Carbon isotope fractionation between CO₂ and carbon in silicate melts at high temperature

3	Hyunjoo Lee ¹ , Yves Moussallam ^{1,2} , Cyril Aubaud ³ , Giada Iacono-Marziano ⁴ , Keiji Hammond ² ,
4	Denton Ebel ² .
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7	¹ Lamont-Doherty Earth Observatory, Columbia University, New York, USA
8	² American Museum of Natural History, Department of Earth and Planetary Sciences, NY 10024, New York,
9	USA
10	³ Université Paris Cité, Institut de Physique du Globe de Paris, CNRS, Laboratoire de Géochimie des
11	Isotopes Stables, 1 rue Jussieu, F-75005 Paris, France
12	⁴ ISTO, UMR 7327 Université d'Orléans-CNRS-BRGM, 1a rue de la Ferollerie, Orléans, France
13	
14	Corresponding author: Hyunjoo Lee; hl3507@columbia.edu
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29 Abstract

30 The isotopic fractionation of carbon between CO₂ gas and silicate melts is a crucial parameter to 31 understand the carbon cycle at the planetary scale that requires accurate quantification. In this 32 study, we conducted experiments to determine the carbon isotope fractionation between CO_2 gas 33 and carbon dissolved in silicate melt at 350-420 MPa and 1160-1225°C, across a range of melt 34 compositions. A linear relationship emerges between the fractionation coefficient and the degree 35 of polymerization of the melt (NBO/T; non-bridging oxygens per tetrahedral cation) with the 36 fractionation coefficient increasing for depolymerized melts (e.g., basalt) and decreasing for 37 polymerized melts (e.g., rhyolite):

1000lnα_{gas-melt}=2.847×NBO/T+0.068 (R²=0.74)

or

 $1000 \ln \alpha_{g-m} = -0.087 \times (SiO_2 + Al_2O_3) wt\% + 7.968 (R^2 = 0.74)$

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41 These equations enable the calculation of carbon fractionation coefficients in silicate melts, 42 providing a mean to interpret δ^{13} C-value measurements in natural volcanic gases and melts through 43 forward and backward modelling of degassing paths from mantle to surface. The relationship is 44 calibrated from NBO/T = 0 to 1, covering most major melt compositions. However, it is expected 45 to stabilize and remain constant at NBO/T > 1. We hypothesize that the ratio of CO_3^{2-}/CO_2 46 dissolved in the melt is the key parameter behind this relationship. Carbon dissolved as CO₂ 47 molecular transfers to the gas phase with a fractionation coefficient of 0 ‰ whilst carbon dissolved as CO_3^{2-} transfers with a fractionation coefficient of 2.9 %. Carbon isotopic fractionation might 48 49 hence offer a window into carbon speciation in natural melts.

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57 **1. Introduction**

58 Dissolved carbon in magmatic systems is an abundant volatile component that plays a crucial 59 role in determining the initiation depth of partial melting (e.g., Dasgupta and Hirschmann, 2010) 60 and physical properties of mantle melts (e.g., Gaillard et al., 2008; Yoshino et al., 2012). Carbon 61 in the mantle exists in various forms, including diamond (e.g., Jaques et al., 1990), graphite (e.g., Pearson et al., 1994), and carbonate inclusions (e.g., Ionov et al., 1993), as well as fluid inclusions 62 63 composed of CO₃²⁻, CH₄, CO, and COS (e.g., Andersen and Neumann, 2001). Under lithospheric 64 conditions (i.e., 0.5 to 4 GPa, below 2000 °C, and FMQ-1 to +3.5; Solomatova et al., 2020), carbon 65 exists primarily in dissolved forms as either carbonate ion (CO_3^{2-}) or CO_2 molecular in silicate 66 melts (e.g., Blank and Brooker, 1994; Brey and Green, 1976; Mysen et al., 1976). In the coexisting 67 exsolved gas phase, carbon is predominantly found as CO₂, with lesser amounts of CO (Wardell 68 et al., 2004).

69 The solubility of CO₂ in silicate melts has a significant positive correlation with 70 depolymerization of the melt and total pressure. To describe the complex relationship between melt composition and CO_2 solubility, researchers have proposed a compositional parameter, Π , 71 72 (Dixon, 1997; Shishkina et al., 2014) and NBO/T (Non-Bridging Oxygens per Tetrahedral Cations; 73 Brooker et al., 2001), which reflects the ratio between the freely diffusing cations and the 74 tetrahedral network, as a way of measuring the degree of melt polymerization. In addition, 75 experiments have consistently shown that the CO₂ solubility decreases with decreasing pressure 76 across a range of compositions, from felsic to ultramafic and from less than a hundred MPa to 77 several GPa, respectively (e.g., Jendrzejewski et al., 1997; Shishkina et al., 2014; Stolper and 78 Holloway, 1988). This positive correlation commonly leads to CO_2 saturation in the magma during 79 its journey toward the surface, with very few examples to date of magmatic systems where CO₂-80 undersaturated melts have been identified (Javoy and Pineau, 1991; Saal et al., 2002; Michael and 81 Graham, 2015; Le Voyer et al., 2017).

82 CO₂ has relatively low solubility among the major volatiles (H₂O-CO₂), for example, the 83 maximum solubility of H₂O is 6 wt% and CO₂ is 0.3 wt% for basaltic melt at 400 MPa (Iacono-84 Marziano et al., 2012), making CO₂ the main driver of degassing and in many cases of magma 85 ascent. Volatile degassing has two end-member processes (e.g., Aubaud, 2022), closed-system 86 degassing and open-system degassing. In closed-system degassing (BED – Batch Equilibrium 87 Degassing, often referred to as equilibrium degassing), the gas phase remains in the system and isotopic equilibrium is maintained (e.g., Graham et al., 2018). In open-system degassing (FED –
Fractional Equilibrium Degassing, often called Rayleigh distillation) exsolved CO₂ escapes from
the system instantaneously (e.g., Taylor, 1986).

91 The carbon stable isotopic ratio in a given phase is expressed with the delta notation defined by 92 Eq. (1) (abbreviation for Equation (1)):

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$$\delta^{13}C = \left(\frac{\left(\frac{1^{13}c}{1^{2}c}\right)sample}{\left(\frac{1^{13}c}{1^{2}c}\right)reference} - 1\right) \times 1000 \tag{1}$$

As CO₂ degasses, carbon isotope fractionation between gas and melt occurs. The fractionation factor ($\alpha_{gas-melt}$) between gas and melt is defined by Eq. (2). It has been often reported as $\Delta^{13}C_{eq}$ using an approximation (Eq. (3)), which is valid only when the absolute values of $\Delta^{13}C_{eq}$, $\delta^{13}C_{gas}$, and $\delta^{13}C_{melt}$ are all less than 10 ‰ (O'Neil, 1986; Sharp, 2017). Hereafter, we will call 1000ln $\alpha_{gas-melt}$ melt and the previously reported $\Delta^{13}C_{eq}$ as the fractionation coefficient.

$$\alpha_{\text{gas-melt}} = \frac{1000 + \delta^{13} C_{gas}}{1000 + \delta^{13} C_{melt}}$$
(2)

(3)

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$$1000 \ln \alpha_{\text{gas-melt}} \approx \Delta^{13} C_{\text{eq}} = \delta^{13} C_{\text{gas}} - \delta^{13} C_{\text{melt}}$$

Degassed CO₂ shows a higher δ^{13} C-value compared to the melt. Several studies on carbonate, 101 102 including experimental work (Chacko et al., 1991; Scheele and Hoefs, 1992; Rosenbaum, 1994; 103 Appora, 1998; Petschnig et al., 2024) and theoretical calculations (Bottinga, 1969; Chacko et al., 104 1991; Deines, 2004), have shown that the fractionation coefficient decreases by 4 ‰ to 2 ‰ with 105 increasing temperatures within the range of 700-1200 °C. The fractionation coefficient has been 106 reported from experimental studies to be close to 0% for rhyolitic melt (Blank, 1993), and anywhere from +1.8 to 4.7 ‰ in basaltic melts (Javoy et al., 1978; Mattey, 1991; Mattey et al., 107 108 1990; Trull et al., 1992). The variation in the fractionation coefficient in the compositional range 109 between rhyolite and basalts has remains unexplored.

110 While carbon isotope fractionation between gas and melt can be used to reconstruct the magma 111 degassing path, the lack of constraints on the fractionation coefficient has led to inconsistency 112 between studies. The higher δ^{13} C-value of degassed CO₂ results in a subsequent decrease in the 113 δ^{13} C-value of the residual melt. As both BED and FED progress, the depletion of δ^{13} C-value of 114 CO₂ within the residual melt and coexisting gas will display a distinct evolutionary path with a 115 steeper depletion of δ^{13} C-value in FED compared to BED. For example, Pineau and Javoy (1983) 116 proposed a two-step degassing model to explain the δ^{13} C-value depletion of carbon dissolved in 117 MORB, with gradual δ^{13} C-value depletion at higher C and steep depletion at lower C. This model 118 has been applied to the δ^{13} C-value variations in MORB (e.g., Cartigny et al., 2001) as well as arc 119 environments (e.g., Shaw et al., 2004). However, due to the lack of agreement in the fractionation 120 coefficient of mafic melts, δ^{13} C-value of volcanic gas and basaltic rocks were modeled to generate 121 degassing paths using either extremes or varying fractionation coefficient. For example, Mattey et 122 al. (1989) used +1 ‰ and +4.4 ‰, Paonita et al. (2012) chose +3 ‰, while several recent studies 123 employed +3.5 ‰ (e.g., Aubaud, 2022; Cartigny et al., 2001).

124 Carbon isotope fractionation in submarine volcanic glasses has been used to reconstruct pre-125 eruptive carbon concentration in the melt, but this estimate is highly dependent on the fractionation 126 coefficient. The procedure measures CO₂ concentration in the gas or melt and assumes a δ^{13} C-127 value of the initial melt (e.g., -4.5 ‰; Cartigny et al., 2001). Pre-eruptive carbon can then be calculated by modelling its degassing path (detailed calculations are provided by Cartigny et al., 128 129 2008). Estimates of Cartigny et al. (2008) (600 – 55000 ppm CO₂) and Aubaud et al. (2004) (2700 130 ppm CO₂) align with pre-degassed carbon estimates obtained through CO₂/Nb ratio and "popping 131 rock", respectively. Macpherson et al. (2005) found that the Kolbeinsey Ridge melt, the northern 132 part of the Mid-Atlantic Ridge, has a pre-eruptive CO₂ concentration (~400 ppm) lower than 133 primitive MORB, suggesting potential degassing before equilibration at 2.7-3.0 km below the 134 seafloor, and indicating a two-stage degassing process. All these calculations rely heavily on 135 knowledge of the fractionation coefficient. Macpherson et al. (2005) applied two fractionation 136 coefficient values of +2.3 ‰ and +4.5 ‰ to their calculations, resulting in a twofold difference in 137 their resulting pre-eruptive CO_2 concentration in the melt (400 and 950 ppm, respectively).

Previous experimental studies on mafic melts yielded inconsistent CO₂ solubility and mass 138 139 balance and sometimes failed to provide critical information such as melt composition or its final 140 water content. In the first attempt to decipher fractionation coefficient, Javoy et al. (1978) reported 141 a range of 4 - 4.6 % for oceanic tholeiite. However, their dissolved CO₂ in the glass at saturation 142 was inconsistent, ranging from 0.73 wt% to 2.56 wt% under identical conditions (750 MPa, 143 1280 °C). At these conditions, a CO₂ solubility of 2.56 wt% is unexpectedly high, which might 144 imply a formation of reduced carbon formation potentially influencing carbon isotope fractionation. 145 Two experiments using different brands of oxalic acid as a carbon source produced inconsistent 146 CO₂ yields, 0.5 and 2.6 wt%. Finally, their step-heating measurement showed a significant contribution of low-temperature carbon released below 600 °C, which could have been from 147

148 reduced carbon form (i.e., graphite) or surface carbon contamination (Mattey et al., 1989). Mattey 149 et al. (1990) conducted experiments on sodamelilite (NaCaAlSi₂O₇) and reported a smaller 150 fractionation coefficient of 2.2 - 2.7 %. However, their step-heating technique only recovered 20 151 -92 % of the expected carbon amount, which might have created additional kinetic isotope 152 fractionation during the step-heating measurements. In a subsequent study, Mattey (1991) 153 conducted an experiment on MORB melt composition obtaining results consistent with their prior 154 research (fractionation coefficient = 1.8 - 2.2 ‰), but suffered CO₂ loss during the experiment as 155 well. Trull et al. (1992) reported fractionation of 1.9 - 4.7 ‰ (average 3.5 ± 1.0 ‰) in experiments 156 on MORB composition but provided limited information on their initial melt composition. The 157 most recent experimental study, Appora (1998), investigated carbon isotope fractionation between 158 CO₂ and carbonate melts, finding a fractionation coefficient of 3.5 ‰, which is probably not 159 directly applicable to silicate melt.

160 One major challenge in interpreting previous experimental studies is the absence of available 161 H₂O measurements on the resulting glasses, except for Blank (1993). The presence of H₂O in the 162 melt plays a critical role in silicate structure and CO₂ behavior (e.g., Moussallam et al., 2016a; 163 Solomatova et al., 2020; Stolper and Holloway, 1988). In one hydrous experiment conducted by 164 Javoy et al. (1978), precise input material quantities were not provided, hindering accurate H₂O 165 content estimation. Even though all of the experiments on mafic melt experiments (except the one 166 above) are nominally H₂O-free, trace amounts of H₂O (a few hundred ppm to a weight percent) 167 can still be present due to ubiquitous atmospheric water and hydrogen diffusion in experiments. 168 Mattey (1991) and Mattey et al. (1990) have detected H_2O peaks in their infrared spectra, 169 emphasizing the importance of measuring H₂O content in the glass, even in nominally H₂O-free 170 experiments.

171 In light of these challenges, this study examines the effect of composition on the carbon isotope 172 fractionation coefficient between gas and melt and fills the compositional gap between rhyolite 173 and basalt. To achieve this goal, we conducted a series of experiments ranging from andesite to 174 basanite composition at 350 - 420 MPa and 1160 - 1225°C to examine the effect of melt structure 175 on the gas-melt isotopic fractionation coefficient.

176 **2. Methods**

177 **2.1. Experimental Methods**

178 The starting materials used in this study included: natural basanite from El Hierro mixed with 179 variable amounts of SiO_2 , and Al_2O_3 to produce compositions ranging from basanite to andesite 180 (NBO series), natural basalt from Mt.Etna (Etna series), and synthetic compositions made of a 181 four-oxide (SiO₂, Al₂O₃, MgO, and CaO) mixtures (Simple series). To ensure homogeneity and 182 remove any initial volatiles present, all starting materials were first fused at 1300 °C in a Pt crucible 183 at 0.1 MPa in air for 2 hours, then quenched. The retrieved glass was crushed and re-melted once 184 more under the same conditions for an additional 2 hours. The fused glasses were examined using 185 Fourier transform infrared spectroscopy (FTIR) to confirm that they were volatile-free (see 2.2.3). 186 Experiments were conducted in two internally heated pressure vessels, one at the Institut des 187 Sciences de la Terre d'Orléans (ISTO) and the other at the American Museum of Natural History 188 (AMNH) (Table 1). The variations in vessel size led to differences in capsule size and material 189 quantity, with ISTO's larger vessel allowing for the use of more materials (350 mg for ISTO and 190 150 mg for AMNH) and larger capsules. Gold-palladium (Au₈₀-Pd₂₀) tubes (5 mm o.d. / 4.5 mm 191 i.d. / 35 mm long for ISTO and 4.0 mm o.d. / 3.5 mm i.d. / 20 mm long for AMNH) were used in 192 all experiments. Cut and annealed capsules were initially welded flat at one end for the ISTO 193 experiment and triple-crimped then welded for AMNH. Then, the capsules were ultrasonically 194 cleaned in dichloromethane for 10 minutes to remove any organic carbon that might have been 195 present on the capsule's surface (Mattey et al., 1989) and then stored at 110 °C prior to use. The 196 capsules were prepared by filling them with starting material, deionized water (1 - 4 wt), and dihydrated oxalic acid (C₂H₂O₄·2H₂O) (1-6 wt% of CO₂). The δ^{13} C-value of the dihydrated oxalic 197 198 acid used is -20.58 ‰ for the Etna series, -26.68 ‰ for the NBO and Simple series and was 199 determined by repeated measurements with an elemental analyzer coupled isotope ratio mass 200 spectrometer (see 2.2.2.) with a standard deviation of ± 0.14 ‰. The final welding process utilized 201 flat welding with tungsten electrodes to minimize the introduction of additional carbon into the 202 capsule.

The detailed experimental conditions are listed in Table 1. Pure argon was used as a pressure medium (leading to oxygen fugacities above FMQ+1, where FMQ is the fayalite-magnetite-quartz buffer; Webster and De Vivo, 2002 for AMNH; Gaillard et al., 2003 for ISTO). This high oxidation state ensured that all carbon exists in oxidized forms, the CO_2 molecular or CO_3^{2-} (Wardell et al., 2004). Most of the experiments were conducted at superliquidus conditions, except for samples
CI_IPGP_NBO_3_6 and CI_IPGP_NBO_5_2. After reaching the target P-T, the experiments were
equilibrated for 18 – 24 hours. Experiments were then quenched isobarically by drop-quench at a
rate of approximately 100 °C/s (Di Carlo et al., 2006).

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Table 1

Conditions of IHPV experiments

	Pressure	Temperature	Duration		
Sample	(MPa)	(°C)	(h)	Run at	
CI_AMNH_NBO_1_3	347	1202	18	AMNH	
CI_AMNH_NBO_2	348	1168	20	AMNH	
CI_AMNH_NBO_2_2	350	1201	22	AMNH	
CI_AMNH_NBO_3	351	1189	48	AMNH	
CI_AMNH_NBO_3_1	351	1182	24	AMNH	
CI_AMNH_NBO_3_2	351	1160	23	AMNH	
CI_AMNH_NBO_3_4	350	1200	24	AMNH	
CI_AMNH_NBO_4	349	1187	24	AMNH	
CI_AMNH_NBO_5	349	1200	24	AMNH	
CI_IPGP_NBO_1_2	400	1225	24	ISTO	
CI_IPGP_NBO_3_6	393	1225	22	ISTO	
CI_IPGP_NBO_5_2	393	1225	22	ISTO	
CI_NBO_2_2	400	1225	24	ISTO	
CI_NBO_3_3	400	1225	24	ISTO	
ETNA-3-2	419	1200	24	ISTO	

ETNA-3-2bis	419	1200	24	ISTO
ETNA-3-3	419	1200	24	ISTO
ETNA-4-1	371	1200	24	ISTO

213 2.2. Analytical Methods

The gas phase was extracted from the sealed capsule and analyzed first, followed by the isotopic analysis of the dissolved CO_2 in the glass using an elemental analyzer coupled to an isotope ratio mass spectrometer. The concentration of volatiles (CO_2 -H₂O) and the major element composition of the glass were analyzed using FTIR and electron microprobe, respectively. The occurrence of crystals within experimental samples was checked by scanning electron microscopy (SEM).

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220 **2.2.1.** δ^{13} C-value analysis in the gas phase

221 The amount of CO₂ in the gas and its isotopic signature ($\delta^{13}C_g$) were analyzed at the Institut de 222 Physique du Globe de Paris (IPGP), France. Post experiment, each sealed capsule was pierced under vacuum in a dedicated extraction line. After piercing, the gas released from the capsule was 223 224 passed over CuO enclosed in a Pt foil to catalyze the oxidation reaction at 450 °C. This process 225 was carried out to ensure the conversion of any CO and H₂ present to CO₂ and H₂O (although the 226 amount of CO and H₂ was negligible). The extracted gas was then captured in a cold trap at -227 196 °C. After 15 minutes of trapping, any remaining gas (mostly N₂) was pumped off. CO₂ was 228 then released by setting the temperature of the cold trap at -135 °C to retain only H₂O. CO₂ was 229 cold trapped again and then released in a calibrated volume with a Baratron capacitance manometer 230 to measure its total amount in the range 0-10 μ mol. For larger amounts (>10 μ mol), CO₂ was 231 measured with a mercury manometer $(\pm 5 \%)$. The gas is finally trapped in a Pyrex® glass tube 232 for isotopic analysis, which was performed using a dual inlet Delta Plus XP isotope ratio mass 233 spectrometer (IRMS) equipped with a microvolume (± 0.03 ‰).

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235 **2.2.2.** δ^{13} C-value analysis in the glass

The isotopic signature of CO_2 dissolved in the glass ($\delta^{13}C_m$) was determined using a Costech elemental analyzer (EA) coupled with a ConFlo IV and Thermo Scientific Delta V plus isotope ratio mass spectrometer (IRMS) at Lamont–Doherty Earth Observatory. Thin glass shards were carefully hand-picked under a microscope to avoid entrapped gas bubbles. Glass shards were then weighed on a microbalance (± 0.001 mg), placed into a 3.2×4 mm tin capsule, and kept in a desiccator until analysis.

The enclosed samples were dropped into a 980 °C oven and then combusted at ~1700 °C with excess oxygen (25 ml/min) over a chromium (III) oxide catalyst. Helium was used as the carrier gas (100 ml/min). Silvered cobaltous/cobaltic oxide inside the quartz combustion tube ensured the complete conversion of sample carbon to CO_2 and the removal of residual halogens or sulfur. After the separation of the CO_2 peaks through a gas chromatography (GC) column (55 °C), each sample was analyzed by IRMS.

The $\delta^{13}C_m$ -values for each sample were calibrated by three-point regression using standards USGS24 ($\delta^{13}C = -16.04 \%$, VPDB), USGS40 ($\delta^{13}C = -26.77 \%$, VPDB), and USGS41 ($\delta^{13}C =$ 37.63 ‰, VPDB) with an average precision of 0.16 ‰. All measurements of carbon isotope standards agree with the accepted values reported by the United States Geological Survey (USGS). To test for instrument performance and linear drift, one standard set was analyzed for every 10 samples. The measurement error of the $\delta^{13}C$ -value, calculated to be $\pm 0.14 \%$, was determined based on repeated measurements of the carbon source, oxalic acid.

The instrument was calibrated using acetanilide (C₈H₉NO) to determine the bulk carbon content (\pm 0.2 %, R² = 0.9999). Additional calibration was performed with a mixture of oxalic acid and SiO₂ containing 70, 20, 5, 2, 1, 0.75, 0.5, and 0.25 wt% of CO₂ (R² = 0.9999) to properly account for the characteristics of the glass material to be analyzed. A set of standards was inserted for each daily run. Linear regression was used to derive the calibrated total carbon content based on the true concentration of the mixture series. Based on the reproducibility of the mixture series, the error in CO₂ estimation was calculated to be \pm 7.3 %.

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263 **2.2.3.** Glass analysis by Fourier transform infrared spectroscopy (FTIR)

H₂O and CO₂ concentrations in the retrieved glasses were measured using a Thermo Nicolet iN10 Fourier transform infrared (FTIR) spectrometer at Lamont–Doherty Earth Observatory. The instrument was purged with dry and CO₂-scrubbed air. The measurements were made on a liquid nitrogen-cooled MCT-A detector.

268 Glass chips were doubly polished with alumina-coated polishing paper. Their thickness ranged 269 from 20 to 100µm. The glass chips were washed with acetone to remove any crystal bond before 270 they were placed on a CaF_2 plate for measurement. Thickness was measured using the reflectance 271 method (\pm 3 µm; Nichols and Wysoczanski, 2007).

- Absorption spectra were collected from each sample in the 400 8000 cm⁻¹ range with 256 scans and 1 cm⁻¹ resolution in transmitted mode. The aperture size was 100 µm in width and height. To confirm the homogeneity of H₂O and CO₂ contents in the glass, a minimum of 3 spots were analyzed for each sample.
- 276 Total H₂O content was calculated using the intensity of the OH⁻ stretching band at around 3550 cm⁻¹. The CO₂ doublet peaks around 1515 cm⁻¹ and 1435 cm⁻¹ were used to determine the CO₂ 277 278 concentration. Baseline correction is performed by subtracting the spectra of chemically matched 279 devolatilized glasses (i.e., our starting compositions fused in air). The absorption coefficient of 280 CO₂ was determined using the Na/(Na+Ca) molar ratio of the glass composition (Dixon and Pan, 281 1995). The absorption coefficient of H₂O was determined to be 63 L/mol/cm (Dixon et al., 1995). 282 To calculate the glass density, both H₂O and CO₂ were taken into account (Bourgue and Richet, 283 2001; Lesher and Spera, 2015).
- 284
- 285 **2.2.4. Electron Microprobe**

286 Electron microprobe analyses were conducted at AMNH using a Cameca SX5-Tactis to 287 determine the major element composition of the experimental glasses, utilizing a 15 kV 288 accelerating voltage, a defocused beam of 10 µm, 4 nA beam current for Na (10 s count time) and 289 10 nA for other elements (20 s count time). Na was analyzed first in order to have a minimum of 290 Na loss during the analysis. The instrument was calibrated using natural and synthetic mineral 291 standards and glasses including: potassium feldspar (Al, Si, and K), rutile (Ti), fayalite (Fe), 292 rhodonite (Mn), olivine (Mg), anorthite (Ca), jadeite (Na), and apatite (P). Ten random spots were 293 analyzed to obtain an average value for each glass and the errors are reported as the standard 294 deviation of the 10 repeated analyses. The samples were imaged with Scattered Electron 295 Microscope (SEM) with Bruker Quantax Quad at AMNH.

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3. Results

298 Quench crystals were not observed under the 500 magnifications of the microscope and SEM 299 images (Fig. S1). IPGP_NBO_3_6 and CI_IPGP_NBO_5_2 included opaque Fe-oxides (less than 300 1 % in volume) that grew during the experiment (Fig. S1). Glass compositions analyzed by electron 301 microprobe are shown in Table 2. CO₂ dissolved in the glass (CO_{2glass}), H₂O dissolved in the glass (H₂O_{glass}), δ^{13} C-value of the gas ($\delta^{13}C_g$) and glass ($\delta^{13}C_m$) are shown in Table 3. CO_{2glass} in Table 302 303 3 is measured by FTIR, except for type "Simple" (CI NBO 2 2 and CI NBO 3 3), in which it 304 was obtained by EA, due to the absence of known absorption coefficient for FTIR on this Si-Al-305 Ca-Mg composition. CO_{2glass} measured by EA (Table S2) is in good agreement with CO_{2glass} 306 measured by FTIR (Fig. 1). CO_{2glass} varies from 500 to 3600 ppm, increasing with the degree of 307 depolymerization (Fig. 2). H₂O dissolved in the glass measured by FTIR varies from 1 to 3 wt%. 308 δ^{13} C-value of the gas and melt diverge from the carbon source as the depolymerization degree 309 increases (Fig. 3). The fractionation coefficient is calculated from the δ^{13} C-value of the melt and 310 of the coexisting gas as $1000 \ln \alpha_{g-m}$ (see introduction for the definition). The fractionation 311 coefficient shows a linear relationship with composition (Fig. 4). The linear correlation derived is 312 below: $1000 \ln \alpha_{g-m} = 2.847 \times NBO/T + 0.068 (R^2 = 0.74)$ 313 (4) 314 $1000 \ln \alpha_{g-m} = -0.087 \times (SiO_2 + Al_2O_3) wt\% + 7.968 (R^2 = 0.74)$ (5) 315 where NBO/T is calculated on an hydrous basis: $X_{\rm H}$ $0 + X_{\rm K}$ $0 + X_{\rm Ne}$ $0 + X_{\rm Ceo} + X_{\rm Meo} + X_{\rm Eeo} + X_{\rm Meo} - X_{\rm Meo} - X_{\rm Meo}$

316 NBO/T = 2 ×
$$\frac{X_{H_20} + X_{K_20} + X_{Na_20} + X_{Ca0} + X_{Mg0} + X_{Fe0} + X_{Mn0} - X_{Al_20_3}}{X_{Si0_2} + 2 \times X_{Al_20_3}}$$
 (6)

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Fig. 1. CO_2 dissolved in the glass (CO_{2glass}) measured by EA versus FTIR in ppm. The dashed line is the 1 to 1 line. Only "Etna" and "NBO" are shown on the plot, as the FTIR result of "Simple" is missing due to the lack of a known absorption coefficient for this composition. Shades represent the average 1-sigma deviation between two measurements. CO_2 dissolved in the glass measured by FTIR and bulk CO_2 measured by EA follow a general 1 to 1 correlation, which confirms that the EA measurements were not contaminated by additional surface carbon, CO_2 in gas bubbles, or other suspicious reduced materials.



Fig. 2. CO_{2glass} in ppm as a function of NBO/T (non-bridging oxygens to tetrahedral cations), which shows a general increase in CO_{2glass} with the depolymerization degree. CO_{2glass} is measured by FTIR except "Simple" series (see result) and associated error is a standard deviation of repeated measurement.



335 Fig. 3. This plot shows the δ^{13} C-value of gas and glass in each experiment relative to the initial

- 336 δ^{13} C-value of the carbon source (dashed line). Only "NBO" and "Simple" are shown on the plot,
- 337 since oxalic acid with a different δ^{13} C-value was used for "Etna" (δ^{13} C=-20.58 ‰).





Table 2

Compositions of the glass (in wt%) analyzed by electron microprobe after the experiment, normalized to a volatile-free basis. The unnormalized raw analytical sum of the averaged measurements is shown in Table S1. The values are an average of 10 repeated analyses. Standard deviations are reported in parentheses.

Sample	Туре	SiO ₂	TiO ₂	Al ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	P_2O_5
CLAMNIL NDO 1	2 NDO	57.77	2.73	18.79	6.86	0.10	3.48	6.12	2.47	1.06	0.63
CI_AMINH_NBO_I_	3 NDO	(0.57)	(0.04)	(0.36)	(0.19)	(0.04)	(0.05)	(0.11)	(0.07)	(0.02)	(0.05)
	NDO	56.18	3.12	16.96	8.03	0.12	3.85	6.85	2.93	1.29	0.67
CI_AMINH_NBO_2	NBU	(0.23)	(0.03)	(0.10)	(0.20)	(0.06)	(0.05)	(0.06)	(0.07)	(0.02)	(0.07)
		55.00	3.17	17.96	7.82	0.12	3.94	7.07	2.95	1.27	0.70
CI_AMINH_NBO_2_	2 NDU	(0.26)	(0.05)	(0.08)	(0.17)	(0.05)	(0.07)	(0.07)	(0.09)	(0.02)	(0.03)
	NDO	52.06	3.61	17.18	8.76	0.15	4.55	8.12	3.31	1.44	0.82
CI_AMNH_NBO_3	NBO	(0.19)	(0.04)	(0.07)	(0.17)	(0.07)	(0.06)	(0.06)	(0.06)	(0.02)	(0.03)
CLAMNILI NDO 2	1 NDO	51.17	3.58	17.69	9.25	0.19	4.50	8.10	3.31	1.40	0.81
CI_AMINH_NBO_5_	INDU	(0.34)	(0.04)	(0.10)	(0.13)	(0.08)	(0.05)	(0.05)	(0.09)	(0.04)	(0.09)
CLAMNILI NDO 2		51.28	3.60	17.51	9.39	0.17	4.47	8.05	3.34	1.41	0.78
CI_AMINH_NBO_5_	2 NDU	(0.22)	(0.06)	(0.10)	(0.23)	(0.05)	(0.07)	(0.08)	(0.12)	(0.02)	(0.04)
CLAMNH NDO 2		51.17	3.56	17.91	9.11	0.11	4.50	8.10	3.31	1.41	0.81
CI_AMINH_NDO_5_	4 NDU	(0.16)	(0.03)	(0.11)	(0.19)	(0.05)	(0.05)	(0.07)	(0.10)	(0.02)	(0.05)
	NDO	49.03	3.98	16.18	10.54	0.19	4.98	8.90	3.70	1.64	0.87
CI_AMINH_NBO_4	NBO	(0.26)	(0.04)	(0.05)	(0.13)	(0.06)	(0.06)	(0.09)	(0.12)	(0.02)	(0.07)
	NDO	45.87	4.51	15.51	11.37	0.20	5.58	10.19	3.98	1.80	0.99
CI_AMINH_NBO_5	NRO	(0.25)	(0.03)	(0.10)	(0.20)	(0.05)	(0.07)	(0.04)	(0.23)	(0.03)	(0.06)

CLIPGP NBO 1-2	NBO	57.94	2.67	18.73	7.06	0.12	3.30	5.95	2.53	1.11	0.59
	NDO	(0.40)	(0.05)	(0.19)	(0.20)	(0.07)	(0.08)	(0.09)	(0.08)	(0.02)	(0.04)
CLIDED NDO 2 6	NPO	54.14	2.62	17.56	6.89	0.14	4.31	8.42	3.65	1.48	0.78
CI_IFOF_NBO_5_0	NBO	(0.24)	(0.11)	(0.28)	(0.14)	(0.03)	(0.04)	(0.17)	(0.15)	(0.03)	(0.07)
CLUBCD NDO 5 2	NDO	48.67	3.60	16.29	7.28	0.20	5.55	10.76	4.58	1.94	1.12
CI_IFOP_NBO_5_2	NBU	(0.25)	(0.04)	(0.05)	(0.27)	(0.06)	(0.04)	(0.07)	(0.10)	(0.03)	(0.06)
CL NDO 2 2	Simula	65.33	0.03	16.75	0.09	0.03	6.62	11.09	0.03	0.02	0.01
CI_NBO_2_2	Simple	(0.36)	(0.01)	(0.06)	(0.06)	(0.04)	(0.09)	(0.07)	(0.01)	(0.01)	(0.01)
CL NDO 2 2	Simple	60.37	0.02	15.45	0.06	0.03	8.97	14.99	0.07	0.02	0.02
CI_NBO_3_3		(0.28)	(0.02)	(0.11)	(0.05)	(0.04)	(0.07)	(0.09)	(0.02)	(0.00)	(0.02)
ETNIA 2 2	Etmo	48.80	1.68	16.66	9.85	0.19	6.43	10.66	3.29	1.91	0.54
EINA-3-2	Etna	(0.25)	(0.08)	(0.15)	(0.07)	(0.03)	(0.08)	(0.12)	(0.19)	(0.03)	(0.04)
ETNIA 2 2hia	Etno	48.68	1.66	16.27	10.63	0.18	6.35	10.53	3.30	1.87	0.53
ETINA-3-2018	Etna	(0.17)	(0.05)	(0.14)	(0.08)	(0.03)	(0.05)	(0.09)	(0.14)	(0.07)	(0.06)
ETNIA 2 2	Etmo	49.22	1.66	16.32	9.87	0.17	6.37	10.64	3.29	1.89	0.56
EINA-3-3	Etna	(0.61)	(0.05)	(0.16)	(0.28)	(0.05)	(0.06)	(0.09)	(0.12)	(0.04)	(0.07)
ETNA 4 1	Etno	51.17	1.70	17.17	5.97	0.18	6.67	11.15	3.45	1.97	0.57
L11NA-4-1	Etna	(0.44)	(0.05)	(0.16)	(0.20)	(0.04)	(0.05)	(0.12)	(0.24)	(0.05)	(0.04)

Table 3

Final CO₂ and H₂O concentrations in the glass, δ^{13} C-value in the glass and gas phases, and resulting isotopic fractionation coefficient. 1-sigma error is shown.

Sample	CO _{2glass}		H_2O_{glass}		$\delta^{13}C_{glass}$		$\delta^{13}C_{gas}$		$1000 \ln \alpha_{g-m}$	
Sample	(ppm) ±		(wt%)±		(‰) ±		(‰) ±		±	
CI_AMNH_NBO_1_3	755	149	2.52	0.44	-27.35	0.17	-26.23	0.03	1.15	0.18

CI_AMNH_NBO_2	1003	113	2.92	0.35	-27.49	0.16	-26.01	0.03	1.52	0.17
CI_AMNH_NBO_2_2	2 1 1 4 4	148	3.01	0.13	-27.32	0.15	-26.30	0.03	1.05	0.15
CI_AMNH_NBO_3	1167	132	2.89	0.19	-27.49	0.07	-26.07	0.03	1.46	0.08
CI_AMNH_NBO_3_1	1397	80	2.68	0.22	-27.68	0.19	-25.98	0.02	1.74	0.20
CI_AMNH_NBO_3_2	2 1 5 0 1	177	2.76	0.05	-27.70	0.19	-26.02	0.02	1.73	0.19
CI_AMNH_NBO_3_4	1357	167	2.94	0.09	-27.89	0.06	-26.17	0.02	1.77	0.06
CI_AMNH_NBO_4	1874	344	2.33	0.14	-27.01	0.15	-25.68	0.03	1.37	0.16
CI_AMNH_NBO_5	3147	301	2.23	0.15	-27.48	0.01	-25.57	0.02	1.97	0.02
CI_IPGP_NBO_1_2	772	71	2.12	0.32	-27.03	0.37	-26.52	0.02	0.53	0.38
CI_IPGP_NBO_3_6	827	139	1.98	0.11	-28.12	0.29	-26.43	0.02	1.74	0.30
CI_IPGP_NBO_5_2	2320	245	1.66	0.15	-28.39	0.05	-25.54	0.03	2.93	0.06
CI_NBO_2_2	561	559	1.36	0.03	-27.61	0.61	-26.52	0.01	1.12	0.62
CI_NBO_3_3	520	322	1.56	0.03	-28.09	0.37	-26.43	0.03	1.71	0.38
ETNA-3-2	3343	101	2.66	0.06	-22.70	0.15	-20.25	0.02	2.50	0.15
ETNA-3-2bis	3335	54	2.81	0.02	-22.19	0.15	-19.58	0.02	2.67	0.15
ETNA-3-3	3580	208	1.73	0.07	-24.23	0.15	-19.65	0.03	4.69	0.16
ETNA-4-1	2396	66	2.84	0.08	-22.31	0.15	-18.88	0.03	3.50	0.16

348

349 **4. Discussion**

350 **4.1. Data reliability**

A critical consideration is whether the experiments reached equilibrium. First, we stress that the experimental systems remained entirely closed with no evidence of open system behavior or leakage. The recovered capsules did not show any weight loss after the experiment (Table S2). Extracted gas was mostly CO₂ (17-276 µmole) with minor traces of H₂O (background level), as 355 expected (Table S2). The recovered glasses exhibited consistent homogeneity, with a relative 356 standard deviation below 5 % in 10 repeated analyses, regardless of their location within the 357 sample (Table 2). Previous experiments under P-T-X conditions similar to ours demonstrated that they reached equilibrium by "reversal test", i.e. first equilibrating at high pressure and then re-358 359 equilibrating at lower pressure for 20 minutes to 2.5 hours (Mattey, 1991; Dixon et al., 1995). 360 Reversal experiments showed CO₂ solubility, δ^{13} C-value of the glass and gas results consistent 361 with forward experiments, indicating that a duration of 20 minutes at 1400 °C is sufficient to reach 362 equilibrium. We are, therefore, confident that the duration of our experiment (18 - 48 hours)363 allowed us to reach elemental and isotopic equilibrium for carbon.

Some of the primary challenges in analyzing the δ^{13} C-value in experimental glasses are the 364 365 presence of sub-solidus carbonate precipitates, CO₂ gas exsolution in the bubbles during quench, 366 or carbon adsorption (Mattey, 1991; Macpherson et al., 1999). Several results support the 367 successful recovery of δ^{13} C-value in the glass. No solid carbonate phases were observed under the 368 microscope and with the SEM. Isobaric drop quenching was used to ensure rapid quenching and 369 minimize post-experiment CO_2 exsolution from the melt to the gas bubble (e.g., Paillat et al., 1992). 370 Few large bubbles were observed in the recovered glasses, due to volatile saturation, but they were 371 easily avoidable by selecting thin shards under the microscope. Although the presence of 372 submicroscopic gas bubbles cannot be excluded, the strong correspondence between dissolved 373 CO₂ contents measured by FTIR and EA (Fig. 1), strongly suggests that the contributions of the 374 solid carbonate, gas bubbles, and surface carbon are minor compared to the predominant dissolved 375 CO_2 in the glass.

376 The question of whether a reduced form of carbon existed in the gas or melt is significant due 377 to its potential to substantially impact solubility, fractionation properties, and carbon interactions 378 with the melt structure. Pure argon was used as the pressure medium for the experiment, which is 379 estimated to induce an oxygen fugacity above FMQ+1 in the IHPV (Webster and De Vivo, 2002; 380 Gaillard et al., 2003). This environment allows carbon to dissolve in the silicate melt in its oxidized form (CO₃²⁻ or CO₂ molecular) (e.g, Wardell et al., 2004). We didn't observe any reduced carbon 381 382 species (e.g., CH₄, CO) in the FTIR spectra, and the good agreement between CO_{2glass} measured 383 by FTIR and EA (Fig.1) suggests the absence of reduced carbon species in the glass. Moreover, 384 our CO₂ solubilities are comparable with those predicted by CO₂ solubility models at the given P-

385

4.2. Compositional effect

as other major carbon species besides CO_3^{2-} in the glass.

389 Our experiments, conducted at 1160 – 1225°C and 350 – 420 MPa, investigate the δ^{13} C-value fractionation coefficient between gas and silicate melt (1000lnag-m) over a range of compositions 390 391 from andesitic basalt to basanite. Fig. 4 illustrates the relationships between melt composition and 392 $1000 \ln \alpha_{g-m}$ derived from both our study and the literature (Mattey et al., 1990; Mattey, 1991; Blank, 393 1993). In reviewing previous experiments, we have excluded the measurement from Javoy et al. 394 (1978) and Trull et al. (1992) because of the absence of melt composition and H₂O_{glass} information, 395 which significantly affects silicate melt structures. In addition, the analyses of Javoy et al. (1978) included low-temperature carbon from the step-heating method for $\delta^{13}C_{glass}$, possibly originating 396 397 from surface carbon or contamination (e.g., Mattey et al., 1989). In CO₂-sodamelilite experiments 398 (Mattey et al., 1990), we excluded the results that showed carbonate precipitation during the 399 experiment to focus solely on the fractionation between CO_2 and silicate melt. Given these 400 considerations, we have excluded these measurements from further examination.

T-X (Iacono-Marziano et al., 2012) (Fig. S2), corroborating the absence of reduced carbon as well

401 Due to the complexity of the various carbon species in the silicate melt and the species-specific 402 fractionation behavior, the gas-melt fractionation factor can be expressed as the sum of the mole 403 fractions by species multiplied by each fractionation factor (Deines, 2004). In the gas, oxidized 404 experimental conditions allowed CH₄ and CO to be excluded and only CO₂ to be included (e.g., Gaillard et al., 2003; Wardell et al., 2004; Webster and De Vivo, 2002). CO2 is dissolved in the 405 406 silicate melt as either carbonate ion (CO_3^{2-}) or CO_2 molecular (Brey and Green, 1976; Mysen et al., 1976; Blank and Brooker, 1994), CO₃²⁻/CO₂ ratios decreasing with the polymerization of the 407 408 melt (detailed in discussion later in this section). Carbon isotopic fractionation in silicate melt is therefore a combination of two fractionations between gas and melt: (1) CO₂ gas-CO₂ molecular 409 in the melt and (2) CO_2 gas- CO_3^{2-} in the melt. 410

It should be noted that equilibrium between volatile gases and silicate melt is achieved by diffusion processes. Reaction of the CO₂ molecular-CO₂ gas is expected to dominate over CO_3^{2-} CO₂ gas in the diffusion process (e.g., Spickenborn et al., 2010). The diffusion coefficient of bulk CO₂ in silicate melts is nearly invariant over a wide range of compositions (e.g., Sierralta et al., 2002). This is because the increase in CO₂ molecular diffusivity from rhyolite to basalt melts is 416 countered by a concurrent decrease in the fraction of CO_2 dissolved as CO_2 molecular with respect 417 to that dissolved as $CO_3^{2^-}$ (Nowak et al., 2004). The diffusion coefficient of CO_2 is one order of 418 magnitude (tholeiite) to four orders of magnitude (rhyolite) higher than that of $CO_3^{2^-}$ (Nowak et 419 al., 2004; Spickenbom et al., 2010). The activation energy of $CO_3^{2^-}$ for diffusion is also more than 420 twice as high (80.6 kJ/mol for $CO_3^{2^-}$ and 34.5 kJ/mol for CO_2 molecular), regardless of the 421 composition range (Guillot and Sator, 2011).

422 In addition, CO_3^{2-} is much more strongly bonded to the silicate structure than CO_2 molecular, in other words, it has a lower mobility in the melt and between the melt and gas phase. The CO₂ 423 424 molecular and CO₃²⁻ are bound to non-bridging oxygen (NBO, bound to one network former or to 425 a network modifier cation) and bridging oxygen (BO, bound to two network formers) in the silicate 426 structure, rather than forming a true carbonate complex with cations (Guillot and Sator, 2011; 427 Moussallam et al., 2016b). According to molecular dynamics simulation for carbon in silicate 428 melts, the distance from the carbon to the nearest oxygen is more than twice as short for CO_3^{2-} 429 (1.25A) than for the CO₂ molecular (2.65A) (Guillot and Sator, 2011).

While CO₃²⁻ exhibits limited mobility due to its lower diffusivity and stronger bonding, a rapid 430 interconversion between the CO_2 molecular and CO_3^{2-} has been observed, surpassing the rate of 431 CO_2 transport in the melt. This is demonstrated by the experiment showing a constant CO_3^{2-}/CO_2 432 433 ratio throughout diffusion profiles measured by FTIR (Sierralta et al., 2002). Molecular dynamics 434 simulations further support this observation, indicating that every diffusing CO₂ molecular into the 435 melt undergoes numerous interconversions before exiting from the melt (Guillot and Sator, 2011). The homogenous equilibrium of the CO_2 gas- CO_3^{2-} reaction is maintained during the diffusion 436 437 process in the silicate melt. Consequently, the carbon exchange between the melt and the gas is controlled by the CO₂ molecular, involving the conversion of CO_3^{2-} to CO_2 molecular in the melt 438 rather than direct diffusion of CO_3^{2-} out of the melt. 439

The δ^{13} C-value fractionation between CO₂ gas and CO₂ molecular in the melt appears to be negligible, as shown by the experiment with a rhyolitic composition that contains only CO₂ molecular in the melt (Fig. 3). Given the similar geometry of CO₂ gas and CO₂ molecular, minimal fractionation is expected. The close correspondence in antisymmetric stretching frequencies observed in IR and NMR studies between the CO₂ molecular in glass (2350cm⁻¹, 125 ppm) and CO₂ gas (2348 cm⁻¹, 124.2 ppm) further supports the similar geometry of the molecular structure between the two (e.g., Kohn et al., 1991). In addition, the comparable activation energy between the molecular CO_2 and Ar suggests that molecular CO_2 behaves as an inert molecule which has no isotopic fractionation between gas and melt at equilibrium (e.g., Marty and Humbert, 1997).

449 In contrast, during $CO_3^{2-}CO_2$ molecular conversion in the melt, ¹³C is enriched in the CO_2 molecular rather than in CO_3^{2-} . This is explained by isotope fractionation rules (Grootes et al., 450 451 1969), which is the preference of the heavy isotope ¹³C for lower coordination numbers and symmetric molecules, CO₂. Considering the lower mobility of CO₃²⁻ and the rapid conversion 452 453 between the two species in the melt (see reasoning above), we expect the isotope fractionation to occur primarily between the CO_3^{2-} and CO_2 molecular in the melt, rather than between CO_3^{2-} and 454 CO_2 gas. Consequently, the bulk fractionation may be indicative of CO_3^{2-}/CO_2 species ratio in the 455 456 melt.

457 Infrared spectroscopy studies of quenched silicate glasses have shown that CO₂ exists exclusively as the CO₂ molecular in rhyolite (Fogel and Rutherford, 1990) and as CO₃²⁻ in basalt 458 459 (Fine and Stolper, 1986; Shishkina et al., 2014). In intermediate compositions, infrared 460 spectroscopy studies showed that dissolved CO_2 in the glass exists as a combination of both with 461 a systematic decrease in CO_3^2 -/CO₂ ratio towards the felsic composition (Brey, 1976; Fogel and 462 Rutherford, 1990; King and Holloway, 2002). However, the speciation in quenched glasses is not 463 an accurate representation of that of silicate melts. Experimental studies (Morizet et al., 2001; 464 Nowak et al., 2003), molecular dynamic studies (Guillot and Sator, 2011; Moussallam et al., 465 2016b), and a synchrotron in situ infrared study (Konschak and Keppler, 2014) all show that, whilst 466 it may not be preserved in the glass, even depolymerized melts such as basalts contain non-467 negligeable amounts of molecular CO₂ at high temperature.

468 Eq. (4) and (5) are calibrated from NBO/T = 0 to 1, covering most major melt compositions at 469 the earth's surface and we interpret these relationships as the result of dissolved carbon speciation going from entirely CO₂ molecular in rhyolite (NBO/T \approx 0) to predominantly CO₃²⁻ in alkalic basalt 470 471 and basanite (NBO/T \approx 1). At NBO/T > 1 however, we expect the relationship to stabilize and remain constant as the majority of the carbon will be dissolved as CO_3^{2-} by then and the CO_3^{2-}/CO_2 472 473 ratio won't increase significantly with further depolymerisation. This interpretation is consistent 474 with the experimental results of Petschnig et al., (2024) on nephelinite-CO₂ fractionation at 1175°C 475 and 0.3 GPa (Fig. 5).

H₂O induces depolymerization of the silicate melt, increasing NBO/T (e.g., Stolper, 1982). An
 increase in fractionation coefficient is simply expected with H₂O addition, however, the paucity of

478 data on the effect of the H_2O content on the carbon speciation hinders a conclusive understanding 479 of its effect on carbon isotope fractionation. Furthermore, the limited range of H_2O content (1.4 – 480 3.0 wt%) in this study makes it difficult to accurately estimate its effect on carbon isotope 481 fractionation.

482



483

Fig. 5. This plot illustrates the carbon isotope fractionation coefficient (1000ln α_{g-m}) between CO₂ 484 gas and carbon dissolved in silicate melts, integrating results from our study with literature data 485 486 from rhyolite (Blank, 1993), MORB (Mattey, 1991), sodamelilite (Mattey et al., 1990), and 487 nephelinite (Petschnig et al., 2024). The graph depicts a scenario where carbon is present in 488 rhyolite only as CO_2 molecular, while the CO_3^{2-}/CO_2 ratio increases with depolymerization of the melt. It also highlights the limitation of CO_3^{2-} incorporation into silicate melts, indicating the 489 attainment of the maximum CO_3^{2-}/CO_2 ratio that silicate melts can have around NBO/T=1. The 490 491 solid line represents Eq. (4), while the dashed line extends from the point of Petschnig et al. (2024). 492

493

494 **4.3. Temperature effect**

Experimental results and literature data are plotted against $10^{6}/T^{2}$ in Fig. 6. Theoretical calculation studies on pure carbonate system predict a decrease in fractionation coefficient with increasing temperature, as expected for an equilibrium process (Chacko et al., 1991; Deines, 2004; Petschnig et al., 2024). This trend is also expected from the changes in species ratios in the melt. A decrease in temperature and an increase in pressure is expected to result in an increase in the 500 CO_3^{2-}/CO_2 ratio (Fine and Stolper, 1985; Guillot and Sator, 2011; Konschak and Keppler, 2014; 501 Morizet et al., 2015), which should lead to an expected increase in fractionation coefficient. 502 However, over the temperature range of our study and previous experiments (Mattey et al., 1990; 503 Mattey, 1991), no trend in fractionation coefficient with temperature at constant composition is 504 discernible.



505

Fig. 6. Isotopic fractionation versus $10^6/T^2$ in K⁻² for this study and literature data. The solid black line represents the carbonate-CO₂ gas fractionation line derived by Petschnig et al. (2024). The shaded area is an error estimated by Petschnig et al. (2024).

509

510 **4.4. Application**

511 The equation derived from this study allows the calculation of fractionation coefficients. Here, 512 comparisons are made with fractionation coefficients derived from natural samples. Graham et al. 513 (2018) estimated the fractionation coefficient in MORB from the East Pacific Rise (EPR). 514 Assuming closed-system degassing, a linear regression slope between δ^{13} C-value and CO₂ fraction of vesicle yields a fractionation coefficient of $3.3 \% (\pm 0.5 \%)$. Using Eq. (4) and (5), fractionation 515 516 coefficient of 2.3 ‰ was calculated based on its basalt composition (Goss et al., 2010). The higher 517 fractionation coefficient in Graham et al. (2018) suggests the potential involvement of open-system 518 degassing, resulting in greater carbon isotope depletion. In the extensively degassed central Lau Basin, most δ^{13} C-values and CO₂ concentrations in the glass data were consistent with a fractional 519

equilibrium degassing (FED) line plotted with a fractionation coefficient of 2.3 % (Macpherson and Mattey, 1994), which is comparable to our calculation of 2.2 % (melt composition and H₂O from Kent et al., 2002 and Pearce et al., 1994).

As the carbon isotope fractionation coefficient varies with melt composition, it should be 523 524 expected to change as magma evolves during fractional crystallization. To investigate the effect of 525 magma evolution on δ^{13} C-value changes during degassing, we performed fractional crystallization 526 simulation using MELTS (Fig. 7). The initial setting was Etna melt inclusion composition 527 (Kamenetsky et al., 2007), 600 MPa, and 1180 °C. The SiO₂ content ranges from 44 wt% to 52 528 wt%, resulting in a decrease in fractionation coefficient from 3.3 ‰ to 1.1 ‰. Both the BED and FED show a steeper decrease in δ^{13} C-value during the early stages of crystallization, with smaller 529 530 isotope fractionation as magma evolves. A higher δ^{13} C-value difference between gas-melt in depolymerized melt further suggests that observing a higher δ^{13} C-value in volcanic gas may not 531 necessarily indicate an increased source δ^{13} C-value but could reflect mafic recharge of the magma 532 533 chamber.



534

535 Fig. 7. A simulation result from rhyolite-MELTS (v.1.2.0) that shows the evolution of isotopic

536 compositions (δ^{13} C) for gas and melt phases under isobaric conditions at 600MPa. The starting 537 composition is from melt inclusions from Etna volcano (Reference) and the temperature is 1200°C.

538 The fractionation coefficient changes from 3.3 % to 1.1 % as the melt composition changes. The

539 $\delta^{13}C_i$ -value is set to be -4 ‰. The dashed line represents the gas $\delta^{13}C$ -value and the solid line

540 represents the melt δ^{13} C-value. BED and FED refer to Batch and Fractional Equilibrium Degassing,

541 respectively. The temperature and corresponding melt fraction remaining in the system are plotted

on the bottom and top axes, respectively.

543 The δ^{13} C-value of carbon dissolved in the melt (δ^{13} C_m) provides valuable insights into 544 identifying the carbon source and determining the initial carbon isotope ratio and carbon 545 concentration by tracing degassing pathways. δ^{13} C_m is influenced by a combination of 1) BED, 2) 546 FED, 3) kinetic fractionation, 4) carbonate precipitation from the melt, 5) mixing with organic 547 sediment, and 6) assimilation with surrounding carbonate (Fig. 8).

+

548
$$\delta^{13}C_{m} = M \times \{\delta^{13}C_{i} + \Delta^{13}C_{eq} (F-1) + \Delta^{13}C_{eq} ln(F/F_{0}) + \Delta^{13}C_{kin} ln(F/F_{0}) + \Delta^{13}C_{carb}(F/F_{0}-1)\}$$
549
$$S \times \delta^{13}C_{org} + L \times \delta^{13}C_{lime} (7)$$

 $\delta^{13}C_i$, the initial $\delta^{13}C$ -value, may differ from the source; most mantle values are estimated in a 550 range of -4 ‰ to -6 ‰ (Cartigny et al., 1998). $\Delta^{13}C_{eq}$ (fractionation coefficient) can be derived 551 552 from Eq. (4) in this study, taking into account the melt composition of each system. F is the remaining CO₂ fraction in the silicate melt relative to the initial, while F₀ is the fraction at which 553 554 Rayleigh distillation or FED or carbonate precipitation begins. BED and Rayleigh distillation 555 induce a decrease in δ^{13} C-values with different slopes. Conversely, kinetic fractionation is a 556 diffusive mass-dependent reaction ($\Delta^{13}C_{kin}$ =-8.7 ‰; Javoy and Pineau, 1991), resulting in the gas having a smaller δ^{13} C-value than the melt, and consequently increasing δ^{13} C_m. Carbonate 557 558 precipitation from the melt has minimal effect on δ^{13} C-values due to the near-zero fractionation ($\Delta^{13}C_{carb}=0.39$ %; Petschnig et al., 2024) between silicate melt and precipitated carbonatite. 559 Organic sediment significantly reduces δ^{13} C-values, whereas marine limestone typically has 560 values close to 0‰ ($\delta^{13}C_{org}$ =-30 ‰ and $\delta^{13}C_{lime}$ =0 ‰; Sano and Marty, 1995). M, S, and L are 561 562 the mole fractions of silicate melts, organic sediment and limestone (or carbonate), respectively with M+S+L=1 (Sano and Marty, 1995). Understanding these processes enables accurate 563 interpretation of δ^{13} C-value for carbon dynamics. 564



Fig. 8. Semi-quantitative assessment of magmatic processes that can affect the δ^{13} C of the melt 566 $(\delta^{13}C_m)$. The fractionation coefficient used here is 2.5 ‰. In a closed system, Batch Equilibrium 567 Degassing (BED) occurs (blue). At point F₀, Fractional Equilibrium Degassing (FED) (orange) 568 569 follows, resulting in a significant decrease in δ^{13} C-value. Kinetic fractionation (green) may occur 570 during degassing, potentially increasing δ^{13} C. The precipitation of carbonatite (red) from the melt has minimal impact on δ^{13} C. The δ^{13} C_m is affected by mixing with carbonate (purple) or organic 571 572 sediment (brown), as indicated by the percentage of mixed carbonate or sediment represented by the tick on its line. 573

- 574
- 575

576 **5.** Conclusions

577 We conducted experiments at 350 - 420 MPa and 1160 - 1225 °C across a range of 578 compositions from basanite to andesite to derived carbon isotope fractionation coefficient between 579 CO₂ gas and carbon dissolved in the silicate melt.

580 We found a positive linear correlation between the carbon fractionation coefficient and the 581 composition or the degree of depolymerization of the melt.

582 583

584

and

 $1000ln\alpha_{gas-melt}=2.847 \times NBO/T+0.068$ (R²=0.74)

 $1000 \ln \alpha_{g-m} = -0.087 \times (SiO_2 + Al_2O_3) \text{ wt} \% + 7.968 \text{ (R}^2 = 0.74)$

585 These equations allow the calculation of the carbon isotope fractionation coefficient for the most 586 commonly occurring silicate melts ranging from basanite to rhyolite. At NBO/T > 1 we expect the 587 isotope fractionation coefficient to remain constant at ~ 2.9 ‰ as we hypothesize that the ratio of 588 CO_3^{2-}/CO_2 dissolved in the melt is the key parameter behind this relationship. Carbon isotopic 589 fractionation might hence offer a window into carbon speciation in natural melts.

590 Melt temperature and H₂O could be of secondary importance to determine the carbon isotope 591 fractionation coefficient compared to melt composition, but their effects should be investigated 592 further.

593

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599 **Data Availability**

600DataareavailablethroughMendeleyDataat601https://data.mendeley.com/preview/vp6ngrjzdy?a=d27af8c4-7fbb-4b02-b771-43723c5d328f

602

603 **CRediT authorship contribution statement**

604 Hyunjoo Lee: Conceptualization, Data curation, Investigation, Methodology, Supervision, 605 Validation, Visualization, Writing – original draft, Writing – review & editing. Yves Moussallam: 606 Conceptualization, Formal Analysis, Funding acquisition, Methodology, Project administration, 607 Resources, Supervision, Validation, Writing - review & editing. Cyril Aubaud: Conceptualization, 608 Formal Analysis, Methodology, Resources, Validation, Writing - review & editing. Giada Iacono-609 Marziano: Conceptualization, Formal Analysis, Methodology, Resources, Validation, Writing -610 review & editing. Keiji Hammond: Methodology, Resources, Validation. Denton Ebel: 611 Resources, Supervision, Writing - review & editing, Validation.

612

613 Appendix A. Supplementary Material

The supplementary material includes Figure S1, which presents SEM images of the experimental glasses, Figure S2, providing a comparison between CO₂ dissolved in the glass and the expected solubility using a model. Table S1 is unnormalized analytical total of major composition data, and Table S2 details the mass difference before and after each experiment, CO_{2glass} measurements by elemental analyzer, and the extracted amounts of CO₂ and H₂O gas.

619	Declaration of generative AI and AI-assisted technologies in the writing process
620	During the preparation of this work the authors used DeepL in order to correct grammatical error
621	and improve readability. After using this tool/service, the authors reviewed and edited the content
622	as needed and takes full responsibility for the content of the publication.
623	
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Supplementary Material

'Carbon isotope fractionation between CO₂ and carbon in silicate melts at high temperature'

Hyunjoo Lee, Yves Moussallam, Cyril Aubaud, Giada Iacono-Marziano, Keiji Hammond, and Denton Ebel

CI_AMNH_NBO_1_3	CI_AMNH_NBO_2	CI_AMNH_NBO_2_2
	1	•
200µm		
CLAMNH NRO 2	CLAMNH NRO 2.4	CLAMNH NBO 2.2
CI_AMINH_NBO_3		
		the second s
CLAMNH NBO 3 4	CLAMNH NBO 4	CLAMNH NBO 5
-		
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6	Cl_IPGP_NBO_5_2
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6	Cl_IPGP_NBO_5_2
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6	CI_IPGP_NBO_5_2
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6	Cl <u>4</u> PGP_NBO_5_2
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6	Cl <u>M</u> PGP_NBO_5_2
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6 Fe-oxide	Cl <u>H</u> PGP_NBO_5_2
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6 Fe-oxide	Cl <u>H</u> PGP_NBO_5_2
CI_IPGP_NBO_1_2	CI_IPGP_NBO_3_6	Cl_IPGP_NBO_5_2
CI_IPGP_NBO_1_2 CI_NBO_2_2	CI_IPGP_NBO_3_6 Fe-oxide CI_NBO_3_3	CI_IPGP_NBO_5_2 Fe-oxide ETNA_3_2
CI_IPGP_NBO_1_2 CI_NBO_2_2	CI_IPGP_NBO_3_6 Fe-oxide CI_NBO_3_3	Cl_IPGP_NBO_5_2 Fe-oxide ETNA_3_2
CI_IPGP_NBO_1_2 CI_NBO_2_2 ETNA_3_2bis	CI_IPGP_NBO_3_6 Fe-oxide CI_NBO_3_3 ETNA 3 3	CI_IPGP_NBO_5_2 Fe-oxide ETNA_3_2 ETNA_4_1
CI_IPGP_NBO_1_2 CI_NBO_2_2 ETNA_3_2bis	CI_IPGP_NBO_3_6 Fe-oxide CI_NBO_3_3 ETNA_3_3	CI_IPGP_NBO_5_2 Fe-oxide ETNA_3_2 ETNA_4_1
CI_IPGP_NBO_1_2 CI_NBO_2_2 ETNA_3_2bis	CI_IPGP_NBO_3_6 Fe-oxide CI_NBO_3_3 ETNA_3_3	Cl_IPGP_NBO_5_2 Fe-oxide ETNA_3_2 ETNA_4_1
CI_IPGP_NBO_1_2 CI_NBO_2_2 ETNA_3_2bis	CI_IPGP_NBO_3_6 Fe-oxide CI_NBO_3_3 ETNA_3_3	CI_IPGP_NBO_5_2 Fe-oxide ETNA_3_2 ETNA_4_1

Fig. S1. Representative Scanning Electron Microscope (SEM) images taken at the AMNH (American Museum of Natural History). Sample ID is shown at top right. The scale is the same for all the images. Except for CI_IPGP_NBO_3_6 and CI_IPGP_NBO_5_2, all samples are crystal-free glass.



Fig. S2. CO₂ that dissolved in the glass modeled by Iacono-Marziano et al., (2012) versus measured by FTIR. Error is given by the model for y-axix error and by one sigma of repeated analysis for x-axis. Dashed line is 1 to 1 line.

Table S1

Raw result of major c	ompositions of th	e glass (in wt%	b) analyzed by	electron micro	probe after
the experiment. The v	alues are an aver	age of 10 repea	ted analyses.		

Sample	Туре	SiO ₂	TiO ₂	Al ₂ O ₃	FeO	MnO	MgO	CaO	Na ₂ O	K ₂ O	P_2O_5	Total
CI_AMNH_NBO_1_3	NBO	56.44	2.67	18.35	6.70	0.10	3.40	5.97	2.42	1.03	0.62	97.7
CI_AMNH_NBO_2	NBO	54.72	3.04	16.52	7.82	0.12	3.75	6.67	2.86	1.26	0.65	97.4
CI_AMNH_NBO_2_2	NBO	53.56	3.08	17.49	7.61	0.12	3.84	6.88	2.87	1.24	0.68	97.4
CI_AMNH_NBO_3	NBO	50.54	3.50	16.68	8.50	0.14	4.42	7.88	3.22	1.40	0.80	97.1
CI_AMNH_NBO_3_1	NBO	49.72	3.48	17.19	8.99	0.18	4.38	7.87	3.22	1.36	0.78	97.2
CI_AMNH_NBO_3_2	NBO	49.89	3.50	17.03	9.13	0.16	4.35	7.83	3.25	1.37	0.76	97.3
CI_AMNH_NBO_3_4	NBO	49.72	3.46	17.40	8.85	0.10	4.37	7.87	3.22	1.37	0.79	97.1
CI_AMNH_NBO_4	NBO	47.67	3.87	15.73	10.24	0.18	4.84	8.65	3.60	1.60	0.85	97.2
CI_AMNH_NBO_5	NBO	44.13	4.34	14.92	10.93	0.20	5.37	9.80	3.83	1.73	0.95	96.2
CI_IPGP_NBO_1_2	NBO	56.57	2.59	16.50	6.89	0.12	3.23	5.81	2.23	1.09	0.56	95.6
CI_IPGP_NBO_3_6	NBO	53.54	2.59	17.36	6.82	0.14	4.26	8.32	3.61	1.46	0.77	98.9
CI_IPGP_NBO_5_2	NBO	47.31	3.50	15.83	7.08	0.20	5.40	10.46	4.45	1.89	1.09	97.2
CI_NBO_2_2	Simple	64.76	0.03	16.61	0.09	0.03	6.56	11.00	0.03	0.02	0.01	99.1
CI_NBO_3_3	Simple	59.70	0.02	15.29	0.06	0.03	8.88	14.84	0.07	0.02	0.02	98.9
ETNA-3-2	Etna	47.85	1.64	16.34	9.66	0.19	6.31	10.45	3.22	1.87	0.53	98.1
ETNA-3-2bis	Etna	47.77	1.63	15.97	10.43	0.17	6.23	10.33	3.23	1.84	0.52	98.1
ETNA-3-3	Etna	48.87	1.65	16.21	9.80	0.17	6.33	10.57	3.26	1.87	0.56	99.3
ETNA-4-1	Etna	49.95	1.66	16.76	5.83	0.17	6.51	10.88	3.37	1.92	0.56	97.6

Table S2

Mass changes of the capsule before and after the experiment, CO ₂ measured by elemental
analyzer (EA) in ppm, and extracted gas μ moles of CO ₂ and H ₂ O.

	Before	aAfter	$CO_{24} = FA$	$CO_{24} = FA$	CO_{2}	H ₂ O
Sample	experiment	experiment	CO2glass_LA	CO2glass_LA	CO ₂ gas	112Ogas
	(g)	(g)	(ppm)	±	(µmol)	(µmol)
CI_AMNH_NBO_1_3	1.02673	1.0278	521	^b 38	26.3	4.9
CI_AMNH_NBO_2	0.99981	0.99992	949	^b 69	20.3	4.6
CI_AMNH_NBO_2_2	1.01559	1.0161	812	^b 59	29.2	8.1
CI_AMNH_NBO_3	0.97511	0.9748	775	°98	21	2.6
CI_AMNH_NBO_3_1	0.9958	0.996	926	°53	23.9	0.9
CI_AMNH_NBO_3_2	0.9872	0.98728	940	^b 69	22.2	6.8
CI_AMNH_NBO_3_4	0.97737	0.9774	1082	°109	23.9	6.5
\overline{CI} AMNH NBO 4	1.01072	1.01074	1857	^b 136	19.9	4
CI_AMNH_NBO_5	0.99472	0.99484	3026	°344	17.5	5.8
CI_IPGP_NBO_1_2	2.1682	2.1685	887	°357	55.9	4.3
CI_IPGP_NBO_3_6	2.2008	2.2013	868	°257	49.8	3.4
CI_IPGP_NBO_5_2	2.2285	2.2291	2637	^b 186	44.4	8.8
CI_NBO_2_2	2.2202	2.2234	561	°559	74.5	2.6
CI_NBO_3_3	2.1489	2.149	520	°322	75	8.3
$\overline{\text{ETNA-3-2}}$	1.2146	1.2145	3726	^b 272	214	1.26
ETNA-3-2bis	1.2913	1.2911	3276	^b 239	276	1.18
ETNA-3-3	1.2684	1.2685	2978	^b 217	54	0
ETNA-4-1	1.147	1.1474	1993	^b 145	174	20.57

^aMass gain may occur by reaction with capsule and Pt thermocouple.

^bDue to the limited amount of samples, there was a lack of repeated measurement. The analytical error of EA, 7.3% of the yielded value, was estimated as an error.

^cError is one sigma of repeated measurements