



# Mineral replacement rate of olivine by chrysotile and brucite under high alkaline conditions

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Romain Lafay, German Montes-Hernandez, Emilie Janots, Rodica Chiriac, Nathaniel Findling, et al.. Mineral replacement rate of olivine by chrysotile and brucite under high alkaline conditions. Journal of Crystal Growth, Elsevier, 2012, 347 (1), pp.62-72. <10.1016/j.jcrysgro.2012.02.040>. <insu-00719186>

## HAL Id: insu-00719186 https://hal-insu.archives-ouvertes.fr/insu-00719186

Submitted on 19 Jul2012

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#### 16 Abstract

Olivine mineral replacement by serpentine is one major alteration reaction of oceanic 17 hydrothermalism. In the present experimental study, olivine grains were replaced by 18 chrysotile and brucite under high alkaline conditions. In our study, olivine replacement 19 implied a spatial and temporal coupling of dissolution and precipitation reactions at the 20 21 interface between olivine and chrysotile-brucite minerals. Coupled dissolution-precipitation led to the alteration of starting olivine grains (so-called primary or parent mineral) to a porous 22 mineral assemblage of chrysotile and brucite with preservation of the initial olivine 23 24 morphology. This mineral replacement reaction of olivine (serpentinization) has been characterized using XRD, FESEM and FTIR measurements. Moreover, a simple and novel 25 method is here proposed to quantify the mineral replacement rate (or serpentinization rate) of 26 olivine by using thermogravimetric (TG) and differential TG (DTG) analyses. 27 Serpentinization extent depends on the grain size: it is complete after 30 days of reaction for 28 the smallest olivine grains (<30µm), after 90 days of reaction for the intermediate olivine 29 grains (30 µm-56 µm) and reaches 55% of olivine replacement after 90 days for the largest 30 fraction (56-150  $\mu$ m). Based on the fitting of the serpentinization extent ( $\xi_t$ ) versus time (t) 31 by using a kinetic pseudo-second-order model, the serpentinization rates vary from  $3.6 \times 10^{-6}$  s<sup>-</sup> 32 <sup>1</sup> to  $1.4 \times 10^{-7}$  s<sup>-1</sup> depending on the olivine grain size. An additional correlation between FTIR 33 spectra analysis and TG measurements is proposed. The mineral replacement reactions 34 frequently observed in natural alteration processes could be a powerful synthesis route to 35 36 design new porous and/or nanostructured materials.

*Keywords:* A1. Serpentinization; A1. Mineral replacement rate; B1. Alkaline medium; B2.
Chrysotile nanotubes; A1. TG analyses

#### 40 **1** Introduction

Serpentine minerals (chrysotile, lizardite and antigorite) are widespread in Earth 41 oceanic lithosphere and are frequently found in chondrites and other extraterrestrial objects. 42 Serpentine mineralization is of great interest in several fields of research. Serpentinized rocks 43 present a great enrichment in trace elements compared to primary mantle rocks [1-6]. 44 45 Serpentine appears as a vector for trace elements between crustal and mantle reservoirs [2,5,7,8]. Experimental studies have tested the influence of major (e.g. Fe, Ni) and/or trace 46 elements (e.g. Li) on the growth of serpentine [9-14]. This kind of synthesis experiments 47 presents a great interest in medical research due to the asbestos toxicity that can be induced by 48 inhalation of magnesium silicates fibers including chrysotile [15-18]. Serpentine minerals are 49 also crucial for sequestration of  $CO_2$  due to its availability and sequestration capacity [19–21]. 50 Indeed a lot of studies are looking for technologies that could possibly contribute to reduce 51 carbon dioxide emissions. Geological sequestration and ex-situ mineralization of CO<sub>2</sub> using 52 53 serpentine could be one of the most efficient methods considering the enormous quantity of serpentine on Earth [22]. 54

In meteorites, serpentine minerals are directly linked to aqueous alteration processes and 55 reaction conditions (e.g. [23] and references therein). In the oceanic lithosphere, serpentines 56 result from interaction between mantle rocks (peridotite composed by olivine and pyroxenes) 57 and hydrothermal fluids, generally with high fracturation dynamic [24,25]. Olivine alters 58 along grain boundaries and fractures to produce a mesh texture that preserves the original 59 olivine morphology at the grain scale [26,27]. This olivine replacement by serpentine is best 60 explained by coupled dissolution-precipitation processes [28–30]. This reequilibration process 61 leads to the replacement of one pristine mineral by a secondary mineral (or assemblage) with 62 a lower solubility in the fluid. Replacement occurs at the fluid/solid interface maintaining the 63 original external grain shape (pseudomorphism). During alteration, a secondary porosity is 64

commonly produced due to volume difference between pristine and secondary minerals, 65 66 material loss during dissolution and grain fracturation [24,25]. Secondary porosity enables the fluid to move interstitially towards the reaction front until the complete mineral replacement 67 reaction. In oceanic lithosphere, peridotite replacement, and consequent element redistribution 68 associated with this alteration, is primarily controlled by the physic-chemical conditions of the 69 hydrothermal fluid (Temperature, Pressure, fluid speciation, pH). Fluids escaping from deep 70 sea hydrothermal vents show a large variety of composition and pH, reflecting a large range 71 of possible physic-chemical conditions. Amongst them, alkaline fluids with high pH were 72 collected in some hydrothermal fields (e.g., [31,32]). 73

Numerous experimental studies were conduct to reproduce serpentinization in hydrothermal context [33–40] and explain serpentine growth [41]. Kinetic appears faster under alkaline conditions [40,42] but few recent studies have addressed the role of pH on serpentinization kinetic, particularly in alkaline conditions.

In the present experimental study, we have investigated the process and kinetics of 78 79 olivine serpentinization in alkaline hydrothermal conditions (pH = 13.5, measured at 25°C). Experimental products were characterized using XRD, FESEM and FTIR. The 80 serpentinization rate was determined using a simple and novel method based on 81 thermogravimetric analyses (TGA/DTGA). This demonstrates that serpentinization is much 82 faster under alkaline conditions refering to previous study at comparable conditions 83 [35,36,43] and can lead to total replacement of  $<30\mu m$  olivine in less than 30 days and 90 84 days for 30<particle size<56 µm. 85

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#### 2 Materials and Methods

87 Millimetric grains of olivine San Carlos (Fo<sub>90</sub>) were crushed by using a Fritsh 88 Pulverisette 7 micro-crusher. Three classes of grain/particle size (particle size $<30\mu$ m, 89 30<particle size $<56\mu$ m and 56<particle size $<150\mu$ m) were isolated by sieving. The samples

90 were washed three times using high-pure water in order to remove the ultrafine particles that 91 possibly stuck at grain surfaces during crushing step. Optical and electron microscopy was 92 performed to control the initial state/appearance of olivine surfaces.

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#### **2.1** Alteration experiments

1.5ml of high alkaline solution (1M of NaOH, pH≈13.5 at 25°C) and 100mg of San 94 Carlos olivine at a given class of particle size were placed in a Teflon cell reaction (cap-cell 95 also in Teflon). Cell reaction and cap-cell were previously washed by an acidic treatment 96 97 followed by washing with high-pure water. This cell reaction was immediately assembled into a steel autoclave without agitation, referred as "static batch reactor" and the closed autoclave 98 was placed in a multi-oven (ten independent-temperature compartments) and heated to 150°C 99 or 200°C (P<sub>sat</sub>~0.5Mpa and P<sub>sat</sub>~1.6Mpa). Various olivine-alkaline solution reaction times 100 from 3h to 90 days were considered in order to determine the mineral replacement rate (or 101 serpentinization rate) of olivine at the investigated hydrothermal conditions. Particularly, the 102 effect of initial particle size (three different classes: particle size<30µm, 30<particle 103 104 size<56µm and 56<particle size<150µm) on the serpentinization rate of olivine was tested. All experimental conditions are summarized in table 1. 105

At the end of the experiment, the autoclave was quenched in cold water. This manipulation avoids precipitation during cooling with no perturbation of the reaction products. Then, the autoclave was disassembled and the fluid was collected for pH and major elements concentration (Mg, Fe and Si) measurements (not shown in this study). After all experiments, pH of the experimental fluid is still around 13.5 (measured at 25°C). The solid product was dried directly in the Teflon cell at 80°C for 24h. The dry solid product was recovered for further solid characterizations described below.

#### 2.2 X-Ray diffraction measurements 113

X-Ray diffraction (XRD) analyses were performed in the ISTerre laboratory. All 114 samples were crushed manually. The powders were carefully placed and manually compacted 115 in borosilicated capillaries of 500µm of diameter, this corresponds to about 5 mg of sample. 116 XRD patterns were recorded with a Bruker D8 powder diffractometer equipped with a SolX 117 118 Si (Li) solid state detector from Baltic Scientific Instruments using CuKa1-Ka2 radiation and a Göbel mirror. Intensities were recorded for an  $2\theta$  interval from 5 to 80° with an 8 s counting 119 time per  $0.024^{\circ} 2\theta$  step for bulk mineralogy determination. 120

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#### 2.3 **FESEM** observations

Grain size and morphology of the experimental products were characterized using 122 secondary or backscattering electrons. Micro-imaging was obtained using Zeiss Ultra 55 Field 123 124 emission gun scanning electron microscopy (FESEM). This enables a spatial resolution of approximately 1nm at 15 kV. Samples were dispersed by ultrasonic treatment in absolute 125 ethanol for at least 5mn in order to disaggregate the particles. One or two drops of dispersion 126 were placed onto an aluminium support and coated with a thin film of platinium for SEM 127 observation. 128

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### 2.4 Thermogravimetric analyses (TGA)

TGA for experimental solid products were performed with a TGA/SDTA 851<sup>e</sup> Mettler Toledo 130 instrument under the following conditions: sample mass of about 10 mg, platine crucible of 131 150  $\mu$ l with a pinhole, heating rate of 10 °C min<sup>-1</sup>, and inert N<sub>2</sub> atmosphere of 50 ml min<sup>-1</sup>. 132 Sample mass loss and associated thermal effects were obtained by TGA/DTGA in a 133 temperature range from 30°C to 1200°C. In order to identify the different mass loss steps, the 134 TGA first derivative (rate of mass loss) was used. TGA apparatus was calibrated in terms of 135 mass and temperature. Calcium oxalate was used for the sample mass calibration. The melting 136 points of three compounds (indium, aluminium and copper) obtained from the DTGA signals 137

were used for the sample temperature calibration. The temperature accuracy of the TGA/SDTA system is about  $\pm 0.25$ °C. The weighting accuracy is around 0.1µg, which correspond to 0.01% for a 10mg sample.

- 141
- 142 **2.4**

#### 2.5 FTIR measurements

FTIR measurements (in transmission mode) were performed using an IR microscope Bruker Hyperion 3000. The IR beam was focused through a 15x lens and the typical size of infrared aperture is  $50*50 \ \mu\text{m}^2$ . The light source is a Globar(TM) and the beam splitter is in KBr. The spectra were measured from 700 to 4000 cm<sup>-1</sup> (4cm<sup>-1</sup>resolution) with a MCT monodetector cooled by liquid nitrogen.

Samples must be thin (less than 100  $\mu$ m) and flat to avoid absorption band saturation or scattering effects. Sample preparation has involved a careful crushing of samples in mortar and manual compaction of fine crushed particles between two K-Br windows. Five spectra per samples were realized; they were fitted using OPUS software.

#### 152 **3 Results and discussion**

#### **3.1 Serpentinization reaction under alkaline conditions**

Secondary minerals were identified by XRD and FESEM (Figs. 1 and 2) and they were quantified by TGA (Fig. 3). Under alkaline conditions, olivine is replaced by chrysotile and brucite, independently on the starting grain size of olivine. No other minerals were detected and/or observed during this alteration reaction.

FESEM micro-imaging has revealed a clear evolution of particle size and morphology of crystal faces during serpentinization advancement (Fig. 2). Serpentine nucleation at olivine-alkaline solution interfaces is observable by FESEM after only few hours of reaction when the starting grain size is  $<30\mu$ m (Fig. 2a). These occur as nanosized nodules forming irregular sub-micrometric rods. They are still observed onto olivine surfaces after few days of

reaction and a more advanced serpentinization reaction (Fig. 2c). The irregular sub-163 micrometric rods were identified as chrysotile and brucite "nucleates" from TGA/DTGA in 164 the solid product (Fig. 3). While brucite is detected by XRD in the first stages of replacement 165 (e.g. 3 hours of reaction for <30µm grains), chrysotile was not detected by XRD, possibly 166 due to its poor crystallinity and/or nanometric size. Chrysotile was detected by XRD after 16h 167 of reaction for the smallest starting grains (<30um), however, several days of olivine-fluid 168 interactions are required when the starting grain size is >30µm (Fig. 1a, b). At advanced 169 alteration stages, serpentinized samples consist of tubular, conical and cylinder-in-cylinder 170 chrysotile with nanometric diameter and microsized hexagonal brucite were clearly observed 171 (Fig. 2e, g, h). Diameter of secondary products depends of the starting material grain size. It 172 increases from  $\sim$ 50 nm for the <30  $\mu$ m, to 200-300 nm for 30-56  $\mu$ m and 56-150  $\mu$ m fractions. 173 Lizardite was not observed. Previous study suggested that lizardite crystallized at lower levels 174 of supersaturation than chrysotile [44]. Here, we confirm that high alkaline pH favors 175 chrysotile precipitation [40,42]. 176

In natural samples, olivine replacement in mesh texture is commonly described as an 177 assemblage of serpentine±brucite±magnetite. Typically, the magnetite amount depends on the 178 Mg# (Mg# = Mg / (Mg + Fe)) in primary olivine. Neither iron oxides (ex. magnetite or 179 hematite) nor oxyhydroxides (ex. goethite) were observed in our experiments. Semi-180 quantitative EDS analyses have revealed that the initial iron contained in olivine was 181 preferentially incorporated as  $Fe^{2+}$  into brucite (Mg#<sub>brucite</sub>=0.84), and slightly sequestered 182 into/onto chrysotile (Mg#chrysotile=0.95) compared to starting San-Carlos olivine. In our 183 conditions a part of Fe<sup>3+</sup> (not determined) might substitute to Si into the chrysotile. In our 184 experiments, one general reaction of San Carlos olivine alteration can be written as: follows: 185

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$$2Mg_{1,8}Fe_{0,2}SiO_4 + 3H_2O = (Mg_{1-Z}Fe_Z)(OH)_2 + (Mg_{1-X}Fe_X)_3Fe_YSi_{2-Y}(OH)_4$$
 (1)

188 This isochemical reaction implies a volume increase of  $\sim 25\%$  (see discussion below).

189 On the XRD patterns (Fig. 1), characteristic peaks of chrysotile appear in the first hours or days of reaction, depending on the initial particle size of olivine (Fig. 1a, b). Secondary 190 191 product growth rate is related to the variation of the crystal size with time. On the XRD patterns the full width at half maximum (FWHM) parameter for specific peaks, corresponding 192 to a crystalline compound, can be directly associated to its coherent domain size when 193 194 instrumental resolution function (IRF) and strain contribution are known. In our case, the FWHM for the plane (002) of chrysotile was used to obtain an estimation of the chrysotile 195 size variation during the serpentinization process. Here, a decrease of FWHM implies an 196 197 increase of coherent domain size as specified in the Scherrer equation. The FWHM variation with the reaction time reported in figure 1c shows a fast crystal growth step followed by a 198 slow crystal growth step for chrysotile from 3h to 30 days when starting grain size is <30µm. 199 200 Conversely, chrysotile coherent domain varies only moderately from 20 to 90 days for starting grain size  $> 30 \,\mu\text{m}$  (Fig. 1c). In this latter case, two explanations are possible: firstly, 201 a preferential growth of other(s) crystal plane(s) than plane (002) is involved, secondly, a 202 dominant nucleation events promoting an increase of particle number concentration with time 203 during mineral replacement of olivine can occur. FESEM observations support the first 204 205 assumption because they have revealed a higher radial growth of tubular chrysotile for larger starting grain size (30-56µm and 56-150 µm) at similar reaction time (Fig. 2f, h). 206

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#### **3.2** Determination of serpentinization rate

The time for complete olivine-to-serpentine transformation or complete 208 serpentinization process depends on the starting grain size. For example, 30 days were 209 required when starting grain size was <30µm while about 90 days were required for the 30-210 56µm fraction (Fig. 1a, b). For 56-150µm starting grain size, around 55% of mineral 211 replacement was reached after 90 days of experiments. 212

Serpentinization rate of olivine was determined by using a simple and novel method 213 214 combining TGA and DTGA. TGA/DTGA were performed to quantify the molecular water (adsorbed or confined in pores) and hydroxyl groups (-OH) (incorporated in brucite and 215 chrysotile) as illustrated in figure 3. In general, molecular water (H<sub>2</sub>O) adsorbed or confined 216 in pores onto mineral assemblage, was released at moderate temperature (<200°C). 217 Conversely, the brucite and chrysotile minerals were dehydroxylated at higher temperatures. 218 Here, a dehydroxylation peak closed to 370°C was associated to brucite. This peak was 219 shifted towards lower temperature, compared to the dehydroxylation of pure brucite peaking 220 at 430°C. This change of thermal stability was related to a significant incorporated amount of 221 iron (Fe<sup>2+</sup>) into brucite structure as recently claimed by Okamoto et al. [45]. Additionally we 222 observed that weight loss ratio between serpentine and brucite is not constant following 223 reaction advancement. Regarding the chrysotile, it was observed that it starts to be 224 dehydroxylated at about 400°C and it was completely dehydroxylated at approximately 700°C 225 for most of the experiments. In several cases, two typical dehydroxylation episodes were 226 measured at 515 and 600 °C, respectively. We note that the starting dehydroxylation step for 227 chrysotile can be overlapped with residual -OH from dehydroxylation of brucite, possibly 228 producing a slight over-estimation of incorporated -OH into the chrysotile phase. This direct 229 230 measurement was used to calculate the serpentinization extent (%) at a given reaction time as follows: 231

232 
$$\xi_t = ((-OH)_{\text{measured}}/(-OH)_{\text{theoretical}})*100$$

where (-OH)<sub>theoretical</sub> is the theoretical weigh (in %) of hydroxyl groups incorporated in chrysotile (400°C to 700°C) with a chemical formula simplified to Mg<sub>3</sub>Si<sub>2</sub>O<sub>5</sub>(OH)<sub>4</sub> for the (-OH)<sub>theoretical</sub> considered. Then, the serpetinization extent was correlated with reaction time ( $\xi_t$ vs. t). These experimental-calculated data is displayed in figure 3. As previously stated, a complete serpentinization extent was determined after 30 days for smaller olivine grains

(2)

 $(<30\mu m)$  and after 90 days for intermediate olivine grains (30<grain size<56). These results clearly show that the serpentinization rate strongly depends on the starting grain size of olivine. This explains why for larger olivine grains (56<grain size<150µm), about 55% of serpertinization extent was only reached after 90 days of olivine-fluid interactions.

Experimental data reported in figure 4 were fitted by using a kinetic pseudo-secondorder model. This simple kinetic model describes a fast serpentinization step followed by a slow serpentinization step until an asymptotic maximum with time. Its integrated form can be expressed as follows:

$$\xi_{t} = \frac{\xi_{\text{max}}t}{t_{1/2} + t}$$
(3)

where  $\xi_t$ , is the serpentinization extent for a given reaction time *t*,  $\xi_{max}$ , the maximum 247 serpentinization extent (close to 100% for a complete serpentinization) and  $t_{1/2}$  the half 248 serpentinization time. The fast serpentinization step can be associated to the fast dissolution of 249 smaller olivine grains initially far from equilibrium, but the dissolution process was spatially 250 and temporally coupled to a fast precipitation of chrysotile and brucite. For this reason, the 251 252 external shape of starting olivine grains (primary or parent mineral) was preserved, leading to a porous granular material composed of chrysotile and brucite minerals when complete 253 mineral replacement of olivine was reached. Finally, the mineral replacement initial-rate  $v_0$ 254 (or serpentinization initial-rate) was defined as the slope of the tangent line when time t tends 255 toward zero on the r vs t curve. Analytically,  $v_0 = (\xi_{\text{max}}/t_{1/2})$  and it varies from  $3.6 \times 10^{-6} \text{ s}^{-1}$  to 256  $1.4 \times 10^{-7}$  s<sup>-1</sup> respectively for  $<30 \mu m$  grain size and 56-150  $\mu m$  grain size. Full serpentinization 257 was not obtain experimentally on 56<particle size<150 µm but the theoretical model 258 presented predict a close to complete serpentinization (>90%) around 2 years. 259

Indeed, the fluid/olivine interface depending of the initial olivine grain size is the main factor controlling the initial rate. These kinetic parameters concerning the three different size classes of olivine and respective correlation factors are summarized in table 2.

The uncertainties of the TG-method result from the calculation method. The main approximation is the systematic consideration of the same temperature range (400-700°C). Refering to all the experiments, a variation of the temperature range of  $\pm$  50°C will only induce a difference of 1- 5% for the final serpentinization advancement calculated. Based on this calculation we assure that the maximum uncertainty of the proposed methods is around 5% as reported in figure 4. The uncertainty on the calculated initial rate is directly depending on the standard deviation obtained for t<sub>1/2</sub> and  $\xi_{max}$  (Table 2).

In a complementary way, the transmission FTIR spectra were obtained. Characteristic 270 stretching bands for hydroxyl group (-OH) in chrysotile-brucite and for Si-O group in olivine-271 chrysotile were used. Particularly, the relative (-OH)/(Si-O) ratio using integrated bands 272 intensities or maximum band intensities was correlated with the serpentinization extent 273 deduced from TG/DTG analyses (Fig. 5). This gives two calibration curves that can be used to 274 determine serpentinization rate using FTIR measurements. However, FTIR measurements 275 should be taken with caution because of the overlapping of –OH (for brucite and serpentine) 276 and Si-O groups (for serpentine and olivine). We could overcome this problem if 277 serpentine/brucite produced molar ratio stayed constant following reaction time, but as 278 described before it is not the case. Here, we conclude that the TG/DTG analyses are a 279 powerful tool to determine the mineral replacement rate of olivine by chrysotile-brucite 280 mineral assemblage. For this specific case, the FTIR measurements provide a first information 281 about the olivine serpentinization extent. 282

This study confirms that the serpentinization rate is enhanced by high alkaline conditions. Serpentinisation rate has been studied for a large variety of experimental

conditions under neutral conditions [33–40,46]. For comparison, serpentinization extent of 50-100  $\mu$ m starting grain size at 200°C reached only ~20% after ~4 months under neutral conditions (Seyfried et al., 2007). This corresponds to a kinetic 5 times slower to our results on the largest starting grain size (56-150  $\mu$ m).

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3.3

## Serpentinization steps and reaction mechanism

Contemporaneously, the first steps of reaction in alkaline medium are characterized by 290 a fast dissolution of olivine associated to the formation of preferential dissolution figures as 291 292 notches and etch pits (Fig. 6a, b, c) [44–46]. Conjointly, precipitation of the first nucleates of brucite and chrysotile occur at the olivine surface. FESEM observations have revealed the 293 preservation of olivine shape after complete serpentinization (Fig. 7). This suggests that 294 olivine alteration invokes coupled dissolution-precipitation processes [28]. Despite ultrasonic 295 treatment and dispersion on ethanol (up to 1h30), which can easily break up the new 296 brucite/chrysotile mineral assemblage, the shape and size of olivine grains are always 297 preserved regardless the completion of the reaction. The few scattered chrysotile grains 298 observed are the result of tearing from olivine grain surfaces and are not the result of 299 precipitation from saturated solution away from olivine grain. Based on these results, we 300 301 assume a higher super-saturation of solution with respect to chrysotile tubes at the olivinesolution interfaces. Supersaturation at olivine-alkaline solution interfaces allows rapid 302 heterogeneous serpentine nucleation as small nodules of chrysotile and brucite (Fig. 2a). This 303 nucleation phase is followed by a phase of epitaxial growth of chrysotile and brucite forming 304 a porous medium (Fig. 6d, e, f). Chrysotile grows as elongated tubes with various morphology 305 (conical, cylinder in cylinder) with a size depending on starting olivine grain size. Largest 306 starting grain size induces smallest reacting surface and thus slower kinetic of alteration and 307 formation of largest chrysotiles particles. Olivine grain size influenced initial dissolution rate 308  $v_0$ , passivation state and general kinetic of alteration. 309

Serpentinization reaction is then favored by the formation of notches connected to a 310 micro-fracturing network inside olivine (e.g. 32 days of reaction for >30 µm grains; Fig. 6b, 311 c). This micro-fracturing network might start following ordered families of fine-scale 312 313 fractures from olivine grains [44]. In the oceanic crust, the microfracturing may be attributed to the stress generated by the volume increase during olivine serpentinization [28]. In our free 314 fluid system, there is no evident expansion during olivine replacement. Volume increase due 315 to olivine serpentinization (25%) is likely accounted by material loss in solution, but it could 316 also cause the microfracturing [28,50]. With the formation of smaller sub-grains, 317 microfracturing allows an easier fluid percolation and increase the total reaction-surface (Fig. 318 319 6e, f). This process permits a continuous and total replacement of olivine by chrysotile and brucite as summarized in figure 8. This is in agreement with natural observation and 320 development of mesh type texture described in natural rocks [26-28,50-52]. Fluid can 321 percolate everywhere without formation of a passivation layer at grain surface. The formation 322 of a protective layer would induce a non-continuous passivation process by blocking intra-323 particles diffusion [53]. Mass transfer will play a major role for the preservation of volume 324 and shape of starting olivine grain. 325

326

#### 327 **4** Conclusion

Olivine alteration was investigated under alkaline conditions for different starting grain sizes at 200 °C. In this study, we were able to follow complete olivine replacement by an assemblage of chrysotile and brucite.

Thermogravimetric analyses were used to investigate the dehydroxylation of hydrated phases and thus the serpentinization extent as a function of time. Based on this innovative approach, we were able to estimate punctually the serpentinization advancement and a kinetic pseudo-second-order model was used to describe experimental data. This could allow a
 prediction of reaction advancement for the largest starting grain size.

We also established two calibration curves between the -OH/Si-O bands measured in FTIR and the degree of serpentinization. This enables to have an estimation of the kinetic rates of serpentinization directly from the FTIR spectra analysis. This must be considered with caution because of the overlapping of –OH bands between brucite and serpentine.

Under our experimental conditions, olivine is replaced by chrysotile and brucite by coupled dissolution-precipitation processes. Fast dissolution of olivine, preferential dissolution features formation and process of grain subdivisions by fracturing are involved to explain a complete mineral replacement of olivine. This implies the preservation of external shape of olivine grains as typically observed in natural systems.

345

### 347 Acknowleggements

348 The authors are grateful to French National Center for Scientific Research (CNRS and

349 University Joseph Fourier (UJF) in Grenoble for providing the financial support. R. Lafay was

350 supported by a Ph.D grant from French education ministry. The authors are grateful to IPAG

institute where infrared measurements where performed.

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#### 504 Figure Captions:

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506 Table 1: Summary of the experimental conditions.

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Table 2: Summary of kinetic parameters obtained from the pseudo-second-order model.

509

Figure 1: XRD patterns following reaction advancement, a) with starting grain size  $<30\mu$ m, b) with starting grain size 30-56  $\mu$ m, c) example of plot for the peak corresponding to bragg angle (20) of 12.1° and miller indice (002) with a gaussian deconvolution and plot of FWHM following reaction time for this peak. O: olivine, B: Brucite and S: serpentine.

Figure 2: Scanning electron microscope imaging of the experimental products. All shown 514 experiments have been performed at pH=13.5 (at 25°C) and 200°C, for <30 µm (a, b, c and 515 e), 30-56 µm (d, f, g) and 56-150µm (h) grains size. a) chrysotile and brucite nodules after 516 3hours of reaction (run no.1), b) chrysotile tube after 30 days of reaction (run 8) and c) 517 518 nodules aggregated in a tubular shape after 3 days of reaction (run no.5), d) chrysotiles "bunch" at olivine grain surface after 10 days of reaction (run no.13), e) characteristic 519 brucite/chrysotile assemblage after 30 days of reaction (run no.6), f,g) characteristic 520 521 brucite/chrysotile assemblage and thick cylinder-in cylinder chrysotile after 90 days of reaction (run no.18), and h) characteristic brucite/chrysotile assemblage after 32 days of 522 reaction (run no.20). Bru: brucite, Ctl: chrysotile, Ol: olivine. 523

524

Figure 3: a) Thermogravimetric analyses (TGA) for experiments of 3 hours to 30 days (run no. 1,3,5,6,7and 8) with fraction grains  $<30 \mu m$ , b) focus and decomposition of TGA and DTG corresponding spectra after 10 days of reaction.

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Figure 4: Serpentinization degree versus time for the hydration reaction at different grain sizes for T=200°C,  $P_{sat}$ =1.6Mpa and fluid/solid=15. Maximum error bars corresponding to 5% of the serpentinization extent calculated was also reported. Data were fitted by using a kinetic pseudo-second-order model.

534 Figure 5: a) Transmission spectra (in absorbance) for starting grain size <30µm after 3 hour (run no.1) and 30 days (run no.8) of reaction with corresponding base line and range 535 employed for treatment. Blue area correspond to -OH band integration A1 with maximum 536 intensity I1, green area correspond to Si-O band integration A2 with maximum intensity I2, b) 537 corresponding -OH/Si-O ratio: A1/A2 and I1/I2 following serpentinization advancement 538 determined by TG and DTG analysis, square, triangle and circles correspond respectively to 539 experiments for <30µm, 30-56µm and 56-150 µm starting grain size fraction (empty for 540 A1/A2 and black for I1/I2) with corresponding standard deviation corresponding, black line 541 correspond to best linear regression with the exclusion of red point. 542

Figure 6: Scanning electron microscope imaging of the experimental products for starting grain size  $<30 \ \mu\text{m}$  (a), 30-56  $\ \mu\text{m}$  (c-f) and 56-150  $\ \mu\text{m}$  (b). a) etch pits at olivine grain surface after 3 days (run no.5), b) dissolution notches after 32 days of reaction (run no. 20), c) dissolution notches after 24 days of reaction (run no. 14), d) chrysotile micro-tubes at olivine grain surface after 3 days of reaction (run no. 12), e) contact between two olivine grains and f) microfracturing in section after 10 days of reaction (run no.13). Red dotted lines represent initial border of the supposed crack. Ctl: chrysotile, OI: olivine.

Figure 7: XRD pattern and corresponding scanning electron microphotography from pure starting San Carlos olivine to close to 100% of replacement by serpentine and brucite. Note the preservation of the olivine grains dimension and shape. S: serpentine, B: brucite, O: olivine.

554 Figure 8: Schematic synthesis presenting the different steps of olivine replacement by 555 chrysotile and brucite.

Run no.	T (°C)	Time	grain size (μm)	Olivine to Chrysotile conversion (%)	products starting-material
1	200	3 h	<30 μm	10.3	O>C>B
2	200	16 h	<30 μm	21.7	O>C>B
3	200	24 h	<30 μm	24.6	O>C>B
4	200	48 h	<30 µm	37.5	O>C>B
5	200	72 h	<30 µm	51.7	O≈C>B
6	200	10 day	<30 μm	73.8	C>O>B
7	200	20 day	<30 μm	88.8	C>B>O
8	200	30 day	<30 μm	98.0	C>B
9	200	90 day	<30 μm	98.2	C>B
10	150	30 day	<30 μm	49.5	O≈C>B
11	150	66 day	<30 μm	not determined	C>O>B
12	200	3day	30-56 μm	19.6	O>C>B
13	200	10 day	30-56 μm	26.2	O>C>B
14	200	24 day	30-56 μm	46.2	O≈C>B
15	200	41 day	30-56 μm	77.1	C>O>B
16	200	51 day	30-56 μm	71.1	C>O>B
17	200	72 day	30-56 μm	88.9	C>B>O
18	200	90 day	30-56 μm	94.4	C>B>O
19	200	13 day	56-150 μm	10.7	O>C>B
20	200	32 day	56-150 μm	26.6	O>C>B
21	200	60 day	56-150 μm	44.2	O>C>B
22	200	90 day	56-150 μm	55.4	C>O>B

556 Table 1

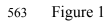
557 All experiments have been performed at pH 13.5 (measured at 25°C). Saturated pressure is

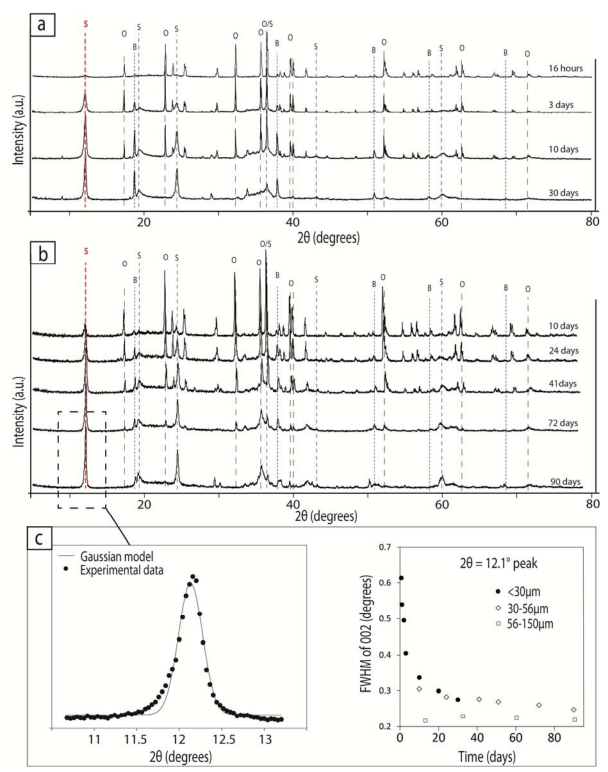
constant: 1.6MPa for 200°C and 0.5 MPa for 150°C reactions. Fluid/rock ratio is always ~15.

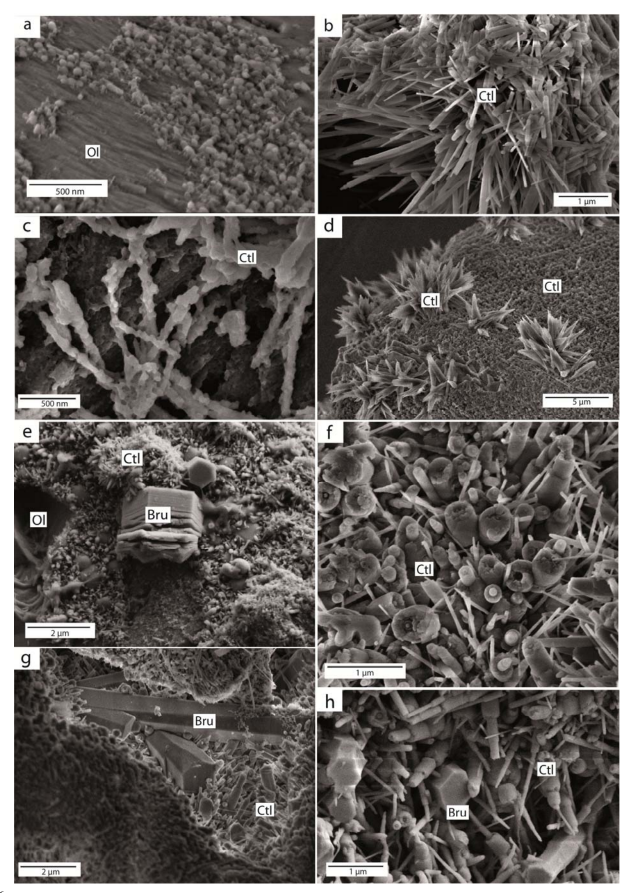
559 O: olivine, C: chrysotile, B: brucite.

## 560 Table 2

Initial particule size	Smax (%)			Initial rate	···· 2
of olivine	Exp.	Calc.	t ½ (days)	(1/s)	fitting r <sup>2</sup>
<30 μm	97.96	100±1.7	3.2±0.3	3.62-06±0.35E-06	0.989
30-56 μm	94.35	100±8.5	23.1±5.3	5.01E-07±0.9E-07	0.947
56-150 μm	55.4	100±16	79±21	1.46E-07±0.2E-07	0.981

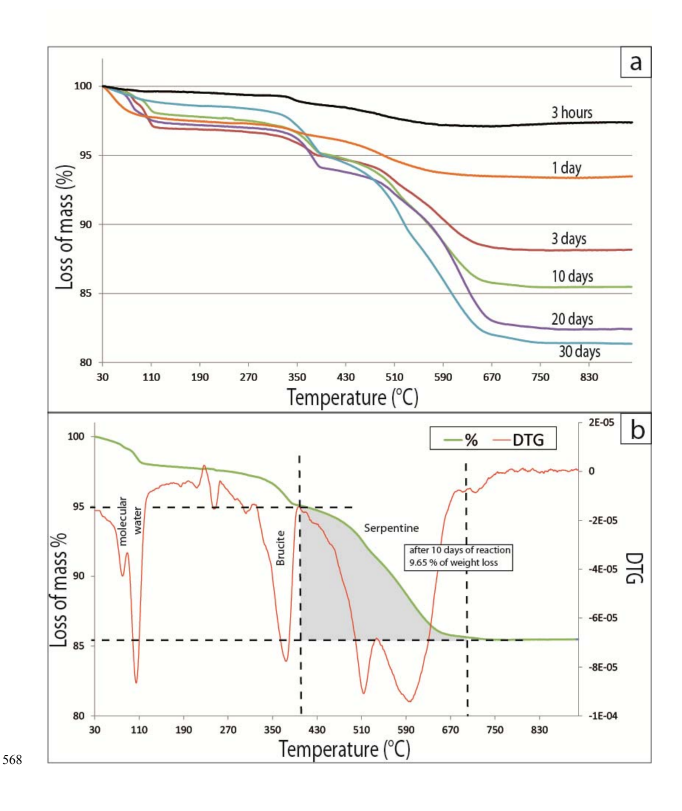


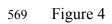






567 Figure 3





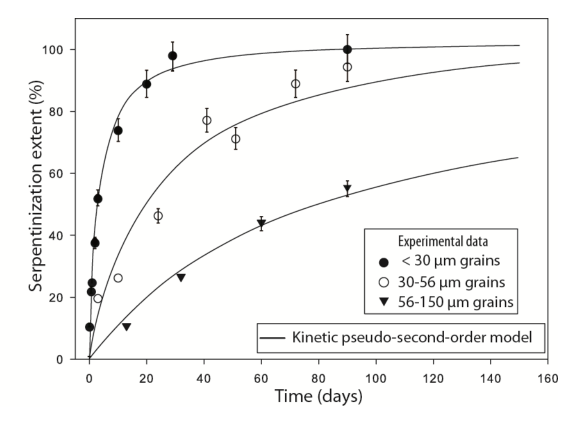
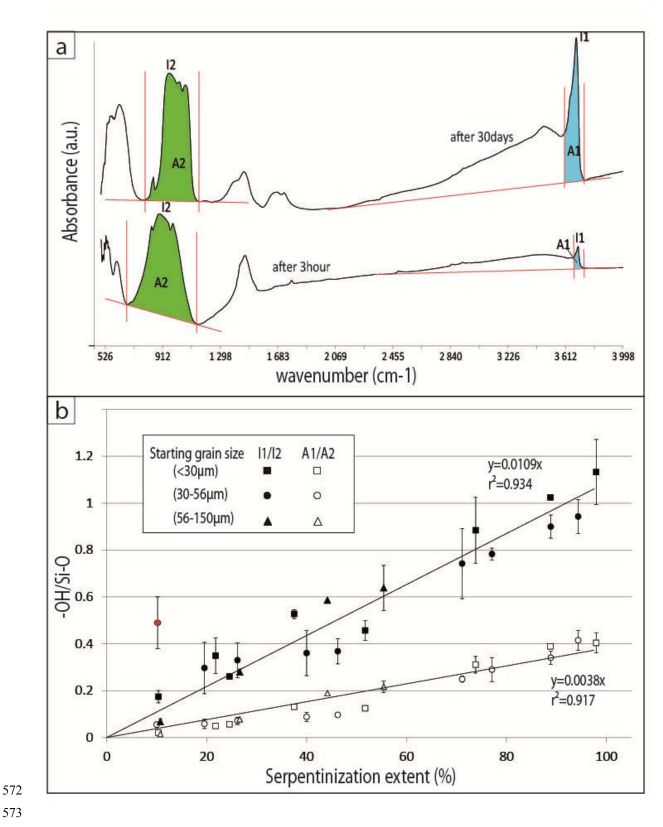


Figure 5 



574 Figure 6

