# <sup>1</sup> Upper mantle mush zones beneath low

## <sup>2</sup> melt flux ocean island volcanoes:

### <sup>3</sup> insights from Isla Floreana, Galápagos

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### 8 ABSTRACT

9 The physicochemical characteristics of sub-volcanic magma storage regions have important implications for magma system dynamics and pre-eruptive behaviour. The architecture of magma 10 storage regions located directly above high buoyancy flux mantle plumes (such as Kīlauea, Hawaii 11 12 and Fernandina, Galápagos) are relatively well understood. However, far fewer constraints exist on the nature of magma storage beneath ocean island volcanoes that are distal to the main zone of mantle 13 upwelling or above low buoyancy flux plumes, despite these systems representing a substantial 14 proportion of global ocean island volcanism. To address this, we present a detailed petrological study 15 16 of Isla Floreana in the Galápagos Archipelago, which is characterised by an extremely low flux of magma into the lithosphere from the underlying mantle plume. Detailed *in situ* major and trace 17 18 element analyses of crystal phases within exhumed cumulate xenoliths, lavas and scoria deposits, 19 indicate that magma storage beneath Floreana is dominated by crystal-rich domains (i.e. mush). Trace 20 element disequilibria between cumulus phases and erupted melts, as well as trace element zoning 21 within the xenolithic clinopyroxenes, reveals that reactive porous flow (previously identified beneath 22 mid-ocean ridges) is an important process of melt transport within these crystal-rich storage regions. 23 In addition, application of three petrological barometers reveal that the Floreana mush zones are 24 located in the upper mantle, at a depth of 23.7±5.1 km. Our barometric results are compared to recent studies of high melt flux volcanoes in the western Galápagos, Hawaii and Iceland, and demonstrate 25 26 that the flux of magma from the underlying mantle source represents a first-order control on the depth 27 and physical characteristics of magma storage beneath ocean island volcanoes.

### 28 **KEY WORDS**

29 Galápagos; magma storage; reactive porous flow; barometry.

### **30 1 INTRODUCTION**

The physicochemical characteristics (such as size, pressure, volatile content and geochemical 31 heterogeneity) of magma storage at volcanic centres located directly above high buoyancy flux mantle 32 plumes (e.g. Kīlauea, Hawaii and Isabela, Galápagos) have been subject to intense study over the past 33 few decades (Bagnardi et al., 2013; Clague and Denlinger, 1994; Geist et al., 1998; Naumann and 34 Geist, 1999; Park et al., 2007; Pietruszka et al., 2015; Poland et al., 2015; Stock et al., 2018; Wieser et 35 36 al., 2020, 2019). Systems such as Kīlauea are characterised by frequent volcanic activity, and geophysical (seismicity, ground deformation) and geochemical (gas emissions) monitoring is 37 38 prevalent. Monitoring data, combined with petrological and geochemical analysis of erupted products 39 (mineral textures, deformation characteristics and chemistry) provide important insights into the 40 architecture and dynamics of their sub-volcanic plumbing systems (Amelung et al., 2000; Davidge et 41 al., 2017; Geist et al., 2014; Hartley et al., 2018; Kilbride et al., 2016). However, these systems 42 (which we term 'high melt flux') represent only one endmember of global plume-derived volcanism. 43 Low melt flux systems, either above low buoyancy flux plumes (e.g. Canary Islands; Longpre et al., 44 2014) or at volcanic systems distal to the centre of mantle melting at high buoyancy flux mantle 45 plumes (e.g. eastern and southern Galápagos; Harpp and Geist, 2018), are the other endmember. 46 Although a substantial number of Holocene hotspot-related volcanic systems are located in regions 47 characterised by a relatively low flux of magma into the lithosphere (e.g. Samoa, Canary Islands, Cape Verde; Global Volcanism Program, 2013), only a small number of eruptions have been observed 48 49 (and recorded) at these systems since the advent of modern volcano monitoring techniques. As a 50 result, few constraints exist on the conditions of magma storage in regions characterised by a low flux 51 of magma into the lithosphere, relative to volcanic centres located above the centre of mantle plumes with a large buoyancy flux (and thus generating a large flux of magma). The flux of mantle-derived 52 53 magma into the lithosphere is important because it is thought to impart a first-order control on the 54 development of large silicic magma bodies (Barker et al., 2020) and the homogeneity of erupted

liquids (Geist et al., 2014). Furthermore, placing constraints on the physicochemical characteristics of magma storage at low melt flux ocean island volcanoes is essential for determining the influence of mantle dynamics and melt generation processes on the structure and physical characteristics of subvolcanic magma plumbing systems. In the absence of detailed monitoring data, petrological and geochemical analyses of volcanic products from past eruptions represent the only available tools for determining the structure and processes operating within these systems.

Isla Floreana in the south-eastern Galápagos Archipelago is currently located ~100 km downstream 61 from where the Galápagos plume impacts on the base of the lithosphere beneath the island of Isabela 62 63 in the western archipelago (Fig. 1; Villagómez et al., 2014). Hence, although the Galápagos plume has a relatively high buoyancy flux (compared to regions such as the Canary Islands; Jackson et al., 64 2017), Floreana's location relative to the main zone of mantle plume upwelling results in an 65 extremely low flux of magma entering the lithosphere and, consequently, very infrequent volcanic 66 67 activity (Harpp et al., 2014a; Harpp and Geist, 2018). In this paper, we present a thorough petrological study of scoria, lava and xenolith samples from Floreana and place constraints of the structure, depth 68 69 and crystallinity of magmatic systems beneath this low melt flux ocean island volcano. We compare 70 our results with more frequently active volcanic centres in the western Galápagos (near the centre of 71 plume upwelling; Geist et al., 1998; Naumann and Geist, 1999; Stock et al., 2018), as well as Iceland 72 and Hawaii (Hammer et al., 2016; Hartley et al., 2018; Poland et al., 2015), to investigate how the flux of magma into the lithosphere influences the depth and crystallinity of sub-volcanic magma 73 74 storage regions.

### 75 2 GEOLOGICAL BACKGROUND

The Galápagos Archipelago in the eastern equatorial Pacific is one of the most volcanically active
regions on Earth, with eruptions typically occurring every 2–3 years (Global Volcanism Program,
2013). Although most historic Galápagos eruptions have taken place on the two westernmost islands
of Isabela and Fernandina (Fig. 1), infrequent volcanic activity has also occurred on several islands in

the eastern and southern Galápagos (e.g. Santiago in 1906 and Marchena in 1991; Global Volcanism
Program, 2013).

Volcanoes in the western Galápagos likely emerged within the last 500 kyr (Naumann and Geist, 82 83 2000), whereas those in the eastern and south-eastern Galápagos are considerably older (eruption ages up to 2.3 Ma and 3.2 Ma have been measured on San Cristobal and Espanola, respectively; Bailey, 84 1976; Geist et al., 1986). In addition, substantial differences in geomorphology and the style of 85 volcanic activity are observed across the archipelago (Geist et al., 1995; Harpp et al., 2014a; Harpp 86 and Geist, 2018). For example, volcanoes in the western archipelago are typified by large summit 87 88 calderas (<700m deep), which are not present on the eastern islands (Chadwick and Howard, 1991; Cleary et al., 2020; Harpp and Geist, 2018). 89 90 Geochemical distinctions between the western and eastern/southern Galápagos islands are also observed, which are primarily related to variations in the composition of the underlying mantle source 91 92 (Geist et al., 1988; Gibson and Geist, 2010; Gleeson et al., 2020; Harpp and White, 2001; White et al., 93 1993) or the volume flux of mantle-derived magma that ascends into the lithosphere (Geist et al., 1995, 2014; Gibson et al., 2016; Harpp and Geist, 2018). For example, variations in the flux of mantle 94 derived magma are hypothesised to influence the geochemical heterogeneity of erupted basalts at each 95 island: volcanoes in the western archipelago typically erupt a very narrow range of basaltic 96 97 compositions over hundreds of millennia during their main shield building phase, whereas basalts 98 erupted from a single island in the eastern and/or south-eastern archipelago, such as Floreana, tend to 99 display far greater compositional heterogeneity (Geist et al., 2014; Harpp and Geist, 2018). 100 Floreana is characterised by numerous scoria cones (up to ~600m elevation at Cerro Pajas) and 101 blocky, heavily vegetated lava flows that can typically be traced to the cone from which they 102 originated (Harpp et al., 2014a). The crustal thickness beneath Floreana is ~16 km, similar to that observed in the western Galápagos 10-18 km (Feighner and Richards, 1994), and the lithospheric 103 thickness beneath the western and south-eastern Galápagos is very similar (~50-60 km; Gibson and 104 Geist, 2010). However, recent work has shown that the volumetric eruption rate on Floreana is 1-10105 106  $m^3 \cdot yr^{-1}$  over the past 1–1.5 Myrs, millions of times lower than the current effusion rate at volcanoes in

the western Galápagos (cf.  $\sim 4.4 \cdot 10^6 \text{ m}^3 \cdot \text{yr}^{-1}$  at Fernandina; Harpp et al., 2014a; Kurz et al., 2014). 107 This variation in volumetric eruption rate likely reflects a substantially lower flux of magma into the 108 lithosphere beneath Floreana than beneath each volcanic centre in the western Galápagos. 109 Additionally, Floreana has a high proportion of pyroclastic deposits compared to the other Galápagos 110 111 islands (Harpp et al., 2014a) and eruption deposits typically contain a large number of cumulate xenoliths (Lyons et al., 2007). The abundance of pyroclastic deposits and xenoliths on Floreana has 112 been interpreted as evidence for very high magma ascent rates (Harpp et al., 2014a). 113 114 Floreana is one of the only Galápagos islands that displays evidence for multiple stages in its volcanic 115 evolution. Submarine parts of the island have isotopic and trace element characteristics that are similar to recent basalts erupted on southern Isabela (e.g. Sierra Negra and Cerro Azul, Fig. 1A), 116 whereas the subaerial material is isotopically distinct (high <sup>206</sup>Pb/<sup>204</sup>Pb ratios; Harpp et al., 2014). The 117 trace element and isotopic variation in the erupted basalts is mirrored in xenoliths found in the 118 119 Floreana lava and scoria deposits: gabbroic xenoliths have radiogenic isotope ratios that are similar to modern Isabela basalts, whereas wehrlitic xenoliths have trace element and isotopic compositions that 120 121 resemble recent Floreana subaerial basalts (Lyons et al., 2007). Variations in the isotopic characteristics of the Floreana lavas (submarine vs subaerial) and xenoliths (gabbros vs wehrlites) are 122 123 thought to indicate a change in the mean composition of magma produced by mantle melting beneath the island at  $\sim 1-1.5$  Ma (Harpp et al., 2014a). In this study, we focus on constraining the depth and 124 125 physical characteristics of magma storage during the most recent period of volcanic activity on Isla 126 Floreana (<1–1.5 Ma) using chemical and textural analysis of crystal phases in lava flows and 127 xenolithic nodules.

### **3 SAMPLES AND PETROGRAPHY**

The Floreana samples analysed in this study were collected during a field campaign to the northern part of the island in June 2017 and consist of lavas (27 samples), scoria (2 samples) and xenoliths (4 wehrlite, 3 dunite and 2 gabbro samples; Fig. 1B). Most lava samples were collected from the unaltered, low vesicularity cores of blocky flows or glassy flow fronts. The scoria samples were

- 133 collected from two separate deposits and comprise rapidly cooled tephra fragments ( $\sim 0.5-2$  cm
- across; 17MMSG16) and scoria bombs (~10 cm across; 17MMSG20). Xenolithic fragments (3–15cm
- across) were sampled from two different scoria cones on the north-east coast of Floreana and are also
- 136 found within most lava flows across the entire island (Fig. 1).
- 137 3.1 Lavas and scoria
- The lava and scoria samples analysed in this study are typically olivine phyric with minor andehralclinopyroxene and very rare orthopyroxene. Except for small plagioclase laths in the microcrystalline
- 140 groundmass, plagioclase crystals are extremely rare in Floreana lavas. Plagioclase macrocrysts are
- only present in one of our lava samples (17MMSG29) where they occur as isolated phenocrysts and in
- 142 plagioclase-olivine crystal clots (Table S.1). Abundant olivine and the absence of plagioclase in the
- 143 Floreana lavas and scoria contrasts with basalts in the central, northern, and western parts of the
- 144 Galápagos Archipelago, where plagioclase-phyric and ultraphyric basalts are common (Geist et al.,
- 145 2002; Gibson et al., 2012; Harpp et al., 2014b).
- 146 Despite their relatively simple mineralogy, Floreana lava and scoria samples contain texturally diverse
- 147 olivine crystals which can be divided into five distinct groups (Fig. 2):
- Group 1 olivines are present in all lava and scoria samples and are the most abundant type of
- 149 olivine found in the Floreana basalts (~60-70% of all crystals). They are characterised by
- 150 homogeneous cores and narrow normally zoned rims (Fig. 2A). Group 1 olivines are
- 151 generally subhedral to euhedral.
- Group 2 olivines are the second most abundant group in the Floreana lava and scoria deposits
  (~20-30%) and display reverse zoning patterns. They are typically euhedral, with occasional
  small embayments (Fig. 2B).
- Group 3 olivines are also reversely zoned, but are distinguished by skeletal overgrowths,
- indicating significant undercooling of the host magma and rapid crystal growth (Fig. 2C;
- 157 Donaldson, 1976; Welsch et al., 2014). Group 3 olivines are less abundant than Groups 1 and
- 158 2 olivines in Floreana lava and scoria deposits (<10%).

159 -	Group 4 olivines are present in low abundance in the Floreana lava and scoria deposits
160	(<10%). They have homogeneous cores and reverse-zoned rims (up to ~300 $\mu m$ thick). The
161	rims have sieve textures, indicating resorption and chemical disequilibrium with their carrier
162	melts (Fig. 2D).
163 -	Group 5 olivines are only found in a minority of samples and are characterised by the
164	presence of 4 compositional zones with alternating high and low forsterite contents (visible in
165	back-scattered electron images; Fig. 2E and F).

### 166 3.2 Xenoliths

167 *3.2.1 Gabbroic xenoliths* 

168 Floreana gabbroic xenoliths predominantly comprise plagioclase (33–66 vol.%), clinopyroxene (28–

46 vol.%) and orthopyroxene (5–15 vol.%), with little or no olivine (Table S.2). Plagioclase and

170 pyroxene crystals are typically  $>500 \mu m$  and grain sizes are relatively constant within a single

171 xenolith sample. Where three plagioclase grains meet at a triple junction, 120° grain boundaries

indicate a high degree of textural equilibrium (Fig. 3A; Holness et al., 2005). Some of the gabbros

display clear clinopyroxene-rich and plagioclase-rich layers (2–5 mm), whereas others have a more

- 174 irregular mineral distribution.
- 175 3.2.2 Dunitic xenoliths
- 176 Cumulus olivine dominates dunitic Floreana xenoliths (>90 vol.%). The olivine crystals are subhedral
- to euhedral and may have undergone partial textural re-equilibration, with some olivine triple
- 178 junctions approaching 120° grain boundaries. Minor intercumulus clinopyroxene is present along
- 179 grain boundaries and between pre-existing olivine crystals (Fig. 3B).

#### 180 3.2.3 Wehrlitic xenoliths

- 181 Floreana wehrlitic xenoliths contain olivine (>50 vol.%), clinopyroxene (20–40 vol.%),
- 182 orthopyroxene (~0–7 vol.%) and minor spinel (<1 vol.%; Table S.1). Clinopyroxene typically occurs
- as large (<5 mm) oikocrysts, which enclose rounded olivine chadacrysts (<500 µm in diameter and
- separated by distances of  $<400 \mu m$ ; Fig. 2C and D). In the most pyroxene-rich samples (e.g.
- 185 17MMSG03a), clinopyroxene crystals contain fine-scale orthopyroxene exsolution lamallae (Fig. 2E

186 and F). Olivine grains that are not enclosed by clinopyroxene are typically larger (>1 mm) and more 187 euhedral than the chadacrysts. In some samples, the boundary between clinopyroxene and olivine crystals is characterised by a thin (<20-30 µm) layer of glass and very fine-grained microcrysts. 188 Orthopyroxene is an intercumulus phase in the wehrlitic xenoliths and has an anhedral morphology, 189 190 infilling the space between earlier formed clinopyroxene and olivine grains. Our observations of 191 dunitic and wehrlitic xenoliths (which have the isotopic signatures of modern day Floreana basalts; 192 Lyons et al., 2007) indicate that the typical order of crystallisation beneath Floreana is olivine, 193 followed by clinopyroxene, with little to no crystallisation of plagioclase.

### **194 4 ANALYTICAL METHODOLOGY**

### 195 **4.1 Electron microprobe analysis**

Glass chips, olivine and clinopyroxene crystals were hand-picked from crushed scoria and lava 196 197 samples, mounted in epoxy or indium, and then ground and polished prior to analysis (crystals 198 mounted in indium were polished individually prior to mounting). Xenolithic crystals were analysed as individual crystals mounted in indium or *in situ* in petrographic thin sections. The major and minor 199 element concentrations of olivine, clinopyroxene and glass were measured using a Cameca SX100 200 electron microprobe in the Department of Earth Sciences, University of Cambridge. Calibrations were 201 202 made using mineral and metal standards prior to each analytical session (see Gleeson and Gibson, 203 2019 for details). Glasses in the two Floreana scoria samples were analysed using a 6 nA, 15 kV, 204 defocused (5 µm) beam for most elements. Na and K were analysed first (10 s peak count time) to 205 avoid alkali migration. Other elements were analysed with peak count times of 10 s (Si), 20 s (Fe), 30 206 s (Al, P, Ca, Mg), 40 s (Mn), or 60 s (Ti). Sulphur was analysed last using a 20 nA beam current and a 207 60 s peak count time.

208 Pyroxene compositions were determined by spot analyses using a 20 nA, 15 kV, focused ( $\sim 1 \mu m$ )

beam, with Na, K and Si analysed first (10 s). Element maps of Cr, Ti, and Al in key clinopyroxene

- 210 crystals from the Floreana xenoliths were created using a 60 nA, 15 kV, focused ( $\sim 1 \mu m$ ) beam, with
- a dwell time of 150 ms. Cr counts were collected on a PET and a LIF crystal, Al counts were

212 collected on two TAP crystals, and Ti counts were collected on a PET crystal. Olivine electron microprobe analysis was carried out using the method outlined in Gleeson and Gibson (2019). 213 214 Analytical uncertainties were tracked through analysis of appropriate Smithsonian Microbeam 215 Standards (Jarosewich et al., 1980). Accuracy is typically between 98 and 102% for all phases.  $2\sigma$ analytical precision of clinopyroxene and olivine analysis is typically better than 2–3% for major 216 elements (>1 wt%) and typically  $\sim$ 5-10% for minor elements (<1 wt%). Similarly, the 2 $\sigma$  precision for 217 glass analysis is typically <3% for major elements,  $\sim5\%$  for Na, and  $\sim10\%$  for K (See Supplementary 218 File). 219

### 220 4.2 Laser ablation Inductively Coupled Plasma Mass Spectrometry

Trace element concentrations were measured in the apparent cores (i.e. as exposed in the 2D plane) of 221 clinopyroxene crystals from scoria and xenolith samples using an ESI193 laser coupled to a Nexion 222 223 350D inductively coupled plasma mass spectrometer in the Department of Earth Sciences, University of Cambridge. Analyses were collected in spot mode using a 20 Hz laser repetition rate, 4 J/cm<sup>2</sup> 224 fluence and 80 µm spot size, or in transect mode using a 10 Hz repetition rate, 3.5 J/cm<sup>2</sup> fluence and 225 30 µm spot size. For transects, individual spots were offset into two (alternating) lines to increase the 226 227 spatial resolution. Laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) data reduction was carried out in Iolite<sup>®</sup>, with NIST 612SRM as the standard reference material and Ca 228 (from electron microprobe analysis) as the internal reference standard. Analytical accuracy was 229 tracked using a USGS glass standard (Jochum et al., 2016) and is between 95% and 105% for most 230 231 elements (See Supplementary File). 2<sup>o</sup> analytical precision of spot analyses was monitored through analysis of an in-house clinopyroxene standard and is 5-10% for the light rare-earth elements 232 (LREE), Y, Sr, and Zr and 10-20% for the heavy rare earth element (HREE). 25 analytical precision 233 is ~10% for all elements of interest (Ce, Y) in transect analyses. 234

### **5 GLASS AND MINERAL CHEMISTRY**

### 236 5.1 Matrix glass compositions

The matrix glass compositions measured in one scoria (17MMSG16) and one glassy lava sample 237 (17MMSG12) from Floreana have very similar MgO concentrations (6.06-6.36 wt%) but exhibit 238 differences in the concentrations of other elements (Fig. 4). For example, sample 17MMSG12 has 239 consistently lower TiO<sub>2</sub> and Al<sub>2</sub>O<sub>3</sub> concentrations than 17MMSG16, which must either reflect 240 heterogeneity in the composition of primary mantle melts or variations in crustal processing (e.g. the 241 extent of plagioclase or clinopyroxene crystallisation; Fig. 4). Differences between our matrix glass 242 major element analyses and previously-published whole-rock data from Floreana (Harpp et al., 2014a) 243 244 are primarily due to olivine accumulation in the whole-rock samples (additional accumulation of clinopyroxene may explain the high CaO content of some whole-rock samples; Fig. 4C). 245 246 Our Floreana matrix glass analyses have higher Al<sub>2</sub>O<sub>3</sub> concentrations than basaltic glass and wholerock measurements from the western Galápagos Archipelago (excluding whole-rock samples with 247 accumulated plagioclase, Fig. 4D; Geist et al., 2002). This indicates substantially lower extents of 248 249 plagioclase fractionation in the Floreana magmatic system and is consistent with the scarcity of 250 plagioclase phenocrysts in erupted Floreana lavas. Reduced plagioclase crystallisation could be due to 251 the major element composition or H<sub>2</sub>O content of primary mantle melts and/or the increased pressure

of magma storage (Asimow and Langmuir, 2003; Neave et al., 2019; Thompson, 1987; Winpenny and
Maclennan, 2011).

### 254 **5.2** Olivine compositions

Olivine crystals in our Floreana lava and scoria samples show large variations in their forsterite contents (Fo = 70–92, where Fo=(Mg/(Mg+Fe<sup>2+</sup>) molar) with histograms showing a primary density peak at Fo~85 (Fig. 5), more primitive than the olivine composition in equilibrium with basaltic glasses from Floreana ( $K_D = 0.30$ ; Roeder and Emslie, 1970). Although there is no clear correlation between Fo and Ca concentration in these crystals, the most forsteritic olivines (Fo>83) have 260 extremely diverse Ca contents (~250 to ~2600 ppm; Fig. 5), whereas crystals with lower Fo contents (<83) have ubiquitously low Ca concentrations (<1500 ppm). Floreana olivines separated from the 261 lava and scoria samples also have a large range of Ni concentrations (~700 to ~3200 ppm), consistent 262 with crystallisation from a peridotite-derived melt (Fig. 5; Gleeson and Gibson, 2019; Herzberg, 263 264 2011; Matzen et al., 2017b, 2017a). All olivine crystals analysed in the wehrlite and dunite xenoliths have a narrow range of Fo contents (83-87) and, crucially, have uniformly low Ca concentrations 265 266 (<1000 ppm) and moderately high Ni contents (~2000 ppm). The range in Ca contents contrasts with 267 olivines from other Galápagos islands where Ca is typically >1000 ppm (Gleeson and Gibson, 2019; 268 Vidito et al., 2013). The Ca and Ni contents of our Floreana olivines are inversely related at a set Fo 269 content (Fig. 5).

270 **5.3 Clinopyroxene compositions** 

### 271 5.3.1 Major elements

The Floreana clinopyroxenes analysed in this study (from scoria and xenolith samples) are augites 272 273  $(En_{48.5}Fs_{8}Wo_{43.5})$  and have a relatively high Mg# (0.85–0.90; Mg# = Mg/(Mg+Fe<sub>t</sub>) molar). The 274 clinopyroxenes typically contain very high Na concentrations (<1.58 wt% Na<sub>2</sub>O, <0.11 Na atoms per formula unit; Fig. 6A) and, correspondingly, up to 10% of the jadeite component. In general, 275 276 clinopyroxene separates from scoria samples display a wide range of Na concentrations, whereas the 277 xenolithic clinopyroxenes have high Na contents (Fig. 6A). The Floreana clinopyroxenes display a large range of Cr contents, ranging from <0.05 wt% in the most evolved crystals to  $\sim1.72$  wt% in the 278 more primitive crystals (Fig. 6B). 279

### 280 5.3.2 Trace elements

Our Floreana clinopyroxenes display a wide range of geochemical enrichment, with LREE to MREE ratios varying from  $[La/Sm]_n \sim 0.2$  to  $[La/Sm]_n \sim 3.1$ . Xenolithic clinopyroxenes typically have more

- enriched trace element ratios (such as  $[La/Sm]_n$  or  $[Ce/Y]_n$ ) than clinopyroxenes from the scoria
- samples (Fig. 7). Furthermore, the melt  $[La/Sm]_n$  ratios calculated to be in equilibrium with
- clinopyroxenes from the scoria and xenolith samples range from ~1 to ~15 (calculated using the

286 elastic strain model of Wood and Blundy (1997) at 1225°C and 700 MPa; Fig. 7B), significantly 287 greater than the range measured in Floreana whole-rock samples (~2–5, with a small number of outliers up to ~7.5; Harpp et al. 2014a). Almost all of the xenolithic crystals, and a large proportion of 288 289 the clinopyroxenes separated from scoria deposits, are too enriched to be in equilibrium with the 290 typical composition of melts erupted on Floreana (Harpp et al., 2014a). In addition, there is a strong 291 correlation between the Na concentrations and highly/moderately incompatible trace element ratios of 292 the Floreana clinopyroxenes, such that crystal with high incompatible trace element concentrations 293 typically contain a high jadeite component (Fig. 7A).

# 6 MAGMA SYSTEM ARCHITECTURE BENEATH ISLA FLOREANA

### **6.1** Mush crystallisation and textural equilibration

Based on pyroxene trace element and radiogenic isotope ratios, Lyons et al. (2007) hypothesised that
gabbroic xenoliths in the Floreana lava and scoria deposits formed in an ancient (>1-1.5 Ma)
magmatic system, compositionally similar to those currently beneath Cerro Azul and Sierra Negra
volcanoes in the western Galápagos Archipelago. In contrast, wehrlite xenoliths preserve isotopic
ratios similar to more recent subaerial lavas on Floreana, suggesting that they are fragments of the
present-day magmatic system (Lyons et al., 2007).

303 Our wehrlitic xenoliths preserve an original igneous texture (clinopyroxene oikocrysts surrounding 304 olivine chadacrysts; Wager et al., 1960) and display no evidence of textural re-equilibration (e.g. 120° 305 dihedral angles) or metamorphic breakdown of plagioclase (e.g. pseudomorphs or relict cores; Lyons et al., 2007). The presence of orthopyroxene lamellae in some clinopyroxenes indicates that they 306 307 cooled relatively slowly (Poldervaart and Hess, 1951). Hence, we suggest that the clinopyroxene 308 crystals in our wehrlitic xenoliths grew within an olivine-dominated cumulate mush (i.e. interstitial growth of clinopyroxene oikocrysts; Wager et al., 1960). The rounded morphology and small size of 309 olivine chadacrysts, relative to olivine crystals that are not enclosed by clinopyroxene oikocrysts (Fig. 310 3D), indicates that clinopyroxene growth may have been at the expense of olivine (e.g. Lissenberg 311

- and MacLeod, 2016). If clinopyroxene growth is principally within a crystal-rich (i.e. relatively
- 313 viscous and immobile) mush zone, this could explain its relatively low abundance in Floreana lava
- and scoria deposits (<5% of separated crystals).
- 315 In contrast with the wehrlites, three-grain plagioclase triple junctions in our gabbroic xenoliths have
- 316 ~120° dihedral angles (Fig. 3A), indicating a high degree of textural equilibration (Holness et al.,
- 317 2019, 2005). Plagioclase textural equilibrium, along with the two-pyroxene phase assemblage,
- 318 suggests that the gabbroic xenoliths represent magmatic cumulates which were stored at high
- temperatures (>900°C) on long timescales (Holness et al., 2006). These petrographic observations are
- 320 consistent with the gabbroic nodules sampling an ancient magmatic system beneath Floreana (>1 Ma;
- 321 Lyons et al., 2007).

### 322 6.2 Mush disaggregation prior to eruption

323 6.2.1 Insights from olivine compositional heterogeneity

324 Olivine crystals separated from the Floreana lava and scoria samples have a wide range of zoning

325 patterns, morphologies, and compositions (Fig. 2). The five olivine groups identified in the Floreana

326 samples have distinct morphologies and zoning patterns (see Section 2 above), suggesting chemically

- 327 heterogeneous magma storage (Holness et al., 2019). In particular, the most evolved crystals (Group
- 4; Fo~70–75) are in equilibrium with melts that are more evolved than the Floreana erupted basalts.
- 329 This is consistent with a recent study which identified highly evolved (rhyolitic) magmas beneath
- 330 basaltic volcanoes in the western Galápagos Archipelago (Stock et al., *in review*).

331 As Fe-Mg interdiffusion in olivine is geologically fast (Costa et al., 2020), preservation of forsterite

332 zoning in the Floreana olivine crystals suggests that multiple magma batches interacted on relatively

333 short pre-eruptive timescales. In Group 5 olivines, for example, four compositional zones are

- preserved over ~100-200 μm near the rim (Fig. 2E and F). Whilst we do not have enough Group 5
- 335 olivine crystals to calculate statistically robust timescales of pre-eruptive magma interactions using
- diffusion chronometry, complex forsterite zoning over  $\sim 100 \,\mu m$  is estimated to last <3 yrs at the
- approximate temperature of basaltic magma storage (~1225°C; using diffusion coefficients from
- 338 Chakraborty, 2010). Therefore, we suggest that the range of crystal morphologies and major element

compositions displayed by the Floreana olivines in lava and scoria deposits indicates mixing and
 amalgamation of chemically heterogeneous magma storage regions over relatively short timescales
 prior to eruption.

342 The minor element chemistry of the olivine crystals allows us to investigate the crystallinity of these chemically diverse magma storage regions. Olivine crystals in our Floreana lava and scoria deposits 343 have an unusually low, and large range of Ca concentrations (~250-2600 ppm compared with ~1500-344 3000 in the eastern Galápagos; Gleeson and Gibson, 2019; Fig. 5A). The lower end of the range in Ca 345 concentrations measured in the Floreana lava and scoria deposits overlaps with those observed in 346 347 cumulate xenoliths (wehrlites) and thus are unlikely to represent mantle olivines (Thompson and Gibson, 2000). Previous studies have shown that the Ca concentration of magmatic olivine is sensitive 348 to several parameters, including: (i) the temperature of the system (Adams and Bishop, 1982; Köhler 349 and Brey, 1990; Shejwalkar and Coogan, 2013); (ii) the  $H_2O$  content of the co-existing melt phase 350 351 (Gavrilenko et al., 2016); and (iii) the major element composition of the co-existing melt phase (Herzberg, 2011). However, the diversity of Ca concentrations in our Floreana olivines is too large to 352 353 be caused by variations in T alone (assuming that the difference between the crystallisation temperature and final storage temperature is <300°C; see Section 6.4). In addition, H<sub>2</sub>O contents of ~3 354 355 wt% would be required to explain the lowest Ca concentrations in the Floreana olivines (according to the hygrometer of Gavrilenko et al., 2016), which is far in excess of that measure in Floreana 356 357 submarine glasses by Peterson et al. (2017). Hence, Ca variations in Floreana olivines likely result 358 from compositional variability in the co-existing melt phase.

Although olivine Ca variations are often interpreted to reflect lithological heterogeneity of the mantle source (resulting in primary melt compositional variability), this is inconsistent with the Ni contents of olivine crystals from Floreana (Fig. 5B; Gleeson and Gibson, 2019). An alternative process is therefore required to explain the variability of olivine Ca concentrations and, specifically, the presence of olivine crystals with <1000 ppm Ca. This process must reduce the Ca concentration of the melt phase (and co-existing olivines), without simultaneously reducing the melt Mg# (as low Ca

365	concentrations are observed across the entire range of forsterite contents in the Floreana olivines; Fig.
366	5A) and may be accompanied by slight variations in T or melt $H_2O$ concentration.
367	Evidence for the origin of the low-Ca olivines in the lava and scoria deposits is present in texture and
368	composition of the wehrlitic xenoliths, which contain uniformly low-Ca olivine crystals (<1000 ppm;
369	Fig. 5A). The petrography of the wehrlitic xenoliths attests to clinopyroxene growth within olivine-
370	dominated mush regions. Clinopyroxene crystallisation within this mush would extract CaO and MgO
371	from the residual melt. However, in an olivine-rich mush, the large reservoir of MgO contained within
372	the cumulus olivine grains would buffer the residual melt at a near-constant Mg# during
373	clinopyroxene crystallisation (Meyer et al., 1989). In contrast, the CaO concentration of the melt is
374	not buffered and decreasing melt CaO contents, due to clinopyroxene crystallisation, will cause the
375	CaO concentration of cumulus olivine grains to decrease (as a result of diffusive re-equilibration).
376	Intercumulus clinopyroxene growth would also increase the H <sub>2</sub> O concentration of the residual melt
377	phase, decreasing the partition coefficient of Ca into olivine (Gavrilenko et al., 2016). Therefore, if
378	these mush-zone crystals are entrained into ascending melts prior to eruption, growth of
379	clinopyroxene within an olivine-dominated mush could explain the presence of anomalously low Ca
380	concentrations in the Floreana olivines separated from lava and scoria deposits.
381	As the majority of olivine analyses from the Floreana lava and scoria deposits have low Ca
382	concentrations (<1000 ppm) that overlap with those in xenolithic nodules, we suggest that a large
383	proportion of the erupted crystal cargo derives from highly crystalline magma storage regions. Higher
384	Ca concentrations (>1500 ppm) are only found in a small number of forsteritic crystals (Fo>83) in the
385	lava and scoria samples (Fig. 5A). As olivine crystals formed from peridotitic melts that have not
386	undergone clinopyroxene growth will typically have Ca concentrations >1500 ppm (Gleeson and
387	Gibson, 2019; Herzberg, 2011), we interpret these crystals to have grown within liquid-rich magma
388	storage regions where magmatic differentiation occurs by fractional crystallisation.
389	However, olivine crystals from the Floreana lava and scoria deposits cannot simply be divided into
390	low Ca (<1000 ppm) and high Ca (>1500 ppm) populations (Fig. 5A). A substantial number of olivine
391	crystals have intermediate compositions ( $Ca = 1000-1500$ ). We interpret these as being sourced from

392	regions where growth of clinopyroxene was ongoing at the time of mush disaggregation. Thus, the
393	olivine crystal cargo of the Floreana magmas is predominantly derived from crystal-rich domains that
394	vary from highly crystalline (Ca <1000 ppm) to moderately crystalline (Ca = 1000–1500 ppm). Only
395	a small number of olivine crystals preserve compositions consistent with fractional crystallisation in
396	liquid-rich storage regions (Ca >1500 ppm).
397 398	6.2.2 Insights from clinopyroxene major element compositions The compositions of clinopyroxene crystals from the Floreana scoria also overlap with those in our
399	xenolith samples, supporting the hypothesis that some of the erupted crystals are derived from
400	disaggregated sub-volcanic mush. We used hierarchical cluster analysis to subdivide our 567
401	clinopyroxene major element analyses from the Floreana scoria and xenolith samples to determine the
402	proportion of material that is derived from each xenolith lithology in the erupted crystal cargo. We
403	find that our clinopyroxene analyses form three distinct groups (Fig. 8):
404	- Group 1 clinopyroxenes are predominantly from the wehlite and dunite xenoliths and include
405	90% of our analyses from these samples. 39% of clinopyroxenes analysed from the scoria
406	samples also fall into this group.
407	- Group 2 clinopyroxenes include all analyses from the gabbroic xenoliths, and ~10% of
408	analyses from crystals separated from the scoria samples.
409	- Group 3 clinopyroxenes are dominantly analyses from scoria derived clinopyroxenes (~50%
410	of analyses from the scoria separates). However, 10% of analyses from the wehrlite and
411	dunite xenoliths also fall into this group.
412	Of our 248 clinopyroxenes analysed from the Floreana scoria samples, approximately half are
413	classified as Group 3 and thus have major element compositions that do not show any affinity to
414	either the wehrlite/dunite or gabbroic cumulates. Therefore, these crystals may represent autocrysts
415	(with growth in liquid-rich magma storage regions). The remainder of clinopyroxene analyses from
416	the scoria samples are either compositionally analogous to those in the wehrlite and dunite xenoliths
417	(Group 1; 39%) or the gabbroic xenoliths (Group 2; 11%); we interpret these as representing
418	disaggregated sub-volcanic mush or wall rock. The high proportion of the clinopyroxene crystal cargo

that is derived from highly crystalline storage regions beneath Floreana is consistent with ourinterpretation of olivine minor element concentrations.

### 421 **6.3 Reactive Porous Flow within a cumulate mush**

422 Whilst the olivine and clinopyroxene major and minor element concentrations show that a large proportion of the erupted crystal cargo is derived from highly crystalline magma storage regions, 423 clinopyroxene trace element concentrations (and zoning) reveal the magmatic processes that operate 424 within these crystal-rich domains. The trace element composition of melts in equilibrium with our 425 clinopyroxene crystals are calculated using the model of Wood and Blundy (1997). The results 426 indicate that many of our clinopyroxene analyses have incompatible trace element ratios (e.g. 427  $[Ce/Y]_n$  which are more enriched than any erupted basalt from Floreana (Harpp et al., 2014a). In fact, 428 429 nearly all clinopyroxenes analyses from our xenolith samples, and ~50% of clinopyroxene analyses from the scoria samples, record trace element disequilibrium with the typical trace element 430 431 composition of the Floreana basalts (Fig. 7 and 9). Over-enriched equilibrium melt signatures are 432 characteristics of Group 1 clinopyroxenes (i.e. chemical affinity to the wehrlitic or dunitic xenoliths), 433 whereas crystals that are near trace element equilibrium with Floreana basalts typically have Group 3 434 major element compositions (i.e. the autocryst group).

Petrographic observations and olivine minor element data indicate that the Floreana sub-volcanic 435 system is characterised by clinopyroxene crystallisation within an olivine-dominated mush. If the 436 clinopyroxene grew from trapped melt within an olivine-dominated mush, progressive crystallisation 437 438 would increase the concentration of highly incompatible trace elements (e.g. Ba, La, Ce) relative to less incompatible trace-elements (e.g. Sm, Y) in the residual melt. A simple fractional crystallisation 439 model indicates that  $\sim 80\%$  crystallisation is required to generate melt  $[Ce/Y]_n$  ratios that are in 440 equilibrium with enriched clinopyroxenes from the scoria samples and even greater extents of 441 442 crystallisation (~90%) would be required to generate the extremely high  $[Ce/Y]_n$  ratios in some of the 443 xenolithic clinopyroxenes (Fig. 9). Such extensive crystallisation would be expected to result in the 444 saturation and crystallisation of plagioclase and other accessory phases (e.g. apatite,

magnetite/ilmenite, quartz), which are observed in more evolved xenoliths from Rabida island in the
central Galápagos (Holness et al., 2019). However, these phases are absent in the Floreana xenoliths,
indicating that another process is responsible for generating the anomalous trace element signatures of
the melts in equilibrium with our clinopyroxenes.

One possible mechanism for generating the observed trace element over-enrichment is reactive porous 449 flow, where clinopyroxene saturated melts ascend through highly-crystalline magmatic systems 450 causing dissolution of the existing crystal framework and precipitation of clinopyroxene (Lissenberg 451 452 and MacLeod, 2016). Reactive porous flow can result in net replacement of olivine by clinopyroxene, 453 leading to enrichment of highly- to moderately-incompatible trace elements in the resulting melt and co-existing crystal phases (Coogan et al., 2000; Gao et al., 2007; Lissenberg et al., 2013; Lissenberg 454 and MacLeod, 2016), and is consistent with the petrography of the Floreana xenoliths. For example, 455 major element maps of clinopyroxene crystals in the Floreana wehrlites show that they are zoned, 456 457 with Ti-rich rims (Fig. 10); equivalent zoning patterns have been attributed to reactive porous flow in plutonic clinopyroxenes from the oceanic crust (e.g. Hess Deep; Lissenberg and MacLeod, 2016). 458 459 Furthermore, reactive porous flow is consistent with the small size and rounded morphology of 460 olivine chadacrysts. In addition, if pre-existing Cr-spinel was dissolved by the reacting melt, then 461 reactive porous flow could also explain the high Cr contents of clinopyroxene in our wehrlitic 462 xenoliths (Fig. 6; Lissenberg and MacLeod, 2016).

To test whether reactive porous flow of clinopyroxene saturated melts through an olivine-dominated mush is consistent with the trace element compositions of melts calculated to be in equilibrium with our Floreana clinopyroxenes, we use the zone refining model of Harris (1957; Fig. 9):

466 
$$\frac{C_l}{C_l^o} = \frac{1}{D} - (\frac{1}{D} - 1)^{-DI}$$

467

468 where *D* is the bulk partition coefficient;  $C_l^o$  and  $C_l$  are the initial and final concentration of that 469 element in the melt phase, respectively; and *I* is the 'equivalent volumes of solid processed by the

(eq. 1)

470 liquid' (Lissenberg and MacLeod, 2016). The model assumes that a migrating melt front depletes solid phases of their incompatible trace elements owing to partial melting, and has previously been 471 employed to investigate geochemical signatures in oceanic gabbros (Lissenberg and MacLeod, 2016). 472 473 The model produces melts with trace element compositions that are comparable with those in 474 equilibrium with our Floreana clinopyroxenes (i.e.  $[Ce/Y]_n \sim 8-13$ ) using I values similar to that 475 invoked in other magmatic settings worldwide (~4–10; Lissenberg and MacLeod, 2016). Hence, 476 reactive porous flow represents a realistic mechanism for generating the geochemical diversity of 477 Floreana clinopyroxenes, including the trace element enriched crystals analysed in the wehrlitic 478 nodules (Fig. 9).

479 In addition, detailed LA-ICP-MS transects of two clinopyroxene grains from the most enriched 480 wehrlitic xenolith analysed in this study (17MMSG02c) show clear trace element zoning (Fig. 11). 481 The core of the larger clinopyroxene crystal has low [Ce] and [Ce/Y]<sub>n</sub> contents that are approximately 482 in equilibrium with Floreana basalts (Harpp et al. 2014a; Fig. 11a), whereas the mantle is 483 characterised by increasing [Ce] and  $[Ce/Y]_n$  contents towards the rim. We interpret this as core crystallisation from a melt with a trace element signature similar to that of erupted Floreana basalts 484 (Harpp et al., 2014a), followed by growth from a melt which became progressively enriched during 485 486 reactive porous flow (Fig. 11).

487 The mantle of the second, smaller xenolithic clinopyroxene shows a similar rim-ward increase in [Ce] 488 (interpreted as progressive melt enrichment during reactive porous flow). However, the [Ce] and  $[Ce/Y]_n$  values of the crystal core are too high to be in equilibrium with erupted Floreana basalts (Fig. 489 490 11C). This is consistent with spot analyses of clinopyroxene cores in other crystals analysed in this study. The high [Ce] and  $[Ce/Y]_n$  values in crystal cores cannot be explained by inward diffusion of 491 492 Ce, owing to significant differences in the diffusivities of Ce and Y and similar [Ce] and [Y] zoning 493 patterns in our two crystal transects (Fig. 11; Van Orman, 2001). Instead, we suggest that the high apparent core [Ce] and [Ce/Y]<sub>n</sub> contents in many of the Floreana clinopyroxenes record crystallisation 494 495 from melts that had already undergone geochemical enrichment via reactive porous flow. However, we cannot discount that our apparent clinopyroxene cores are fragments of larger oikocrysts that have 496

497 been broken during mush disaggregation or sample crushing and, as a result, do not represent the true498 core compositions of each crystal.

499 Nevertheless, our clinopyroxene major and trace element data, as well as petrographic observations of
500 the wehrlitic xenoliths, provide substantial evidence that reactive porous flow is an important
501 mechanism of melt migration and melt differentiation in highly crystalline magma storage regions
502 beneath Floreana. Although reactive porous flow has been identified as an important process in MOR
503 gabbros, this is the first study to identify reactive porous flow in an ocean island setting.

### 504 6.4 Petrographic estimates of magma storage pressures

Petrological and geophysical constraints on magma storage depths exist for several recently active 505 volcanoes in the western Galápagos Archipelago (Bagnardi et al., 2013; Case et al., 1973; Geist et al., 506 507 1998; Stock et al., 2018; Vigouroux et al., 2008). However, in the absence of geophysical data (owing 508 to a paucity of recent eruptions), there are far fewer constraints on the structure of magma storage 509 regions in the eastern and south-eastern archipelago. To date, the only investigation of magma storage depths beneath these volcanoes is by Geist et al. (1998), who undertook a visual comparison between 510 whole-rock lava compositions and the MORB olivine + plagioclase + augite + melt pseudoinvariant 511 512 point, parameterised by Grove et al. (1992). This approach is subject to substantial uncertainty, but the authors suggest that Floreana magmas consistently equilibrate at >5 kbar (typically >7 kbar) at a 513 depth >16 km, within the upper mantle. 514

515 We used three petrological barometers to provide improved constraints on magma storage depths 516 beneath Floreana. First, we applied the clinopyroxene-only barometer of Putirka (2008), in which 517 pressure and temperature are solved iteratively based solely on the clinopyroxene major element composition (primarily the jadeite component; standard error of estimate [SEE] =  $\pm 310$  MPa). 518 Second, we applied the clinopyroxene-melt barometer of Neave and Putirka (2017), which uses the 519 520 composition of a co-existing melt phase and the proportion of the Jadeite component in clinopyroxene to calculate the pressure of crystallisation (SEE =  $\pm 140$  MPa; pressure is solved iteratively with 521 temperature using the clinopyroxene-melt thermometer of Putirka, 2008). Third, for the xenolithic 522

523 nodules, we estimate the final pressure and temperature of storage using the two-pyroxene

524 thermobarometer of Putirka (2008) (SEE =  $\pm 260$  MPa).

525 Taken at face value, initial application of the clinopyroxene-only barometer to all clinopyroxene 526 analyses from the scoria and xenolith samples gives a range of pressure estimates between ~450 MPa and ~1800 MPa. However, reactive porous flow has a substantial influence on the compositions of the 527 Group 1 (and Group 2) clinopyroxenes, which may influence the barometric results. Specifically, 528 crystals that show evidence for reactive porous flow also have elevated Na concentrations, leading to 529 530 an anomalously high jadite component. Therefore, we filter our dataset to remove crystals that show a 531 chemical signature indicative of reactive porous flow and only use Group 3 clinopyroxenes that have trace element compositions in equilibrium with the Floreana basalts (using the whole-rock data from 532 Harpp et al. 2014) in our barometric calculations (n=78). Results indicate that crystallisation beneath 533 534 Floreana occurs at a pressure of 766  $\pm$ 322 MPa (2 $\sigma$  of calculated pressures), which equates to a depth of 25.2 ±9.9 km (using the crustal density estimate of Putirka (1997) and a mantle density estimate of 535 3300 km/m<sup>3</sup>; Fig. 12). 536

537 Application of the Neave and Putirka (2017) clinopyroxene-melt barometer requires identification of equilibrium clinopyroxene-liquid pairs. We achieve this using an automated melt-matching algorithm 538 (as in Winpenny and Maclennan, 2011, Neave and Putirka, 2017, Stock et al. 2018), with  $K_{\rm D}$  (Fe-Mg), 539 540 diopside-hedenbergite, enstatite-ferrosillite and calcium Tschermak's equilibrium tests ( $K_D$ (Fe-Mg)) within ±0.03 other components within 2 SEE; Putirka, 1999, Putirka, 2008, Mollo et al., 2013). We 541 used the whole-rock data of Harpp et al. (2014a) and basaltic glass analyses from this study as 542 potential equilibrium liquids. Input crystal compositions were again filtered to remove analyses that 543 544 showed evidence of reactive porous flow (i.e. only Group 3 clinopyroxenes in equilibrium with the Floreana whole-rock were used). In total, 70 of the 78 input clinopyroxene analyses returned at least 545 one equilibrium match to either the basaltic glass or whole-rock compositions. Where clinopyroxene 546 547 compositions produced an equilibrium match with more than one equilibrium melt, an average melt 548 composition was used in the barometric model. Results from this barometer indicate that magma 549 crystallisation occurred at 717  $\pm$ 165 MPa (23.7  $\pm$ 5.1 km) and 1224  $\pm$ 33°C (Fig. 12).

Clinopyroxene-orthopyroxene thermobarometry records the final storage conditions of the cumulate xenoliths, rather than the crystallisation conditions of clinopyroxene autocrysts (orthopyroxene is only found as an intercumulus phase). Temperature and pressure estimates were only calculated from orthopyroxene-clinopyroxene pairs in wehrlite and dunite xenoliths that passed the  $K_D$ (Fe-Mg) equilibrium test of Putirka (2008; within ±0.14). Results suggest that the cumulates were stored at ~975–1100°C and 600–900 MPa, with a mean storage pressure of 712 ±200 MPa (23.7 ±6.4 km; Fig. 12).

557 The depths of magma storage calculated from our three petrological barometers show an excellent

agreement within the model uncertainties. These new data provide robust evidence that magma

storage beneath Floreana occurs below the Moho (~16 km; Feighner and Richards, 1994), in the upper
mantle.

# 7 IMPLICATIONS FOR MAGMATIC PLUMBING SYSTEMS BENEATH LOW MELT FLUX OCEAN ISLAND VOLCANOES

Our new petrological and geochemical data show that magma storage beneath Floreana occurs in 564 mush-dominated regions in the upper mantle (Fig. 13). Mineral chemistry (such as low olivine Ca 565 566 concentrations and clinopyroxene major elements) reveal that a substantial portion of the erupted crystal cargo is derived from disaggregated mush and wall rock material which has been entrained 567 into the ascending magmas. During ascent, magmas may entrain coherent nodules (xenoliths) as well 568 as disaggregated mush (Fig. 13). Coherent nodules represent areas of the magmatic system beneath 569 570 Floreana that have undergone cooling to temperatures <1100°C (compared to the clinopyroxene 571 crystallisation temperatures of  $\sim 1225^{\circ}$ C) and may represent material from the border of the active 572 mush zone or older, almost completely solidified magma storage regions that are intersected during 573 magma ascent (Fig. 13).

574 Petrographic observations and clinopyroxene trace element chemistry from both the xenolith and
575 scoria samples reveal that clinopyroxene growth occurs via reactive porous flow in the mush-

576 dominated areas beneath Floreana. Reactive porous flow causes distinct trace element enrichment in the percolating melt phase and crystallising clinopyroxene, which can explain the trace element 577 disequilibrium between the erupted Floreana basalts and their clinopyroxene cargo. Nevertheless, the 578 presence of some clinopyroxene crystals with major and trace element compositions in equilibrium 579 580 with erupted Floreana basalts indicates that at least some crystallisation occurs in liquid-rich subvolcanic storage regions, likely situated as localised melt pockets within the larger mush zone (Fig. 581 582 13). Transport of melts modified by reactive porous flow into these melt pockets could impact the 583 LREE enriched signature of the resultant hybridised melts. This could explain the LREE-enriched 584 signature in the Floreana basalts, which is not seen in other regions of the Galápagos Archipelago 585 (Harpp et al., 2014a).

Our results indicate substantial differences in the architecture of the magmatic systems beneath 586 587 Floreana and the frequently active shield volcanoes in the western Galápagos Archipelago. For 588 example, previous petrological and geophysical studies have identified that western Galápagos magmatic systems are characterised by crustal magma storage, often with a large storage region in the 589 590 mid-to-lower crust (~7 km depth) and a smaller storage region at shallow levels, within the volcanic 591 edifice (~1 km depth; Geist et al. 1998; Bagnardi et al. 2013; Bagnardi and Hooper, 2018; Stock et al., 592 2018; Fig 12). In contrast, our barometric data indicate that magmas beneath Floreana ascend directly 593 from the upper mantle and undergo no detectable crustal storage. Although mush-rich regions have 594 been inferred beneath the western Galápagos shield volcanoes (based on whole-rock data and the 595 presence of gabbroic glomerocrysts; Chadwick et al., 2011; Geist et al., 1995, 2014; Stock et al., 596 2018), magmatic differentiation appears to be driven by simple fractional crystallisation and mixing 597 of chemically diverse magmas (Geist et al., 1995; Naumann and Geist, 1999; Stock et al., in review). One major factor that differentiates Floreana from shield volcanoes in the western archipelago (on 598 599 Isabela and Fernandina) is the flux of magma into the lithosphere, as evidenced by variations in the 600 volumetric eruption rate (Harpp et al., 2014a; Harpp and Geist, 2018; Kurz et al., 2014). Hence, we 601 suggest that the greater pressure of magma storage and prevalence of reactive porous flow beneath 602 Floreana, relative to volcanoes in the western archipelago, are related to the substantially lower flux of

603 magma into the lithosphere from the underlying mantle source (and thus the thermal structure of the 604 lithosphere). For example, the magma flux entering the lithosphere beneath Wolf volcano (northern Isabela) has been substantially greater than that beneath Floreana for several 100,000s of years. The 605 high magma flux beneath Wolf maintains the average temperature of the mid-to-lower crust at 606 607 ~1125°C ( $\Delta T$  ~22 °C), with only small-scale thermal and compositional heterogeneities present in the sub-volcanic plumbing system (Geist et al., 2014, 2005; Stock et al., 2018, in review). In contrast, the 608 609 flux of magma entering the magmatic system beneath Floreana is much lower and the temperature of 610 the mid-crust is likely to be  $<<800^{\circ}$ C (i.e. significantly cooler than the lowest temperature recorded by 611 the Floreana xenoliths; Fig. 12). As the flux of magma (and heat) from the mantle is insufficient to 612 maintain an elevated crustal geotherm beneath Floreana, magmas that stall in the crust are likely to rapidly crystallise, increase their viscosity, and become uneruptable. Therefore, eruptions must be fed 613 614 by melts ascending from much deeper storage regions (~700-750 MPa) where super-solidus melts can 615 persist over long time periods.

Our results have global implications for the architecture and dynamics of magma storage regions 616 beneath ocean island volcanoes that are characterised by a relatively low flux of magma from the 617 underlying mantle. These include off-axis volcanic systems (e.g. Halekala, Hawaii) and those above 618 619 low buoyancy flux mantle plumes (e.g. El Hierro, Canary Islands). Figure 14 shows a compilation of published barometric estimates from 'high' and 'low' melt flux systems in some of the most active 620 621 ocean island systems on Earth (Hawaii, Iceland and Galápagos; Hammer et al., 2016; Hartley et al., 622 2018; Neave and Putirka, 2017; Poland et al., 2015; Stock et al., 2018). Central volcanic systems from 623 each region typically display relatively low-pressure magma storage compared to regions that are 624 characterised by a comparatively low magma flux from the convecting mantle (i.e. off-axis volcanic 625 systems; Fig. 14). This observation is consistent with the high pressures of magma storage identified 626 at other ocean island volcanoes that are located above mantle plumes with a relatively low buoyancy 627 flux (e.g. >15-25 km at El Hierro in the Canary Islands; Longpre et al., 2014; Taracsák et al., 2019). Hence, we speculate that high-pressure magma storage is characteristic of low melt flux ocean island 628 629 volcanoes globally.

### 630 8 CONCLUSIONS

Petrographic and geochemical analyses of lava, scoria and xenolith samples from Floreana in the 631 632 south-eastern Galápagos Archipelago provide new insights into the architecture and dynamics of 633 magma storage beneath low melt flux ocean island volcanoes. Comparison of olivine and clinopyroxene major, minor and trace element contents between our different sample types reveals 634 that a substantial portion of the erupted crystal cargo is derived from mush-dominated magma storage 635 regions beneath Floreana. Mineral textures, highly enriched clinopyroxene trace element signatures 636 637 and trace element zoning in the xenoliths reveals that reactive porous flow is an important process of chemical differentiation and melt transport within these mush-dominated regions. Mixing between 638 melts that have been geochemically enriched by reactive porous flow and those in overlying liquid-639 rich storage regions could explain the anomalous LREE enriched signature of the Floreana basalts, 640 641 which is absent in other parts of the Galápagos Archipelago where reactive porous flow has not been identified. 642

643 Application of independent petrological barometers to crystals in Floreana scoria and xenolith samples indicates that magmas are stored in the upper mantle ( $\sim 23.7 \pm 5.1$  km). Floreana is in a distal 644 location to the Galápagos plume where the melt flux entering the lithosphere is low; the depth of 645 magma storage beneath Floreana contrasts with more proximal, higher melt flux volcanoes in the 646 western archipelago where magmas are stored in the crust (Geist et al., 1998; Stock et al., 2018). 647 Comparing our new data with ocean island volcanoes globally (e.g. Hawaii, Iceland and the Canary 648 649 Islands) reveals that the Galápagos is not unique and that magma storage is ubiquitously shallower in proximal magmatic systems above high buoyancy flux plumes than in off-axis systems, or above low 650 buoyancy flux plumes. We therefore suggest that the flux of mantle-derived magma entering the 651 652 lithosphere imparts a first-order control on the depth of magma storage beneath ocean island 653 volcanoes.

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### 919 **FIGURES**

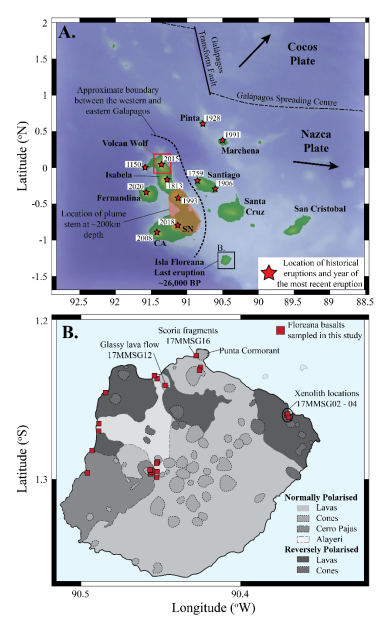
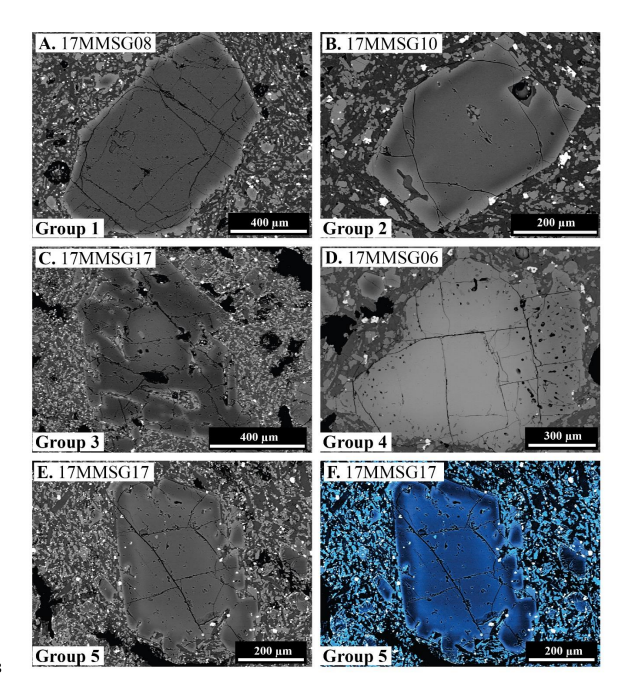


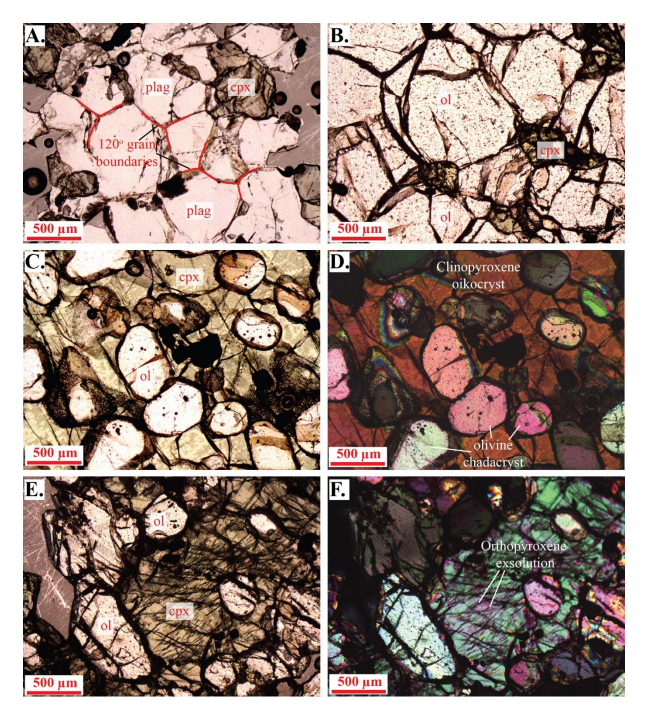


Figure 1 - A. Regional map of the Galápagos Archipelago highlighting the location of Isla Floreana,
Cerro Azul (CA), Sierra Negra (SN) and Wolf volcanoes. Dates show the most recent eruptions at
historically active volcanic centres. Black arrows show the direction of plate motion for the Nazca and
Cocos tectonic plates, respectively. B. Geological map of Floreana adapted from Harpp et al. (2014a).
Dashed lines delineate monogenetic scoria cones. Normally and reversely polarised lava flows are
shown along with the largest (Cerro Pajas) and most recent (Alayeri; ~26,000 years) eruptions on
Floreana.



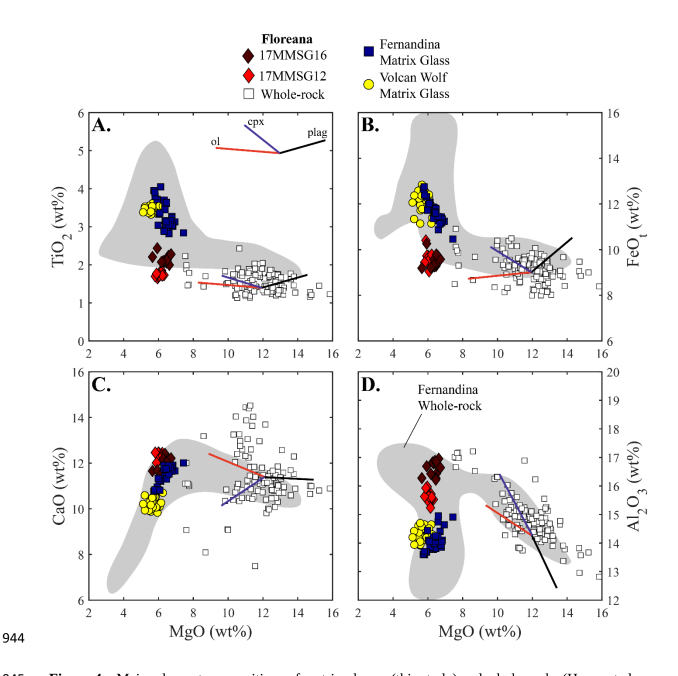
928

Figure 2 - SEM images of A. Group 1 olivines – euhedrual to subhedral crystal morphologies with
large, unzoned, crystal cores and narrow, normally-zoned rims. B. Group 2 olivines – subhedral to
euhedral crystals with clear, reverse-zoning profiles. C. Group 3 olivines – skeletal crystals with high
forsterite overgrowths on low forsterite cores. D. Group 4 olivines – anhedral crystals with sieved
textured, reverse zoned rims. E. (greyscale) and F. (false colour) Group 5 olivines – crystals preserve
at least 4 composition zones over ~100-200 µm.



936

Figure 3 - Plane Polarised Light (A. – C. and E.) and Crossed Polarised Light (D., F.) images of
Floreana xenoliths. A. – gabbroic xenolith (17MMSG04b), highlighting near 120° grain boundaries at
monomineralic plagioclase (plag) triple junctions. B. – dunitic xenolith (17MMSG04c) with
intercumulus clinopyroxene growth. C. and D. – wehrlitic xenolith (17MMSG02c) showing a large
clinopyroxene (cpx) oikocryst surrounding olivine (ol) chadacrysts. E. and F. – wehrlitic xenolith
(sample 17MMSG03a) showing olivine chadacrysts within a clinopyroxene oikocryst. Orthopyroxene
exsolution lamallae are visible within the clinopyroxene.



945Figure 4 – Major element compositions of matrix glasses (this study) and whole-rocks (Harpp et al.,9462014a) from Floreana, as well as glasses from Fernandina (Peterson et al., 2017) and Wolf volcano947(Stock et al., 2018) in the western Galápagos Archipelago. Lines show trajectories of liquid948compositional evolution olivine (ol; red), clinopyroxene (cpx; blue) and plagioclase (plag; black)949crystallisation. The grey field shows whole-rock data from Isla Fernandina in the western Galápagos950(Allan and Simkin, 2000; Geist et al., 2006). The  $2\sigma$  precision of our matrix glass analyses is smaller951than the symbol size.

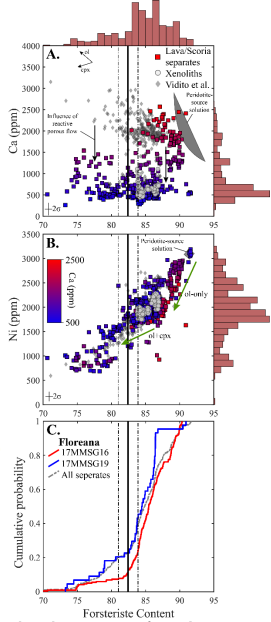
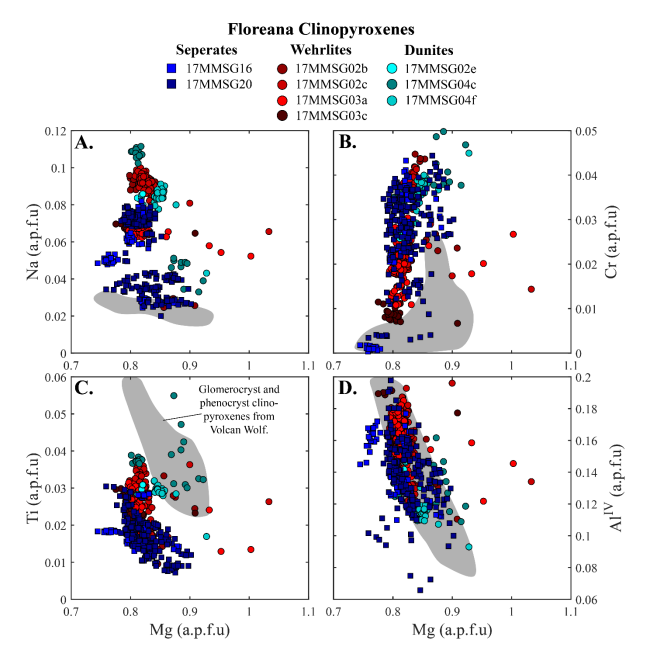


Figure 5 – Major and minor element compositions of olivine crystals from the Isla Floreana basalts. A. Fo vs. Ca and **B** Fo vs. Ni in Galápagos olivine crystals with analyses from our lava/scoria separates and xenolith samples, as well as a compilation of available olivine data from Floreana (Vidito et al. 2013). Our lava/scoria analyses are coloured according to their Ca concentration (see colour scale in **B**). The histograms above and to the right of the plots show the data distributions (excluding in situ analyses of xenolithic olivines). Peridotite source solutions are taken from Herzberg (2011) and Matzen et al. (2017a). Black arrows in A. show the trajectory of crystal compositional evolution during olivine (ol) and clinopyroxene (cpx) crystallisation (taken from Gleeson and Gibson, 2019) and the hypothesised influence of reactive porous flow. The green lines in

971 **B.** show the trajectories of crystal compositional evolution during olivine only, followed by olivine + 972 clinopyroxene fractional crystallisation (from Gleeson and Gibson, 2019). C. Cumulative probability 973 of forsterite in olivine separates from our lava (17MMSG19) and scoria (17MMSG16) samples. A 974 two-sample Kolmogorov-Smirnov test is used to assess the similarity of the forsterite distributions of 975 the two samples (and all analyses from lava and scoria deposits in this study). Results indicate that the 976 olivine populations from the two samples are drawn from the different distribution (p=0.0021). The 977 vertical black lines show the forsterite compositions of crystals calculated to be in equilibrium with the matrix glass composition of tephra sample 17MMSG16 ( $K_d = 0.30 \pm 0.03$  after Roeder and Emslie, 978 979 1970).



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981 **Figure 6** – Major element composition of Floreana clinopyroxenes from our scoria samples and 982 wehrlite and dunite xenoliths. The grey field shows the compositions of clinopyroxenes from Wolf 983 volcano in the western Galápagos Archipelago (from Stock et al. 2018). The  $2\sigma$  precision of our 984 clinopyroxene analyses is smaller than the symbol size.

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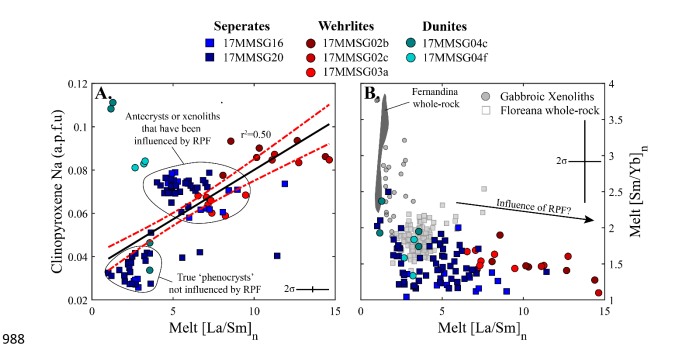


Figure 7 – A.  $[La/Sm]_n$  vs. Na in clinopyroxenes from our scoria samples and wehrlite and dunite 989 xenoliths. The black line shows a regression through the data ( $r^2 = 0.50$ ) and the red dashed lines 990 show the 95% confidence limits on the regression. **B.**  $[La/Sm]_n$  vs.  $[Sm/Yb]_n$  of melts calculated to be 991 in equilibrium with our Floreana clinopyroxenes using the model of Wood and Blundy (1997). The 992 black arrow shows the approximate trend of crystal compositional evolution hypothesised to occur as 993 994 a result of reactive porous flow. The grey field shows whole-rock compositions from Fernandina (Geist et al., 2006; White et al., 1993). B additionally shows the whole-rock compositions of erupted 995 Floreana lavas (Harpp et al., 2014a) and analyses of the gabbroic xenoliths from Floreana (this study). 996 997 Error bars show the fully propagated  $2\sigma$  precision of our analyses.

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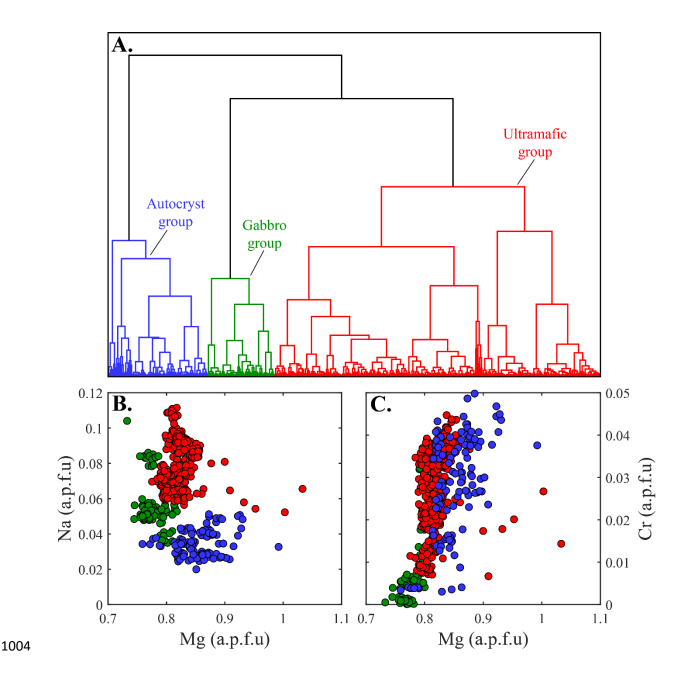
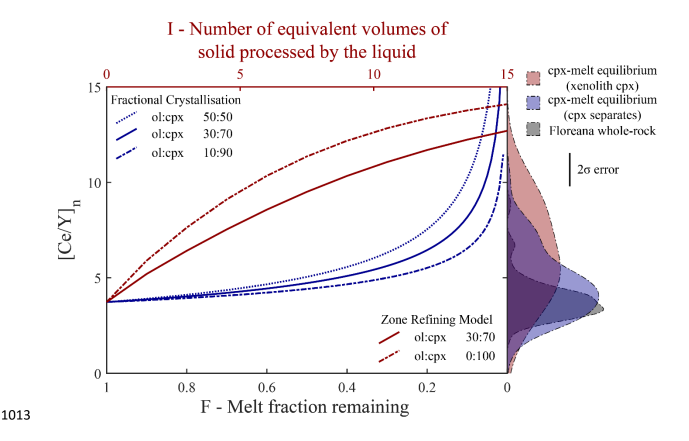


Figure 8 – A. Hierarchical cluster analysis of our clinopyroxene major element analyses. Colours
show the high-level division of crystal compositions into three groups: Group 1 is predominantly
comprised of crystals from wehrlite and dunite xenoliths (red), Group 2 is predominantly comprised
of crystals from gabbroic xenoliths (green) and Group 3 (blue) is dominated by crystals separated
from scoria samples. B Na vs Mg and C Cr vs Mg in our clinopyroxene analyses from the scoria and
xenoliths, coloured by their group.



1014 Figure 9 – Models showing the evolution of melt  $[Ce/Y]_n$  ratios as a function of the melt fraction 1015 remaining (F) during fractional crystallisation and as a function of the number of equivalent volumes 1016 of solid processed by the liquid (I) during reactive porous flow (red; using the zone-refining model of 1017 Harris, 1957). Line types show how models vary with different ratios of crystallising olivine (ol) and 1018 clinopyroxene (cpx). The kernel density plots to the right show the  $[Ce/Y]_n$  density distributions of liquids calculated to be in equilibrium with clinopyroxene crystals in our wehrlite and dunite xenoliths 1019 1020 (red), scoria samples (blue; 2<sup>o</sup> error for the clinopyroxene analyses is shown by the black bar), and 1021 whole-rock analyses of erupted Floreana basalts (grey; Harpp et al., 2014a). Partition coefficients were calculated for clinopyroxene using the method of Wood and Blundy (1997), the mean major 1022 1023 element composition of the group 3 clinopyroxenes from this study, a temperature of 1225°C, and a 1024 pressure of 700 MPa. Initial Ce and Y concentrations for both models were taken as the mean values 1025 from the whole-rock dataset of Harpp et al. (2014a). Calculation increments are 0.01 in F and 0.1 in I. 1026

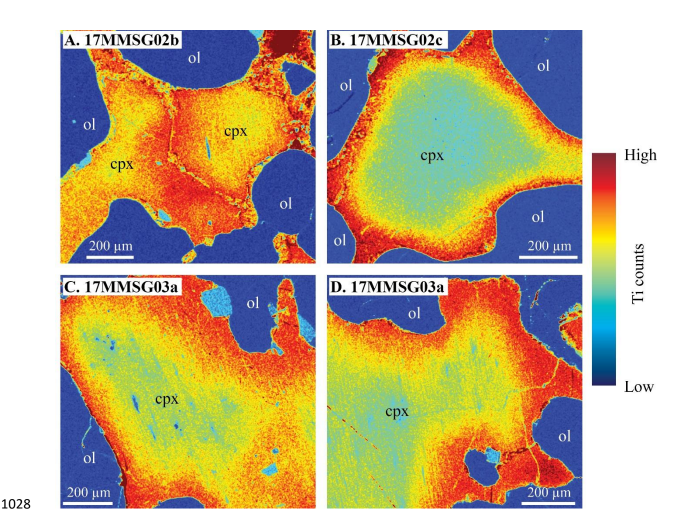


Figure 10 – Ti maps of key clinopyroxene crystals in the Floreana wehrlite xenoliths. These maps
display clear zoning in the xenolithic clinopyroxenes with Ti-poor cores and Ti-rich rims. The dark
blue regions surrounding clinopyroxene grains are olivine crystals.

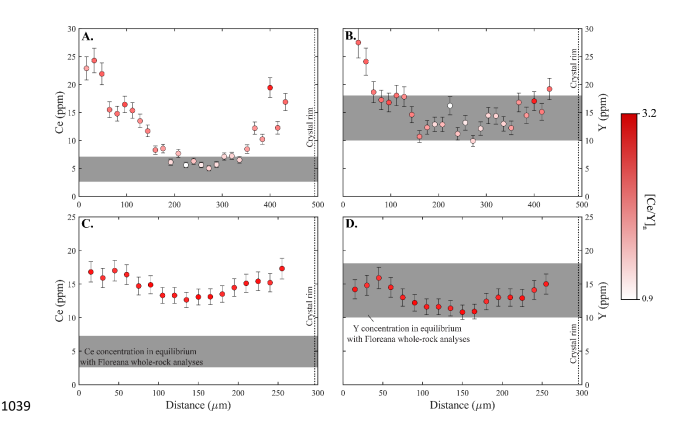


Figure 11 – LA-ICP-MS transects showing Ce (A, C) and Y (B, D) zoning across two clinopyroxene grains in a wehrlitic xenolith (17MMSG02c). A. and B. show a transect across the core of a large clinopyroxene oikocryst. C. and D. show a transect across the apparent core of a smaller clinopyroxene oikocryst. Points are coloured according to their [Ce/Y]<sub>n</sub> ratio (see colour scale). The grey bars show the crystal compositions calculated to be in equilibrium with whole-rock analyses of erupted Floreana basalts (Harpp et al., 2014a). 

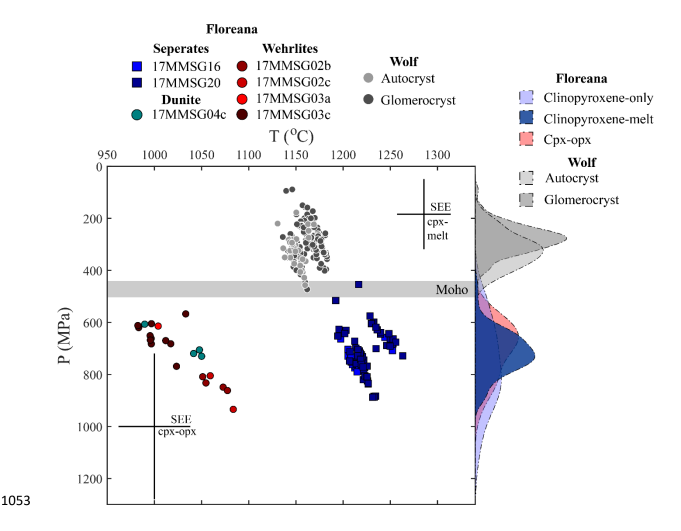
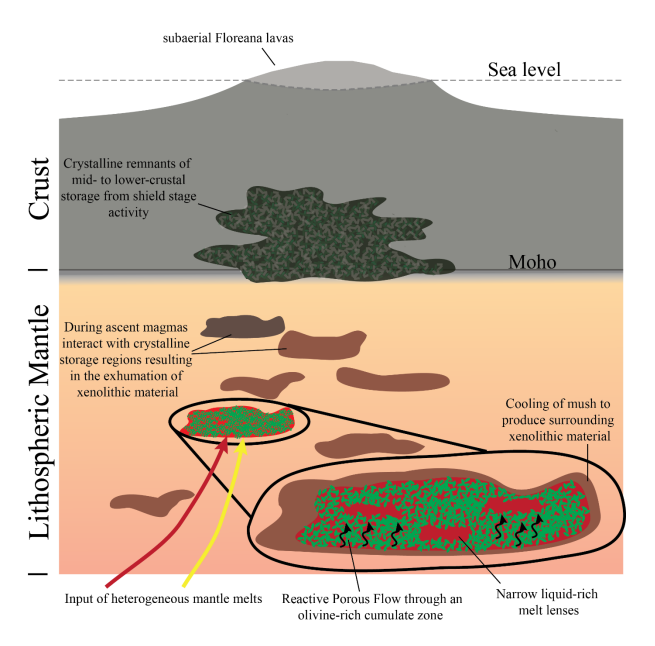
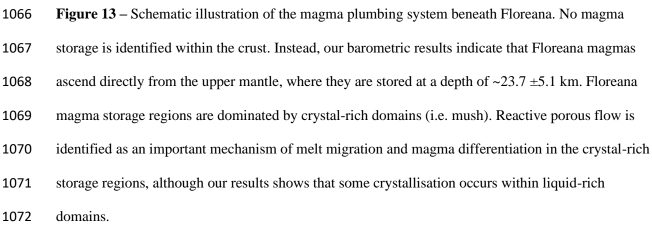


Figure 12 – Petrological thermobarometry results. The kernel density plots to the right show the
density distributions of barometric results from different models (light blue – clinopyroxene-only,
Putirka (2008); dark blue – clinopyroxene-melt, Neave and Putirka (2017); red – clinopyroxeneorthopyroxene Putirka (2008)). The grey bar shows the Moho depth beneath Fernandina (from
Feighner and Richards, 1994) and the grey points and kernel density estimates show clinopyroxenemelt thermobarometric results for autocrysts and glomerocrysts from Wolf volcano for comparison
(from Stock et al., 2018).





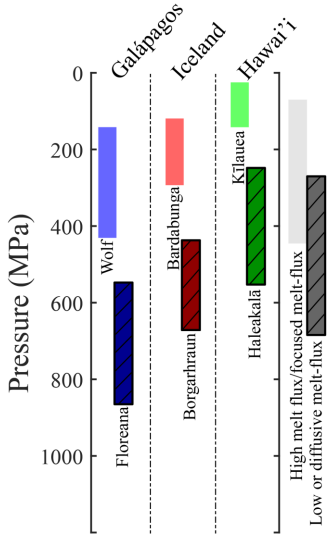


Figure 14 - Compilation of barometric estimates from different hotspot localities. Boxes without boarders show magma storage pressures at central volcanic systems, proximal to the centre of high buoyancy plumes (i.e. those with a high flux of mantlederived magma), and boxes with black boarders and dashes show magma storage pressures at volcanic systems that are off-axis (i.e. those with a low flux of mantle-derived magma). The pressure of magma storage is greater in regions of Hawai'i, Iceland, and the Galápagos that are characterised by a relatively low flux of mantle-derived magma. Barometric estimates are taken from: Poland

1091 et al. (2015) for Kīlauea, using multiple available data streams to create a model of summit magma 1092 storage (~25-150 MPa); Hammer et al. (2016) for Haleakalā using clinopyroxene melt barometry 1093 (~400  $\pm$ 160 MPa); Hartley et al. (2018) for Bardabunga using OPAM (210  $\pm$ 70 MPa); Neave and 1094 Putirka (2017) for Borgarhraun using clinopyroxene-melt barometry (570  $\pm$ 120 MPa); Stock et al. 1095 (2018) for Volcan Wolf using OPAM and clinopyroxene-melt barometry (280  $\pm$ 140 MPa; OPAM 1096 results), and this study for Floreana using clinopyroxene-melt barometry (717  $\pm$ 165 MPa).