# An experimental study to explore hydrogen diffusion in clinopyroxene at low temperatures (195 - 400 °C) and consequences for re-equilibration at near surface conditions

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#### Abstract

Studying diffusion of hydrogen in nominally anhydrous minerals (NAMs), like clinopyroxene, at low temperatures is a challenging task due to experimental and analytical difficulties. We applied a combination of hydrogen implantation to produce concentration gradients in natural diopside crystals with Nuclear Resonance Reaction Analysis (NRRA) measurements of nanoscale diffusion profiles. Thereby, we were able to conduct experiments at temperatures between 195 - 400 °C. Obtained diffusion rates show a consistent Arrhenius relation  $D_{\rm H} = 5.47 \ (\pm 13.98) \cdot 10-8 \cdot \exp(-115.64 \ (\pm 11.5) \ kJ \ mol^{-1}/RT) \ m^2 s^{-1}$ . Notably, our results lie well within the range of extrapolations from high temperature experiments ([?] 600 degC) of previous studies. This implies that fast diffusion model that simulates the ascent of crystals (0.5, 1.0, and 2.0 mm) along two representative geotherms (oceanic and continental) from 600 to 100 degC, to assess potential re-equilibration of H contents in clinopyroxene at low temperatures. Our model highlights the need to carefully consider boundary conditions, which are a function of P-T-?O2, that control the concentration gradient at the crystal's rim. The results from this model allow an assessment when re-equilibration in dependence of crystal size and cooling rate must be considered. Fast ascent (e.g., kimberlitic melt) preserves initial hydrogen contents even in 0.5 mm size clinopyroxene crystals. However, dwelling at low temperatures (e.g., 300 °C) for several thousands of years (e.g., serpentinization) leads to extensive re-equilibration in 2 mm crystals.

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13	Key Points:
14 15 16 17 18 19 20 21	<ul> <li>Diffusion rates of hydrogen in clinopyroxene in the low temperature range (195 – 400 °C) were quantified for the first time</li> <li>Diffusion coefficients at low temperatures lie within the range of extrapolations from high temperature experiments</li> <li>Non-isothermal modeling was applied to evaluate potential re-equilibration of clinopyroxene crystals at low temperatures</li> </ul>
22	Abstract

23 Studying diffusion of hydrogen in nominally anhydrous minerals (NAMs), like clinopyroxene, 24 at low temperatures is a challenging task due to experimental and analytical difficulties. We applied a combination of hydrogen implantation to produce concentration gradients in natural 25 26 diopside crystals with Nuclear Resonance Reaction Analysis (NRRA) measurements of 27 nanoscale diffusion profiles. Thereby, we were able to conduct experiments at temperatures between 195 – 400 °C. Obtained diffusion rates show a consistent Arrhenius relation  $D_H =$ 28 5.47 (± 13.98)  $\cdot 10^{-8} \cdot exp$  (-115.64 (±11.5) kJ mol<sup>-1</sup> /RT) m<sup>2</sup>s<sup>-1</sup>. Notably, our results lie well 29 30 within the range of extrapolations from high temperature experiments ( $\geq 600$  °C) of previous 31 studies. This implies that fast diffusion of hydrogen (compared to other elements) extends to 32 low temperatures. We used these results in a non-isothermal diffusion model that simulates the ascent of crystals (0.5, 1.0, and 2.0 mm) along two representative geotherms (oceanic and 33 continental) from 600 to 100 °C, to assess potential re-equilibration of H contents in 34 35 clinopyroxene at low temperatures. Our model highlights the need to carefully consider

boundary conditions, which are a function of  $P-T-fO_2$ , that control the concentration gradient at the crystal's rim. The results from this model allow an assessment when re-equilibration in dependence of crystal size and cooling rate must be considered. Fast ascent (e.g., kimberlitic melt) preserves initial hydrogen contents even in 0.5 mm size clinopyroxene crystals. However, dwelling at low temperatures (e.g., 300 °C) for several thousands of years (e.g., serpentinization) leads to extensive re-equilibration in 2 mm crystals.

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# 43 **1 Introduction**

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45 The presence of hydrogen (often simplistically termed "water") in nominally anhydrous 46 minerals (NAMs) has considerable implications for our understanding of Earth's dynamics. Properties such as melting (Hirth and Kohlstedt 1996), electrical conductivity (Karato 1990; 47 48 Fei et al. 2020), seismic response (Karato and Jung 1998), phase transitions (Ohtani and Litasov 49 2006), and mechanical features (Chen et al. 2006) of a mineral are affected already by trace 50 amounts of hydrogen. Although measured concentrations of H in NAMs are very low (e.g., a 51 few to several thousand ppm H<sub>2</sub>O, typically; e.g., Bell and Rossman 1992; Warren and Hauri 52 2014; Le Roux et al. 2021) compared to those in common hydrous phases, when the hydrogen 53 contents of a single mineral are scaled to a planetary scale it becomes obvious that the amount 54 of hydrogen that might be stored in the mantle would make a large contribution to the earth's 55 overall water budget and cycle (e.g., Hirschmann 2006; Hirschmann and Kohlstedt 2012). Thus, 56 it is not surprising that great efforts have been invested in further exploring the role of hydrogen 57 in NAMs since the early works of Martin and Donnay (1972), as reflected in Volume 62 of 58 Reviews in Mineralogy and Geochemistry (Keppler and Smyth 2006) and a vast number of 59 more recent studies (e.g., Padrón-Navarta et al. 2014; Tollan et al. 2017; Ferriss et al. 2018; 60 Reynes et al. 2018; Thoraval et al. 2019; Liu and Yang 2020; Moine et al. 2020; Demers-61 Roberge et al. 2021; Jollands et al. 2021).

However, a legitimate question that remains is whether hydrogen concentrations that are measured in NAMs collected on Earth's surface really represent the original H contents established in the mantle, or if these contents were modified at some later stage e.g., by alteration, metasomatism (Kilgore *et al.* 2020; Tang *et al.* 2020; Peslier *et al.* 2015) or mineralmelt interaction (Lynn and Warren 2021; Le Roux *et al.* 2021). In spite of the growing number of experimental studies that reveal new details on hydrogen in NAMs and systematic studies about hydrogen contents in these minerals (Warren and Hauri 2014; Kumamoto *et al.* 2019), 69 studies that treat natural mantle xenoliths seem to yield contrasting results in terms of what the 70 hydrogen content actually recorded (Gose et al. 2011; Wang et al. 2021). Since diffusion controls the redistribution of hydrogen in a crystal it is crucial to determine rates of diffusive 71 72 transport as a function of various parameters (e.g., temperature, pressure,  $fO_2$ , or mineral 73 composition) to assess potential modifications of hydrogen contents at different conditions. 74 These quantities are known only by extrapolation of high temperature data because most 75 experimental data for H diffusion in NAMs have been collected only over a limited range of 76 high temperatures (800-1200°C for olivine, 600-1000°C for pyroxenes; e.g., Demouchy and 77 Mackwell 2006; Demouchy et al. 2016; Ferriss et al. 2018; Mackwell and Kohlstedt 1990; Ferriss et al. 2016; Hercule and Ingrin 1999; Ingrin et al. 1995; Stalder and Skogby 2003; 78 79 Woods et al. 2000; Stalder and Behrens 2006) and may lead to large uncertainties on 80 extrapolation (Lynn and Warren 2021).

81 To quantify the loss / retention of hydrogen at low temperatures (e.g., <<600°C) it is necessary 82 to know the rates of diffusion at those conditions. We combined two experimental methods, ion 83 implantation and Nuclear Resonance Reaction Analysis (NRRA), to develop a method for the 84 determination of diffusion rates of H in NAMs at low temperatures. Hereby we were able to quantify diffusion rates of hydrogen in clinopyroxene at 195-400 °C, temperatures that are 85 associated with metasomatism, prolonged cooling of lava flows or eruption deposits, and 86 87 serpentinization and that are relevant for applications such as ascent chronometry (e.g., 88 Newcombe et al. 2020; Jollands et al. 2020). We use non-isothermal diffusion modeling with 89 the newly obtained diffusion rates to assess the feasibility of hydrogen re-equilibration during 90 ascent of clinopyroxene from the mantle.

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92 2 Methods
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- 93
- 94 **2.1 Sample preparation**
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96 For our diffusion experiments we used gem quality single crystals of diopsidic clinopyroxene. 97 These had to have homogeneous surface areas that were free of cracks (to avoid adsorbed water 98 causing artefacts) and could accommodate the ion beam used for NRRA analysis (spot  $\emptyset$  1-2 99 The had of mm). crystal used an average formula  $Na_{0.03}Ca_{0.95}Mg_{0.88}Fe^{2+}0.08Fe^{3+}0.02Cr_{0.01}Al_{0.04}Si_{1.98}O_{6}$  (standard deviation of each formula unit 100

101 < 0.01). The composition was determined using a *Cameca SX Five FE* electron microprobe 102 operated at 15 keV and 15 nA using wavelength dispersive spectrometers (Crystals used: LTAP for Na, Mg, TAP for Al, Si, LPET for K, Ti, PET for Ca, Cr and LLIF for Mn, Fe). Fe<sup>2+</sup>/Fe<sup>3+</sup> 103 104 ratios were estimated following Droop (1987). The standards used were jadeite (Na), San Carlos 105 Olivine (Mg, Si), orthoclase (Al, K), diopside (Ca), rutile (Ti), synthetic Cr<sub>2</sub>O<sub>3</sub> (Cr), spessartine 106 (Mn), and fayalite (Fe). The crystallographic orientation was verified by Laue analysis before 107 large prismatic crystals (few cm in length) were cut into several slabs (thickness ~ 3 mm) with 108 the surface perpendicular to the crystallographic c-axis [001]. This was followed by embedding 109 the samples in epoxy, grinding, and a final polishing (to  $0.25 \,\mu m$ ) using diamond paste. Cuboids with a surface area of about 4x4 mm were produced from these (Figure 1). Last, the samples 110 111 were intensively cleaned using a routine involving deionized water, acetone, and ethanol and 112 checked for the quality of polishing in an optical microscope.

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# 2.2 Hydrogen implantation

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Hydrogen was implanted at the 500 kV accelerator at the Central Unit for Ionbeams and 115 116 Radionuclides (RUBION) at the Ruhr-University Bochum, Germany. The depth of the implantation depends on the ion beam energy and the material-specific stopping-power, s, 117 118 which is an expression for the energy loss of the incident ion per distance unit (e.g., keV/nm). Concentration profiles were simulated using the software SRIM (Ziegler et al. 2010) prior to 119 120 the actual implantation to determine the best choice of ion beam energy. Ideally, the latter is 121 high enough to produce a concentration peak at depths far from the sample surface so that the 122 profile after diffusion experiments is not (partially) superimposed by the surface peak that is 123 related to adsorption of contaminants on the sample surface. Notably, we observed in another 124 study that the process of preparation can introduce excess hydrogen in the near surface region 125 in olivine (~ first 100 nm, Figure S3 supplementary material). Additionally, beam energies 126 should be small enough to deposit hydrogen at depths that are still easily accessible using NRRA (max. depth ~ 2-3  $\mu$ m). Fluences of 1.48  $\cdot$  10<sup>16</sup> at/cm<sup>2</sup> were implanted at an ion energy 127 128 of 60 keV and a beam intensity of 15 µA, while the implantation area was about 1 cm in 129 diameter. The hydrogen source was a duoplasmatron and the ion beam was moved across the 130 sample in a scanning mode during implantation to ensure homogeneous lateral distribution of



**Figure 1:** Illustration of the experimental procedure. (1) Diopside crystals were oriented and cuboids with a surface area of ca. 4x4 mm were cut with their surfaces perpendicular to the crystallographic c-axis (2) Polished sample surfaces were implanted by a hydrogen beam at 60 keV to produce concentration peaks at around 650 nm depth and 1 at% H (1650 H<sub>2</sub>O ppm wt%) concentration maxima. The beam width is larger than the sample size and the implantation is operated in a scanning mode (yellow line). (3) An analysis of the implantation profile by NRRA to precisely define the initial diffusion setup (t<sub>0</sub>) was followed by diffusion anneals at 195 °C, 270 °C, and 400 °C in a gas mixing furnace (100 vol% CO<sub>2</sub>). Resulting diffusion profiles were finally measured with NRRA. The non-destructive nature of the experimental design allows to obtain snapshots (profiles t<sub>1</sub> and t<sub>2</sub>) of the proceeding homogenization with time. Dimensions are not to scale.

hydrogen (Figure 1). Implanted concentration maxima were located at around 650 nm from thesample surface.

#### 133 **2.3** Nuclear Resonance Reaction Analysis (NRRA)

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Hydrogen concentration profiles that result from implantation and diffusion experiments were
measured using Nuclear Resonance Reaction Analysis (NRRA). We limit the information about

137 this method to the essentials and refer the reader to Becker and Rogalla (2016) for a thorough 138 description and to Bissbort et al. (2021) for an explanation in context of a diffusion study. NRRA utilizes the nuclear reaction  ${}^{1}H({}^{15}N,\alpha\gamma){}^{12}C$  to detect hydrogen by bombarding the sample 139 140 with <sup>15</sup>N ions. Emitted y-rays have a specific energy of 4.4 MeV, which is characteristic for this 141 nuclear reaction. The  $\gamma$ -rays are counted using a NaI(Tl) borehole detector with a high detection 142 efficiency. The probability for the nuclear reaction depends strongly on the <sup>15</sup>N energy (width of the resonance window is 1.8 keV (Maurel and Amsel 1983)) with a resonance at  $6393.6 \pm$ 143 144 1.3 keV (Osipowicz *et al.* 1987) or at 6399.1  $\pm$  2.9 keV (Becker *et al.* 1995). <sup>15</sup>N at the resonance 145 energy will introduce the nuclear reaction in the very first atomic layers of the sample surface 146 i.e., probe the hydrogen content at the surface. An increase in beam energy shifts the nuclear 147 reaction to greater depths, since <sup>15</sup>N must loose energy by ion-sample-interaction first to attain 148 the resonance energy. Hence, entire hydrogen depth profiles can be obtained by increasing the 149 beam energy by small increments (e.g., 20 keV) while maintaining the depth resolution of a 150 few nm due to the strong energy-sensitivity of the nuclear reaction. The detected  $\gamma$ -rays are 151 normalized to the incident beam ions (= yield) and are a direct measure of the number of 152 hydrogen atoms, thus making NRRA a standard-less technique. A conversion of the beam 153 energy to a distance unit requires the knowledge of the material-specific stopping-power s, 154 which is a function of the sample composition and density. We used the software SRIM (Ziegler 155 et al. 2010) to calculate the stopping-power for <sup>15</sup>N in clinopyroxene of the relevant composition. Dividing the difference between beam energy and resonance energy ( $\Delta E$ ) by the 156 157 stopping power yields the distance. Typical beam currents during analysis were  $\sim 40$  nA. NRRA 158 is a nuclear method that is independent of chemical bonds (which depend on outer electrons), 159 thus the total hydrogen content is detected irrespective of its nature of bonding (e.g., H<sub>2</sub>, H<sub>2</sub>O, 160 (OH)<sup>-</sup>). One initial calibration is sufficient, and no standards are needed in contrast to alternative 161 methods (e.g., SIMS or IR-spectroscopy) that are often used to measure hydrogen concentration 162 profiles. Most notably, NRRA is a non-destructive technique that allowed us to measure the 163 initial profile prior to experiments and to observe the development of diffusion profiles with 164 time within one sample.

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#### 2.4 Hydrogen diffusion experiments

167 Implantation of hydrogen produced concentration profiles that are approximately described by 168 a normal distribution with depth (depth of implantation maxima ~650 nm), with a minor excess 169 of hydrogen along the surface-facing side of the concentration peak (Figure 2a). Hydrogen 170 concentration depth profiles were obtained by NRRA in each sample after implantation and 171 before diffusion experiments, thereby allowing us to characterize the initial diffusion setup.

- 172 Diffusion experiments were performed in a gas-mixing furnace using a K-type thermocouple
- 173 (NiCr-Ni). The furnace was flushed with 100 vol% CO<sub>2</sub> to establish non-oxidizing conditions
- 174 during experiments (it is not possible to control  $fO_2$  at a specific value at the low temperatures
- 175 of this study by mixing CO and CO<sub>2</sub>). Experiments were terminated by dropping the samples
- 176 in an actively cooled zone for fast quench.
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# 178 **3** Modeling of hydrogen diffusion

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180 Diffusion profiles were fitted using an explicit numerical solution. Concentration changes with 181 time at a specific location are described by Fick's second law (equation 1), with *c* being the 182 concentration, *t* being time, *x* being the distance, and *D* being the diffusion coefficient. A finite 183 difference scheme was used to solve the diffusion equation (equation 2), with  $\Delta t$  being the time 184 step size,  $\Delta x$  being the distance step size, *c* being

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$$\frac{\partial c(x,t)}{\partial t} = D \frac{\partial^2 c(x,t)}{\partial x^2} \qquad (eq.1)$$

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$$c_{i,j+1} = c_{i,j} + \frac{D \cdot \Delta t}{\Delta x^2} \cdot \left[ c_{i+1,j} - 2c_{i,j} + c_{i-1,j} \right]$$
(eq. 2)

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190 
$$\sigma_{fit} = \frac{1}{N} \sqrt{\sum (c_{ms}(x) - c_{cal}(x))^2}$$
 (eq. 3)

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192 the concentration with the two subscripts *i* and *j* being indices for space and time, respectively. 193 The geometry of the diffusion problem in this study involves a concentration gradient (i.e., 194 implantation peak) that is located far from the crystal surface (Figure 2). Thus, the system 195 boundaries are unaffected by diffusion (i.e., infinite medium) and are modelled to be open at 196 both ends of each profile. The reader is referred to Costa et al. (2008) for a detailed description 197 of the numerical solution. Notably, the non-destructive NRRA allows us to measure 198 concentration profiles prior to experiments that are used as an initial condition (t = 0) in 199 diffusion calculations (i.e., providing  $c_{i,0}$  values). This approach provides an ideal description 200 of the diffusion geometry, in contrast to an assumed initial profile (e.g., Gaussian). Finally, this

allows to model much smaller changes in hydrogen distribution, and it eliminates the necessityto consider convolution effects.

203 Diffusion of hydrogen at experimental temperatures led to a broadening of the implanted 204 concentration peak while its maximum decreased. The resulting concentration profile could be 205 satisfactorily fitted with a constant diffusion coefficient D along the profile distance at a given 206 temperature, which indicates that there is no observable dependence of diffusion rates on



**Figure 2:** (a) Example for a hydrogen concentration depth profile produced by ion implantation and measured by NRRA (open circles). The shape of the peak can be described a Gaussian distribution (grey area). (b) The initial hydrogen implantation profile (open circles) broadened and flattened due to diffusion at 195 °C for 9.5 days (filled circles). The red line is the best fit of the numerical solution. (c) A time-series was performed in sample Cpx25\_3 at 270 °C. The obtained diffusion coefficients from best fits display consistency over time. (d) The concentration profile in Cpx25\_1 was strongly modified after only 15 minutes at 400 °C. The grey circles illustrate how a depth profile is affected if a fissure loaded with adsorbed atmospheric water interferes with NRRA analysis (i.e., the ion beam spot).

207 hydrogen concentration. Experiments were conducted at three different temperatures, 195 °C,

208 270 °C, and 400 °C (Figure 2b-c). A timeseries of 3 experiments within one sample (Cpx25\_3)

at 270 °C gives insight on how the diffusion profile develops within one sample with increasing



**Figure 3:** Diffusion coefficients that were obtained from fitting concentration profiles from the time-series at 270 °C yield constant values over time. The range of scatter in  $\log_{10}$ D is about 0.24 log units and is larger than error estimates calculated from error propagation. For further details see Appendix A in Faak *et al.* (2013).

experimental duration. It was also used to assess the suitability of our method for obtaining diffusion coefficients, which is supported by almost constant diffusion coefficients ( $\Delta \sim 0.24$ log units, Figure 3) in this time-series. D values were constrained by minimizing the misfit ( $\sigma_{fit}$ ) between each measured concentration,  $c_{ms}(x)$ , and the corresponding calculated concentration,  $c_{cal}(x)$  at the same location (equation 3). A summary of the experiments, determined diffusion coefficients, and misfit  $\sigma_{fit}$  is provided in Table 1.

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Table 1: Diffusion experiments at 195°C, 270 °C, and 400 °C. For the timeseries the total duration is the sum of
 the dwell time.

Sample_experiment	T (°C)	<b>Total duration</b>	D (m <sup>2</sup> /s)	σfit
Cpx25_1_EXP1	400	15 min	3.50E-17	0.015
Cpx25_2_EXP1	195	9.5 days	3.71E-21	0.006
Cpx25_3_EXP1	270	40 min	4.95E-19	0.007
Cpx25_3_EXP2	270	100 min	5.15E-19	0.010
Cpx25_3_EXP3	270	220 min	8.50E-19	0.006

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- 222 **4** Comparison with previous studies
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Diffusion rates that were determined in our experiments describe an Arrhenius relation  $D_H = 5.47 (\pm 13.98) \cdot 10^{-8} \cdot exp (-115.64 (\pm 11.5) kJ mol^{-1} /RT) m^2 s^{-1}$  (purple solid line, Figure 4). Our experimental method allowed us to explore diffusion of hydrogen in diopside ( $X_{Fe} = Fe/(Fe+$  Mg) = 0.102) at low temperatures (195 – 400 °C). However, a comparison of our results with those of other studies in clinopyroxene is restricted to higher temperatures ( $\geq 600$  °C).

Ingrin *et al.* (1995) conducted dehydration experiments in Russian gem quality diopside ( $X_{Fe} =$ 0.036) crystals at 700 to 1000 °C in air. They studied diffusion along three crystallographic directions, [001], [100]\*, and [010] using FTIR but did not find anisotropy in diffusion rates. Thus, they propose that diffusion of hydrogen is isotropic in diopside. The temperature dependence, which is valid for all orientations, is expressed by an Arrhenius relation  $D_H = 5.04$  $(\pm 14.12) \cdot 10^{-7} \cdot exp (-136 \pm 27 kJ mol^{-1} /RT) m^2/s$  (grey solid line, Figure 4).

235 Instead of dehydration experiments, Hercule and Ingrin (1999) performed extractionincorporation experiments in diopside crystals similar to those used in Ingrin et al. (1995) at 236 237 700 – 1000 °C at a pH<sub>2</sub> of 0.1 atm and 1 atm. Similar to Ingrin et al. (1995) they observe that the kinetics are independent of crystallographic orientation, but also of the partial pressure of 238 H<sub>2</sub> and oxygen fugacity  $fO_2$ . They determined an Arrhenius law  $D = 2.00 (\pm 5.05) \cdot 10^{-7} \cdot exp$  (-239  $126 \pm 24 \text{ kJ mol}^{-1}/\text{RT}$ ) m<sup>2</sup>/s for this set of experiments (green dotted line, Figure 4). The authors 240 241 also carried out hydrogen-deuterium exchange experiments in the same diopside at 600 - 900 242 °C along [001], [100]\*, and [010] at 1 atm. They observe that diffusion along [001] and [100]\* 243 is two orders of magnitude faster than the incorporation rate from the other experimental setup. The Arrhenius relation is  $D_H = 3.98 (\pm 7.33) \cdot 10^{-4} \cdot exp (-149 \pm 16 \text{ kJ mol}^{-1} / RT) m^2/s$  (green 244 245 dashed line, Figure 4). Diffusion along [010] is slower than in the other directions but one magnitude faster than the H uptake. The related Arrhenius law is  $D_H = 1.00 (\pm 3.91) \cdot 10^{-5} \cdot exp$ 246  $(-143 \pm 33 \text{ kJ mol}^{-1} / RT) \text{ m}^2/\text{s}$  (green solid line, Figure 4). They conclude that incorporation of 247 hydrogen is coupled to oxidation-reduction of iron  $Fe^{3+} + O^{2-} + 1/2H_2(g) = Fe^{2+} + OH^2$ . Hence, 248 the rate of hydrogen uptake is a function of the crystals Fe-content (Hercule and Ingrin 1999). 249

Woods et al. (2000) used Jaipur diopside crystals ( $X_{Fe} = 0.069$ ) for dehydration experiments 250 between 700 and 850 °C and at  $fO_2$  of 10<sup>-14</sup> bar. A comparison of diffusion rates between 251 crystallographic orientations shows that rates along [100] and [001]\* are fastest with the 252 253 Arrhenius relations  $D_H = 7.94 (\pm 34.75) \cdot 10^{-3} \cdot exp (-181 \pm 38 \text{ kJ mol}^{-1} / RT) m^2/s$  (vellow dashed line, Figure 4) and  $D_H = 3.98 \ (\pm 14.67) \ \cdot 10^{-4} \ \cdot \ exp \ (-153 \pm 32 \ kJ \ mol^{-1} \ /RT) \ m^2/s$  (yellow solid 254 255 line, Figure 4), respectively. The relatively narrow experimental temperature range did not 256 allow them to obtain a temperature-relation for diffusivity along [010]. However, their 257 experiments indicate that diffusion along [010] is an order of magnitude slower than in the other 258 two directions. The observed anisotropy agrees with the results from Hercule and Ingrin (1999). 259 Woods et al. (2000) point out that mm-size diopside crystals will be affected by re-equilibration 260 at temperatures as low as 800 °C.



**Figure 4:** A comparison of published diffusion coefficients from high temperature experiments ( $T \ge 600 \text{ °C}$ ) and associated Arrhenius relations (coloured lines) with results from this study (purple data, error bars smaller than symbols) confirm that latter lie well within the log<sub>10</sub>D range predicted by extrapolation from high temperatures. F16 = Ferriss et al. (2016); H&199 = Hercule & Ingrin (1999); W00 = Woods et al. (2000); S09 = Sundvall et al. (2009); I95 = Ingrin et al. (1995).

- Sundvall *et al.* (2009) used Fe-free diopside ( $X_{Fe} \approx 0.0$ ) that was synthesized at water-saturated 261 conditions. Their results from FTIR analysis of dehydration experiments at 800 - 1000 °C 262 indicate that diffusion rates along [010] and [100]\* are similar within error limits, which 263 264 contradicts previous studies mentioned above. The Arrhenius relation for [010] is  $D_H = 7.94$  (±  $(42.07) \cdot exp (-331 \pm 50 \text{ kJ mol}^{-1} /RT) \text{ m}^2/\text{s}$  (pink solid line, Figure 4). Diffusion rates along 265 266 [100]\* exhibit an Arrhenius relation  $D_H = 3.16 (\pm 17.48) \cdot exp (-312 \pm 55 \text{ kJ mol}^{-1} /RT) \text{ m}^2/\text{s}$ 267 (pink dashed line, Figure 4). These diffusion rates are several orders of magnitude slower than 268 in natural Fe-bearing samples. This difference also applies to diffusion rates we obtained in our 269 natural diopside crystals when one extrapolates the high temperature data to low temperatures 270 (pink lines, Figure 4).
- 271 Ferriss *et al.* (2016) studied dehydration in Kunlun diopside ( $X_{Fe} = 0.025$ ), Jaipur diopside ( $X_{Fe}$ = 0.075), and augite ( $X_{Fe}$  = 0.180) at 800 – 1000 °C at an  $fO_2$  at the QMF buffer. They present 272 273 an Arrhenius relation for isotropic hydrogen diffusion in Kunlun diopside  $D_H = 1.59 (\pm 31.02)$  $\cdot 10^{-9} \cdot exp (-100.4 \pm 2.7 \text{ kJ mol}^{-1}/\text{RT}) \text{ m}^2/\text{s}$  (blue solid line, Figure 4). Experiments in the Jaipur 274 275 diopside and augite, which are more Fe-rich than the Kunlun diopside, show faster diffusion of 276 hydrogen at the same temperature. This suggests that higher Fe contents cause faster diffusion 277 rates of hydrogen. The authors also compare diffusion rates of hydrogen between 278 clinopyroxenes of various Fe contents, including other studies. There is an increase in diffusion 279 coefficients with increasing Fe-contents, but this increase diminishes towards high Fe contents. 280 As pointed out by Ferriss et al. (2016), a compositional dependence of hydrogen diffusion in 281 clinopyroxene might not be adequately described by the Fe content, instead Al could also play 282 a significant role as has been shown for orthopyroxene by Kumamoto et al. (2019). 283 Unfortunately, they do not provide Arrhenius relations for the Jaipur diopside and augite, which 284 precludes a direct comparison of results from these samples with results of our study. However, 285 an extrapolation of the Arrhenius law that was obtained for the Kunlun diopside is in excellent 286 agreement with our results, although  $X_{Fe}$  is much higher in our diopside (0.102 vs. 0.025). The 287 conformity of diffusion rates in our diopside and the Kunlun diopside despite substantially 288 different Fe contents ( $X_{Fe}$  of our sample lies between that of the Jaipur diopside and augite) also 289 indicates a more complex composition-diffusivity relationship.

A comparison of our diffusion rates with extrapolations from previous studies at high temperatures, highlights that we were able to resolve the outstanding issue whether diffusion rates can be extrapolated from high temperatures towards low temperatures. Our results lay well within the range that is constrained by extrapolation of Arrhenius relationships from high temperature experiments for Fe-bearing diopside to temperatures relevant in this study (Figure 4). Although, we did not constrain diffusion rates as a function of crystal orientation, previous studies (e.g., Woods *et al.* 2000; Hercule and Ingrin 1999) suggest that diffusion coefficients along [001] are greatest and like those along [100]. Therefore, it is evident that the high diffusivity of hydrogen (compared to other elements) in clinopyroxene extends continuously towards low temperatures.

300 Further, diffusion profiles in our experiments are well fitted by a simple (i.e., concentration-301 independent) numerical diffusion model (solving Fick's second law, equation 1) and diffusion 302 rates as a function of temperature describe an Arrhenius relation. The consistency of our results 303 resolves two concerns that might arise when implanted concentration distributions are to be 304 used for diffusion studies. (i) Does the damage introduced by ion implantation affect diffusion? 305 (ii) Is the implanted hydrogen bonded to the mineral matrix like "natural" hydrogen? First, 306 hvdrogen is a light element and therefore causes less damage during implantation compared to 307 heavy elements at low energies (Zhang et al. 2009). Further evidence for a negligible damage 308 of the sample is provided by results of diffusion experiments. Sample damaging (i.e., 309 introduction of defects) due to irradiation follows a distribution similar to that of the hydrogen 310 concentration. Thus, the region between the sample surface and the end of the implantation 311 peak is affected by irradiation damage, but not so the rest of the sample at greater depth. If 312 irradiation damage is thought to be intense enough to affect diffusion to a non-negligible extent, 313 one would expect this to be reflected in an asymmetric development of the concentration 314 profile, caused by a different diffusion rate and/or mechanism in the damaged zone compared 315 to the pristine crystal. This was not observed in our diffusion profiles (Figure 2). Regarding (ii), 316 IR-measurements of implanted hydrogen in olivine confirm that all the implanted hydrogen is 317 present as OH, similar to the bonding state of natural hydrogen (Schaible and Baragiola 2014). 318 In this work we assume that clinopyroxene behaves the same way. Hypothetically, a change 319 from H or H<sub>2</sub> to OH during annealing in diffusion experiments would be expected to be 320 accompanied by a change in diffusion rate with time as the proportions of different H-species 321 change. However, time-series experiments feature a constant diffusion rate within uncertainties (Figure 3). Last, implanted hydrogen fluences of 1.48.10<sup>16</sup> at/cm<sup>2</sup> are relatively low (peak 322 323 maximum ~ 1 at% H; 1650 wt. ppm H<sub>2</sub>O) but close to natural concentrations in mantle pyroxenes (e.g., Warren and Hauri 2014). 324

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# 327 5 Evaluation of re-equilibration of H in clinopyroxene at temperatures 328 below 600 °C.

329

#### 330 5.1 A non-isothermal diffusion model

331

The newly determined diffusion rates at low temperatures offer us the opportunity to model changes in hydrogen concentrations in clinopyroxene due to diffusion at conditions that are relevant for late-stage hydration and dehydration. Diffusive flux occurs when there is a cause i.e., chemical disequilibrium, caused, for example, in response to changes in temperature, pressure, oxygen fugacity ( $fO_2$ ), the speciation of accompanying fluids, and solubility of hydrogen in clinopyroxene while a crystal of clinopyroxene ascends from the mantle to Earth's



**Figure 5:** A continental geotherm (CT), an oceanic geotherm (OC), and solubilities of hydrogen in clinopyroxene that correspond to conditions along these PT paths (indicated by number labels and colour) were calculated following the procedure described in the Appendix. Diffusion rates in our model change during ascent according to the Arrhenius relationship obtained in this study (i.e., non-isothermal). Boundary concentrations that induce a concentration gradient change according to the solubility. Temperature boundaries of the model are indicated by dashed lines (600  $^{\circ}$ C to 100  $^{\circ}$ C).

surface, for example in volcanic rocks such as kimberlites or during the process of emplacement of an ophiolite. We have developed a model that simulates diffusion of hydrogen in clinopyroxene during its ascent, while the system variables change according to temperaturepressure paths and associated conditions (Figure 5).

342 Diffusion is calculated as a temperature dependent process (non-isothermal). The diffusion equation is solved using a finite difference model as described in section 3. The diffusion 343 344 coefficient  $D_H$  changes during the calculation according to the change in temperature during ascent and is derived from our Arrhenius relation  $D_H = 5.47 (\pm 13.98) \cdot 10^{-8} \cdot exp$  (-115.64 345  $(\pm 11.5)$  kJ mol<sup>-1</sup>/RT) m<sup>2</sup>s<sup>-1</sup>. Possible effects of pressure and fO<sub>2</sub> on the diffusion coefficient are 346 not included as they are expected to be weak but are not constrained experimentally. However, 347 348 the variation of pressure and  $fO_2$  in the surroundings plays an important role in the calculation 349 through their influence on the boundary conditions. This is considered in our model (with  $fO_2$ 350 set to vary along the OMF buffer), as these variables control, together with temperature, the 351 speciation of a C-O-H fluid (carbon saturated) that may be present. The associated  $fH_2O$  affects 352 the solubility of OH in clinopyroxene (Keppler and Bolfan-Casanova 2006). Boundary 353 concentrations are adjusted to the appropriate values of hydrogen concentration at the rim 354 corresponding to the P-T- $fO_2$  condition at each time step. The initial condition is taken to be a 355 homogeneous hydrogen distribution in the crystal at equilibrium at the P-T-fO2 condition of the 356 starting depth.

The degree of re-equilibration was calculated using equation 4 where  $c_{initial}$  is the initial concentration,  $c_{model\ core}$  is the maximum concentration after cooling (usually in the core), and  $c_{eq}$  corresponds to the final equilibrium concentration (i.e., the concentration at the rim of the crystal) (Chakraborty and Ganguly 1991; Costa *et al.* 2003; Lynn *et al.* 2017).

361

362 
$$re - equilibration (\%) = \frac{c_{initial} - c_{model core}}{c_{initial} - c_{eq}} \times 100$$
 (eq.4)

363

The results provide an impression of how the hydrogen budget and its distribution in clinopyroxene is modified and how this depends on the exhumation history and crystal size. We aim to clarify what the possibilities are in terms of hydrogen re-equilibration, demonstrated through some representative settings, rather than to reproduce and resolve specific cases. Therefore, two geotherms were calculated that are meant to constitute a continental (CT, 534

°C/GPa) and an oceanic (OC, 226 °C/GPa) setting. A detailed description of the development 369 370 of this model is provided in Appendix I. Non-isothermal diffusion of hydrogen was modelled 371 for clinopyroxene crystals of 2 mm, 1 mm, and 0.5 mm from 600 °C to 100 °C and the cooling 372 rate (i.e., ascent rate) was varied by orders of magnitude (1000 to 0.1 °C/yr) to study its effect 373 on the progress of equilibration. Figure 5 displays both geotherms in P-T-space whereas the 374 modelled temperature range is indicated by dashed lines. The geotherm that reflects a 375 continental lithosphere (CT) yields higher temperatures compared to the oceanic geotherm 376 (OC) at same pressures, due to the assignment of higher radiogenic heat production to the 377 former and a thin oceanic crust of 10 km to the latter (instead of 40 km, see Appendix I for 378 more details).

379 Although diffusion rates that were derived from the experimentally established Arrhenius 380 relationship are the same for both geotherms at similar temperature (D = f(T)), the pressure and 381 therefore the solubility of OH in the model clinopyroxene is different. Consequently, the 382 starting point at 600 °C corresponds to a higher pressure of 2.35 GPa for the oceanic and a 383 lower pressure of 1.11 GPa for the continental model geotherm, which corresponds to 70.1 and 384 45.5 wt. ppm H<sub>2</sub>O, respectively. This difference is reflected in the initial state of our models (t 385 = 0) by a homogeneous concentration plateau and boundary concentrations at these values 386 (Figure 6, dotted lines). A comparison of the evolution of hydrogen diffusion profiles in crystals 387 of similar size that followed different P-T-paths reveals that at the same cooling rate the 388 hydrogen content in the clinopyroxene that travelled along CT is more re-equilibrated than one 389 that followed OC. This contrast is a consequence of different OH solubilities along the two 390 paths that define the boundary concentrations (i.e., a compositional gradient), but not a result 391 of different diffusivities. To further test the effect of solubility of OH on the progress of re-392 equilibration, we changed the boundaries to a fixed low concentration (= 0 wt. ppm) right from 393 the start of diffusion until the final state is attained. We have done this for a cooling rate of 394 100°C/yr in 1 mm crystals along both geotherms (Figure 6, dashed lines). The obtained 395 concentration profiles display an overall different shape compared to the original model that 396 includes variable boundary concentrations. Steeper flanks towards the crystal surface are caused 397 by a steep concentration gradient, which is present from the initial state, and which promotes a strong diffusive flux. Most notably, the extent of re-equilibration proceeded much more when 398 399 the model did not include solubility-controlled boundary concentrations. Thus, a consideration



**Figure 6:** Diffusion profiles calculated from our model for the continental geotherm (CT, left side) and the oceanic geotherm (OC, right side) for crystal sizes of 0.5, 1.0, and 2.0 mm (top to bottom) using various cooling/ascent rates. The initial homogeneous OH concentration corresponds to the solubility of OH in clinopyroxene at the starting condition ( $T_0 = 600 \, ^\circ$ C and corresponding pressure) and is indicated by the dotted line (~46 wt. ppm for CT, ~70 wt. ppm for OC). Temperature labels below and above the profiles are apparent closure temperatures as a function of the cooling rate (taken from equilibrium temperatures corresponding to concentrations at 15 µm from the surface). The grey dashed lines in the 1.0 mm models are calculated concentration profiles for a cooling rate of 100 °C/yr, but the boundary concentrations were in this case set to the final concentration (at T = 100 °C) right from the beginning. The substantial differences in shape and extent of re-equilibration compared to the changing boundary concentration model emphasizes the influence of concentration gradients on the rate at which concentration profiles change.

401 of not only diffusivities, but of the system variables such as fluid composition (or, more 402 generally, P-T- $fO_2$ - $fH_2O$  conditions in the surrounding medium) that finally control OH 403 solubility in the mineral is necessary to obtain trustworthy estimates on ascent rates or to assess 404 the degree of re-equilibration. On the other hand, details of the profile shapes contain 405 information on how the boundary conditions evolved with time – this may be helpful in setting 406 the boundary conditions for modelling natural profiles.

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- 408 409

# 5.2 Geological implications

Hydrogen concentration profiles that result from non-isothermal diffusion modelling illustrate 410 411 that below some threshold in cooling rate for a given grain size the initial core concentration is 412 no longer preserved. Small crystals at slow ascent rates show advanced re-equilibration, which 413 is reflected by flat concentration profiles at low values (Figure 6). These should not be 414 misinterpreted to be a sign for low water contents of the mantle, but to be results of a 415 combination of relatively fast diffusion of hydrogen in clinopyroxene even at low temperatures 416 and low H solubilities at those conditions. The dependence of re-equilibration on crystal size 417 and cooling rate follows a systematic behaviour that can be described by equation 5 for each 418 geotherm with the crystal size sz in mm, the cooling rate cr in °C/yr, and the fitting parameters 419 pla, plb, qla, qlb, q2a, and q2b (full description in Appendix II). This equation provides a 420 tool to estimate the progress of re-equilibration for the range of crystal sizes and cooling rates 421 for a given geotherm (Figure 7).

422

423 
$$re-equilibration (\%) = \frac{p1a \times sz^{p1b}}{cr^2 + q1a \times sz^{q1b} \times cr + q2a \times sz^{q2b}}$$
 (eq. 5)

424

These calculations have important geological implications, which we discuss below in the context of (1) ascent chronometry and (2) water contents in NAMs as signature of a mantle source.

(1) Fast diffusion of hydrogen in NAMs has been identified to be a promising tool to model
ascent/decompression rates of magma (Newcombe *et al.* 2020) or to estimate eruption
timescales in volcanic settings (Jollands *et al.* 2020). The principles of the method are that once
an initial equilibrium water concentration was determined (either from water concentrations in

432 melt inclusions and  $K_D(NAM/melt)$  or directly from preserved hydrogen concentration plateaus 433 in crystal cores) and concentration profiles that result from dehydration are measurable, 434 diffusion modelling can be applied to constrain ascent rates. Our model indicates that even very 435 small clinopyroxene crystal sizes (< 0.1 mm) potentially preserve original water contents at 436 their cores at the lower boundary values for estimates on typical ascent rates in volcanoes (e.g., 437 0.05 MPa/s) and temperatures below 600 °C. This implies, that the hydrogen distribution that 438 results from diffusion in NAMs that travel along an adiabat (as an approximation to transport



**Figure 7:** Re-equilibration (req) mapped as a function of crystal size and cooling rate/ascent rate for the continental geotherm (left) and the oceanic geotherm (right). Contours in both maps represent the extent of re-equilibration along these lines. The colour gradient visualizes the re-equilibration (blue = OH content preserved; white = completely re-equilibrated). The plotted lines at the top are re-equilibration profiles for 0.1 mm (green), 0.5 mm (blue), 1.0 mm (yellow), and 2.0 mm (pink) obtained from the model (equation 5). Symbols are the re-equilibration values calculated from the modelled diffusion profiles.

in a melt during ascent), are "frozen in" at relatively high temperatures (Figure 8, left). A modification of these at shallow depths are therefore less likely, and the measured profiles can be interpreted to record ascent rates at high temperatures. However, the uncertainty that is introduced by the choice of boundary concentrations that induce a concentration gradient can be very large. For example, a simplification of the problem to a scenario in which dehydration corresponds to a large concentration gradient from the beginning (e.g., 0 ppm) produces

overestimates on the order of an order of magnitude or so in cooling rates (see Figure 6, dashed 445 446 lines in 1 mm profiles, Figure 8, right). Our model highlights that when changing boundary 447 concentrations are applied, which are a better approximation to natural cases, the uncertainty in 448 the determined cooling rates can be substantially reduced (Figure 8). Our model is set up for 449 calculating ascent in any system (e.g., volcanoes or emplacement of massive lherzolites), but 450 the boundary conditions in the model would need to be set according to the nature of the 451 surrounding medium, and if it is not a C-O-H fluid then equation (5) would not apply (new 452 calculations would be required to produce the equivalent of Eq. (5)).

453 (2) The finding that NAMs from mantle xenoliths contain hydrogen led to the conclusion that 454 these contents represent hydrogen contents that are in equilibrium with a mantle source (Bell 455 and Rossman 1992). Although NAMs incorporate only trace amounts of hydrogen, their major 456 abundance in the mantle implies a large contribution to Earth's overall hydrogen budget and 457 cycle. The effect hydrogen has on various properties of minerals (e.g., rheology and melting 458 temperature) highlights the importance of carefully evaluating its abundance in mantle phases. 459 Our results provide a basis to assess whether original hydrogen contents associated with a 460 mantle source can be preserved during ascent or might be modified, or even completely erased. 461 This helps to evaluate whether measured hydrogen concentrations in these natural samples can 462 be interpreted as a mantle signature. Figure 7 shows that mantle xenoliths that were transported 463 to the surface in a kimberlitic magma probably preserved the original hydrogen content due to 464 the fast ascent rates (e.g., 5 - 37 m/s, Peslier *et al.* 2008). This means that OH concentrations 465 that were measured for example by Bell and Rossman (1992) from a kimberlite source and by 466 Peslier and Bizimis (2015) from Hawaiian peridotite probably reflect those of a mantle source. 467 However, slow cooling or dwelling at temperatures that are typical for serpentinization ( $\sim 300$ 468 °C, Guillot et al. 2015) can erase a mantle signature in large crystals (2 mm) in a few thousand 469 years (Figure 9, left). Results from modelling hydrogen diffusion in clinopyroxene at 300 °C 470 illustrate the development of the hydrogen distribution at temperatures associated with 471 serpentinization, a process that was first suggested by Lynn and Warren (2021) and is supported 472 by the newly obtained low-temperature diffusivity constraints here. As an example, after 2000 473 years at 300 °C, diffusion strongly modifies the distribution from rim to core, while the core 474 content is almost preserved (dependent on the boundary concentration). Although, a mantle 475 signature could have been preserved in the core, the result highlights the requirement for a 476 careful analysis of core contents. Accurate measurements should prevent integration of



**Figure 8:** Left – freezing temperature as a function of cooling rate. Clinopyroxene crystals that were transported in a melt during ascent in volcanic settings are expected to experience fast cooling (> 1000 °C/yr) and therefore preserve a hydrogen distribution that was established at higher temperatures. Slow exhumation (i.e., slow cooling) allows the modification of hydrogen distribution within NAMs in response to lower equilibrium contents at the mineral boundary at low temperatures. Right – Differences in re-equilibration between the two geotherms for different crystal sizes and as a function of cooling rate (lines). In general, differences in the degree of re-equilibration between crystals of the same size but that travelled along different geotherms are smaller for larger crystals. However, this is complicated by a dependence on cooling rate. At low cooling rates larger crystals display greater differences in the degree of re-equilibration between both geotherms. Notably, setting the boundary concentration to a constant value of 0 ppm (instead of a variable boundary concentration) introduces large differences in the degree of re-equilibration (symbols).



**Figure 9:** Modelling of isothermal diffusion at 300 °C for a 2 mm crystal with constant boundary concentrations that mimics dwelling at a temperature that is characteristic for serpentinization. The initial OH concentration (dotted line) is set to 80 OH ( $H_2O$  wt ppm) Using different fixed boundary concentrations (numbers above and below profiles) highlight their influence on the progress of re-equilibration and change in profile shape. Left - Notably, the core concentration is in almost all cases preserved after 2,000 years although boundary concentrations are very high (e.g., 500 wt ppm). However, concentrations in the rim have substantially changed. Right - The original hydrogen content almost completely re-equilibrated to the boundary concentration after 20,000 years at 300 °C.

- 478 300 °C the original hydrogen concentration is almost completely homogenized (Figure 9, right);
- these effects are stronger in smaller crystals / diffusion domains (note that the distance to nearest
  source of hydration / dehydration, e.g. a crack, is relevant in this context rather than the size of
- 481 a grain).
- 482

# 483 6 Conclusion

484

485 Diffusion experiments at low temperatures (195 - 400 °C) and non-isothermal modelling has
486 resulted in the following conclusions:

487

488 (1) We applied an experimental and analytical procedure that enables us to study diffusion 489 of hydrogen in clinopyroxene at low temperatures ( $\leq 400$  °C). It was possible to 490 circumvent previous hindrances by using ion implantation to introduce an artificial 491 hydrogen concentration gradient. Concentration profiles on a nm-scale and changes in 492 these due to diffusion at experimental temperatures were measured using Nuclear 493 Resonance Reaction Analysis (NRRA). Taking advantage of the non-destructive nature 494 of NRRA allowed us to observe the evolution of diffusion profiles with time within one 495 sample, and thereby resolve small differences in concentration profile shapes before and 496 after diffusion. Diffusivities are constant over time and show no concentration 497 dependence. The diffusion-temperature relationship is described by an Arrhenius law  $D_H = 5.47 \ (\pm 13.98) \ \cdot 10^{-8} \ m^2/s \ \cdot \ exp \ [-115.6 \ (\pm 11.5) \ kJ/mol/RT].$ 498

(2) A determination of diffusion coefficients for hydrogen in NAMs was possible only at
high temperatures (> 600 °C) up to now. However, re-equilibration at shallow depths
far from the mantle source would occur at lower temperatures. It was therefore crucial
to experimentally verify whether diffusion rates from high temperatures follow an
Arrhenius relation to lower temperatures. Our experimental results from experiments
between 195 and 400 °C prove that diffusion coefficients of hydrogen in clinopyroxene
can be extrapolated to low temperatures.

(3) Non-isothermal diffusion modelling using the newly obtained diffusion coefficients at
 low temperatures provides information on the degree of re-equilibration as a function
 of crystal size and cooling rate for a specific geotherm. We modelled the ascent of

509	clinopyroxene (0.5, 1.0, 2.0 mm) in a temperature range of 600 to 100 °C while the
510	concentration boundaries are determined by the solubility of hydrogen in clinopyroxene
511	at the respective P-T- $fO_2$ conditions. Small clinopyroxene crystals of 0.5 mm preserve
512	original water contents at cooling rates > 1,000 – 10,000 °C/yr ( $1 \cdot 10^{-3} - 1 \cdot 10^{-2}$ m/s).
513	However, at slow cooling rates (< 10 °C/yr) hydrogen contents in even large crystals of
514	2.0 mm size get substantially modified, thus erasing the hydrogen content equilibrated
515	at mantle depths.
516	
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522	
523 524	Data Availability Statement
525	Programs used in this work are available in the Supplementary Material.
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# 540 Appendix I

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### 542 Development of a non-isothermal diffusion model with variable system boundaries

We modeled diffusion of hydrogen in clinopyroxene of different sizes during their ascent along two geotherms (oceanic and continental) in a low temperature range of 600°C to 100 °C in the presence of a C-O-H fluid that is carbon saturated. A calculation of OH concentration profiles that result from these scenarios required a derivation of parameters that affect diffusion, which is outlined in the following. A temperature-depth relationship was derived after Sclater *et al.* (1980), equations A1-A3 and Table A 1. The quantities in equation A1-A3 are explained in the table caption.

550

551 
$$RHP = \frac{q_{surface} - q_{base}}{h_r} \qquad (eq.A1)$$

552

553 
$$T_{crust} = T_0 + \frac{q_{base^Z}}{k_1} + \frac{(q_{surface} - q_{base}) \times h_r}{k_1} \left(1 - e^{-\frac{Z}{h_r}}\right) \qquad (eq. A2)$$

554

555 
$$T_{mantle} = T_{crust\_base} + \frac{(z - z_{crust})}{k_2} \times q_{base} \qquad (eq.A3)$$

556

**Table A 1**: Parameters that were used to compute geotherms associated with an oceanic (OC) and a continental (CT) setting.  $d_c$  = thickness of the crust,  $\rho_c$  = density of the crust,  $q_{surface}$  = surface heat flow, RHP = radiogenic heat production,  $k_c$ : thermal conductivity of the crust,  $h_r$  = ratio of heat flow to heat production,  $\rho_m$  = density of the mantle,  $q_{base}$  = heat flow base of crust,  $k_m$  = thermal conductivity mantle. 1 = Philpotts and Ague (2009), 2 = Carlson and Herrick (1990), 3 = Christensen and Mooney (1995), 4 = Sclater *et al.* (1980), 5 = Dziewonski and Anderson (1981).

	crust					mantle			
setting	dc	ρc	qsurface	RHP	kc	hr	ρm	<b>q</b> base	km
	(km)	(kg/m <sup>3</sup> )	(mW/m²)	(µW/cm³)	(W/K	(km)	(kg/m <sup>3</sup> )	$(mW/m^2)$	(W/K
					m)				m)
OC	10 <sup>1</sup>	2900 <sup>2</sup>	344	0.89	2.51 <sup>4</sup>	10	3380 <sup>5</sup>	25.1 <sup>4</sup>	3.35 <sup>4</sup>
СТ	$40^{1}$	2830 <sup>3</sup>	46 <sup>4</sup>	1.20	2.514	10	3380 <sup>5</sup>	$34.0^4$	3.35 <sup>4</sup>

The lithostatic pressure along the geotherms was calculated by integrating the densities with 564 565 depth followed by multiplication with the constant for accelearation due to gravity. The oxygen fugacity was assumed to be continuously buffered by a QMF assemblage. Whether hydration 566 567 or dehydration occurs also depends on the solubility of hydroxyl in clinopyroxene at respective conditions and the availability of a hydrogen source (in this case a fluid). The solubility of 568 569 hydrogen in NAM is a function of pressure, temperature, and  $fH_2O$ , but is also affected by contents of certain elements (e.g., Al) and fO2 (Liu and Yang 2020). We used the solubility law, 570 571 equation A4, and experimentally determined values by Liu and Yang (2020) and Bromiley et 572 al. (2004) (dH = 21.2 kJ/mol, dV = 7.3 cm<sup>3</sup>/mol, A = 7.144 ppm/bar, n = 0.5) to describe 573 solubility of hydroxyl (cwater in wt. ppm H2O) in clinopyroxene as a function of temperature, 574 pressure, and  $H_2O$  fugacity (*fH*<sub>2</sub>*O*). The latter is controlled by the speciation of the present fluid, 575 which was determined from thermodynamic modelling of a C-O-H (carbon saturated and fixed 576  $fO_2$  at QMF) fluid as a function of temperature, pressure, and  $fO_2$  following the procedure in 577 Huizenga (2005). The database of fugacity coefficients of the species used in Huizenga (2005) had to be extended by extrapolation in P-T space (between 100 °C and 300 °C), which is a 578 579 sufficient approximation for our purpose. OH contents that were derived from the solubility law 580 were used as boundary concentrations that were constantly changing (i.e., decreasing) and 581 thereby introducing a concentration gradient at the system boundaries.

582

583 
$$c_{water} = AfH_2O^n \exp\left(-\frac{dH^{1bar} + dV^{solid}P}{RT}\right) \qquad (eq.A4)$$

584

585 In the final model, the rate of diffusion is affected by changes in temperature and was thus 586 computed as non-isothermal process by allowing the diffusion coefficient to change according 587 to the respective temperature. The diffusion coefficients were derived through the Arrhenius relationship  $D_H = 5.47 \ (\pm 13.98) \ \cdot 10^{-8} \ m^2/s \ \cdot exp \ [-115.6 \ (\pm 11.5) \ kJ/mol/RT]$  which was obtained 588 589 through our experiments. Effects of pressure and  $fO_2$  on hydrogen diffusion in clinopyroxene 590 are experimentally weakly or not constrained but are expected to play a minor role compared 591 to temperature. The actual diffusion modelling was performed by approximating Fick's second 592 law (equation A5) by an explicit numerical solution (finite difference, equation A6).

593

594 
$$\frac{\partial c(x,t)}{\partial t} = D \frac{\partial^2 c(x,t)}{\partial x^2} \qquad (eq. A5)$$

596 
$$c_{i,j+1} = c_{i,j} + \frac{D \cdot \Delta t}{\Delta x^2} \cdot \left[ c_{i+1,j} - 2c_{i,j} + c_{i-1,j} \right] \qquad (eq.A6)$$

The initial condition (t = 0) for the diffusion model is a homogeneous hydrogen distribution in the crystal. This initial concentration corresponds to the solubility of *OH* in clinopyroxene at the starting P-T-*fH*<sub>2</sub>*O* condition. During the diffusion calculation the diffusion rate  $D_H$  is continuously changed according to the temperature at that time step. All codes that were used in these calculations are provided in the Supplementary Material.

# 625 Appendix II

626

## 627 *Fitting re-equilibration as a function of crystal size and cooling rate*

628 Re-equilibration is defined by equation A7 with  $c_{initial}$  being the initial OH concentration (t = 0), *c*<sub>model core</sub> being the OH concentration in the crystal core after diffusion, *c*<sub>equilibrium at T(final)</sub> 629 being the equilibrium concentration at the final temperature (here 100 °C). The degree of re-630 equilibration is calculated for each crystal size, cooling rate, and geotherm. The re-equilibration 631 632 - cooling rate relationship (Figure S1 and S2 in supplementary information) is then fitted using equation A8 to obtain parameters p1, q1, q2 for each geotherm and crystal size. MATLAB curve 633 634 fitting tool and the implemented non-linear least square procedure (Levenberg-Marquardt 635 algorithm) was used for all fitting procedures that are described in this chapter.

636

637 
$$re - equilibration (\%) = \frac{c_{initial} - c_{model core}}{c_{initial} - c_{equilibrium at T(final)}} \times 100 \qquad (eq. A7)$$

638

639 
$$re - equilibration (\%) = \frac{p_1}{cr^2 + q_1 \times cr + q_2} \qquad (eq.A8)$$

640

641 Table A 2: Parameters *p1*, *q1*, *q2* obtained by fitting re-equilibration – cooling rate relationships for each crystal
 642 size and geotherm.

Geotherm / crystal size	p1	q1	q2	R <sup>2</sup>
CT / 0.5 mm	9.972E+6	1.955E+3	1.118E+5	0.9826
CT / 1.0 mm	9.492E+5	7.668E+2	1.081E+4	0.9955
CT / 2.0 mm	8.962E+4	2.665E+2	1.060E+3	0.9958
OC / 0.5 mm	1.889E+10	1.307E+7	2.067E+8	0.9668
OC / 1.0 mm	4.902E+5	9.293E+2	6.008E+3	0.9899
OC / 2.0 mm	9.555E+4	7.263E+2	1.220E+3	0.9990

643

644 The fitting parameters p1, q1, and q2 were then fitted as a function of crystal size for both

645 geotherms using simple power-law equations (eq. A9-A11).

646

$$p1 = p1a \times sz^{p1b} \qquad (eq.A9)$$



Table A 3: Parameters obtained by fitting re-equilibration as a function of crystal size (sz) and cooling rate (cr)
 for the two geotherms CT and OC.

parameter	СТ	OC
pla	9.4663E+5	9.6171E+6
p1b	-3.399	-8.8003
R <sup>2</sup>	1.00	1.00
qla	7.3650E+2	2.0662E+4
q1b	-1.4375	-7.0677
R <sup>2</sup>	0.99	1.00
q2a	1.0861E+4	1.1485E+5
q2b	-3.3604	-8.6851
R <sup>2</sup>	1.00	1.00

657 Substituting parameters p1, q1, and q2 in equation A8 by the power-law equations A9-A11 658 above yields the following equation A12 that was used to calculate re-equilibration in % as a 659 function of crystal size *sz* in mm and cooling rate *cr* in °C/yr for both geotherms.

661 
$$re - equilibration (\%) = \frac{p_{1a \times sz^{p_{1b}}}}{cr^2 + q_{1a \times sz^{q_{1b}} \times cr} + q_{2a \times sz^{q_{2b}}}}$$
(eq.A12)

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