 1200°C (adapted

424 from Wang et al., 2016). (b-d) Successive crystallizations of orthopyroxene, clinopyroxene, amphibole, and

425 plagioclase during the step-cooling.

Mg# and NiO content in olivine are governed by extent of peridotite melting. The high 426 degree of peridotite melting promoted by hydrous melt infiltration at 1200°C accounts for the 427 higher Mg# and NiO contents in olivine from the hydrous experiments than those from the 428 anhydrous ones (Fig. 6a). The hydrous reaction experiments were run at relatively low 429 430 temperatures (1200°C with a cooling to 880°C) compared to the anhydrous ones (1300-1320°C with a cooling to 1200°C or 1050°C or without), which gives rise to the lower CaO in olivine, 431 the lower CaO in orthopyroxene, and higher CaO in clinopyroxene from the hydrous 432 433 experiments than those in the anhydrous ones (e.g., Brey and Köhler, 1990; Köhler and Brey, 1990, Figs. 6 and 7). Such relationship between temperature and CaO in olivine and pyroxenes 434 has been also observed in previous melt-rock reaction studies (e.g., Wang et al., 2013, 2016; 435 Saper and Liang, 2014). Step-cooling expands the temperature ranges of the runs and hence the 436 larger ranges of Mg# in pyroxenes and Cr# in spinel along the melt-rock reaction trends in Figs. 437 7 and 8. 438

The variations in amphibole composition from the experimental studies can be attributed to the differences in temperature, pressure, and melt composition. From the low melt-to-rock

ratio reaction experiments (Sen and Dunn, 1994; Mallik et al., 2015; Mandler and Grove, 2016) 441 442 to the peridotite and orthopyroxenite regions to the gabbronorite region in the present experiments, Mg# in amphibole decreases, in accordant with the difference in melt composition 443 from which amphibole is crystallized (Fig. 9). The variation trends of TiO₂ and MnO versus Mg# 444 in amphibole from the melt-rock reaction experiments are broadly similar to those from the 445 phase equilibrium experiments (Figs. 9d and 9f), whereas Al₂O₃ and Na₂O in amphibole from the 446 reaction experiments are generally higher than those from the phase equilibrium experiments. 447 Al₂O₃ in amphibole increases with temperature (for tetrahedral Al) and pressure (for octahedral 448 Al) (e.g., Spear, 1981; Ernst and Liu, 1998; Larocque and Canil, 2010), and Na₂O in amphibole 449 increases with pressure (e.g., Ernst and Liu, 1998; Niida and Green, 1999). Amphibole from 450 experiments run at high temperatures (> 850° C) or at high pressures (> 0.7 GPa) has relatively 451 high Al_2O_3 content, and amphibole from experiments run at high pressures (> 0.7 GPa) has 452 relatively high Na₂O content (Fig. 9). These temperature- and pressure-dependent Al₂O₃ and 453 Na₂O variations are consistent with the grain-scale variations of amphibole composition: as 454 temperature and pressure decrease during step-cooling, the amphibole grows with decreasing 455 Al₂O₃ and Na₂O contents in the rim (Fig. S3). In spite of the numerous factors that control 456 amphibole composition during melt-rock reaction, the laboratory melt-rock reaction experiments 457 establish general variation trends (blue arrows in Figs. 9 and 12). As melt-rock reaction or 458 diffusive exchange between reacting melt and minerals becomes extensive, Mg# and Na₂O in 459 amphibole decrease, whereas MnO, TiO₂, and Al₂O₃ in amphibole increase. 460

Composition of plagioclase can be affected by temperature, pressure, and melt 461 composition (Longhi et al., 1976; Namur et al., 2012; Faak et al., 2013). As demonstrated by 462 Saper and Liang (2014), reaction of MORB with peridotite at a moderate pressure delays the 463 crystallization of plagioclase. Presence of water also delays plagioclase crystallization in basalts 464 (Takagi et al., 2005; Borghini et al., 2010; Namur et al., 2012). In our hydrous reactive 465 crystallization experiments, plagioclase is the last mineral to crystallize. This explains the 466 relatively low An# in plagioclase from these experiments, compared with that from the phase 467 equilibrium studies (Fig. 10a). Because of the late-stage crystallization (i.e., from an evolved 468 residual melt at low P and T), FeO and MgO contents in plagioclase from the reaction 469 experiments are low, comparable to those from the subsolidus phase equilibrium experiments of 470 Borghini et al. (2010) and Fumagalli et al. (2017) (Figs. 10b and 10c). In contrast, FeO and MgO 471 contents in plagioclase from the phase equilibrium experiments of Takagi et al. (2005) and 472 Koepke et al. (2005) are high (Figs. 10b and 10c), as it is the near-liquidus mineral in these 473 experiments. 474

475 **4. Geological applications**

The experiments reported here and those in Tursack and Liang (2012), Saper and Liang 476 (2014), and Wang et al. (2016) underscore the importance of water in the reacting melt in 477 determining the lithology and mineral compositional variations produced by melt-rock reaction 478 in the mantle. Hydrous basalts-peridotite reaction and in situ crystallization can produce a 479 spatially associated amphibole-bearing peridotite and amphibole-bearing orthopyroxenite 480 sequence. Because of the simplified experimental setup, the mass transfer mechanism may be 481 different between melt-peridotite interaction in the laboratory experiments and those in nature 482 (e.g., diffusion vs. advection). Extensive water infiltration into the powdered starting peridotite 483 lead to partial melting and faster reaction. The rate of cooling in the experiments is likely faster 484 than that takes place in nature. Nevertheless, the kinetics and compositional variation trends are 485

486 quite similar between the laboratory experiments and those in nature, because they are

determined by chemical potential gradients of the major components and phase relationships in

the hydrous basalt-peridotite system. The fast reaction and crystallization rates manifest grain-

489 scale processes and help to develop chemical variations on the laboratory time scale and length 490 scale. To assess the effect of hydrous melt-peridotite interaction in modifying composition and

490 scale. To assess the effect of hydrous melt-peridotite interaction in modifying composition and 491 mineralogy of the lithospheric mantle, we compare mineral compositions obtained from

491 Inhieratogy of the https://energiene.compare inhierat compositions obtained from492 laboratory melt-rock reaction experiments with those from the field in Figs. 6-9, and 12.

493 Calculation of chemical formula and nomenclature of the amphibole supergroup follow the

494 method described in Leake et al. (1997). We prefer to use this Si-Al-Na-K based amphibole

495 classification in this study because abundances and substitution of these elements are important

496 indications of melt composition and temperature and pressure conditions.

497

4.1. Olivine, pyroxenes and spinel in amphibole-bearing mantle rocks

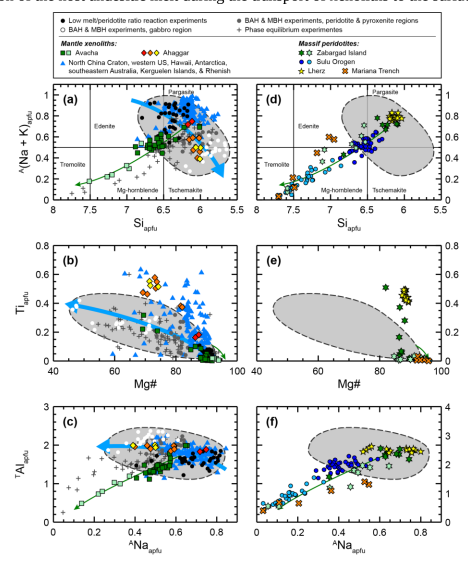
498 Figures 6, 7 and 8 compare compositions of olivine, pyroxenes, and spinel obtained from laboratory melt-peridotite reaction experiments with those in the amphibole-bearing mantle rocks 499 from the field. Compositional variations of these minerals from the field samples broadly follow 500 the trends established by the melt-peridotite reaction experiments, although clinopyroxenes from 501 the reaction experiments are lower in Mg# than the field data. The mismatch in Mg# between the 502 experimental data and the field data are also observed in the amphibole compositions (Figs. 12b 503 and 12e). These are in part due to the evolved nature of reacting melts and high melt-to-504 peridotite ratios in the experiments and the subsolidus state of the natural samples. In previous 505 506 experimental studies (e.g., Morgan and Liang, 2005; Van den Bleeken et al., 2010; Tursack and Liang, 2012; Wang et al., 2013; Saper and Liang, 2014), common mismatches in CaO 507 concentrations in olivine and pyroxenes were observed between field data and melt-rock reaction 508 experimental data: clinopyroxenes are higher in CaO while olivines and orthopyroxenes are more 509 depleted in CaO for the field data. These are attributed to subsolidus reequilibration of the field 510 samples and the high temperature nature of the dissolution experiments (>1200°C). Figures 6 511 and 7 show that CaO contents in olivine and pyroxenes from the hydrous reaction experiments 512 are close to those from the field observations, consistent with the low reaction and crystallization 513 514 temperatures in these experiments (1200°C to 880°C).

515

4.2. Amphiboles from mantle xenoliths

Figures 12a-12c compare compositions of amphibole obtained from the laboratory 516 experiments with those from mantle xenoliths. Amphiboles in the mantle xenoliths are mainly 517 pargasite and edenite with a small population being Mg-hornblende and tremolite. The main 518 composition range of amphibole in the mantle xenoliths overlaps that obtained from the melt-519 rock reaction experiments (gray fields). Harzburgite xenoliths in andesites from Avacha, 520 Kamchatka (Bénard and Ionov, 2013) contain amphiboles in different lithological regions, 521 including the host harzburgite, orthopyroxene-rich veins that have reaction texture with host 522 harzburgite, and orthopyroxene-rich veins without reaction texture. The reaction texture between 523 the orthopyroxene-rich veins and the host harzburgite is similar to that observed around the 524 orthpyroxenite-peridotite interface in our experiments (Fig. 3). Amphiboles in the host 525 harzburgite and those in the veins with reaction texture (dark green squares) are mainly pargasite 526 and subordinately enenite and Mg-hornblende (Fig. 12a). Compositions of these amphiboles are 527 mainly in the range defined by the melt-rock reaction experiments (gray fields). The textural and 528 529 compositional features suggest that these veins were formed by reaction of peridotite and

- hydrous melt in the lithospheric mantle (Bénard and Ionov, 2013). Amphiboles in the host
- harzburgite were likely crystallized from reacting melt that infiltrated into the peridotite. In
- contrast, the other type of orthopyroxene-rich veins crosscut the host harzburgite, and they have
- a straight contact with the host harzburgite. Amphiboles in these veins (light green squares) are
- mainly Mg-hornblende (Fig. 12a). They have low T-site Al, A-site Na, Ti abundances, and high
- 535 Mg#, varying along a trend defined by some amphiboles from phase equilibrium experiments 536 (thin green arrows in Fig. 12). The above evidence indicates that these veins are formed by direct
- 537 crystallization of the host andesitic melt during the transport of xenoliths to the surface.



539 Figure 12. Plots of atomic compositions for amphiboles from the experiments and field observations of amphibole-540 bearing mantle xenoliths (a-c) and massif peridotite (d-f). Calculation of chemical formula and division of compositional boundaries (in a and d) follow Leake et al. (1997). The experimental data are shown as black dots 541 542 (low melt-to-rock ration reaction experiments, shown in Fig. 9), gray dots (peridotite and pyroxenite regions in 543 experiments from the present study), white dots (gabbronorite regions in experiments from the present study), and 544 gray plus symbols (phase equilibrium experiments, shown in Fig 9). The gray fields show composition range of amphibole from melt-rock reaction experiments, and the thick blue arrows indicate the melt-rock reaction trend of 545 546 amphibole compositional variation. Source of the field data are the same as in Fig. 6. Amphiboles in peridotite 547 xenoliths and orthopyroxene-rich veins that have reaction texture with the host peridotite from Avacha are shown as 548 dark green squares, and those in orthopyroxene-rich veins without reaction texture are shown as light green squares.

549 Amphiboles in peridotite xenoliths, amphibole-rich veins and amphibolite xenoliths, and amphibole megacrysts from Ahaggar are shown as red, orange, and vellow diamonds, respectively. Amphiboles from the other mantle 550 551 xenoliths are shown as blue triangles. Amphiboles in spinel peridotites and the amphibole-bearing bands associate 552 with secondary minerals in massif peridotites from Zabargad Island, are shown as dark green hexagons, and 553 amphiboles in the mylonitic share zones are shown as light green hexagons. Amphibole cores and rims from the 554 Sulu peridotites are shown as dark and light circles, respectively. Amphiboles in lherzolites and amphibolite veins 555 from the Lherz massif are shown as yellow pentagrams, and those in serpentinized peridotites from Mariana Trench 556 are shown as orange crosses. The thin green arrows indicate the compositional variation trend of amphiboles that 557 were likely formed by crystallization of host magma of mantle xenolith or by hydrothermal alteration at shallower 558 depths.

Different occurrences of amphiboles are also observed in a suit of mantle xenoliths and 559 megacrysts entrained by basalts from Ahaggar, southern Algeria (Dautria et al., 1987). 560 Amphiboles are present in peridotites with amphibole-rich veins, amphibolites, and as 561 megacrysts in the host basalt. Amphiboles are anhedral grains interstitial to anhydrous minerals 562 in the peridotites and oikocrysts that have chadacrysts of anhydrous minerals in the amphibole-563 rich veins and amphibolites. Although amphibolite is not formed in the hydrous melt-rock 564 reaction experiments, the poikilitic texture of amphibole in the Ahaggar xenoliths is similar to 565 that observed in the peridotite region in the experiments (Fig. 2). Furthermore, amphiboles in the 566 Ahaggar xenoliths with different occurrences show a compositional variation along the trend 567 defined by the melt-rock reaction experiments (thick blue arrows in Fig. 12). Amphiboles in the 568 peridotites (red diamonds) are high in Mg# and low in Ti (Fig. 12b), similar to those in 569 ultramafic regions from our experiments (gray dots) and the low melt-to-rock ratio experiments 570 (black dots). Amphiboles in the amphibole-rich veins and amphibolites (orange diamonds) have 571 low Mg# and high Ti abundance (Fig. 12b), similar to the amphibole megacrysts in the host 572 basalt (yellow diamonds) and those in the gabbronorite region of our experiments (white dots). 573 The variations in occurrence and composition suggest that amphiboles in the Ahaggar peridotites 574 and amphibolites were formed by reactive crystallization when hydrous melts infiltrate 575 peridotite, and the amphibole megacrysts were crystallized from the reacted melt (Dautria et al., 576 1987). The reaction trend is also observed in the amphibole-bearing lherzolites from Victoria 577 Land, Antarctica: amphibole disseminated in peridotite has a higher Mg# and a lower Ti 578 579 abundance than that in the amphibole veins, suggestive of melt-rock reaction origin of the amphiboles with different melt-to-rock ratios (Coltorti et al., 2004). 580

581 4.3. Amphiboles from massif peridotites

Figures 12d-12f compare the experimental amphibole compositions with those from 582 massif peridotites. The massif peridotites from Zabargad Island, Red Sea contain amphiboles 583 interstitial to olivines in spinel peridotite, in bands associated with secondary spinel between 584 bands of olivines, and in mylonitic shear zones (Agrinier et al., 1993). Massif peridotites from 585 the Sulu Orogenic Belt, eastern China have amphiboles with core-to-rim compositional variation, 586 and inclusions of olivine, serpentine and spinel (Li et al., 2018). The peridotite sample from the 587 Lherz massif has an amphibolite vein. Modal abundance of amphibole increases from the host 588 peridotite towards the vein (Zanetti et al., 1996). Massif peridotite samples from the island arc at 589 Mariana Trench are serpentinized and contain tremolite, hornblende, and enenite (Fig. 12d) 590 (Ohara and Ishii, 1998). Compositionally, amphiboles in the spinel peridotite and the secondary 591 bands from Zabargad (dark green hexagons in Figs. 12d-12f), amphibole cores from Sulu (dark 592 blue circles), and amphiboles from Lherz (yellow pentagrams) are mainly in the composition 593 range of the melt-rock reaction experiments. This is consistent with the occurrences that these 594

- amphiboles are interstitial or associated with secondary minerals, and supports the origin that
- they are from by reaction of peridotite with hydrous melts (e.g., Agrinier et al., 1993; Zanetti et
- al., 1996). Amphiboles in the mylonitic shear zones in the massif peridotites from Zabargad
 (light green hexagons in Figs. 12d-12f), amphibole rims in the Sulu orogenic peridotite (light
- blue circles in Figs. 12d-12f), and amphiboles in the serpentinized peridotites from Mariana
- Trench (orange crosses, Ohara and Ishii, 1998) vary along the trend that is different from that
- defined by the melt-rock reaction experiments (thin green arrows in Fig. 12d-12f). The
- 602 compositional variation and the occurrences of these amphiboles indicate that they were formed
- by processes such as hydrothermal alteration (fluid-rock reaction) when the peridotites have
- emplaced or exhumed at shallow depths (Agrinier et al., 1993; Ohara and Ishii, 1998; Li et al.,
- 605 2018).

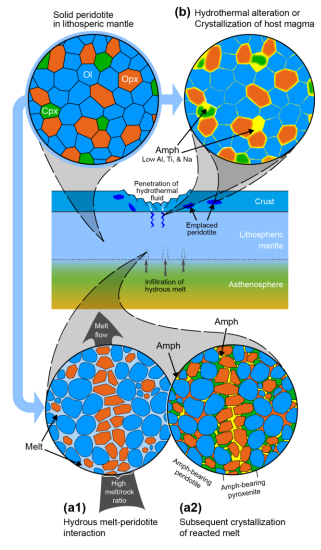


Figure 13. Schematic diagrams illustrating scenarios where amphiboles are formed in the lithospheric mantle. (a1) and (a2) show the processes leading to the formation of amphibole-bearing peridotite and pyroxenite by hydrous

- 609 melt-peridotite interaction. Hydrous melt infiltration initiates partial melting of the peridotite in lithospheric mantle.
- Reaction between the hydrous melt and the peridotite forms orthopyroxene-depleted peridotite and orthopyroxene
- 611 vein or dike (a1). As the peridotite cooled, the interstitial melts crystallize clinopyroxene and amphibole with
- 612 composition varying along the trends established by the melt-rock reaction experiments forming amphibole-bearing 613 peridotite and pyroxenite (a2). (b) shows the formation of amphibole in peridotite by metamorphism. Hydrothermal

fluids penetrate peridotite at the crust-mantle boundary or in the lower crust after peridotite emplacement forming amphibole with high Mg#, SiO₂, and CaO and low Al₂O₃, TiO₂, and Na₂O contents (magenta trends in Fig. 13). Ol =

- 616 olivine; Opx = orthopyroxene; Cpx = clinopyroxene; Amph = amphibole.
- 617 4.4. Formation mechanisms of amphibole-bearing mantle rocks

The amphibole-bearing pyroxenite and amphibole-bearing peridotite that formed in the 618 hydrous melt-peridotite reaction experiments are similar to a number of field observations in 619 terms of texture, lithology, and mineral compositions, indicating that these field amphiboles are 620 likely formed by hydrous melt-peridotite interaction. The schematic diagram in Fig 13 621 summarizes mechanisms through which various amphiboles are formed in the lithospheric 622 mantle. Infiltration of hydrous melts may be focused either by means of hydro-fracturing or 623 reaction induced infiltration instability similar to the one that produces replacive dunite in the 624 mantle section of ophiolite (e.g., Kelemen et al., 1997). When a hydrous melt (or fluid) infiltrate 625 the lithospheric mantle along pressure gradients or fractures, it may induce partial melting of the 626 surrounding peridotite. Further reaction between the hydrous melt and the partially molten 627 peridotite produces a melt-bearing orthopyroxenite vein (Fig. 13a1, Wang et al., 2016). A new 628 generation of clinopyroxene and amphibole then precipitate from the trapped interstitial melts as 629 the peridotite is cooled or water in melt is released (by diffusion or change of pressure). The 630 precipitation may occur in the partially molten peridotite and the melt-bearing orthopyroxenite, 631 forming an amphibole-bearing peridotite and an amphibole-bearing pyroxenite, respectively 632 (Fig. 13a2). It can be inferred that the precipitation may form an amphibole-rich lithology in 633 reaction zone where the proportion of hydrous melt is high. The amphibolites from Ahaggar 634 (Dautria et al., 1987) and amphibole veins in lherzolites from Victoria Land (Coltorti et al., 635 2004) may represent such examples. 636

However, not all amphiboles in mantle rocks are formed by melt-rock reaction. 637 638 Compositions of some amphiboles in natural samples are different from those produced by the melt-rock reaction experiments (Fig. 12). These amphiboles grow as rims of amphiboles formed 639 by melt-rock reaction (e.g., Li et al., 2018), occur in veins that show not textural evidence of 640 reaction with the host peridotites (e.g., Agrinier et al., 1993; Bénard and Ionov, 2013), or present 641 in peridotite with evidence of hydrothermal alteration (e.g., Ohara and Ishii, 1998). They are 642 likely formed by processes such as crystallization of host magma of mantle xenoliths or 643 644 hydrothermal alteration after emplacement or exhumation of peridotite massifs at crustal depths (Fig. 13b). 645

646 **5 Conclusions**

The role of hydrous melt-peridotite reaction in the formation of amphibole-bearing 647 mantle rocks is examined by reacting spinel lherzolite with a hydrous basaltic andesite and a 648 hydrous MORB (with 4 wt% water in the melts) in Au-Pd capsules using the reaction couple 649 method. The experiments include a reaction at 1200°C and 1 GPa and a further 650 reaction/crystallization through a series of step cooling to 880°C and 0.8 GPa. Results of the 651 hydrous melt-peridotite reaction experiments together with those conducted under anhydrous 652 conditions are used to deduce processes leading to the formation of amphibole in mantle rocks. 653 The main conclusions of this study are as follows: 654

- Reaction between hydrous basaltic melt and lherzolite followed by in situ crystallization
 produces an amphibole gabbronorite, amphibole-bearing orthopyroxenite, and amphibole bearing peridotite sequence.
- A high-porosity orthopyroxenite-dunite sequence is formed in the reaction at 1200°C.
 Crystallization of orthopyroxene continues at the early stages of cooling, followed by
 clinopyroxene, and finally amphibole in the peridotite and orthopyroxenite.
- Lithology, texture, and mineral compositional variation are key features that can be used to
 deduce the origin of amphiboles in peridotites and pyroxenites.
- 4. Similarities in these key features between the experiments reported in this study and those
 observed in amphibole-bearing mantle xenoliths and massif peridotites underscore the
 importance of hydrous melt-peridotite reaction and in situ crystallization in the formation of
 amphiboles in these mantle rocks.
- 5. Texture and compositional variations of some amphiboles from the field are distinct from
 those obtained from the melt-peridotite reaction experiments, including the low Ti, Al, and
 Na abundances. These amphiboles are likely formed by crystallization of xenoliths' host
 magma and hydrothermal alterations at shallow depths.

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Supporting Information for

Formation of Amphibole-Bearing Peridotite and Amphibole-Bearing Pyroxenite through Hydrous Melt-Peridotite Reaction and In Situ Crystallization: An Experimental Study

Chunguang Wang^{1, 2, *}, Yan Liang², and Wenliang Xu¹

1. College of Earth Sciences, Jilin University, Changchun 130061, China

2. Department of Earth, Environmental and Planetary Sciences, Brown University, Providence, RI 02912, USA

*Corresponding author: Chunguang Wang (c_wang@jlu.edu.cn)

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Introduction

This file includes figures and data supporting the main text:

- Figure S1 displays BSE images for BAH11;
- Figure S2-S3 show core-to-rim compositional variations in orthopyroxene, amphibole, and plagioclase grains;
- Figure S4-S8 plot olivine, clinopyroxene, and spinel compositions as a function of distance from the gabbronorite-orthopyroxenite interface;
- Tables S1-S3 present electron microprobe data of experimental results.

Oxide abundances are in wt%, and distances are in µm; Total Fe as FeO; NA = not analyzed; ND = not detected; X = distance from gabbronorite-orthopyroxenite interface (X > 0 in the orthopyroxenite and peridotite regions, X < 0 in the gabbronorite region); D = distance from the center of mineral grain. Mg# = $100 \times Mg/(Mg + Fe)$, in molar; Cr# = $100 \times Cr/(Cr + AI)$, in molar; An# = $100 \times Ca/(Ca + Na)$, in molar.

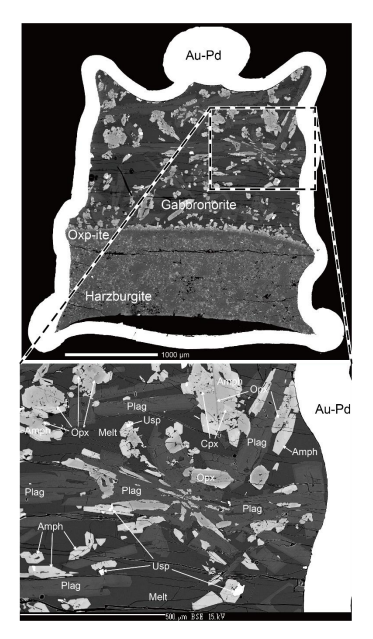


Figure S1. BSE images showing the experimental charge (upper) and a close-up view of the gabbronorite region (lower) of BAH11 (hydrous basaltic andesite vs. Iherzolite, 3 h reaction). Opx-ite = orthopyroxenite; Opx = orthopyroxene; Cpx = clinopyroxene; Amph = amphibole; Plag = plagioclase; Usp = Ulvöspinel.

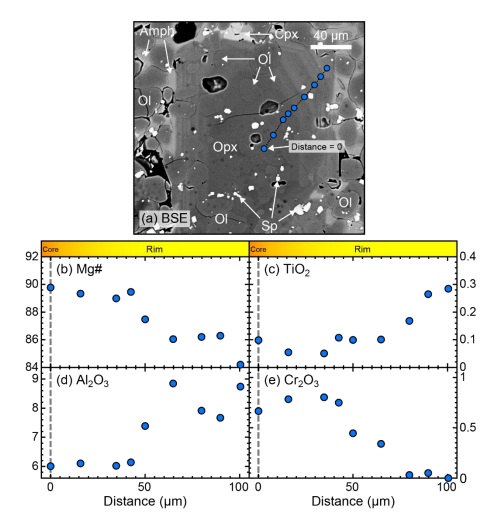


Figure S2. BSE image and electron microprobe data showing the chemical zoning of an orthopyroxene grain. (a) Core-to-rim zoned orthopyroxene in peridotite from BAH13. (b-e) Variations of Mg# and oxide abundances (in wt%) in orthopyroxene as a function of distance from the orthopyroxene core. Blue circles in (a) mark positions of the probe analyses plotted in (b-e).

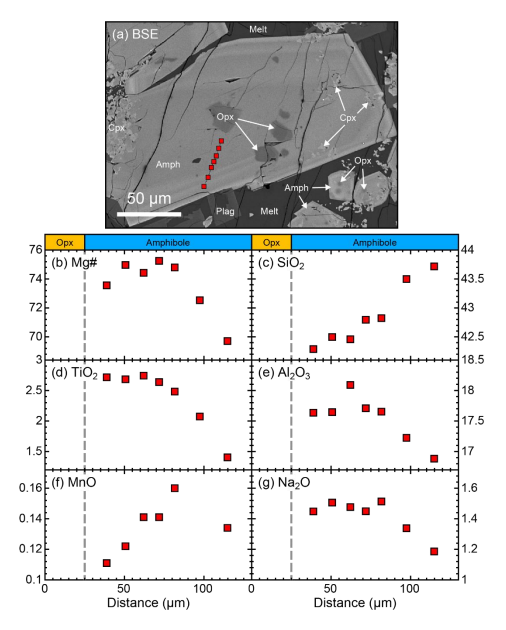


Figure S3. BSE image and electron microprobe data showing the chemical zoning of an amphibole grain. (a) Core-to-rim zoned amphibole in peridotite from MBH12. (b-g) Variations of Mg# and oxide abundances (in wt%) in amphibole as a function of distance from the amphibole core. Red squares in (a) mark positions of the probe analyses plotted in (b-g). Vertical dashed red lines in (b-g) mark the position of orthopyroxene-amphibole boundary. Note inclusions of orthopyroxenes in amphiboles.

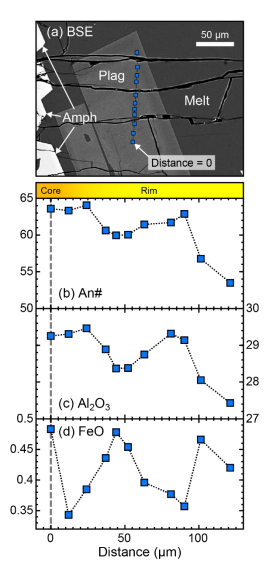


Figure S4. BSE image and electron microprobe data showing the chemical zoning of plagioclase grain. (a) Core-to-rim zoned plagioclase in amphibole gabbronorite from BAH13. (b-d) Variations of An# and oxide abundances (in wt%) in plagioclase as a function of distance from the plagioclase core. Blue squares in (a) mark positions of the probe analyses plotted in (b-d).

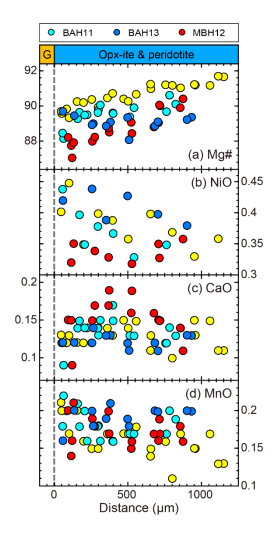


Figure S5. Plots of Mg# and oxide abundances (in wt%) in olivine as a function of distance from the gabbronorite-orthopyroxenite interface (dashed lines). Cyan, blue, red, and yellow circles are data from BAH11, BAH13, BAH12, and BAH8 (Wang et al. 2016), respectively.

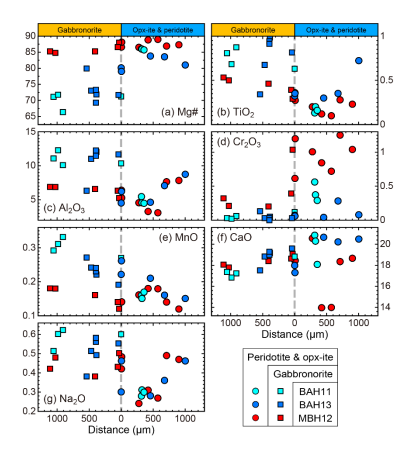


Figure S6. Plots of Mg# and oxide abundances (in wt%) in clinopyroxene as a function of distance from the gabbronorite-orthopyroxenite interface (dashed lines). Circles represent clinopyroxene in the orthopyroxenite and peridotite regions, and squares represent those in the gabbronorite regions.

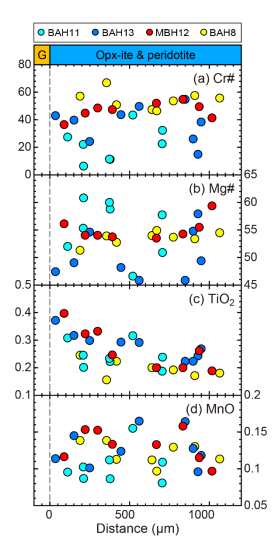


Figure S7. Plots of Cr#, Mg#, and oxide abundances (in wt%) in spinel as a function of distance from the gabbronorite-orthopyroxenite interface (dashed lines).

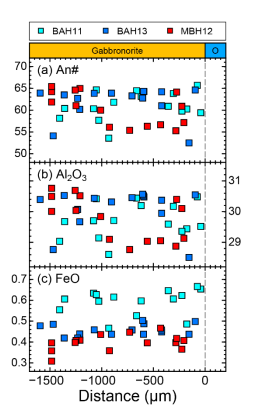


Figure S8. Plots of An# and oxide abundances (in wt%) in plagioclase as a function of distance from the gabbronorite-orthopyroxenite interface (dashed lines).

Table 5.	L-I. Elect		phone da	ta or amp		DAILIT					
SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	Х
42.11	1.93	13.14	0.05	16.68	0.37	10.79	10.55	1.74	97.36	53.56	4
42.00	2.36	13.61	0.00	17.05	0.39	10.71	9.93	1.78	97.83	52.82	-40
39.97	3.74	14.37	0.02	18.28	0.39	9.70	9.34	2.16	97.96	48.61	-5
41.67	1.47	16.78	0.05	7.21	0.10	17.10	10.46	2.79	97.63	80.87	127
42.75	0.83	15.87	0.04	6.96	0.08	17.65	10.90	2.66	97.73	81.89	197
44.10	0.47	15.31	0.04	7.15	0.12	18.04	10.68	2.38	98.27	81.81	298
42.48	1.19	15.63	0.14	6.80	0.10	17.51	10.82	2.67	97.34	82.11	402
44.49	0.37	14.28	0.31	6.63	0.14	18.53	10.76	2.30	97.79	83.28	843
40.14	3.38	14.01	0.00	18.23	0.40	9.27	9.80	1.93	97.17	47.55	-300
40.35	3.31	14.04	0.00	18.33	0.43	9.37	9.55	2.03	97.40	47.68	-924
39.83	3.14	15.02	0.01	18.47	0.41	8.97	9.33	1.83	97.02	46.40	-1007

Table S1-1. Electron microprobe data of amphibole in BAH11

 Table S1-2.
 Electron microprobe data of olivine in BAH11

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SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na₂O	NiO	Total	Mg#	Х
40.34	0.02	0.05	0.05	11.41	0.22	47.60	0.09	ND	NA	99.76	88.15	62
40.56	0.01	0.04	0.06	10.43	0.21	48.71	0.14	ND	NA	100.18	89.28	160
40.68	0.01	0.06	0.04	10.19	0.16	48.71	0.14	ND	NA	99.99	89.50	223
40.46	0.01	0.03	0.05	9.93	0.18	48.89	0.13	0.01	NA	99.69	89.77	300
40.66	ND	0.13	0.06	9.63	0.16	48.96	0.15	0.04	NA	99.79	90.06	313
40.53	ND	0.08	0.05	10.08	0.16	48.76	0.17	ND	NA	99.83	89.61	397
40.43	0.01	0.04	0.06	10.33	0.17	48.81	0.13	0.02	NA	100.00	89.39	512
40.65	ND	0.04	0.05	9.15	0.20	49.58	0.14	ND	NA	99.82	90.62	782
40.86	0.01	0.03	0.05	9.68	0.19	49.50	0.13	ND	NA	100.44	90.11	825
40.65	ND	0.06	0.03	11.12	0.18	47.84	0.12	ND	0.44	100.45	88.46	58
40.89	0.01	0.03	0.07	10.09	0.18	48.99	0.15	ND	0.40	100.81	89.64	172
40.76	0.01	0.03	0.05	10.19	0.20	48.61	0.14	0.01	0.35	100.35	89.48	206
40.74	ND	0.05	0.03	10.11	0.18	48.98	0.14	ND	0.38	100.62	89.62	297
40.92	0.01	0.02	0.07	10.17	0.16	49.05	0.15	ND	0.37	100.91	89.58	400
40.88	0.01	0.06	0.08	9.85	0.17	49.17	0.13	0.01	0.33	100.68	89.90	542
40.78	0.01	0.03	0.05	10.06	0.17	48.98	0.12	ND	0.35	100.55	89.67	763

Table 31	- J . LICCU		probe dat		pyrozene						
SiO ₂	TiO ₂	Al ₂ O ₃	Cr_2O_3	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	х
48.53	0.20	10.72	0.06	14.66	0.33	24.37	1.35	0.05	100.28	74.77	29
50.94	0.13	7.34	1.00	11.14	0.29	27.79	1.42	0.05	100.10	81.64	42
52.46	0.10	6.78	0.78	7.90	0.19	31.03	1.17	0.03	100.45	87.50	60
52.16	0.09	7.11	0.84	7.89	0.19	30.84	1.24	0.05	100.41	87.45	83
54.93	0.06	3.43	0.36	7.17	0.18	32.30	1.63	0.04	100.11	88.93	103
52.69	0.14	3.57	0.04	14.63	0.40	26.52	1.50	0.04	99.54	76.37	1
52.56	0.14	4.78	0.10	13.33	0.35	26.99	1.94	0.09	100.27	78.30	21
53.59	0.08	4.08	0.58	10.54	0.29	29.45	1.41	0.05	100.07	83.28	40
52.77	0.10	6.38	0.72	7.59	0.18	31.09	1.39	0.04	100.27	87.95	94
54.96	0.05	3.46	0.28	7.25	0.18	32.23	1.82	0.06	100.30	88.79	99
55.44	0.05	2.70	0.57	6.49	0.18	32.98	1.65	0.05	100.10	90.06	485
54.38	0.09	4.28	0.09	7.73	0.18	32.09	1.39	0.03	100.24	88.10	472
51.80	0.07	6.50	0.25	8.52	0.24	31.36	1.11	0.01	99.86	86.77	470
51.82	0.12	7.44	0.23	8.23	0.19	30.46	1.75	0.04	100.27	86.84	781
48.05	0.35	10.30	0.05	15.96	0.35	23.40	1.47	0.10	100.02	72.33	-202
48.44	0.38	7.63	0.05	18.43	0.48	21.85	2.11	0.04	99.41	67.88	-264
46.78	0.35	12.00	0.02	17.18	0.42	22.05	1.56	0.10	100.46	69.58	-875
49.05	0.26	9.05	0.06	14.74	0.38	24.20	1.99	0.09	99.80	74.53	-1003

 Table S1-3.
 Electron microprobe data of orthopyroxene in BAH11

 Table S1-4.
 Electron microprobe data of clinopyroxene in BAH11

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SiO ₂	TiO ₂	AI_2O_3	Cr_2O_3	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	Х
46.82	0.63	10.34	0.12	9.33	0.27	12.96	18.80	0.60	99.86	71.23	18
50.63	0.13	5.45	0.56	4.91	0.16	16.97	20.85	0.28	99.94	86.04	335
51.18	0.20	4.53	0.37	5.11	0.15	17.51	20.20	0.31	99.55	85.93	346
51.81	0.16	4.47	0.29	5.64	0.17	19.05	18.06	0.30	99.95	85.76	375
46.31	0.68	12.23	0.02	9.39	0.31	13.39	16.78	0.60	99.69	71.77	-966
46.38	0.87	10.04	0.06	11.44	0.33	12.67	17.15	0.62	99.55	66.38	-891
46.35	0.80	10.99	0.03	9.67	0.29	13.36	17.23	0.51	99.24	71.12	-1037

Table	JI-J. LICCI		probe dat	a or spine	III DAILI.	6					
SiO ₂	TiO ₂	Al ₂ O ₃	Cr_2O_3	FeO	MnO	MgO	CaO	Total	Mg#	Cr#	Х
0.13	12.35	5.12	0.07	76.17	0.27	1.38	0.07	95.56	3.14	0.91	-789
0.11	10.52	6.01	0.02	76.00	0.25	1.45	0.05	94.40	3.28	0.22	-1324
0.12	9.75	7.32	0.05	75.93	0.23	1.60	0.07	95.07	3.62	0.46	-577
0.11	10.37	6.65	0.03	76.32	0.23	1.58	0.05	95.34	3.56	0.30	-367
0.11	11.64	5.26	0.07	77.34	0.27	1.37	0.07	96.14	3.07	0.89	-322
0.13	9.22	8.02	0.02	75.98	0.22	1.66	0.06	95.30	3.75	0.17	-75
0.17	10.25	6.00	0.03	75.96	0.22	1.49	0.10	94.22	3.37	0.34	-6
0.12	0.31	36.70	20.55	25.87	0.10	15.73	0.11	99.49	52.01	27.31	112
0.12	0.24	39.77	16.75	24.06	0.10	16.72	0.15	97.91	55.33	22.03	210
0.17	0.20	52.92	5.28	21.01	0.09	18.30	0.10	98.06	60.82	6.28	212
0.24	0.22	49.16	9.34	22.00	0.09	17.60	0.14	98.78	58.78	11.30	377
0.13	0.23	47.68	9.19	22.21	0.11	18.68	0.07	98.30	59.98	11.45	373
0.13	0.31	26.32	30.04	26.84	0.15	13.15	0.05	96.98	46.61	43.37	520
0.13	0.23	33.12	23.52	24.75	0.11	14.40	0.13	96.39	50.92	32.27	706
0.11	0.19	41.03	17.75	22.56	0.08	17.30	0.11	99.12	57.74	22.49	704

Table S1-5. Electron microprobe data of spinel in BAH11

 Table S1-6.
 Electron microprobe data of plagioclase in BAH11

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SiO ₂	TiO ₂	Al ₂ O ₃	FeO	MgO	CaO	Na₂O	K ₂ O	Total	An	х
53.99	0.04	29.66	0.49	0.11	12.02	4.29	0.14	100.75	60.76	-177
53.94	0.02	29.69	0.63	0.09	12.19	4.43	0.15	101.13	60.33	-224
53.81	0.03	29.80	0.66	0.11	11.97	4.42	0.15	100.95	59.95	-41
52.30	0.04	30.67	0.67	0.08	13.06	3.69	0.12	100.62	66.17	-75
52.93	0.01	30.35	0.61	0.09	12.60	3.86	0.13	100.59	64.34	-307
53.63	0.03	29.77	0.65	0.05	12.12	4.21	0.14	100.60	61.40	-357
52.82	0.01	30.36	0.60	0.07	12.58	3.97	0.13	100.54	63.65	-619
52.63	0.05	30.69	0.53	0.06	12.90	3.86	0.11	100.82	64.87	-663
53.70	0.01	29.92	0.62	0.09	12.16	4.05	0.15	100.71	62.40	-883
55.48	0.02	28.82	0.47	0.02	10.73	5.01	0.20	100.74	54.20	-932
54.17	0.05	29.38	0.60	0.09	11.71	4.65	0.15	100.81	58.19	-1029
53.72	0.04	29.99	0.64	0.04	12.08	4.29	0.15	100.96	60.88	-1079
52.37	0.01	30.56	0.63	0.05	12.90	3.82	0.12	100.47	65.11	-1052
54.29	0.02	29.32	0.56	0.15	11.86	4.62	0.15	100.98	58.65	-1407
53.54	0.03	29.88	0.61	0.08	12.11	4.29	0.15	100.69	60.94	-1359

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SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	Total	Mg#	х
65.24	0.32	15.56	0.07	3.46	0.12	0.79	4.31	1.97	1.18	93.03	28.93	-36
64.64	0.39	15.77	0.05	3.50	0.16	0.80	4.53	2.28	1.29	93.49	28.95	-236
64.87	0.37	15.71	0.13	3.61	0.18	0.81	4.43	2.45	1.37	93.98	28.57	-397
64.99	0.36	15.63	0.05	3.52	0.12	0.80	4.35	2.39	1.31	93.51	28.83	-645
64.62	0.44	15.54	0.03	3.38	0.13	0.81	4.50	1.99	1.29	92.76	29.93	-693
63.81	0.56	16.09	0.00	3.48	0.12	0.80	4.60	2.22	1.28	92.95	29.07	-651
63.78	0.43	16.19	0.10	3.38	0.19	0.83	4.66	2.45	1.32	93.40	30.45	-389
64.22	0.44	15.70	0.04	3.47	0.14	0.81	4.56	2.42	1.31	93.11	29.38	-622
64.89	0.38	15.68	0.02	3.22	0.15	0.84	4.51	2.53	1.24	93.48	31.74	-82
63.93	0.49	15.85	0.25	3.46	0.17	0.80	4.61	2.39	1.30	93.24	29.19	-494
64.94	0.39	15.48	0.19	3.60	0.15	0.76	4.43	2.45	1.30	93.75	27.34	-956
64.59	0.45	15.72	0.00	3.61	0.14	0.78	4.47	2.09	1.28	93.17	27.81	-1057
64.89	0.34	15.72	0.00	3.53	0.15	0.82	4.36	3.08	1.37	94.25	29.28	-1032

Table S1-7. Electron microprobe data of melt in BAH11

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SiO ₂	TiO ₂	AI_2O_3	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	х
41.30	2.42	14.63	0.02	13.59	0.22	12.79	10.54	2.21	97.71	62.65	-395
40.73	2.34	15.23	0.01	13.79	0.23	12.65	10.27	2.34	97.58	62.05	-380
40.28	2.38	15.79	0.00	14.05	0.21	12.22	10.17	2.26	97.36	60.79	-385
39.10	2.30	17.89	0.01	13.63	0.21	11.76	10.26	2.25	97.42	60.60	-36
39.68	2.10	17.31	0.01	13.55	0.22	12.12	10.33	2.43	97.76	61.46	-127
41.53	1.80	16.41	0.02	7.95	0.14	16.36	10.59	2.46	97.26	78.58	194
43.10	0.54	15.75	0.12	7.25	0.16	17.35	10.88	2.24	97.40	81.01	436
44.33	0.54	13.94	0.26	7.28	0.15	18.16	10.47	2.05	97.18	81.64	701
42.40	1.59	16.12	0.05	7.75	0.14	17.04	10.13	2.52	97.75	79.67	956

Table S2-1. Electron microprobe data of amphibole in BAH13

 Table S2-2.
 Electron microprobe data of olivine in BAH13

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SiO ₂	TiO ₂	AI_2O_3	Cr_2O_3	FeO	MnO	MgO	CaO	Na ₂ O	NiO	Total	Mg#	Х
40.60	0.00	0.01	0.06	10.18	0.20	48.42	0.12	ND	NA	99.60	89.45	136
40.69	0.02	0.02	0.06	10.32	0.20	48.67	0.13	ND	NA	100.09	89.37	933
40.53	0.01	0.02	0.06	10.76	0.20	48.11	0.12	0.02	NA	99.84	88.85	684
40.49	ND	0.02	0.04	11.52	0.18	47.75	0.11	0.04	NA	100.15	88.08	508
40.59	0.01	0.03	0.05	10.55	0.21	48.33	0.13	0.01	NA	99.91	89.09	383
40.60	ND	0.02	0.05	10.65	0.18	48.31	0.14	ND	NA	99.94	88.99	266
40.58	0.01	0.01	0.05	10.43	0.20	48.45	0.13	ND	0.38	100.24	89.22	902
40.76	ND	ND	0.06	10.60	0.20	48.58	0.11	ND	0.40	100.72	89.09	705
40.67	ND	ND	0.06	10.38	0.19	48.85	0.12	ND	0.43	100.70	89.35	499
40.71	ND	0.02	0.07	10.78	0.19	48.23	0.13	ND	0.39	100.52	88.86	352
40.60	0.01	0.03	0.05	10.69	0.20	48.20	0.12	ND	0.44	100.35	88.93	258
40.74	ND	0.01	0.07	9.96	0.16	48.60	0.12	0.01	0.42	100.07	89.69	59

i8 1.67 0.03 100.30 83.49 -330 Core i3 1.77 0.04 100.26 83.23 -295 Core i4 1.62 0.08 100.32 83.19 -380 Core i5 1.63 0.05 100.12 83.44 -325 Core i1 1.16 ND 100.37 89.20 126 Core i0 1.16 0.02 100.21 89.29 145 Core i1 0.07 100.17 85.41 40 Rim i4 1.39 0.07 100.22 86.82 29 Rim			III D/ III J	pyroxene		oprobe at			Tuble 5
1.77 0.04 100.26 83.23 -295 Core 1.62 0.08 100.32 83.19 -380 Core 1.63 0.05 100.12 83.44 -325 Core 1.16 ND 100.37 89.20 126 Core 0.01 1.16 0.02 100.21 89.29 145 Core 0.02 100.25 89.42 125 Core 0.02 100.17 85.41 40 Rim 0.4 1.39 0.07 100.22 86.82 29 Rim	Na	CaO	MgO	MnO	FeO	Cr ₂ O ₃	Al ₂ O ₃	TiO ₂	SiO ₂
24 1.62 0.08 100.32 83.19 -380 Core 95 1.63 0.05 100.12 83.44 -325 Core 81 1.16 ND 100.37 89.20 126 Core 80 1.16 0.02 100.21 89.29 145 Core 82 1.15 0.02 100.25 89.42 125 Core 99 1.41 0.07 100.17 85.41 40 Rim 84 1.39 0.07 100.22 86.82 29 Rim	0.0	1.67	28.68	0.26	10.11	0.34	7.31	0.15	51.77
95 1.63 0.05 100.12 83.44 -325 Core 81 1.16 ND 100.37 89.20 126 Core 80 1.16 0.02 100.21 89.29 145 Core 82 1.15 0.02 100.25 89.42 125 Core 99 1.41 0.07 100.17 85.41 40 Rim 84 1.39 0.07 100.22 86.82 29 Rim	0.0	1.77	28.43	0.27	10.21	0.26	7.81	0.18	51.30
A11.16ND100.3789.20126CoreA01.160.02100.2189.29145CoreA21.150.02100.2589.42125CoreA91.410.07100.1785.4140RimA41.390.07100.2286.8229Rim	0.0	1.62	28.74	0.28	10.35	0.23	7.04	0.16	51.81
301.160.02100.2189.29145Core321.150.02100.2589.42125Core791.410.07100.1785.4140Rim441.390.07100.2286.8229Rim	0.0	1.63	28.95	0.26	10.24	0.20	6.78	0.16	51.85
321.150.02100.2589.42125Core791.410.07100.1785.4140Rim441.390.07100.2286.8229Rim	N	1.16	32.81	0.20	7.08	0.44	3.12	0.06	55.51
19 1.41 0.07 100.17 85.41 40 Rim 14 1.39 0.07 100.22 86.82 29 Rim	0.0	1.16	32.80	0.19	7.01	0.43	3.03	0.06	55.49
4 1.39 0.07 100.22 86.82 29 Rim	0.0	1.15	32.82	0.18	6.92	0.43	3.18	0.06	55.48
	0.0	1.41	29.79	0.25	9.07	0.87	6.19	0.13	52.39
0 1.64 0.06 99.84 85.88 8 Rim	0.0	1.39	31.44	0.22	8.51	0.52	3.08	0.08	54.91
	0.0	1.64	30.50	0.26	8.94	0.26	3.56	0.10	54.51
2 1.36 0.05 100.48 88.88 913 Core	0.0	1.36	31.82	0.16	7.10	0.79	5.65	0.07	53.47
7 1.31 0.01 100.60 87.15 911 Rim	0.0	1.31	30.87	0.17	8.11	0.52	7.11	0.11	52.40
1 1.19 0.04 100.07 88.62 580 Core	0.0	1.19	31.31	0.17	7.17	0.97	6.48	0.07	52.69
1 1.49 0.05 100.31 86.35 623 Rim	0.0	1.49	30.21	0.20	8.51	0.19	7.49	0.16	52.00
3 1.29 0.03 99.93 87.63 624 Rim	0.0	1.29	30.83	0.20	7.76	0.50	6.77	0.12	52.41
i6 1.18 0.02 99.90 88.70 624 Core	0.0	1.18	31.66	0.20	7.19	0.79	5.47	0.09	53.30

Table S2-3. Electron microprobe data of orthopyroxene in BAH13

 Table S2-4.
 Electron microprobe data of clinopyroxene in BAH13

SiO2	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	Х
50.22	0.34	6.30	0.13	7.58	0.27	16.93	17.44	0.38	99.59	79.93	-541
45.51	0.98	11.98	0.05	8.73	0.22	12.51	19.07	0.49	99.54	71.87	-387
46.33	0.67	10.94	0.03	8.67	0.24	13.19	18.70	0.51	99.28	73.06	-470
46.15	0.94	11.44	0.03	8.44	0.23	12.98	18.94	0.56	99.71	73.27	-395
45.27	0.91	12.21	ND	9.39	0.24	11.88	19.23	0.58	99.71	69.28	-395
51.31	0.32	4.51	0.07	7.62	0.22	17.23	17.88	0.30	99.45	80.12	-48
50.21	0.35	6.18	0.06	7.92	0.26	16.74	17.19	0.46	99.38	79.03	-12
45.66	0.81	11.60	0.03	8.70	0.19	12.38	19.47	0.55	99.39	71.72	-43
51.16	0.30	4.61	0.04	5.77	0.21	16.77	20.62	0.28	99.77	83.82	456
49.43	0.35	7.06	0.28	5.66	0.16	16.11	20.12	0.36	99.53	83.54	676
47.86	0.72	8.69	0.08	6.21	0.15	14.86	20.38	0.46	99.42	81.01	1001

			<u> </u>								
SiO ₂	TiO ₂	AI_2O_3	Cr_2O_3	FeO	MnO	MgO	CaO	Total	Mg#	Cr#	Х
0.20	0.36	26.85	30.22	26.05	0.11	13.20	0.03	96.28	47.45	43.02	36
0.14	0.31	28.57	28.03	25.55	0.14	13.81	0.08	95.88	49.06	39.70	151
0.12	0.29	39.15	18.65	24.15	0.10	16.32	0.08	97.93	54.64	24.22	250
0.13	0.28	26.70	30.88	25.65	0.12	13.38	0.17	96.58	48.19	43.68	446
0.10	0.28	23.39	34.41	26.36	0.16	12.52	0.15	96.72	45.84	49.67	561
0.09	0.22	21.14	37.97	25.76	0.16	12.25	0.09	97.04	45.87	54.65	848
0.97	0.24	46.74	12.18	22.41	0.10	17.29	0.16	99.06	57.89	14.89	928
0.35	0.26	29.74	27.51	25.76	0.12	14.10	0.09	97.15	49.38	38.29	950
0.12	0.22	38.47	20.06	23.23	0.13	15.78	0.11	97.22	54.77	25.92	900

Table S2-5. Electron microprobe data of spinel in BAH13

 Table S2-6.
 Electron microprobe data of plagioclase in BAH13

Table SZ	-o. Electro	on micropi	obe data	i oi piagic	clase in BA	NH 13				
SiO ₂	TiO ₂	AI_2O_3	FeO	MgO	CaO	Na ₂ O	K ₂ O	Total	An	Х
52.23	0.01	30.67	0.50	0.06	12.97	3.84	0.13	100.41	65.11	-96
55.42	0.01	28.68	0.44	0.03	10.65	5.23	0.15	100.60	52.95	-156
52.79	0.02	30.49	0.45	0.06	12.65	3.81	0.13	100.39	64.72	-421
53.19	0.04	30.11	0.46	0.04	12.31	4.27	0.12	100.55	61.44	-421
52.62	0.03	30.64	0.49	0.08	12.73	3.84	0.11	100.54	64.69	-593
52.12	0.04	30.69	0.44	0.07	13.15	3.96	0.11	100.58	64.73	-596
52.35	0.02	30.97	0.51	0.05	13.09	4.22	0.11	101.32	63.16	-604
53.52	0.02	29.74	0.44	0.08	12.00	4.31	0.12	100.24	60.61	-1213
52.19	0.02	30.58	0.42	0.03	12.83	4.01	0.10	100.18	63.87	-1364
54.94	0.04	29.07	0.49	0.03	11.19	5.14	0.16	101.06	54.61	-1470
52.66	0.01	30.54	0.48	0.08	12.71	3.89	0.11	100.48	64.36	-1596
53.02	0.04	30.14	0.42	0.08	12.41	3.99	0.13	100.22	63.22	-1233
52.66	0.03	30.54	0.46	0.07	12.60	3.87	0.11	100.34	64.28	-1073
52.65	0.04	30.49	0.44	0.05	12.85	3.94	0.11	100.58	64.32	-906
52.30	0.02	30.60	0.46	0.08	12.87	4.05	0.12	100.49	63.72	-708

	n Liceti		obe dutu	of ment in								
SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	Total	Mg#	х
63.07	0.35	16.59	0.00	2.70	0.13	1.02	4.57	2.18	1.17	91.83	40.24	-142
63.05	0.16	16.97	0.00	2.72	0.13	1.10	4.60	2.88	1.24	92.90	41.89	-82
62.15	0.40	16.88	0.00	3.17	0.10	1.06	4.81	2.53	1.18	92.28	37.35	-639
62.12	0.42	16.78	0.03	3.26	0.12	1.01	4.76	2.37	1.17	92.06	35.58	-805
61.24	0.46	17.06	0.00	3.43	0.13	1.04	4.95	2.55	1.21	92.07	35.09	-1118
61.67	0.44	17.34	0.00	3.35	0.12	1.04	5.01	2.75	1.21	92.94	35.62	-938
61.81	0.52	16.95	0.00	3.51	0.11	1.04	4.91	2.82	1.18	92.86	34.56	-935
61.22	0.55	17.13	0.00	3.38	0.14	1.03	4.90	2.39	1.17	91.91	35.20	-1148
61.45	0.46	17.18	0.23	3.42	0.12	1.01	4.96	2.59	1.19	92.60	34.49	-1307
62.49	0.34	16.64	0.15	3.28	0.17	1.03	4.75	2.60	1.19	92.64	35.89	-1593
61.91	0.56	16.98	0.16	3.31	0.13	0.99	4.85	2.53	1.13	92.59	34.77	-1399
62.65	0.32	16.36	0.06	3.32	0.14	0.99	4.65	2.57	1.18	92.24	34.71	-1566

 Table S2-7.
 Electron microprobe data of melt in BAH13

Table S2-8. Electron microprobe data of traverse on an orthopyroxene grain in BAH13 used in Figure S3

							17	0			0		
	SiO ₂	TiO ₂	AI_2O_3	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	NiO	Total	D
-	53.19	0.10	5.92	0.66	6.30	0.09	31.05	1.08	0.03	0.02	0.14	98.57	0
	53.12	0.05	6.06	0.78	6.61	0.24	31.02	1.19	0.03	0.02	0.12	99.24	16
	52.72	0.05	5.97	0.80	6.92	0.18	31.35	1.11	0.02	0.00	0.06	99.18	35
	53.11	0.11	6.06	0.74	6.50	0.11	30.92	1.02	0.01	0.03	0.13	98.73	42
	51.94	0.10	7.29	0.44	7.58	0.12	29.70	1.29	0.05	0.01	0.10	98.62	50
	50.66	0.10	8.77	0.34	8.38	0.13	28.98	1.61	0.00	0.00	0.14	99.10	65
	50.97	0.17	7.82	0.03	8.46	0.18	29.65	1.45	0.02	0.00	0.11	98.85	80
	51.36	0.26	7.59	0.05	8.48	0.13	29.96	1.04	0.00	0.01	0.15	99.03	90
_	50.73	0.28	8.70	0.00	9.64	0.32	28.81	0.92	0.02	0.00	0.10	99.52	100

TUDIC 32													
SiO2	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	Total	An	D	
52.93	0.00	29.26	0.01	0.48	0.00	0.06	13.01	4.12	0.15	100.01	63.58	0	
52.91	0.13	29.13	0.00	0.34	0.00	0.05	12.66	4.05	0.13	99.39	63.33	12	
52.37	0.13	29.26	0.05	0.38	0.06	0.04	12.91	4.00	0.13	99.31	64.06	24	
51.01	0.05	27.41	0.04	0.41	0.03	0.06	11.58	4.16	0.12	94.88	60.62	37	
54.17	0.00	28.35	0.00	0.48	0.02	0.07	12.19	4.50	0.14	99.92	59.97	44	
54.22	0.00	28.52	0.00	0.46	0.01	0.05	12.53	4.61	0.11	100.50	60.05	52	
53.80	0.00	28.72	0.00	0.40	0.00	0.05	12.46	4.32	0.15	99.90	61.44	63	
53.02	0.08	29.32	0.00	0.38	0.01	0.05	12.68	4.35	0.14	100.01	61.71	81	
53.80	0.00	29.30	0.04	0.36	0.01	0.05	12.74	4.15	0.11	100.55	62.89	90	
54.93	0.05	28.22	0.02	0.47	0.05	0.05	11.75	4.94	0.14	100.61	56.78	101	
55.92	0.03	27.62	0.00	0.42	0.00	0.01	11.12	5.34	0.20	100.67	53.50	121	

 Table S2-9.
 Electron microprobe data of traverse on a plagioclase grain in BAH13 used in Figure S5

Table 33														
SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	х			
41.11	1.82	17.54	0.14	10.34	0.18	13.70	10.76	2.66	98.24	70.25	-137			
40.46	2.57	17.61	0.15	10.09	0.15	14.17	10.40	2.74	98.33	71.46	-111			
41.79	0.78	18.01	0.09	9.29	0.18	14.92	10.37	2.41	97.85	74.11	92			
40.88	2.98	17.26	0.04	7.41	0.11	16.46	10.01	2.85	98.00	79.84	114			
41.39	1.81	16.78	0.13	10.44	0.18	13.87	10.76	2.47	97.83	70.31	-90			
40.86	2.05	17.41	0.06	10.08	0.15	14.27	10.58	2.66	98.11	71.62	-79			
41.42	1.61	17.55	0.01	9.28	0.17	14.71	10.49	2.48	97.71	73.86	96			
42.37	2.15	16.41	0.02	7.04	0.12	17.33	9.55	2.96	97.95	81.44	260			
41.67	2.53	16.72	0.03	6.88	0.14	17.17	9.89	2.83	97.84	81.65	391			
42.72	2.55	15.84	0.03	6.70	0.15	17.65	9.74	2.75	98.12	82.44	560			
43.18	0.86	16.44	0.40	6.15	0.10	18.12	10.21	2.78	98.25	84.01	981			
39.59	2.75	18.11	0.04	10.42	0.18	13.60	10.55	2.65	97.88	69.94	-371			
41.19	2.24	17.02	ND	10.21	0.17	14.26	10.40	2.69	98.17	71.34	-342			
39.94	2.70	18.21	0.02	9.59	0.13	14.84	9.84	2.73	98.00	73.39	-1067			
40.77	2.27	16.94	0.08	10.12	0.14	14.17	10.90	2.52	97.91	71.40	-1057			
39.96	2.80	17.96	0.01	9.32	0.14	14.65	10.49	2.78	98.11	73.70	-1260			
40.99	2.03	17.04	0.02	10.16	0.17	14.10	10.66	2.53	97.71	71.21	-1085			

 Table S3-1.
 Electron microprobe data of amphibole in MBH12

 Table S3-2.
 Electron microprobe data of olivine in MBH12

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SiO ₂	TiO ₂	Al ₂ O ₃	Cr_2O_3	FeO	MnO	MgO	CaO	Na ₂ O	NiO	Total	Mg#	Х
40.04	0.02	0.02	0.02	12.45	0.16	46.94	0.09	0.03	NA	99.77	87.05	122
40.24	0.01	0.03	0.03	11.35	0.20	47.67	0.15	ND	NA	99.68	88.22	95
40.34	0.01	0.03	0.08	11.61	0.19	47.71	0.15	ND	NA	100.13	87.99	259
40.42	ND	0.04	0.06	11.11	0.17	48.00	0.19	0.05	NA	100.04	88.51	374
40.46	0.01	0.02	0.07	10.62	0.15	48.50	0.16	0.01	NA	99.99	89.06	523
40.77	ND	0.02	0.06	9.76	0.17	49.31	0.15	0.01	NA	100.26	90.01	719
40.52	ND	0.04	0.06	10.87	0.17	48.40	0.16	ND	NA	100.22	88.81	677
40.65	0.01	0.01	0.06	9.85	0.18	49.21	0.14	0.03	NA	100.14	89.90	856
40.21	ND	0.01	0.04	11.82	0.14	47.50	0.15	0.02	0.32	100.22	87.75	116
40.35	0.02	ND	0.04	11.54	0.21	47.14	0.22	0.03	0.35	100.08	87.92	133
40.48	0.01	0.02	0.05	11.38	0.18	47.88	0.17	0.01	0.34	100.52	88.24	276
40.62	0.01	0.03	0.05	10.90	0.20	48.25	0.17	ND	0.33	100.55	88.75	370
40.67	ND	0.04	0.05	11.21	0.16	48.13	0.19	0.02	0.32	100.80	88.44	528
40.55	0.01	0.03	0.04	10.57	0.16	48.19	0.15	ND	0.35	100.06	89.04	709
40.96	0.01	0.03	0.07	9.73	0.19	49.41	0.15	0.01	0.33	100.89	90.05	715
40.93	0.01	0.02	0.05	9.40	0.16	49.68	0.11	ND	0.36	100.72	90.40	876

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SiO2	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	Х	Comment
52.47	0.22	7.04	0.55	7.85	0.17	30.21	1.89	0.04	100.43	87.28	34	
54.48	0.12	4.96	0.40	6.39	0.15	32.62	0.96	0.10	100.18	90.10	2	
50.91	0.34	8.89	0.02	9.16	0.19	29.07	1.69	0.06	100.35	84.98	37	
52.53	0.20	6.61	0.49	7.93	0.14	30.49	1.69	0.04	100.12	87.27	28	
51.74	0.32	8.21	0.06	7.88	0.19	29.89	1.65	0.03	99.97	87.12	695	
51.83	0.29	8.29	0.12	7.98	0.16	30.23	1.30	0.06	100.27	87.10	665	
51.21	0.26	9.70	0.10	7.67	0.17	29.81	1.54	0.05	100.50	87.39	856	
52.17	0.22	7.41	0.15	8.55	0.23	30.10	1.49	0.03	100.34	86.26	856	
49.44	0.46	10.07	0.02	11.57	0.23	26.85	1.81	0.06	100.52	80.53	-1067	In center part of amph
49.79	0.33	9.83	0.04	10.94	0.21	27.48	1.22	0.04	99.89	81.74	-1305	In center part of amph

 Table S3-3.
 Electron microprobe data of orthopyroxene in MBH12

 Table S3-4.
 Electron microprobe data of clinopyroxene in MBH12

SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	Total	Mg#	Х
51.09	0.29	5.23	1.03	4.37	0.12	17.93	18.98	0.50	99.53	87.97	-44
50.56	0.36	6.39	0.61	4.94	0.14	17.89	18.36	0.42	99.66	86.59	-7
50.86	0.39	6.30	0.39	4.89	0.14	17.69	18.56	0.43	99.64	86.57	-66
51.05	0.27	5.23	1.19	4.40	0.14	18.27	18.34	0.48	99.37	88.10	-5
50.81	0.20	4.51	1.00	4.78	0.16	17.21	20.43	0.24	99.34	86.52	266
53.36	0.12	3.25	0.84	5.07	0.18	22.69	13.92	0.31	99.74	88.86	408
53.92	0.10	3.11	0.72	5.02	0.18	22.85	14.01	0.27	100.18	89.03	560
49.93	0.28	7.63	1.25	4.66	0.14	17.35	18.36	0.49	100.08	86.91	697
49.91	0.23	7.83	1.04	4.48	0.12	17.24	18.66	0.47	99.98	87.28	891
50.53	0.46	6.55	0.20	5.41	0.16	17.56	18.31	0.38	99.56	85.26	-421
50.47	0.50	6.87	0.21	5.70	0.18	17.82	17.80	0.48	100.03	84.79	-1041
50.29	0.53	6.85	0.32	5.43	0.18	17.60	17.98	0.42	99.57	85.25	-1124
	51.09 50.56 50.86 51.05 50.81 53.36 53.92 49.93 49.91 50.53 50.47	51.090.2950.560.3650.860.3951.050.2750.810.2053.360.1253.920.1049.930.2849.910.2350.530.4650.470.50	51.090.295.2350.560.366.3950.860.396.3051.050.275.2350.810.204.5153.360.123.2553.920.103.1149.930.287.6349.910.237.8350.530.466.5550.470.506.87	51.090.295.231.0350.560.366.390.6150.860.396.300.3951.050.275.231.1950.810.204.511.0053.360.123.250.8453.920.103.110.7249.930.287.631.2549.910.237.831.0450.530.466.550.2050.470.506.870.21	51.090.295.231.034.3750.560.366.390.614.9450.860.396.300.394.8951.050.275.231.194.4050.810.204.511.004.7853.360.123.250.845.0753.920.103.110.725.0249.930.287.631.254.6649.910.237.831.044.4850.530.466.550.205.4150.470.506.870.215.70	51.090.295.231.034.370.1250.560.366.390.614.940.1450.860.396.300.394.890.1451.050.275.231.194.400.1450.810.204.511.004.780.1653.360.123.250.845.070.1853.920.103.110.725.020.1849.930.287.631.254.660.1449.910.237.831.044.480.1250.530.466.550.205.410.1650.470.506.870.215.700.18	51.090.295.231.034.370.1217.9350.560.366.390.614.940.1417.8950.860.396.300.394.890.1417.6951.050.275.231.194.400.1418.2750.810.204.511.004.780.1617.2153.360.123.250.845.070.1822.6953.920.103.110.725.020.1822.8549.930.287.631.254.660.1417.3549.910.237.831.044.480.1217.2450.530.466.550.205.410.1617.5650.470.506.870.215.700.1817.82	51.090.295.231.034.370.1217.9318.9850.560.366.390.614.940.1417.8918.3650.860.396.300.394.890.1417.6918.5651.050.275.231.194.400.1418.2718.3450.810.204.511.004.780.1617.2120.4353.360.123.250.845.070.1822.6913.9253.920.103.110.725.020.1822.8514.0149.930.287.631.254.660.1417.3518.3649.910.237.831.044.480.1217.2418.6650.530.466.550.205.410.1617.5618.3150.470.506.870.215.700.1817.8217.80	51.090.295.231.034.370.1217.9318.980.5050.560.366.390.614.940.1417.8918.360.4250.860.396.300.394.890.1417.6918.560.4351.050.275.231.194.400.1418.2718.340.4850.810.204.511.004.780.1617.2120.430.2453.360.123.250.845.070.1822.6913.920.3153.920.103.110.725.020.1822.8514.010.2749.930.287.631.254.660.1417.3518.360.4949.910.237.831.044.480.1217.2418.660.4750.530.466.550.205.410.1617.5618.310.3850.470.506.870.215.700.1817.8217.800.48	51.090.295.231.034.370.1217.9318.980.5099.5350.560.366.390.614.940.1417.8918.360.4299.6650.860.396.300.394.890.1417.6918.560.4399.6451.050.275.231.194.400.1418.2718.340.4899.3750.810.204.511.004.780.1617.2120.430.2499.3453.360.123.250.845.070.1822.6913.920.3199.7453.920.103.110.725.020.1822.8514.010.27100.1849.930.287.631.254.660.1417.3518.360.49100.0849.910.237.831.044.480.1217.2418.660.4799.9850.530.466.550.205.410.1617.5618.310.3899.5650.470.506.870.215.700.1817.8217.800.48100.03	51.090.295.231.034.370.1217.9318.980.5099.5387.9750.560.366.390.614.940.1417.8918.360.4299.6686.5950.860.396.300.394.890.1417.6918.560.4399.6486.5751.050.275.231.194.400.1418.2718.340.4899.3788.1050.810.204.511.004.780.1617.2120.430.2499.3486.5253.360.123.250.845.070.1822.6913.920.3199.7488.8653.920.103.110.725.020.1822.8514.010.27100.1889.0349.930.287.631.254.660.1417.3518.360.49100.0886.9149.910.237.831.044.480.1217.2418.660.4799.9887.2850.530.466.550.205.410.1617.5618.310.3899.5685.2650.470.506.870.215.700.1817.8217.800.48100.0384.79

Table														
SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Total	Mg#	Cr#	Х			
0.13	0.39	34.09	29.26	20.32	0.12	14.59	0.06	98.07	56.14	36.54	89			
0.13	0.33	26.34	36.90	20.93	0.15	13.80	0.18	97.98	54.02	48.45	300			
0.14	0.32	28.45	34.45	20.74	0.15	13.67	0.20	97.34	54.02	44.82	222			
0.14	0.24	26.99	36.11	20.55	0.13	13.40	0.13	96.92	53.76	47.30	392			
0.14	0.20	24.38	39.53	20.47	0.13	13.23	0.02	97.35	53.53	52.09	669			
0.13	0.20	23.17	41.91	19.61	0.16	13.05	0.14	97.65	54.26	54.82	835			
0.18	0.26	26.50	38.60	20.29	0.12	14.18	0.11	99.40	55.47	49.42	938			
0.18	0.19	31.63	33.06	18.32	0.10	14.99	0.04	97.59	59.33	41.21	1017			

Table S3-5. Electron microprobe data of spinel in MBH12

 Table S3-6.
 Electron microprobe data of plagioclase in MBH12

SiO ₂	TiO ₂	Al ₂ O ₃	FeO	MgO	CaO	Na₂O	K ₂ O	Total	An	Х
54.77	0.02	29.35	0.41	0.06	11.43	4.73	0.01	100.76	57.18	-206
53.53	0.01	30.46	0.37	0.03	12.38	4.39	0.01	101.19	60.91	-226
55.08	0.01	29.08	0.40	0.03	11.12	4.97	0.01	100.71	55.29	-283
54.85	0.01	29.34	0.40	0.06	11.46	4.91	0.01	101.04	56.33	-559
55.46	0.01	28.97	0.45	0.04	10.91	4.85	0.02	100.71	55.42	-730
54.74	0.02	29.24	0.36	0.02	11.24	4.86	0.01	100.49	56.10	-925
53.73	0.02	30.11	0.44	0.05	12.10	4.46	ND	100.91	59.99	-1011
52.51	0.02	30.62	0.41	0.07	12.87	3.83	0.01	100.34	65.00	-1211
53.13	0.04	30.22	0.41	0.07	12.36	4.35	0.02	100.60	61.09	-1250
52.18	0.01	30.82	0.40	0.04	13.03	3.94	0.01	100.43	64.63	-1258
52.88	0.04	30.66	0.36	0.09	12.63	3.88	0.01	100.55	64.27	-1485
52.30	0.03	30.93	0.31	0.09	13.06	3.81	0.01	100.55	65.45	-1485
53.52	0.03	30.30	0.40	0.04	12.44	4.24	ND	100.97	61.85	-1485
52.95	0.01	30.62	0.42	0.04	12.73	3.92	0.02	100.73	64.22	-276
54.63	0.02	29.26	0.47	0.05	11.45	4.84	0.01	100.71	56.66	-431

SiO ₂	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	Total	Mg#	х
60.32	0.06	18.37	ND	2.21	0.05	1.76	5.60	0.58	0.05	89.00	58.67	41
58.94	0.08	18.10	0.25	2.39	0.09	1.73	5.72	2.32	0.05	89.67	56.34	136
61.99	0.21	16.52	0.03	2.07	0.06	1.32	5.31	2.29	0.05	89.84	53.20	-27
62.27	0.28	17.09	ND	2.12	0.06	1.32	5.57	2.52	0.06	91.34	52.60	-80
62.08	0.26	16.73	ND	2.37	0.10	1.28	5.70	2.33	0.06	90.91	49.05	-285
62.18	0.27	16.82	ND	2.35	0.07	1.31	5.72	2.67	0.06	91.45	49.84	-378
62.01	0.30	16.92	ND	2.45	0.11	1.30	5.70	2.48	0.05	91.32	48.61	-593
62.10	0.29	16.92	ND	2.51	0.10	1.27	5.71	2.79	0.05	91.77	47.42	-716
61.97	0.24	17.08	ND	2.48	0.10	1.31	5.76	2.45	0.06	91.46	48.50	-810
61.91	0.26	17.12	0.14	2.34	0.10	1.30	5.69	2.40	0.06	91.33	49.76	-1005
61.47	0.34	16.77	0.02	2.78	0.08	1.25	5.81	2.43	0.06	91.05	44.49	-1105
61.97	0.30	16.56	0.07	2.44	0.02	1.26	5.74	2.25	0.06	90.74	47.93	-1329
61.37	0.33	16.67	ND	2.62	0.09	1.22	5.81	2.42	0.04	90.57	45.36	-1358
60.98	0.35	17.27	ND	2.82	0.08	1.26	5.96	2.33	0.05	91.11	44.33	-1510

 Table S3-7.
 Electron microprobe data of melt in MBH12

Table S3-8. Electron microprobe data of traverse on an amphibole grain with orthopyroxene in the center part in MBH12 used in Figure S4

SiO2	TiO ₂	Al ₂ O ₃	Cr ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	NiO	Total	Mg#	D	Mineral
50.54	0.31	9.72	0.06	10.95	0.14	26.78	1.27	0.06	0.03	0.04	99.90	81.34	0.00	Орх
40.05	2.57	16.70	0.07	9.13	0.11	14.26	10.37	1.37	0.02	0.05	94.70	73.57	38.83	Amph
40.25	2.54	16.72	0.00	8.59	0.12	14.44	10.60	1.43	0.04	0.00	94.72	74.97	50.70	Amph
40.01	2.59	17.05	0.00	8.69	0.13	14.44	10.16	1.39	0.02	0.00	94.24	74.43	62.10	Amph
40.41	2.49	16.72	0.04	8.38	0.13	14.29	10.59	1.37	0.01	0.00	94.44	75.25	71.84	Amph
40.03	2.32	16.50	0.03	8.50	0.15	14.15	10.35	1.41	0.01	0.03	93.48	74.80	81.61	Amph
40.95	1.95	16.22	0.02	9.54	0.00	14.13	10.02	1.26	0.01	0.04	94.14	72.54	97.42	Amph
41.64	1.34	16.08	0.03	10.60	0.13	13.68	10.62	1.13	0.00	0.00	95.25	69.71	114.79	Amph