

CO 2 Solubility in Kimberlite melts CO 2 Solubility in Kimberlite melts

Yves Moussallam, Yann Morizet, Malcolm Massuyeau, Mickaël Laumonier,

Fabrice Gaillard

• To cite this version:

Yves Moussallam, Yann Morizet, Malcolm Massuyeau, Mickaël Laumonier, Fabrice Gaillard. CO 2 Solubility in Kimberlite melts CO 2 Solubility in Kimberlite melts. Chemical Geology, Elsevier, 2015, 418, pp.198-205. <10.1016/j.chemgeo.2014.11.017>. <i sub-01092334>

HAL Id: insu-01092334 https://hal-insu.archives-ouvertes.fr/insu-01092334

Submitted on 9 Dec 2014

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution - NonCommercial - NoDerivatives 4.0 International License

1 CO₂ Solubility in Kimberlite melts

2

3 <u>Yves Moussallam¹, Yann Morizet^{1,2}, Malcolm massuyeau¹, Mickael Laumonier^{1,3}, Fabrice</u>

- 4 <u>Gaillard¹</u>
- 5

6 ¹ISTO, 7327 Université d'Orléans-CNRS-BRGM, 1A rue de la Férollerie, 45071 Orléans cedex 2, France

7 ² Université de Nantes, Nantes Atlantique Universités, Laboratoire de Planétologie et Géodynamique de Nantes

8 (LPGN) UMR CNRS 6112

³Bayerisches Geoinstitut, University of Bayreuth, 95440 Bayreuth, Germany

10 Corresponding author: Yves Moussallam: <u>yves.moussallam@cnrs-orleans.fr</u>

11 ABSTRACT

12 Carbon dioxide is the most abundant volatile in kimberlite melts and its solubility exerts a 13 prime influence on the melt structure, buoyancy, transport rate and hence eruption dynamics. 14 The actual primary composition of kimberlite magma is the matter of some debate but the 15 solubility of CO₂ in kimberlitic melts is also poorly constrained due to difficulties in 16 quenching these compositions to a glass that retains the equilibrium CO₂ content. In this study we used a range of synthetic, melt compositions with broadly kimberlitic to carbonatitic 17 18 characteristics which can, under certain conditions, be quenched fast enough to produce a glass. These materials are used to determine the CO₂ solubility as a function of chemical 19 20 composition and pressure (0.05-1.5 GPa). Our results suggest that the solubility of CO_2 21 decreases steadily with increasing amount of network forming cations from ~30 wt% CO₂ at 12 wt% SiO₂ down to ~3 wt% CO₂ at 40 wt% SiO₂. For low silica melts, CO₂ solubility 22 23 correlates non-linearly with pressure showing a sudden increase from 0.1 to 100 MPa and a 24 smooth increase for pressure >100 MPa. This peculiar pressure-solubility relationship in low 25 silica melts implies that CO₂ degassing must mostly occur within the last 3 km of ascent to

the surface having potential links with the highly explosive nature of kimberlite magmas and
some of the geo-morphological features of their root zone. We present an empirical CO₂
solubility model covering a large range of melt composition from 11 to 55 wt% SiO₂
spanning the transition from carbonatitic to kimberlitic at pressures from 1500 to 50 MPa.
Keywords: kimberlite; carbonatite; CO₂ solubility; transitional melt

31

1. INTRODUCTION

Experimental investigation of the solubility of CO₂ in kimberlite melt has been limited due to 32 33 the difficulty in quenching such melt fast enough to form a pure glass (e.g. Brey and 34 Ryabchikov, 1994). Previous studies have therefore estimated the CO₂ solubility based on: (i) 35 extrapolation from trend defined in silicate melts (Brooker et al., 2001) (ii) solubility of melt in partially to highly crystallised experiments (Brooker et al., 2011) (iii) solubility at 0.1 MPa 36 37 of simple, four components, synthetic compositions considered as analogue to the natural system (Russell et al., 2012) (iv) molecular dynamics simulation studies (Guillot and Sator, 38 2011). While these studies all suggest that the solubility of CO_2 decreases as the melt 39 becomes enriched in silica, the effect of pressure has remained speculative. Our current 40 41 knowledge from experimental studies on silicate melt suggests that CO₂ solubility increases 42 with pressure (Blank and Brooker, 1994; Iacono-Marziano et al., 2012; Morizet et al., 2002, 43 2014) while in pure carbonate melts, CO₂ solubility is more related to satisfying stoichiometric requirements and therefore less dependent on pressure as exemplified by 44 Oldoinyo Lengai's alkali carbonatite lava (Kervyn et al., 2008). However, experiments from 45 Wyllie (1989) and Brooker & Kjardgaard (2011) suggest that there is some 'excess' CO₂ 46 solubility in pure CaCO₃ melt, apparently causing a decrease in liquidus temperature. 47

The composition of primary kimberlite melts in term of volatile and major element chemistryis poorly constrained. Not only are natural kimberlite laden with xenolith and xenocryst

Moussallam et al.

50 (Mitchell, 2008), but the strong alteration by external fluids they typically endured, in particular serpentinization, strongly affects the volatile and major element composition 51 (Brooker et al., 2011; Sparks et al., 2009). Latest attempt to back-calculate the kimberlite 52 53 melts chemistry by removal of the serpentinization process points towards an original melt 54 being richer in CO₂ and CaO and poorer in SiO₂, water and MgO than calculated from "uncontaminated" kimberlites (Brooker et al., 2011; Sparks et al., 2009). This findings are 55 corroborated with the fact that the Mg number in kimberlite is too high for equilibrium with a 56 57 mantle sources (Kopylova et al., 2007; Price et al., 2000) and that liquidus temperature for "uncontaminated" kimberlite composition of 1400°C (Brooker et al., 2011) are higher than 58 59 predicted from mineral thermometry (Fedortchouk et al., 2002). We therefore chose here to 60 follow from the work of Brooker et al (2011) who argued that kimberlite melts are originally 61 transitional, that is, they have a composition intermediate between a carbonate and a silicate liquid, and investigated several melt compositions with SiO₂ contents between 11 and 32 62 wt%. 63

64

In this contribution, we present results on the first experimental kimberlitic glasses synthesised using an internally heated pressure vessel (IHPV) equipped with an optimised quench configuration. We produced a series of melt compositions across the "transitional" melt field (SiO₂ ~18 to ~32 wt%) to assess the effect of silica activity on CO₂ solubility. The effect of pressure was then investigated by equilibrating melts of similar composition at different pressures (50 to 1500 MPa).

Moussallam et al.

72

2. EXPERIMENTAL METHODOLOGY

73 2.1 Starting material

74 Starting materials were produced by mixing powders from a natural lamproite from Torre Alfina, Italy (table 1, see Peccerillo et al., 1988) with various amounts of synthetic powders 75 76 of pure oxides and natural dolomite. In order to ensure homogeneity and remove the volatiles 77 present in the Torre Alfina rock, it was fused twice in air at 1400°C and quenched to glass. 78 The composition of all mixtures used as starting materials of all experiments is reported in 79 table 1. The melt compositions we have synthesized here are very Ca-rich when compared to 80 the more Mg-rich typical "contamination free" kimberlite (Mitchell, 1986) but later compare 81 our findings to experiments with more Mg-rich composition. Therefore, an underlying 82 supposition of our work is that Mg and Ca cations behave in a similar way with respect to CO₂ solubility in the melt (see Brey et al., 1991 and Brey and Ryabchikov, 1994 for an 83 investigation of the effect of Mg-Ca substitution on CO₂ solubility in kimberlite melts and see 84 85 discussion section 5). The Al content of the composition presented here might also be considered slightly high by some authors while the Fe content would be considered low. 86 Notwithstanding these differences, the compositions we explored here provide a close 87 88 analogue to kimberlite melts (in term of NBO/T for instance) for which the true composition 89 remains unknown and debated (see review by Sparks 2013).

90

91 2.2 High pressure experiments; internally heated pressure vessels and piston 92 cylinder

To investigate the solubility of CO_2 in transitional melts at crustal pressure, we performed a series of solubility experiments in the pressure range 50 to 350 MPa at relatively constant temperature (1225 to 1270°C) in relatively dry and oxidized conditions (log fO_2 = FMQ+3). We used internally heated pressure vessels at the ISTO-CNRS laboratory in Orléans, which can reach pressures of up to 400 MPa (\pm 3 MPa) under controlled temperature up to 1300°C (\pm 2°C). The vessel was pressurised using argon gas as the pressure medium. A two-Mowinding vertical furnace was used, creating a 5cm isothermal (\pm 5°C) "hot-spot" zone and controlled by two S-type thermocouples located on both side of the sample.

101 Experimental charges consisted of anhydrous sample powder (30 to 150 mg) loaded in gold-102 palladium (Au₈₀Pd₂₀) or platinum capsules (2 to 4 cm in length, 2.5 mm inner diameter and 103 2.9 mm outer diameter). The capsules were welded shut. Although the initial powder 104 mixtures were stored at 120°C, the strongly hydroscopic character of some oxides (in 105 particular MgO, NaO and K₂O) absorbing atmospheric H₂O within the 15 min necessary to 106 load the capsules, made it impossible to obtain fully water-free experimental charges, hence lowering slightly the XCO₂ in the fluid in equilibrium with the melt at P & T. For each 107 108 experiment, one to three capsules were hanged by a thin Pt wire. The temperature gradient 109 along the "hot-spot" zone, where the capsules were located, was maintained at about 10°C; the hot zone was always on the top end of the sample. This gradient, although inducing an 110 111 uncertainty in the run temperature, was part of the quench optimisation strategy as it imposed 112 a thermal profile in the furnace that ensures a strong thermal contrast over a small vertical 113 length located directly below the hanged capsule. Rapid quenching was achieved by passing 114 an electrical current to the holding Pt wire so that the sample dropped into the cold, bottom part of the furnace. The cooling rate was estimated to $>100-200^{\circ}C \text{ s}^{-1}$ although no precise 115 116 estimate could be obtained. After each experiment, capsules were weighed (i) before and (ii) 117 after, opening in order to (i) verify that no leakage had occurred, and (ii) determine the 118 amount of gas not incorporated in the glass. Capsules were then opened and a fragment of the 119 charge was embedded in an epoxy resin and polished for SEM and microprobe analyses. We 120 conducted a total of 18 solubility experiments. The strategy was first to produce a complete

121 isobaric section at 350 MPa, for a range of melt chemistry spanning SiO₂ from 11 to 38 wt% 122 (expressed as nominal starting composition, Table 1). We then explored the effect of pressure on several compositions (between 17 and 38 wt% SiO₂; expressed as nominal starting 123 124 composition) from 50 to 350 MPa at temperatures of 1225 to 1270°C. While most 125 experiments successfully produced a pure glass (Fig. 1 and Fig.S1), experiments performed at 126 50 MPa (run number: TA6_1_6 and TA7_1_3), one experiment at 100 MPa (run number: TA10_1_2) and the experiment with lowest SiO_2 (run number TA15_1_1) could not be 127 128 quenched fast enough and quench crystals were present throughout the experimental charge 129 (these crystals could not be analysed free of the glass component).

130

131 A single piston cylinder experiment was performed at 1.5 GPa and 1300°C in a ¹/₂ inch 132 piston-cylinder apparatus. Experimental charge consisted of natural anhydrous sample powder (30 mg) loaded in gold-palladium (Au₈₀Pd₂₀) capsules (1 cm in length, 2.5 mm inner 133 diameter and 2.9 mm outer diameter). The capsule was introduced in a talc-pyrex-graphite 134 135 furnace assembly and surrounded by MgO. A B-type thermocouple was located at ~1 mm 136 atop of the capsule and the run temperature should be considered as a minimum value. The quench of the piston cylinder assembly was not fast enough to prevent the pervasive growth 137 138 of quench crystals (Fig. 1; Fig. S1; experiment TA6_1_5).

139 **3. ANALYTICAL TECHNIQUES**

140 **3.1 EMPA**

All experimental products were examined by optical microscope and scanning electronic microscope (SEM) to check for the presence of quench crystals (Fig. S1). Electron microprobe analyses (EMPA) were performed on a Cameca SXFive at the ISTO-CNRS laboratory in Orléans. Glasses were analysed using an accelerating voltage of 15 kV, a beam 145 current of 6 nA and a defocused beam of 10 μ m. Na and K were analysed first in order to 146 minimise alkali loss during analysis. The shortfall in the totals for analysed glasses was 147 broadly consistent with the CO₂ content estimated by other techniques, but systematically 148 higher (Fig. S2).

149

150 **3.2 Micro-Raman**

151 We used an Innova 300-5W Argon ion laser (Coherent©) operating at 514 nm as the light 152 source to produce Raman scattering. Spectra were collected by a Jobin-Yvon Labram 153 spectrometer (focal distance = 300 mm) equipped with a 2400 grooves/mm CCD detector. 154 Analyses were performed in confocal mode (hole= $500 \mu m$, slit = $200\mu m$) and using a x50 Olympus objective reducing the analysed volume size to a few um³. Spectra were acquired 155 over the aluminosilicate network and CO_3^{2-} region between 200 and 1350 cm⁻¹. In a few cases 156 spectra were also acquired in the OH region between 2500 and 3900 cm⁻¹. The acquisition 157 158 time was of 10 x 60 sec and acquisition depth was optimised in order to obtain the highest 159 Raman signal (Mercier et al., 2009). At least three spectra were acquired for each sample. 160 The CO₂ content in the glass was quantify using the calibration method of Morizet et al. (2013), which provided a reproducibility on the CO_2 measurements of ~5% relative, 161 corresponding to an estimated absolute error on the CO₂ measurements less than 1 wt% for 162 the investigated samples. 163

164

165 3.3 Elemental Analyser

We used a Thermo Scientific, Flash 2000, CHNS elemental analyser to measure water (as H) and CO₂ (as C) in all experimental products. About 1 mg of sample was loaded together with 1 mg of vanadium pentoxide in a tin capsule folded and analysed by combustion at 1800°C. The analyser was calibrated directly prior to analysis and reproducibility on external 170 standards (marble, dolomite and hydrated basalt) was found to be better than 2% for C and 171 10% for H, giving an estimated error on the CO_2 measurement of < 1 wt%.

172

173 **3.4 Gravimetric Weight loss**

174 After experiments the capsules were weighed using a high precision balance before and after 175 puncture, to measure the mass of CO_2 (\pm H₂O) in the excess fluid phase from which, the mass of dissolved CO₂ in the glass can be calculated by mass-balance. The error associated with 176 177 this technique depends largely on the initial amount of material loaded in the capsule. 178 Considering a maximum error on the weight measurements of 0.5 mg, for a minimum amount 179 of loaded material of 30 mg, the error on the subsequent CO_2 estimate by this method is < 2180 wt%. There may also be an under-estimation of fluid phase mass if some is retained in 181 enclosed bubbles. However, most vesicles are found at the capsule wall (see Fig 1); as a 182 result, most of the fluid must be released on piercing.

4. RESULTS

184 All experimental charges were CO₂ saturated. This was attested by the presence of bubbles in 185 all investigated charges (Fig.1) and by weight loss after piercing the capsules. All results from solubility experiments are reported in Table 2 with CO₂ content estimated from 186 elemental analyser, weight loss (after capsule puncture), EMPA and micro-Raman 187 188 spectroscopy. A comparison of the four different methods used to measure the CO₂ content of 189 the experimental charges is presented in Fig. S2. This figure shows that the agreement 190 between the Raman, elemental analyser and gravimetric weight loss methods is very good. 191 Estimates from EMPA shortfall are consistently overestimating the amount of CO₂ even after 192 correction for H₂O (determined by elemental analyser), suggesting that using EMPA shortfall 193 to estimate the melt CO₂ content might be inaccurate. Unless stated otherwise, all CO₂

contents reported in the following figures are from measurements using the elemental 194 analyser. The results of CO₂ solubility experiments at 350 MPa for a range of compositions 195 196 are reported in Fig. 2 together with a data point from an earlier study using a similar composition (Iacono-Marziano et al., 2012). Fig.2 clearly shows a decrease in CO₂ solubility 197 with increasing amount of network forming cations (Si^{4+} and Al^{3+}). This decrease appears to 198 199 be smooth and continuous across the transitional field. The effect of pressure, that is 200 nominally the pressure of CO₂, on solubility is shown in Fig. 3 for starting compositions at 18 201 to 41 wt% SiO₂ (SiO₂ + Al₂O₃ of 21 to 51 wt% respectively, expressed with volatiles at 202 saturation at 350 MPa). Since experiments at 50 MPa could not be quenched fast enough to 203 prevent the growth of quench crystals, CO₂ solubility was estimated from the amount of gas 204 released after capsule puncture and their CO₂ contents are therefore associated with a larger 205 error (estimated at $\pm 2 \text{ wt\%}$) than those determined by elemental analyser (estimated at < 1wt%). At 18 wt% SiO₂ (at the edge of the carbonate field) a pressure drop from 1500 to 180 206 207 MPa seems to have a limited effect on the solubility of CO₂ (25 to 20 wt% CO₂) and most of the solubility drop takes place from 100 to 0.1 MPa (assuming no CO₂ remains in the melt at 208 209 0.1 MPa). This results in a strongly non-linear dependence of CO₂ solubility with pressure for 210 melt compositions with SiO₂ contents in the range 18-30 wt%. At SiO₂ content higher than 30 211 wt% the solubility dependence on pressure (from 350 to 0.1 MPa) is linear, as expected in 212 silicate melts for the pressure range considered.

213

5. DISCUSSION

214 Evidence for a genetic connection between kimberlite and carbonatite magmas goes beyond 215 their, long noticed, spatial and temporal occurrence (e.g. Janse, 1975; Mitchell, 2005; White et al., 1995). Kimberlite and carbonatite melts can both be produced by very low degree 216 partial melting of a carbonated mantle source as inferred from their high concentration of 217

Moussallam et al.

incompatible trace elements (Nelson et al., 1988) and from high pressure melting experiments 218 219 of carbonated peridotite phase assemblages (Dalton and Presnall, 1998; Dasgupta and 220 Hirschmann, 2006; Ghosh et al., 2009; Gudfinnsson and Presnall, 2005; Rohrbach and 221 Schmidt, 2011). In addition kimberlite and carbonatite melts have been argued to be related 222 by fractionation processes (Dalton and Presnall, 1998; Dawson and Hawthorne, 1973; Larsen and Rex, 1992). This clear link between the two types of magma supports the hypothesis of 223 224 Brooker et al. (2011) that kimberlite primary magmas were originally more transitional or 225 carbonatitic in composition. The set of transitional synthetic super-liquidus melt 226 compositions that have been produced here should therefore yield a robust approximation of 227 the solubility law for CO₂ in primary kimberlitic melts.

228

229 **5.1 Empirical model of CO₂ solubility in carbonated melts**

Based on the solubility experiments presented here and literature data we developed a purely empirical model of CO_2 solubility as a function of the amount of network forming cations (SiO₂ + Al₂O₃) and pressure. We emphasised that it is the first modelling effort addressing such low silica melt compositions. The simple model is defined as a third order polynomial linking the amount of CO_2 dissolved in the melt to the SiO₂ + Al₂O₃ content and in which each coefficient is in turn a function of pressure as follow:

236
$$CO_{2_{Wt\%}} = A \times (SiO_2 + Al_2O_3)^3 + B \times (SiO_2 + Al_2O_3)^2 + C \times (SiO_2 + Al_2O_3) + D$$
 (1)

Where:

 $A = a \times P^b$

- $B=c\times P^d$
- $C = e \times P^f$

With *D*, *a*, *b*, *c*, *d*, *e* and *f* representing adjusted parameters reported in Table 3. The rationale behind the formulation of equation (1) is to simulate a pressure-dependent process of dilution

from a pure carbonate end-member (D=50) with the addition of network-forming cations 240 $(SiO_2 \text{ and } Al_2O_3)$. In order to adjust the parameters in equation (1), we used CO₂ solubility 241 data for basalt (Mattey, 1991; Pawley et al., 1992; Shishkina et al., 2010; Stolper and 242 Holloway, 1988), haplo-phonolite (Morizet et al., 2002), phonotephrite (Behrens et al., 2009), 243 244 melilitite (Brooker et al., 2001) in addition to selected data from Brey and Ryabchikov, 245 (1994); Brooker, (1995); Brooker et al., (2011); Brooker and Kjarsgaard, (2011); Iacono-Marziano et al., (2012); Morizet et al., (2013); Morizet et al., (2014) and this study (all shown 246 247 in the subsequent figures). Equation (1) reproduces our experimental data well (standard deviation of 0.9 wt% CO₂ between the modelled and measured values) and allow us to 248 249 explore a larger P- $X_{(SiO2 + Al2O3)}$ space. Fig. 4 shows the modelled solubility of CO₂ as a 250 function of pressure for compositions ranging from $SiO_2 + Al_2O_3$ of 0 to 51 wt% and pressure from 1500 to 10 MPa. Fig.4 shows the gradual change in the CO₂ solubility behaviour from 251 carbonate to silicate end-member melts as a function of pressure. We note that from Fig.4, a 252 253 typical transitional melt enters the kimberlite "root zone" (see the classical model from 254 Hawthorne, 1975) with a CO₂ content of 10 to 18 wt%, consistent with the "target value" of 255 15 wt% determined by Brooker et al. (2011) for kimberlites.

256

257 The range of experiments presented here is not extensive enough to incorporate the effect of 258 temperature and alkali content in equation (1). For this, we considered the available from the 259 literature. Fig. 5 compares data from a range of studies and shows the evolution of CO₂ 260 solubility as a function of network forming cations $(SiO_2+Al_2O_3)$ for a large range of composition and pressure from 2000 to 50 MPa. The predicted evolution from equation (1) is 261 also shown for pressure between 2000 and 20 MPa. The melt composition of all data shown 262 in Fig 5 is reported in Table S2, where one can notice the wide range of compositions 263 explored by these various studies. The variety of methods used to estimate the CO₂ content in 264

these studies (EMP shortfall, Raman, FTIR, elemental analyser and gas chromatography), 265 makes the comparison of dataset not ideal. In addition, while the present study, that of 266 Iacono-Marziano et al., (2012), Morizet et al., (2014), Brooker et al., (2001) and Brey and 267 Ryabchikov, (1994) are from super-liquidus experiments, data from Brooker et al., (2011) are 268 269 from patches of glass in mostly crystallised charges while data from Brooker and Kjarsgaard 270 (2011) and Brooker (1995) are from silicate melts coexisting with an immiscible carbonate 271 melt. Perhaps most importantly, the CO₂/H₂O ratio varies widely over the reported dataset 272 with volatile component in experiments from Iacono-Marziano et al., (2012) and Morizet et al., (2014) being dominated by H₂O. Keeping in mind these limitations in the comparison, we 273 274 can see that the set of data from Morizet et al. (2014) at 350 and 100 MPa and Morizet et al. 275 (2013) at 1000 MPa are well reproduced by our model. A similar agreement is noted with the 276 data at 2000 MPa from Brooker et al., (2001), at 200 MPa from Brooker et al (2011) and (to a lesser extent) at 1500 MPa from Brooker (1995). We also note that data at 1000 MPa from 277 278 Brey and Ryabchikov (1994) (we only report experimental data at 1000 MPa where they reported clear glass) are also well reproduced despite the significant difference in terms of 279 280 chemical composition (MgO up to 27wt%). Fig 6, shows a plot of experimentally determined 281 vs. calculated CO₂ solubility from which it appears that the highest deviation from the model 282 are for data by Brooker et al (2011) and Brooker and Kjarsgaard (2010). We made several 283 attempt to link the deviation from the modelled value to the melt composition (Na₂O, K₂O, 284 Na_2O+K_2O , SiO_2/Al_2O_3) but could not find any robust correlations. There is however a hint 285 that melt compositions rich in alkali can incorporate more CO₂ than comparatively Ca and Mg-rich compositions (see in Fig. 5 the isolated experiment from Brooker (1995) at 1500 286 MPa with 15.6 wt% CO₂ and a composition of 0 wt% CaO and 37 wt% Na₂O). As additional 287 experimental data become available, future models should be able to explore the full effect of 288 composition (in term of Mg, Ca, Na, K, Fe cations) on CO₂ solubility. 289

Moussallam et al.

290 A strong correlation was found between the method used to determine CO₂ content and deviation from our model value. Indeed, 70% of the data located further than two standard 291 292 deviations away from their calculated value had CO₂ content determined by shortfall of 293 EMPA analyses. Comparing our own EMPA shortfall measurements to CO₂ determined by 294 bulk analyser (Fig. S2), we observe that determining the CO₂ content by EMPA shortfall can 295 lead to an overestimation of 2 to 6 wt% CO₂. We therefore conclude (i) that most of the literature data yielding CO₂ solubility values that deviates from our model can be attributed to 296 297 the large error associated with estimating CO₂ content from EMPA shortfall and (ii) that from 298 the current, limited, dataset there is no identifiable systematic deviations related to the melt 299 composition. Equation (1) therefore provides a good first order approximation of the CO_2 300 content of carbonated melt over a large range of composition (SiO₂ < 55 wt%; Na₂O+K₂O <301 22 wt%). The entrapment pressure of recently reported CO₂-rich melt inclusions from 302 Oldoinyo Lengai (de Moor et al., 2013) for instance, can be estimated using equation (1) 303 (yielding an entrapment depth of 265 to 2085 MPa for inclusions containing 3 to 8 wt% 304 CO_2).

305

306 **5.2 Implications for Kimberlite ascent and volcanism**

307 As hypothesised by Brooker et al (2011) our results indicate that CO₂ solubility decreases with increasing amount of network forming cations over the carbonate-silicate transitional 308 309 field. This finding supports the conclusions of Russell et al., (2012) that an ascending 310 carbonate melt, assimilating mantle orthopyroxene (OPX) and becoming progressively 311 enriched in silica, will release an increasing amount of CO₂ into the gas phase and promote 312 magma ascent by keeping it buoyant in spite of increasing xenocryst loading. The effect of 313 pressure (shown in Fig. 3 and 4) on transitional melts appears to be a function of their composition; melts with a composition closer to the carbonate end-member retain a large 314

315 quantity of CO₂ until very shallow (3 km) depth while slightly more silicate-rich melts show 316 a continuous and progressive degassing profile with CO₂ being equally released from mantle 317 depth to the surface. The solubility-pressure relationships shown by the transitional melt 318 compositions in Fig. 3 and 4 are intermediate between the behaviour of a carbonate melt 319 which will show little pressure dependence on solubility, and that of a typical silicate melt 320 which will show a mostly linear relationship in the pressure range considered (Fig. 3, 4,e.g. 321 Papale et al., 2006; Iacono-Marziano et al 2012).

322

The behaviour of transitional melts, retaining more than 18 wt% CO₂ up until very shallow 323 324 depth should lead to a rapid increase in ascent velocity while reaching the last 3 km depth 325 where important quantities of CO_2 will be released (Fig. 4). This acceleration will be 326 exacerbated by the fact that the volume fraction of gas will reach values higher than 77%, corresponding to the point at which the bubble foam will collapse (Woods, 1995) changing 327 the continuous phase from a bubbly liquid magma to an ash-laden gas and therefore 328 329 considerably increasing the flow speed as the frictional force become much smaller and the 330 viscosity becomes closer to that of the gas phase. This ascent scenario can readily explain 331 some of the typical morphological features of kimberlite pipes. (i) The widening upward 332 geometry of the pipe, typical of eruptions happening at depth, can be explained by the high 333 amount of CO₂ released over a, most probably, very small time increment leading to 334 overpressure and explosion below the surface. (ii) The appearance of breccias and irregularly 335 shaped blocks at depth of ~3 km, in the "root zone" (see the classical model from Hawthorne, 336 1975) is consistent with large amount of CO₂ starting to be released at this depth and the associated rapid volumetric expansion (Fig. 4). The occurrence of hypabyssal intrusions of 337 338 kimberlites however suggests that some kimberlite stall during their ascent through the crust. 339 Although we cannot confidently assign a reason for this, possibilities include the loss of volatiles, segregating from the magma during ascent or the progressive cooling of the magma
(Kavanagh and Sparks, 2009), both process inducing crystallisation and greatly increasing
magma viscosity. Experiments on these compositions suggest that crystallisation is extremely
rapid, as illustrated by the speed at which quench crystals formed in some of our experiments.

While CO_2 is the most abundant volatile in kimberlite magmas, an unknown but probably significant amount of water is also dissolved in these melts (up to 10 wt%; Kopylova et al., 2007; also see discussion in Sparks et al., 2009 and experimental investigation by Keppler, 2003). Water degassing is therefore likely to exert a large influence on the ascent dynamics of Kimberlite melts. It remains unclear however whether this process would enhance, limit or displace the depth at which the CO_2 -exsolution-triggered acceleration of kimberlite melts that we predict takes place.

352

353 **6. CONCLUSIONS**

354 We have determined CO₂ solubility laws for kimberlite melt and have also provided a general solubility model for carbonated melt composition (SiO₂ < 55 wt%) at pressure between 50 355 and 1500 MPa. We found that (i) CO₂ solubility decreases steadily with increasing amount of 356 network forming cations (ii) The effect of pressure on CO₂ solubility is a strong function of 357 composition with low SiO₂ melts retaining large amount (~15 wt%) of CO₂ up to very 358 shallow pressure (~100 MPa) while more silicated melts have a near linear CO₂ solubility 359 360 dependence on pressure. We propose that this peculiar pressure-solubility dependency should 361 result in a strong acceleration of an ascending kimberlite magma in the last 3 km of crust 362 where CO₂ starts to exsolve in large quantities. This behaviour can explain morphological features of kimberlite pipes, notably the depth extent of the root zone and the widening-363

upward shape of the conduit and can account for the highly explosive character of Kimberliteeruptions.

366

367 ACKNOWLEDGEMENTS

368 We are very grateful to Rémi Champallier and Leïla Hashim for their help with various

369 instruments. Special thanks also go to Ida Di Carlo for help with the electron probe analyses

and to Marielle Hatton for help with the elemental analyser. Funding for this work was

371 provided by the European Research Council (ERC grant number 279790) and the Agence

372 Nationale de la Recherche (ANR-10-BLAN-62101). Valuable reviews by K. Russell and R.

- 373 Brooker improved the quality of the manuscript. We also thanks the guest Editors of this
- 374 special issue.

375 **REFERENCES**

- Behrens, H., Misiti, V., Freda, C., Vetere, F., Botcharnikov, R.E., Scarlato, P., 2009.
 Solubility of H (sub 2) O and CO (sub 2) in ultrapotassic melts at 1200 and 1250
 degrees C and pressure from 50 to 500 MPa. Am. Mineral. 94, 105–120.
 doi:10.2138/am.2009.2796
- Blank, J.G., Brooker, R.A., 1994. Experimental studies of carbon dioxide in silicate melts;
 solubility, speciation, and stable carbon isotope behavior. Rev. Mineral. Geochem. 30,
 157–186.
- Brey, G.P., Kogarko, L., Ryabchikov, I., 1991. Carbon dioxide in kimberlitic melts. N Jb
 Min. Mh H4, 159–168.
- Brey, G., Ryabchikov, I., 1994. Carbon dioxide in strongly undersaturated melts and origin
 of kimberlitic magmas. N Jb Min. Mh H10, 449–463.
- Brooker, R.A., 1995. Carbonatite genesis; the role of liquid immiscibility to 25 kb (PhD thesis). University of Manchester.
- Brooker, R.A., Kjarsgaard, B.A., 2011. Silicate–Carbonate Liquid Immiscibility and Phase
 Relations in the System SiO2–Na2O–Al2O3–CaO–CO2 at 0·1–2·5 GPa with
 Applications to Carbonatite Genesis. J. Petrol. 52, 1281–1305.
 doi:10.1093/petrology/egq081
- Brooker, R.A., Sparks, R.S.J., Kavanagh, J.L., Field, M., 2011. The volatile content of
 hypabyssal kimberlite magmas: some constraints from experiments on natural rock
 compositions. Bull. Volcanol. 73, 959–981. doi:10.1007/s00445-011-0523-7
- Brooker, R.., Kohn, S.., Holloway, J.., McMillan, P.., 2001. Structural controls on the
 solubility of CO2 in silicate melts: Part I: bulk solubility data. Chem. Geol. 174, 225–
 239. doi:10.1016/S0009-2541(00)00353-3

- 399 Dalton, J.A., Presnall, D.C., 1998. The Continuum of Primary Carbonatitic–Kimberlitic Melt
 400 Compositions in Equilibrium with Lherzolite: Data from the System CaO–MgO–
 401 Al2O3–SiO2–CO2 at 6 GPa. J. Petrol. 39, 1953–1964. doi:10.1093/petroj/39.11402 12.1953
- Dasgupta, R., Hirschmann, M.M., 2006. Melting in the Earth's deep upper mantle caused by
 carbon dioxide. Nature 440, 659–662. doi:10.1038/nature04612
- 405 Dawson, J.B., Hawthorne, J.B., 1973. Magmatic sedimentation and carbonatitic
 406 differentiation in kimberlite sills at Benfontein, South Africa. J. Geol. Soc. 129, 61–
 407 85. doi:10.1144/gsjgs.129.1.0061
- De Moor, J.M., Fischer, T.P., King, P.L., Botcharnikov, R.E., Hervig, R.L., Hilton, D.R., 408 409 Barry, P.H., Mangasini, F., Ramirez, C., 2013. Volatile-rich silicate melts from Oldoinyo Lengai volcano (Tanzania): Implications for carbonatite genesis and 410 Planet. 411 eruptive behavior. Earth Sci. Lett. 361. 379-390. 412 doi:10.1016/i.epsl.2012.11.006
- Fedortchouk, Y., Canil, D., Carlson, J.A., 2002. Intensive Variables in Primary Kimberlite
 Magmas (Lac de Gras, N.W.T., Canada) and Application for Diamond Preservation.
 AGU Fall Meet. Abstr. -1, 1348.
- Ghosh, S., Ohtani, E., Litasov, K.D., Terasaki, H., 2009. Solidus of carbonated peridotite
 from 10 to 20 GPa and origin of magnesiocarbonatite melt in the Earth's deep mantle.
 Chem. Geol., Volatiles and Volatile-Bearing Melts in the Earth's Interior 262, 17–28.
 doi:10.1016/j.chemgeo.2008.12.030
- Gudfinnsson, G.H., Presnall, D.C., 2005. Continuous Gradations among Primary
 Carbonatitic, Kimberlitic, Melilititic, Basaltic, Picritic, and Komatiitic Melts in
 Equilibrium with Garnet Lherzolite at 3–8 GPa. J. Petrol. 46, 1645–1659.
 doi:10.1093/petrology/egi029
- Guillot, B., Sator, N., 2011. Carbon dioxide in silicate melts: A molecular dynamics
 simulation study. Geochim. Cosmochim. Acta 75, 1829–1857.
 doi:10.1016/j.gca.2011.01.004
- 427 Hawthorne, J.B., 1975. Model of a kimberlite pipe. Phys. Chem. Earth 9, 1–15.
 428 doi:10.1016/0079-1946(75)90002-6
- Iacono-Marziano, G., Morizet, Y., Le Trong, E., Gaillard, F., 2012. New experimental data and semi-empirical parameterization of H2O–CO2 solubility in mafic melts.
 Geochim. Cosmochim. Acta 97, 1–23. doi:10.1016/j.gca.2012.08.035
- Janse, A.J.A., 1975. Kimberlite and related rocks from the Nama Plateau of South-West
 Africa. Phys. Chem. Earth 9, 81–94. doi:10.1016/0079-1946(75)90009-9
- Kavanagh, J.L., Sparks, R.S.J., 2009. Temperature changes in ascending kimberlite magma.
 Earth Planet. Sci. Lett. 286, 404–413. doi:10.1016/j.epsl.2009.07.011
- 436 Keppler, H., 2003. Water solubility in carbonatite melts. Am. Mineral. 88, 1822–1824.
- Kervyn, M., Ernst, G.G.J., Klaudius, J., Keller, J., Kervyn, F., Mattsson, H.B., Belton, F.,
 Mbede, E., Jacobs, P., 2008. Voluminous lava flows at Oldoinyo Lengai in 2006:
 chronology of events and insights into the shallow magmatic system. Bull. Volcanol.
 70, 1069–1086. doi:10.1007/s00445-007-0190-x
- Kopylova, M.G., Matveev, S., Raudsepp, M., 2007. Searching for parental kimberlite melt.
 Geochim. Cosmochim. Acta 71, 3616–3629. doi:10.1016/j.gca.2007.05.009
- Larsen, L.M., Rex, D.C., 1992. A review of the 2500 Ma span of alkaline-ultramafic, potassic
 and carbonatitic magmatism in West Greenland. Lithos 28, 367–402.
 doi:10.1016/0024-4937(92)90015-Q
- Mattey, D.P., 1991. Carbon dioxide solubility and carbon isotope fractionation in basaltic
 melt. Geochim. Cosmochim. Acta 55, 3467–3473. doi:10.1016/0016-7037(91)905083

- Mercier, M., Di Muro, A., Giordano, D., Métrich, N., Lesne, P., Pichavant, M., Scaillet, B.,
 Clocchiatti, R., Montagnac, G., 2009. Influence of glass polymerisation and oxidation
 on micro-Raman water analysis in alumino-silicate glasses. Geochim. Cosmochim.
 Acta 73, 197–217. doi:10.1016/j.gca.2008.09.030
- 453 Mitchell, R.H., 1986. kimberlites mineralogy geochemistry and petrology. Plenum, NY.
- Mitchell, R.H., 2005. Carbonatites and Carbonatites and Carbonatites. Can. Mineral. 43,
 2049–2068. doi:10.2113/gscanmin.43.6.2049
- Mitchell, R.H., 2008. Petrology of hypabyssal kimberlites: Relevance to primary magma
 compositions. J. Volcanol. Geotherm. Res. 174, 1–8.
 doi:10.1016/j.jvolgeores.2007.12.024
- Morizet, Y., Brooker, R.A., Iacono-Marziano, G., Kjarsgaard, B.A., 2013. Quantification of
 dissolved CO2 in silicate glasses using micro-Raman spectroscopy. Am. Mineral. 98,
 1788–1802. doi:10.2138/am.2013.4516
- Morizet, Y., Brooker, R.A., Kohn, S.C., 2002. CO2 in haplo-phonolite melt: solubility,
 speciation and carbonate complexation. Geochim. Cosmochim. Acta 66, 1809–1820.
 doi:10.1016/S0016-7037(01)00893-6
- Morizet, Y., Paris, M., Gaillard, F., Scaillet, B., 2014. Carbon dioxide in silica-undersaturated
 melt Part I: The effect of mixed alkalis (K and Na) on CO2 solubility and speciation.
 Geochim. Cosmochim. Acta.
- 468 Nelson, D.R., Chivas, A.R., Chappell, B.W., McCulloch, M.T., 1988. Geochemical and
 469 isotopic systematics in carbonatites and implications for the evolution of ocean-island
 470 sources. Geochim. Cosmochim. Acta 52, 1–17. doi:10.1016/0016-7037(88)90051-8
- 471 Papale, P., Moretti, R., Barbato, D., 2006. The compositional dependence of the saturation
 472 surface of H2O+CO2 fluids in silicate melts. Chem. Geol. 229, 78–95.
 473 doi:10.1016/j.chemgeo.2006.01.013
- Pawley, A.R., Holloway, J.R., McMillan, P.F., 1992. The effect of oxygen fugacity on the
 solubility of carbon-oxygen fluids in basaltic melt. Earth Planet. Sci. Lett. 110, 213–
 225. doi:10.1016/0012-821X(92)90049-2
- 477 Peccerillo, A., Poli, G., Serri, G., 1988. Petrogenesis of orenditic and kamafugitic rocks from
 478 central Italy. Can. Mineral. 26, 45–65.
- Price, S.E., Russell, J.K., Kopylova, M.G., 2000. Primitive Magma From the Jericho Pipe,
 N.W.T., Canada: Constraints on Primary Kimberlite Melt Chemistry. J. Petrol. 41,
 789–808. doi:10.1093/petrology/41.6.789
- 482 Rohrbach, A., Schmidt, M.W., 2011. Redox freezing and melting in the Earth/'s deep mantle
 483 resulting from carbon-iron redox coupling. Nature 472, 209–212.
 484 doi:10.1038/nature09899
- 485 Russell, J.K., Porritt, L.A., Lavallée, Y., Dingwell, D.B., 2012. Kimberlite ascent by
 486 assimilation-fuelled buoyancy. Nature 481, 352–356. doi:10.1038/nature10740
- Shishkina, T.A., Botcharnikov, R.E., Holtz, F., Almeev, R.R., Portnyagin, M.V., 2010.
 Solubility of H2O- and CO2-bearing fluids in tholeiitic basalts at pressures up to
 500 MPa. Chem. Geol. 277, 115–125. doi:10.1016/j.chemgeo.2010.07.014
- 490 Sparks, R.S.J., Brooker, R.A., Field, M., Kavanagh, J., Schumacher, J.C., Walter, M.J.,
 491 White, J., 2009. The nature of erupting kimberlite melts. Lithos 112, Supplement 1,
 492 429–438. doi:10.1016/j.lithos.2009.05.032
- 493 Stolper, E., Holloway, J.R., 1988. Experimental determination of the solubility of carbon
 494 dioxide in molten basalt at low pressure. Earth Planet. Sci. Lett. 87, 397–408.
 495 doi:10.1016/0012-821X(88)90004-0
- White, S.H., de Boorder, H., Smith, C.B., 1995. Structural controls of kimberlite and
 lamproite emplacement. J. Geochem. Explor. 53, 245–264. doi:10.1016/03756742(94)00033-8

Wyllie, P.J., 1989. Origin of carbonatites; evidence from phase equilibrium studies. Unwin
Hyman : London, United Kingdom, United Kingdom, pp. 500–545.

502 **TABLES**

503 **Table 1**: Starting oxide and natural rock powder mix compositions. The composition of a

natural lamproite from Torre Alfina, Italy, (Peccerillo et al., 1998), fused twice at 1400°C and

505 used to prepare the oxide-mix compositions is reported at the top of the table.

506

```
507 Table 2: Run conditions, EMPA analyses (averaged from 20), H<sub>2</sub>O determined by elemental
```

analyser (Flash) and CO_2 determined from: (i) EMPA (by subtracting H₂O content to EMPA

509 shortfall), (ii) Elemental analyser (Flash), (iii) Micro-Raman spectroscopy (following the

510 method of Morizet et al., (2013) and (iv) Weight loss measured by puncturing the capsule

(allowing the CO_2 to escape) after each run. Run duration was of 2 hours in all experiments.

512

Table 3: Adjusted parameters (and corresponding standard error on multiple non-linear
regression fit) to calculate CO₂ solubility from Eq. (1)

515

516 **FIGURES**

Figure 1: Optical microphotograph showing the results from various supra-liquidus experimental run producing a pure glass in all cases excepted for sample TA6_1_5 (lower right) in which quench crystals give a "milky" aspect to the otherwise glassy product. All samples shown are synthesised in IHPV apparatus excepted sample TA6_1_5 which was synthesised with piston cylinder. Corresponding SEM images are shown in Fig S1.

Figure 2: CO_2 solubility measurements for a series of compositions across the silicate to carbonate melt transitions equilibrated at 350 MPa. The SiO₂+Al₂O₃ content is reported including volatiles. For most experiments, CO_2 content is estimated from elemental analyser and associated error is estimated at ± 1 wt%. CO_2 content from Iacono-Marziano et al., (2012) is estimated from FTIR analyses.

528

Figure 3:CO₂ solubility as a function of pressure for three Kimberlite/Transitional and two silicate melt compositions spanning SiO₂ content of 18 to 41 wt% and SiO₂ + Al₂O₃ content of 21 to 51 wt% respectively (calculated including volatiles and corresponding to SiO₂ + Al₂O₃ contents of 27 to 54 wt% on a volatile-free basis). The light-brown to green colour part of the plot represents the upper 3 km of crust, corresponding to the typical "root zone" of Kimberlites. Dashed lines are fitted by eye through the data points.

535

Figure 4: Modelled evolution of CO_2 solubility as a function of pressure using equation (1) for melt compositions ranging from 0 to 41 wt% SiO₂ and 0 to 51 wt% SiO₂ + Al₂O₃. Experiments of corresponding chemistry are shown as squares of matching colour.

539

Figure 5: CO_2 solubility measurements from this study and the literature for compositions across the silicate to carbonate melt transition, equilibrated at 2000, 1500, 350, 200, 100 and 50 MPa. Literature data are from Brooker and Kjarsgaard, 2011; Brooker et al., 2011; Iacono-Marziano et al., 2012; Brooker et al., 2001; Brooker 1995; Morizet et al., 2013 Morizet et al., 2014 and Brey and Ryabchikov, 1994. Note that literature data are reported using the experiment total pressure and not the P_{CO2}. Model prediction values from equation (1) are shown for a range of pressure (2000 to 20 MPa) as dotted black curves.

Figure 6: Experimentally determined vs. calculated CO_2 solubility using equation (1). Doted red line represent the 1:1 line while continuous red lines on either sides represent the $\pm 2\sigma$ deviation.

551

552 SUPPLEMENTARY INFORMATION

TableS1: Standard deviation for EMP analyses reported in Table 2 (n=20).

554

555 **Table S2**: Run conditions, EMPA analyses and CO₂ determined from Raman or by bulk CO₂

556 LECO analyser for experiments from Brooker et al., (2011); Brooker and Kjarsgaard, (2011);

557 Iacono-Marziano et al., (2012); Brooker (1995) and Morizet et al., (2014).

558

Figure S1: Scanning electron microscopy (SEM) images of experimental charges shown in Fig. 1. All run produced a pure glass, excepted sample TA6_1_5 (lower right) in which quench crystals are pervasive.

562

Figure S2:Comparison between CO_2 content in experimental charges determined by (i) EMPA (by subtracting H₂O content to EMPA shortfall), (ii) Elemental analyser (Flash), (iii) Micro-Raman spectroscopy (following the method of Morizet et al., 2013), (iv) Weight loss measured by puncturing the capsule (allowing the CO_2 to escape) after each run and (v) EMPA shortfall without correction.

Name	SiO ₂	TiO ₂	AI_2O_3	FeO	MgO	CaO	Na ₂ O	K ₂ O	P_2O_5	CO ₂	Total
TA	55.74	1.34	13.05	5.83	9.43	5.45	0.96	7.68	0.51	0.00	100
TA15	11.01	0.09	2.34	0.41	8.45	36.36	0.27	2.14	0.04	38.90	100
TA6	16.60	0.20	3.85	0.86	6.93	35.48	0.28	2.27	0.07	33.47	100
TA7	20.08	0.25	4.66	1.08	8.16	32.50	0.34	2.74	0.10	30.09	100
TA8	21.66	0.30	5.03	1.30	8.21	31.30	0.37	2.96	0.11	28.75	100
TA9	23.11	0.34	5.37	1.49	8.26	30.20	0.40	3.16	0.13	27.53	100
TA10	23.79	0.36	5.53	1.58	8.29	29.68	0.41	3.26	0.14	26.95	100
TA11	30.18	0.56	7.04	2.43	8.52	24.84	0.52	4.14	0.21	21.56	100
TA12	37.70	0.79	8.81	3.43	8.78	19.14	0.65	5.18	0.30	15.22	100

Table 1: Starting oxide and natural rock powder mix compositions. The composition of a natural lamproite from Torre Alfina, Italy, (Peccerillo et al., 1998), fused twice at 1400°C and used to prepare the oxide-mix compositions is reported at the top of the table.

Experiment	Pressure (MPa)	Temperature (°C)	SiO ₂	TiO ₂	Al ₂ O ₃	FeO	MnO	MgO	CaO	Na₂O	K ₂ O	P_2O_5	Total	Shortfall	H₂O (Flash)	CO ₂ (EMPA)	CO ₂ (Flash)	CO ₂ (Raman)	CO ₂ (weight loss)
TA6_1_1	357.2	1225	18.44	0.2	2.05	0.74	0.03	6.87	45.25	0.3	2.3	0.04	76.22	23.8	1.54	22.24	21.84	21.4	24.4
TA6_1_4	178	1220-1260	17.97	0.19	2.63	0.81	0.04	6.75	44.8	0.41	2.19	0.04	75.83	24.2	1.19	22.98	20.07	23.4	18.9
TA7_1_1	340.2	1190	23.58	0.29	4.27	1.12	0.03	9.91	37.22	0.3	3.26	0.07	80.07	19.9	1.09	18.84	16.28	18.2	nd
TA9_1_1	336.4	1225	26.07	0.35	5.7	1.49	0.04	8.18	36.87	0.45	3.28	0.09	82.52	17.5	1.57	15.91	14.95	14.7	13.4
TA10_1_1	336.4	1225	26.77	0.35	5.8	1.64	0.05	8.52	36.24	0.4	3.39	0.09	83.25	16.8	1.73	15.02	13.80	14.4	12.0
TA11_1_1	347.8	1225	34.48	0.6	7.81	2.85	0.06	10	29.05	0.66	4.42	0.12	90.04	10.0	0.99	8.97	7.50	4.7	7.8
TA12_1_1	347.8	1225	40.97	0.82	9.74	3.68	0.06	9.38	22.96	0.82	5.58	0.19	94.19	5.8	0.59	5.22	2.79	3.5	3.7
TA6_1_5	1500	1300						Crys	stallise	d					1.36		24.85		nd
TA6_1_6	59.3	1225						Crys	stallise	d					0.84		15.87		12.7
TA7_1_2	179.2	1225	21.50	0.30	4.00	1.20	0.10	8.00	39.70	0.40	4.30	0.10	79.60	20.40	0.77	19.63	16.36	13.3	15.3
TA7_1_3	48.9	1215-1290						Crys	stallise	d					0.54		12.86		9.1
TA6_2_1	105.4	1270	17.80	0.20	2.40	0.70	0.10	8.10	45.20	0.20	2.50	0.10	77.20	22.80	0.79	22.01	16.47	16.3	18.3
TA7_1_4	100.7	1225	20.40	0.40	3.00	1.20	0.00	7.10	43.30	0.50	3.80	0.10	79.80	20.20	0.76	19.44	15.09	17.1	13.0
TA9_1_2	100.7	1225	24.50	0.50	4.40	1.90	0.00	8.10	39.80	0.50	4.20	0.10	84.20	15.80	0.59	15.22	10.12	10.8	11.0
TA15_1_1	351	1225						Crys	stallise	d					1.60		28.63		31.4
TA10_1_2	101	1225						Crys	stallise	d					0.71		9.00	10.18	8.59
TA11_1_2	101	1225	34.68	0.96	9.87	3.95	0.05	8.20	26.50	0.67	6.31	0.22	91.42	8.58	0.50	8.08	2.58	2.37	2.09
TA12_1_3	101	1225	43.16	0.85	9.82	3.80	0.06	9.85	22.09	0.96	5.74	0.23	96.57	3.43	0.56	2.87	1.02	1.35	1.55

Table 2: Run conditions, EMP analyses (averaged from 20), H₂O determined by elemental analyser (Flash) and CO₂ determined from: (i) EMPA (by subtracting H₂O content to EMPA shortfall), (ii) Elemental analyser (Flash), (iii) Micro-Raman spectroscopy (following the method of Morizet *et al.*, 2013) and (iv) Weight loss measured by puncturing the capsule (allowing the CO₂ to escape) after each run. Run duration was of 2 hours in all experiments.

	а	b	С	d	е	f	D
Value	-0.001271	-0.476116	0.132494	-0.322187	-4.501117	-0.163249	50
Std. Err.	0.000244	0.040612	0.016065	0.020635	0.281876	0.008332	1.4

Table 3: Adjusted parameters (and corresponding standard error on multiple non-linear regression fit) to calculate CO₂ solubility from Eq. (1). Regression was calculated from over 160 data points (detail in text).



Figure 1. Moussallam et al.



Figure 2, Moussallam et al.



*(by elemental analyser and weight loss)

Figure 3, Moussallam et al.



Figure 4, Moussallam et al.



Figure 5, Moussallam et al.



(calulated from equation 1)

Figure 6, Moussallam et al.



Figure S1. Moussallam et al.



Figure S2, Moussallam et al.