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Revision 3 INTRINSIC CONDITIONS OF MAGMAS FROM THE LUNAR CRATER VOLCANIC FIELD (NEVADA): IMPLICATIONS FOR INTERNAL PLUMBING AND MAGMA ASCENT. Joaquín A. Cortés^{1,2*}, Eugene I. Smith³, Greg A. Valentine¹, Racheal Johnsen³, Christine Rasoazanamparany⁴, Elisabeth Widom⁴, Mai Sas^{3,5}, Dawn Ruth¹. ¹ Department of Geology, 411 Cooke Hall, Buffalo, NY 14260, University at Buffalo, State University of New York, USA. ² School of GeoSciences, The University of Edinburgh, Grant Institute, The King's Buildings, West Mains Road, Edinburgh EH9 3JW, UK. ³ Department of Geoscience, 4505 S. Maryland Parkway, Las Vegas, NV 89154, University of Nevada Las Vegas, USA. ⁴ Department of Geology, 114 Shideler Hall, Oxford, OH 45056, Miami University, USA. ⁵ Department of Geology, 516 High Street, Bellingham, WA 98225-9080, Western Washington University, USA. *Corresponding Author, caco@buffalo.edu American Mineralogist, special issue "Volcanic Rocks"

44 ABSTRACT

The northern part of the Lunar Crater Volcanic Field (central Nevada, USA) contains more than one hundred Quaternary basaltic cones and maars and related eruptive products. We focused on four informal units of different ages and locations in the field to test the compositional variability and magma ascent processes within the time span of an individual eruption and the variability between very closely spaced volcanoes with different ages. Based in whole-rock chemistry, mineral chemistry and the calculation of intrinsic properties (pressure, temperature and oxygen fugacity) we found that individual magma batches were generated in the asthenospheric mantle from a heterogeneous garnet lherzolite/olivine websterite source by ~3-5% partial melting. Each magma batch and temporary deep reservoir was a separate entity rather than part of a continuous long-lived reservoir. Magmas ascended relatively fast, stalled and crystallized in the uppermost several kilometres of the mantle near the base of the crust and also stalled at mid-crustal levels with minor or no geochemical interaction with surrounding rocks. Our data also suggest that volcanoes erupting within certain time windows have similar source characteristics and ascent processes whether they are located within a few hundred meters of each other or are separated by many kilometres.

Keywords: Lunar Crater Volcanic Field, monogenetic volcanism, whole-rock chemistry, mineral chemistry, geothermobarometry.

INTRODUCTION

Most of Earth's volcanism occurs at plate boundaries, in association with rifting or subduction, or in high magma-flux intraplate hotspots. However, substantial volcanism also occurs within continents and ocean basins, forming intraplate volcanic fields with relatively low magma fluxes and

commonly consisting of small-volume mafic volcanoes with alkaline affinities. Continental intraplate volcanic fields can consist of one to several hundred individual volcanoes, most of which are monogenetic (erupt in a single episode lasting weeks to years), and have lifespans of a few million years (Connor and Conway, 2000). Unlike "hotspot" systems, these fields tend to have a diffuse spatial distribution of eruptive centers and in many cases have no clear migration of eruptive activity with time. Volcanic landforms in intraplate fields are dominated by scoria cones, spatter/agglomerate ramparts, lava fields, maars, and tuff cones, in proportions that depend upon the relative dominance of explosive versus effusive activity and the local hydrologic environment (Valentine and Gregg, 2008; White and Ross, 2011; Brown and Valentine, 2013).

Petrologic studies of intraplate volcanism have tended to focus on "big picture" questions such as the nature of the mantle, or the broad evolution of regional magmatic activity over millions of years; however, recent studies have begun to explore the details of volcanic fields with much higher spatial and temporal resolution, including the complexities often recorded within individual monogenetic volcanoes. An advantage of a study of low magma-volume monogenetic volcanoes is that they may display compositional complexities that are closely related to mantle source characteristics, which may be totally overprinted by crustal reservoir processes in higher volume-flux, long-lived volcanoes. Geochemical and volcanological data from the Plio-Pleistocene Southwest Nevada Volcanic Field in the western U.S.A. for example, suggest that individual volcanoes tap domains of partial melt that are progressively decreasing in volume and degree of partial melting with time (Valentine and Perry, 2007). The domains are inferred to reflect local enrichments in volatile components within the lithospheric mantle source that has been subjected to repeated metasomatic events over ~1 Ga. Valentine and Perry (2006, 2007) suggested that the length scales of the partial melt domains are on the order of kilometers in size and the size of each volcano is proportional to its melt domain size. Brenna et al. (2012) showed how the Jeju Volcanic Field (South Korea) records deepening, increasing degree of melting, and increased melt volumes through time. Detailed studies of individual volcanoes within

the Jeju field indicate that each eruption tapped a domain of partial melt on the order of a few kilometers in size and each with a slightly unique composition that may vary within individual eruptions, reflecting mantle heterogeneity (Brenna et al., 2010). The Auckland Volcanic Field (New Zealand) shows similar behavior, with the volumes of individual volcanoes being correlated positively with the degree and spatial scale of partial melting in the asthenospheric source, but with large-volume melt batches interacting more extensively with lithospheric melts compared to smaller batches (McGee et al., 2013). Within the Auckland field, detailed data from the Motukorea volcanic center illustrates complex magma source history even within an individual monogenetic eruption that would commonly be assumed to be a "single batch." Other detailed studies support the perspective that despite their relatively small volumes (typically less than $\sim 1 \text{ km}^3$) and short lives; monogenetic volcanoes are accompanied by relatively complex processes related to magma sources, ascent, and temporary storage (e.g. Paricutín: Rowe et al., 2011; Erlund et al., 2009; Pioli et al., 2008; Luhr and Simkin, 1993; Jorullo: Johnson et al., 2008). Questions remain, however, as to whether the temporal and spatial scales of mantle source domains and the subsequent ascent and eruption of magmas have consistent relationships within and between volcanic fields.

In this work we focus on the products of four monogenetic volcanoes in the Lunar Crater Volcanic Field (central Nevada, USA) to explore two questions. (1) What are the compositional variability and magma ascent processes within the time span of an individual eruption? (2) What is the variability between very closely spaced volcanoes with different ages? Trace element data presented here suggest some degree of small-scale heterogeneity in the asthenospheric mantle, consistent with isotopic data (Rasoazanamparany et al., in review). Thermodynamic calculations of intrinsic variables show that parent melts ascended through two types of magma feeding systems: one in which magma ponded at or just below the base of the crust, and a second where magmas also ponded at mid-crustal levels. The volcanoes were fed by small magma batches that had essentially no chemical interaction with the crust, and ascended rapidly between storage levels and to the surface. In the cases studied,

two early eruptions tapped slightly variable mantle source rocks, and the two younger eruptions tapped a different source but one that was relatively homogeneous in space over a length scale of several kilometres and across a time span of several tens of thousands of years. Differences in composition and feeding systems between individual volcanoes seems to be more sensitive to time windows than to spatial variability in the source or crustal plumbing systems.

GEOLOGICAL SETTING AND PREVIOUS WORK

The Lunar Crater Volcanic Field (LCVF) lies at the northern end of a belt of scattered, Plio-Pleistocene basaltic fields that extends north-northeast from Death Valley (California) into central Nevada (Vaniman et al., 1982). The field is in the Basin and Range Province, a broad area of extensional tectonics, but is located centrally within the Province while most other young volcanic fields are found around its edge (e.g., Smith and Leudke, 1984; Brown et al. 2010; Gazel et al., 2012). The LCVF contains more than 100 individual cones with attendant lava fields, and four maars (Valentine et al., 2011; Hintz and Valentine, 2012; Valentine and Cortés, 2013), covering an area of 80 by 20 km extending from the Reveille Range to the Pancake Range (Figure 1).

Previous work in the LCVF has focused on geochemical and petrologic evolution of the field as a whole or of major parts of it, differently from our focus on more local scales (Scott, 1969; Scott and Trask, 1971; Bergman et al., 1981; Bergman, 1982; Lum, 1986; Kargel, 1986); see summary by Foland and Bergman (1992). Several authors used isotopic compositions of LCVF rocks along with samples from many volcanic fields in the western U.S.A. to define regional mantle isotopic domains (Lum et al., 1989; Menzies, 1989; Farmer et al., 1989). In general, isotopic, trace element, and xenolith geothermometry data (Smith, 2000) suggest that the LCVF magmas are sourced in the asthenosphere with characteristics of ocean island basalt (OIB) (Foland and Bergman, 1992), and with evidence of early contamination by crustal rocks (mainly carbonates beneath the early Pliocene Reveille Range

volcanoes; Yogodzinski et al., 1996) and by lithospheric mantle (potentially veinlets of intermediate composition in the younger Pancake Range volcanoes; Dickson, 1997).

FIELD DESCRIPTION, STRATIGRAPHY AND PETROGRAPHY

We focus on four different basaltic eruptive units (red area in Figures 1, 2), which are here informally named (from youngest to oldest): Marcath (a.k.a. Black Rock, with cosmogenic exposure age of 38 ± 10 ka; Shepard et al., 1995), Giggle Springs and Mizpah (40 Ar/ 39 Ar ages of $<81 \pm 5$ ka and 620-740 ka, respectively; Heizler, 2013) and Hi Desert basalts. Note that we infer the Giggle Springs age to be close to its 81 ka maximum based on the degree of lava surface modification. Field relationships indicate that Hi Desert basalt is older than Giggle Springs, but it is uncertain whether it predates or postdates the Mizpah eruption. The vents for the Giggle Springs, Mizpah, and Hi Desert basalts are located within \sim 500-600 m of each other, and allow us to assess variations between closely spaced, but different aged, volcanoes as well as potential variations within individual units. The vent for Marcath is \sim 6 km southeast of these three, allowing to test for variability in magmatic processes that might be related to time rather than co-location.

Hi Desert Basalt (Qhib, age undetermined)

The Hi Desert basalt (Figure 2) crops out as remnants of two small agglomerate cones, 10-20 m high and ~150 m diameter, that are surrounded and partly buried by later lavas and sediments. Hi Desert products are basalts to trachybasalts (Figure 3) notable in the field for the abundant euhedral plagioclase megacrysts, ~1-1.5 cm in size and rarely up to 5 cm. In thin section (neglecting megacrysts), the rock contains ~20% phenocrysts of subhedral olivine (2 and 3 mm in size), 5-10% of

phenocrysts of subhedral clinopyroxene (0.5 to 2 mm), and 10% euhedral plagioclase (1 to 2 mm), in a groundmass of plagioclase and ferromagnesian minerals.

Mizpah Basalt (Qmzb; 620-740 ka)

The Mizpah products form an elongate (~2.5 km long, ~800 m wide, 10-15 m thick) lava field with a subdued surface that extends westward from its small, 40 m-high scoria cone source (Figure 2). Mizpah rocks are trachybasalts (Figure 3) and typically porphyritic, containing < 10% subhedral olivine (0.1-0.2 mm) with sparse ~2 mm olivine phenocrysts; 5% subhedral to euhedral clinopyroxene (0.1-0.2 mm); and <10% euhedral plagioclase phenocrysts (0.3-0.6 mm) in some cases with sieve textures. The groundmass has a trachytic texture containing plagioclase and ferromagnesian minerals.

Giggle Springs basalt (Qgsb; $<81 \pm 5$ ka)

Giggle Springs products consist of two tephrite/basanite lava fields (5 km and 1.6 km long, 2-5 m thick; Figures 2, 3) that emanate from the ends of an agglomerate rampart (fissure vent), and partly overlie and partly surround Mizpah lava. The two Giggle Springs lava fields have moderately modified lava surfaces, where rafts of proximal agglomerate are preserved as mounds but areas between mounds have significant accumulations of eolian sediment and desert pavement. Lava levees and a drained lava channel are well preserved within a few hundred meters of the vent area. Megacrysts of olivine and clinopyroxene (up to ~2 cm) and plagioclase (up to ~1 cm) are abundant and ubiquitous in Giggle Springs products. Samples of the lavas contain (neglecting the megacrysts) <5% phenocrysts of subhedral olivine and clinopyroxene (0.1 - 0.2 mm) in an intergranular groundmass of plagioclase and ferromagnesian minerals.

Marcath/Black Rock Basalt (Qm; 38 ± 10 ka)

Marcath volcano consists of a ~150 m high, elongate scoria/agglomerate cone with a basal diameter of 900 × 500 m. The cone partly buries a >900 m-long agglomerate rampart that likely records an early fissure-fed phase of the eruption, and is open to the west where aa lavas flowed outward to form two compound lava fields as flows diverted around an older cone onto the adjacent valley floor (Figure 2). The 10-25 m-thick lava fields are 3.2 and 3 km long, and have maximum widths of 1.7 and 1.4 km, respectively. Tephra fall deposits extend south and northeast from the volcano, with thicknesses >4 m near the cone; the northeast lobe can be traced continuously up to 5 km from the vent (Johnson et al., 2014); local patches of ash suggest that it originally extended farther, but wind and water reworking have made it impossible to define the original outer edge of the deposit. The Marcath products are basanites to trachybasalts (Figure 3), ubiquitously containing centimeter-sized megacrysts of plagioclase, olivine, clinopyroxene and amphibole. Under the microscope the samples (neglecting megacrysts) are porphyritic with ~10% of phenocrysts of subhedral olivine (0.1 and 0.2 mm), and ~10% of euhedral (0.1-0.3 mm) plagioclase. The groundmass is microcrystalline with microlites of plagioclase and ferromagnesian minerals.

ANALYTICAL TECHNIQUES

A suite of 24 samples (Table 1) was collected from the four units that are the focus of the study for petrography, whole-rock chemistry, mineral chemistry and geochronology. Samples were also collected for isotopic analyses (Rasoazanamparany et al., in review.). Petrography was determined using the petrographic microscope combined with the study of back-scattered electron images taken with a Hitachi S-4000 scanning electron microscope (SEM) at the University of Buffalo, equipped with

an energy dispersive X-ray spectrometer to qualitatively determine the composition of the main mineral phases in the samples.

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Whole rock chemistry samples were chipped using a Bico Badger and powdered in a Bico Shatterbox for three minutes. Approximately 20 g of powdered samples were weighed in porcelain crucibles and placed in a furnace at 110 °C for two hours, cooled and weighed again, and then placed in a 1000 °C furnace for an additional 1.5 hours in order to measure the loss on ignition. For majorelement analyses with x-ray fluorescence spectrometry (XRF), sample powders were mixed with a 50-50 mix of Li-tetraborate and Li-metaborate, melted, and rapidly cooled to form glass disks. The glass disks were ground on 30-micron diamond abrasion pads to expose a fresh and flat analytical surface and then cleaned with isopropanol. Sample powders were prepared for trace element analysis on the XRF by mixing 3 g of binder and 12 g sample powder. Samples were then placed on a Buehler Specimen Mount Press and pressed to pellets. A Panalytical Axios wavelength dispersive XRF was used for both major and trace element analyses at University of Nevada Las Vegas (UNLV). Elements analyzed with the glass fusion disks include SiO₂, Al₂O₃, TiO₂, Fe₂O₃(t), MgO, Na₂O, K₂O, MnO, CaO, and P₂O₅, all reported in weight percent. Elements analyzed with the pressed pellets include Sc, V, Ni, Cu, Ga, Rb, Sr, Y, Zr, Nb, Ba, La, Hf, Pb, Th, and U, all reported in parts per million (ppm). The detection limits for most elements is <5 ppm, except La (30 ppm) and Nb (10 ppm).

Rare earth elements were analyzed at Activation Laboratories in Ancaster, Ontario, using a Litetraborate-Li-metaborate fusion method on an ICP-MS (see www.actlabs.com for analytical procedure and detection limits)

Main mineral phases were analysed using the JEOL JXA-8900 electron microprobe at UNLV and at the Cornell Center for Materials Research (CMMR) at Cornell University. Conditions of the analysis on both instruments were 15 kV current, a 5-µm beam for ferromagnesian phases, and a 10-µm beam for plagioclase. In the latter, sodium was analysed first in order to avoid loss in the counts of this element. Consistent with hand specimen and petrographic studies, important phenocrysts phases

are olivine, clinopyroxene, plagioclase and some amphibole, as well as gabbroic and ultramafic enclaves from the Marcath unit. Analyzable glass was not found in the groundmass of the samples. Melt inclusions were not detected in any mineral phases; therefore, calculations of the intrinsic conditions of the different systems are based on the equilibrium between mineral phases, rather than between mineral phases and glass (Putirka, 2008).

ANALYTICAL DATA AND RESULTS

Major and Trace Elements

Samples from the four volcanoes are tephrites/basanites and basalts to trachybasalts (Table 2, Figure 3) with SiO₂ varying from 43 to 50 wt. %, MgO from 6 to 12 wt. %, and CaO from 8 to 11 wt. %; this range occurs across the four volcanoes as well as within two of the individual volcanoes. Although the Hi Desert basalt seems to be slightly more evolved than the other units, overall no clear trends can be observed in a set of element-versus-MgO variation diagrams (Figure 4).

Rare-earth elements compared to OIB basalts strongly suggest an OIB source for the melts (Table 2, Figures 5, 6; Sun and McDonough, 1989). Slight differences in the slope of the REE plots (Figure 5) are well illustrated in a plot of La/Yb vs. Zr/Nb (Figure 7a); Marcath and Giggle Springs units are similar and relatively tightly clustered, while Hi Desert and Mizpah have broader, overlapping fields. Consistent with the main crystallizing phases (olivine, orthopyroxene, clinopyroxene), we have selected Th and Ta as incompatible elements and Cr as compatible (Table 4a) to test for degree of partial melting and further mineral fractionation of these melts. In a plot of Th/Ta vs. Cr/Ta (Figure 7a), no significant differences in the degree of fractionation (Th/Ta) of the source are observed although the figure suggests a variable degree of partial melting (Cr/Ta) in the units (Ta is slightly more incompatible than Th; Tables 4a,b). In a plot of Dy/Yb vs La/Yb (Figure 7b) to test for garnet or spinel in the source,

no clear differences in the ratio Dy/Yb are observed (~0.08 to 0.18). Following Barth et al., (2000), we have also plotted the ratio La/Nb vs Nb [ppm] (Figure 8a) and the ratio Nb/Ta vs Nb [ppm] (Figure 8b) to test for crustal contamination. Overall, the samples cluster close to the chondritic ratio although the amount of Nb is several orders of magnitude higher than the sources proposed by Barth et al. (2000).

Mineral Chemistry

Olivine phenocrysts occur in all the units, both as phenocrysts and in the groundmass, and have a range of compositions from Fo₅₇ to Fo₈₇ for Hi Desert, Fo₆₃ and Fo₈₇ for Mizpah, Fo₆₇ and Fo₈₇ for Giggle Springs and for Marcath (Table 3a, Figure 9a), although intermediate-composition olivine is less abundant in Marcath units.

Clinopyroxene phenocrysts and groundmass crystals are generally less common than the olivine phase, but occur in all samples. The main pyroxene phase is diopside in Hi Desert and Mizpah samples, and both diopside and augite (based on the classification of Morimoto, 1989) in Giggle Springs and Marcath. The fact that several of the analyses plot above the diopside limit (Table 3b, Figure 9b) is due to a relatively high Al³⁺ content in their formula, as they contain amounts of the jadeite end-member (NaAlSi₂O₆), equivalent to a subtle omphacitic component in clinopyroxenes from Hi Desert, Giggle Springs, and Marcath. This suggests relatively higher-pressure crystallization conditions compared to a normal Ca-Mg-Fe²⁺ clinopyroxene (Morimoto, 1989) and requires the charge balance proposed by Papike et al., (1974) and Cameron and Papike (1981) to estimate iron speciation in the pyroxene formula.

Plagioclase (Table 3c, Figure 9c) has a relatively narrow range of labradoritic compositions in Mizpah, Giggle Springs, and Marcath rocks (An₆₁₋₆₈, An₆₀₋₆₉, and An₅₇₋₇₂, respectively), with a few phenocrysts of an andesine-oligoclase-like phase (~An₃₀). A similar andesine-oligoclase phase was observed in Hi Desert, however plagioclase phenocrysts range in composition between An₄₈ and An₇₁;

a wider range than found in the other three units.

Amphibole was found exclusively in products from Marcath volcano as a phenocryst phase and also as the main phase of <4-cm mafic inclusions that are common around the Marcath edifice and in the proximal areas of Marcath lava flows (Figure 2). Marcath amphibole compositions vary from pargasite to magnesiohastingite and kaersutite (Table 3d, Figure 9d) based on the Leake et al. (1997) cation allocation scheme and classification.

Intrinsic conditions of crystallization

The methods used here to calculate pressure, temperature and oxygen fugacity are based on mineral equilibria of the main crystallizing phases. There are two fundamental aspects considered in these calculations: (1) whether the pairs of mineral phases are in thermodynamic equilibrium (a non-trivial issue in volcanic rocks), and (2) how to determine one value of a given intrinsic condition, based on the calculation of such a value from a data set that is often composed of hundreds microprobe analyses.

Thermometry. Crystallization temperature was estimated using an olivine – clinopyroxene thermometer. There are several formulations of this equilibrium, originally developed by Powell and Powell (1974), all of them based of the interchange of Mg-Fe between olivine and augite:

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$$Mg_2SiO_4 + CaFeSi_2O_6 = Fe_2SiO_4 + CaMgSi_2O_6$$
 (1)
315 ol cpx ol cpx

One of its current formulations was proposed by Loucks (1996), wherein the equilibrium constant of Equation (1) depends on temperature through a non-linear relationship that can be solved with a

numerical method. In order to assess whether the two mineral phases are in equilibrium we rely on the equilibrium window proposed by Cortés et al. (2005). In their model, since the equilibrium olivinemelt can be equated with the equilibrium clinopyroxene-melt assuming the same melt, the value of the equilibrium constant of the reaction (Equation 1) is constrained if the mineral phases are in equilibrium. According to Cortés et al. (2005) the equilibrium constant, expressed as iron-magnesium ratios, varies between 0.64 and 1.88 with a mean of 1.22. We also rely on the petrography of the samples and assume that equilibrium is attained between two mineral phases in physical contact, although the calculation is not performed on analyses on the interface between the two grains, to avoid Ostwald ripening and elemental diffusion issues between the involved phases.

The composition of each mineral phase in Equation 1, using a large number of analyses, should be normally distributed due to the inherent randomness in crystallization. Because expressing the free energy at equilibrium *implies the calculation of a ratio* between two random variables in which each follow a normal distribution (e.g. the chemical analyses of the mineral phases), the result of the calculation will be a random variable that follows a Cauchy-Lorentz distribution. Since the expected value and variance of a Cauchy-Lorentz distribution are undefined, the best estimation of a measure of central tendency is the median while half the sample interquartile range (HIQR) is a robust estimator of the spread of the statistical sample (e.g. all the calculations performed). The procedure is then used to first calculate all the possible outcomes of temperature for mineral pairs that are assumed in equilibrium, and then to select the median and HIQR of the calculated temperatures. Resulting crystallization temperatures for Hi Desert and Mizpah rocks are between 1050-1100°C, compared to 900-1050°C for the younger Giggle Springs and Marcath units.

Pressure. The lack of glass also precludes crystallization geobarometric calculations based upon melt-mineral phase equilibrium (e.g., clinopyroxene-melt; Putirka, 2008). Nimis and Ulmer

(1998) and Nimis (1999) suggest an alternative geobarometer that is based on the structural lattice of clinopyroxene formed in equilibrium with a basaltic melt, such that its composition is not explicitly needed. This geobarometer is relatively insensitive to temperature in alkaline compositions such as those studied here. Because the calculation is performed directly from the composition of clinopyroxene, it is desirable that the distribution of pressures follows a normal distribution if there is one population of clinopyroxene. With this condition, it is straightforward to calculate the mean and standard deviation of the pressure values. Calculated Hi Desert and Mizpah crystallization pressures range between 0.9-1.3 GPa (~30-45 km). The younger units (Giggle Springs and Marcath) have bimodal crystallization pressures with values between 1-1.3 GPa and between 0.3-0.6 GPa (~35-45 km and 10-20 km, respectively).

Empirical observations of amphiboles from calc-alkaline intrusions (Hammarstron and Zen, 1986) show that the amount of total aluminium in amphiboles is proportional to pressure of crystallization (see Ridolfi et al., 2010, for a summary of the different calibrations). Recently, Ridolfi and Renzulli (2012) and Simakin et al. (2012a) have extended the original calibration to mafic products and mafic enclaves, however more work remains to be done in order to have a robust geobarometer based on this mineral phase. The accuracy of the Ridolfi et al. (2010) geobarometer is \pm 0.3 GPa (Simakin, pers. comm.) while the Simakin et al., (2012a) model has an accuracy of \pm 0.05 GPa. We applied the Ridolfi and Renzulli (2012) model to a set of amphibole phenocrysts found in mafic inclusions within the Marcath lava flow (Table 3d). According to this model, crystallization pressures vary between 0.69-0.75 GPa, overlapping the lower range of values of the clinopyroxene-based crystallization pressures described above. On the other hand, the Simakin et al., (2012a) model did not produce meaningful results in our analyses (negative values), hence we favour the Ridolfi and Renzuli (2012) model, conscious that it is an over-estimation of the pressure with an error of ~25%. The Ridolfi and Renzulli (2012) model also provides temperature estimates between 1037-1070 °C (consistent with the olivine-clinopyroxene-based crystallization temperatures stated above, taking into account the large

error in the calculation), oxygen fugacity between ΔNNO -0.2 and ΔNNO +0.5, and estimated water content in the melt between 5-7 wt%. This range of water content is comparable to those estimated for trachybasalts in the Southwest Nevada Volcanic Field, the nearest Quaternary volcanic field, ~200 km to the south (Nicholis and Rutherford, 2004), and with the higher part of the water content range determined for the Big Pine field, ~250 km west-southwest of LCVF (Gazel et al., 2012).

Oxygen fugacity. Oxygen fugacity estimates are based on the speciation of iron in the clinopyroxene unit formula (Cortés et al., 2006, Simakin et al., 2012b). If the speciation of iron in the clinopyroxene is not known (as it is always the case from a typical electron microprobe analysis), such speciation is calculated based on the charge balance proposed by Papike et al., (1974) and Cameron and Papike (1981). The calculation must also include the contribution of elements such as Na, Ti, Fe³⁺ and Cr, which are not considered in the pyroxene quadrilateral. The intrinsic oxygen fugacity of the system is then estimated assuming equilibrium between the clinopyroxene formula and the melt, although the composition of the melt is not directly needed for the calculation (Cortés et al., 2006). If the temperature is calculated using an independent method, it is possible to additionally determine the absolute oxygen fugacity of the system. As with the barometry described above, one population of clinopyroxene phenocrysts ought to produce values of intrinsic oxygen fugacity that are normally distributed. Cortés et al. (2006) estimated that this approach has a sensitivity of ± 1 QFM unit.

Oxygen fugacity calculations using both Cortés et al. (2006) and Simakin et al. (2012b) methods give comparable values of the oxidation state in the samples. For the older units (Hi Desert and Mizpah) intrinsic oxygen fugacity relative to QFM buffer is estimated between Δ NNO+2 and Δ NNO+4. For the younger magmas (Giggle Springs and Marcath) are relatively reduced with values of intrinsic fugacity between Δ NNO+1 and Δ NNO+2, comparable with amphibole oxygen geobarometry results.

Depth of partial melting and origin of the basaltic melts. Depths of partial melting were estimated using two approaches: a silica-melt barometer (Lee et al., 2009), and an orthopyroxene-liquid geobarometer (Putirka, 2008); both calculations are performed using the whole-rock composition of the basaltic products as a proxy of the equilibrium melt. The Lee et al. (2009) technique applies a silica-based barometer that is based solely on the activity of silica in the melt. As a result, it is more reliable and less sensitive to incompatible element variability and mantle composition than earlier barometers (e.g. Wang et al., 2001). The Lee et al. (2009) barometer was calibrated using 433 basalt compositions in equilibrium with olivine and orthopyroxene over a range of temperatures from 1110 to 1800 °C and pressures of 1 atmosphere to 7 GPa. Barometer calibration yielded an uncertainty of \pm 0.20 GPa (Lee et al., 2009). The thermometer generated is consistent with the barometer and has an uncertainty of \pm 3% (Lee et al., 2009). When using this barometer, it is critical that the primary magma contains both olivine and pyroxene. Further, the barometer is not calibrated for silica-undersaturated rocks with < 40 wt. % SiO₂, and it is best to choose the most primitive basalt with MgO > 8.0 wt.% for pressure and temperature calculations.

The orthopyroxene-barometer (Putirka, 2008) is calibrated for basalts with SiO₂ contents as low as 35 wt. %, will work for basalt with olivine and both clinopyroxene and orthopyroxene, and is independent of source composition (as long as pyroxene and olivine are in equilibrium with the source at the time of partial melting). Using clinopyroxene + olivine results in almost no increased error relative to olivine + orthopyroxene (Putirka et al., 2012).

Using the method of Lee et al. (2009) in the most primitive whole-rock chemical analyses from our dataset produces temperatures of partial melting of the mantle of the order of 1500-1700°C with pressures between 3-6 GPa, which, for an average density of ~3000 kg/m³, implies depths of the partial melting between ~100-200 km. The Putirka (2008) calibration indicates that partial melting occurred at temperatures ranging from 1460 to 1650 °C and pressures between 1.7-3.8 GPa, which implies depths of ~60-130 km, consistent with the depth of the asthenospheric mantle in the region (e.g. Fisher et al.,

2010; Kumar et al., 2012).

An important caveat for the Lee et al. (2009) approach is that to apply the model, the source rock is assumed to be a peridotite. In order to assess this condition for the magmas of interest here, we estimate the mineralogy of the mantle sources based on olivine mineralogy (Sobolev et al., 2000). In Figure 10a, we plotted Fe²⁺/Mn in olivine vs. %Fo, while in Figure 10b we plotted 100xCa/Fe²⁺ vs 100xMn/Fe²⁺ for the olivine analyses with %Fo > 85, following Sobolev et al. (2007). Based on these plots, we conclude that the most likely source for these magmas has a modal mineralogy that lies between the pyroxenite and peridotite (lherzolite) fields as defined by Sobolev et al. (2007), although nickel content in olivine reported by Sobolev et al., (2007) is higher compared with LCVF olivine. Source identification based on olivine composition is consistent with the approach of Hirschmann (2000) and Petermann and Hirschmann (2003) based on the whole-rock composition, because FeO/MnO_(melt) is proportional to the ratio Fe²⁺/Mn_(ol). Thus because the Lee et al. (2009) method assumes a peridotite source, Putirka's (2008) method, which can be applied to a variety of sources as long as they are olivine and pyroxene saturated is favored here.

MELTS modelling. Geothermometry and geobarometry calculations were compared with the MELTS and pMELTS thermodynamic models (Ghiorso and Sack, 1995; Asimow and Ghiorso, 1998; Ghiorso et al., 2002) as a preliminary, but rough validation, since the LCVF conditions are indeed beyond the working limits defined by Ghiorso et al. (2002). We specifically aimed for a preliminary estimation of liquidus temperatures at the calculated pressures by the silica activity barometer, and at the crystallization pressures calculated using the clinopyroxene barometer described above. We also used these thermodynamic models to estimate the potential crystallizing mineralogy to compare with products of the four volcanoes. No calculations of the liquid line of descent were performed because evolved magmas are not present at Lunar Crater. Instead, compositions cluster around SiO₂ ~44-46 wt%.

Liquidus temperatures at the pressures of interest were calculated for the most primitive sample of our dataset (sample LC10-20; $SiO_2 = 44.02$ wt%; Table 5), as well as, for comparison, for a reference pyroxenite sample (Table 2) from the Earth Reference database (http://earthref.org); oxygen fugacity $\Delta QFM+2$ was assumed in both cases in order to constrain the iron speciation in the samples, and 2% of H_2O was assumed in sample LC10-20 consistent with a conservative "wet" liquidus. The temperatures predicted using the Lee et al. (2009) or Putirka (2008) models are comparable to pMELTS-calculated liquidus temperatures (1584°C-1664°C) for both the basalt and pyroxenite given estimated partial melting pressures not higher than 3.8 GPa. The pMELTS model also predicts that the crystallizing phases at 35-45 km depth (equivalent to the crystallization pressures calculated above for Mizpah and Hi Desert, and the deeper of two crystallization pressures that were calculated for Giggle Springs and Marcath) are plagioclase, olivine, clinopyroxene, and orthopyroxene, as well as amphibole and biotite. Except for biotite, all these phases were observed in the volcanic products.

Crystal Size Distribution (CSD)

Crystal size distributions (CSD; Marsh, 1988, 1998; Higgins, 2006) of olivine, clinopyroxene, and plagioclase were measured from back-scattered electron images of samples from the Mizpah, Giggle Spring and Marcath units. Note that olivine CSD was not measured for Marcath products because olivine is rare in these units (Figure 10a). Images were imported into ImageJ freeware (http://rsb.info.nih.gov/ij/) in which long and short axes of the different mineral phases were measured; a total of 200 measurements were performed on each mineral phase in order to have a representative sample of each CSD (Mock and Jerram, 2005; Morgan and Jerram, 2006). Mean crystal aspect ratios were determined using the CSDSlice method (Morgan and Jerram, 2006), fabric was considered massive (e.g. with no foliation), and measurements were not corrected for crystal roundness. All these values as well as the measurements were exported into CSDCorrections 1.4 (Higgins, 2006) in order to

calculate the crystal size distribution of each mineral phase. The total slide area, volumetric phase abundance and vesicularity of the samples were determined using built-in features of ImageJ and estimated from the petrography of the samples.

Proximal products of Giggle Springs volcano have smaller sizes than the more distal lava, and there is a partial overlap of the CSDs of the main phenocryst phases although groundmass CSDs are different. The CSD of the olivine phase in Giggle Springs and Mizpah has an inflexion point in the distribution at approximately 0.1 mm for some samples (Figure 11a); olivine phenocrysts in both units have similar distributions except at the smallest sizes. The slope of the CSD of olivine phenocrysts is \sim -16 [mm⁻⁴] (r² \sim 0.96) and \sim -58 [mm⁻⁴] (r² \sim 0.95) for groundmass microlites (cut off at 0.1 mm), suggesting residence time of few hours for the phenocrysts using growth rates of 0.2-0.4 mm per day reported for olivines in Hawaiian tholeiites (Maaløe, 2011).

The clinopyroxene CSDs (Figure 11b) are similar between samples from Giggle Springs and Mizpah, but different from Marcath samples, which has consistently larger crystals. The plagioclase CSDs for the different units agree in an intermediate range of sizes between ~0.1 and ~0.3. While this is the upper range for Giggle Springs and Mizpah, it is the lower range for the samples from Marcath (Figure 11c). Oxides (Figure 11d) are substantially different in all the samples (even those from the same unit), suggesting late crystallization events.

486 DISCUSSION

Melting depths and source composition

Independent of the age and location of the four studied volcanoes, the geochemical data strongly indicate sources located in the asthenosphere, at depths between 60 and 130 km based on the Putirka (2008) orthopyroxene barometer. Considering Fe²⁺/Mn and Ca/Mg in olivine phenocrysts with %Fo > 85, the source material is likely to have a modal mineralogy between pyroxenite and peridotite (Petermann and Hirschmann, 2003; Tuff et al., 2005; Sobolev et al., 2007; Herzberg, 2011). The most

likely source is an olivine websterite, although the nickel content in the LCVF olivine is lower (less than 2000 ppm) than the reported values by Sobolev et al., (2007) for pyroxenite or peridotite (between 2000 and 4000 ppm). The lower concentration of nickel in olivine suggests a low-Ni source (Herzberg, et al., 2013), although Ni content in whole-rock analyses is within reasonable values (120 to 330 ppm). A possible explanation for this difference might be related to the oxidation state of the magmas. Oxidation state of the depleted and enriched mantle is thought to be close to QFM and NNO+1 respectively (Balhaus, 1993). Our estimation of the oxygen fugacity by the clinopyroxene and amphibole formulas seems to indicate more oxidizing conditions for these basalts thus implying changes in oxygen fugacity during ascent and the main crystallization event. The more oxidizing conditions might have affected Ni³⁺ concentrations in the basaltic melt, a species that cannot enter in the olivine formula due to its charge, although the specific relation of the oxygen fugacity and the speciation of nickel is not yet completely understood. A plausible explanation for the more oxidizing conditions during crystallization is likely to be related to the water content of these magmas. The presence of amphibole seems to indicate a relatively high amount of water, which, depending on its speciation, has been linked to changes in the oxidation state of melts (e.g. Barker and Rutherford, 1996).

Rare-earth element ratios are helpful for estimating the depth of partial melting as well as the mineralogy and chemistry of the source, because of the different solid/melt distributions of REE in spinel vs. garnet bearing peridotite/pyroxenite. Heavy rare-earth elements like Yb are highly compatible in garnet while light-rare earth elements like La are incompatible; therefore partial melting of peridotite or pyroxenite containing garnet will produce melts with high La/Yb. Variable degrees of partial melting will result in changes in La/Yb such that the higher degree of melting, the lower La/Yb. Further, middle-rare earth element (MREE)-heavy rare earth element (HREE) ratios are sensitive to whether garnet exists as a residual phase in the source, because HREEs like Yb are retained by garnet during a melting event. This produces large changes in MREE/HREE in magmas generated by melting

of a garnet-bearing source, a change that is especially prominent with larger degrees of partial melting. Garnet is a relatively refractory mineral and does not significantly contribute to the melt phase until larger degrees of melting. At small melt fractions, melting of olivine or pyroxene results in little to no change in in MREE/HREE (Dy/Tb) because of similar partition coefficients for Dy and Tb in olivine and pyroxene (e.g., D₀Dy/D₀Yb=0.75 to 1.5 for clinopyroxene; Chazot et al, 1996). This results in melting trajectories that are relatively flat (or with slightly decreasing Dy/Yb) until about 30% melting. For higher degrees of melting, Dy/Yb decreases rapidly as garnet is melted. In the case of the Lunar Crater volcanoes La/Yb varies from 3 to 15 with the higher ratios suggesting a garnet source (Figures 7a,c, 12a,b). The mode of a typical garnet lherzolite is 60% olivine, 20% orthopyroxene, 8% clinopyroxene and 12% garnet (Thirwall et al, 1994). This source produces melts that are too high in Dy/Yb to match Lunar Crater magmas (Figure 12a). If, however, garnet is not as abundant in the source (5 to 8%), modelled melts provide a much better match to the sample set. We also modelled the melting of a spinel lherzolite with 57.8% olivine, 27% orthopyroxene, 11.9% clinopyroxene and 3.3% spinel. Although this source produced melts with relatively constant Dy/Yb, the value of the ratio was much to low to match the dataset. Based on Figure 10b and following Sobolev et al., (2007), we have also modelled the source as an olivine websterite (i.e. a composition between lherzolite and pyroxenite) with 30-35% olivine, 30-40% orthopyroxene, 20-30% clinopyroxene, and 5-11% garnet (Table 4b). Results using this source are similar to the lherzolite models (Figure 12b) except that the final melt has a lower Dy/Yb. Our conclusion is that the source of Lunar Crater magma has a composition between a garnet lherzolite and an olivine websterite containing 5 to 8% garnet. Consistency between this result and that of the independent method based upon olivine Fe²⁺/Mn and 100xCa/Mg ratios reinforces this interpretation. Note that we did not calculate major element models involving partial melting because of the sensitivity of melt chemistry to source composition. Meaningful major element models would require knowledge of major element chemistry of the source. This information is at present not known well enough for Lunar Crater.

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Magma evolution

The variation of Th/Ta is used here to determine the degree of fractionation because both elements are incompatible, however Ta is more incompatible than Th (Table 4b) while Cr/Ta varies with different degrees of partial melting in a source containing an olivine websterite component (Cr is highly compatible in both ortho- and clinopyroxene). The narrow range of Th/Ta values relative to Cr/Ta (Figure 7b) indicates variable differences in the degree of partial melting within units, while the trace element variability (e.g., Figure 6) is consistent with small variations in source composition. On the other hand, there is little evidence for systematic fractionation processes in the resulting magma batches as evidenced by nearly constant Th/Ta ratios (Figure 7b).

In order to generate the basaltic compositions similar to those reported in Table 2, the required amount of partial melting of a olivine websterite/lherzolite source is ~3-5% based on the batch model calculations presented in Figures 12a,b.

Source Heterogeneity

Source heterogeneity is evident between different, but closely spaced, volcanoes (e.g., Giggle Springs compared to Hi Desert; Figure 7a) and within individual volcanoes (e.g., Mizpah, and possibly Hi Desert). The spatial scale of subtle mantle variability must be similar to the scale of the partial melt domains tapped by the older Hi Desert and Mizpah feeding systems. These observations suggest lateral length scales of ~500 m or less for compositional heterogeneity in the source, if the magmas were sourced at similar depths. In contrast, the two younger volcanoes (Giggle Springs and Marcath) had essentially identical sources based upon the data presented here, even though the volcanoes are ~6 km apart and have age differences of several tens of thousands of years; furthermore, their sources were homogeneous on the scale of the individually tapped melt domains. We realize that melts can travel laterally from source to surface, implying that in some cases, source areas may not lie directly below

vents. If this is the case for the northern Lunar Crater volcanoes, our stated spatial scale of source heterogeneity should be regarded as a minimum estimate. Figures 8a,b suggest that although the samples are close to chondritic Nb/Ta and La/Nb, Nb [ppm] is several order of magnitude higher than the mantle or the upper crust and is especially enriched in the younger units (up to 90 ppm in Marcath). This result strongly suggests the addition of an additional crustal (oceanic) component. Isotopic data (Sr Nd Pb, Hf and Os) also indicate a heterogeneous source over very small spatial scales, likely related to ancient subduction processes and consistent with our interpretation (Rasoazanamparany et al. in review).

Magma Ascent and lithospheric contamination

Crystallization occurred at depths of 30-45 km beneath the four volcanoes, with additional crystallization at 10-20 km for the two younger units. Crustal thickness in the area is estimated at 30-35 km (Gilbert and Sheehan, 2004). Thus, enriched melts generated in the asthenospheric mantle travelled relatively unimpeded to near or just beneath the base of the crust, where all four magma batches ponded and underwent up to 50% crystallization as determined from MELTS modelling. The intrinsic variables and the lack of evidence for crustal contamination (Rasoazanamparany et al., in review) indicate that magmas feeding the two older volcanoes, Mizpah and Giggle Springs, ascended rapidly from near the base of the crust to the surface without intermediate ponding.

In contrast, the younger Giggle Springs and Marcath magmas apparently stalled at mid-crustal depths and underwent further crystallization. Fundamental differences in the CSD of their main mineral phases suggest that their mid-crustal reservoirs were not connected to each other, which is also consistent with the likely age difference between the two (38±10 ka and 81±5 ka for Marcath and Giggle Springs, ~620-740 ka for Mizpah). Amphibole crystallized in Marcath's crustal reservoir, and is present in the eruptive products as megacrysts. This phase has only been observed at Marcath, implying either that: (1) there was a slight difference between the crystallization conditions of Marcath and

Giggle Springs such as higher water content in the former, while pressure, temperature and oxygen fugacity were similar; or (2) that the amphibole efficiently separated from or was completely assimilated by the soon-to-be-erupted Giggle Springs magma. Arguing against complete assimilation is the observation that resorption of >1 cm amphiboles during the short time scale of final ascent might not be feasible. Mafic enclaves at both Marcath and Giggle Springs volcanoes likely represent crystallization products of these or other ponded magma batches that had not yet solidified; in the latter case, temporary stalling allowed for thermal and chemical interactions that facilitated entrainment of the enclaves (e.g., Valentine and Hirano, 2010).

Despite the mid-crustal ponding of Giggle Springs and Marcath magmas, the lack of evidence of crustal contamination (supported both by the major and trace element chemistry, and by isotopic data; Rasoazanamparany et al., in review) indicates that ascent between storage levels, and between the crustal reservoir and the surface, was rapid. Rapid ascent is a common feature of intraplate, small volume alkali-basalt volcanoes (e.g., Luhr et al., 1995; Valentine and Perry, 2007; Smith et al., 2008; Muffler et al., 2011). The lack of contamination, even though the magma did pond in the mid-crust, is consistent with the volcanoes being fed by small, single batches of magma. When a small batch of magma ponds it is expected that the margins will rapidly chill, increasing in local viscosity and potentially solidifying. This isolates of the inner part of the intrusion from country rocks both chemically and mechanically, until crystallization causes the volatile content to increase sufficiently in the melt to drive dike propagation and final magma ascent to the surface. In contrast, relatively large and repeated magma batches might convect vigorously and retard development of a thick chilled margin and provide enough heat to partly melt and assimilate wall rocks.

An alternative explanation of the deeper-derived phenocrysts is that the Giggle Springs and Marcath magmas did not stall at the deeper 30-45 km level, but simply entrained crystals from that depth that had formed during ponding of earlier (e.g., Mizpah, Hi Desert) magmas. We view this as unlikely because if younger magmas were entraining material as they migrated upwards we would

expect them to sample a variety of upper mantle and crustal rocks, not just the products of earlier magma batches.

The final stage on the evolution of these volcanic products is eruption after the ponded magmas have crystallized olivine, clinopyroxene, plagioclase and traces of amphibole. The composition of the amphibole crystals (kaersutite to pargasite/magnesiohastingite) and its stability suggest that this phase is derived either from a metasomatized lithospheric mantle or crystallized from an alkaline melt derived from a peridotite + olivine websterite source (Mayer et al., 2014). Our data are consistent with the latter, phenocrystic origin, for this phase.

The CSDs of one the younger volcanic products (Giggle Springs lava) suggest short crustal residence time, and the further crystallization and growth of olivine, clinopyroxene and an oxide phase, during eruption. Most of the samples have well developed groundmasses rather than glassy matrixes, indicating that subsolidus temperatures were reached slowly after the lava was emplaced in its final location.

633 IMPLICATIONS

The original motivation for this study was to address two questions: (1) what are the compositional variability and magma ascent processes within the time span of an individual eruption? The results presented here indicate two main behaviors of the four studied volcanoes. Individual magma batches were fed by slightly heterogeneous source material for the two older volcanoes, but the younger of the two volcanoes' source rocks were relatively homogeneous even over distances of several kilometres and time differences of tens of thousands of years. Each magma batch stalled and crystallized in the uppermost several kilometres of the mantle near the base of the crust. Even for the closely spaced volcanoes (Mizpah, Hi Desert, and Giggle Springs), the age differences of 10s to 100s

ka and lack of substantial fractionation suggest that each magma batch and temporary deep reservoir was a separate entity rather than part of a continuous long-lived reservoir. Magmas feeding the Marcath and Giggle Springs volcanoes also stalled at mid-crustal levels. From the evidence at hand it appears that ascent rates from source to storage levels, and from final storage to eruption, were relative rapid with no geochemical interaction with surrounding rocks. (2) What is the variability between very closely spaced volcanoes with different ages? The Mizpah and Hi Desert magmas have subtle compositional variability, but time is a major factor as well. Volcanoes erupted within certain time windows (an older time window for Mizpah and Hi Desert, and a younger 0-100 ka window for Giggle Springs and Marcath) have similar source characteristics and ascent processes whether they are located within a few hundred meters of each other or are separated by many kilometres. If each monogenetic volcano is fed by a single small-volume batch of magma, with source and feeder system length scales on the order of hundreds of meters to a few kilometres (e.g., Valentine and Perry, 2006, 2007; Valentine and Keating, 2007; Brenna et al., 2012), it is not clear why volcanoes that are several kilometres and tens of thousands of years apart would have such similar magma dynamics. Preliminary data from other parts of the Lunar Crater Volcanic Field are suggesting that this time window-dependent behavior might be general over larger distances; testing this and developing a fieldscale conceptual model are topics of current research.

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Figure Captions Figure 1: Location of the Lunar Crater Volcanic Field in the western U.S.A. and the State of Nevada showing the Pancake and Reveille Range within the volcanic field. Dashed lines are the main roads in the area. Red box encompasses the study area (see Figure 2). Figure 2: Simplified geological map of the northern part of the Lunar Crater Volcanic Field, highlighting the four volcanoes studied in this work. Outline of tephra deposits from Johnson (2013). Figure 3: Total Alkali versus silica diagram according to Le Maitre et al. (2002). Hi Desert basalt (Ohib) open circle, Mizpah basalt (Omzb) open square, Giggle Spring basalt (Ogsb) asterisk, Marcath volcano products (Qm) crosses. All whole-rock figures plotted using PINGU (Cortés and Palma, 2011). Figure 4: Whole-rock major element variation diagrams versus MgO for the samples of this study. Symbols as in Figure 3. Figure 5: Rare Earth Element diagram normalized according to Sun and McDonough (1989). Symbols as in Figure 3. Figure 6: Spider diagrams normalized according to OIB from Sun and McDonough (1989). Symbols as in Figure 3. Figure 7: a) La/Yb vs. Zr/Nb b) Rb/Zr vs. Ni/Zr c) La/Yb vs. Dy/Yb. Symbols as in Figure 3.

Figure 8: a) La/Nb vs Nb [ppm] b) Nb/Ta vs Nb [ppm]. Fields as defined in Barth et al., (2000).

Symbols as in Figure 3.

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Figure 9: Mineral classification based on mineral chemistry for a) olivine (Deer et al., 1996) b)

pyroxenes (Morimoto, 1989) c) plagioclase (Deer et al., 1996) and d) amphiboles (Leake, 1997) from

electron microprobe analyses in the studied samples. Symbols of the units as in Figure 3.

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Figure 10: (a) Fe^{2+}/Mn vs. %Fo=Mg/(Mg+Fe²⁺+Mn) and (b) $100xMn/Fe^{2+}$ vs. $100xCa/Fe^{2+}$ (%Fo>85)

in mineral chemistry analyses of olivine recalculated as cation per formula unit from electron

microprobe analyses. Symbols of the units as in Figure 3.

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Figure 11. Natural logarithm of the density population (mm⁻⁴) vs. size (mm) plots of the crystal size

distribution for a) olivine b) clinopyroxene c) plagioclase and d) spinel in the studied samples.

Minimum crystal sizes that can be measured using ImageJ in an SEM image are ~10-20 pixels long,

equivalent to ~5% of the average scale bar of ~300 pixels (around ~0.01-0.02 mm in most of our

images except images for measuring oxides with minimum sizes of 0.001-0.002 mm). Black asterisk:

Giggle Spring basalt (Qgsb) sampled at the vent. Light grey asterisk: Giggle Spring basalt (Qgsb) lava.

Black open squares: Mizpah basalt (Qmzb), Black crosses: Marcath volcano products (Qm).

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Figure 12: La/Yb vs. Dy/Yb plots with different models of partial melting of a mantle source. Data

points are normalized with C1 chondrite (Sun and McDonough 1989), symbols as in Figure 3. (a)

Partial melting of lherzolite source: grey line and filled triangles is a model of batch melting of a garnet

lherzolite with 11.5 % of garnet; black line with filled circles is a model of batch melting of a garnet

lherzolite with 6% of garnet, black line with open squares is a model of batch melting of a garnet

lherzolite with 2% of garnet, black line with open triangles is a model of batch melting of a spinel

lherzolite. Tick marks on all curves are at 5% melting intervals. (b) Partial melting of pyroxenite source: black line and filled triangles is a model of batch melting of a garnet pyroxenite with 11% garnet; black line with filled circles is a model of batch melting of a garnet pyroxenite with 9% garnet, black line with open squares models batch melting of a garnet pyroxenite with 5% garnet. Tick marks on all curves are at 5% melting intervals.















































