1	Dissolution susceptibility of glass-like carbon versus crystalline graphite in high-		
2	pressure aqueous fluids and implications for the behavior of organic matter in		
3	subduction zones		
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7	Author names:		
8	Simone Tumiati ^a , Carla Tiraboschi ^{a,b} , Francesca Miozzi ^{a,c} , Alberto Vitale-Brovarone ^{c,d} , Craig E.		
9	Manning ^e , Dimitri Sverjensky ^f , Sula Milani ^a , Stefano Poli ^a		
10			
11	Affiliations:		
12	^a Dipartimento di Scienze della Terra, Università degli Studi di Milano, via Mangiagalli 34, 20133		
13	Milano, Italy; simone.tumiati@unimi.it (corresponding author)		
14	^b Institut für Mineralogie, Universität Münster, Correnstrasse 24, 48149 Münster, Germany		
15	^c Sorbonne Université, Muséum National d'Histoire Naturelle, UMR CNRS 7590, IRD, Institut de		
16	Minéralogie, de Physique des Matériaux et de Cosmochimie, IMPMC, 75005 Paris, France		
17	^d Dipartimento di Scienze della Terra, Università degli Studi di Torino, via Valperga Caluso 35,		
18	10125 Torino, Italy		
19	^e Department of Earth, Planetary and Space Sciences, University of California, Los Angeles,		
20	California 90095-1567, USA		
21	^f Department of Earth & Planetary Sciences, Johns Hopkins University, Baltimore, MD 21218, USA		
22			
23	Abstract:		
24	Organic matter, showing variable degrees of crystallinity and thus of graphitization, is an important		
25	source of carbon in subducted sediments, as demonstrated by the isotopic signatures of deep and		
26	ultra-deep diamonds and volcanic emissions in arc settings. In this experimental study, we		
27	investigated the dissolution of sp^2 hybridized carbon in aqueous fluids at 1 and 3 GPa, and 800°C,		
28	taking as end-members i) crystalline synthetic graphite and ii) X-ray amorphous glass-like carbon.		
29	We chose glass-like carbon as an analogue of natural "disordered" graphitic carbon derived from		
30	organic matter, because unlike other forms of poorly ordered carbon it does not undergo any		
31	structural modification at the investigated experimental conditions, allowing approach to		
32	thermodynamic equilibrium. Textural observations, Raman spectroscopy, synchrotron X-ray		
33	diffraction and dissolution susceptibility of char produced by thermal decomposition of glucose		
34	(representative of non-transformed organic matter) at the same experimental conditions support this		

- 35 assumption. The redox state of the experiments was buffered at $\Delta FMQ \approx -0.5$ using double 36 capsules and either favalite-magnetite-quartz (FMQ) or nickel-nickel oxide (NNO) buffers. At the investigated $P-T-fO_2$ conditions, the dominant aqueous dissolution product is carbon dioxide, 37 38 formed by oxidation of solid carbon. At 1 GPa and 800°C, oxidative dissolution of glass-like 39 carbon produces 16–19 mol% more carbon dioxide than crystalline graphite. In contrast, fluids 40 interacting with glass-like carbon at the higher pressure of 3 GPa show only a limited increase in CO_2 (fH_2^{NNO}) or even a lower CO_2 content (fH_2^{FMQ}) with respect to fluids interacting with 41 crystalline graphite. The measured fluid compositions allowed retrieving the difference in Gibbs 42 43 free energy (ΔG) between glass-like carbon and graphite, which is +1.7(1) kJ/mol at 1 GPa-800°C and +0.51(1) kJ/mol (fH_2^{NNO}) at 3 GPa–800°C. Thermodynamic modeling suggests that the decline 44 in dissolution susceptibility at high pressure is related to the higher compressibility of glass-like 45 46 carbon with respect to crystalline graphite, resulting in G-P curves crossing at about 3.4 GPa at 47 800°C, close to the graphite-diamond transition. The new experimental data suggest that, in the 48 presence of aqueous fluids that flush subducted sediments, the removal of poorly crystalline 49 "disordered" graphitic carbon is more efficient than that of crystalline graphite especially at shallow 50 levels of subduction zones, where the difference in free energy is higher and the availability of 51 poorly organized metastable carbonaceous matter and of aqueous fluids produced by 52 devolatilization of the downgoing slab is maximized. At depths greater than 110 km, the small 53 differences in ΔG imply that there is minimal energetic drive for transforming "disordered" 54 graphitic carbon to ordered graphite; "disordered" graphitic carbon could even be energetically 55 slightly favored in a narrow *P* interval.
- 56

57 **1. Introduction**

58 The carbon isotopic signature of the upper mantle, transition zone and lower mantle (Stachel et al., 59 2002; Cartigny et al., 2004; Palot et al., 2014), and of gaseous CO₂ emitted from arc volcanoes 60 (Mason et al., 2017) suggests that organic matter subducted within sediments displays a major role in the deep carbon cycle (Hayes and Waldbauer, 2006). The dissolution of graphitic carbon in 61 62 aqueous fluids due to oxidation or reduction processes (Connolly and Cesare, 1993; Connolly, 1995; Zhang et al., 2018; Tumiati and Malaspina, 2019b) is of primary importance as it governs the 63 64 removal of organic matter from the sediments flushed by fluids released from the dehydrating 65 subducted plate (Schmidt and Poli, 2013). In contrast to carbonates (e.g., Kelemen and Manning, 66 2015), graphite has long been considered to represent a refractory sink of carbon in the subducting 67 slab (Plank and Manning, 2019), showing low solubility in metamorphic fluids (Connolly and 68 Cesare, 1993) and silicate melts (Duncan and Dasgupta, 2017). However, recent thermodynamic

69 models and experiments suggest that graphite can be readily dissolved in subduction fluids

70 (Manning et al., 2013), stressing for instance the importance of pH (Sverjensky et al., 2014) and of

71 dissolved silica (Tumiati et al., 2017). However, it is still not known how graphite crystallinity

72 might affect the compositions of COH fluids in subduction zones.

73 Carbonaceous material in sedimentary rocks metamorphosed under temperature and pressure

74 conditions characteristic of subduction zones exhibit a progressive increase in crystallinity (e.g.,

75 Beyssac et al., 2002). Besides exceptional preservations of amorphous-like carbon in some

76 metamorphic rocks (Bernard et al., 2007), carbonaceous material metamorphosed under prograde

77 temperature increase to about 550 °C is characterized by a variety of turbostratic structures, from

78 poorly crystallized to almost crystalline, that are commonly referred to as disordered graphitic C

79 (Beyssac and Rumble, 2014; Vitale Brovarone et al., 2013; Bollinger et al., 2004)). Carbonaceous

80 material in high-temperature and ultra-high-pressure terranes is instead characterized by rather

81 crystalline graphite (Beyssac et al., 2002). Thus, disordered graphitic carbon should represent the

82 most common form of carbonaceous material under forearc to sub-arc metamorphic conditions

83 where large amounts of aqueous fluids are released from the slab.

84 Crystalline (ordered) graphite is crystallographically defined by an interplanar *d* value of exactly

85 3.35 Å and a long-range crystalline order with crystallite size of at least a few dozen nanometers

86 (Luque et al., 1998). The chemical bonds within the layers of hexagonally arranged carbon atoms

87 are covalent with sp^2 hybridization (Fitzer et al., 1995; Langenhorst and Campione, 2019). The

88 thermodynamic properties of carbon without long-range crystalline order and of poorly crystallized

89 and/or defect-rich ("disordered") graphitic carbon are not available at high-pressure and high-

90 temperature conditions, hence thermodynamic models generally assume perfect ordering and well-

91 developed crystallinity of graphite. Although some studies suggested that "disordered" graphite and

92 well crystallized graphite may display different dissolution behavior in aqueous fluids (Ziegenbein

and Johannes, 1980; Connolly, 1995; Luque et al., 1998), others concluded that their

94 thermodynamic properties must be very similar (McCollom, 2013).

95 In this study, we provide the first experimental results on the high-pressure/high-temperature

96 dissolution at relatively oxidized conditions of i) ordered crystalline graphite (with crystallite size

97 around 50 nm determined by synchrotron X-ray diffraction and small Raman "disorder" *D* peak)

98 and ii) X-ray amorphous glass-like carbon, by measuring the CO₂ content of aqueous fluids in

99 equilibrium with these two sp^2 -hybridized carbon forms. Glass-like carbon is indeed a typical

- 100 disordered sp^2 bonded carbon (like graphitic carbon) consisting of randomly distributed curved
- 101 graphene layer (also called disordered multilayer graphene, or DMLG) fragments dispersed in an
- amorphous matrix (Hu et al., 2017). Glass-like carbon is non-crystalline (X-ray amorphous) and

103 non-graphitizing, meaning that it resists the development of graphite crystals up to 3000°C at room 104 pressure, and up to 45 GPa at room temperature (Shiell et al., 2018). From a thermodynamic point 105 of view, glass-like carbon can be considered as a quasi-amorphous or crypto-crystalline solid with 106 crystallite dimensions smaller than 5–8 nm (Guencheva et al., 2001). In light of these properties, we 107 used glass-like carbon as an analogue of poorly organized organic matter. The advantage of using 108 glass-like carbon instead of other poorly crystalline disordered carbon allotropes lies in its 109 metastable persistence at the investigated experimental conditions (1-3 GPa, 800°C, 12-24h). Other 110 disordered carbon materials, such as saccharose-based char, are known to recrystallize promptly at 111 high-pressure/high-temperature conditions (Beyssac et al., 2003), which would prevent 112 equilibration between fluids and solid carbon in days-long runs, and therefore the retrieval of 113 thermodynamic parameters. The choice of using glass-like carbon as an analogue for natural 114 disordered carbon is also supported by comparison of glass-like carbon with char produced by 115 thermal decomposition of glucose at the same experimental conditions. 116 The experimental results allowed retrieving the difference in Gibbs free energy (G) of glass-like 117 carbon compared to that of crystalline graphite at high pressures and temperatures, demonstrating

- 118 that, although small, this difference can lead to substantial changes in dissolved carbon content
- predicted by available thermodynamic models that assume a perfectly ordered crystalline state ofgraphitic carbon.
- 121

122 **2. Methods**

In this study, we use the recommended terminology of IUPAC (Fitzer et al., 1995) for the
description of carbon as a solid, which is summarized in the Appendix.

125

126 2.1 Bulk compositions, starting materials and buffering strategy

127 Fluids in equilibrium with crystalline and disordered sp^2 carbon were generated starting from oxalic

acid dihydrate (OAD; H₂C₂O₄·2H₂O; Sigma-Aldrich), highly ordered graphite powder (from

129 spectroscopic-pure rods, gently hand-ground in boron carbide mortar) and glass-like carbon

130 spherical powder (Alfa Aesar; type I, i.e., produced by firing polymeric precursors at temperatures

131 below 2000°C). Crystalline graphite and glass-like carbon were characterized by scanning-electron

- 132 microscopy, micro-Raman spectroscopy and synchrotron X-ray diffraction. Additionally, two
- 133 experiments have been replicated using pure water (MilliQ, boiled while flushed with N₂ to remove
- 134 dissolved atmospheric CO₂) instead of OAD to check for possible discrepancies. Experiments were
- 135 buffered using the double-capsule technique (Eugster and Skippen, 1967) to prevent the direct
- 136 contact with the buffering assemblages, with an inner H_2 -permeable $Au_{60}Pd_{40}$ capsule (OD = 2.3

- 137 mm) and an outer Au capsule (OD = 4.5 mm). The outer capsule contained H₂O and either the
- 138 fayalite-magnetite-quartz (FMQ) or the nickel-nickel oxide (NNO) buffers (Fig. 1). Fayalite and
- 139 magnetite have been synthesized at 1100°C in a gas-mixing furnace under reducing atmosphere
- 140 (CO₂:CO=10:1), starting from stoichiometrically weighted reagent-grade Fe₂O₃ (Sigma-Aldrich)
- 141 and amorphous SiO₂ from hydrolyzed tetraethyl orthosilicate (Sigma-Aldrich). Natural hyaline
- 142 quartz powder has been used to build up the final FMQ buffering assemblage. NNO buffer was
- 143 prepared by mixing Ni metal powder (Sigma-Aldrich) and green NiO nanopowder (Sigma-Aldrich),
- 144 previously sintered at 1300°C for 24 h under oxidizing atmosphere to prevent grain size-induced
- discrepancies with the accepted free-energy values of the NNO buffer (Mattioli and Wood, 1988;
- 146 O'Neill and Pownceby, 1993). At equilibrium conditions, as long as all the buffering phases are
- 147 present, the chemical potential of hydrogen is expected to be homogeneous in the inner and in the
- 148 outer capsules. In the outer capsule, the hydrogen fugacity (fH_2) is constrained through the
- 149 reactions:

150
$$Ni + H_2O = NiO + H_2$$
 (Eq. 1)

- 151 3 $Fe_2SiO_4 + H_2O = 2 Fe_3O_4 + 3 SiO_2 (quartz) + 2 H_2 (1 GPa, 800°C)$ (Eq. 2)
- 152 3 $FeSiO_3 + H_2O = Fe_3O_4 + 3 SiO_2$ (coesite) + H_2 (3 GPa, 800°C; cf. Supplementary Fig. 1) (Eq. 3)
- 153 In the inner capsule, the equilibration of the COH fluid is accomplished by the fH_2 -dependent 154 coupled reactions (Tumiati et al., 2017):

155
$$C + H_2 + O_2 = CO_2 + H_2O$$
 (Eq. 4)

156 and

157
$$C + 2 H_2O = CO_2 + 2 H_2$$

As a consequence, the initial fluid composition ($H_2O:CO_2 = 1:1$ starting from OAD) adjusts its CO_2 fraction until equilibrium in fH_2 is reached between the inner and the outer capsule. The oxygen

(Eq. 5)

- 160 fugacity (fO_2) in the inner capsule (containing mixed H₂O–CO₂ fluids instead of pure water) is
- 161 constrained indirectly and it will be slightly lower compared to the fO_2 constrained in the outer
- 162 capsule (containing pure H₂O) by FMQ and NNO (cf. Luth, 1989) and it is calculated by
- 163 thermodynamic modeling (see Section 2.5 and Table 2), ranging from ΔFMQ (= log fO_2^{sample} log 164 log fO_2^{FMQ}) –0.5 to –0.8.
- 165 High-pressure pyrolysis of glucose to char has been conducted in unbuffered single Au capsules
- 166 (OD = 3 mm) welded shut after filling with anhydrous glucose ($C_6H_{12}O_6$, or CH_2O) produced by
- 167 dehydration at 70°C (2 h) of α-D-glucose monohydrate (Sigma Aldrich) (Ponschke and House,
- 168 2011).
- 169
- 170 2.2 Experimental conditions and apparatus

Experiments were performed at 1 and 3 GPa at 800°C using an end-loaded piston-cylinder 171 172 apparatus. Capsules were embedded in MgO rods (Norton Ceramics) and inserted in graphite 173 furnaces surrounded by NaCl and borosilicate glass (Pyrex). At the top of the assembly, a 174 pyrophyllite-steel plug was placed to ensure the electrical contact. Temperatures were measured with K-type thermocouples and are considered accurate to \pm 5°C. An alumina disk was placed at the 175 176 top of the capsule to avoid the direct contact with the thermocouple. Pressure calibration of the 177 apparatus is based on the quartz to coesite transition according to Bose and Ganguly (1995) 178 (accuracy ± 0.01 GPa). Samples were first pressurized at run pressure, then heated to $T = 800^{\circ}$ C, 179 with a ramp of 100°C/min. Experiments were quenched by turning off the power supply, resulting 180 in a rate of temperature decline of $> 40^{\circ}$ C/sec. The double capsules were prepared by peeling off the outer capsule, then heated at 110°C in a vacuum oven (> 2 h) to remove any residual water 181 182 trapped in the buffer. After the analysis of volatiles (see below), double capsules were mounted in 183 epoxy resin and polished for scanning electron microscopy and micro-Raman spectroscopy. The 184 persistence of the buffering assemblages was always verified by means of electron microprobe 185 analyses. Single capsules containing char derived from the decomposition of anhydrous glucose 186 were analyzed for volatiles, then opened to collect the sample, which has been analyzed by 187 scanning electron microscopy, micro-Raman spectroscopy and synchrotron X-ray diffraction.

188

189 2.3 Analysis of solids

190 Solid carbon in both buffered and unbuffered runs has been characterized by scanning electron 191 microscopy, micro-Raman spectroscopy and synchrotron X-ray diffraction. Graphite and glass-like 192 carbon were analyzed both as starting materials and after quench from run P-T conditions. 193 Quantitative analyses and back-scattered electron imaging of the experimental products were 194 performed to check the integrity of the buffering mineral assemblages, using a JEOL 8200 195 wavelength-dispersive (WDS) electron microprobe, at 15 kV accelerating potential, 5 nA sample 196 current and 1 µm beam size. Standards used were favalite (Fe), niccolite (Ni) and grossular (Si). A 197 counting time of 30 s (10 s background) was used for all the elements. Secondary electron imaging 198 of pyrolytic carbon has been performed at 15 kV and 0.05 nA sample current. 199 Micro-Raman spectra were acquired using the integrated micro/macro-Raman LABRAM HRVIS

200 (Horiba Jobin Yvon Instruments) of the Interdepartmental Center "G. Scansetti" (Department of

201 Earth Sciences, University of Torino, Italy). Excitation lines at 532 nm (solid-state Nd laser and 80

202 mW of emission power) were used with Edge filter and a grating of 600 grooves/mm. Calibration

was performed using the 520.6 cm^{-1} Si band. The laser power on the sample was set upon the

- 204 measured materials at 2 mW by the addition of filters. Acquisition times were set at 25 s for 3
- 205 accumulations with a laser spot of 2 μ m.
- 206 X-ray Powder diffraction (XRD) data were obtained at XRD1 beamline (Elettra, Trieste, Italy). The
- samples were placed in glass capillaries and mounted onto the head of the Huber Kappa
- 208 Goniometer installed at the beamline. The data were collected with a monochromatic radiation
- 209 (λ =0.7000 Å), using a Dectris Pilatus 2M detector. The beam size at the sample was of 0.2×0.7 mm
- and the calibration of the set-up geometry was checked with a LaB₆ pattern. Successively, the
- 211 images were integrated with Fit2D software package (Hammersley et al., 1995; Hammersley, 1997)
- and analyzed using the GSAS EXPGUI software (Toby, 2001).
- 213

214 2.4 Analysis of volatiles

- 215 For the analysis of quenched volatiles in the capsules (Table 1) we employed the capsule-piercing 216 technique (Tiraboschi et al., 2016). Volatiles, heated to $T \approx 80^{\circ}$ C to allow liquid water to be 217 converted into water vapor, were extracted from the capsules in a Teflon reactor and conveyed to a 218 quadrupole mass spectrometer (EXXTORR 0–200 amu, Mod. XT 200, equipped with secondary 219 electron multiplier) using ultrapure Ar as carrier gas. The pressure conditions in the reactor were 220 monitored through high-resolution sensor gauges (± 1 mbar precision). The temperature in the 221 reactor was monitored with K-type thermocouple. Gas mixtures of known compositions and 222 ultrapure water were utilized for the calibration of the quadrupole mass spectrometer. The technique 223 enabled retrieval of micromolar quantities of the volatiles H₂O, CO₂, CO, CH₄, H₂ and O₂ with 224 uncertainties of ~ 1 mol% (10 mol% for CO, due to the interference with atmospheric N_2 on the 28 225 m/z channel). The periodic analysis of test capsules filled with 1 mg of oxalic acid dihydrate, 226 thermally decomposed at 250°C to H₂O, CO, CO₂ and H₂ (Tiraboschi et al., 2016), ensures the 227 stability and the reproducibility of the analyses over time.
- 228

229 2.5 Thermodynamic modeling

- 230 The volatile composition of carbon-saturated COH fluids, and in particular their *X*CO₂
- 231 $[=CO_2/(H_2O+CO_2)_{molar}]$ in relatively oxidized systems, is dependent on the redox state of the system
- 232 (cf. the review of Tumiati and Malaspina, 2019), which can be controlled indirectly in experiments
- by fixing the hydrogen fugacity in double capsules (e.g., Eugster and Skippen, 1967).
- The fugacities of oxygen and hydrogen in the outer and in the inner capsules at NNO and FMQ
- conditions were calculated using the Perple_X package (Connolly, 2005), using the thermodynamic
- dataset of Holland and Powell (Holland and Powell, 1998) revised by the authors in 2004
- 237 (hp04ver.dat), the Perple_X water equation of state (EoS) "H–O HSMRK/MRK hybrid", and using

238 the Excel spreadsheet GFluid (Zhang and Duan, 2010) with the COH fluid EoS of Zhang and Duan 239 (2009), which is based on the equilibrium constants $K_{\rm P}$ of the following reactions:

240	$C + O_2 = CO_2$	(Eq. 6)
241	$C + 0.5 O_2 = CO$	(Eq. 7)
242	$H_2 + 0.5 O_2 = H_2O$	(Eq. 8)
243	$C + 2 H_2 = CH_4$	(Eq. 9)

Tumiati et al. (2017) showed that the composition of graphite-saturated COH fluids at 1-3 GPa and 244 800°C are reproduced best using a modified version of Zhang and Duan's model, implemented with 245 246 fH_2 coefficients (γH_2) changing dynamically as a function of X(O) [= $O_2/(H_2+O_2)_{molar}$], taken from the EoS of Connolly and Cesare (1993). Therefore, in this study we refer to this modified model to 247 248 i) predict the composition of buffered fluids in equilibrium with crystalline graphite (Table 2); ii) 249 compare it with the measured composition of fluids in equilibrium with either crystalline graphite 250 or glass-like carbon (Fig. 2; Table 1) and iii) retrieve the difference in free energy between graphite 251 and glass-like carbon (ΔG), by iteratively changing the Zhang and Duan's K_{PS} (cf. Eqs. 4–7) to 252 account for the measured XCO_2 of fluids in equilibrium with the latter phase (Table 3). The experimental ΔG was compared with the ΔG predicted by thermodynamic modeling, performed 253 254 with thermodynamic parameters retrieved from literature added to the database of Holland and Powell (hp04ver.dat) (Table 4), thus making feasible with Perple X the calculation of G–P and log 255 256 fO_{2} -(P, T) phase diagrams involving glass-like carbon in addition to crystalline graphite. 257 Recent studies outlined the importance of non-volatile charged carbon species (not detectable with 258 the capsule-piercing technique) and/or organic dissolved compounds in high-pressure COH fluids at certain P-T-fO₂-pH conditions (Sverjensky et al., 2014; Pan and Galli, 2016; Tiraboschi et al., 259 2018; Huang and Sverjensky, 2019). In particular, Sverjensky et al. (2014) showed that at high pH 260 261 values aqueous bicarbonate and carbonate species become dominant instead of molecular CO₂ and 262 CH₄. In addition, organic dissolved species (e.g., formates and acetates) may form at ultra-high 263 pressures (e.g., 5 GPa at 600°C in Sverjensky et al., 2014). Therefore, conventional thermodynamic 264 models (Connolly and Cesare, 1993; Zhang and Duan, 2009), which consider neutral molecular 265 species only, are not always adequate to predict carbon speciation in subduction fluids. It is an open 266 question whether the available thermodynamic models are still valid in the presence of COH fluid 267 immiscibility, suggested at P = 1.5-2.5 GPa and T = 600-700°C in graphite-saturated slightly saline 268 fluids (Li, 2016) and in low-temperature/high-pressure hydrocarbon fluids (Huang et al., 2017). In 269 particular, Li (2016) found at 2.5 GPa and 700°C at Re–ReO₂ redox conditions ($\approx \Delta FMQ = +2$) 270 mixed H₂O–CO₂ fluids in equilibrium with almost pure CO₂. As the capsule-piercing technique 271 used in our study only allows measurement of the bulk volatile components, it cannot be used to

272 investigate fluid immiscibility. However, on the basis of the results given in our study and in

- 273 previous works (Matveev et al., 1997; Carla Tiraboschi et al., 2016; Tumiati et al., 2017), the
- 274 conventional models are still able to predict the *bulk* composition (but not necessarily the
- $275 \qquad \text{speciation) of high-pressure fluids in terms of bulk CO_2/(H_2O+CO_2) and CH_4/(H_2O+CH_4), although a special of the special of th$
- 276 they could fail at certain $P-T-fO_2$ -pH conditions where species other than H₂O, CO_{2(aq)} and CH_{4(aq)}
- become dominant. In order to justify our experimental approach we used the Deep Earth Water
- thermodynamic model (Sverjensky et al., 2014) to confirm that molecular $CO_{2(aq)}$ is by far the
- dominant carbon-bearing fluid species at our experimental conditions, and to estimate the pH of thefluids (Table 2).
- 281
- **3. Results**
- 283
- 284 *3.1 Characterization of solid carbon*
- 285 3.1.1 Crystalline graphite
- 286 Back-scattered electron observation of graphite powder used as starting material did not reveal any 287 foreign material. The powder consists of homogeneous flakes with grain size of 10–100 µm 288 (Supplementary Fig. 2 a). X-ray powder diffraction showed that the powder is monomineralic and crystalline, with a sharp diffraction peak at $d_{002} = 3.3672$ Å (Supplementary Fig. 3 - graphite). 289 290 Fitted pattern showed that the averaged crystallite size of the sample is around 50 nm. Micro-Raman spectroscopy of unpolished graphite powder showed a sharp graphite (G) peak at 1600 cm⁻¹ 291 and only a little disorder (D) peak at 1350 cm⁻¹, confirming that this material is highly ordered 292 (Ferrari, 2007) (Supplementary Fig. 4). The G' peak (also called 2D in Ferrari, 2007) at about 2700 293 294 cm^{-1} is also well developed. Raman spectra of polished graphite showed a marked increase of the D peak owing to mechanical modifications during the polishing (Pasteris, 1989), and was therefore 295 296 not considered. Ouenched graphite samples did not show any evidence of substantial modification 297 with respect to starting graphite powder, within the uncertainties caused by polishing.
- 298
- 299 3.1.2 Glass-like carbon
- 300 Glass-like carbon spheres used as starting material appeared homogenous at the scanning electron
- 301 microscope. They are about 100 µm in size, and show sparse closed porosity (Supplementary Fig. 2
- b). X-ray powder diffraction of the spherical powder showed broad peaks, typical of amorphous
- 303 materials (Supplementary Fig. 3). Micro-Raman spectra show broad G and D peaks, which
- 304 characterize amorphous sp^2 carbon (cf. Ferrari and Robertson, 2001), with D peak > G peak (Fig.

305 2). The *G*' peak is poorly developed. Glass-like carbon retrieved after runs showed Raman spectra
306 identical to the starting material.

307

308 3.1.3 Char from high-pressure pyrolysis of anhydrous glucose

309 Secondary electron imaging of solid carbon formed by decomposition of glucose (char) at 1–3 GPa 310 and 800°C show microtextures that are dependent on the run duration. In the 12-h run at 1 GPa, 311 char comes as a loose spherical powder with an average grain size of 1 µm (Figs. 3 a, b). In the 24-h 312 runs at 1 and 3 GPa, spherical elements are not visible, and char is characterized by a glass-like 313 appearance with conchoidal fracture and absence of cleavage (Figs. 3 c-f). X-ray powder 314 diffractions of char (Supplementary Fig. 3) show that the diffraction angle of the most intense peak is lower compared to that of crystalline graphite, while the full width at half maximum (FWHM) is 315 316 higher, meaning a decline of the crystallite size. The crystallite size of char synthesized at the 317 investigated high-pressure conditions is of about 2-4 nm typical of nano-crystalline materials. The 318 Raman spectra of char synthesized at 1 and 3 GPa in 24-h long runs are very similar, with broad G 319 and D peaks (D > G) (Fig. 4). Compared to glass-like carbon, the D peak is slightly higher, but 320 broadening of the peaks is a little less pronounced. The G' peak is comparable to that of glass-like 321 carbon. The Raman spectrum of char synthesized at 1 GPa in the shorter 12-h run is characterized 322 by similar G and D peaks, but remarkably higher G' peak (Supplementary Fig. 5), which

- 323 characterizes crystalline graphite (Fig. 4).
- 324

325 *3.2 Characterization of the fluid phase*

Volatiles in all the experimental runs have been measured by means of the capsule-piercing technique (see details in Tiraboschi et al., 2016) (Table 1; Fig. 5; Supplementary Fig. 6). Total measured volatiles range from 5.51 to 49.5 micromoles in double capsules where fluids were interacting with either crystalline graphite or glass-like carbon, and from 23.21 to 60.74 micromoles in single capsules where fluids were produced by complete dehydration of anhydrous glucose due to the reaction:

$$332 CH_2O = C + H_2O (Eq. 10)$$

- 333
- 334 3.2.1 Buffered COH fluids interacting with crystalline graphite
- 335 Fluids synthesized at 1–3 GPa and 800 $^{\circ}$ C contain mostly H₂O and CO₂, with only traces of CO,
- 336 CH₄ and H₂ (typically close to or below the detection limit). At P = 1 GPa, XCO_2 (= CO₂/H₂O+CO₂
- 337 molar) ranges from $0.674 \pm 0.007 (fH_2^{FMQ})$ to $0.731 \pm 0.008 (fH_2^{NNO})$. Because the FMQ and NNO
- buffers cross at about 1.5 GPa (cf. Supplementary Fig. 1), meaning that the NNO buffer is more

- oxidizing than FMQ at P < 1.5 GPa and less oxidizing at P > 1.5 GPa, fluids at P = 3 GPa buffered
- by NNO are expected to show lower *X*CO₂ compared to fluids buffered by FMQ (Table 2). At
- fH_2^{FMQ} , the run COH76, containing oxalic acid dihydrate (OAD) as starting source of fluid, displays
- 342 $XCO_2 = 0.74 \pm 0.02$, and run COH105, containing water instead of OAD, displays a similar XCO_2
- of 0.714 ± 0.05 . Instead, at fH_2^{NNO} conditions the fluid phase is dominated by water, resulting an
- 344 XCO_2 of 0.094 ± 0.001 in run COH103.
- 345
- 346 3.2.2 Buffered COH fluids interacting with glass-like carbon
- 347 Fluids interacting with glass-like carbon at 1–3 GPa and 800°C, buffered by either FMQ or NNO,
- 348 contain only traces of CO, CH₄ and H₂, as in the case of graphite-saturated fluids (Table 1).
- 349 However, the fluids with glasslike carbon display different *X*CO₂ ratios to the graphite-saturated
- ones (Fig. 5). In particular, fluids at 1 GPa are display a higher CO₂ content, and are characterized
- 351 by XCO₂ of 0.801 ± 0.009 at fH_2^{FMQ} and 0.850 ± 0.006 at fH_2^{NNO} , corresponding to a higher content
- 352 of +19 and +16 mol% CO₂, respectively, if compared to graphite-saturated fluids displaying XCO₂
- 353 = 0.674 at fH_2^{FMQ} and 0.731 ± at fH_2^{NNO} (see above). However, the higher CO₂ compared with
- 354 graphite-saturated fluids declines dramatically at 3 GPa. At this pressure, fluids saturated with
- 355 glass-like carbon buffered at fH_2^{NNO} show indeed a XCO_2 of 0.099 ± 0.001, corresponding to only
- $+5 \text{ mol}\% \text{ CO}_2$ higher content relative to fluids saturated with graphite displaying $X \text{CO}_2 = 0.094$.
- Fluids buffered at fH_2^{FMQ} conditions show CO₂ content that is even lower than that characterizing
- 358 graphite-saturated fluids. In fact, run COH112 is characterized by $XCO_2 = 0.57 \pm 0.01 (0.50 \pm 0.03)$
- in run COH108, with H₂O instead of OAD as starting source of fluid), corresponding to a lower
- 360 CO₂ content of $-24 \mod \%$ CO₂ relative to fluids saturated with graphite (*X*CO₂ = 0.74; see above).
- 361
- 362 3.2.3 Unbuffered COH fluids generated by high-pressure thermal decomposition of glucose
- 363 Fluids produced by high-pressure dehydration of anhydrous glucose display variable compositions,
- 364 changing as a function of run time and pressure conditions (Table 1; Supplementary Fig. 6). Fluids
- 365 at 1 GPa show high contents of CH₄, especially in run COH122 characterized by a relatively short
- 366 runtime of 12 h. In this run, XCO_2 is 0.072 ± 0.001 and XCH_4 [=CH₄/(H₂O+CH₄)_{molar}] = 0.261.
- 367 Fluid in run COH124, characterized by a longer duration of 24 h, displays a higher XCO_2 of 0.240 ±
- 368 0.003 and a lower CH₄ content (XCH₄ = 0.056). Fluid at 3 GPa (24 h) is nearly pure water with
- $369 \quad XCO_2 = 0.0098 \pm 0.0003 \text{ and } XCH_4 = 0.001.$
- 370

4.Discussion

372 4.1 Comparison between experimental results and available thermodynamic models of fluids in

373 equilibrium with crystalline graphite

- 374 The measured compositions of COH fluids interacting with crystalline graphite at 1 GPa and 800°C
- 375 (Table 1) is in excellent agreement, in both FMQ- and NNO-buffered experiments, with the
- 376 compositions predicted at equilibrium conditions by the modified thermodynamic model of Zhang
- and Duan (2009) (Table 2; see details of the model in Section 2.5). At 3 GPa, while the fH_2^{FMQ} -
- buffered fluid matches the composition predicted by the model, the measured composition of the
- 379 fluid buffered externally by NNO is characterized by a lower content of CO₂. The measured *X*CO₂
- 380 would be consistent with an inner-capsule $\log fO_2$ of -12.56, instead of the predicted value of -
- 381 11.99 (Table 2). At this stage, we can only speculate that this could be ascribed to uncertainties
- associated to the nickel-nickel oxide buffer (c.f. O'Neill and Pownceby, 1993), which could affect
- 383 the estimates of fO_2 and fH_2 imposed by the NNO buffer at 3 GPa.
- 384 Some recent studies underlined the importance of the variable pH in governing the abundance of
- 385 dissolved organic species and charges species (e.g., bicarbonates, carbonates) instead of molecular
- 386 species (Sverjensky et al., 2014; Pan and Galli, 2016). Therefore, we used the Deep Earth Water
- 387 (DEW) model (Sverjensky et al., 2014) to draw pH vs fO₂ diagrams showing the loci of points
- 388 where graphite is stable in COH fluids at saturation conditions at 1GPa–800°C (Fig. 6 a) and 3
- 389 GPa-800°C (Fig. 6 b) (black solid lines). The oxygen fugacities expected in COH fluids buffered at
- 390 fH_2^{FMQ} and fH_2^{NNO} conditions are shown for reference. The intersection of these oxygen fugacities
- 391 with the graphite-saturation curve represents the investigated experimental conditions (black dots),
- and thus provides an estimation of the pH value expected in the synthesized fluids, i.e. 3.5–3.6 at 1
- 393 GPa-800°C (neutral pH = 4.02) and 2.22 at 3 GPa-800°C (neutral pH = 3.09). The model predicts
- that at the investigated $P-T-fO_2-fH_2$ conditions the equilibrium between graphite and COH fluids is
- thus reached in the $CO_{2(aq)}$ stability field and at acidic conditions, which prevent the stability of
- 396 carbonate and bicarbonate ions. $CO_{2(aq)}$ is therefore expected to be by far the dominant dissolved
- 397 carbon-bearing species, while other C–O–H species should occur in very minor amount, the most
- abundant being ethane ($\approx 10^{-4}$ mol%) at 1 GPa–800°C and formic acid ($\approx 10^{-3}$ mol%) at 3 GPa–
- 399 800°C. Other organic and/or charged species display even lower abundances. This validates the
- 400 methodological approach of this study, aiming to retrieve the composition of quenched fluids in
- 401 terms of volatile molecular species, assuming that they represent the speciation at run conditions.
- 402 As a corollary, the diagrams in Figure 6 suggest that when redox conditions are imposed on a
 403 petrological system (redox-buffered systems), the pH becomes merely a dependent variable, as long
- 404 as graphite-saturation conditions persist. Conversely, in pH-buffered systems, the redox state would
- 405 be controlled by pH.

406

428

- 407 4.2 Experimental fluids in equilibrium with glass-like carbon and retrieval of its thermodynamic 408 properties at high-pressure conditions
- 409 The CO₂ content of fluids interacting with glass-like carbon is different with respect to the CO₂
- 410 content of fluids in equilibrium with crystalline graphite (Fig. 5; Table 1). Fluids at 1 GPa are
- 411 considerably enriched in CO₂, while fluids at 3 GPa are only slightly enriched or even depleted with
- respect to graphite-saturated fluids. The observed difference is +19 mol% CO₂ at 1 GPa- fH_2^{FMQ} , 412
- +16 mol% CO₂ at 1 GPa- fH_2^{NNO} , +5 mol% at 3 GPa- fH_2^{NNO} and -24 mol% CO₂ at 3 GPa- fH_2^{FMQ} . 413
- 414 These different fluid compositions are uniquely ascribable to difference between the
- 415 thermodynamic properties of glass-like carbon and crystalline graphite. Taking as example
- Equation 6 ($C + O_2 = CO_2$), the equilibrium constant of the reaction is: 416

417
$$K_p(T) = \frac{|CO_2|}{|O_2|},$$
 (Eq. 11),

- 418 where brackets indicate equilibrium concentrations of fluid phases over solid carbon.
- 419 At 1 GPa and 800°C, involving perfectly crystalline graphite, the equilibrium constant of the reaction following the thermodynamic model of Zhang and Duan (2009) is 3.416×10^{19} (ln K_p = 420 44.98). By changing K_p , the fluid CO₂ content predicted by the model changes accordingly. By 421 422 means of iterative calculation, it is possible to find a K_p that fits the measured CO₂ value in runs 423 where fluids reacted with glass-like carbon instead of graphite at the same P-T conditions. The 424 equilibrium constant of a heterogeneous reaction with the participation of glass-like carbon (gl) 425 instead of graphite (graph) can be expressed as:

426
$$K_{p\,\text{gl}}(T) = K_{p\,\text{graph}}(T) \exp\left(\frac{\Delta G(T)}{RT}\right)$$
 (Eq. 12)

where $\Delta G(T)$ is the difference in Gibbs free energy between glass-like carbon $G_{gl}(T)$ and graphite-427

428
$$G_{\text{graph}}(T)$$
, and *R* is the gas constant. This difference can be made explicit, resulting in the equation:
429 $\Delta G(T) = RT \ln \left(\frac{K_{p\,\text{gl}}(T)}{K_{p\,\text{graph}}(T)} \right)$ (Eq. 13)

430 As this difference in Gibbs free energy at fixed $P-T-fH_2$ must be the same for all the reactions 6, 7 431 and 9, by means of simulation analysis performed with the Solver tool in the Excel spreadsheet 432 provided by Zhang and Duan (2010) we changed iteratively all the pertaining equilibrium constants 433 simultaneously, imposing the mathematical constraint that the resulting ΔG is identical for all the 434 equations, until the model converges to the measured XCO₂. The retrieved ΔG and the equilibrium 435 constants retrieved by simulation analysis at 1 and 3 GPa are shown in Table 3. ΔG is almost 436 coincident in runs performed at NNO and FMQ conditions at 1 GPa-800°C, where fluids 437 equilibrated with glass-like carbon contain higher CO₂ fractions compared to graphite-saturated fluids. At these *P*–*T* conditions, ΔG is equal to +1.6(1) at fH_2^{NNO} and +1.7(1) kJ/mol at fH_2^{FMQ} . The 438

- 439 difference in Gibbs free energy between graphite and glass-like at 1 GPa agree with previous
- studies performed at room pressure, where ΔG is +1.8 kJ/mol at 800°C (Guencheva et al., 2001;
- 441 Gutzow et al., 2005). At 3 GPa and 800°C, fluids in equilibrium with glass-like carbon buffered at
- 442 fH_2^{NNO} conditions display a slightly higher CO₂ content compared to graphite-saturated fluids, and
- 443 ΔG is equal to +0.5(1) kJ/mol. Fluids buffered at fH_2^{FMQ} conditions show a CO₂ content which is
- 444 even lower than fluids in equilibrium with graphite, corresponding to a negative ΔG of -2.3(2)
- 445 kJ/mol.
- 446

447 4.3 Thermodynamic modeling of glass-like carbon and implications for its stability towards 448 graphite and diamond

We are not aware of previous Gibbs free energy of glass-like carbon at high pressures, even though
layers of glass-like carbon spheres have long been employed in experimental petrology as melt

451 traps at pressure conditions ranging from 1 to 5 GPa (Robinson et al., 1998; Wasylenki, 2003;

- 452 Dasgupta et al., 2005; Falloon et al., 2008; Spandler et al., 2008). Therefore, we derived the G(P, T)
- 453 curve of glass-like carbon at pressures up to 3 GPa (Fig. 7), by using the following thermodynamic
 454 parameters (Table 4; Supplementary Table 1):
- the standard Gibbs free energy of formation, retrieved from the G-T data of Guencheva et al. (2001) and Gutzow et al. (2005) at room pressure and T = 298 K ($\Delta_f G^0 = 2.357$ kJ/mol);
- the standard entropy (S_{298}^{0}) , retrieved by fitting the G-T data at room pressure of Gutzow et al. (2005). The retrieved $S_{298}^{0} = 6.6$ J/mol/K agrees well with the standard entropy of 6.2 J/mol/K resulting from the integration of low-temperature heat capacity $[C_P(T)]$ measurements (Cappelletti et al., 2018), following:

 $S_{298}^0 = S_0^0 + \int_0^{298} \left[\frac{C_P^0(T)}{T} \right] dT$

(Eq. 14),

taking into account that $S_{0,i}^{0}$, i.e. the residual entropy at 0 K, is non-zero in amorphous solids; 462 the following parameters were assumed to be the same as for graphite (Holland and Powell, 463 • 1998; Day, 2012), as they have been reported to be very similar in glasslike carbon: i) the 464 thermal expansion (cf. Cowlard and Lewis, 1967), ii) the coefficients of the heat capacity 465 466 function (cf. Takahashi and Westrum, 1970; Yokoyama et al., 1971) and iii) the standard molar volume, relying on the fact that the disordered multilayer graphene component in 467 glass-like carbon has a density close to that of graphite (2.25 g/cm^3) and the interplanar d-468 spacings are in broad agreement with those of graphite (Supplementary Fig. 3; cf. also Zhao 469 et al., 2015; Hu et al., 2017). Models with molar volume increased by 50%, reflecting the 470 apparently lower bulk density of glass-like carbon (≈ 1.5 g/cm³; Cowlard and Lewis, 1967; 471

Zhao et al., 2015) imputable to the high closed porosity, are also provided for comparison in
Figure 7 b-c.

- the isothermal bulk modulus under standard conditions (K^0) and its first pressure derivative 474 • (K) were retrieved by linear regression of the data of type-I glass-like carbon reported by 475 476 Zhao et al. (2015) (green dots in Fig. 7 a), where K^0 is the intercept at 0.0001 GPa (1 bar) at 25°C and K' is the slope of the K–P curves. However, the estimation of glass-like carbon K^0 477 478 and K' is not straightforward. Glass-like carbon displays a very high compressibility 479 compared to graphite (cf. bulk modulus data in Day, 2012; Table 4), with a more than halved K^0 decreasing abnormally with increasing pressure up to 1 GPa (i.e., negative K' = -480 2.4, corresponding to a K^0 intercept of 15 GPa; purple thick dashed lines in Fig. 7), followed 481 482 by a reversal to positive pressure dependence, accounting to a K' of 2.9 in the 1–3 GPa pressure range (extrapolated $K^0 = 9$ GPa; red thick lines in Fig. 7; preferred model as it is 483 consistent with the pressures investigated in this study and with K^0 of other forms of non-484 graphitic sp^2 -type carbon; cf. fullerene in Sundqvist and Olabi, 2016) and K = 7.8 at 485 pressures above 3 GPa (extrapolated $K^0 = -6$ GPa; grey dashed lines in Fig. 7). In figure 7b, 486 we show how the choice of different K^0 and K' values affects the $G(P, 800^{\circ}\text{C})$ of glass-like 487 488 carbon with pressure increasing from 0 to 5 GPa. The predicted difference in Gibbs free 489 energy at 800°C versus pressure between glass-like carbon and graphite (ΔG) using the 490 different models is also shown (Fig. 7c).
- 491

492 At 800°C, the G–P curves of glass-like carbon (preferred model) and of graphite cross at 3.4 GPa, 493 showing a continuous decline of ΔG with increasing pressure, which is 1.5 kJ/mol at 1 GPa and 0.29 kJ/mol at 3 GPa. These ΔG values match well our experimental results at fH_2^{NNO} conditions 494 (Table 3). Runs buffered at fH_2^{FMQ} conditions agree with the model at 1 GPa, while at 3 GPa the 495 decline observed experimentally, characterized by a negative ΔG , is more pronounced compared to 496 497 the model. Models assuming a molar volume coherent with the apparent bulk density of glass-like carbon (1.5 g/cm³) are inconsistent with experimental results, as they show ΔG values that would 498 499 increase by increasing pressures (red dashed line in Fig. 7). Negative ΔG values at 3 GPa can be reproduced only by models that assume very high compressibility ($K^0 = 2$ GPa; orange dashed line 500 501 in Figure 7). In this case, however, the intersection point with graphite is shifted at lower pressures 502 (\approx 1.5 GPa), so that the *G*-*P* curve of glass-like carbon would always lie below that of diamond, 503 meaning that, above the intersection point, glass-like carbon would be the stable form of carbon relative graphite and diamond, which is unlikely. On the other hand, the preferred model does 504 505 intersect the G-P curve of diamond, although it occurs at 4.2 GPa, i.e., above the intersection point

- 506 with graphite at 3.4 GPa and above the graphite-diamond transition at 3.7 GPa, too. This would 507 imply that in the pressure window 3.4–3.7 glass-like carbon could be more stable than graphite and 508 between 3.7 and 4.2 it could even be more stable than diamond (yellow field in Figure 7). However, 509 because of the similar slopes of glass-like carbon- and graphite G-P curves, small fluctuations in the chosen K^0 can result in large uncertainties on the position of the intersection point with graphite 510 511 relative to the graphite-diamond transition. In fact, a glass-like carbon G-P curve calculated assuming $K^0 = 12$ GPa (red dashed lines in Figure 7) instead of $K^0 = 9$ GPa (preferred model) would 512 intersect the G-P curve of diamond at 3.4 GPa and the G-P curve of graphite at 4 GPa, implying 513 514 metastability of glass-like carbon at any pressure towards both graphite and diamond, without 515 having a marked effect on the predicted decline of ΔG with increasing pressure. 516 The estimated thermodynamic parameters of glass-like carbon allow also the calculation of fO_2-T 517 phase diagrams at high pressures. In Figure 8, the boundaries C–CO₂ (often referred to as CCO) and 518 enstatite + magnesite = olivine + C (EMOG) are compared considering crystalline graphite and glass-like carbon. At 1 GPa, CCO_(glass-like carbon) and EMOG_(glass-like carbon) are shifted below 519 $CCO_{(graphite)}$ and $EMOG_{(graphite)}$ by 0.08 log fO_2 units. At 3.5 GPa, the possible reversed stability of 520 521 glass-like carbon over graphite is shown, with CCO_(glass-like carbon) and EMOG_(glass-like carbon) located above CCO_(graphite) and EMOG_(graphite) by 0.01 log *f*O₂ units. In both cases, the difference of CCO 522 523 curves pertaining to the two considered types of carbon is very small in terms of absolute fO_2 524 values. Nevertheless, as shown by our experimental results, these differences are enough to induce 525 large variations in the composition of COH fluids interacting with crystalline versus glass-like solid 526 carbon.
- 527

528 4.4 Comparison of glass-like carbon and glucose-derived char

529 Char synthesized for 24 h at 1 and 3 GPa, 800°C starting from glucose displays conchoidal 530 fractures (Fig. 3 c-f) and absence of cleavage, similar to glass-like carbon. Moreover, char and 531 glass-like carbon show broadly similar Raman spectra (Fig. 4), X-ray diffraction patterns (Supplementary Fig. 3) and nano-sized crystallite dimensions, supporting our experimental strategy 532 533 in which glass-like carbon was chosen as the best analogue for disorganized, poorly crystalline 534 carbonaceous organic matter. In this study, we observed a marked difference in Raman spectra 535 acquired for char synthesized in 12-h and 24-h runs performed at 1 GPa and 800°C. In particular, the graphite G' peak at about 2700 cm⁻¹, well developed in the 12-h run, becomes only hinted in the 536 537 24-h run, suggesting that the size of the graphite crystallites decreased with time. This implies, in agreement with Beyssac et al. (2003), that the structure of char, although somewhat similar to glass-538 539 like carbon, is not stable at high-pressure/high-temperature conditions, and so this type of materials

540 would be not suitable for long experiments at static equilibrium conditions. In this study, however, 541 we observed transient near-equilibrium conditions between char and aqueous fluids generated by 542 dehydration of glucose. Ideally, in perfectly closed systems, the thermal decomposition of 543 anhydrous glucose (CH₂O) should produce carbon and pure water; however, this has been observed 544 only in the run performed at 3 GPa and 800°C, containing almost pure water ($\approx 99 \text{ mol}\% \text{ H}_2\text{O}$). In fact, dehydration of glucose at 1 GPa produced CH₄-rich, CO₂-bearing fluids after 12 h and CO₂-545 546 rich, CH₄-bearing fluids after 24 h. However, if the measured composition of these fluids is 547 compared with those predicted by the modified model of Zhang and Duan (2009) implemented with 548 the retrieved glass-like carbon equilibrium constants, both XCO₂ and XCH₄ ratios match redox conditions of $\Delta FMQ = -1.8$, suggesting near-equilibrium conditions and a close thermodynamic 549 550 affinity between glass-like carbon and char. In longer runs (24 h) at the same conditions of 1 GPa 551 and 800°C, measured XCO₂ and XCH₄ ratios correspond also in this case to consistent redox states 552 $(\Delta FMQ = -1.2, \text{ if } XCO_2 \text{ is considered}; \Delta FMQ = -1.0 \text{ if } XCH_4 \text{ is taken into account})$. The measured 553 fluid compositions suggest that the interaction with char in unbuffered single gold capsules evolves 554 over time from reduced conditions, likely buffered by glucose (Hawkins, 1929; Kunz et al., 2011), 555 to more oxidized conditions, likely constrained by the MgO-graphite-NaCl-Pyrex furnace 556 assembly (close to FMQ; cf. Olafsson and Eggler, 1983). In agreement with previous studies (e.g., Truckenbrodt et al., 1997; Truckenbrodt and Johannes, 1999; Matjuschkin et al., 2014), we 557 558 observed that unbuffered single gold capsules cannot therefore be considered systems perfectly 559 closed to H₂O and H₂, so the buffering of the system (for instance using double capsules) is 560 mandatory to constrain the H₂ (and, indirectly, the O₂) chemical potential.

561

562 4.5 Implications for organic matter dissolution at subduction zones

563 Organic matter can be an important constituent of oceanic sediments (Mayer et al., 1992), and on 564 average it accounts for less than 1 wt.% (Kelemen and Manning, 2015). Nevertheless, organic 565 matter in deep-sea fans can dominate the carbon input flux at some margins (Plank and Manning, 566 2019). The proportion of organic to inorganic carbon (i.e., marine carbonates) subducted globally is 567 about 20% (Plank and Manning, 2019) and the total amount of organic carbon subducted in modern 568 active subduction zones is estimated >11 Mt C/y (Clift, 2017). Once subducted and heated, poorly 569 organized organic matter is progressively transformed into crystalline graphite through a multitude 570 of intermediate stages generally referred to disordered graphitic carbon (Beyssac and Rumble, 571 2014; Buseck and Beyssac, 2014). In addition, graphite can also form by reduction of carbonates 572 during subduction (Galvez et al., 2013; Vitale-Brovarone et al., 2017) and by precipitation from 573 subduction C-O-H fluids (Luque et al., 1998).

574

- 575 The oxidation susceptibility and therefore the dissolution of graphite in aqueous fluids varies as a 576 function of P, T and fO₂ conditions (e.g., Connolly, 1995; Tumiati and Malaspina, 2019). In 577 general, low-temperature and high-pressure conditions characterizing subduction zones are thought 578 to promote the stability of graphite, thus fluids interacting with this mineral should contain very low 579 amounts of carbon and are essentially nearly pure water (Schmidt and Poli, 2013). However, we 580 show in Figure 9 a that this is expected only at forearc conditions. In fact, at *P*–*T* conditions 581 characterizing the slab surface (Syracuse et al., 2010), nearly pure water is expected only up to 582 around 2 GPa and 450°C. At greater depths, graphite-saturated fluids become progressively more enriched in CO₂ as FMQ and CCO buffers get very close (Fig. 9 a), with maximum CO₂ contents 583 $(XCO_2 = 0.55)$ at subarc conditions (3 GPa-700°C), where the two buffers nearly converge. These 584 585 fluid compositions are predicted assuming a perfectly crystalline and ordered state of graphite. 586 Several previous studies suggested that the poorly ordered graphite might behave differently, in 587 particular showing a higher solubility in aqueous fluids (Ziegenbein and Johannes, 1980; Connolly, 588 1995; Luque et al., 1998). Although our investigated *P*–*T* conditions are not strictly comparable 589 with "normal" subduction regimes, especially at low pressures, we demonstrate that glass-like 590 carbon is characterized by a marked difference in free energy ($\approx 2 \text{ kJ/mol}$) with respect to 591 crystalline graphite at low pressures, diminishing with increasing pressures and with a possible sign 592 reversal close to graphite-diamond transition (located at 3.6 GPa and 760°C in Fig. 9 a). Because 593 we showed that the difference in free energy between graphite and glass-like carbon is 594 predominantly due to their different compressibility behavior, the relative difference in carbon 595 solubility is only slightly depending on temperature. Therefore, we can speculate that similar 596 differences in fluid carbon content (in terms of $CO_{2(aq)}$, but possibly also of $CH_{4(aq)}$, HCO^{3-} , CO_{3}^{2-} 597 and other dissolved C–O–H species at appropriate $P-T-fO_2-fH_2$ –pH conditions) can be expected at 598 lower temperatures characterizing the subduction zones and in particular the subduction surface 599 where sediments containing organic matter can be abundant. 600 This implies that disordered graphitic carbon is more prone to oxidation if compared with ordered 601 crystalline graphite down to about 110 km, resulting in fluids that are enriched in CO_2 compared to 602 current estimates: the lower the pressure the higher the differential dissolution susceptibility. 603 Therefore, because it is unlikely that organic carbon can persist in its disordered state at great 604 depths and consequent relatively high temperatures (Beyssac et al., 2002), and because the 605 difference in free energy is higher at low pressures, we infer that the most important effect on
- 606 carbon recycling concerns the most shallow levels of subduction zones, where disordered organic
- 607 carbon could be far more reactive than expected for graphite towards aqueous fluids (Fig. 9 b).

608 Obviously an important source of uncertainty arises from the assumption that glass-like carbon can 609 be considered an analogue of disordered natural carbonaceous matter. Carbon materials are very complex because their properties change extensively with structural defects and impurities, and the 610 type of hybridization, in particular sp^2 (graphite-like carbon) vs. sp^3 (diamond-like carbon) (e.g., 611 Robertson, 2002; Langenhorst and Campione, 2019). Moreover, the activity and the reactivity of 612 these materials can be modified by bringing them to nanosize dimensions. For instance, Guencheva 613 614 et al. (2001) and Gutzow et al. (2005) showed that nanodispersed (10 nm) glass-like carbon displays 615 a difference in Gibbs free energy compared to graphite of +12 kJ/mol at standard conditions, which 616 is substantially higher relative to bulk glass-like carbon (+2.4 kJ/mol; Table 4) and which would 617 result in dissolution susceptibility higher than that provided in our study. In addition, natural highly 618 disordered carbon, as synthetic amorphous and graphitic carbon, is likely characterized by a higher 619 kinetic reactivity because of the presence of active immobilized free radicals (dangling bonds) and defects which make these materials efficient catalysts (e.g., Jüntgen, 1986). The results provided in 620 621 our study therefore represent a first attempt to provide a conservative minimum estimate of the 622 enhanced dissolution of disordered carbon with respect to crystalline graphite in natural systems.

623

624 **5.** Conclusions

Well-ordered crystalline graphite and X-ray amorphous glass-like carbon display different 625 626 dissolution susceptibility in aqueous fluids, because of their different thermodynamic properties. On the basis of our experimental observations concerning this type of disordered sp^2 carbon, and aware 627 of the uncertainties arising from the choice of glass-like carbon as an analogue materials for 628 629 naturally occurring poorly organized carbon derived from the graphitization of organic matter, we 630 may speculate at this stage that also natural poorly organized graphitic carbon could behave in a 631 different manner compared to perfectly crystalline graphite. In particular, disordered carbonaceous 632 matter could be more prone to dissolve in aqueous fluids compared to well-crystallized graphite 633 especially at pressures corresponding to the forearc region of subduction zones. High fluxes of water coming from the dehydration of the down-going slab would therefore induce an effective 634 635 removal of organic matter from its sedimentary cover, prompting metasomatism of the mantle 636 wedge (Sieber et al., 2018) and contributing to the global deep carbon cycle. As for glass-like 637 carbon, the differential dissolution susceptibility declines with increasing pressure corresponding to 638 110 km depth, below which "disordered" carbon could be even less soluble than graphite. The 639 stability of glass-like carbon over diamond, although predicted by the suggested model in the 3.7-640 4.2 GPa pressure window, relies on thermodynamic parameters that are affected by uncertainties 641 that are currently unquantifiable. The occurrence of nano-crystalline disordered graphitic carbon

and amorphous sp^2 and sp^3 carbon has been indeed reported in micro- and nano-sized diamonds from Cignana Lake in the Western Alps (Frezzotti et al., 2014; Frezzotti, 2019) and glass-like X-ray amorphous carbon has been obtained experimentally at diamond-stable conditions (7.7 GPa and 1000°C; Yamaoka et al., 2002). However, more investigations are required to confirm this hypothesis.

647

648 Appendix: terminology used for the description of solid carbon

649 In this study, the terminology used for the description of solid carbon follows the recommendations650 of IUPAC (Fitzer et al., 1995).

- Graphite: an allotropic form of the element carbon consisting of layers of hexagonally
 arranged carbon atoms in a planar condensed ring system (graphene layers). The layers are
 stacked parallel to each other in a three-dimensional crystalline long-range order. The
 chemical bonds within the layers are covalent with *sp*² hybridization.
- Graphitic carbon: all varieties of substances consisting of the element carbon in the
 allotropic form of graphite irrespective of the presence of structural defects. The use of the
 term is justified if three-dimensional hexagonal long-range order can be detected in the
 material by diffraction methods, independent of the volume fraction and the homogeneity of
 distribution of such crystalline domains.
- 660 • Glass-like carbon: in this study we use this term although the commercial/trademark terms "glassy carbon" and "vitreous carbon" are still widely used in experimental petrology and 661 materials sciences papers. Glass-like carbon is agranular (i.e., homogenous microstructure 662 with structural elements undistinguishable by optical microscopy) and non-graphitizable 663 664 carbon (i.e., it does not convert into graphitic carbon upon heat treatment to 2500–3300 K) 665 with a very high isotropy. Although its structure is not comparable to silicate glasses, the fracture surfaces have a pseudo-glassy (conchoidal) appearance. It consists of curved two-666 dimensional structural elements (graphene layers, i.e., single carbon layers of the graphite 667 structure) dispersed in an X-ray amorphous matrix, but it does not exhibit dangling bonds 668 that characterize the so-called "amorphous carbon". In fact, the term "amorphous carbon" is 669 670 restricted to the description of carbon materials which, in addition to a lack of long-range crystalline order and to deviations of the interatomic distances with respect to graphite 671 672 lattice as well as to the diamond lattice, show deviations in the bond angles because of the 673 presence of dangling bonds. Amorphous carbon is disordered even on the atomic scale and have a fraction of sp^3 bonds ranging from a few ("graphite-like" structure) to almost 100% 674 ("diamond-like" structure) (Sundqvist and Olabi, 2016) 675

- Char: a solid decomposition product of a natural or synthetic organic material. In this study,
- 677 char is produced by carbonization (pyrolysis) of glucose. The term pyrolytic carbon has
- been avoided because it is restricted to carbon materials deposited from gaseous
- 679 hydrocarbon compounds by chemical vapor deposition.
- 680

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- 913

914 Tables

- 915 <u>Table 1</u>: Run table of the experiments and volatiles measured using the capsule-piercing technique.
- 916 <u>Table 2</u>: Thermodynamic modeling of graphite-saturated fluids buffered at the investigated P-T-917 *f*H₂ conditions.
- 918 <u>Table 3</u>: Equilibrium constants (K_P) and difference in Gibbs free energy of glassy carbon with 919 respect to crystalline graphite (ΔG), retrieved from measured fluid composition.
- 920 Table 4: Thermodynamic properties of graphite and glass carbon (preferred model) at different
- 921 pressures and temperatures. See also Supplementary Table 1 for Perple_X formatted version
- 922 including diamond.
- 923

924 Figure captions:

- 925 <u>Figure 1</u>: Experimental setup. (A, B): sketch of the double capsule system. The inner Au–Pd
- 926 capsule, permeable to H₂, contains oxalic acid dihydrate (OAD), which decomposes at $T > 200^{\circ}$ C
- 927 to CO₂, H₂O and H₂, and either crystalline graphite (A) of glass-like carbon (B). The outer Au
- 928 capsule contains the inner capsule and the fH_2 buffer, either fayalite-magnetite-quartz-H₂O or
- 929 nickel-nickel-oxide- H_2O . The fH_2 constrained by the buffer is expected to be homogenous in the
- 930 inner and in the outer capsule. (C): back-scattered electron image of the representative sample
- 931 COH62 (P = 1 GPa and T = 800°C) across the inner Au–Pd capsule boundary, showing glass-like
- 932 carbon spheres on the left and the fayalite-magnetite-quartz buffer on the right.
- 933 Figure 2: Micro-Raman spectra of glass-like carbon as starting material (unpolished: blue; polished:
- 934 purple) and as quenched product from 1 GPa–800°C (green) and from 3 GPa–800°C (red).
- 935 Figure 3: Secondary electron images of char synthesized from glucose at 800°C and high-pressure
- 936 conditions. (A, B): P = 1 GPa, runtime 12 h; (C, D): P = 1 GPa, runtime 24 h; (E, F): P = 3 GPa,
- 937 runtime 24 h.
- <u>Figure 4</u>: Raman spectra of quenched char (1 GPa: blue; 3 GPa: purple) compared with glass-like
 carbon (green) and crystalline graphite (red).
- 940 Figure 5: Fluid compositions at the investigated $P-T-fH_2$ measured by using the capsule-piercing
- 941 technique (C. Tiraboschi et al., 2016), plotted on ternary C–O–H diagrams. Green triangles: fluids
- 942 in equilibrium with crystalline graphite. Blue dots: fluids in equilibrium with glass-like carbon.
- 943 Yellow squares: composition of graphite-saturated fluids according to the modified model of Zhang
- and Duan (2009) (see text for details). Grey dots: analytical uncertainty cloud estimated by Monte
- 945 Carlo method using the standard deviations provided in Table 1.

- 946 <u>Figure 6</u>: $\log fO_2 vs$ pH diagrams at 1 GPa–800°C (A) and 3 GPa–800°C (B), calculated using the
- 947 Deep Earth Water model, showing the COH-fluid graphite saturation curve (thick black). Black 948 dots: experimental conditions at fH_2^{FMQ} and fH_2^{NNO} .
- 949 Figure 7: Thermodynamic properties of glass-like carbon. (A) glass-like carbon bulk modulus and
 950 its pressure dependence, compared with graphite. Measurements from Zhao et al. (2015) (green
- dots) are fitted with different K^0 and K'. The preferred model (red thick) assumes $K^0 = 9$ GPa and K'
- 952 = 2.9. See text for other details. (B) Gibbs free energy versus pressure at 800°C, calculated using
 953 different thermodynamic models of glassy carbon. Red arrows indicate the shift of the preferred
- 954 model assuming either an increased molar volume (i.e., lower density) or an increased
- 955 compressibility (lower K^0). (C) difference in Gibbs free energy between glass-like carbon and
- 956 graphite (ΔG), plotted as a function of pressure at $T = 800^{\circ}$ C. Red thick (preferred model): $K^0 = 9$
- 957 GPa, K = 2.9, density (d) = 2.2.5 g/cm³. Red dashed: $K^0 = 12$ GPa, K = 2.9, d = 2.25 g/cm³. Purple
- 958 dashed: $K^0 = 15$ GPa, K = -2.4, d = 2.25 g/cm³. Orange dashed: $K^0 = 0.1$ GPa, K = 6.3, d = 2.25
- 959 g/cm³. Green dashed: $K^0 = 9$ GPa, K' = 2.9, d = 1.5 g/cm³. Blue: diamond. Black thick: graphite.
- 960 Light yellow field: thermodynamic stability of glass-like carbon (preferred model) over graphite961 and diamond.
- 962 Figure 8: Calculated T-log fO_2 diagrams of the univariant equilibria C + O₂ = CO₂ (CCO) and
- $MgSiO_3 + MgCO_3 = Mg_2SiO_4 + C + O_2$ (EMOG) involving graphite and glass-like carbon close to 963 $T = 800^{\circ}$ C at P = 1 GPa and at P = 3.5 GPa, using the thermodynamic properties reported in Table 964 965 4. FMQ: fayalite-magnetite-quartz or ferrosilite (fs)-magnetite (mt)-coesite (coes) oxygen buffer. 966 Figure 9: Fate of organic matter in subduction. (A) Stability of COH fluids (grey field) calculated as a function of fO_2 along a subduction P-T gradient consistent with the average thermal model of slab 967 surface after Syracuse et al. (2010). COH fluids are stable between the two boundaries $C + O_2 =$ 968 CO_2 (CCO) and C + 2 H₂ = CH₄. At this scale, differences between graphite and amorphous carbon 969 970 are negligible, but according to the preferred model in Figure 7 glass-like carbon would be the 971 stable carbon polymorph at P > 3.4 GPa. The calculation of fluid isopleths ($XCO_2 = CO_2/H_2O+CO_2$; $XCH_4 = CH_4/H_2O+CH_4$) has been performed using the Perple X and the EoS of Connolly and 972 973 Cesare (1993). Reference buffers FMQ, hematite-magnetite (HM), wustite-magnetite (WM), iron-974 wustite (IW) and quartz-iron-fayalite (QIF) are shown for reference. Fluids in equilibrium with graphite buffered at FMQ conditions become increasingly enriched in CO₂, which reaches its 975 976 maximum concentration at about 100 km depth, where FMQ and CCO almost converge. (B)
- 977 Cartoon showing the fate of organic matter in subduction zones. Disordered organic matter
- 978 contained in marine sediments undergoes partial to complete graphitization by increasing
- 979 subduction temperature. Experimental results and thermodynamic models presented in this study

- 980 indicate that disordered carbon is more prone to oxidation with respect to crystalline carbon
- 981 especially at low pressures, characterizing the forearc region. At these depths, an intense flush of
- 982 water would be able to dissolve selectively disordered organic matter from the subducted
- 983 sediments, while graphite behaves in a more refractory manner. This differential dissolution
- 984 susceptibility is expected to progressively decrease as subduction proceeds, vanishing at about 100
- 985 km depth where the difference in free energy between graphite and disordered carbon tends to zero.
- 986

987 Electronic Annex

- 988 <u>Supplementary Table 1:</u> thermodynamic data for graphite (gph), glass-like carbon type I (GC;
- 989 preferred model) and diamond (diam), formatted for Perple_X package
- 990 (http://www.perplex.ethz.ch) and to be copy-pasted in the used thermodynamic database file. Heat-
- 991 capacity function and thermal expansion of glass-like carbon assumed identical to graphite.
- 992 Diamond data after Day (2012). See also Table 4 and text for details.
- 993 <u>Supplementary Figure 1</u>: Thermodynamic model of FMQ and NNO buffers, calculated with
- 994 Perple_X and the hp04ver.dat database.
- 995 <u>Supplementary Figure 2</u>: Electron microscope images of the starting materials. (a) back-scattered
- 996 electron image of graphite; (b) back-scattered electron image of glass-like carbon.
- 997 <u>Supplementary Figure 3</u>: Synchrotron X-ray powder diffraction of graphite, char and glass-like
 998 carbon.
- 999 <u>Supplementary Figure 4</u>: Raman spectra of polished and unpolished graphite used as starting
- 1000 material, and of polished graphite quenched from experimental conditions.
- 1001 <u>Supplementary Figure 5</u>: Raman spectra of char obtained from high-pressure decomposition of
- 1002 glucose at 1 GPa–800°C after 12h and 24 h.
- 1003 <u>Supplementary Figure 6</u>: Composition of the fluids from decomposed glucose, measured by using
- 1004 the capsule-piercing technique and plotted on ternary C–O–H diagram.