1	High pressure trace element partitioning between clinopyroxene and alkali basaltic melts
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3	Barbara Bonechi ^a , Cristina Perinelli ^a , Mario Gaeta ^a , Alessandro Fabbrizio ^b , Maurizio Petrelli ^c ,
4	Ladislav Strnad ^d
5	
6	a: Dipartimento di Scienze della Terra, Sapienza Università di Roma, P. le Aldo Moro 5, 00185,
7	Rome, Italy
8	b: Institute of Petrology and Structural Geology, Faculty of Science, Charles University, Albertov 6,
9	12843 Prague, Czech Republic
10	c: Department of Physics and Geology, University of Perugia, Piazza Università, Perugia 06100, Italy
11	d: Laboratories of the Geological Institutes, Charles University, Albertov 6, Prague 2, CZ-12843,
12	Czech Republic
13	
14	
15	
16	Corresponding author:
17	Barbara Bonechi
18	Dipartimento di Scienze della Terra, Sapienza Università di Roma,
19	P. le Aldo Moro, 5 00185 Rome, Italy
20	E-mail: barbara.bonechi@uniroma1.it
21	ORCID: 0000-0002-0242-4320
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27 Abstract

28 We present new experimental data on major and trace element partition coefficients (D) between 29 clinopyroxene and a K-basaltic melt from Procida Island (Campi Flegrei Volcanic District, south 30 Italy). Time-series experiments were conducted at 0.8 GPa and 1080-1250 °C aiming to investigate 31 the role of the crystallization kinetics on trace elements partitioning behaviour at a pressure relevant 32 for deep magmatic reservoirs. Results indicate that large ion lithophile elements (LILE) are 33 incompatible (e.g., $D_{Sr} \le 0.15$), light rare elements (LREE; e.g., $D_{La} \le 0.20$) are always more 34 incompatible than heavy rare elements (HREE), which in some cases result to be compatible with clinopyroxene (e.g., $D_{Dy} = 1.40$); high field strength elements (HFSE) are generally incompatible 35 36 $(D_{\text{HFSE}} \le 0.8)$, while transition elements (TE) range from slightly incompatible (e.g., $D_{\text{V}} = 0.6$) to highly compatible (e.g., $D_{Cr} = 63$). The calculated D values for LILEs, REEs, HFSEs, and TEs tend 37 38 to decrease with the increase of temperature and to increase with increasing tetrahedrally-coordinated 39 aluminium content, in agreement with the previous studies. Moreover, we observed the influence of the growth rate on the partition coefficients, with the highest D_{REE} values calculated in the runs with 40 the highest growth rate ($\sim 10^{-7}$ cm s⁻¹), due to the less efficient rejection of incompatible elements 41 42 during rapid crystal growth, that in this study is not linked to disequilibrium conditions, but to the 43 presence of pre-existing nuclei. Additionally, the apparent increase in D_{REE} values with time observed 44 in some runs is not referable to a change in time but rather to the different degrees of polymerization, 45 expressed as the ratios NBO/T of these melts, strictly related to a loss of Fe occurred during the experiments, and thus to a different melt viscosity. Finally, the application of the experimental 46 47 clinopyroxene/melt partition coefficients highlights that the deepest step of the magmatic 48 differentiation in the Campi Flegrei Volcanic District is represented by the fractionation of about 20-49 30% of a clinopyroxenitic mineral assemblage from a basaltic parental magma.

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Keywords: clinopyroxene; trace element partition coefficient; lattice strain model; order-disorder
 crystal growth; fractional crystallization

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55 **1. Introduction**

A central factor in the evolution of an igneous body is the partitioning of elements between the melt 56 57 and the crystallizing minerals. The partition coefficient of an element i (D_i) has been defined as its 58 concentration in a mineral divided by the concentration of that element in the coexisting melt (Beattie 59 et al., 1993) and describes the chemical distribution between coexisting phases under equilibrium 60 conditions. Most partition coefficients, indeed, are usually investigated presupposing equilibrium 61 conditions, assuming thus that natural systems are characterized by chemical homogeneity of melt 62 and crystals and that these systems evolve slowly enough to allow for continuous re-equilibration 63 between melt and growing crystals (e.g., Onuma et al., 1968; Higuchi and Nagasawa, 1969; Philpotts 64 and Schnetzler, 1970; Schnetzler and Philpotts, 1970). However, disequilibrium conditions are not 65 unusual in natural systems, potentially affecting both absolute and relative partition coefficients, as observed by Albarede and Bottinga (1972). The resulting disequilibrium partition coefficients can be 66 used as sensitive monitors of magmatic processes such as fractional crystallization, assimilation, 67 68 mixing and partial melting of crustal or mantle rocks (e.g., Leeman and Phelps, 1981; Mahood and 69 Hildreth, 1983; Drexler et al., 1983; White, 2003; Henderson and Pierozynski, 2012; Arzilli et al., 70 2018). Indeed, as shown in several studies, in some natural igneous systems, events of rapid crystal 71 growth (driven by the undercooling degree) can produce an excessive accumulation or depletion of 72 trace elements, depending on their degree of compatibility or incompatibility, at the crystal-melt 73 interface with a consequent disequilibrium element uptake during crystal growth (e.g., Albarede and 74 Bottinga, 1972; Watson, 1996; Morgan and London, 2003; Watson and Müller, 2009; Mollo et al., 75 2011, 2013; Arzilli et al., 2018) with respect to equilibrium conditions (e.g., Morgan and London, 76 1996; Lofgren et al., 2006; Mollo et al., 2011, 2013). Therefore, trace elements partitioning can be 77 controlled by kinetic effects such as diffusive re-equilibration and undercooling degree (Arzilli et al.,

78 2018; Shea et al., 2019) as well as by temperature and pressure (Shimizu, 1974; Green and Pearson, 79 1983; Ray et al., 1983; Green and Pearson, 1985; Gallahan and Nielsen, 1992; Jones, 1995; Blundy 80 and Wood, 2003; McDade et al., 2003; Sun and Liang, 2012; Yao et al., 2012; Bédard, 2014; Mollo 81 et al., 2018), mineral composition (Yagi and Onuma, 1967; Onuma et al., 1968; Colson et al., 1988; 82 Blundy and Wood, 1991, 1994; Skulski et al., 1994; Lundstrom et al., 1998; Hill et al., 2000; Bédard, 83 2010; Mollo et al., 2016), melt composition (Mysen and Virgo, 1980; Ray et al., 1983; Nielsen, 1985; 84 Gallahan and Nielsen, 1992; Bennett et al., 2004; Gaetani, 2004), and redox conditions (Aigner-85 Torres et al., 2007; Cartier et al., 2014; Dygert et al., 2020). Among the works focused on the 86 influence of the crystallization kinetics on partition coefficients (Albarede and Bottinga, 1972; 87 Morgan and London, 2003; Lofgren et al., 2006; Milman-Barris et al., 2008; Mollo et al., 2011, 2013; 88 Arzilli et al., 2018; Shea et al., 2019), only a few have focused on clinopyroxene (Shimizu, 1981; 89 Lofgren et al., 2006; Mollo et al., 2013). The work of Shimizu (1981) investigates kinetic effects on 90 sector-zoned augite phenocrysts showing a clear case of disequilibrium crystallization. Lofgren et al. 91 (2006) performed equilibrium and dynamic crystallization experiments at cooling rates in the range 92 5-1000 °C/h to explore the effects of kinetics on D-values. Mollo et al. (2013) determined trace 93 element partition coefficients at the interface between clinopyroxene rims and co-existing glasses 94 resulting from isothermal and cooling rate experiments. The study of the factors influencing trace 95 element partitioning is of paramount importance since the latter are frequently used in petrological 96 and geochemical studies to better understand magmatic processes, like fractional or equilibrium 97 crystallization, assimilation, and partial melting (e.g., Mollo et al., 2016). This is because 98 clinopyroxene has crystal-melt REE partition coefficients approximately up to 4 orders of magnitude 99 higher than other mafic rock-forming minerals (e.g., olivine and orthopyroxene; Chen and Zhang, 100 2009; Sun, 2018). Clinopyroxene therefore, exerts a strong control on the overall partitioning of the 101 crystallising assemblage. In this study, we present a new set of partition coefficient data derived from 102 synthetic clinopyroxenes in equilibrium and in disequilibrium with a K-basaltic composition from the 103 Campi Flegrei Volcanic District (CFVD) as a function of temperature and time. The investigated

104 pressure of 0.8 GPa represents the depth of the crustal-mantle boundary in the CFVD area (~25 km 105 of depth; Fedi et al., 2018). Complementing our experimental data, we also measured the trace 106 element concentrations in some natural clinopyroxene phenocrysts belonging to scoria clasts from 107 Procida island in the CFVD. The main aim of this work, indeed, is to address: i) the scarcity of 108 quantitative experimental data on the role of the crystallization kinetics on clinopyroxene-melt trace 109 elements partitioning behaviour at high-pressure conditions relevant for deep magmatic reservoirs; 110 *ii*) the absence of clinopyroxene-melt partition coefficient experimental data for primitive 111 compositions in the CFVD area. Based on the evidence relating to the stages of deep crystallization of the Campi Flegrei Volcanic District parental magmas (Di Stefano et al., 1999; Piochi et al., 2005; 112 113 Chiarabba et al., 2008; Mormone et al., 2011; Moretti et al., 2013; Fedi et al., 2018), we explored the 114 mineral assemblage involved in the magmatic differentiation at the deepest ponding level, applying 115 the experimental results of this study.

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117 **2. Methods**

118 2.1. Samples description

119 Synthetic samples were obtained through crystallization experiments performed by using the ¹/₂ inch 120 end loaded piston cylinder apparatus at the HP-HT Laboratory of the Earth Sciences Department 121 (Sapienza University, Rome, Italy) at 0.8 GPa, 1080-1250 °C, $1 \le H_2O \le 4$ wt.%, with durations of 122 0.25, 3, 6 and 9 hours. For these experiments we used two different starting materials: the APR16 123 powder and the APR16GL glass (Table 1), both obtained from a natural primitive K-basalt (APR16 124 sample) found as scoria clast in the deposits of the Solchiaro hydromagmatic eruption (~22 ka ago; 125 (Morabito et al., 2014) located in Procida Island (Campi Flegrei, Italy). This K-basalt is characterized 126 by 12 vol.% of forsteritic olivine and diopsidic clinopyroxene phenocrysts dispersed in a groundmass 127 made of olivine, clinopyroxene, plagioclase, Ti-magnetite, alkali feldspar and glass (D'Antonio et al., 1999; De Astis et al., 2004). The APR16 powder, previously used as starting material in the 128 129 experimental runs of Bonechi (2020) (Table 2), was obtained by grinding the natural APR16 rocky

130 sample in an agate mortar under acetone. The APR16GL, instead, used as starting material in the runs 131 of Bonechi et al. (2020b) (Table 2), was prepared by melting for 15 min the APR16 natural powder in a 1 atm gas-mixing (CO-CO₂) furnace installed at the Bayerisches Geoinstitut (Bayreuth, 132 133 Germany) at 1400 °C and oxygen fugacity corresponding to the NNO (nickel-nickel oxide) buffer. 134 Then, the obtained glassy starting material was grinded in an agate mortar under acetone. 135 Compositions of synthetic clinopyroxenes were compared with natural clinopyroxene phenocrysts 136 occurring in the SCE and SOG scoria clasts, belonging to the Solchiaro eruption. The SCE and SOG 137 samples (Table 1 and Supplementary Material 1) are porphyritic, scarcely vesiculated rocks 138 containing ~10-20 vol.% of medium-grained olivine and clinopyroxene phenocrysts 139 (olivine/clinopyroxene ratio ~1). A detailed description of these samples is reported in Bonechi et al. 140 (2020b).

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142 **2.2.** Experimental techniques

The detailed experimental procedure is described by Perinelli et al. (2019) and by Bonechi et al. (2020a, 2020b). Here we report the main aspects. Assemblies made of CaF₂ outer sleeve, graphite furnace and crushable magnesia spacers were used. The friction correction factor associated with our assembly was determined to be ~6% through the calibration based on the reaction fayalite + quartz = ferrosilite at the fixed temperature of 1000 °C; at this temperature, the reaction takes place at approximately 1.4 GPa (Bohlen et al., 1980).

The experimental charges were prepared loading Au₇₅Pd₂₅ capsules with ~40 mg of sample powder. For hydrous experiments we added water in known amounts by using a 1.0 μ L microsyringe (Table 2). Loss of H₂O by vaporization during welding was prevented by freezing the crucible with nitrogen. The welded capsule was weighed, stored in a furnace at 110 °C, and then weighed again to confirm the absence of water leaks. Moreover, to minimise possible loss of H₂ during the experiments the charges were surrounded by pyrophyllite powder (Freda et al., 2001). Temperature was controlled by a factory calibrated W₃Re₉₇-W₂₅Re₇₅ (type D) thermocouple and maintained with an uncertainty of ±

156 5 °C. For the experimental run-up we used the "hot piston-out" procedure (Johannes et al., 1971; 157 Presnall et al., 1978) modified consisting in the first cold pressurization of the assembly to a value ~ 158 10% over the target pressure and then the sample was heated to the final run temperature. This 159 technique allows to closely approach the run pressure through the stress relaxation within the 160 assembly upon heating, requiring only minor upward pressure adjustments during the experiment in 161 most cases (Watson et al., 2002). The heating rate was imposed at 150 °C/min. The sample was 162 maintained at the pressure and temperature for a given duration (nominal duration), and then 163 quenched by turning off the power. The temperature drop below 500 °C was of <5 s (average cooling rate of 150 °C/s). All the runs were self-buffered; an estimation of experimental oxygen fugacity (fO₂) 164 165 was made through the equation of Kress and Carmichael (1991). To use this equation, where possible, we first assessed the FeO content of experimental residual glass, and then its Fe²⁺/Fe_{tot} mole ratio; to 166 167 do this we used the value of iron and magnesium partitioning between olivine and liquid (K_D Fe-Mg 168 ol/liq) calculated according to the procedure proposed by Toplis (2005). Therefore, the application of 169 Kress and Carmichael (1991) equation residual glass of each experiment, yielded fO_2 values between 170 of -1.9 and +0.9 log-bar units with respect the nickel-nickel oxide buffer. As a whole the estimated 171 range of fO₂ value is in agreement with those estimated for similar furnace assemblages (Conte et al., 172 2009; Weaver et al., 2013; Perinelli et al., 2019; Bonechi et al., 2020a). Finally, we determined the 173 effective undercooling ($\Delta T_{\rm eff}$) that is the difference between the equilibrium clinopyroxene liquidus 174 temperature (TCpx_L) and the experimental temperature (Bonechi et al., 2020b, 2020a; Bonechi, 175 2020). For calculations, TCpxL value of APR16-NWA experiments was taken from Procida K-basalt 176 experimental data (i.e., 1270 °C; see Table 2 in Perinelli et al., 2019) whereas TCpx_L values of 177 APR16GL experiments were calculated by the MELTS software (Ghiorso and Sack, 1995; Asimow 178 and Ghiorso, 1998) using 0, 2 and 4 wt% as initial water contents (i.e., H_2O_i in Table 2).

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180 2.3. Analytical techniques

181 2.3.1. Textural investigations and major element determinations

182 Back-scattered electron (BSE) images used to investigate textural aspects of both natural and 183 experimental samples were collected by Scanning Electron Microscopy (SEM) using a FEI-quanta 400 equipped for microanalysis with an EDAX Genesis system at the Earth Sciences Department, 184 185 Sapienza University of Rome. Major elements were analysed at the CNR-Istituto di Geologia 186 Ambientale e Geoingegneria di Roma, with a Cameca SX50 electron microprobe equipped with five 187 wavelength dispersive spectrometers (WDS). Quantitative analyses were performed using 15 kV 188 accelerating voltage and 15 nA beam current. As standards we employed metals for Mn and Cr, 189 Jadeite for Na, Wollastonite for Si and Ca, Orthoclase for K, Corundum for Al, Magnetite for Fe, 190 Rutile for Ti, Periclase for Mg, Apatite for P. Counting times for all elements were 20 s on peak and 191 10 s on both backgrounds. Light elements (Na, K) were counted first to prevent loss by volatilization. 192 The PAP correction method was used. Minerals were analysed using a beam diameter of 1 µm 193 whereas to minimize alkali loss during glass analysis, the beam was defocused to 15 µm. In order to 194 evaluate the accuracy of the analyses, repeated analyses of three international secondary standards 195 (Kakanui augite, Iceladic Bir-1 and rhyolite RLS132 glasses from USGS) were made prior to any 196 series of measurements. The mean precision from the standard value was about 1% for SiO₂, 2% for 197 Al₂O₃, 5% for K₂O, CaO and FeO, and 8-10% for other elements. The analytical precision (2 sigma 198 error) is $\leq 1\%$ for elements in the concentration range >10 wt% oxide, 5% for elements in the range 199 2–10 wt% oxide and better than 10% for elements in the range 0.5–2 wt% oxide. The water contents 200 of experimental glasses were estimated according to the by-difference method (Devine et al., 1995; 201 Humphreys et al., 2006). We are aware that this method is not fully reliable given the estimation is 202 affected by i) element concentrations not measured by EMPA and ii) surface charge effects (Hughes 203 et al., 2019) leading to an overestimation of H₂O concentration. However, the obtained H₂O values show a good linear correlation (correlation coefficient $R^2 = 0.97$) with are close to those determined 204 205 by micro-Raman spectroscopy measurements on residual melts of equilibrium experiments 206 performed on the same composition, at the same P, T and initial hydrous conditions (see Perinelli et 207 al., 2019, for details).

210 LA-ICP-MS analyses were performed on both natural (SCE and SOG samples) and experimental 211 samples. LA-ICP-MS analyses of SCE and SOG natural samples were performed on a quadrupole-212 based ICP-MS iCAP-Q instrument (Thermo Fisher Scientific, Bremen) coupled to a modified 213 NewWave UP 213 laser microprobe (NewWave, USA) installed at Charles University, Prague. 214 Analyses of natural clinopyroxene crystals were performed using a circular laser beam of 100 µm 215 diameter, a frequency of 10 Hz and a laser density on the sample surface of 3.5 J/cm². External calibration of the laser ablation analyses was done using Standard Reference Materials NIST 216 217 (National Institute of Standards and Technology, USA) 612 with internal standardisation using Si, based on electron microprobe measurements. The concentration values for NIST of all measured 218 219 elements were taken from (Pearce et al., 1997). The external reproducibility of this method was 220 checked on repeated analyses of USGS BCR-2G glass as reference material (e.g., Strnad et al., 2005 221 and reference therein; Table S1 in Supplementary Material 2). The isotopes used were selected with 222 respect to their most abundant species, free from isobaric overlap and minimum interferences. 223 Formation of oxides (MO⁺/M⁺) was monitored using U in NIST 612 directly from ablation and the measured ratios $(^{254}UO^{+/238}U^{+})$ was below 10^{-2} . Further details about the analytical protocol and 224 225 correction strategy have been described by Strnad et al. (2005) and Skála et al. (2009). Detection 226 limits for all elements were calculated as 3σ level of the gas blank (Table S1 in Supplementary 227 Material 2). LA-ICP-MS analyses of the experimental samples were performed at the Department of 228 Physics and Geology, University of Perugia (Italy). The instrumentation consisted of a 229 Teledyne/Photon Machine G2 LA device equipped with a Two-Volume ANU HelEx 2 cell coupled 230 with a Thermo Fisher Scientific quadrupole-based iCAP-Q ICP-MS. Analyses of clinopyroxene 231 crystals and glasses were performed using a circular laser beam of 8-20 µm diameter, a frequency of 10-15 Hz and a laser density on the sample surface of 3.5 J/cm². Oxide formations was measured on 232 the NIST SRM 612 monitoring and maintaining the ratio ThO/Th below 0.005. The NIST SRM-610 233

234 standard reference material was used as calibrant and Ca, previously analysed by EPMA, as internal 235 standard. Under these operating conditions precision and accuracy are better than 10% for most of the elements (Petrelli et al., 2007, 2008, 2016). In detail, at beam sizes equal or larger than 15 µm, 236 237 precision (measured as one sigma) and accuracy (expressed as relative deviation from the reference value) are of the order of 10% (Petrelli et al., 2016). At a smaller beam size of 8 µm, precision 238 decreases to 15% for concentration above 1.7 μ g g⁻¹ (Petrelli et al., 2016). Accuracy is better or equal 239 to 10% and 13% at beam sizes of 15 and 10 µm, respectively (Petrelli et al., 2016). When the spatial 240 241 resolution is increased to 8 µm, accuracy remains better than 15% (Petrelli et al., 2016).

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243 **3. Results**

244 3.1 Phase relations

245 Experimental samples investigated in this study are described in detail in previous works dealing with 246 phase equilibria and crystallization kinetics in the APR16 K-basalt (Perinelli et al., 2019; Bonechi et al., 2020b, 2020a; Bonechi, 2020). Here we briefly report a description of the paragenesis of these 247 248 samples given the importance of the coexistence of several phases for the fractionation of some trace 249 elements, like the presence of plagioclase for the fractionation of Eu. Overall, the experimental 250 products consist of glass and microlites of clinopyroxene (Cpx) \pm olivine (Ol) \pm plagioclase (Plg) \pm 251 oxide (Ox). In particular, the APR16-1C and APR16-35 runs (1225 °C), performed using the natural 252 starting material APR16, are characterized by the presence of glass (GI) and Cpx plus restitic Ol and 253 Ox. In the runs performed at 1250 °C (APR16-C1Ac and APR16-C1Ad), the dominant phase is Cpx 254 while olivine (Ol) crystallizes only in the APR16-C1Ac run. The mineral phases crystallized in the 255 runs at 1220 °C (APR16-C2Aa, APR16-C2Ab, APR16-C2Ac, APR16-C2Ad) are Cpx + Ol + Plg + 256 Ox. Clinopyroxene and Ol are ubiquitous in these runs, whereas Plg is present only in the runs at 3 257 and 6 hours (i.e., APR16-C2Ab, APR16-C2Ac) and Ox is absent in the run APR16-C2Ac. The runs 258 at 1170 °C (APR16-C2Bc and APR16-C2Bd) show a higher degree of crystallization and the mineral assemblage is $Cpx + Ol + Plg \pm Ox$. Finally, the hydrous run at 1080 °C (APR16-C3Ac; 6.5 wt.% 259

 H_2O H₂O) shows a degree of crystallization similar to that of the runs at 1170 °C and the phases crystallized are Cpx and Ol.

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263 **3.2.** Texture and chemical composition of experimental samples

264 *Clinopyroxene*. Chemical compositions of Cpx are given in Table S2 in Supplementary Material 2. 265 Overall synthetic Cpx crystals are characterized by homogeneous composition and euhedral habits 266 (Fig. 1a-c). In particular, in the runs performed using the glassy starting material (i.e., APR16GL) 267 habit and sizes vary with time: subhedral with sizes of ~5-10 µm in the shortest run (e.g., APR16-268 C2Aa; 0.25 h; Fig.1a) and euhedral with sizes up to ~20 μ m in the longest run (\geq 3 h; Fig. 1b). The 269 runs APR16-1C and APR16-35, performed using the natural starting material (i.e., APR16 natural 270 powder), instead, are characterized by euhedral habit and sizes up to ~50 µm (Fig. 1c) (Bonechi, 271 2020). These Cpx crystals are augitic (Wo₄₂En₄₁Fs₁₈) and diopsidic (Wo₄₇En₄₃Fs₁₀) in composition 272 according to the classification scheme of Morimoto et al. (1998) (Fig. 2a). Overall, the Mg# of Cpx 273 (calculated as the molar MgO/(MgO+FeOtot) ratio) varies between 0.81 and 0.92, the TiO₂ contents 274 are between 0.58 and 1.18 wt.% while those of Al₂O₃ are between 5.00 and 9.49 wt.%. The Fe-Mg 275 distribution coefficients between clinopyroxene and melt $(K_{\rm D}({\rm Fe-Mg})^{Cpx-liq})$ = $Fe_{Cpx} \times Mg_{Iig}/Mg_{Cpx} \times Fe_{Iig}$, calculated assuming all Fe as Fe^{2+} in both phases), for the longest runs ≥ 3 276 277 hours lie in the equilibrium range of 0.28±0.08 (Putirka, 2008) whereas the shortest experiment 278 (APR16-C2Aa) shows significant deviation from the equilibrium (Table S2 in Supplementary 279 Material 2).

Glass. Chemical compositions of experimental glasses are given in Table S3 in Supplementary Material 2. Glass composition of anhydrous and hydrous experiments varies from K-basalt to Ktrachybasalt up to shoshonite compositions with decreasing temperature (Fig. 2b). In particular, in the anhydrous runs, before reaching the temperature at which plagioclase significantly crystallizes, the glass composition is controlled by clinopyroxene crystallization thus leading to a very limited SiO₂ increase in the residual glasses (Perinelli et al., 2019). In the hydrous experiments, the earlier crystallization of olivine \pm spinel causes a slightly more marked increase of SiO₂ abundance in the residual glasses (from ~52 to ~53 wt.%; Table S3 in Supplementary Material 2).

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289 3.3. Trace and REE content

Average compositions of trace elements concentrations analysed by LA-ICP-MS technique insynthetic clinopyroxene and coexisting glass are reported in Table 3.

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293 3.3.1. Clinopyroxene

Figure 3 shows the trace elements concentrations in synthetic and natural clinopyroxene normalized 294 295 to chondrite C1 after Anders and Grevesse (1989) for LILEs (Large-Ion Lithophile Elements), REEs 296 + Y (Rare Earth Elements plus Yttrium), HFSEs (High Field Strength Elements) and TEs (Transition 297 Elements). In this diagram some LIL elements (e.g., Rb and Ba) of synthetic Cpx crystals display 298 different relative abundances depending on experimental conditions unlike the REEs that instead, 299 show quite uniform high contents. In particular, the chondrite-normalised REE patterns have a general 300 L/HREE (Light/Heavy rare earth element) enrichment (i.e., $La_N/Yb_N = 0.85-1.20$, where the subscript 301 N denotes chondrite-normalized values) with a maximum at Sm_N (~20-37). The REE profiles from 302 the crystallized cpx are comparable to each other with the exception of the clinopyroxene in the 303 APR16-C2Bd charge that shows an evident Eu negative anomaly (Eu/Eu* = 0.58; Eu/Eu* = 304 $Eu_N/(Sm_N \times Gd_N)^{1/2}$), likely due to the crystallization of plagioclase in this run. Indeed, the lack of 305 plagioclase in the hydrous experiment APR16-C3Ac, justifies the absence of troughs at Eu (Eu/Eu* 306 = 0.95), despite both crystallization degree and amount of Cpx crystallized in this experiment are 307 similar to that of APR16-C2Bd run (40 wt.% and 43 wt.%, and 33 wt.% and 34 wt.%, respectively; 308 Table 2). Finally, tetravalent HFSEs (i.e., Zr and Hf) have a higher concentration than pentavalent 309 cations (i.e., Ta and Nb), while as regards TEs, it is possible to see evident troughs at Co and Ni.

310 3.3.2 Glass

311 Figure 4 shows the trace elements concentration in the experimental glasses normalized to chondrite 312 C1 after Anders and Grevesse (1989) for LILEs, REEs + Y, HFSEs and TEs. Generally, all the 313 experimental glasses match the trend of the APR16 sample (Mazzeo et al., 2014), with evident 314 troughs at Pb, Co and Ni. C1-normalized trace element diagrams are L-MREE (Light/Middle rare earth element) enriched ($La_N = 58-90$, $La_N/Sm_N = 2.09-2.66$, $La_N/Yb_N = 5.30-8.58$) and nearly flat in 315 316 the HREE region (H_{0N}/Yb_{N} 1.01-1.48). All the considered glasses have absent or slightly positive Eu 317 anomaly (Eu/Eu* 0.95-1.10; inset of Fig. 4). Eu/Eu* ratio decreases with increasing degree of 318 chemical evolution with the lowest values observed in APR16-C2Bd run (see TAS diagram in Fig. 319 2b). Moreover, pentavalent HFSEs (i.e., Ta and Nb) are more abundant than tetravalent cations (i.e., 320 Zr and Hf), unlike what was observed for Cpx crystals.

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322 3.4. Partition coefficients

323 The clinopyroxene-melt trace element *D*-values for the reported experiments are given in Table 4. 324 Figure 5 shows the partition coefficients between clinopyroxene and melt for LILEs, REEs + Y, 325 HFSEs and TEs. LILEs are incompatible (e.g., $D_{Sr} \le 0.15$, $D_{Ba} \le 0.05$). LREEs (e.g., $D_{La} \le 0.20$) are 326 always more incompatible than HREEs, which in some cases result to be compatible with cpx (e.g., D_{Tb} and $D_{\text{Dy}} = 1.30-1.40$ for APR16-C2Bd and APR16-C2Ad experiments). There is no clear 327 328 relationship between D_i variation and dwell time for LREEs. In contrast, for MREE and HREE 329 longest duration experiments (e.g., APR16-35 and APR16-C1Ad) are characterized by lower partition 330 coefficients. In some cases, it was possible to note evident differences of D_i values as function of 331 experimental time, as shown for instance by Dy behaviour in the runs at 1250 °C, where it results a compatible element in the 6 h-experiment (APR16-C1Ac; $D_{Dy} = \sim 1$) while it is moderately 332 333 incompatible in the run at 9 hours (APR16-C1Ad; $D_{Dy} = \sim 0.6$). Furthermore, we observed that, except for the APR16-C3Ac run (the most hydrated run), the Eu behaves to a more incompatible extent than 334 335 the adjacent REEs (i.e., Sm and Gd, Fig. 5). Given the low plagioclase abundance, this behaviour reflects only the Eu^{3+}/Eu^{2+} ratio variation caused by changing the redox conditions. As regards 336

HFSEs, tetravalent and pentavalent-charged cations are always incompatible (D_{Nb} and $D_{Ta} \le 0.1$; D_{Zr} and D_{Hf} between 0.2 and 0.8), except for D_{Hf} measured in the cpx of the APR16-C2Bd run ($D_{Hf} \sim 1.2$). Finally, trace element partition coefficients of TEs (i.e., D_{Co} , D_{Cr} , D_{Ni} , D_{Sc} and D_{V}) range from slightly incompatible to highly compatible. In particular, D_{Co} and D_{V} vary between 0.7 and 3, whereas D_{Sc} , D_{Ni} and D_{Cr} are between 2 and 63.

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343 **4. Discussion**

344 4.1. Lattice strain model (LSM)

Assuming equilibrium, the cpx/melt partition coefficient is quantitatively described by the nearparabolic relationship (Brice, 1975; Blundy and Wood, 1994; Wood and Blundy, 1997; Sun and Liang, 2012):

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$$D_j^{cpx-melt} = D_0 exp \left[\frac{-4\pi E N_A}{RT} \left(\frac{r_0}{2} \left(r_0 - r_j \right)^2 - \frac{1}{3} \left(r_0 - r_j \right)^3 \right) \right]$$
 (1),

350

where D_0 is the clinopyroxene-melt partition coefficient for the strain-free substitution; r_0 is the optimum radius for the lattice site; r_j is the ionic radius of the element of interest in the appropriate co-ordination taken from Shannon (1976) (Table S4 in Supplementary Material 2); *E* is the apparent Young's modulus for the lattice site (GPa); *R* is the gas constant (8.3144 J mol⁻¹ K⁻¹); N_A is Avogadro constant and *T* is temperature in K. The apparent Young's modulus *E* controls the tightness of the parabola, and D_0 determines the apex of the parabola with corresponding ideal lattice site r_0 .

357 Cpx crystals are characterized by the general formula XYT_2O_6 . The T site could contain tetrahedrally 358 coordinated Si⁴⁺ and ^{VI}Al³⁺, whereas X and Y refer to two octahedral sites, a larger M2 site and a 359 smaller M1 site (Sun, 2018). The M1 site is occupied by small cations in VI-fold coordination, 360 including divalent Ni, Mg, Co and V, trivalent Al, Cr and Sc, tetravalent Ti, Hf and Zr, and 361 pentavalent cations (e.g., Nb⁵⁺ and Ta⁵⁺) (Blundy and Wood, 2003; Sun and Liang, 2012; Sun, 2018; 362 Baziotis et al., 2019). The M2 site, instead, could host larger cations in VIII-fold coordination like 363 monovalent Na, K and Rb, divalent Fe, Ca, Eu, Sr, Pb and Ba, trivalent REE (La to Lu) and Y, and 364 tetravalent U and Th (Blundy and Wood, 2003). To apply the lattice strain model (LSM) to both M1 and M2 sites and fit the experimentally determined trace element partition coefficients simultaneously 365 366 we used the DOUBLE FIT program (Dalou et al., 2018), a lattice strain model fit by a differentialevolution-constrained algorithm (Storn and Price, 1997) adapted to be error weighted. When errors 367 are large, we individually fitted the M1 and M2 sites using the SIMPLE FIT program (Dalou et al., 368 369 2018). In presence of a low number of measured trace elements, the program failed to produce good 370 fits. Therefore, following (Dalou et al., 2018), we reduced the parameter space from the default range $(D_0^{M1-M2}min and/or D_0^{M1-M2}max; E^{M1-M2}min and/or E^{M1-M2}max; r_0^{M1-M2}min and/or r_0^{M1-M2}max)$ to 371 372 overcome this limitation. Best-fit parabolas determined for divalent, trivalent, and tetravalent cations are plotted in Figure 6 and Figure S1 in Supplementary Material 3. Fit parameters E, D_0 and r_0 for 373 trivalent cations in M2 site are reported in Table 5, together with E and r calculated following 374 375 equations from Wood and Blundy (1997):

376

377
$$r_{0(M2)}^{3+} = 0.974 + 0.067 X_{Ca}^{M2} - 0.051 X_{Al}^{M1}(\text{\AA})$$
 (2),

378

$$379 \quad E_{M2}^{3+} = 318.6 + 6.9P - 0.036T (GPa) \tag{3},$$

380

where X_{Ca}^{M2} and X_{Al}^{M1} refer to the atomic fractions of Ca and Al on the clinopyroxene M2 and M1 sites, *P* is in GPa and *T* in K. Fit parameters *E*, D_0 and r_0 for divalent cations in M1 and M2 site, and for trivalent and tetravalent cations in M1 site are reported in Table 5 and compared with *E* and *r* calculated using the model of Wood and Blundy (1997) for M2²⁺ and that of Hill et al. (2011) for M1⁴⁺. Following a method reported in the literature (Aigner-Torres et al., 2007; Padilla and Gualda, 2016), D_{Eu}^{3+} and D_{Eu}^{2+} were calculated using the LSM, based on the measured partition coefficients of other REE³⁺, ideally close in ionic radius to Eu³⁺ (i.e., Gd and Sm), and Sr, respectively. These

data were included in the data set of divalent (D_{Eu}^{2+}) and trivalent (D_{Eu}^{3+}) cations used for the fitting. 388 For the fitting of trivalent cations in M1 site, Al in the M1 site was calculated as the difference 389 between total Al and the Al^{IV} completing the occupancy of the tetrahedral site. As regards divalent 390 cations, to understand their distribution between M1 and M2 sites, we performed a preliminary fitting 391 with DOUBLE FIT program, observing that *i*) Mn, Ca, Eu^{2+} , Sr, Pb and Ba partition only in M2 site; 392 ii) Ni partitions only in M1 site; *iii*) Mg, Co and Fe, instead, partition in both M1 and M2 sites. At 393 this point we fitted divalent cations entering only in M2 site (i.e., Mn, Ca, Eu^{2+} , Sr and $\pm Pb$). The 394 395 obtained lattice strain parameters were used to calculate $D_{(Fe,Mg,Co)M2}$, that is the *D* value in M2 site of 396 the elements entering both the M1 and M2 sites (i.e., Fe, Mg, Co). Then, using the concentration of 397 these elements in the melt $(X_{(Fe,Mg,Co)melt})$ we calculated the concentrations of the elements in M2 $(X_{(Fe,Mg,Co)M2})$ and by difference in M1 $(X_{(Fe,Mg,Co)M1})$. Finally, we obtained $D_{(Fe,Mg,Co)M1}$ for these 398 399 elements (Table 5) dividing $X_{(Fe,Mg,Co)M1}$ with $X_{(Fe,Mg,Co)melt}$. In the fitting of the divalent cations in M1 400 site we constrained r_0 between 0.6 and 0.65 Å, based on the work of Purton et al. (1996).

401

402 4.1.1. Onuma diagrams for divalent, trivalent and tetravalent cations

The partitioning behaviour for cations (D_i) with the same valence is conveniently shown in Onuma diagrams as a function of the ionic radius (Onuma et al., 1968; Leeman and Phelps, 1981; Blundy and Wood, 1994), indicating which elements are compatibles/incompatibles in a given site or which ones are fractionated between two distinct sites.

407 *Divalent cations.* Values of D_0 , r_0 and E for divalent cations in M1 and M2 site as fitted to the 408 experimental data are given in Table 5. For divalent cations in M1 site we obtained $D_0^{2+} = 6.07 \pm$ 409 2.02, $E^{2+} = 499 \pm 45$ GPa, and $r_0^{2+} = 0.62 \pm 0.01$ Å (Fig. 6 and Fig. S1 in Supplementary Material 3). 410 V^{2+} was excluded from the fitting since it falls outside the parabola, probably due to its presence as 411 V^{5+} . The LSM parameters for the M2 site are $D_0^{2+} = 2.50 \pm 0.37$, $E^{2+} = 256 \pm 26$ GPa, and $r_0^{2+} = 1.08$ 412 ± 0.01 Å (Fig. 6 and Fig. S1 in Supplementary Material 3). The partitioning of divalent cations into 413 M2 provides a good illustration of the control on partitioning by site size energetics. The linearized

regressions for Mn, Ca, Eu²⁺, Sr and Pb produce good fits. Among these divalent cations, Ca²⁺, that 414 415 dominantly occupies the M2 site in clinopyroxene (Wood and Blundy, 2001), is the cation closer to the parabolas' apex and Sr is more compatible than Ba. However, the Ba position is far from the fit 416 417 curve of divalent cations. This anomalous behaviour is probably due to the large ionic radius that 418 produces a displacement of the nearest-neighbour ions in the charged region with an electrostatic 419 work of substitution ($\neq 0$) in spite that of Ba is a fully charge-balanced substitution in the M2 site. A 420 minor amount of melt contamination during crystal analysis can cause a large offset from the lattice 421 strain curve.

Trivalent cations. Among trivalent cations TEs (i.e., Cr and Sc) are compatible, LREEs are all 422 423 incompatible, while HREEs are both compatible and incompatible. Among REEs Dy and Er show 424 the highest distribution coefficients, that tend to decrease with increasing the ionic radius (towards 425 La). Because the HREEs (Dy through Lu) are more compatible than LREEs, clinopyroxene is slightly enriched in HREEs. The LSM parameters for the M1 and M2 sites are $D_0^{3+} = 47 \pm 18$ and 0.85 \pm 426 $0.27, E^{3+} = 1456 \pm 113$ and 311 ± 10 GPa, and $r_0^{3+} = 0.66 \pm 0.01$ and 1.03 ± 0.01 Å, respectively. The 427 values of r_0^{3+} and E^{3+} in the M2 site match within the range of previous studies on basaltic 428 compositions ($r_0 = 1-1.05$ Å, E = 325-441 GPa; Hill et al., 2011; Dalou et al., 2012; Mollo et al., 429 2016; Baudouin et al., 2020). Similarly, D_0^{3+} and E for the M1 site display similar values compared 430 431 to other basaltic compositions ones (Hill et al., 2000; Dalou et al., 2012). Figure 6 and Figure S1 in 432 Supplementary Material 3 show that our apparent partition coefficients for REEs lie on parabola-like curves of Onuma diagrams (Onuma et al., 1968), similar to the trajectories found by Pappalardo et 433 434 al. (2008), Fedele et al. (2009) and Mollo et al. (2016) for differentiated magmas at Campi Flegrei. 435 From these diagrams, also, it is possible to note the different partitioning behaviour of Eu, which occurs as both 2+ and 3+ cations, depending on fO_2 . Particularly, if all Eu is present as Eu³⁺ D_{Eu} lies 436 along the parabola defined by other REE³⁺, whereas if Eu is present as Eu²⁺, D_{Eu} lies along the 2+ 437 parabola defined by Mn, Ca, Sr, \pm Pb (Aigner-Torres et al., 2007). In all the runs of this study D_{Eu} lies 438 along or close to the parabola defined by other REE^{3+} . In particular, D_{Eu} lies along the parabola in the 439

440 APR16-C3Ac run (Fig. 6c), slightly below the parabola in APR16-1C, APR16-35 and APR16-C1Ac 441 runs (Fig. 6a, b and Fig. S1 in Supplementary Material 3), and far below the parabola in the APR16-442 C2Bd runs (Fig. 6d). The Eu-anomaly is due to the presence of some Eu²⁺ in the melt, which is much 443 less compatible in the M2 site than Eu³⁺ because of its larger size (Shearer et al., 2006). Moreover, in 444 the APR16-C2Bd run, a clear difference is observed difference between D_{Eu3+} calculated and D_{Eu} 445 measured (Fig. 6d). The latter indeed is much lower (~0.6) than D_{Eu3+} (~1) due to the crystallization 446 of plagioclase in this run.

447 The height of the parabola varies as a function of the crystal composition (Blundy and Wood, 1994, 448 2003) and the physical conditions of the system (Wood and Blundy, 2001; Sun and Liang, 2012). In 449 the experimental runs the peak position can be alternatively higher or lower than 1 as a function of Al^{IV} content and temperature, in agreement with the observation that these parameters have opposing 450 451 effects on trace element partitioning (Sun and Liang, 2012). For instance, we noted that the increase of Al^{IV} content, at a given temperature and under similar water content conditions, causes an increase 452 of D_0 , and a decrease of r_0 and E (Fig. 7). As a consequence, the parabolic patterns of D_{REE} move 453 454 upwards and shift toward left, resulting in a more open parabola (Fig. 7a). This suggests that the increase in Al^{IV} content has a greater effect on the D_{HREE} than on D_{LREE} . In fact, HREEs are more 455 456 easily accommodated in clinopyroxene than LREEs (McKay et al., 1994; Lofgren et al., 2006; Mollo 457 et al., 2013), since their smaller ionic radius can better compensate for the enlargement of the 458 tetrahedral site induced by the replacement of Si with Al (Hill et al., 2000). Consequently, partition 459 coefficients of HREEs, which are on the left side of the parabola, increase with the increase in Al^{IV} content, while partition coefficients of light and middle REEs (LREE and MREE), which are on the 460 461 right side of the parabola, decrease slightly as the parabola shifts to the left, but increase as the 462 parabola becomes more open, as previously observed by Sun and Liang (2012). Furthermore, even if it was not possible to discriminate the only effect of the temperature in our runs, we believe that for 463 464 a constant composition an upward movement of the parabola (e.g., APR16-C3Ac and APR16-C2Ac, 465 APR16-C1Ad and APR16-C2Bd; Fig. 7b) may be due to a decrease in temperature, following the

466 observations of Sun and Liang (2012). Moreover, together with tetrahedrally coordinated Al and 467 temperature, also the undercooling ($\Delta T = T_{\text{liquidus}} - T_{\text{experiment}}$) influence the partitioning of trace 468 element, as shown in Figure 7b. In particular, the increase of the undercooling moves the parabola 469 upwards as observed for Al^{IV} content.

470 *Tetravalent cations.* The partition coefficients for Ti, Hf and Zr have been used to obtain E, D_0 , and r_0 for the tetravalent cations. Best fits to the data yield $D_0^{4+} = 3.35 \pm 1.54$, $E^{4+} = 2600 \pm 536$ GPa, and 471 $r_0^{4+} = 0.657 \pm 0.004$ Å (Fig. 6 and Table 5). The inferred values for *E* (~1950-3100 GPa) are consistent 472 with the large values expected for small, highly charged cation sites (~800-2400 GPa; (Lundstrom et 473 al., 1998; Hill et al., 2011; Baudouin et al., 2020). Th⁴⁺ and U⁴⁺ fall off the Ti-Hf-Zr regression 474 suggesting that their ionic radii (1.05 and 1.00 Å, respectively) are too large for incorporation in the 475 M1 site (Lundstrom et al., 1998). The r_0^{4+} of the M1 site is very similar to that calculated in previous 476 studies $(r_0^{4+} \sim 0.65 - 0.67 \text{ Å})$ (Hill et al., 2000, 2011; Mollo et al., 2016, 2018; Baudouin et al., 2020). 477 478

479 **4.2.** Influence of kinetics processes on the trace element partitioning

As general rule, partition coefficients tend to decrease with the increase of temperature (Wood and 480 481 Blundy, 1997) and the effect of pressure at pressures < 2 GPa may be negligible for trace element 482 partitioning between clinopyroxene and mafic melts (Sun and Liang, 2012; Mollo et al., 2016). As 483 regards the influence of the chemical composition, it has been observed that in Cpx the tetrahedrally-484 coordinated aluminium content has a great influence on trace element partitioning (Lindstrom, 1976; Ray et al., 1983; Hart and Dunn, 1993; Forsythe et al., 1994; Lundstrom et al., 1994; Skulski et al., 485 486 1994; Lundstrom et al., 1998; Blundy et al., 1998; Hill et al., 2000; Wood and Trigila, 2001; Francis 487 and Minarik, 2008; Sun and Liang, 2012; Yao et al., 2012; Mollo et al., 2013; Sun and Liang, 2013; 488 Scarlato et al., 2014; Mollo et al., 2016). Nevertheless, the magnitude of the T, P and composition 489 effects on the partitioning of trace elements between minerals and melts is not correctly assessed 490 without considering the kinetic effect.

491 In the literature, the few works investigating the influence of kinetics processes on D_i values are 492 generally focused on the effect of the cooling rates (Albarede and Bottinga, 1972; Watson, 1996; 493 Lofgren et al., 2006; Mollo et al., 2013) and time (Arzilli et al., 2018), showing an increase of REE 494 partition coefficients with increasing cooling rates or with decreasing crystallization times and, 495 consequently, with increasing growth rates. Thus, since growth rates are fast at the beginning of the 496 crystallization process and then decrease with increasing time (Vetere et al., 2013; Arzilli and Carroll, 497 2013; Iezzi et al., 2014; Vetere et al., 2015; Arzilli et al., 2018; Bonechi et al., 2020b, 2020a; Bonechi, 498 2020), we would expect a decrease of D_i values with increasing time until equilibrium conditions are 499 reached. In this study, at static conditions (i.e., no cooling rate), D_{REE} values seem to increase with 500 increasing time (Fig. 8), with D_{REE} values calculated in long-duration experiments higher than those 501 calculated in short duration ones, showing thus a behaviour apparently opposite to what would have 502 been expected. This is noticeable considering the group of runs APR16-1C/APR16-C1Ac/APR16-503 C2Ad (respectively 3, 6 and 9 hours long), characterized by similar T and ΔT conditions and similar Al^{IV} and H₂O contents (Table 2 and Table S2 in Supplementary Material 2). In particular, we observed 504 505 an increase in D_{REE} values from 3 (APR16-1C) to 9 hours (APR16-C2Ad) experiments, more evident 506 in the right side of the three parabolae (i.e., where the LREEs fall). Along this side, the parabolae 507 tend to widen to the right and upwards as time increases, as confirmed by the increase of r_0 (from 1.0297 to 1.0301 Å; Table 5) and the decrease of E (from 325 to 304 GPa; Table 5) with increasing 508 509 time. Conversely, there are no appreciable differences for the apical zone (where the HREEs fall) in 510 the 3 and 6 hours experiments. This variation in the shape of the parabolae makes the effect of time 511 on the D values stand out more clearly for the LREEs that fall on the flanks rather than for the HREEs 512 that fall in the apical area. Indeed, D_{HREE} values of the runs at 3 and 6 hours are very similar despite their different duration, and this similarity can be explained by considering their different growth 513 514 rates. Notably, the APR16-1C run (3 hours) is characterized by a growth rate $(1.6 \cdot 10^{-7} \text{ cm s}^{-1})$ higher than that of the other runs with the same duration (e.g., $3 \cdot 10^{-8}$ cm s⁻¹), as a consequence of the use of 515 516 natural powder containing pre-existing nuclei as starting material, as described in the work of Bonechi 517 (2020). Therefore, the high growth rate of this run yields D_{REE} values higher than those expected for 518 runs at 3 hours, but similar to those measured in the run at 6 hours (APR16-C1Ac). This agrees with 519 observations made in previous studies (Watson, 1996; Lofgren et al., 2006), according to which, 520 during rapid crystal growth the less efficient rejection of incompatible elements leads to higher D-521 values. Moreover, the greater similarity in D_{HREE} than in D_{LREE} in APR16-1C and APR16-C1Ac runs 522 is due to their different diffusivities, that decrease as the REE size decreases (Zhang et al., 2010), 523 since slow-moving elements (HREE) result more enriched compared with fast moving elements 524 (LREE), as observed for instance during disequilibrium growth (Arzilli et al., 2018). The increase in 525 D_{REE} values with time observed in this work, which would seem to contrast with what was observed 526 in the literature, actually is not due to a change in time but rather to the different non-bridging oxygens 527 per tetrahedrally co-ordinated cation ratio (NBO/T; Mysen et al., 1982, 1985) of the melts in the 528 considered experiments, that seems to counteract the effect of increasing time which would tend to 529 decrease the partition coefficient values. As stated in the literature, melt composition affects the melt 530 structure, and thus trace element partition coefficients, that generally decrease with decreasing melt 531 polymerization (i.e., increasing the NBO/T ratio) (Mysen and Virgo, 1980; Gaetani, 2004; Huang et 532 al., 2006; Mollo et al., 2016; Michely et al., 2017), while in some cases this influence seems to be 533 negligible in comparison to the effect of mineral chemistry (Beard et al., 2019; Baudouin et al., 2020). 534 Indeed, as observed by Gaetani (2004), the influence of melt structure is significantly evident when 535 NBO/T < 0.49. In this study, despite the NBO/T ratio varies between 0.6 and 1 (Table 2), we observed 536 the influence of this parameter on REE partition coefficients. As shown in Figure 9, there is a negative 537 correlation between D_{REE} and NBO/T ratio, described by the following equations for a representative 538 LREE (4), MREE (5) and HREE (6) element:

539

540
$$D_{Nd} = -1.54 * \frac{NBO}{T} + 1.72 \ (R^2 = 0.98)$$
 (4)

541

542
$$D_{Eu3+} = -2.45 * \frac{NBO}{T} + 2.73 \ (R^2 = 0.92)$$
 (5),

544
$$D_{Dy} = -4.54 * \frac{NBO}{T} + 4.58 (R^2 = 0.99)$$
 (6),

545

where R^2 is the coefficient of determination. In particular, the highest D_{REEs} are those measured in 546 547 the APR16-C2Ad run characterized by the lowest NBO/T (0.70). The NBO/T ratio represents the degree of polymerization of a melt and thus yields information about its viscosity (Giordano and 548 549 Dingwell, 2003). The lower NBO/T ratio of the APR16-C2Ad run with respect to that of the other 550 two runs (0.82 and 0.78 for APR16-1C and APR16-C1Ac, respectively) suggests its higher viscosity. 551 High viscosity values imply a lower diffusion of the elements with a consequent increase of D values 552 due to the inability of diffusion to keep pace with the crystal growth rate (Lofgren et al., 2006). 553 Moreover, the low NBO/T ratio calculated for the APR16-C2Ad run can be explained by considering 554 its higher loss of Fe (-27 % Δ Fe; Table 2) occurred during the experiment with respect to that occurred 555 in the APR16-C1Ac and APR16-1C runs (-20 and -5 % Δ Fe, respectively; Table 2). This difference, 556 that suggests a different melt viscosity between the runs, highlights the strong influence of this latter 557 parameter on the partitioning of trace elements between cpx and basaltic melt.

558

559 4.3 Deep differentiation of primitive parental K-basaltic magmas of the CFVD

The partitioning of trace elements between crystals and melts provides an important petrogenetic tool for understanding magmatic processes. Trace elements in volcanic rocks tend to fractionate into specific minerals and therefore are useful in formulating models for magmatic differentiation. Nevertheless, phenocrysts formed in deeper magmatic reservoirs are frequently cannibalized during their ascent in the plumbing system (e.g., Tecchiato et al., 2018b, 2018a). Our study shows that the mineral assemblage obtained from high pressure experiments is similar to that obtained from mafic volcanic products of Campi Flegrei (Table 2). This petrological evidence allows us to validate 567 geochemical models. In the Campi Flegrei Volcanic District mafic products (shoshonitic basalts, 568 trachybasalts, basanites and tephrites) are poorly represented with respect to the differentiated 569 compositions (trachyte or phonolite) that are the largest part of the magma volume erupted (several 570 hundreds of cubic kilometers; e.g., Orsi et al., 1992; Fedele et al., 2003). This suggested that a huge 571 volume of magmas should be differentiated by ponding at variable depths within the crust, producing 572 cumulates joint to the most evolved magmas that have fed the volcanism of the Campania Plain over 573 the past ca. 1.8 Ma (Fedi et al., 2018 and references therein). A first step of differentiation has been 574 supposed at upper mantle-crustal boundary (i.e., at the Moho = ~ 25 km of depth; Di Stefano et al., 575 2011) on the basis of the results of melt inclusion studies (Mormone et al., 2011; Moretti et al., 2013), 576 seismic data interpretations (Di Stefano et al., 1999; Piochi et al., 2005; Chiarabba et al., 2008) and 577 gravimetric and petrological modelling (Fedi et al., 2018). As shown in Perinelli et al. (2019), a stage 578 of high-pressure crystallization is also revealed by the estimate of T-P crystallization conditions of 579 clinopyroxenes phenocrysts from the less differentiated CFVD products indicating crystallization 580 temperatures and pressures between 1140 and 1220 °C and between 0.6 and 0.9 GPa, respectively. 581 Fedi et al. (2018) combining gravimetric data with petrological modelling suggest that a first step of 582 differentiation of the Campi Flegrei primary magmas may occur as a result of the underplating at the 583 Moho level. In this deep reservoir K-trachybasalt melts could be originated after about 30 wt% 584 fractionation of gabbrous cumulates. On the other hand, our experimental results along with those 585 reported in Perinelli et al. (2019) indicate that at pressure corresponding to the Moho depth 586 plagioclase has a limited role in the differentiation of K-trachybasalts from parental APR16-like 587 magmas. Indeed, residual melts of trachybasalt can formed after the fractionation of 20-30% of a 588 clinopyroxenitic mineral assemblage (Table 2 of this study and Table 2 in Perinelli et al., 2019) in a 589 range of temperature that depending on water content (1 up to 3 wt%), varies from 1250 and 1150 590 °C. To examine the role of plagioclase in the deep differentiation process of a primitive parental K-591 basaltic magma we calculated the concentration of selected elements in residual trachybasaltic melts 592 due to crystal-liquid fractionation process (FC). To model the FC process, we used the Excel©

593 spreadsheet program "FC-AFC-FCA and mixing modeler" (Ersoy and Helvaci, 2010) considering: 594 1) the partition coefficients clinopyroxene/melt determined in this study from the longest experiments 595 for which clinopyroxene is in equilibrium with the residual melt (Table S2 in Supplementary Material 596 2) and 2) a fractionate mineral assemblage formed by clinopyroxene + olivine \pm plagioclase (Fig. 9). Modelling results indicate that significant amounts of plagioclase crystallization (>10 wt%) did not 597 598 fully reproduce the composition of CFVD trachybasalts, particularly for the HREEs (Fig. S2 in 599 Supplementary Material 3). The latter increase also using the highest HREE partition coefficients 600 between clinopyroxene and glass obtained in the longest or hydrated experiments (i.e., APR16-C2Bd 601 and APR16-C3Ac, Table 2 and Table 4). Particularly, by increasing the percentage of plagioclase in 602 the fractionated solid, a slight increase in incompatible elements is observed, including the HREEs. 603 Moreover, it is clearly seen that the increase in plagioclase fails to model the Sr abundances. This is 604 a key aspect since Sr is compatible in plagioclase and compatibles elements are much more sensitive 605 than incompatible ones to low degrees of fractionation. On the contrary, the fractionation of 15-26 606 wt% of clinopyroxene + 2-6 wt% of olivine, corresponding to results of the crystallization 607 experiments in which the residual melts had a trachybasalt composition (Table 2 of this study and 608 Table 2 in Perinelli et al., 2019), well mimics the trace element pattern of natural trachybasalts (Fig. 609 9).

610

611 5. Conclusions

This study provides new experimental data on trace element partition coefficients between clinopyroxene and a K-basaltic melt, showing the influence of temperature, chemical composition and also kinetic effects like growth rate and order-disorder crystal growth. Particularly, the rapid growth of clinopyroxene under disequilibrium conditions and/or in presence of pre-existing nuclei under near-equilibrium conditions causes less efficient rejection of incompatible elements (e.g., REEs), since disordered crystal growth is the preferred process during the crystallization, yielding higher D_{REE} than those calculated in presence of lower growth rates, generally under equilibrium 619 conditions. Since growth rates are fast at the beginning of the crystallization process and then decrease 620 with increasing time, D_i values should decrease with increasing time. Therefore, the increase in D_{REE} values with time observed in some runs of this work is not related to a change in time but rather to 621 622 the different NBO/T ratio. The low NBO/T ratio calculated in the run with longer duration (9 hours) 623 is linked to its higher loss of Fe with respect to the other run at 3 and 6 hours, pointing out the strong 624 influence of melt structure and thus of melt viscosity on the partitioning behaviour of trace elements 625 between cpx and basaltic melt. Finally, by using the partition coefficients clinopyroxene/melt determined in this study, geochemical data modelling applied to natural less evolved products of 626 Campi Flegrei Volcanic District, indicate that the differentiation process at mantle-crustal boundary 627 628 depth is controlled by fractionation of clinopyroxene >> olivine mineral assemblage and that K-629 trachybasalts can derive from parental APR16-like magmas after the fractionation of about 20-30% 630 of a clinopyroxenitic cumulate.

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632

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995 **Figures captions**

Figure 1 Backscattered (BSE) images of experimental samples. Details of the experimental conditions are reported at the bottom of each image. Figures a) and b) show the variation of habit and sizes with time in the runs performed using the glassy starting material (i.e., APR16GL): a) subhedral Cpx with sizes of ~5-10 μ m in the shortest run APR16-C2Aa (0.25 hours) and b) euhedral Cpx with sizes up to ~20 μ m in the longest run APR16-C2Ad (9 hours); c) Cpx with euhedral habit and size up to ~50 μm crystallized in the APR16-35 run performed using the natural starting material (i.e., APR16
natural powder). Abbreviation: Cpx: clinopyroxene; OI: olivine; GI: glass.

1003 Figure 2 a) Clinopyroxene compositions plotted into the pyroxene classification diagram (Morimoto 1004 et al., 1998). Wo: wollastonite; En: enstatite; Fe: ferrosilite. The inset show a magnification of the 1005 area of interest. Synthetic samples are indicated with different colours: in orange the runs at 1250 °C, 1006 in yellow the runs at 1225 °C, in light blue the runs at 1220 °C, in blue the runs at 1170 °C, in light 1007 green the run at 1080 °C. Duration of the experimental runs is indicated with different symbols: circle 1008 for 0.25 hours, triangle for 3 hours, diamond for 6 hours and square for 9 hours. b) Composition of 1009 the experimental glasses plotted in the TAS diagram (Le Maitre et al., 2002). Coloured bar indicates 1010 the variation of temperature of the runs that decreases from red to green. Symbols are as in Figure 2a. 1011 Crosses indicate the APR16 (in black) and the APR16GL (in red) starting materials.

Figure 3 C1-normalized (Anders and Grevesse, 1989) trace elements abundances for synthetic clinopyroxenes. Symbols are as in Figure 2. For comparison we reported C1-normalized trace elements abundances for natural Cpx (SCE and SOG samples; Supplementary Material 1), indicated with the grey field.

Figure 4 C1-normalized (Anders and Grevesse, 1989) trace elements abundances for analysed synthetic glasses. Symbols are as in Figure 2. Black crosses indicate data for the APR16 natural samples from Mazzeo et al. (2014). The inset shows a magnification of REE + Y abundances. Yellow field represents REE abundances for natural glasses of Campi Flegrei Volcanic District (CFVD) products (data from GEOROC database; http://georoc.mpch-mainz.gwdg.de/georoc/Start.asp).

1021Figure 5 Calculated $^{Cpx/L}D$ values for trace elements in experimental samples. Symbols are as in1022Figure 2. Yellow field represents $^{Cpx/L}D_{REE}$ values for natural Campi Flegrei Volcanic District (CFVD)1023products (data from Pappalardo et al. (2008), Fedele et al. (2009) and Mollo et al. (2016)).

Figure 6 Onuma diagrams for clinopyroxene-liquid partition coefficients for mono-, di-, tri-, and tetravalent cations versus ionic radii (in Å; Shannon, 1976) for VI- and VIII-fold coordinated sites along with the results of the lattice strain fits in M1 and M2 sites for samples (a) APR16-1C (1225 1027 °C, 3 hours), (b) APR16-35 (1225 °C, 6 hours), (c) APR16-C3Ac (1080 °C, 6 hours) and (d) APR16-C2Bd (1170 °C, 9 hours). The dotted curves represent the fits of the lattice strain model to $^{Cpx/L}D_{i}$ 1028 values while symbols represent measured $C_{px/L}D_i$ values. Cations are indicated as follow: monovalent 1029 1030 with orange triangles, divalent with yellow squares, trivalent with blue circles and tetravalent with 1031 green diamonds. Ba and V (yellow empty square) have not been used in the fit of the LSM for divalent 1032 cations. Yellow full squares indicate D_i calculated for M1 and M2 sites. Blue full circle represents $D_{\rm Eu}$ measured in this study, plotted together with the calculated $D_{\rm Eu3+}$ and $D_{\rm Eu2+}$ values (see text for 1033 1034 details).

Figure 7 Onuma diagrams showing the variation of predicted REE partitioning between cpx and basaltic melt as function of a) composition (Al^{IV}), and b) composition (Al^{IV}), temperature and undercooling. a) The increase in Al^{IV} content shifts the parabola upward (D_0 increases) and towards left (r_0 decreases); b) the increase in undercooling moves the parabola upward like the Al^{IV} content, while the increase in temperature shifts the parabola downward.

Figure 8 Plot of clinopyroxene-liquid partition coefficients for REEs vs ionic radius (Å). The curves 1040 represent the fits of the lattice strain model to $^{Cpx/L}D_i$ values while symbols represent measured $^{Cpx/L}D_i$ 1041 1042 values for Dy (HREE), Eu³⁺ (MREE) and Nd (LREE) in the APR16-C2Ad (1220 °C; in light blue), APR16-C1Ac (1250 °C; in orange) and APR16-1C (1225 °C; in yellow) runs. Horizontal arrow 1043 1044 indicates the diffusivity decease from LREEs to HREEs, while vertical arrow indicates NBO/T 1045 increase and viscosity decrease with decreasing time (from APR16-C2Ad to APR16-1C). On the right side of the figure are reported plots of partition coefficients for Dy, Eu³⁺ and Nd vs NBO/T, in which 1046 1047 it is possible to see the negative correlation between D values and NBO/T.

Figure 9 Primitive Mantle normalized (McDonough and Sun, 1995) trace elements patterns for primitive basaltic (APR22; D'Antonio et al., 1999) and trachybasaltic (APR19; D'Antonio et al., 1999) rocks, and the predicted composition of differentiated melts obtained after 20 and 28 % of fractional crystallization of APR22 basalt assumed as primitive parental K-basaltic magma. The reported degrees of fractionation (FC) reflect the crystallized phase assemblage and proportions

1053	determined for runs APR16-C2Ad ($Cpx_{(17)}+Ol_{(3)}$) and AR16-C2Ab ($Cpx_{(23)}+Ol_{(4)}+Plg_{(1)}$) that
1054	produced trachybasaltic residual glass. The modelled compositions were calculated by using the
1055	Excel© spreadsheet program of Ersoy and Helvaci (2010) (see §4.3 for details). Normalized patterns
1056	for Campi Flegrei Volcanic District basalts and trachybasalts (D'Antonio et al., 1999; Pappalardo et
1057	al., 1999; De Astis et al., 2004), indicated with red and green field, respectively, are shown for
1058	comparison.

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- 1060 Tables captions
- Table 1 Composition (wt.%) of APR16 rock sample, APR16GL starting material, SCE and SOG rock
 samples
- 1063 **Table 2** Experimental conditions of the experimental samples
- 1064 Table 3 Major and trace element concentrations (ppm) for synthetic clinopyroxene crystals and1065 glasses
- Table 4 Major and trace elements partition coefficients between clinopyroxene and glass for
 experimental samples
- 1068 **Table 5** Best fits for E, r_0 and D_0 obtained by regression of clinopyroxene/liquid partition coefficients
- 1069 of divalent, trivalent and tetravalent cations in M1 and M2 sites
- 1070
- 1071 Supplementary materials
- 1072 Supplementary Material 1 contains information about natural SCE and SOG samples
- 1073 Supplementary Material 2 contains supplementary tables
- 1074 Supplementary Material 3 contains supplementary figures

Sample	APR16	APR16GL	SCE	SOG
SiO ₂	48.89	49.30 (0.46)	47.62	47.78
TiO ₂	1.23	1.33 (0.03)	1.23	1.23
Al ₂ O ₃	15.65	15.31 (0.14)	15.33	15.43
FeO _{tot}	8.08	7.79 (0.13)	8.19	8.24
MnO	0.14	0.14 (0.05)	0.14	0.15
MgO	8.89	9.31 (0.07)	9.78	9.75
CaO	11.64	12.02 (0.09)	11.89	11.59
Na ₂ O	2.88	2.86 (0.05)	2.69	2.83
K ₂ O	1.52	1.60 (0.02)	1.58	1.67
P2O5	0.31	0.55 (0.02)	0.29	0.29
Total	99.99	99.31 (0.29) [§]	99.65	99.88
L.O.I.	0.61	-	0.26	0.28
Mg# 1	0.66	0.68	0.68	0.68

Table 1 Composition (wt.%) of APR16 rock sample, APR16GL startingmaterial, SCE and SOG rock samples

Notes: 1 [MgO/(MgO + FeO_{tot})] molar, assuming all Fe²⁺ and Fe³⁺ as FeO_{tot}. Chemical composition of APR16 is from D'Antonio et al. (1999); APR16GL composition is from Perinelli et al. (2019); SCE and SOG from Bonechi et al. (2020b). § For APR16GL the analysis is normalized to 100 % but the original total is reported. Standard deviation is reported in parenthesis.

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Runs	Starting material	P (GPa)	T (°C)	ΔT (°C)	Time (h)	G _{LCpx} (cm s ⁻¹)	H ₂ O _i (wt.%)	H ₂ O _f (wt.%)	NBO/ T	Phases		% ∆Fe§	References
APR16- 1C APR16- 35	APR16 APR16	0.8 0.8	1225 1225	55 55	3 6	1.59·10 ⁻⁷ 8.34·10 ⁻⁸	NWA NWA	0.96±0.29 1.7±0.42	0.82 0.80	$Gl(94) + Cpx(6) + Ol^{\#} + Ox^{\#}$ $Gl(96) + Cpx(4) + Ol^{\#}$	0.38 0.14	-5 -22	Bonechi (2020)
APR16- C1Ac APR16-	APR16GL	0.8	1250	45	6	2.06.10-8	NWA	2.03±0.43	0.78	Gl(83) + Cpx(15) + Ol(2)	0.83	-20	
C1Ad APR16- C2Aa	APR16GL APR16GL	0.8 0.8	1250 1220	45 50	9 0.25	1.73·10 ⁻⁸ 3.24·10 ⁻⁷	NWA 2	1.95±0.38 1.92±0.34	0.85 0.92	Gl(90) + Cpx(10) Gl(91) + Cpx(4) + Ol(1) + Ox(4)	0.65 0.02	-19 -25	
APR16- C2Ab APR16-	APR16GL APR16GL	0.8 0.8	1220 1220	50 50	3 6	$3.04 \cdot 10^{-8}$ $2.67 \cdot 10^{-8}$	1.9 2.1	1.56±0.36 2.47±0.35	0.61 0.60	$Gl(72) + Cpx(23) + Ol(4) + Plg(1) + Ox^*$ Gl(65) + Cpx(27) + Ol(5) + Plg(3)	0.66 0.93	-17 -25	Bonechi et al. (2020b)
APR16- C2Ad	APR16GL	0.8	1220	50	9	1.93.10-8	1.9	1.3±0.38	0.70	$Gl(80) + Cpx(17) + Ol(3) + Ox^*$	0.25	-26	
C2Bc APR16- C2Bd	APR16GL APR16GL	0.8 0.8	1170 1170	100 100	6 9	$2.77 \cdot 10^{-8}$ $2.34 \cdot 10^{-8}$	2 3.77 ± 0.1 0.67 $Gl(58) + Cpx(33) + Ol(5) + Plg(1)$ 0 1.8 2.02 ± 0.46 0.61 $Gl(57) + Cpx(34) + Ol(2) + Plg(3) + Ox(4)$ 0		0.73 0.56	-27 -4			
APR16- C3Ac	APR16GL	0.8	1080	144	6	1.31.10-8	3.8	6.5±0.35	1.00	Gl(60) + Cpx(37) + Ol(3)	0.62	-10	Bonechi et al. (2020a)

Table 2 Experimental conditions of the experimental samples

Notes: Gl = glass; Cpx = clinopyroxene; Ol = Olivine; Plg = plagioclase; Ox = oxide; NWA = No Water Added. ΔT is the difference between the Cpx liquidus temperature and the nominal temperature of the runs; G_{LCpx} is the Cpx growth rate taken from Bonechi et al. (2020b); H₂O_i is nominal content (wt.%) of water added to the charge; H₂O_f = water content in the melt determined by the by-difference calculation method (respect to the total of EMP analyses); NBO/T is the ratio of non-bridging oxygens to number of structural tetrahedra in the liquid calculated as in Mysen (1985) considering the H₂O_f of the runs. Σr^2 values from mass balance calculation. § Percent of relative Fe loss from starting material based on difference between the starting material and calculated bulk composition of experiment obtained from mass-balance calculations.* crystals present in trace; # pre-existing crystals.

Run#	APR16-1C	APR16-35	APR16-C1Ac	APR16- C1Ad	APR16- C2Aa	APR16-C2Ab	APR16-C2Ac	APR16-C2Ad	APR16-C2Bc	APR16-C2Bd	APR16- C3Ac
Phase	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cpx	cpx
1 huse	glass	glass	glass	glass	glass	glass	glass	glass	glass	glass	glass
Na	2560 (838)	2950 (307)	3709 (351)	3709 (265)	3041 (302)	4525 (452)	6083 (433)	3561 (363)	bdl	5638 (502)	4154 (341)
INA	23215 (275)	22163 (225)	25074 (213)	23591 (203)	21365 (211)	23665 (198)	23294 (231)	22107 (220)	-	28339 (223)	27226 (207)
к	279 (102)	103 (52)	581 (103)	498 (98)	581 (92)	1909 (232)	1826 (312)	415 (105)	bdl	1079 (343)	913 (123)
K	14562 (291)	14693 (204)	15772 (206)	14942 (232)	13033 (243)	17599 (221)	20754 (209)	16270 (214)	-	21916 (225)	15275 (213)
Ca	150086 (5800)	150086 (6100)	129450 (5980)	147514 (4800)	137400 (6400)	145155 (5100)	146086 (4350)	152874 (5210)	bdl	146585 (4600)	153017 (3980)
	73870 (1250)	74268 (1130)	71242 (3850)	80403 (2100)	86335 (4120)	76620 (3560)	58748 (3400)	77187 (2980)	-	56533 (3230)	67039 (4550)
	38643 (4451)	22399 (2350)	32569 (2310)	34357 (1890)	35134 (1930)	34823 (2100)	34357 (1850)	31248 (2210)	bdl	42130 (1900)	45317 (2530)
Fe	61647 (1208)	30246 (903)	48193 (950)	48193 (860)	44696 (810)	42519 (950)	42674 (890)	34901 (730)	-	47183 (1010)	46950 (980)
	84353 (6092)	96961 (6310)	90573 (11665)	95300 (7600)	81600 (4300)	108000 (16000)	87000 (8200)	110000 (13000)	5900 (110)	83000 (7900)	86933 (4600)
Mg	43253 (507)	45112 (466)	46575 (1800)	46456 (1540)	44060 (1480)	45660 (1840)	47100 (3125)	32183 (1225)	28075 (2869)	33100 (1925)	33280 (1580)
4.1	25520 (6890)	28709 (7650)	59765 (15843)	41540 (5760)	22900 (2200)	28700 (160)	41070 (3000)	36500 (1700)	5750 (95)	36300 (3300)	37783 (1700)
Al	83792 (2340)	83050 (3050)	94219 (3525)	92220 (3100)	73560 (3080)	128750 (11675)	105150 (4800)	96220 (6920)	102650 (3830)	103575 (5925)	99238 (3920)
Mn	880 (130)	915 (82)	900 (110)	1100 (199)	1080 (320)	1140 (170)	1143 (95)	1210 (120)	bdl	1054 (92)	1271 (97)
	1151 (10)	1127 (13)	1211 (12)	1296 (30)	1136 (21)	1299 (35)	1310 (172)	1359 (43)	1185 (107)	1307 (42)	1254 (28)
T :	4370 (259)	4257 (1041)	2684 (29)	3275 (1136)	3570 (910)	3510 (570)	4440 (650)	6410 (670)	bdl	8050 (870)	5270 (139)
11	7296 (92)	7395 (113)	7070 (40)	7180 (90)	6432 (100)	8253 (388)	6405 (61)	7906 (205)	7575 (367)	8625 (128)	7526 (150)
7.	34 (11)	31 (2)	18 (3)	33 (10)	bdl	26 (9)	33 (7)	60 (12)	bdl	82 (12)	40 (2)
Zr	103 (2)	105 (2)	111 (5)	111 (2)	99 (2)	129 (7)	198 (9)	120(1)	133 (5)	143 (2)	128 (2)
NIL	0.11 (0.07)	0.20 (0.04)	bdl	1.1 (0.6)	bdl	2.4 (0.3)	bdl	bdl	bdl	bdl	0.7 (0.1)
ND	13.1 (0.3)	13.1 (0.3)	13.7 (0.5)	12.7 (0.6)	10.2 (0.6)	15(1)	26.2 (0.1)	14(1)	17 (2)	18(1)	14.1 (0.4)
Цf	1.6 (0.4)	1.6 (0.2)	bdl	bdl	bdl	bdl	1.2 (0.2)	2(1)	bdl	3.9 (1.4)	1.8 (0.2)
п	2.5 (0.2)	2.6 (0.1)	2.8 (0.1)	2.85 (0.03)	2.2 (0.1)	2.8 (0.3)	3.8 (0.2)	2.7 (0.3)	2.9 (0.3)	3.3 (0.4)	3.3 (0.4)
То	0.05 (0.01)	0.04 (0.01)	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl
1a	0.68 (0.04)	0.67 (0.04)	0.7 (0.2)	0.7 (0.1)	0.5 (0.1)	0.85 (0.05)	1.5 (0.2)	0.8 (0.1)	0.99 (0.13)	1.04 (0.16)	0.9 (0.1)
Th	0.06 (0.01)	0.06 (0.02)	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	0.20 (0.06)
111	2.9 (0.1)	2.9 (0.1)	3.4 (0.2)	3.3 (0.2)	2.5 (0.1)	3.6 (0.4)	6.3 (0.8)	3.2 (0.3)	4.1 (0.4)	4.9 (0.3)	3.7 (0.3)
II	0.01	0.03 (0.01)	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl
0	0.93 (0.05)	0.94 (0.04)	1.0 (0.1)	1.2 (0.1)	0.8 (0.1)	1.1 (0.1)	2.24 (0.04)	1.1 (0.2)	1.4 (0.1)	1.5 (0.2)	1.2 (0.2)
Rb	bdl	bdl	bdl	11 (3)	bdl	bdl	8 (4)	bdl	bdl	bdl	2.2 (0.3)
KU	57 (1)	56 (1)	64 (2)	66 (1)	53.1 (0.1)	71 (2)	120 (4)	69 (4)	78 (6)	95 (1)	70 (3)
Sr	54 (8)	52 (2)	59 (1)	83 (8)	55 (16)	71 (18)	87 (16)	84 (13)	bdl	60 (8)	78 (6)
51	503 (6)	499 (5)	531 (2)	540 (6)	442 (4)	631 (13)	698 (15)	594 (8)	652 (21)	693 (26)	605 (3)
Cs	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl	bdl
05	2.4 (0.1)	2.4 (0.1)	2.7 (0.2)	3.0 (0.2)	2.2 (0.3)	3.3 (0.3)	4.8 (0.1)	2.9 (0.1)	3.3 (0.5)	3.9 (0.2)	3.0 (0.3)
Ba	0.41 (0.01)	0.76 (0.37)	12 (7)	30 (13)	16 (9)	bdl	43 (22)	15 (7)	bdl	bdl	20.8 (0.6)
2	534 (9)	532 (7)	536 (7)	553 (17)	450 (2)	670 (39)	1013 (43)	589 (4)	773 (4)	749 (29)	606 (1)

 Table 3 Major and trace element concentrations (ppm) for synthetic clinopyroxene crystals and glasses

Dh	0.12 (0.15)	0.14 (0.05)	bdl								
FU	1.1 (0.5)	11.7 (0.4)	bdl	bdl	0.9 (0.1)	bdl	bdl	bdl	bdl	bdl	bdl
S a	119 (21)	113 (5)	80 (10)	87 (8)	87 (27)	110 (13)	88 (5)	112 (19)	bdl	155 (14)	93 (3)
30	28 (1)	30(1)	40(1)	40 (2)	40(1)	27 (3)	23 (1)	43 (2)	19(1)	22 (1)	26(1)
V	248 (23)	764 (86)	120 (16)	174 (49)	133 (17)	237 (43)	263 (41)	385 (36)	bdl	505 (50)	199 (8)
v	190 (2)	162 (4)	201 (5)	194 (4)	184 (2)	182 (4)	180 (6)	217 (4)	157 (2)	198 (1)	224 (3)
Cr	3229 (708)	4460 (605)	4445 (345)	2390 (1259)	2210 (44)	1740 (600)	1100 (120)	1200 (190)	bdl	1400 (230)	1617 (92)
CI	80 (12)	96 (9)	77 (23)	57 (1)	318 (2)	35 (17)	31 (11)	163 (3)	bdl	28 (12)	26 (13)
Co	28 (2)	15 (3)	22 (5)	33 (2)	26 (11)	13 (7)	19 (4)	21 (9)	bdl	43 (5)	36(1)
CO	34 (2)	12(1)	24 (1)	25 (1)	27 (1)	19(1)	17 (1)	17 (1)	10(1)	15(1)	21 (10)
Ni	139 (12)	44 (18)	37 (17)	60 (6)	bdl	51 (18)	bdl	bdl	bdl	137 (18)	177 (10)
INI	43 (9)	7 (1)	15 (2)	26 (2)	33 (1)	12(1)	8 (2)	9(1)	bdl	9(1)	12(1)
v	15 (2)	17 (1)	17 (2)	bdl							
1	19.9 (0.3)	20.2 (0.3)	19(1)	bdl							
Iа	1.9 (0.4)	1.8 (0.2)	bdl	1.5 (0.5)	bdl	2.6 (1.1)	bdl	3.9 (1.7)	bdl	4.7 (1.2)	3 (0.2)
La	15.6 (0.3)	16.01 (0.24)	17.1 (0.5)	17.9 (0.3)	14.7 (0.2)	20.9 (0.9)	33 (2)	19 (1)	22 (1)	25 (2)	20(1)
Ce	7.3 (1.7)	6.7 (0.7)	6.7 (0.3)	5.8 (3.4)	bdl	7.8 (2.3)	bdl	11.2 (3.5)	bdl	13 (2)	10.1 (0.5)
ce	34.5 (0.5)	35.1 (0.6)	36 (1)	37 (1)	30(1)	42 (1)	67 (4)	37 (1)	45.2 (0.5)	49 (2)	41 (1)
Pr	1.4 (0.3)	1.3 (0.1)	1.8 (0.3)	1.1 (0.4)	bdl	1.6 (0.7)	1.4 (0.2)	2.1 (0.3)	bdl	2.6 (0.5)	1.9 (0.1)
	4.4 (0.1)	4.5 (0.1)	4.5 (0.2)	5.2 (0.1)	3.9 (0.3)	5 (0.5)	7.8 (1.2)	4.9 (0.2)	5.1 (0.2)	5.9 (0.3)	5.2 (0.3)
Nd	8.5 (1.5)	7.9 (0.7)	11 (1)	7.2 (2.1)	bdl	8.3 (4.2)	7.2 (0.9)	14 (5)	bdl	16 (3)	11.7 (0.4)
itu	19(1)	19.4 (0.5)	20(1)	23 (2)	17 (1)	24 (2)	35 (4)	21 (2)	22 (1)	25 (2)	24 (1)
Sm	3.02 (0.61)	3.03 (0.4)	3.4 (0.6)	3.8 (0.2)	bdl	bdl	bdl	bdl	bdl	5.5 (2.2)	4.4 (1.1)
SIII	4.5 (0.3)	4.7 (0.3)	4.89 (0.03)	5.9 (0.6)	4.4 (0.6)	5.4 (1)	7.7 (0.8)	5.3 (0.6)	5.4 (0.4)	6.1 (1.5)	5.8 (0.4)
Eu	0.9 (0.1)	0.8 (0.1)	0.9 (0.2)	bdl	bdl	bdl	0.8 (0.1)	bdl	bdl	1.1 (0.4)	1.5 (0.2)
Eu	1.5 (0.1)	1.5 (0.1)	1.6 (0.2)	1.9 (0.6)	1.5 (0.2)	1.8 (0.2)	2.5 (0.2)	1.9 (0.3)	1.6 (0.5)	1.9 (0.2)	1.7 (0.1)
Gd	3.7 (0.4)	3.5 (0.3)	3.1 (0.2)	2.7 (0.2)	bdl	bdl	bdl	5.1 (1.6)	bdl	5.8 (2.1)	4.9 (0.9)
ou	4.5 (0.2)	4.4 (0.3)	4.6 (0.4)	5.5 (0.6)	4.1 (0.4)	4.8 (0.7)	6.3 (0.3)	5.3 (0.3)	4.4 (0.5)	5.8 (1.2)	5.1 (0.8)
Tb	0.53 (0.07)	0.57 (0.05)	bdl	0.44 (0.1)	bdl	bdl	0.38 (0.05)	bdl	bdl	1.1 (0.3)	0.68 (0.05)
10	0.63 (0.04)	0.66 (0.05)	0.66 (0.01)	0.74 (0.16)	0.59 (0.08)	0.86 (0.09)	1.02 (0.23)	0.76 (0.13)	0.72 (0.11)	0.88 (0.17)	0.73 (0.11)
Dv	3.3 (0.4)	3.52 (0.05)	3.9 (1.2)	2.6 (0.8)	bdl	bdl	bdl	5.8 (0.4)	bdl	4.7 (1.7)	4.4 (0.5)
29	3.8 (0.2)	3.9 (0.2)	3.8 (0.1)	4.5 (0.4)	3.6 (0.3)	4.9 (0.4)	5.6 (0.2)	4.1 (0.4)	4.2 (0.6)	4.2 (0.2)	4.2 (0.3)
Ho	0.64 (0.07)	0.66 (0.05)	0.76 (0.21)	bdl	bdl	0.59 (0.33)	bdl	bdl	bdl	0.89 (0.39)	0.84 (0.12)
110	0.75 (0.05)	0.77 (0.04)	0.81 (0.1)	0.92 (0.18)	0.65 (0.08)	0.93 (0.16)	1.13 (0.22)	0.81 (0.14)	0.80 (0.14)	0.90 (0.12)	0.90 (0.12)
Er	1.59 (0.27)	1.75 (0.18)	1.53 (0.34)	1.12 (0.11)	bdl	bdl	bdl	bdl	bdl	2.62 (0.95)	2.14 (0.12)
21	2.12 (0.13)	2.04 (0.15)	2.04 (0.19)	2.31 (0.29)	1.94 (0.18)	2.22 (0.19)	3.07 (0.6)	2.14 (0.2)	2.09 (0.35)	2.15 (0.3)	2.11 (0.34)
Tm	0.20 (0.03)	0.22 (0.03)	0.19 (0.08)	bdl	0.32 (0.04)						
TIII	0.27 (0.02)	0.29 (0.04)	0.28 (0.02)	0.36 (0.05)	0.25 (0.03)	0.32 (0.03)	0.37 (0.04)	0.31 (0.04)	bdl	0.30 (0.05)	0.28 (0.09)
Yb	1.24 (0.22)	1.43 (0.16)	bdl	1.73 (0.29)							
10	1.78 (0.15)	1.87 (0.21)	1.91 (0.38)	2.34 (0.17)	1.87 (0.32)	2.07 (0.44)	2.65 (0.07)	2.01 (0.27)	2.01 (0.57)	2.26 (0.3)	1.78 (0.68)
Lu	0.17 (0.02)	0.21 (0.01)	bdl	0.26 (0.03)							
Lu	0.27 (0.03)	0.27 (0.02)	0.29 (0.06)	0.36 (0.06)	0.28 (0.08)	0.33 (0.03)	0.49 (0.12)	0.33 (0.04)	0.31 (0.03)	0.4 (0.09)	0.31 (0.05)

Notes: cpx = clinopyroxene; standard deviation is reported in brackets.

For single analysis Int2SE (i.e., analytical error) is reported also in brackets but in italic; bdl = below detection limit.

Run#	APR16-1C	APR16-35	APR16- C1Ac	APR16- C1Ad	APR16-C2Aa	APR16- C2Ab	APR16- C2Ac	APR16-C2Ad	APR16-C2Bd	APR16-C3Ac
Na	0.11 (0.01)	0.13 (0.02)	0.15 (0.01)	0.16 (0.02)	0.14 (0.02)	0.19 (0.02)	0.26 (0.03)	0.16 (0.01)	0.20 (0.03)	0.15 (0.01)
K	0.019 (0.001)	0.007 (0.001)	0.04 (0.01)	0.03 (0.01)	0.04 (0.01)	0.11 (0.01)	0.088 (0.005)	0.026 (0.005)	0.05 (0.01)	0.042 (0.002)
Ca	2.03 (0.14)	2.02 (0.03)	1.82 (0.15)	1.83 (0.15)	1.59 (0.12)	1.89 (0.12)	2.47 (0.25)	1.98 (0.15)	2.59 (0.25)	2.28 (0.13)
Fe	0.59 (0.07)	0.61 (0.08)	0.66 (0.08)	0.69 (0.08)	0.77 (0.08)	0.77 (0.08)	0.78 (0.08)	0.88 (0.08)	0.87 (0.08)	0.97 (0.09)
Fe _{M1}	0.293	0.367	0.329	0.421		0.410	0.411	0.45	0.535	0.672
Fe _{M2}	0.297	0.247	0.329	0.268		0.358	0.372	0.43	0.338	0.293
Mg	1.94 (0.04)	1.92 (0.07)	2.23 (0.24)	2.12 (0.72)	1.85 (0.12)	2.41 (0.37)	2.55 (0.38)	2.03 (0.14)	2.5 (0.3)	2.68 (0.27)
Mg _{M1}	1.809	1.811	2.074	1.991		2.232	2.382	1.809	2.353	2.552
Mg _{M2}	0.135	0.106	0.158	0.129		0.18	0.167	0.219	0.143	0.129
Al	0.42 (0.12)	0.38 (0.12)	0.33 (0.17)	0.29 (0.06)	0.36 (0.03)	0.36 (0.04)	0.46 (0.02)	0.35 (0.02)	0.49 (0.04)	0.37 (0.02)
Al_T	0.225	0.185	0.181	0.155	0.187	0.167	0.222	0.199	0.257	0.199
Al_{M1}	0.191	0.192	0.153	0.137	0.169	0.189	0.242	0.148	0.233	0.171
Mn	0.72 (0.36)	0.63 (0.21)	0.75 (0.41)	0.62 (0.27)		0.74 (0.20)	0.91 (0.25)	0.91 (0.37)	0.85 (0.28)	0.74 (0.30)
Ti	0.56 (0.15)	0.58 (0.09)	0.38 (0.09)	0.46 (0.12)	0.55 (0.14)	0.42 (0.07)	0.69 (0.07)	0.81 (0.11)	0.93 (0.12)	0.70 (0.06)
Zr	0.33 (0.11)	0.30 (0.02)	0.16 (0.03)	0.3 (0.1)		0.20 (0.01)	0.16 (0.01)	0.500 (0.004)	0.57 (0.01)	0.31 (0.02)
Nb	0.008 (0.005)	0.015 (0.003)		0.087 (0.004)			0.094 (0.001)			0.05 (0.01)
Hf	0.67 (0.17)	0.62 (0.08)					0.31 (0.01)	0.73 (0.08)	1.19 (0.13)	0.56 (0.09)
Та	0.07 (0.01)	0.06 (0.01)								
Th	0.020 (0.005)	0.02 (0.01)								0.055 (0.001)
U	0.011 (0.001)	0.03 (0.01)								
Rb				0.166 (0.003)			0.068 (0.003)			0.031 (0.003)
Sr	0.11 (0.02)	0.104 (0.004)	0.111 (0.002)	0.153 (0.002)	0.122 (0.001)	0.106 (0.002)	0.125 (0.003)	0.141 (0.002)	0.086 (0.003)	0.13 (0.01)
Ba	0.001 (0.001)	0.001 (0.001)	0.0232 (0.0003)	0.054 (0.002)	0.0349 (0.0002)		0.042 (0.002)	0.0248 (0.0002)		0.034 (0.001)
Pb	0.11 (0.05)	0.012 (0.004)								
Sc	4.24 (0.74)	3.74 (0.19)	2.01 (0.26)	2.18 (0.23)	2.20 (0.08)	4.09 (0.40)	3.84 (0.17)	2.58 (0.12)	7.06 (0.24)	3.60 (0.18)

Table 4 Major and trace elements partition coefficients between clinopyroxene and glass for experimental samples

V	1.30 (0.12)	4.71 (0.54)	0.597 (0.083)	0.892 (0.252)	0.722 (0.008)	1.31 (0.03)	1.46 (0.04)	1.77 (0.03)	2.55 (0.01)	0.886 (0.037)
Cr	41 (11)	47 (8)	57 (18)	42 (22)	6.9 (0.1)	50	35	7.3 (1.2)	50	63 (4)
Со	0.83 (0.07)	1.25 (0.23)	0.92 (0.21)	1.32 (0.08)	0.95 (0.40)	0.68 (0.38)	1.12 (0.24)	1.21 (0.52)	2.87 (0.35)	1.71 (0.78)
Сом1	0.64	1.11	0.71	1.15		0.46	0.9	0.93	2.67	1.54
Co _{M2}	0.18	0.14	0.2	0.17		0.22	0.22	0.28	0.19	0.17
Ni	3.21 (0.71)	6.29 (2.81)	2.38 (1.17)	2.33 (0.28)		4.31 (0.43)			15.6 (2.3)	15 (2)
Y	0.77 (0.10)	0.85 (0.03)	0.89 (0.12)							
La	0.12 (0.03)	0.11 (0.01)		0.082 (0.001)		0.124 (0.005)		0.20 (0.01)	0.19 (0.01)	0.15 (0.09)
Ce	0.21 (0.05)	0.19 (0.02)	0.19 (0.01)	0.16 (0.09)	0.19 (0.01)			0.28 (0.01)	0.25 (0.01)	0.25 (0.01)
Pr	0.33 (0.07)	0.29 (0.03)	0.4 (0.1)	0.213 (0.005)		0.31 (0.03)	0.18 (0.03)	0.42 (0.01)	0.44 (0.02)	0.36 (0.03)
Nd	0.45 (0.08)	0.41 (0.03)	0.53 (0.07)	0.31 (0.09)		0.35 (0.02)	0.21 (0.02)	0.64 (0.05)	0.66 (0.04)	0.49 (0.03)
Sm	0.67 (0.14)	0.65 (0.09)	0.69 (0.13)	0.64 (0.06)					0.90 (0.03)	0.77 (0.17)
Eu	0.64 (0.1)	0.54 (0.06)	0.59 (0.13)				0.31 (0.02)		0.57 (0.06)	0.84 (0.13)
Eu^{2+}	0.137	0.132	0.140	0.193				0.179	0.111	0.169
Eu^{3+}	0.756	0.729	0.772	0.718				1.071	1.032	0.877
Gd	0.82 (0.13)	0.78 (0.09)	0.67 (0.06)	0.49 (0.05)				0.95 (0.05)	1.01 (0.20)	0.97 (0.23)
Tb	0.83 (0.12)	0.87 (0.11)		0.59 (0.13)			0.38 (0.08)		1.29 (0.25)	0.93 (0.15)
Dy	0.86 (0.11)	0.90 (0.05)	1.02 (0.32)	0.59 (0.05)				1.40 (0.13)	1.12 (0.05)	1.04 (0.14)
Но	0.85 (0.10)	0.86 (0.08)	0.94 (0.28)			0.64 (0.11)			0.99 (0.13)	0.93 (0.18)
Er	0.75 (0.13)	0.86 (0.08)	0.75 (0.18)	0.49 (0.06)					1.22 (0.17)	1.01 (0.17)
Tm	0.75 (0.12)	0.77 (0.15)	0.68 (0.29)							1.14 (0.39)
Yb	0.70 (0.13)	0.76 (0.12)								0.97 (0.40)
Lu	0.62 (0.11)	0.75 (0.07)								0.83 (0.16)

Notes: standard deviation is reported in brackets. D_{M1} and D_{M2}, and Eu²⁺ and Eu³⁺ partition coefficients were recalculated following the procedure described in section 4.1.

Run#	APR16-1C	APR16-35	APR16-C1Ac	APR16-C1Ad	APR16- C2Aa	APR16- C2Ab	APR16- C2Ac	APR16-C2Ad	APR16-C2Bd	APR16-C3Ac
M2 ²⁺	2 28+0 20	2 28 - 0 40	2 25 . 0 22	2.02.0.71	2 21 0 07	2 27 . 0 40	2 07 0 47	246-028	2 10 0 10	2 (8+0.22
D_0	2.38±0.29 1.081±0.005	2.38±0.49 1.083±0.002	2.25±0.33 1.079+0.005	2.02 ± 0.71 1.088+0.003	2.31 ± 0.07 1 059+0 001	2.27 ± 0.40 1.077±0.006	3.07 ± 0.47 1.081±0.006	2.46±0.38 1.078+0.005	3.19±0.49 1.080+0.003	2.08 ± 0.32 1 084+0 004
r_0 (WB97)	1.081	1.082	1.082	1.083	1.037±0.001	1.079	1.076	1.084	1.078	1.084
E	271±22	287±20	260±24	246±31	200±20	252±29	274±27	236±22	286±19	249±18
E (WB97)	180	180	179	179		180	180	180	181	183
N#12+										
MI ⁻	5 34+1 35	7 07+3 11	4 64+1 84	3 74+1 01		6 55+1 27	6 50+2 65	3 47+0 78	9 95+0 07	7 37+1 92
r_0	0.616±0.014	0.610 ± 0.011	0.621 ± 0.013	0.639±0.013		0.614 ± 0.013	0.617 ± 0.012	0.646±0.009	0.634±0.004	0.634±0.009
E	504±133	461±60	484±72	523±148		473±82	468±54	530±74	596±5	455±77
M2 ³⁺	0.960 0.006	0.957.0.002	0.870 0.001	0 (14) 0 001		0 (52 0 005	0 272 0 009	1 1 65 0 012	1 128 0 002	1 120 0 000
D_0	0.860 ± 0.006 1 0297 ± 0 0002	1.0257 ± 0.002	1.0300 ± 0.001	1.022 ± 0.001		1.033 ± 0.005 1.033 ±0.001	1.033 ± 0.008 1.033±0.001	1.165 ± 0.015 1.0301 ± 0.0003	1.128 ± 0.002 1.0323 ± 0.0003	1.139 ± 0.008 1.0292 ± 0.0005
r_0 (WB97)	1.0208	1.0221	1.0220	1.023		1.019	1.016	1.0239	1.0177	1.0245
E	325±6	326±3	322±1	307±4		305±3	304±3	304±1	310±4	300±2
E (WB97)	270	270	269	269		270	270	270	272	275
M13+										
D ₀	40.05+0.07	40.07+0.10	40 12+0 28	40.05+0.11	11 01+0 55	40 07+0 11	40.03+0.06	20.04+0.05	42 21+1 99	40 11+0 51
r_0	0.664±0.008	0.665±0.006	0.654±0.004	0.655±0.017	0.664±0.002	0.661±0.002	0.660±0.01	0.666±0.001	0.672±0.001	0.663±0.001
E	1499±2	1499±1	1499±1	1499±5	1202±48	1499±1	1499±1	1499±1	1499±4	1499±1
MI**	2 60 1 20	2 50 1 21					1.09+0.92	2.52+0.00	6 21 1 22	2.21 ± 0.04
\mathbf{D}_0	0.659+0.001	0.659 ± 0.001					0.650 ± 0.005	2.52 ± 0.99 0.658+0.002	0.21 ± 1.82 0.660+0.001	2.31 ± 0.94 0.656+0.002
r_0 (H11)	0.649±0.007	0.649±0.007	0.649±0.007	0.649±0.007	0.649±0.007	0.649±0.007	0.649±0.007	0.649±0.007	0.649±0.007	0.649±0.007
E	3077±559	3115±634					2404±703	1967±623	3014±518	2024±578
E (H11)	2936±722	2936±722	2936±722	2936±722	2936±722	2936±722	2936±722	2936±722	2936±722	2936±722

Table 5 Best fits for *E*, *r*₀ and *D*₀ obtained by regression of clinopyroxene/liquid partition coefficients of divalent, trivalent and tetravalent cations

in M1 and M2 sites

Note: D_0 : strain-compensated partition coefficient. *E*: Young Modulus of the lattice site (GPa). r_0 : optimum site radius (Å). D_0 , r_0 and *E* were calculated using the SIMPLE and DOUBLE FIT program (Dalou et al., 2018). Values in italic are the lattice strain parameters $r_0^{2+,+3}$ and $E^{2+,+3}$ for the M2 site calculated by the relations of Wood and Blundy (1997), and the average r_0^{4+} and E^{4+} values obtained for the M1 site of clinopyroxene by Hill et al. (2011).























Supplementary Material 1

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