

Clinopyroxene–Liquid Equilibria and Geothermobarometry in Natural and Experimental Tholeiites: the 2014–2015 Holuhraun Eruption, Iceland

David A. Neave (1) 1,2*, Enikő Bali³, Guðmundur H. Guðfinnsson³, Sæmundur A. Halldórsson³, Maren Kahl^{3,4}, André-Sebastian Schmidt¹ and François Holtz¹

¹Leibniz Universität Hannover, Institut für Mineralogie, Callinstraße 3, Hannover 30167, Germany; ²Department of Earth and Environmental Sciences, The University of Manchester, Oxford Road, Manchester M13 9PL, UK; ³NordVulk, Institute of Earth Sciences, University of Iceland, Askja, Sturlugata 7, Reykjavík 101, Iceland; ⁴Institut für Geowissenschaften, Ruprecht-Karls-Universität Heidelberg, Im Neuenheimer Feld 234–236, Heidelberg 69120, Germany

*Corresponding author. Telephone: +44 (0)161 306 6589. E-mail: david.neave@manchester.ac.uk
Received November 22, 2018; Accepted September 4, 2019

ABSTRACT

Clinopyroxene-liquid geothermobarometry is a widely used tool for estimating the conditions under which mafic magmas are stored before they erupt. However, redox variability, sector zoning and disequilibrium crystallization present major challenges to the robust estimation of magma storage conditions. Moreover, most recent studies seeking to address these challenges have focused on clinopyroxenes from alkalic systems and are thus of limited use for understanding clinopyroxenes from the tholeiitic systems that dominate global magma budgets. Here we combine observations on natural clinopyroxenes from the 2014-2015 Holuhraun lava in Iceland with observations on experimental clinopyroxenes synthesized during high-pressure, high-temperature experiments on the same lava in order to investigate clinopyroxene-liquid equilibria in tholeiitic systems and optimize of geothermobarometric strategies. Natural clinopyroxenes from the 2014–2015 Holuhraun lava are sector zoned, with {111} hourglass sectors being enriched in the enstatite-ferrosillite component at the expense of all other components with respect to $\{hk0\}$ prism sectors. In contrast with observations on clinopyroxenes from alkalic systems, sector zoning in clinopyroxenes from the 2014-2015 Holuhraun lava is characterized by differences in Ca and Na contents as well as in Ti and Al contents. The products of crystallization experiments performed at 100-600 MPa and 1140-1220 °C on a powdered starting glass at two sets of melt H₂O contentoxygen fugacity conditions (~0.1 wt % H₂O and close to the graphite-oxygen redox buffer, and 0.5-1.0 wt % H₂O and approximately one and half log units above the guartz-favalite-magnetite redox buffer) demonstrate that clinopyroxene crystals from nominally equilibrium experiments can preserve strongly disequilibrium compositions. The compositional systematics of experimental clinopyroxenes are consistent with the presence of sector zoning. Furthermore, the magnitude of compositional variability increases with decreasing melt H₂O content and increasing deviations of experimental temperatures below clinopyroxene liquidus temperatures (i.e. degrees of undercooling sensu lato), indicating that kinetic processes play a key role in controlling clinopyroxene compositions, even under notionally equilibrium conditions. Few published analyses of experimental clinopyroxene crystals may thus represent truly equilibrium compositions. Stoichiometric calculations on natural and experimental clinopyroxenes show that Fe³⁺ is a major constituent of clinopyroxenes from tholeiitic magmas under naturally relevant oxygen fugacity conditions. They also show that Fe $^{3+}$ is most likely incorporated as Ca- and Al- bearing Ca–Fe-Tschermak's component rather than Na-bearing aegirine component at oxygen fugacities up to one and a half log units above the quartz–fayalite–magnetite buffer. Elevated oxygen fugacities are thus less likely to compromise clinopyroxene–liquid geothermobarometry than previously thought. Guided by our experimental results, we combined published descriptions of clinopyroxene–liquid equilibria with geothermobarometric equations to develop an internally consistent and widely applicable method for performing geothermobarometry on tholeiitic magmas that does not require equilibrium zones to be selected *a priori*. Applying this method to natural clinopyroxene crystals from the 2014–2015 Holuhraun lava that formed under low but variable degrees of undercooling (perhaps 25 °C or less) returns values in excellent agreement with those from independent methods (232 \pm 86 MPa, 1161 \pm 11 °C). Robust estimates of magma storage conditions can thus be obtained by performing clinopyroxene–liquid geothermobarometry on tholeiitic magmas when disequilibrium is suitably accounted for.

Key words: geothermobarometry; clinopyroxene; tholeiite; experimental petrology; Iceland

INTRODUCTION

Clinopyroxene is a key constituent of many mantle, igneous and metamorphic rocks. Its stability across diverse pressure-temperature-composition (P-T-X) conditions derives from its ability to incorporate a large range of cations in both octahedral and tetrahedral coordination states (Morimoto et al., 1988). Although this compositional plasticity makes it challenging to model clinopyroxene chemistry (e.g. Sack & Ghiorso, 1994; Green et al., 2012), it does mean that clinopyroxene can record more information about the conditions under which it formed than chemically simpler minerals such as olivine (e.g. Putirka, 2008). For example, the P-sensitive incorporation of Na into clinopyroxene can be exploited as a geobarometer for estimating the depths at which volcanic products were stored prior to eruption (Putirka et al., 1996). Such depth estimates are vital for reconstructing magmatic plumbing system architectures and hence understanding mechanisms of crustal accretion (e.g. Maclennan et al., 2001) and interpreting geophysical signals of magma movement before and during volcanic eruptions (e.g. Longpré et al., 2014). However, obtaining reliable values from clinopyroxene geobarometers is not always straightforward: variations in oxygen fugacity (fO₂), sector zone development, disequilibrium crystallization and uncertainties in geobarometric equations can all lead to spurious results, even when geochemical analyses are of superior quality (Mollo et al., 2013; Hammer et al., 2016; Neave & Putirka, 2017). Here we use observations on natural clinopyroxenes from an archetypal tholeiitic basalt emplaced during the 2014-2015 Holuhraun eruption in Iceland to identify sources of uncertainty in magma storage depth estimates. Observations on natural samples are complemented by experimental observations designed to test the reliability of Neave & Putirka's (2017) clinopyroxene-liquid geobarometer and facilitate the optimisation of methods for determining the conditions under which the tholeiitic magmas that dominate global magma budgets crystallize.

Clinopyroxene in mafic magmas

Tholeiitic basalts erupt at mid-ocean ridges, many ocean islands and some volcanic arcs. They are typically characterized by the presence of olivine, plagioclase and augitic clinopyroxene (e.g. Bryan, 1983; Tormey et al., 1987), though low-Ca pyroxene (pigeonite or orthopyroxene) may also occur (e.g. Lindsley, 1983; Grove & Juster, 1989). Although augite compositions can be summarised with reference to the pyroxene quadrilateral bounded by the diopside-hedenbergite (Di-Hd; CaMgSi₂O₆-CaFe²⁺Si₂O₆) and enstatite-ferrosillite (En–Fs; Mg₂Si₂O₆–Fe₂²⁺Si₂O₆) solid solutions, natural crystals often contain appreciable quantities of Ti, Al, Cr, Fe³⁺ and Na in non-quadrilateral components such as Ca-Tschermak's component (CaTs; CaAlAlSiO₆). The P-dependent incorporation of Na into clinopyroxene as the jadeite component (Jd; NaAlSi₂O₆) is of particular petrological interest because of its use in Jd-inclinopyroxene geobarometery (e.g. Putirka et al., 1996). However, the incorporation of the aegirine component (Ae; NaFe³⁺Si₂O₆) alongside the Jd component under oxidizing conditions can complicate relationships between P and clinopyroxene Na contents (Blundy et al., 1995), potentially limiting the reliability of Jd-inclinopyroxene geobarometry when magma Fe³⁺ contents are high (Neave & Putirka, 2017). Although Mössbauer spectroscopy on basanite-hosted clinopyroxenes from the Canary Islands confirms that clinopyroxenes in oxidized ocean-island magmas can contain significant quantities of Fe³⁺ (Fe³⁺/ Σ Fe \sim 0.37–0.51; Weis et al., 2015), a paucity of comparable data from more reduced magmas and the products of experiments performed at controlled fO2 conditions means that the effects of magma redox state on the performance of Jd-in-clinopyroxene geobarometry remain largely unquantified. By carrying out experiments at two different redox conditions, we aim to start exploring the effects of fO2 on augite chemistry and clinopyroxene-liquid geothermobarometry; a comprehensive assessment is however beyond the scope of this study.

Observations from natural and experimental clinopyroxenes

Augitic clinopyroxenes often exhibit sector zoning that manifests as compositional variability between families of crystallographically-related sectors (Strong, 1969; Hollister & Gancarz, 1971; Downes, 1974; Dowty, 1976; Skulski et al., 1994; Ubide et al., 2019). Chemical differences between sectors reflect the ability of different crystal faces to incorporate cations in equilibrium proportions during crystal growth (Nakamura, 1973; Leung, 1974; Dowty, 1976). For example, Nakamura & Coombs (1973) observed that {100} sectors of augite crystals from the tholeiitic Tawhiroko intrusion in New Zealand are characterized by disequilibrium componentry, with En and Fs components being enriched at the expense of Di and Hd components when compared with {010} and {110} sectors from the same grains. However, the sense and magnitude of sector zoning depends strongly on the degree of undercooling (i.e. the difference between the liquidus T and the magmatic or experimental T at which crystallization occurs) experienced (Kouchi et al., 1983). Thus, simply assuming that low-EnFs (i.e. Ti-, Al- and Na-rich) sectors record equilibrium growth conditions may be inappropriate. For instance, in their recent study of ankaramites from Haleakala, Hawaii, Welsch et al. (2016) described complex sector zoning resulting from the rapid, dendritic crystallization of Ti-, Al- and Na-rich {111} hourglass sectors at high degrees of undercooling, followed by the subsequent infilling of Ti-, Al- and Na-poor {hk0} prism sectors at lower degrees of undercooling. In this case, {hk0} sectors were interpreted as recording equilibrium growth conditions and hence thought to yield reliable geobarometric results (Hammer et al., 2016). However, despite Welsch et al. (2016) and Hammer et al. (2016) making compelling cases for critically investigating how sector zoning influences crystal compositions and geothermobarometric calculations, it is currently unclear whether their observations on alkalic systems provide a reliable framework for interpreting data from tholeiitic systems (Neave & Putirka, 2017; Ubide et al., 2019).

Although the degree of undercooling mediates the nature and degree of sector zone development during crystallization (Kouchi et al., 1983; Welsch et al., 2016), it also affects the composition of crystals independently of sector zone development (Lofgren et al., 2006; Mollo & Hammer, 2017). That is, clinopyroxene bulk compositions depend on the degree of undercooling as well as on the prevailing P-T-X conditions. Augite crystals formed at high undercooling are characterized by an excess of non-quadrilateral components (Lofgren et al., 2006; Mollo & Hammer, 2017), potentially resulting in the calculation of spurious storage conditions by clinopyroxene-liquid geothermobarometry, as demonstrated in dynamic crystallization experiments with high cooling rates (Mollo et al., 2010). Filtering clinopyroxenes for multicomponent equilibrium nevertheless provides a way to exclude disequilibrium compositions

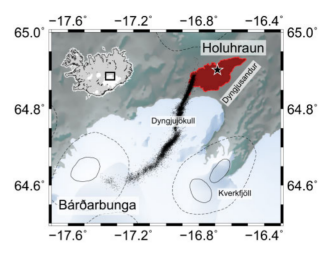


Fig. 1. Map showing the location of the 2014–2015 Holuhraun lava in central Iceland. The extent of the lava flow is shown in red. Black points show the epicentres of earthquakes associated with dyke propagation from the Bárðarbunga central volcano towards the eruption site (Ágústsdóttir et al., 2016). Calderas and central volcanoes are outlined with solid and dashed lines respectively. Sample MSR161014 was collected at the location marked with a black star.

from geothermobarometric calculations (e.g. Mollo et al., 2013) and has enabled the calculation of coherent magma storage depths in a number of settings including Iceland (e.g. Halldórsson et al., 2018). However, such multicomponent filtering has yet to be validated with experiments on tholeitic compositions, much of the work carried out to date has focussed on alkali basalts from Mt. Etna, Italy (e.g. Mollo et al., 2011, 2013, 2018). We thus aim to develop an experimentally validated strategy for robustly identifying equilibrium clinopyroxene–liquid pairs in tholeitic magmas that will be of widespread use for studying systems beyond the 2014–2015 Holuhraun lava.

The 2014–2015 Holuhraun eruption

The 2014-2015 Holuhraun volcano-tectonic event in central Iceland was one of the best monitored basaltic fissure eruptions ever to have taken place (Sigmundsson et al., 2015; Ágústsdóttir et al., 2016; Guðmundsson et al., 2016). Approximately 1.2 km³ of lava (dense rock equivalent) was emplaced on the Dyngjusandur plain north of the Vatnajökull icecap over the course of the eight-month long eruption (Fig. 1; Pedersen et al., 2017; Bonny et al., 2018), making it the volumetrically largest eruption in Europe since the 1783-1784 Laki Fires (Thordarson & Self, 1993). Subsidence within the Bárðarbunga caldera was recorded from the onset of dyke intrusion (Sigmundsson et al., 2015), and continued at a rate commensurate with the decaying mass flux at the vent until the end of the eruption (Guðmundsson et al., 2016). A magmatic connection between the vent and a steadily deflating reservoir beneath the Bárðarbunga caldera thus appears to have been maintained throughout the course of the eruption.

Geodetic observations place a putative source reservoir at a depth of 8–12 km beneath the Bárðarbunga central volcano (Guðmundsson *et al.*, 2016), consistent with depths from carefully implemented, if not experimentally verified, petrological geobarometry based on clinopyroxene–liquid geothermobarometry, olivine–plagioclase–augite–melt (OPAM) thermal minima positions and melt and fluid inclusion entrapment pressures (Bali *et al.*, 2018; Halldórsson *et al.*, 2018; Hartley *et al.*, 2018).

The 2014–2015 Holuhraun lava is a moderately evolved and strikingly homogenous tholeiite similar in composition to magmas erupted across Iceland and the mid-ocean ridge system more broadly (Halldórsson et al., 2018). Crystals of olivine, plagioclase and augitic clinopyroxene were erupted throughout, but are present in sufficiently low abundances (<5 vol.%) that some whole-rock compositions approximate those of magmatic liquids; whole-rock compositions overlap with melt inclusion compositions (Hartley et al., 2018). The presence of clinopyroxene in an otherwise modestly porphyritic basalt makes the 2014-2015 Holuhraun lava an ideal subject for evaluating clinopyroxene geothermobarometry in tholeiitic systems because most clinopyroxene crystals are likely to have grown from liquids directly parental to their carrier melts (i.e. whole $rock \approx carrier liquid + equilibrium crystal assemblage);$ note that this is not the case for some primitive olivine and plagioclase compositions (Halldórsson et al., 2018). Crystal-poor whole-rock samples can thus be used as starting materials for experiments designed to mimic the final stages of magmatic evolution. Here we focus on poorly porphyritic MSR161014 (≤5 vol.% macrocrysts; Halldórsson et al., 2018) from the main phase of the 2014-2015 Holuhraun eruption. This sample was collected on day 50 of the eruption at 64-902° N, 16-686° W by the 2014-2015 Holuhraun Eruption Team of the University of Iceland (Fig. 1). It has been thoroughly characterized geochemically and is thus well-suited to the focussed investigation we present here.

METHODS

Experimental methods

A starting glass powder for high-*P*, high-*T* experiments was produced by fusing 10 g of powder from sample MSR161014 in a muffle furnace at the Institut für Mineralogie of the Leibniz Universität Hannover, Germany. By using a quenched glass powder we sought to minimise reactions prior to crystallizing an equilibrium assemblage in order to mimic crystallization from super-liquidus conditions (Mollo & Hammer, 2017). The MSR161014 powder was melted at 1600 °C in a large-volume Pt crucible under air to prevent Fe loss. A homogenous glass powder was then obtained by quenching the crucible in H₂O after 1 hour and grinding the resulting glass chips in an agate disc mill. The resulting glass powder was then melted, quenched and ground a second time prior to storage in a desiccator.

The composition of the starting glass powder is within analytical uncertainty of published whole-rock compositions for MSR161014 (Supplementary Material; supplementary data are available for downloading at http://www.petrology.oxfordjournals.org; Halldórsson *et al.*, 2018).

High-P, high-T experiments were carried out under different fO2 conditions by using different capsule designs. Experimental redox conditions close to (or below) the graphite-oxygen (CCO) redox buffer and one log unit above the quartz-fayalite-magnetite (QFM) redox buffer were chosen to bracket those recorded in the 2014-2015 Holuhraun lava (maximum $fO_2 \sim QFM + 0.5$; Bali et al., 2018). Reducing conditions were achieved by loading ~20 mg of dried starting glass powder into graphite inner capsules that were then enclosed within Pt outer capsules. In this case, the graphite inner capsules theoretically buffered experimental samples at an fO2 at or below CCO (i.e. at or below \sim QFM -1.7), with excess O₂ reacting to form CO₂ (Holloway et al., 1992; Jakobsson & Óskarsson, 1994; Médard et al., 2008). In a second suite of capsules designed to run under moderately oxidizing conditions, \sim 60 mg of dried starting glass powder was loaded into Au₈₀Pd₂₀ (AuPd) capsules that had been pre-saturated with ~ 0.2 wt % Fe (e.g. Gaetani & Grove, 1998; Balta et al., 2011). With the intention of buffering samples at approximately QFM+1, no H₂O was added (e.g. Husen et al., 2016). Small quantities of H_2O (~ 0.5 wt %) were nevertheless expected to form by the reaction of trace H₂ in the pressure medium with O₂ liberated from the reduction of Fe₂O₃ in the starting glass (Botcharnikov et al., 2005; Husen et al., 2016).

Experimental runs were performed by suspending either graphite-Pt double capsules or AuPd capsules from thin Pt wires in the hot zone of an internally heated pressure vessel (IHPV) at the Institut für Mineralogie of the Leibniz Universität Hannover, Germany (e.g. Berndt et al., 2002). Experiments were performed at 100, 300 and 600 MPa using an Ar pressure medium; no 600 MPa experiments were performed using graphite-Pt double capsules. Pressure was continuously monitored with a strain gauge manometer and did not vary more than the measurement uncertainty (5 MPa) during experimental runs. Experiments were performed at 1140, 1160 and 1180 °C for all P conditions in order to bracket magma storage conditions estimated in earlier work (Halldórsson et al., 2018); additional higher-T experiments were performed at 1200 and 1220 °C for some PfO₂ combinations (Table 1). Experimental samples were heated to 20 °C below the target T at a rate of 50 °C/min, and then to the target T at a rate of 10 °C/min in order to avoid overshooting; no high-T treatments were applied at the start of any runs. After holding samples at the target T for one hour, thermal oscillations of ± 10 °C were imposed for the first third of each experimental run to promote the formation of large crystals and large melt pools (Erdmann & Koepke, 2016). Temperature was monitored across the vessel's 25 mm-tall hot zone with

Table 1: Summary of experimental conditions

Experiment number	P (MPa)	T (°C)	Capsule	Duration (hours)	Fe loss/ gain (%)	fO ₂ estimates ¹				H ₂ O (wt.%) ⁶	Phase proportions ⁷				ΔT _{cpx} ⁸ (°C)
						ΔQFM (S08) ²	ΔQFM (BG10) ³	ΔQFM (T05) ⁴	ΔQFM (B05) ⁵	(VVI./0)	gl	ol	plg	срх	(0)
synthesis		1600		1+1											
B0188-3	100	1140	AuPd	120	-14		0.51		2.70	1.06	1.00				
Y0185-11	100	1160	AuPd	120	-9	1.29	0.76		2.08	0.75	1.00				
Y0186-2	100	1180	AuPd	72	-1	1.47	1.12		1.76	0.61	1.00				
Y0187-5	100	1200	AuPd	48	6	1.51	1.03		1.44	0.50	1.00				
Y0205-2	300	1140	AuPd	72	3		1.42				0.65		0.15	0.20	20-40
Y0201-2	300	1160	AuPd	75	1		0.77				0.94			0.06	<20
Y0174-7	300	1180	AuPd	90	1	1.55	1.09		1.28	0.64	1.00				
Y0166-10	300	1200	AuPd	48	-6	1.77	0.80		1.38	0.68	1.00				
B0187-8	600	1140	AuPd	96	-9		0.82				0.64		0.10	0.26	>60
B0183-4	600	1160	AuPd	96	-14		0.33				0.87			0.13	>40
B0167-1	600	1180	AuPd	72	-6		0.67				0.90			0.10	>20
B0184-10	600	1200	AuPd	48	-6		0.62				0.99			0.01	>0
Y0272-1	100	1140	C-Pt	96	-3			-0.70			0.66	0.04	0.18	0.12	20-40
Y0287-1	100	1160	C-Pt	72	1			-0.96			0.82	0.02	0.10	0.06	<20
Y0281-1	100	1180	C-Pt	72	-1				-2.72	0.04	1.00				
Y0204-3	300	1140	C-Pt	72	-1			-1.00			0.50	0.00	0.24	0.26	40-60
Y0200-3	300	1160	C-Pt	72	1						0.66		0.15	0.18	20-40
Y0206-3	300	1180	C-Pt	72	0						0.97			0.03	<20
Y0216-3	300	1200	C-Pt	71	-3				-2.08	0.08	1.00				
Y0218-3	300	1220	C-Pt	66	-1				-2.08	0.08	1.00				

¹Note that all methods of fO_2 estimation used are associated with uncertainties in the order of 0.5-1.0 log units.

four unsheathed S-type (Pt₉₀Rh₁₀) thermocouples and was typically within 5 °C of the target temperature (no measured temperature deviated more than 10°C from the target temperature). Run durations varied from 48 hours for the hottest experiments to 96-120 hours for the coolest. Capsules were quenched rapidly at the end of each run by fusing the Pt wires on which they were suspended, dropping them into a cold zone at the vessel's base.

Analytical methods

After the capsules had been weighed to validate their integrity during high-P, high-T runs, experimental samples (including capsule pieces) were mounted in epoxy resin and polished for imaging and microanalysis. Sample pieces from super-liquidus experiments were mounted in epoxy resin and double-polished for spectroscopic analyses to monitor experimental H₂O contents. Further sample fragments from super-liquidus experiments were conserved for determining glass $Fe^{3+}/\Sigma Fe$ values by colorimetry to monitor experimental fO₂ conditions. Natural clinopyroxenes were investigated in a polished thin section cut from a block of MSR161014.

Both natural and experimental samples were imaged using a JEOL JSM-7610F field emission gun scanning electron microscope (FEG-SEM) at the Institut für Mineralogie of the Leibniz Universität Hannover, Germany. Glass, mineral and AuPd-capsule compositions were determined by electron probe microanalysis (EPMA) at the Institut für Mineralogie of the Leibniz Universität Hannover, Germany using a Cameca SX100 instrument, Silicon, Ti, Al, Cr, Fe, Mn, Mg, Ca, Na, K and P were measured in glasses with a beam size of 12 μm, an accelerating voltage of 15 kV and a current of 10 nA. Silicon, Ti, Al, Cr, Fe, Mn, Mg, Ca, Na and K were measured in minerals with a beam size of 1 µm, an accelerating voltage of 15 kV and a current of 15 nA. Gold, Pd and Fe were measured in capsules with a beam size of $1 \mu m$, an accelerating voltage of 15 kV and a current of 40 nA. Elements were counted on peak for 20 s, with the exceptions of Si and Na that were counted on peak for 10 s to minimise drift and migration respectively. Background counting times were half on-peak counting times. The following standards were used for calibration: wollastonite (Si and Ca), TiO2 (Ti), jadeite (AI), Cr2O3 (Cr), Fe₂O₃ (for Fe in silicates), Fe metal (for Fe in capsules), Mn₃O₄ (Mn), MgO (Mg), albite (Na), orthoclase (K), apatite (P), Au metal (Au) and Pd metal (Pd).

To ensure consistency across multiple analytical sessions, analyses of unknowns were normalized to secondary standards as follows: glass analyses were normalized to VG-2 basaltic glass (NMNH 111240-52; using the preferred MgO content); clinopyroxene and plagioclase analyses were normalized to Kakanui augite

²Oxygen fugacity estimated by colorimetry following Schuessler et al. (2008).

³Oxygen fugacity estimated from AuPd-capsule compositions following Barr & Grove (2010).

⁴Oxygen fugacity estimated from glass Fe³⁺/ΣFe contents determined using K_D^{Mg-Fe}_{ol-liq} values calculated following Toplis (2005).

⁵Oxygen fugacity estimated from aH₂O following Botcharnikov et al. (2005).

 $^{^6}$ Glass $\mathrm{H}_2\mathrm{O}$ contents determined by FTIR on superliquidus runs.

⁷gl, glass; ol, olivine; plg, plagioclase; cpx, clinopyroxene. phase proportion rounded to two significant figures.

⁸Difference between experimental T and clinopyroxene liquidus T (i.e. degree of undercooling sensu lato).

(NMNH 122142; using preferred values); olivine analyses were normalized to San Carlos olivine (NMNH 11312–44) (Jarosewich *et al.*, 1980). Accuracy and precision were monitored by measuring the following additional secondary standards: A-99 basaltic glass (NMNH 113498), Ney County Cr-augite (NMNH 164905) and Lake County plagioclase (NMNH 115900) (Jarosewich *et al.*, 1980, 1987). Major (>1 wt %) and minor (<1 wt %) elements were typically determined with accuracies better than 2% and 10%, and 1σ precisions better than 2% and 15% respectively. Typical standard analyses are provided alongside analyses of our experimental products in the Supplementary Data; supplementary data are available for downloading at http://www.petrology.oxfordjournals.org.

Water contents of super-liquidus experiments were determined by Fourier-transform infrared (FTIR) spectroscopy using a Bruker IFS88 instrument at the Institut für Mineralogie of the Leibniz Universität Hannover, Germany, and an absorption coefficient for the 3500 cm⁻¹ H_2O_t band given by Shishkina *et al.* (2010). The speciation of iron (i.e. $Fe^{3+}/\Sigma Fe$) in super-liquidus experiments was determined by colorimetric analysis following the methods of Schuessler *et al.* (2008). The precision of $Fe^{3+}/\Sigma Fe$ determinations were monitored by measuring an inhouse andesite standard (PU-3; Borisov *et al.*, 2018). Super-liquidus glass H_2O contents and $Fe^{3+}/\Sigma Fe$ values are given in Table 1.

PETROGRAPHIC RESULTS

Natural samples

MSR161014 is a fresh, poorly porphyritic basalt (≤5 vol.% macrocrysts) containing, in order of abundance, plagioclase, clinopyroxene and olivine macrocrysts in a crystalline groundmass (Fig. 2). Note that we simply define macrocrysts as crystals that are >100 μm in length and texturally distinct from the groundmass (e.g. Thomson & Maclennan, 2013). Plagioclase and clinopyroxene macrocrysts are typically 100-400 μm in length, while olivine macrocrysts are slightly smaller, with lengths of 100-200 μm Macrocrysts occur as individual grains and within glomerocrystic clusters. clinopyroxene and plagioclase macrocrysts frequently occur in a sub-ophitic arrangement, although monomineralic glomerocrysts of plagioclase and clinopyroxene are also present. There are no clear textural differences between isolated macrocrysts and macrocrysts in glomerocrysts. Concentric and sector zoning textures are evident in plagioclase and clinopyroxene, respectively, when viewed under crossed polars (Fig. 2b). Fe-Ti oxides are restricted to the groundmass.

Backscattered electron (BSE) imaging reveals that clinopyroxene macrocrysts cores are generally equant, sub- to euhedral and enveloped by complex rims intergrown with the sample's crystalline groundmass (Fig. 3). Two compositionally distinct families of sector zones can be identified on the basis of their different BSE intensities. Combining SEM observations with

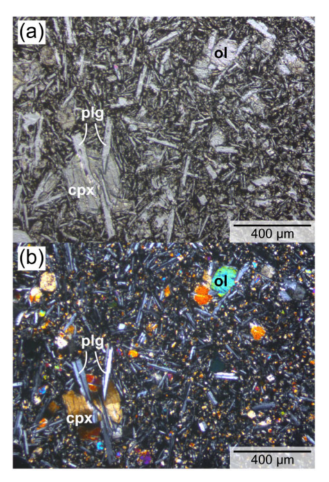


Fig. 2. Photomicrographs of sample MSR161014 from the 2014–2015 Holuhraun lava under (a) plane and (b) crossed polars. Macrocrysts of olivine (ol), plagioclase (plg) and clinopyroxene (cpx) are visible within a crystalline groundmass. The labelled clinopyroxene is in sub-ophitic arrangement with plagioclase and shows sector zoning under crossed polars.

optical microscopy indicates that low BSE intensities are associated with {111} hourglass sectors. For example, the macrocryst shown in Fig. 3a (cpx-21) has been cut parallel to (010) such that [100] and [001] lie within the plane of the thin section and are separated by a β -angle of \sim 105°. High-BSE-intensity sectors in this image thus formed by growth on cleavage-parallel {110} faces, and low-BSE-intensity sectors by growth on cleavage-oblique {111} faces (e.g. Downes, 1974). A small high-BSE-intensity zone associated with growth on {010} faces is visible in the centre of the macrocryst. In contrast, the macrocryst shown in Fig. 3b (cpx-22) has been cut between (100) and (001) such that only [010] lies close to the plane of the thin section. Low-BSE-intensity sectors in this macrocryst are still associated with growth on {111} faces; high-BSE-intensity sectors reflect growth on {110} and {010} faces. However, the plane of the thin section must lie close to the interface between {111} and {100} sectors because (111) and (111) faces cannot be distinguished from each other (Leung, 1974; Welsch et al., 2016). Further macrocrysts in the sample were cut close to (001),

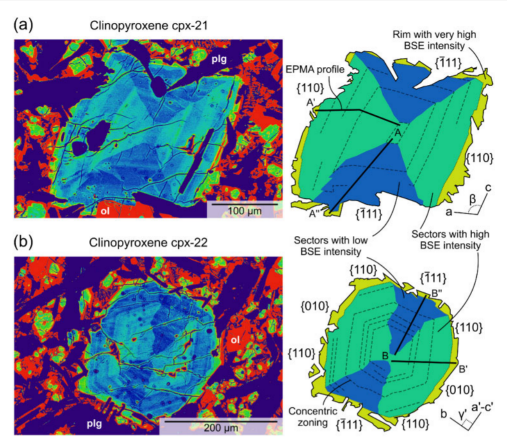


Fig. 3. Backscattered electron (BSE) images of clinopyroxene macrocrysts in sample MSR161014 from the 2014–2015 Holuhraun lava. Olivine (ol) and plagioclase (plg) macrocrysts are indicated. Interpretive sketches are shown to highlight patterns of sector and concentric zoning. Compositional profiles through pairs of sectors measured by electron probe microanalysis (EPMA) are marked on the sketches.

confirming the association of high BSE intensities with {100}, {010} and {110} prism sectors (Supplementary Data; supplementary data are available for downloading at http://www.petrology.oxfordjournals.org).

Concentric zoning is present in all of the clinopyroxene macrocrysts imaged (Fig. 3). Although variations in BSE intensity associated with concentric zoning are comparable with those associated with sector zoning, there are, with the exception of very high-BSE-intensity rims in contact with the groundmass, no consistent changes in BSE intensity along core-to-rim profiles. Importantly, concentric zones can be traced across sector zone boundaries, indicating that all sectors grew concurrently (cf. Welsch et al., 2016).

Experimental samples

Experimental conditions are summarised in Table 1 and BSE images of typical experimental products are shown in Fig. 4. Crystals are typically 10–40 μm , but occasionally up to 200 μm in length, and are separated by pools of quenched and visually homogenous glass. Most crystals are euhedral; only olivine crystals show consistently rounded morphologies. No compositional zonation was observed in experimental olivine and plagioclase. Although the texture of experimental clinopyroxene crystals varies considerably between the products of

different experiments, two overarching trends can nevertheless be identified. Firstly, the largest clinopyroxene crystals grew in high-T, high-melt-fraction (high-F) experiments (Fig. 4a and c). Specifically, large (>20 μm diameter) clinopyroxene crystals only grew in the highest-T sub-liquidus experiments performed under any combination of P and fO2 conditions. Some large clinopyroxenes also show evidence for internal zonation that is reminiscent of the complex zoning observed in natural crystals (Fig. 4a and d). No large clinopyroxene crystals were produced in our 600 MPa experiments, though we did not bracket the liquidus under these conditions (Fig. 4b). Secondly, clinopyroxene crystals in low-T, low-F experiments typically form chains of small ($<10\,\mu m$ diameter), intergrown grains (Fig. 4d).

MICROANALYTICAL RESULTS

Natural samples

The composition of the experimental starting glass prepared from sample MSR161014 is shown in Fig. 5 (black star) and is comparable with published whole-rock analyses of this sample (Halldórsson *et al.*, 2018). MSR161014 is slightly more primitive (6-82 wt % MgO) than matrix glass samples collected during the eruption

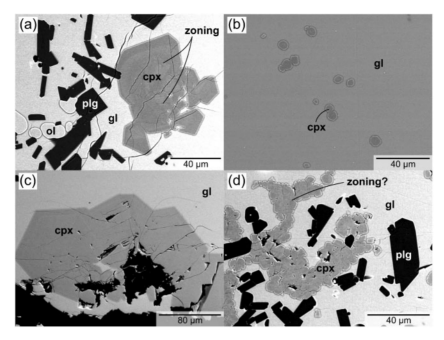


Fig. 4. Backscattered electron (BSE) images of typical products from experiments on sample MSR161014. (a) Olivine (ol), plagioclase (plg) and clinopyroxene (cpx) crystals in a glass (gl) matrix produced at 1160 °C and 300 MPa in a graphite-Pt double capsule. Zoning is highlighted in the largest clinopyroxene crystal. (b) Small, euhedral clinopyroxene crystals produced at 1200 °C and 600 MPa in an AuPd capsule. (c) Large, euhedral clinopyroxene crystals produced at 1180 °C and 300 MPa in a graphite-Pt double capsule. (d) Chains of small clinopyroxene crystals associated with somewhat larger plagioclase crystals produced at 1140 °C and 300 MPa in an AuPd capsule.

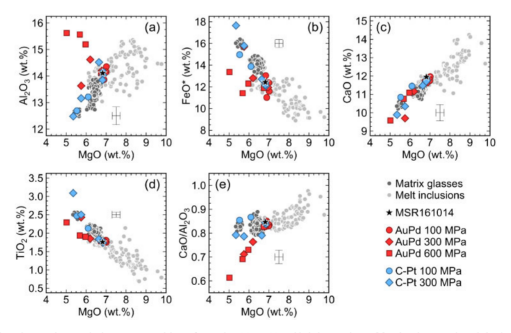


Fig. 5. Natural and experimental glass compositions from the 2014–2015 Holuhraun lava. Matrix glass and melt inclusion compositions from Halldórsson *et al.* (2018) and Hartley *et al.* (2018) are shown as dark and light grey circles respectively. Note that published matrix glasses with <6 wt % MgO preserve an imprint of syn-emplacement crystallization and are unrepresentative of magmatic liquids at depth (Halldórsson *et al.*, 2018). The experimental starting glass synthesized from sample MSR161014 is shown as a black star. Mean experimental glass compositions produced at different temperatures are shown with coloured symbols. Red symbols indicate experiments in AuPd capsules and blue symbols experiments in graphite-Pt (C-Pt) double capsules. Circles, diamonds and squares indicate glasses produced at 100, 300 and 600 MPa, respectively. All Fe is assumed to be Fe²⁺ (i.e. FeO*). Characteristic 2σ analytical uncertainties are shown.

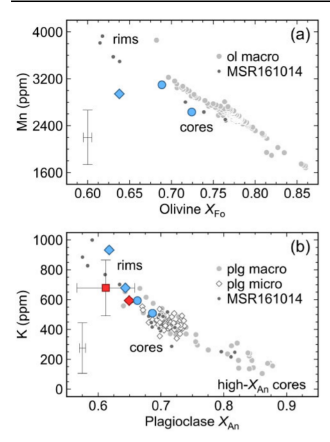


Fig. 6. Summary of natural and experimental olivine (ol) and plagioclase (plg) compositions from the 2014-2015 Holuhraun lava. Mean experimental compositions are shown using the same symbols as in Fig. 5. Core and rim compositions are annotated. Characteristic 2σ analytical uncertainties are shown. (a) Olivine compositions are summarised in terms of their forsterite [X_{Fo} , where $X_{Fo} = Mg/(Mg+Fe)$ on a molar basis] and Mn contents. Olivine macrocryst compositions from Halldórsson et al. (2018) are shown as light grey circles; natural compositions from sample MSR161014 are shown as dark grey circles. (b) Plagioclase compositions are summarised in terms of their anorthite [X_{An} , where $X_{An} = Ca/(Ca+Na+K)$ on a molar basis] and K contents. Plagioclase micro- and macrocryst compositions from Halldórsson et al. (2018) are shown as outlined white diamonds and light grey circles, respectively; compositions from sample MSR161014 are shown as dark grey circles. Experimental plagioclase crystals produced at 600 MPa are more heterogeneous than those produced under other conditions, as reflected in the greater uncertainty shown for this experiment.

(5·34–6·74 wt % MgO; Halldórsson *et al.*, 2018) and is instead most similar to the composition of evolved melt inclusions (Hartley *et al.*, 2018), consistent with its slightly primitive and porphyritic nature (Fig. 2). Note that published matrix glasses with <6 wt % MgO record syn-emplacement crystallization and are unrepresentative of magmatic liquids at depth (Halldórsson *et al.*, 2018).

Olivine and plagioclase macrocryst compositions from MSR161014 are compared with published crystal compositions in Fig. 6 (Halldórsson *et al.*, 2018). Olivine macrocryst cores have similar forsterite contents [$X_{Fo} \sim 0.76$, where $X_{Fo} = Mg/(Mg+Fe^*)$ on a molar basis and all Fe is assumed to be Fe²⁺ (i.e. Fe*)] to the mode of

published data ($X_{Fo} \sim 0.77$; Fig. 6a). Olivine rim compositions extend to lower X_{Fo} values than previously reported ($X_{Fo} \sim 0.62$ vs $X_{Fo} \sim 0.70$), probably because MSR161014 records an interval of crystallization during lava flow emplacement. Most plagioclase macrocryst cores have similar anorthite contents [$X_{An} \sim 0.70$, where $X_{An} = \text{Ca/(Ca+Na+K)}$ on a molar basis] to previously reported microcryst and evolved macrocryst cores. Some macrocrysts have high-anorthite cores ($X_{An} \sim 0.81$), as observed in other samples (Halldórsson et al., 2018; Hartley et al., 2018). Plagioclase rims in MSR161014 also extend to more evolved compositions than previously reported ($X_{An} \sim 0.58$ vs $X_{An} \sim 0.63$).

Clinopyroxene macrocryst compositions are summarised in Fig. 7. Macrocrysts from MSR161014 are compositionally similar to the main macrocryst population observed in lavas emplaced throughout the eruption (Halldórsson et al., 2018). Specifically, clino pyroxene macrocryst cores have Mg numbers [Mg#cpx/ where $Mg\#_{cox} = Mg/(Mg+Fe^*)$ on a molar basis and all Fe is assumed to be Fe²⁺ (i.e. Fe*)] in the range 0.75-0.82 (Fig.7a); a few rim analyses extend to Mg#cpx < 0.75. We did not observe any of the Cr-rich, highcompositions reported previously Halldórsson et al. (2018). Despite exhibiting only modest variability in Mg#cpx, clinopyroxenes show considerable correlated variability in TiO₂ and Al₂O₃ (Fig. 7b): high- and low-TiO2 (and Al2O3) populations with approximately >0.5 and <0.5 wt % TiO₂ respectively can be identified at broadly consistent Mg#cpx values.

Variability in clinopyroxene major element contents is illustrated in Fig. 8. Most clinopyroxene analyses from MSR161014 define a population characterized by low Mg but high Ti, Al, Ca and Na contents. Chromium contents are also somewhat elevated in this low-Mg population. A subordinate population of analyses form a tail to distinctly higher Mg contents. These high-Mg analyses are also characterized by low Ti, Al, Ca and Na contents. Integrating spatial information from imaging with compositional information from EPMA on grains cpx-21 and cpx-22 demonstrates that low- and high-Mg analyses are located systematically within high- and low-BSE-intensity sectors, respectively (Figs 3 and 8). Specifically, high-Mg analyses are from {111} hourglass sectors, and low-Mg analyses are from {100}, $\{010\}$ and $\{110\}$ (i.e. $\{hk0\}$) prism sectors.

Transects through high-BSE-intensity sectors are characterized by lower Mg# $_{cpx}$ contents than transects though low-BSE-intensity sectors (Figs 8 and 9). Importantly, transects from pairs of $\{hk0\}$ and $\{\bar{1}11\}$ sectors show correlated variations in Mg# $_{cpx}$, indicating that they grew concurrently. When clinopyroxene compositions are converted into clinopyroxene components following Putirka (2008), it becomes clear that low-BSE-intensity $\{\bar{1}11\}$ hourglass sectors are considerably depleted in DiHd and CaTs components with respect to high-BSE-intensity $\{hk0\}$ prism sector zones. Jadeite component contents are also generally lower in $\{\bar{1}11\}$ sectors. This sense of compositional variability shares

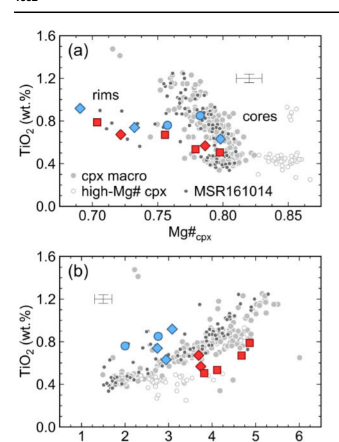


Fig. 7. Summary of natural and experimental clinopyroxene (cpx) compositions from the 2014–2015 Holuhraun lava. Clinopyroxene macrocryst compositions from Halldórsson et al. (2018) are shown as light grey circles. High-Mg#_{cpx} clinopyroxene macrocrysts [where Mg#_{cpx} = Mg/(Mg+Fe) on a molar basis where all Fe is assumed to be Fe²+] are shown as outlined white circles. Compositions from sample MSR141016 are shown as dark grey circles. Mean experimental clinopyroxene compositions are shown using the same symbols as in Fig. 5; see Fig. 10 for more detailed information about variability in experimental clinopyroxene compositions. Core and rim compositions are annotated. Characteristic 2σ analytical uncertainties are shown.

 Al_2O_3 (wt.%)

some similarities with recently described variability in Etnean clinopyroxenes that can be expressed in terms of exchange between $\{hk0\}$ and $\{\bar{1}11\}$ sectors (Ubide *et al.*, 2019):

$$\begin{split} \left[\text{Si}^{4+} + \text{Fe}^{2+} + \text{Mg}^{2+}\right]_{\left\{\bar{1}11\right\}} &\rightleftharpoons \\ \left[\text{Ti}^{4+} + \text{AI}^{3+} + \text{Fe}^{3+}\right]_{\left\{\hbar k0\right\}} \,. \end{split} \tag{1}$$

Moreover, the incorporation of Na and Ti into the $\{hk0\}$ sectors is consistent with high CaTs component contents facilitating complex coupled substitutions (Bennett *et al.*, 2004):

$$\label{eq:califold} \text{Na}^+ + \text{Ti}^{4+} + \text{CaAIAISiO}_6 {\rightleftharpoons} \text{Ca}^{2+} + \text{AI}^{3+} + \text{NaTiAISiO}_6. \tag{2}$$

However, we note that clinopyroxenes from the tholeiitic 2014–2015 Holuhraun lava also differ from those

described in recent contributions on alkalic systems (cf. Welsch *et al.*, 2016; Ubide *et al.*, 2019). Specifically, sector zoning is well developed in Ca as well as in Ti, Al and Na such that $\{\bar{1}11\}$ hourglass sectors are not only depleted in non-quadrilateral components but in DiHd component as well. In other words, $\{\bar{1}11\}$ hourglass sectors appear to be enriched in EnFs component at the expense of all other components. In line with previous observations (Nakamura, 1973; Neave & Putirka, 2017), we propose that clinopyroxene sector zoning in tholeiitic systems can thus be generalised using a modified version of equation (1):

$$\left[Si^{4+} + Fe^{2+} + Mg^{2+}\right]_{\{\overline{1}11\}} \rightleftharpoons$$

$$\left[Ti^{4+} + AI^{3+} + Fe^{3+} + Ca^{2+} + Na^{+}\right]_{\{hk0\}}$$
(3)

Furthermore, compositional differences between $\{hk0\}$ and $\{\bar{1}11\}$ sectors are generally larger than those within individual sectors, suggesting that only one family of sectors is likely to have faithfully recorded the conditions of clinopyroxene formation (e.g. Welsch *et al.*, 2016).

Experimental samples

Phase proportions, H_2O contents and fO_2 conditions

Experimental phase proportions were estimated by mass balance, including metallic Fe as a phase in order to evaluate the degree of Fe exchange with capsule materials. Melt fraction decreases steadily as a function of decreasing T for all combinations of P and capsule type (Table 1), though experiments performed in AuPd capsules have higher melt fractions than those performed in graphite-Pt double capsules at the same P-T conditions. With the exception of 100 MPa experiments in graphite-Pt double capsules, clinopyroxene was the liquidus phase in all experimental series in which crystals formed; no crystals grew in AuPdcapsule experiments at 100 MPa. Fe exchange between experimental samples and capsule materials was generally minimal: only two experiments in AuPd capsules experienced >10 % relative losses of Fe. No Fe was lost from experiments in graphite-Pt double capsules (Table 1).

Experimental redox conditions were verified using a range of approaches (Table 1). Experiments in graphite-Pt double capsules were theoretically buffered at or below CCO (i.e. at or below \sim QFM-1-7; Holloway *et al.*, 1992; Jakobsson & Óskarsson, 1994). H₂O contents of super-liquidus glasses (0·04–0·08 wt % H₂O; Table 1) are consistent with reducing conditions of \sim QFM-2·7 and \sim QFM-2·1 at 100 and 300 MPa, respectively, according to the relationship between aH_2O and fO_2 in closed-capsule IHPV (Botcharnikov *et al.*, 2005). Glass Fe $^{3+}/\Sigma$ Fe values in olivine-bearing runs estimated using $K_{Dol-liq}^{Mg-Fe}$ values calculated with the model of Toplis (2005) indicate slightly more oxidizing conditions of QFM-0·7 to QFM-0·9 and QFM-1·0 at 100 and 300 MPa, respectively,

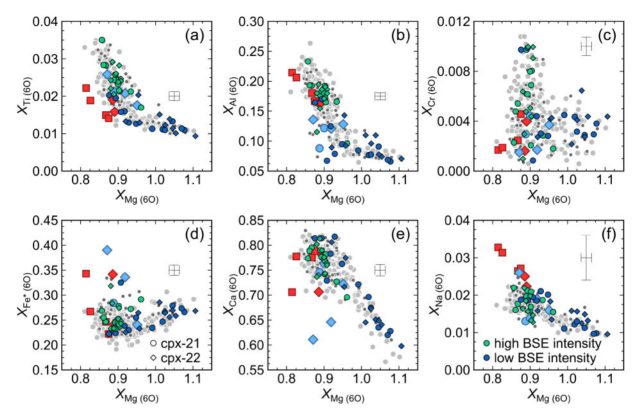


Fig. 8. Summary of natural and experimental clinopyroxene (cpx) major element compositions from the 2014–2015 Holuhraun lava. Compositions are expressed as cations calculated on a 6-oxygen basis (e.g. Putirka, 2008). Spot analyses from compositional profiles through cpx-21 and cpx-22 (profiles A–A' and A–A'', and B–B' and B–B'', respectively; Fig. 3) are shown with circles and diamonds, respectively. Low- and high-BSE-intensity sector zones are shown with blue and turquoise symbols, respectively. All Fe is assumed to be Fe²⁺ (i.e. Fe*). Mean experimental compositions (n= 16–46) are shown using the same symbols as in Fig. 5; see Fig. 10 for more detailed information. Characteristic 2 σ analytical uncertainties are shown.

when converted into fO_2 with the model of Kress & Carmichael (1991). However, both aH_2O - and olivine-based methods of fO_2 estimation have uncertainties of at least half a log unit and, in the case of the latter, even small overestimations of $K_{D_0I-liq}^{Mg-Fe}$ (e.g. by 0.01) can inflate fO_2 estimates by more than a log unit. It is thus likely that our graphite-Pt-double-capsule experiments were successfully buffered close to CCO.

Based on previous work, we expected super-liquidus experiments in AuPd capsules to be buffered close to QFM + 1 given that \sim 0.5 wt % H₂O is typically produced at high-P, high-T conditions with this capsule design (Botcharnikov et al., 2005; Husen et al., 2016). Using the approach described above for graphite-Pt double-capsule experiments, super-liquidus glass H₂O contents of 0.50-1.05 wt % (100 MPa) and 0.64-0.68 wt % (300 MPa), generated by the reduction of Fe₂O₃ in the starting glass powder, indicate that slightly more oxidizing experimental fO_2 conditions of QFM + 1.5 to QFM + 2.7 and QFM + 1.3 to QFM + 1.4 were achieved in AuPd capsules at 100 and 300 MPa, respectively (Table 1). Converting glass $Fe^{3+}/\Sigma Fe$ values determined by colorimetry into fO2 with the model of Kress & Carmichael (1991) returns similar fO₂ estimates: super-liquidus glass Fe³⁺/ ΣFe values of 0.23-0.24 and 0.23-0.25 are indicative of

 fO_2 conditions of QFM + 1.3 to QFM + 1.5 and QFM + 1.5 to QFM + 1.8 at 100 and 300 MPa, respectively. Oxygen fugacities were also estimated from capsule compositions (e.g. Grove, 1981), allowing us to evaluate the conditions prevailing during sub-liquidus experiments. However, redox conditions estimated for super-liquidus experiments with the AuPdFe solution model of Barr & Grove (2010) are systematically \sim 0.6 log units lower than those obtained from colorimetric analyses. Given the coherence between fO2 conditions estimated from glass H_2O and $Fe^{3+}/\Sigma Fe$ values, we conclude that superliquidus experiments were performed close to QFM + 1.5 and that the model of Barr & Grove (2010) underestimates fO2 at the conditions we investigated, possibly because it was calibrated on lower-Fe systems than the 2014-2015 Holuhraun lava. Capsule compositions nevertheless suggest that low-F experiments may have attained fO2 conditions about half a log unit higher than those at super-liquidus conditions, consistent with the crystallization of anhydrous phases resulting in higher residual glass H₂O concentrations (e.g. Whitaker et al., 2007). Overall, we estimate that most of our AuPd-capsule experiments equilibrated at \sim QFM + 1.5 under slightly more oxidizing conditions than initially planned.

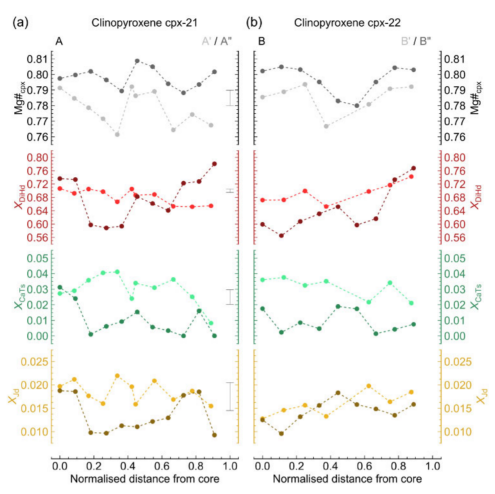


Fig. 9. Compositional profiles through pairs of sector zones from the 2014–2015 Holuhraun lava. Low-BSE intensity sectors are shown with dark symbols and high-BSE intensity sectors with light ones. $Mg\#_{cpx}$ is Mg/(Mg+Fe) on a molar basis where all Fe is assumed to be Fe^{2+} . Diopside–hedenbergite (DiHd), Ca-Tschermak's (CaTs) and jadeite (Jd) component contents were calculated following Putirka (2008). Characteristic 2σ analytical uncertainties are shown. (a) Profiles through low- and high-BSE intensity sector zones in cpx-21 (profiles A–A' and A–A'', respectively; Fig. 3a). (b) Profiles through low- and high-BSE intensity sector zones in cpx-22 (profiles B–B' and B–B'', respectively; Fig. 3b).

Glass, olivine and plagioclase compositions

Experimental glass compositions are shown in Fig. 5 alongside published matrix glass and melt inclusion compositions (Halldórsson et al., 2018; Hartley et al., 2018). Compositional variability in glass analyses from individual capsules is comparable with estimates of analytical uncertainty, indicating that equilibrium was equally well approached throughout our experimental samples (Supplementary Data; supplementary data are available for downloading at http://www.petrology. oxfordjournals.org). Down-T compositional trends defined by experimental glasses depend strongly on P and capsule type. In general, low-P experiments reproduce natural matrix glass compositions better then high-P experiments, with the notable exception that no crystals were produced in AuPd-capsule experiments at 100 MPa. Experiments in graphite-Pt double capsules also replicate natural compositions better than those in AuPd capsules: glasses produced in AuPd capsules at 300 and 600 MPa are enriched in Al₂O₃ with respect to natural compositions, whereas those produced in graphite-Pt double capsules at 100 and 300 MPa are not (Fig. 5a and e). Magmatic liquid compositions (i.e. matrix glasses with >6 wt % MgO) were approximated best by experiments at >1160 °C.

Experimental olivine and plagioclase compositions are shown in Fig. 6 alongside published compositions (Halldórsson et al., 2018). As for experimental glasses, olivine and plagioclase crystals from individual capsules exhibit generally limited compositional variability (Supplementary Data; supplementary data are available for downloading at http://www.petrology.oxford journals.org). There are, however, two exceptions. Firstly, some plagioclase crystals formed at 600 MPa and 1140°C in AuPd capsules are compositionally variable, possibly as a result of disequilibrium crystallization. Secondly, some plagioclase analyses return variably elevated FeO* contents as a result of secondary fluorescence from surrounding Fe-rich glasses polished plagioclase crystals are often thinner than the excitation volume of typical EPMA spots (i.e. $<2 \mu m$).

Olivine was only produced during lower-T runs in graphite-Pt double capsules (Fig. 4a); no olivine formed in AuPd capsule experiments. Although natural macrocryst cores are within the range of compositions predicted to be in equilibrium with the starting glass powder at experimental P-fO $_2$ conditions according to the $K_{D_{\text{ol-liq}}}^{\text{Mg-Fe}}$ model of Toplis (2005) ($X_{\text{Fo}} = 0.74$ -0.79), experimentally produced olivines are typically more evolved and have X_{Fo} contents comparable with natural macrocryst rims ($X_{\text{Fo}} \sim 0.64$ -0.72). Olivines produced at 100 MPa are more similar to natural olivine cores than those produced at 300 MPa (Fig. 6a).

Although plagioclase crystals were produced under diverse experimental conditions (Fig. 6b), they exhibit only modest compositional variability ($X_{An} = 0.61-$ 0.69). Plagioclase stability nonetheless depends strongly on P in AuPd capsule experiments, with plagioclase joining the crystallizing assemblage at >1180°C, >1140°C and <1140°C at 600, 300 and 100 MPa, respectively. This probably reflects the effect of small amounts of H₂O on plagioclase stability (Table 1; Almeev et al., 2012). No comparable P dependence was observed in graphite-Pt double-capsule experiments: plagioclase was stable at >1160 °C at both 100 and 300 MPa, reflecting the uniformly low H₂O content of these runs (Supplementary Data; supplementary data are available for downloading at http://www.petrology. oxfordjournals.org; Almeev et al., 2012). According to the model of Namur et al. (2012), the starting glass powder would be in equilibrium with plagioclase compositions similar to those observed in microcrysts or evolved macrocryst cores from natural samples ($X_{\rm An} \sim$ 0.75; Fig. 6b; Halldórsson et al., 2018). However, most experimental plagioclase crystals are more evolved than predicted compositions, and are instead comparable with natural macrocryst rims (Fig. 6b). Plagioclase crystals produced at 1160 °C and 100 MPa in graphite-Pt double capsules most closely approach microcryst and evolved macrocryst core compositions ($X_{An} \sim 0.68$; Fig. 5b; Halldórsson et al., 2018).

Clinopyroxene compositions

With the exception of experiments performed at 100 MPa in graphite-Pt double capsules, clinopyroxene was the liquidus phase for all combinations of P and fO_2 under which crystals formed (Figs 4b, c, 7 and 8). Mean clinopyroxene compositions span the Mg#cpx range 0.69-0.80. The highest-Mg experimental clinopyroxenes have Mg#cpx contents comparable to those in natural macrocryst cores (Mg# $_{cpx} \sim$ 0.80; Halldórsson et al., 2018). In contrast, the lowest-Mg clinopyroxenes have similar compositions to macrocryst rims (Mg# $_{cox}$ \sim 70). However, simple comparisons between natural and experimental clinopyroxenes are otherwise impeded by the complex dependencies of clinopyroxene compositions on P, T and fO₂. For example, mean clinopyroxene Na contents depend on P (e.g. Blundy et al., 1995; Putirka et al., 1996), with crystals produced at 100 and

600 MPa containing, respectively, less and more Na than natural crystals (Fig. 8f); natural Na contents are best approximated by the products of 300 MPa experiments at 1180 °C. In contrast, mean clinopyroxene Ca and Fe contents appear to be more sensitive to T (or F) than P: crystals formed during our lowest-F runs, i.e. those from low-T runs in graphite-Pt double capsules at 300 MPa and AuPd capsules at 600 MPa, have lower Ca but higher Fe contents than natural macrocryst cores (Fig. 8d and e). Finally, clinopyroxenes from graphite-Pt double-capsule experiments are consistently richer in Ti but poorer in Al than those from AuPd capsule experiments performed under similar conditions (Fig. 8a and b). Overall, mean experimental clinopyroxene compositions lie closer to natural clinopyroxene analyses from $\{hk0\}$ prism sectors than $\{111\}$ hourglass sectors. However, mean compositions provide only partial descriptions of the true chemical complexity present in experimental clinopyroxenes.

Although phase equilibria studies on tholeiitic systems have generally reported mineral compositions as mean values of multiple spot analyses, it has been noted for some decades that clinopyroxene compositions are typically more heterogeneous than can be explained by analytical uncertainty alone (e.g. Grove & Bryan, 1983; Grove et al., 1992; Villiger et al., 2004; Berndt et al., 2005; Ulmer et al., 2018). We, therefore, report all the spot analyses we made on clinopyroxenes (n=16-46 per sample) in order to document the full compositional diversity present in our experimental samples (Fig. 10 and Supplementary Data; supplementary data are available for downloading at http://www. petrology.oxfordjournals.org). For example, clinopyroxene Al and Ca contents are much more variable than corresponding analytical uncertainties, regardless of experimental conditions (Fig. 10a-f). In the case of the 300 MPa experiment carried out in a graphite-Pt double capsule at 1180 °C, the range of experimental clinopyroxene compositions is broadly comparable with that observed in natural samples (Fig. 10c, f and i). Clinopyroxenes from the lowest-F experiment performed in a graphite-Pt double capsule at 300 MPa and 1140 °C show the greatest compositional variability (expressed as a mean-normalized standard deviation) in geobarometry-critical Na contents (Fig. 11a). Moreover, clinopyroxenes from graphite-Pt double-capsule experiments have consistently more variable Al₂O₃ contents than those produced in AuPd capsule experiments (Fig. 11b).

Mean clinopyroxene Fe³⁺ contents and Fe³⁺/ Σ Fe values estimated using the approach of Lindsley (1983) are shown in Fig. 12. Although Fe³⁺/ Σ Fe values estimated by stoichiometry can be subject to absolute uncertainties of up to 25% (e.g. McGuire *et al.*, 1989), it was still possible for us to resolve compositional differences between clinopyroxenes produced under different P-fO₂ conditions. Clinopyroxenes from experiments performed under reducing conditions in graphite-Pt double capsules contain no consistently resolvable Fe³⁺,

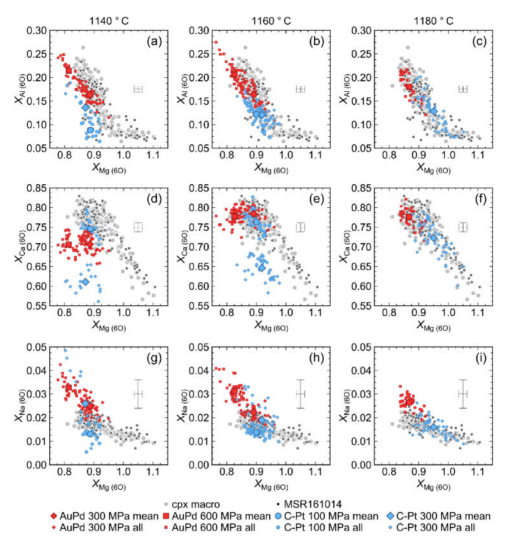


Fig. 10. Experimental clinopyroxene compositions from different run temperatures. Mean clinopyroxene compositions from each run are shown using the same symbols as in Fig. 5. Individual spot analyses from clinopyroxene crystals are shown with smaller symbols. Data are shown for the following temperatures: 1140 °C (a, d and g), 1160 °C (b, e and h) and 1180 °C (c, f and i). Natural data are shown using the same symbols as in Fig. 7. Characteristic 2σ analytical uncertainties are shown.

resulting in estimated Fe³⁺/ Σ Fe values of 0.00 \pm 0.04(1 σ). In contrast, clinopyroxenes from AuPd capsule experiments performed under moderately oxidizing conditions contain appreciable amounts of Fe³⁺: clinopyroxenes produced at 300 MPa return estimated Fe³⁺/ Σ Fe values of 0.18 \pm 0.01(1 σ), whereas those produced at 600 MPa return slightly lower Fe³⁺/ Σ Fe values of 0.10 \pm 0.02(1 σ). For comparison, we estimate that natural clinopyroxenes from sample MSR161014 have Fe³⁺/ Σ Fe values of approximately 0.05–0.25 using the same approach (Fig. 12b).

DISCUSSION

Clinopyroxene in phase equilibria experiments

Compositional variability and sector zoning Experimental clinopyroxenes produced in this study are often too small for zoning to be resolved clearly, even by FEG-SEM (Fig. 4). Nevertheless, many experimental clinopyroxenes show heterogeneous BSE intensities. Although some crystals show hints of concentric zoning, potentially formed during thermal cycling (Fig. 4a and c), patchy zoning is more pervasive (Fig. 4a and d), suggesting that sector zoning is mainly responsible for the chemical systematics we observe rather than concentric zoning (e.g. Ulmer et al., 2018). Moreover, the compositional variability within clinopyroxenes from individual experiments is more consistent with sector zoning than concentric zoning (Nakamura, 1973; Ubide et al., 2019): compositional arrays lie between two endmembers that are Mg-poor, but Al-rich (i.e. rich in nonquadrilateral components) on the one hand and Mgrich, but Al-poor (i.e. rich in quadrilateral components), on the other (Fig. 10). Clinopyroxene compositions from some high-F experiments (e.g. 300 MPa and 1180 °C in a graphite-Pt double capsule) also occupy a similar region of geochemical space to natural crystals with welldeveloped sector zoning (Figs 8 and 10). Unfortunately, the exact nature of clinopyroxene variability in our

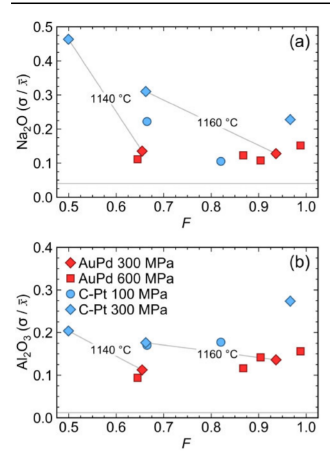


Fig. 11. Mean-normalized standard deviations (σ/\bar{x}) of clinopyroxene (a) Na₂O and (b) Al₂O₃ contents as functions of melt fraction (F). Data are shown using the same symbols as in Fig. 5. Grey horizontal lines show variability in the Kakanui augite standard (NMNH 122142) as an indication of analytical uncertainties. Tie lines join 300 MPa experiments performed at the same temperatures.

experimental samples remains unclear. Do the compositional arrays observed reflect complex internal zoning, or do they represent variably mixed analyses of otherwise compositionally distinct zones? It is nevertheless certain that large numbers of spot analyses (>20 and ideally >40) are required to meaningfully characterize experimental clinopyroxenes.

Disequilibrium in nominally equilibrium experiments

Dynamic crystallization experiments (i.e. those performed under variable *T* conditions) demonstrate that clinopyroxene compositions depend strongly on the degree of undercooling they experience during crystallization, as well as on absolute *P-T-X* conditions (Grove & Bence, 1977; Lofgren *et al.*, 2006; Mollo *et al.*, 2018). Specifically, EnFs, CaTs and Jd component contents in augitic clinopyroxenes increase at the expense of DiHd component contents with increasing undercooling (Mollo *et al.*, 2010). This deviation of mean clinopyroxene compositions away from equilibrium values has important implications for geothermobarometry: calculations performed on crystals grown at high degrees of

undercooling overestimate true P-T conditions because of the incorporation of excess Jd component (Mollo et al., 2010, 2013). Although Mollo et al. (2010) did not report compositional zoning in the products of their dycrystallization experiments, others have described strong correlations between the degree of nominal undercooling (i.e. the difference between the liquidus T and the experimental T, ΔT) and the intensity of compositional zoning in isothermal crystallization experiments. For example, Kouchi et al. (1983) observed that {111} sectors in Al-rich clinopyroxenes grown at $13^{\circ}C < \Delta T < 25^{\circ}C$ are depleted in Al with respect to $\{hk0\}$ sectors. At $\Delta T > 25$ °C they observed that the sense of compositional zoning reversed, highlighting clinopyroxene's sensitivity to undercooling. Moreover, their observations indicate that the incorporation of EnFs, CaTs and Jd components during disequilibrium growth of clinopyroxene may partially reflect an intensification of sector zoning rather than a simple shift in bulk composition.

Our experiments were designed to reproduce equilibrium crystallization conditions. That is, after using low amplitude (±10°C) thermal cycling to promote the growth of large crystals (Erdmann & Koepke, 2016), experiments were held at their final target T until quenching. However, both the texture and compositional variability of experimental clinopyroxenes depend on T and F, implying that they reflect the deviation of T below the clinopyroxene liquidus T (i.e. the degree of undercooling sensu lato) experienced at the onset of crystallization. For example, small numbers of large clinopyroxene crystals were produced in high-T, high-F experiments (Fig. 4c), whereas numerous small crystals were produced in low-T, low-F experiments (Fig. 4d), a relationship consistent with changes in the relative influences of nucleation and growth with increasing undercooling (Kirkpatrick, 1976; Cashman, 1993; Hammer, 2008; Mollo & Hammer, 2017). That is, nucleation appears to have dominated over growth in experiments performed well below the clinopyroxene liquidus (low-F) when thermodynamic barriers to nucleation could be readily overcome, though competition for chemical nutrients may have also contributed towards restricting crystal sizes (Mollo & Hammer, 2017). In contrast, when experiments were performed close to the clinopyroxene liquidus (high-F), growth dominated over nucleation. Importantly, the same magnitude of thermal cycling was applied to all samples, meaning that even if cycling affected the details of crystal nucleation and growth in individual experiments, it is unlikely to account for systematic differences in clinopyroxene properties as a function of T or F. Indeed, other studies reporting compositional heterogeneity in experimental clinopyroxenes from tholeiitic systems did not apply thermal cycling (e.g. Grove et al., 1992; Berndt et al., 2005).

The compositional variability in clinopyroxenes produced in our lowest-*F* experiment performed in a graphite-Pt double capsule at 300 MPa and 1140 °C

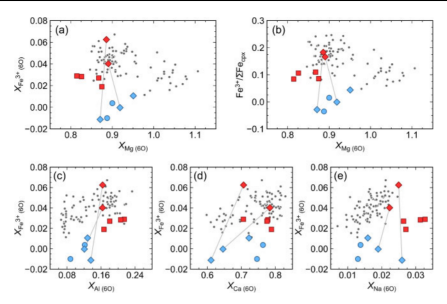


Fig. 12. Ferric iron systematics in natural and experimental clinopyroxene crystals. (a) Mean experimental clinopyroxene Fe³⁺ contents estimated following Lindsley (1983) as a function of Mg content. Data are shown using the same symbols as in Fig. 5. (b) Mean clinopyroxene Fe³⁺/ΣFe values as a function of Mg content. (c and d) Mean clinopyroxene Fe³⁺ contents as functions of (c) AI, (d) Ca and (e) Na contents. Tie lines join 300 MPa experiments performed at the same temperature.

 $(F \sim 0.5)$ highlights that clinopyroxenes from so-called equilibrium experiments can sometimes preserve strongly disequilibrium compositions (Fig. 11). How ever, highly variable compositions are restricted to nominally dry experiments in graphite-Pt double capsules. Given that H₂O acts to reduce chemical disequilibrium around growing crystals by increasing elemental diffusivities, decreasing melt viscosities and reducing the degree of undercooling by depressing liquidi (Hammer, 2008), we suggest that the presence of small amounts of H₂O can explain the differences in clinopyroxene compositional variability we observe. For example, the presence of 1 wt % H₂O may increase Si and Mg diffusivities in silicate melts by factors of approximately three and five, respectively (Zhang et al., 2010). We thus propose that the 0.5-1.0 wt % H₂O present in the melts of our AuPd capsule experiments was sufficient to prevent the crystallization of disequilibrium clinopyroxene compositions like those observed in the products of nominally anhydrous graphite-Pt double capsule experiments $(\sim 0.1 \text{ wt } \% \text{ H}_2\text{O}; \text{ Fig. 11})$ by allowing chemical nutrients to be delivered to growing crystal faces at sufficiently fast rates to maintain near-equilibrium conditions.

Oxygen fugacity and clinopyroxene $Fe^{3+}/\Sigma Fe$ values

Experiments in graphite-Pt double capsules at 300 MPa are thought to have attained fO_2 conditions close to the CCO buffer—Fe³⁺/ Σ Fe \sim 0.08 according to calculations following Kress & Carmichael (1991)—within a few hours (e.g. Médard *et al.*, 2008). Equivalent calculations for 300 MPa AuPd capsule experiments buffered at approximately QFM + 1.5 predict melt Fe³⁺/ Σ Fe values of \sim 0.25 that are consistent with colorimetric analyses of

super-liquidus experiments (Table 1). This difference in melt $Fe^{3+}/\Sigma Fe$ values is also reflected in clinopyroxene $Fe^{3+}/\Sigma Fe$ values estimated by stoichiometry (Lindsley, 1983), which are $0.00 \pm 0.04(1\sigma)$ for 300 MPa experiments in graphite-Pt double capsules $0.18 \pm 0.01(1\sigma)$ for 300 MPa experiments in AuPd capsules (Fig. 12a and b). Ferric iron is thus an important constituent of clinopyroxenes formed at P-T-fO2 conditions relevant to the evolution of natural tholeiites; simply assuming that all Fe in clinopyroxenes occurs as Fe²⁺ is invalid for tholeiitic systems, a situation likely to be even worse for oxidized alkalic and calc-alkaline systems (e.g. Kelley & Cottrell, 2012; Longpré et al., 2017).

High-Fe³⁺ clinopyroxenes produced in AuPd capsules are richer in Al and Ca than low-Fe³⁺ clinopyroxenes produced in graphite-Pt double capsules under the same P-T conditions (Fig. 12c and d). Although plagioclase co-crystallization affects clinopyroxene compositions to some degree, comparing clinopyroxenebearing experiments performed at 300 MPa and similar F conditions to mitigate the effects of other phases reveals similar overall trends. In contrast, there is no clear relationship between redox conditions and clinopyroxene Na contents in our experimental data (Fig. 12e). Within the range of investigated fO2 conditions (approximately CCO to QFM + 1.5), our results thus suggest that Fe3+ is incorporated into augitic clinopyroxene alongside Ca and Al as Ca-Fe-Tschermak's component (CaFeTs; CaFe³⁺AlSiO₆; Luth & Canil, 1993; Blundy et al., 1995; Mollo et al., 2018). We find no evidence for the significant incorporation of Fe³⁺ as the Na-bearing Ae component. Therefore, assuming that all Na in clinopyroxene is present as the Jd component appears to be valid in tholeiitic magmas as oxidized as QFM + 1.5. Whether this assumption still holds in

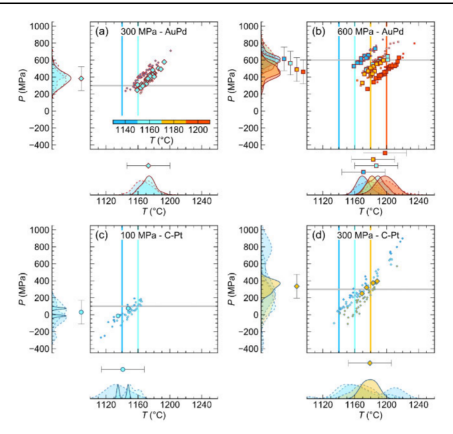


Fig. 13. Pressure (*P*) and temperature (*T*) estimates from experimental clinopyroxene–liquid pairs calculated iteratively with equation 1 of Neave & Putirka (2017) and equation 33 of Putirka (2008). Small symbols show *P*–*T* values calculated from all clinopyroxene analyses. Large symbols show *P*–*T* values calculated from filtered clinopyroxene analyses whose observed diopside-hedenbergite (DiHd), enstatite–ferrosilite (EnFs) and Ca-Tschermak's (CaTs) component contents calculated following Putirka (2008) were within one standard error of estimate (SEE) of values predicted with equations 6 and 7 of Mollo *et al.* (2013) (DiHd and EnFs) and equation 3.4 of Putirka (1999) (CaTs). Kernel density estimates (KDEs) show *P* and *T* distributions calculated for both all and filtered subsets of clinopyroxene analyses (dashed and solid lines respectively). Mean *P* and *T* estimates from filtered clinopyroxene analyses are shown beside the KDEs; error bars indicate model SEEs. (a) Experiments performed in AuPd capsules at 300 MPa. (b) Experiments performed in graphite-Pt (C–Pt) double capsules at 100 MPa. (d) Experiments performed in graphite-Pt double capsules at 300 MPa.

oxidized alkalic systems such as the Canary Islands, where clinopyroxene Fe³⁺/ Σ Fe values can exceed 0.5 (Weis *et al.*, 2015), remains to be addressed.

Geothermobarometry and clinopyroxene-liquid equilibria

If geothermobarometric models are well calibrated, then they should be able to reproduce imposed *P-T* conditions when applied to experimental products. By applying various models to the products of experiments performed at crustal *P* conditions (100–800 MPa), Neave & Putirka (2017) observed that previously published Jdin-clinopyroxene geobarometers consistently overestimated experimental *P* conditions for runs performed at ≤700 MPa. This led them to propose a new geobarometer optimised for mafic and alkali-poor magmas stored at crustal depths. However, they only tested their model using mean compositions of experimentally produced clinopyroxenes that, as demonstrated above, offer only partial descriptions of clinopyroxene compositions. In order to provide a more complete test of the

geothermobarometric approach recommended by Neave & Putirka (2017) — iteratively solving their equation 1 with equation 33 from Putirka (2008) – we applied it to all clinopyroxene–liquid pairs measured in our experimental products (Fig. 13).

For any given set of P-T-fO2 conditions, geothermobarometric calculations on individual clinopyroxene-liguid pairs return positively correlated and broadly distributed P-T arrays, reflecting the interdependence of geobarometric and geothermometric models. In some cases, such as for experiments performed in AuPd capsules at 300 MPa (Fig. 13a), P-T estimates are dispersed to an extent commensurate with the prediction errors of geothermobarometric models: calibrations of equation 1 from Neave & Putirka (2017) and equation 33 from Putirka (2008) are associated with standard errors of estimate (SEE) of ±140 MPa and ±27 °C respectively. In other cases, P-T estimates vary well beyond model SEEs. For example, low-T experiments in graphite-Pt double capsules are associated with large ranges of estimated P-T conditions (Fig. 13c and d). In the most extreme case, clinopyroxene

analyses from the 300 MPa experiment at 1140 °C return bimodal P and T distributions with absolute ranges greater than 100-900 MPa and 1140-1210 °C, respectively (Fig. 13d). The deviation of calculated P-T conditions from imposed conditions correlates with the degree of geochemical variability within clinopyroxene populations. Given that clinopyroxene variability is linked to both the melt H₂O content and the deviation below the clinopyroxene liquidus T, spreads in estimated P-T values are thus linked to the degree of disequilibrium experienced during crystallization (e.g. Kouchi et al., 1983; Mollo et al., 2010). Despite this, mean estimated P-T conditions are almost always within 1SEE of experimental values; all mean P-T conditions are well within 2SEE. Therefore, experiments performed at 100, 300 and 600 MPa can be readily distinguished from each other based on the results of geobarometric calculations. Pressure conditions estimated from AuPd capsule experiments also show no systematic P differences with respect to those performed in graphite-Pt double capsules, lending further credence to our suggestion that Fe³⁺ is incorporated as the CaFeTs component, rather than the Ae component under the conditions studied.

The wide ranges of P-T values estimated from clinopyroxene compositions obtained at fixed experimental conditions reinforce previous suggestions that using only $K_{D_{cpx-liq}}^{Mg-Fe}$ values to check for clinopyroxene-liquid equilibrium in natural rocks is inadequate (Mollo et al., 2010, 2013; Winpenny & Maclennan, 2011; Neave & Putirka, 2017). However, the fact that >65% of clinopyroxene analyses return P values within 1SEE of the imposed conditions does indicate that experimental conditions were recorded faithfully in some cases; the low-F, 300 MPa experiment performed in graphite-Pt double capsules at 1140 °C represents a possible exception to this. The departure of individual clinopyroxene analyses from equilibrium was evaluated using equations 6 and 7 from Mollo et al. (2013) that describe DiHd and EnFs component equilibria, and equation 3.4 from Putirka (1999) that describes CaTs component equilibrium. We note that these descriptions of clinopyroxeneliquid equilibria have been calibrated on mean experimental clinopyroxene compositions and without considering the effects of Fe³⁺. However, they represent the most complete empirical descriptions of clinopyroxene-liquid equilibria currently available and have been validated using experiments performed at low degrees of undercooling that are unlikely to have experienced appreciable disequilibrium crystallization (Mollo et al., 2013).

Tests for DiHd component equilibrium are shown in Fig. 14; equivalent plots for EnFs and CaTs components are provided in the Supplementary Data; supplementary data are available for downloading at http://www.petrology.oxfordjournals.org. Clinopyroxene analyses from most experiments performed in AuPd capsules are close to being in DiHd component equilibrium (Fig. 14a and b), though some analyses from 600 MPa

experiments have DiHd component contents slightly more than 1SEE above predicted values. In contrast, almost all clinopyroxene analyses from graphite-Pt double capsule experiments contain DiHd component contents more than 1SEE below predicted values (Fig. 14c and d). Such deviations are frequently greater than 2SEE in the cases of clinopyroxenes formed at 1140 and 1160 °C. The only experiment in graphite-Pt double capsules to return a meaningful number of clinopyroxene analyses in DiHd component equilibrium with their surrounding melt was performed under high-F conditions at 1180 °C and 300 MPa.

Internally consistent clinopyroxene geothermobarometry

With the aim of developing an internally consistent approach for performing clinopyroxene-liquid geothermobarometry on natural tholeiites, we tested the effect of filtering the results of our geothermobarometric calculations on experimentally produced clinopyroxene-liquid pairs for multicomponent equilibrium (e.g. Hildner et al., 2012; Neave et al., 2013; Neave & Putirka, 2017; Halldórsson et al., 2018; Shane & Coote, 2018). Glass and clinopyroxene compositions were considered to be in equilibrium if observed DiHd, EnFs and CaTs component values were within 1SEE of predicted values (i.e. within ± 0.06 , ± 0.05 and ± 0.03 for DiHd, EnFs and CaTs components, respectively; Putirka, 1999; Mollo et al., 2013). Thresholds for equilibrium were set at 1SEE for two reasons. Firstly, filters based on calibration statistics account for model uncertainties explicitly, whereas filters based on arbitrary thresholds do not. Indeed, some previous studies have over-filtered for CaTs component equilibrium when using arbitrary filters (e.g. Neave & Putirka, 2017). Secondly, adopting a conservative approach (i.e. setting thresholds at 1SEE instead of 2SEE) reduces the risk of returning false positives.

The results of filtering experimental clinopyroxeneliquid pairs are shown alongside unfiltered data in Fig. 13. Two key trends can be identified: firstly, high-F, high-T experiments are more likely to pass equilibrium filters than low-F, low-T experiments; and secondly, experiments in AuPd capsules return a greater number of reliable P-T estimates than experiments in graphite-Pt double capsules. The likelihood of clinopyroxene-liquid pairs passing equilibrium tests thus correlates inversely with melt H₂O content and the deviation from clinopyroxene liquidus T, two properties related to the ability of crystals to crystallize under near-equilibrium conditions (see above; Hammer, 2008). That is, the greatest number of equilibrium pairs are found in experimental products synthesized close to the clinopyroxene liquidus in the presence of >0.5 wt % H₂O. Encouragingly, mean P conditions recorded by clinopyroxene-liquid pairs passing multicomponent equilibrium tests all lie within 1SEE of imposed experimental conditions. Filtered P estimates are much less dispersed than unfiltered P estimates and mean T conditions are

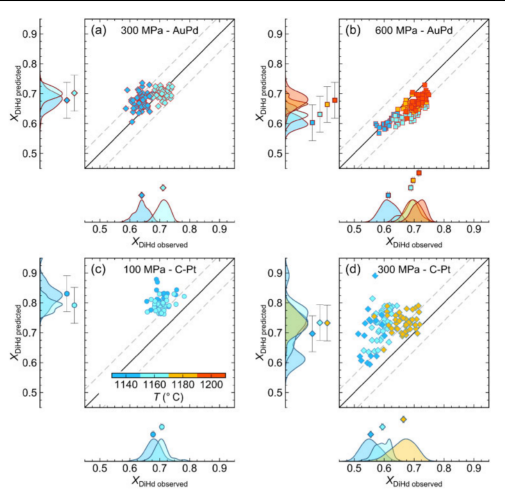


Fig. 14. Comparison of observed and predicted diopside–hedenbergite (DiHd) component contents in experimental clinopyroxenes. Observed DiHd component contents were calculated following Putirka (2008). Predicted DiHd component contents were calculated by iteratively solving equations 6 and 7 of Mollo *et al.* (2013) using the *P-T* conditions presented in Fig. 13. One-to-one relationships between observed and predicted DiHd component contents are shown as solid black lines. Model uncertainties (standard errors of estimate) are shown as envelopes delimited by dashed grey lines. (a) Experiments performed in AuPd capsules at 300 MPa. (b) Experiments performed in AuPd capsules at 100 MPa. (d) Experiments performed in graphite-Pt double capsules at 300 MPa.

mostly within 1SEE of experimental values, especially for higher T experiments. Our findings nonetheless suggest that clinopyroxene geothermometer calibrations could be improved by re-evaluating relationships between T and X in Fe³⁺-bearing systems; obtaining precise estimates of clinopyroxene Fe³⁺/ Σ Fe values will be vital if clinopyroxene geothermobarometry is to be developed further.

Filtering for multicomponent equilibrium thus provides an efficient way of eliminating disequilibrium clinopyroxene compositions from geothermobarometric calculations on tholeitic as well as alkalic samples (cf. Mollo et al., 2013). Importantly, no a priori filtering of clinopyroxene data is required to obtain coherent *P-T* estimates. In other words, there is no need to assume that one family of sector zones records equilibrium crystallization while another does not (e.g. Neave et al., 2015). When performing calculations on natural clinopyroxene-liquid pairs, we therefore recommend checking for equilibrium in DiHd, EnFs and CaTs

components as well as ensuring that $K_{D_{cpx-liq}}^{Mg-Fe}$ values are within a sensible range [i.e. within 0.03 of values from equation 35 of Putirka (2008)]. We include a spreadsheet in the Supplementary Data; supplementary data are available for downloading at http://www.petrology.oxfordjournals.org to facilitate such calculations.

Implications for the 2014–2015 Holuhraun eruption

Geothermobarometry

Clinopyroxene–liquid geothermobarometry was performed on natural clinopyroxenes from MSR161014 using an iterative melt-matching algorithm that refines storage conditions over multiple cycles of equilibrium testing and geothermobarometry, that is described in detail by Neave et al. (2013) and Neave & Putirka (2017). Clinopyroxene compositions were tested for equilibrium against the matrix glass dataset reported by Halldórsson et al. (2018) and were considered to be in

equilibrium if: (1) $K_{D_{cpx-liq}}^{Mg-Fe}$ values were within 0.03 of equilibrium values calculated assuming a melt Fe³⁺/ΣFe value of 0.15 (Putirka, 2008; Bali et al., 2018); and (2) DiHd-, EnFs- and CaTs-component contents were within 1SEE of predicted values (Putirka, 1999; Mollo et al., 2013). The R code used to perform these calculations is available from the authors on request. We also performed geothermobarometric calculations with the spreadsheet provided in the Supplementary Data; supplementary data are available for downloading at http:// www.petrology.oxfordjournals.org in order to test if a simpler approach would yield comparable results to those from the more rigorous melt-matching algorithm. In this case, we paired clinopyroxene compositions with the mean matrix glass composition reported by Halldórsson et al. (2018).

The results of geothermobarometric calculations are summarised in Fig. 15. In the case of calculations with the melt-matching algorithm (Fig. 15a), we report P-T estimates for each clinopyroxene analysis to which equilibrium liquids could be matched (4-274 matches per clinopyroxene; median = 59). When multiple matches were found for individual analyses, we report mean P-T estimates. In the case of spreadsheet calculations (Fig. 15b), we report P-T estimates for clinopyroxene analyses that passed equilibrium tests and distinguish results from clinopyroxene rims (but not outermost rims) that are likely to have been in textural equilibrium with carrier liquids (i.e. matrix glass) during their crystallization (e.g. Klügel et al., 2005). Individual P-T estimates calculated with both methods form tight arrays, confirming that matched clinopyroxenes do indeed represent a single macrocryst population. The reduced scatter in Fig. 15a with respect to that in Fig. 15b results from the use of iterative calculations and is not of geological significance, however. Mean P-T estimates lie in the centre of calculated ranges for both approaches and are almost identical to previous estimates (Halldórsson et al., 2018): 232 MPa and 1161°C for calculations with the melt-matching algorithm and 208 MPa and 1161°C for spreadsheet calculations (Fig. 15). P-T estimates are distributed in proportion with the prediction errors of the geothermobarometric models used (±140 MPa and ±27 °C), albeit with a slight skew towards lower P-T conditions for calculations with the melt-matching algorithm. The only significant difference between the approaches is that the spreadsheet calculations failed to exclude two compositions that returned spurious, negative pressures. Mean P-T estimates from rims in textural equilibrium with their carrier melts are indistinguishable from those calculated using the full dataset. However, in agreement with previous observations (Halldórsson et al., 2018), we note that P-T conditions obtained from these could be less dispersed than those obtained from the dataset as a whole. Combining textural constraints with tests for multicomponent chemical equilibrium is thus likely to return the most precise P-T estimates from volcanic samples.

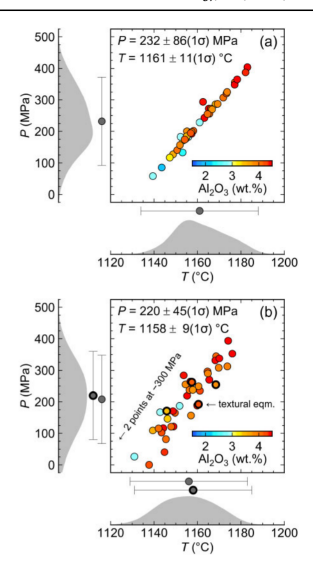


Fig. 15. Pre-eruptive storage conditions of the 2014-2015 Holuhraun lava estimated with natural clinopyroxene macrocrysts from sample MSR141016. Iterative geothermobarometric calculations were performed using equation 1 of Neave & Putirka (2017) and equation 33 of Putirka (2008). (a) Pressure (P) and temperature (T) estimates calculated using an iterative melt-matching algorithm adapted from Neave et al. (2013) and Neave & Putirka (2017). Putative equilibrium melts were sourced from the matrix glass data of Halldórsson et al. (2018). Clinopyroxene-liquid pairs were considered to be in equilibrium when observed diopside-hedenbergite (DiHd), enstatiteferrosilite (EnFs) and Ca-Tschermak's (CaTs) component contents were within one standard error of estimate (SEE) of values predicted with equations 6 and 7 of Mollo et al. (2013) (DiHd and EnFs) and equation 3-4 of Putirka (1999) (CaTs), and observed $K_D^{Mg-Fe}_{cpx-liq}$ values were within 0-03 of values predicted with equation 35 of Putirka (2008), assuming a melt $Fe^{3+}/\Sigma Fe$ value of 0.15 (Bali et al., 2018). Kernel density estimates (KDEs) showing P and T distributions are shown alongside mean P and T values; error bars indicate model SEEs. (b) Storage conditions estimated with the spreadsheet included in the Supplementary Data; supplementary data are available for downloading at http://www.petrology.oxfordjournals.org by pairing clinopyroxene analyses with the mean matrix glass composition of Halldórsson et al. (2018) and applying the same equilibrium criteria used in calculations with the melt-matching algorithm. Results with thick outlines are from analyses located within the rims (but not outermost rims) of clinopyroxene macrocrysts, most likely to have been in textural equilibrium with their carrier liquids at depth (e.g. Klügel et al., 2005).

Clinopyroxene macrocrysts analysed in this study appear to have formed within a narrow P interval centred at ~230 MPa based on calculations using our meltmatching algorithm. That is, the dispersion in estimated P-T values reflects a combination of analytical uncertainties, geothermobarometric uncertainties and disequilibrium crystallization; we see no evidence for polybaric crystallization in clinopyroxene macrocrysts MSR161014. Importantly, the 2014-2015 Holuhraun lava's extreme compositional homogeneity (Halldórsson et al., 2018) suggests that our findings can be justifiably extrapolated to the eruption as a whole. Our refined storage pressure estimates are wholly consistent with independent petrological estimates based on modelling OPAM thermal minima (210 \pm 70(1 σ) MPa; Hartley et al., 2018). Moreover, assuming a mean oceanic crustal density of 2860 kg m⁻³ (Carlson & Herrick, 1990), our preferred storage P estimate of $230 \pm 140(1SEE)$ MPa corresponds to a depth of 8.3 ± 5.0 km, which overlaps with geodetic and seismic signals of magma storage and movement in the Bárðarbunga volcanic system (8-12 km; Sigmundsson et al., 2015; Ágústsdóttir et al., 2016; Guðmundsson et al., 2016).

Geothermobarometric results from the meltmatching algorithm are compared with results from spreadsheet calculations performed without applying equilibrium tests in Fig. 16. Most clinopyroxene analyses for which equilibrium liquids could be found are from {hk0} prism sectors (Figs 3 and 16). When equilibrium liquids could be found for analyses from {111} hourglass sectors, the compositional divergence from analyses in {hk0} prism sectors was minimal. Inspecting unfiltered geothermobarometric results from spreadsheet calculations highlights the effect of disequilibrium crystallization on P estimates: low DiHd component contents in {111} sectors are associated with low Jd component contents, shifting clinopyroxene-liquid Jd component exchange coefficients away from equilibrium, resulting in the calculation of geologically implausible negative pressures. Thus, disequilibrium crystallization has the opposite effect on P estimates from clinopyroxene macrocrysts from the 2014-2015 Holuhraun lava when compared with that predicted from dynamic crystallization studies (Baker & Grove, 1985; Mollo et al., 2010, 2013). Specifically, geobarometric calculations on disequilibrium compositions from our samples are likely to underestimate rather than overestimate true storage pressures. This is because disequilibrium manifests primarily in sector zone development rather than the bulk modification of clinopyroxene compositions typically reported in previous work.

Magma assembly

Clinopyroxene sector zoning theoretically records information about the degree of undercooling experienced during crystallization (e.g. Welsch et al., 2016).

However, too few experimental data are currently available to robustly invert clinopyroxene sector zoning patterns for cooling histories in all but the simplest systems (Kouchi et al., 1983). Nevertheless, comparing the findings of Kouchi et al. (1983) with the compositional systematics observed in our natural clinopyroxenes suggests that crystallization probably occurred at low degrees of undercooling (perhaps on the order of 25 °C or less). It is thus probable that crystallization was driven purely by cooling with no contribution from H₂O exsolution (cf. Applegarth et al., 2013), which is consistent with the magma's modest pre-eruptive H₂O content (<0.6 wt %; Bali et al., 2018): H₂O would have only exsolved during final ascent (<40 MPa; Hartley et al., 2018), too late to affect the crystallization of the clinopyroxene macrocrysts considered here.

Compositional heterogeneity in clinopyroxene macrocrysts is not restricted to the development of sector zoning: concentric zoning within {hk0} sectors is characterized by variability in Ti, Al, Cr and Fe contents at relatively constant Mg contents (Figs 7, 8 and 16). Specifically, some concentric zones are enriched in non-quadrilateral components, consistent with disequilibrium crystallization described by the following kinetically controlled exchange (e.g. Mollo et al., 2013):

$$[Di + Hd]_{cpx} + melt \implies (4)$$

$$[Jd + CaTs + CaFeTs + CaTiTs + En]_{cpx}$$

Given that concentric zoning can be traced through both $\{hk0\}$ and $\{111\}$ sectors (Fig. 3), we suggest that clinopyroxene crystallization took place at low but nonetheless variable degrees of undercooling, leading not only to the development of {111} sectors with disequilibrium compositions, but also to the growth of some concentric zones with disequilibrium compositions, even within $\{hk0\}$ sectors. There are numerous ways that variations in undercooling could have arisen. For example, punctuated recharge of a clinopyroxenesaturated reservoir could have driven cyclical changes in undercooling (Petrone et al., 2016; Ubide & Kamber, 2018). Alternatively, convective stirring within a meltrich storage region, consistent with the extreme homogeneity of erupted liquids (Halldórsson et al., 2018), could have transported clinopyroxene macrocrysts through a range of thermally distinct environments (Couch et al., 2001; Ginibre et al., 2002). Indeed, the euhedral form of many clinopyroxene crystals suggests that they grew in a melt-rich environment rather than within a static crystal mush (e.g. Holness et al., 2019). Therefore, although compositional variability within clinopyroxene macrocrysts complicates geothermobarometric calculations, it does encode information about magma reservoir dynamics.

Clinopyroxene textures, compositions and geother-mobarometry suggest that the magma was stored at $\sim\!1160\,^\circ\text{C}$ prior to eruption, conditions under which olivine, plagioclase and clinopyroxene crystallized simultaneously at the basaltic thermal minimum. The sense

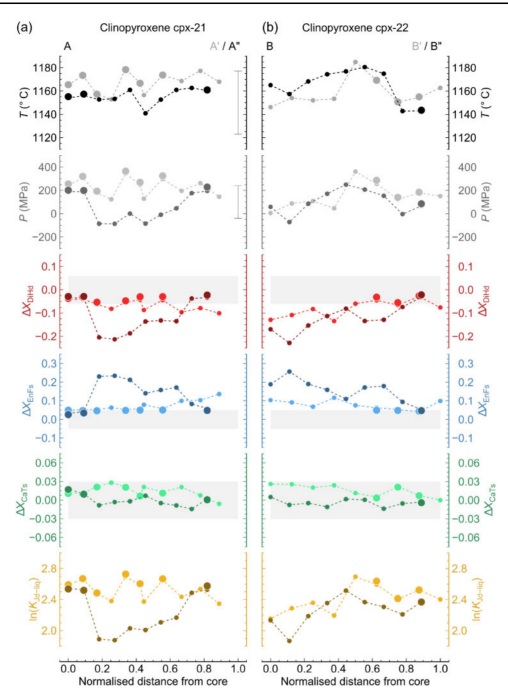


Fig. 16. Results of geothermobarometric calculations performed on sector zone pairs from the 2014–2015 Holuhraun lava. Results from low-BSE intensity sectors are shown with dark symbols and results from high-BSE intensity sectors with light ones. Dashed lines and small circles show results obtained by pairing all clinopyroxene analyses with the mean matrix glass composition from Halldórsson *et al.* (2018) and applying no filtering. Large circles show the results of calculations with an iterative melt-matching algorithm to which equilibrium filtering was applied (Fig. 15a). Deviations from equilibrium are expressed as differences between observed and predicted diopside–hedenbergite (DiHd), enstatite–ferrosilite (EnFs) and Ca-Tschermak's (CaTs) component contents (i.e. $\Delta_{\text{component}}$ values). DiHd and EnFs component contents were predicted with equations 6 and 7 of Mollo *et al.* (2013) and CaTs component contents with equation 3·4 of Putirka (1999). Jadeite-liquid equilibrium constants are also presented as values of $\ln(K_{\text{Jd-liq}})$. (a) Profiles through low- and high-BSE intensity sector zones in cpx-21 (profiles A–A' and A–A' respectively; Fig. 3a). (b) Profiles through low- and high-BSE intensity sector zones in cpx-22 (profiles B–B' and B–B'' respectively; Fig. 3b).

of sector zoning suggests that the degree of undercooling was low (perhaps 25 °C or less) and that the magma could not have cooled from temperatures much above the clinopyroxene liquidus (~1180 °C at ~230 MPa

according to our experiments) upon arrival in the shallow crust. Clinopyroxene macrocrysts nevertheless record fluctuations in storage conditions, resulting in the development of complex concentric zoning that is

mirrored in the oscillatory zoning of cogenetic plagioclase macrocryst rims (Halldórsson et al., 2018). However, our geothermobarometric results and an absence of progressive core-to-rim zoning (with the exception of outermost rims formed during lava emplacement) imply that pre-eruptive storage conditions did not diverge consistently from 230 \pm 140(1SEE) MPa and 1161 \pm 27(1SEE) $^{\circ}$ C during the final stages of storage in the magma reservoir. However, not all macrocrysts carried by the 2014–2015 Holuhraun lava formed at these conditions; primitive macrocryst record earlier phases of evolution in different magmatic environments.

Primitive macrocrysts of high- X_{Fo} olivine, high- X_{An} plagioclase and rare high-Mg#cpx clinopyroxene record crystallization at higher temperatures than the macrocryst assemblage formed immediately prior to eruption (Figs 5, 6 and 7; Halldórsson et al., 2018), However, the relationship between evolved and primitive macrocrysts is not straightforward. Melt inclusions hosted in primitive macrocrysts record the entrapment of compositionally diverse melts that cannot be related to erupted liquids and evolved macrocrysts by differentiation along a simple single liquid line of descent (Hartley et al., 2018). Instead, primitive macrocrysts are likely to have been entrained by melts parental to the erupted magma during their ascent through crystal mushes beneath the Bárðarbunga volcanic system (Halldórsson et al., 2018; Hartley et al., 2018), as has been suggested for other Icelandic systems (Hansen & Grönvold, 2000; Halldórsson et al., 2008; Passmore et al., 2012; Neave et al., 2013, 2014). Moreover, observations on Icelandic lavas suggest that evolved and primitive assemblages are typically juxtaposed within weeks of eruption (Hartley et al., 2015, 2016, 2018; Pankhurst et al., 2018), placing a maximum limit on the time taken to crystallize evolved macrocrysts.

Integrating our new observations with published work enables us to propose the following simplified model for the assembly of the 2014-2015 Holuhraun magma. Firstly, melts directly parental to the erupted liquid segregated efficiently from crustal mushes at pressures in excess of 230 MPa. These melts entrained primitive and geochemically diverse macrocrysts during ascent towards their final storage in a reservoir at ~230 MPa (Hartley et al., 2018). Importantly, it appears that no primitive macrocrysts are unambiguously cogenetic with the erupted melt, meaning that the origins of this melt remain frustratingly obscure (Halldórsson et al., 2018). Mixing during ascent may have facilitated the homogenisation of parental melts that subsequently cooled from temperatures close to the clinopyroxene liquidus (\sim 1180°C) to \sim 1160°C, resulting in the crystallization of olivine, plagioclase and sector-zoned clinopyroxene at low, but variable, degrees of undercooling. This crystallization produced the erupted liquid which ultimately carried two discrete macrocryst assemblages to the surface, an equilibrium assemblage formed at ~230 MPa and ~1160 °C and an entrained assemblage formed at higher pressures and temperatures.

CONCLUSIONS

Natural clinopyroxene macrocrysts from the tholeiitic 2014–2015 Holuhraun lava are characterized by both sector and concentric zoning. In contrast with recent work on alkalic systems (e.g. Welsch *et al.*, 2016; Ubide *et al.*, 2019), we observe that sector zoning is characterized by differences in Ca and Na, as well as in Ti and Al, with {111} hourglass sectors being enriched in the EnFs component with respect to {*hk*0} prism sectors. We therefore caution against extrapolating findings from well-studied, but not universally representative, alkalic systems to geologically more widespread tholeiitic systems.

By quantifying the degree of compositional variability in clinopyroxene crystals produced in experiments on the 2014-2015 Holuhraun lava, we demonstrate that crystals from nominally equilibrium experiments can preserve strongly disequilibrium compositions. The magnitude of clinopyroxene compositional variability in any given experiment depends on both the melt H₂O content and the degree to which the experimental T deviated from the clinopyroxene liquidus T, indicating that clinopyroxene compositions are controlled by kinetic processes within the melt as well as at crystal-melt interfaces. In future, separating the effects of H₂O and major element composition will help to reconcile seemingly divergent observations on clinopyroxene crystallization kinetics in different magma types and make sector zoning an effective probe of crystallization conditions.

Although clinopyroxene $Fe^{3+}/\Sigma Fe$ values estimated by stoichiometry are subject to considerable uncertainties, estimated values up to ~ 0.2 in our natural and experimental samples demonstrate that Fe^{3+} is a major constituent of augitic clinopyroxenes at naturally relevant fO_2 conditions. The compositional systematics of our experimental clinopyroxenes suggest that Fe^{3+} is incorporated as the Ca- and Al-bearing CaFeTs component with minimal incorporation as the Na-bearing Ae component. Therefore, assuming that all Na in clinopyroxene occurs as the Jd component is justified in tholeitic systems when fO_2 is below QFM + 1.5. Elevated fO_2 conditions are thus less likely to compromise Jd-inclinopyroxene geothermobarometry than previously envisaged (Neave & Putirka, 2017).

By combining geothermobarometric models with empirical descriptions of clinopyroxene–liquid equilibria we present an internally consistent and experimentally verified method for performing Jd-inclinopyroxene geothermobarometry. Our approach avoids having to select equilibrium zones *a priori* by performing geothermobarometry and filtering for multicomponent equilibrium concurrently. Applying our approach to natural samples from the 2014–2015 Holuhraun lava returns *P-T* estimates in excellent

agreement with published values from numerous independent methods: $232 \pm 140(1SEE)$ MPa and $1161 \pm 27(1SEE)$ °C. Our findings illustrate how reliable estimates of storage conditions can be obtained from complex clinopyroxene crystals carried by tholeitic magmas when sufficiently robust equilibrium filtering is undertaken.

ACKNOWLEDGEMENTS

We thank the 2014–2015 Holuhraun Eruption Team (Institute of Earth Sciences, University of Iceland) for sampling during the eruption. J. Feige prepared samples, R. Balzer and S.A. Linsler assisted with IHPV experiments, and R.A. Almeev and L.A. Fischer assisted with EPMA. We thank S. Mollo and A. Klügel and one anonymous reviewer for their detailed and constructive reviews that greatly improved our manuscript. We also thank V. Troll and M. Wilson for their editorial handling. Most of the 300 MPa experiments initially formed part of an MSc thesis (Hannover) undertaken by A.-S.S. under the supervision of D.A.N. and F.H.

FUNDING

This work and D.A.N. were supported by the Alexander von Humboldt Foundation, the German Research Foundation (NE2097/1–1) and a Presidential Fellowship from the University of Manchester. The involvement of S.A.H. was partly in relation to H2020 project EUROVOLC, funded by the European Commission (Grant 731070).

SUPPLEMENTARY DATA

Supplementary data are available at *Journal of Petrology* online.

REFERENCES

- Ágústsdóttir, T., Woods, J., Greenfield, T., Green, R. G., White, R. S., Winder, T., Brandsdóttir, B., Steinthórsson, S. & Soosalu, H. (2016). Episodic propagation of the 2014 Bárðarbunga-Holuhraun dike intrusion, central Iceland. *Geophysical Research Letters* **43**, 1495–1499.
- Almeev, R. R., Holtz, F., Koepke, J. & Parat, F. (2012). Experimental calibration of the effect of H_2O on plagioclase crystallization in basaltic melt at 200 MPa. *American Mineralogist* **97**, 1234–1240.
- Applegarth, L. J., Tuffen, H., James, M. R., Pinkerton, H. & Cashman, K. V. (2013). Direct observations of degassing-induced crystallization in basalts. *Geology* 41, 243–246.
- Baker, M. B. & Grove, T. L. (1985). Kinetic controls on pyroxene nucleation and metastable liquid lines of descent in a basaltic andesite. *American Mineralogist* 70, 279–287.
- Bali, E., Hartley, M. E., Halldórsson, S. A., Guðfinnsson, G. H. & Jakobsson, S. (2018). Melt inclusion constraints on volatile systematics and degassing history of the 2014–2015 Holuhraun eruption, Iceland. *Contributions to Mineralogy and Petrology* 173, 9.

- Balta, J. B., Beckett, J. R. & Asimow, P. D. (2011). Thermodynamic properties of alloys of gold-74/palladium-26 with variable amounts of iron and the use of Au-Pd-Fe alloys as containers for experimental petrology. *American Mineralogist* 96, 1467–1474.
- Barr, J. A. & Grove, T. L. (2010). AuPdFe ternary solution model and applications to understanding the fO₂ of hydrous, high-pressure experiments. *Contributions to Mineralogy and Petrology* **160**, 631–643.
- Bennett, S. L., Blundy, J. & Elliott, T. (2004). The effect of sodium and titanium on crystal-melt partitioning of trace elements. *Geochimica et Cosmochimica Acta* **68**, 2335–2347.
- Berndt, J., Koepke, J. & Holtz, F. (2005). An experimental investigation of the influence of water and oxygen fugacity on differentiation of MORB at 200 MPa. *Journal of Petrology* 46, 135–167.
- Berndt, J., Liebske, C., Holtz, F., Freise, M., Nowak, M., Ziegenbein, D., Hurkuck, W. & Koepke, J. (2002). A combined rapid-quench and H₂-membrane setup for internally heated pressure vessels: description and application for water solubility in basaltic melts. *American Mineralogist* 87, 1717–1726.
- Blundy, J. D., Falloon, T. J., Wood, B. J. & Dalton, J. A. (1995). Sodium partitioning between clinopyroxene and silicate melts. *Journal of Geophysical Research: Solid Earth* **100**, 15501–15515.
- Bonny, E., Thordarson, T., Wright, R., Höskuldsson, A. & Jónsdóttir, I. (2018). The volume of lava erupted during the 2014 to 2015 eruption at Holuhraun, Iceland: a comparison between satellite- and ground-based measurements. *Journal of Geophysical Research: Solid Earth* 123, 1–15.
- Borisov, A., Behrens, H. & Holtz, F. (2018). Ferric/ferrous ratio in silicate melts: a new model for 1 atm data with special emphasis on the effects of melt composition. *Contributions to Mineralogy and Petrology* **173**, 98.
- Botcharnikov, R. E., Koepke, J., Holtz, F., McCammon, C. & Wilke, M. (2005). The effect of water activity on the oxidation and structural state of Fe in a ferro-basaltic melt. *Geochimica et Cosmochimica Acta* **69**, 5071–5085.
- Bryan, W. B. (1983). Systematics of modal phenocryst assemblages in submarine basalts: petrologic implications. *Contributions to Mineralogy and Petrology* **83**, 62–74.
- Carlson, R. L. & Herrick, C. N. (1990). Densities and porosities in the oceanic crust and their variations with depth and age. *Journal of Geophysical Research* 95, 9153–9170.
- Cashman, K. V. (1993). Relationship between plagioclase crystallization and cooling rate in basaltic melts. *Contributions to Mineralogy and Petrology* **113**, 126–142.
- Couch, S., Sparks, R. S. J. & Carroll, M. R. (2001). Mineral disequilibrium in lavas explained by convective self-mixing in open magma chambers. *Nature* **411**, 1037–1039.
- Downes, M. J. (1974). Sector and oscillatory zoning in calcic augites from M. Etna, Sicily. Contributions to Mineralogy and Petrology 47, 187–196.
- Dowty, E. (1976). Crystal structure and crystal growth: II. sector zoning in minerals. *American Mineralogist* **61**, 460–469.
- Erdmann, M. & Koepke, J. (2016). Experimental temperature cycling as a powerful tool to enlarge melt pools and crystals at magma storage conditions. *American Mineralogist* **101**, 960–969.
- Gaetani, G. A. & Grove, T. L. (1998). The influence of water on melting of mantle peridotite. *Contributions to Mineralogy* and Petrology 131, 323–346.
- Ginibre, C., Wörner, G. & Kronz, A. (2002). Minor- and trace-element zoning in plagioclase: implications for magma chamber processes at Parinacota volcano, northern Chile. Contributions to Mineralogy and Petrology 143, 300–315.

- Green, E. C. R., Holland, T. J. B., Powell, R. & White, R. W. (2012). Garnet and spinel Iherzolite assemblages in MgO-Al₂O₃-SiO₂ and CaO-MgO-Al₂O₃-SiO₂: Thermody namic models and an experimental conflict. *Journal of Metamorphic Geology* 30, 561–577.
- Grove, T. L. (1981). Use of FePt alloys to eliminate the iron loss problem in 1 atmosphere gas mixing experiments: Theoretical and practical considerations. *Contributions to Mineralogy and Petrology* 78, 298–304.
- Grove, T. L. & Bence, A. E. (1977). Experimental study of pyroxene-liquid interaction in quartz-normative basalt 15597. In: *Proceedings of the Lunar Science Conference 8*. Houston, pp. 1549–1579.
- Grove, T. L. & Bryan, W. B. (1983). Fractionation of pyroxene-phyric MORB at low pressure: an experimental study. Contributions to Mineralogy and Petrology 84, 293–309
- Grove, T. L. & Juster, T. C. (1989). Experimental investigations of low-Ca pyroxene stability and olivine-pyroxene-liquid equilibria at 1-atm in natural basaltic and andesitic liquids. *Contributions to Mineralogy and Petrology* **103**, 287–305.
- Grove, T. L., Kinzler, R. J. & Bryan, W. B. (1992). Fractionation of Mid-Ocean Ridge Basalt (MORB). In: Morgan, J. P., Blackman, D. K. & Sinton, J. M. (eds), Mantle Flow and Melt Generation at Mid-Ocean Ridges, Geophysical Monograph 71. Washington DC: American Geophysical Union, pp. 281–310.
- Guðmundsson, M. T., Jónsdóttir, K., Hooper, A., Holohan, E. P., Halldórsson, S. A., Ófeigsson, B. G., Cesca, S., Vogfjörd, K. S., Sigmundsson, F., Högnadóttir, T., Einarsson, P., Sigmarsson, O., Jarosch, A. H., Jónasson, K., Magnússon, E., Hreinsdóttir, S., Bagnardi, M., Parks, M. M., Hjörleifsdóttir, V., Pálsson, F., Walter, T. R., Schöpfer, M. P. J., Heimann, S., Reynolds, H. I., Dumont, S., Bali, E., Gudfinnsson, G. H., Dahm, T., Roberts, M. J., Hensch, M., Belart, J. M. C., Spaans, K., Jakobsson, S., Gudmundsson, G. B., Fridriksdóttir, H. M., Drouin, V., Dürig, T., Aðalgeirsdóttir, G., Riishuus, M. S., Pedersen, G. B. M., van Boeckel, T., Oddsson, B., Pfeffer, M. A., Barsotti, S., Bergsson, B., Donovan, A., Burton, M. R. & Aiuppa, A. (2016). Gradual caldera collapse at Bárdarbunga volcano, Iceland, regulated by lateral magma outflow. Science 353, aaf8988.
- Halldórsson, S. A., Bali, E., Hartley, M. E., Neave, D. A., Peate, D. W., Guðfinnsson, G. H., Bindeman, I., Whitehouse, M. J., Riishuus, M. S., Pedersen, G. B. M., Jakobsson, S., Askew, R., Gallagher, C. R., Guðmundsdóttir, E. R., Gudnason, J., Moreland, W. M., Óskarsson, B. V., Nikkola, P., Reynolds, H. I., Schmith, J. & Thordarson, T. (2018). Petrology and geochemistry of the 2014–2015 Holuhraun eruption, central Iceland: compositional and mineralogical characteristics, temporal variability and magma storage. *Contributions to Mineralogy and Petrology* 173, 64.
- Halldórsson, S. A., Óskarsson, N., Grönvold, K., Sigurdsson, G., Sverrisdottir, G. & Steinthórsson, S. (2008). Isotopic-heterogeneity of the Thjorsa lava-implications for mantle sources and crustal processes within the Eastern Rift Zone, Iceland. Chemical Geology 255, 305–316.
- Hammer, J. E. (2008). Experimental studies of the kinetics and energetics of magma crystallization. *Reviews in Mineralogy and Geochemistry* **69**, 9–59.
- Hammer, J. E., Jacob, S., Welsch, B., Hellebrand, E. & Sinton, J. M. (2016). Clinopyroxene in postshield Haleakala ankaramite 1. Efficacy of thermobarometry. Contributions to Mineralogy and Petrology 171, 7.
- Hansen, H. & Grönvold, K. (2000). Plagioclase ultraphyric basalts in Iceland: the mush of the rift. *Journal of Volcanology and Geothermal Research* 98, 1–32.

- Hartley, M. E., Bali, E., Maclennan, J., Neave, D. A. & Halldórsson, S. A. (2018). Melt inclusion constraints on petrogenesis of the 2014–2015 Holuhraun eruption, Iceland. Contributions to Mineralogy and Petrology 173, 10.
- Hartley, M. E., Morgan, D. J., Maclennan, J., Edmonds, M. & Thordarson, T. (2016). Tracking timescales of short-term precursors to large basaltic fissure eruptions through Fe-Mg diffusion in olivine. *Earth and Planetary Science Letters* 439, 58–70.
- Hartley, M. E., Neave, D. A., Maclennan, J., Edmonds, M. & Thordarson, T. (2015). Diffusive over-hydration of olivinehosted melt inclusions. *Earth and Planetary Science Letters* 425, 168–178.
- Hildner, E., Klügel, A. & Hansteen, T. H. (2012). Barometry of lavas from the 1951 eruption of Fogo, Cape Verde Islands: implications for historic and prehistoric magma plumbing systems. *Journal of Volcanology and Geothermal Research* 217–218, 73–90.
- Hollister, L. S. & Gancarz, A. (1971). Compositional sector-zoning in clinopyroxene from the Narce area, Italy. American Mineralogist 56, 959–979.
- Holloway, J. R., Pan, V. & Guðmundsson, G. B. (1992). High-pressure fluid-absent melting experiments in the presence of graphite: oxygen fugacity, ferric/ferrous ratio and dissolved CO₂. European Journal of Mineralogy 4, 105–114.
- Holness, M. B., Stock, M. J. & Geist, D. (2019). Magma chambers versus mush zones: constraining the architecture of sub-volcanic plumbing systems from microstructural analysis of crystalline enclaves. *Philosophical Transactions of the Royal Society A* 377, 1–28.
- Husen, A., Almeev, R. R. & Holtz, F. (2016). The effect of H₂O and pressure on multiple saturation and liquid lines of descent in basalt from the Shatsky Rise. *Journal of Petrology* **57**, 309–344.
- Jakobsson, S. & Óskarsson, N. (1994). The system C-O in equilibrium with graphite at high pressure and temperature: an experimental study. Geochimica et Cosmochimica Acta 58, 9–17.
- Jarosewich, E., Gooley, R. & Husler, J. (1987). Chromium augite
 a new microprobe reference sample. Geostandards and Geoanalytical Research 11, 197–198.
- Jarosewich, E., Nelen, J. A. & Norberg, J. A. (1980). Reference samples for electron microprobe analysis. *Geostandards* and *Geoanalytical Research* 4, 43–47.
- Kelley, K. A. & Cottrell, E. (2012). The influence of magmatic differentiation on the oxidation state of Fe in a basaltic arc magma. Earth and Planetary Science Letters 329–330, 109–121.
- Kirkpatrick, R. J. (1976). Towards a kinetic model for the crystallization of magma bodies. *Journal of Geophysical Research* 81, 2565–2571.
- Klügel, A., Hansteen, T. H. & Galipp, K. (2005). Magma storage and underplating beneath Cumbre Vieja volcano, La Palma (Canary Islands). Earth and Planetary Science Letters 236, 211–226.
- Kouchi, A., Sugawara, Y., Kashima, K. & Sunagawa, I. (1983). Laboratory growth of sector zones clinopyroxenes in the system CaMgSi₂O₆ – CaTiAl₂O₆. *Contributions to Mineralogy and Petrology* **83**, 177–184.
- Kress, V. C. & Carmichael, I. S. E. (1991). The compressibility of silicate liquids containing Fe₂O₃ and the effect of composition, temperature, oxygen fugacity and pressure on their redox states. *Contributions to Mineralogy and Petrology* **108**, 82–92.
- Leung, I. S. (1974). Sector-zoned titanaugites: morphology, crystal chemistry, and growth. *American Mineralogist* **59**, 127–138.

- Lindsley, D. H. (1983). Pyroxene thermometry. *American Mineralogist* **68**, 477–493.
- Lofgren, G. E., Huss, G. R. & Wasserburg, G. J. (2006). An experimental study of trace-element partitioning between Ti-Al-clinopyroxene and melt: equilibrium and kinetic effects including sector zoning. *American Mineralogist* 91, 1596–1606.
- Longpré, M.-A., Klügel, A., Diehl, A. & Stix, J. (2014). Mixing in mantle magma reservoirs prior to and during the 2011–2012 eruption at El Hierro, Canary Islands. *Geology* **42**, 315–318.
- Longpré, M.-A., Stix, J., Klügel, A. & Shimizu, N. (2017). Mantle to surface degassing of carbon- and sulphur-rich alkaline magma at El Hierro, Canary Islands. *Earth and Planetary Science Letters* 460, 268–280.
- Luth, R. W. & Canil, D. (1993). Ferric iron in mantle-derived pyroxenes and a new oxybarometer for the mantle. *Contributions to Mineralogy and Petrology* **113**, 236–248.
- Maclennan, J., McKenzie, D., Grönvold, K. & Slater, L. (2001). Crustal accretion under Northern Iceland. Earth and Planetary Science Letters 191, 295–310.
- McGuire, A. V., Dyar, M. D. & Ward, K. A. (1989). Neglected Fe³⁺/Fe²⁺ ratios a study of Fe³⁺ content of megacrysts from alkali basalts. *Geology* **17**, 687–690.
- Médard, E., McCammon, C. A., Barr, J. A. & Grove, T. L. (2008). Oxygen fugacity, temperature reproducibility, and H₂O contents of nominally anhydrous piston-cylinder experiments using graphite capsules. *American Mineralogist* 93, 1838–1844.
- Mollo, S., Blundy, J., Scarlato, P., De Cristofaro, S. P., Tecchiato, V., Di Stefano, F., Vetere, F., Holtz, F. & Bachmann, O. (2018). An integrated P-T-H₂O-lattice strain model to quantify the role of clinopyroxene fractionation on REE+Y/HFSE patterns of mafic alkaline magmas: application to eruptions at Mt. Etna. *Earth-Science Reviews* **185**, 32–56.
- Mollo, S., Del Gaudio, P., Ventura, G., Iezzi, G. & Scarlato, P. (2010). Dependence of clinopyroxene composition on cooling rate in basaltic magmas: implications for thermobarometry. *Lithos* 118, 302–312.
- Mollo, S. & Hammer, J. E. (2017). Dynamic crystallization in magmas. In: Heinrich, W. & Abart, R. (eds), Mineral Reaction Kinetics: Microstructures, Textures, Chemical and Isotopic Signatures. European Mineralogical Union and the Mineralogical Society of Great Britain & Ireland, Jena, Germany; Twickenham, UK, pp. 378–418.
- Mollo, S., Lanzafame, G., Masotta, M., lezzi, G., Ferlito, C. & Scarlato, P. (2011). Cooling history of a dike as revealed by mineral chemistry: a case study from Mt. Etna volcano. *Chemical Geology* 288, 39–52.
- Mollo, S., Putirka, K. D., Misiti, V., Soligo, M. & Scarlato, P. (2013). A new test for equilibrium based on clinopyroxene-melt pairs: clues on the solidification temperatures of Etnean alkaline melts at post-eruptive conditions. *Chemical Geology* 352, 92–100.
- Morimoto, N., Fabries, J., Ferguson, A. K., Ginzburg, I. V., Ross, M., Seifert, F. A., Zussman, J., Aoki, K. & Gottardi, G. (1988). Nomenclature of pyroxenes. *American Mineralogist* 73, 1123–1133.
- Nakamura, Y. (1973). Origin of sector-zoning of igneous clinopyroxenes. *American Mineralogist* **58**, 986–990.
- Nakamura, Y. & Coombs, D. S. (1973). Clinopyroxenes in the tawhiroko tholeiitic dolerite at Moeraki, north-eastern Otago, New Zealand. Contributions to Mineralogy and Petrology 42, 213–228.
- Namur, O., Charlier, B., Toplis, M. J. & Vander Auwera, J. (2012). Prediction of plagioclase-melt equilibria in anhydrous silicate melts at 1-atm. *Contributions to Mineralogy and Petrology* 163, 133–150.

- Neave, D. A., Maclennan, J., Hartley, M. E., Edmonds, M. & Thordarson, T. (2014). Crystal storage and transfer in basaltic systems: the Skuggafjöll eruption, Iceland. *Journal of Petrology* 55, 2311–2346.
- Neave, D. A., Maclennan, J., Thordarson, T. & Hartley, M. E. (2015). The evolution and storage of primitive melts in the Eastern Volcanic Zone of Iceland: the 10 ka Grímsvötn tephra series (i.e. the Saksunarvatn ash). Contributions to Mineralogy and Petrology 170, 21.
- Neave, D. A., Passmore, E., Maclennan, J., Fitton, J. G. & Thordarson, T. (2013). Crystal-melt relationships and the record of deep mixing and crystallization in the AD 1783 Laki eruption, Iceland. *Journal of Petrology* 54, 1661–1690.
- Neave, D. A. & Putirka, K. D. (2017). A new clinopyroxene-liquid barometer, and implications for magma storage pressures under Icelandic rift zones. *American Mineralogist* 102, 777–794.
- Pankhurst, M. J., Morgan, D. J., Thordarson, T. & Loughlin, S. C. (2018). Magmatic crystal records in time, space, and process, causatively linked with volcanic unrest. *Earth and Planetary Science Letters* 493, 231–241.
- Passmore, E., Maclennan, J., Fitton, J. G. & Thordarson, T. (2012). Mush disaggregation in basaltic magma chambers: evidence from the AD 1783 Laki eruption. *Journal of Petrology* 53, 2593–2623.
- Pedersen, G. B. M., Höskuldsson, A., Dürig, T., Thordarson, T., Jónsdóttir, I., Riishuus, M. S., Óskarsson, B. V., Dumont, S., Magnusson, E., Gudmundsson, M. T., Sigmundsson, F., Drouin, V. J. P. B., Gallagher, C., Askew, R., Gudnason, J., Moreland, W. M., Nikkola, P., Reynolds, H. I. & Schmith, J. (2017). Lava field evolution and emplacement dynamics of the 2014–2015 basaltic fissure eruption at Holuhraun, Iceland. *Journal of Volcanology and Geothermal Research* 340, 155–169.
- Petrone, C. M., Bugatti, G., Braschi, E. & Tommasini, S. (2016).

 Pre-eruptive magmatic processes re-timed using a non-isothermal approach to magma chamber dynamics.

 Nature Communications 7, 12946.
- Putirka, K. D. (1999). Clinopyroxene + liquid equilibria to 100 kbar and 2450 K. *Contributions to Mineralogy and Petrology* 135, 151–163
- Putirka, K. D. (2008). Thermometers and barometers for volcanic systems. *Reviews in Mineralogy and Geochemistry* **69**, 61–120.
- Putirka, K. D., Johnson, M., Kinzler, R. J., Longhi, J. & Walker, D. (1996). Thermobarometry of mafic igneous rocks based on clinopyroxene-liquid equilibria, 0–30 kbar. *Contributions to Mineralogy and Petrology* **123**, 92–108.
- Sack, R. O. & Ghiorso, M. S. (1994). Thermodynamics of multicomponent pyroxenes: I. Formulation of a general model. Contributions to Mineralogy and Petrology 116, 277–286.
- Schuessler, J. A., Botcharnikov, R. E., Behrens, H., Misiti, V. & Freda, C. (2008). Oxidation state of iron in hydrous phonotephritic melts. *American Mineralogist* 93, 1493–1504.
- Shane, P. & Coote, A. (2018). Thermobarometry of Whangarei volcanic field lavas, New Zealand: Constraints on plumbing systems of small monogenetic basalt volcanoes. *Journal of Volcanology and Geothermal Research* 354, 130–139.
- Shishkina, T. A., Botcharnikov, R. E., Holtz, F., Almeev, R. R. & Portnyagin, M. V. (2010). Solubility of H2O- and CO2-bearing fluids in tholeitic basalts at pressures up to 500 MPa. *Chemical Geology* 277, 115–125.
- Sigmundsson, F., Hooper, A., Hreinsdóttir, S., Vogfjörd, K. S., Ófeigsson, B. G., Heimisson, E. R., Dumont, S., Parks, M., Spaans, K., Gudmundsson, G. B., Drouin, V., Árnadóttir, T., Jónsdóttir, K., Gudmundsson, M. T., Högnadóttir, T., Fridriksdóttir, H. M., Hensch, M., Einarsson, P., Magnússon,

- E., Samsonov, S., Brandsdóttir, B., White, R. S., Ágústsdóttir, T., Greenfield, T., Green, R. G., Hjartardóttir, Á. R., Pedersen, R., Bennett, R. A., Geirsson, H., La Femina, P. C., Björnsson, H., Pálsson, F., Sturkell, E., Bean, C. J., Möllhoff, M., Braiden, A. K. & Eibl, E. P. S. (2015). Segmented lateral dyke growth in a rifting event at Bárðarbunga volcanic system, Iceland. *Nature* **517**, 191–195.
- Skulski, T., Minarik, W. & Watson, E. B. (1994). High-pressure experimental trace-element partitioning between clinopyroxene and basaltic melts. *Chemical Geology* **117**, 127–147.
- Strong, D. F. (1969). Formation of the hour-glass structure in augite. *Mineralogical Magazine* **37**, 472–479.
- Thomson, A. & Maclennan, J. (2013). The distribution of olivine compositions in Icelandic basalts and picrites. *Journal of Petrology* 54, 745–768.
- Thordarson, T. & Self, S. (1993). The Laki (Skaftár Fires) and Grímsvötn eruptions in 1783–1785. *Bulletin of Volcanology* **55**, 233–263.
- Toplis, M. J. (2005). The thermodynamics of iron and magnesium partitioning between olivine and liquid: criteria for assessing and predicting equilibrium in natural and experimental systems. *Contributions to Mineralogy and Petrology* 149, 22–39.
- Tormey, D. R., Grove, T. L. & Bryan, W. B. (1987). Experimental petrology of normal MORB near the Kane Fracture Zone: 22°–25°N, mid-Atlantic ridge. *Contributions to Mineralogy and Petrology* **96**, 121–139.
- Ubide, T. & Kamber, B. S. (2018). Volcanic crystals as time capsules of eruption history. *Nature Communications* 9, 326.
- Ubide, T., Mollo, S., Zhao, J.-X., Nazzari, M. & Scarlato, P. (2019). Sector-zoned clinopyroxene as a recorder of magma

- history, eruption triggers, and ascent rates. *Geochimica et Cosmochimica Acta* **251**, 265–283.
- Ulmer, P., Kaegi, R. & Müntener, O. (2018). Experimentally derived intermediate to silica-rich arc magmas by fractional and equilibrium crystallization at 1.0 GPa: an evaluation of phase relationships, compositions, liquid lines of descent and oxygen fugacity. *Journal of Petrology* 59, 11–58.
- Villiger, S., Ulmer, P., Müntener, O. & Thompson, A. B. (2004). The liquid line of descent of anhydrous, mantle-derived, tholeiitic liquids by fractional and equilibrium crystallization—an experimental study at 1.0 GPa. *Journal of Petrology* **45**, 2369–2388.
- Weis, F. A., Skogby, H., Troll, V. R., Deegan, F. M. & Dahren, B. (2015). Magmatic water contents determined through clinopyroxene: examples from the Western Canary Islands, Spain. *Geochemistry, Geophysics, Geosystems* 16, 2127–2146.
- Welsch, B., Hammer, J. E., Baronnet, A., Jacob, S., Hellebrand, E. & Sinton, J. M. (2016). Clinopyroxene in postshield Haleakala ankaramite 2. Texture, compositional zoning, and supersaturation in the magma. *Contributions to Mineralogy* and Petrology 171, 6.
- Whitaker, M. L., Nekvasil, H., Lindsley, D. H. & Difrancesco, N. J. (2007). The role of pressure in producing compositional diversity in intraplate basaltic magmas. *Journal of Petrology* 48, 365–393.
- Winpenny, B. & Maclennan, J. (2011). A partial record of mixing of mantle melts preserved in Icelandic phenocrysts. *Journal of Petrology* **52**, 1791–1812.
- Zhang, Y., Ni, H. & Chen, Y. (2010). Diffusion data in silicate melts. *Reviews in Mineralogy and Geochemistry* **72**, 311–408.