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Dissolution-precipitation processes governing the carbonation and silicification of the serpentinite sole of the New Caledonia Ophiolite

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Marc Ulrich^{1,2,#,*}, Manuel Muñoz², Stéphane Guillot², Michel Cathelineau¹, Christian Picard³,

- 5 Benoit Quesnel⁴, Philippe Boulvais⁴, Clément Couteau⁵.
- 6 ¹Laboratoire Géoressources, CNRS, UMR 7566, Université de Lorraine, France.
 - ² Institut des Sciences de la Terre, CNRS, UMR 5275, Université de Grenoble 1, France.
- 7
 ³ Laboratoire Chrono-environnement, UMR 6249, Université de Franche-Comté, France.
- 8 ⁴ Géosciences Rennes, CNRS, UMR 6118, Université de Rennes 1, France.
- 9 ⁵ Service géologique, Koniambo Nickel SAS, Nouvelle Calédonie.
- [#] present location: IPGS-EOST, UMR 7516, Université de Strasbourg, France
- 10 * corresponding author: <u>mulrich@unistra.fr</u>
- 11

12 ABSTRACT

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14 The weathering of mantle peridotite tectonically exposed to the atmosphere leads commonly 15 to natural carbonation processes. Extensive cryptocrystalline magnesite veins and stock-work are 16 widespread in the serpentinite sole of the New Caledonia ophiolite. Silica is systematically 17 associated with magnesite. It is commonly admitted that Mg and Si are released during the 18 laterization of overlying peridotites. Thus the occurrence of these veins is generally attributed to a 19 per descensum mechanism that involves the infiltration of meteoric waters enriched in dissolved 20 atmospheric CO₂. In this study we investigate serpentinite carbonation processes, and related 21 silicification, based on a detailed petrographic and crystal-chemical study of serpentinites. The 22 relationships between serpentine and alteration products are described using an original method for 23 the analysis of μ -XRF images performed at the centimeter scale.

Our investigations highlight a carbonation mechanism, together with precipitation of amorphous silica and sepiolite, based on a dissolution-precipitation process. In contrast with the *per descensum* Mg/Si-enrichment model that is mainly concentrated in rock fractures, dissolutionprecipitation process is much more pervasive. Thus, although the texture of rocks remains relatively preserved, this process extends more widely into the rock, and may represent a major part of total carbonation of the ophiolite.

- 30
- Keywords: Serpentine, magnesite, carbonation, silicification, New Caledonia ophiolite
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INTRODUCTION

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Carbon dioxide is currently one of the primary greenhouse gases having an impact on global 3 warming. Therefore, numerous recent studies have focused on the potential of sequestration of CO₂ 4 by mineral carbonation, either through ex situ (e.g. Bobicki et al., 2011; Power et al., 2011; 5 Renforth et al., 2011; Badlucan and Dlugogorski, 2013; Harrison et al., 2013) or in situ processes 6 (e.g. Cipolli et al., 2004; Hansen et al., 2005; Teir et al., 2007; 2009; Andreani et al., 2009; Rudge 7 et al., 2010; Kelemen et al., 2011; Klein and Garrido, 2011). Among the various mineral species 8 that may undergo carbonation reaction, Mg-bearing minerals (as well as Ca-carbonates) are of a 9 great interest as they are very common at Earth surface and thus represent an important reservoir for 10 CO₂. In addition, magnesite (MgCO₃) has a long term stability, contrarily to alkali carbonates, 11 which are readily soluble in water (Lackner et al., 1995). Basically, magnesite is formed by the 12 reaction between a Mg-rich source and CO₂-rich fluids (Bain, 1924) and can integrate variable 13 amount of cations (mainly divalent, e.g. Ca and Fe) by substituting Mg in the crystal structure. 14 Based on the study of numerous magnesite deposits, Abu-Jaber and Kimberley (1992b) 15 distinguished vein-type and massive-type magnesite, the second type forming deeper than the 16 former. Numerous other parameters play a role in magnesite formation, such as temperature, the 17 origin of the components or the mechanisms of precipitation. The temperature of magnesite 18 formation extends from ambient to ~400°C (Halls and Zhao, 1995; Wilson et al., 2009; Klein and 19 Garrido, 2011). The origin of CO_2 is variable as it may be related to either weathering (atmospheric 20 CO₂), metamorphic (deep seated CO₂) or magmatic (magmatic CO₂) sources (Abu-Jaber and 21 Kimberley, 1992b). The source of magnesium is usually local at the outcrop scale but can also be 22 distant, e.g. coming from the weathering of magnesian rocks at the Earth surface and transported 23 downward by meteoric water infiltration (Podwojewski, 1995; Jurković et al., 2012). Mechanisms 24 of magnesite precipitation are also variable. Abu-Jaber and Kimberley (1992b) reported two main 25 ways of magnesite precipitation: i) precipitation through a reaction that involves CO₂-rich fluid and 26 Mg-rich rock, or alternatively Mg-rich fluids and CO₂-rich rock (*i.e.* the most common way to form 27 magnesite); ii) the oversaturation of the fluid with respect to magnesite may be enhanced by the 28 fluid evaporation and/or degassing (Dabitzias, 1980; Fallick et al., 1991; Zedef et al., 2000; 29 Ghoneim *et al.*, 2003). In both cases, hydrated Mg-carbonate species may precipitate alternatively 30 or in association to magnesite (e.g. Zedef et al., 2000; Beinlich and Austrheim, 2012) Detailed 31 studies of the precipitation mechanisms can thus provide major clues for the understanding of 32 natural CO₂ sequestration. 33

Among the various rocks that have the potential to react with CO₂, ultramafic rocks, and 1 particularly serpentinites, are probably the most efficient feedstock material for long-time storage 2 through the formation of magnesite (Dabitzias, 1980; Jedrysek and Halas, 1990; Pohl, 1990; Fallick 3 et al., 1991; Abu-Jaber and Kimberley, 1992b; Sherlock and Logan, 1995; Goff and Lackner, 1998; 4 Gerdemann et al., 2003; Ghoneim et al., 2003; Cipolli et al., 2004; Schulze et al., 2004; Hansen et 5 al., 2005; Teir et al., 2007, 2009; Kelemen and Matter, 2008; Rudge et al., 2010; Klein and Garrido, 6 2011; Jurković et al., 2012). Serpentine carbonation onsets by the dissolution of atmospheric CO₂ 7 into water, where CO_2 forms different species as a function of pH. At pH < 6.5, carbonic acid 8 9 (H₂CO₃) dominates, at pH between 6.5 and 10.5, bicarbonate (HCO₃⁻) dominates and at higher pH, 10 carbonate anion (CO₃²⁻) dominates. Serpentine dissolution and magnesite precipitation are also pH-11 dependent (Klein and Langmuir, 1987; Guthrie et al., 2001; Tneir et al., 2007; Prigiobbe et al., 12 2009; Teir et al., 2009; Krevor and Lackner, 2011). Experimentally, Teir et al. (2007) have shown 13 that the best efficiency for serpentine to carbonate conversion is obtained in the pH range of 8-11, 14 with an optimum at pH 9. Roughly similar pH conditions were measured for the optimum 15 carbonation of olivine (Prigiobbe *et al.*, 2009). At pH > 8, bicarbonate starts to dissociate into H⁺ 16 and CO_3^{2-} ions: 17

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Interaction between CO₂-rich water and serpentine gives rise to the exchange of H⁺ and Mg²⁺ cations on the mineral surface. This reaction produces silica and water, while free Mg²⁺ cations react with CO_3^{2-} anions to form magnesite:

 $Mg^{2+} + CO_3^{2-} \rightarrow MgCO_3$ (3)

 $\mathrm{HCO}_{3}^{-} \rightarrow \mathrm{H}^{+} + \mathrm{CO}_{3}^{2-}(1)$

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- $Mg_3Si_2O_5(OH)_4 + 6H^+ \rightarrow 3Mg^{2+} + 2H_4SiO_4 + H_2O(2)$
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²⁹ The overall reaction can be summarized as follow:

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- 31 $Mg_{3}Si_{2}O_{5}(OH)_{4} + 3(2H^{+} + CO_{3}^{2-}) \rightarrow 3MgCO_{3} + 2SiO_{2} + 5H_{2}O \quad (4)$ 32 Some entries + CO = ... Macmanita + Silica = + Water
 - Serpentine + $CO_{2(aq)} \rightarrow Magnesite + Silica_{(aq)} + Water$
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Worldwide, carbonation of serpentinite commonly leads to the formation of magnesite 2 deposits in association with ophiolitic bodies, e.g. in California (Bodenlos, 1950; Sherlock and 3 Logan, 1995), Egypt (Ghoneim et al., 2003), Greece (Dabitzias, 1980), Italy (Cipolli et al., 2004), 4 Norway (Beinlich et al., 2012) or Oman (Kelemen and Matter, 2008). In New Caledonia, similar 5 magnesite deposits have been described in the serpentine sole of the ophiolite, but few studies have 6 focused on the origin of this magnesite (Glasser, 1904; Trescases, 1973; Ulrich, 2010; Quesnel et 7 al., 2013). Quesnel et al. (2013) propose that magnesite veins in the serpentinite sole formed 8 tectonically, at temperatures between 40 and 80°C. On the basis of stable isotope analyses (O and 9 C), these authors propose that the fluid from which magnesite formed was originally meteoric water 10 and that the carbon source is mostly atmospheric, with a possible biogenic contribution. They also 11 suggest that magnesium originated from the dissolution of the peridotite at the top of the ophiolite 12 during laterization and was transferred down to the sole by the infiltration of meteoric waters under 13 tectonically active conditions. The per descensum model is largely invoked to explain magnesite 14 deposits in lateritic environments (Dabitzias, 1980; Pefrov et al., 1980; Pohl, 1990; Abu-Jaber and 15 Kimberley, 1992a; 1992b; Foster et al., 2002; Jurković et al., 2012; Oskierski et al., 2012). It 16 requires a well-drained system where meteoric waters, charged with atmospheric CO₂, dissolve 17 serpentine (and other Mg-silicates), a process that releases magnesium and silicon into solution. 18 Meteoric waters then percolate downward thanks to a microfracturing permeability system, thus 19 dissolving more magnesium. Within this frame, precipitation of magnesite is due to supersaturation 20 of fluids, which are generated by the neutralization of carbonic acid by serpentine dissolution, as 21 shown by reactions (2) and (3) (Bodenlos, 1950; Pohl, 1990; Fallick et al., 1991; Abu-Jaber and 22 Kimberley, 1992b; Giammar et al., 2005; Kelemen et al., 2011). Alternatively, the ultramafic pile 23 may become a partially closed system, and therefore limit the exchange between atmospheric CO₂ 24 and meteoric water (Jurković et al., 2012). Progressive dissolution of serpentine during the fluid 25 migration downwards consumes H⁺ ions, as shown by reaction (3). As CO₂ is not freely available 26 anymore, such consumption of H⁺ ions causes the increase of pH, enhancing magnesite 27 precipitation (Jurković et al., 2012). 28

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29 Considering the origin of magnesium as deriving from the laterization rises the question about 30 the mechanism of magnesite precipitation in New Caledonia. In such a case, and similarly to 31 magnesite deposits from Euboea (Greece, Boydell, 1921), magnesite formation may be the result of 32 direct precipitation from the fluids without interacting with hosted rocks of the deposit site. This 33 hypothesis was favored by the composition of New Caledonia waters passing through peridotites

and serpentinites (sampled in la Coulée river, Mont-Dore), showing that they are enriched in 1 magnesium and CO_3^{2-} ions (Barnes *et al.*, 1978). Alternatively, field investigations highlight a close 2 3 relationship between serpentinite, magnesite and silica, suggesting that the two latter phases may 4 originate from the dissolution of the former. In this case, magnesite would precipitate by in situ 5 replacement of the serpentine, using magnesium released during the serpentine dissolution. In this 6 study we provide evidences of magnesite formation through such a process on the basis of mineral 7 characterization using an original analytical method for the interpretation of micro-X-Ray 8 Fluorescence (µ-XRF) images. We particularly focus on the formation process of magnesite veins 9 coupled to intense silicification observed in the serpentinite sole of the New Caledonia ophiolite.

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New Caledonia is located in the SW Pacific, 2000 km east of the Australian coasts (Figure 1). It is composed of several islands that belong to the Norfolk ridge (La Grande Terre, Island of Pines, Belep Islands) and to the Loyalty ridge (Loyalty Islands). The main island, la Grande Terre, consists of a patchwork of terranes reflecting the geodynamic evolution of the SW Pacific region from late

1. GEOLOGICAL SETTINGS AND SAMPLE DESCRIPTIONS

17 Permien to Eocene (e.g. Cluzel and Meffre, 2002; Cluzel et al., 2001; 2012). Among these terranes, 18 the ophiolite is the most prominent as it covers more than 25% of the island. The so-called 19 Peridotite Nappe is composed of a large and continuous massif located south of the island and some 20 isolated klippes widespread along the west coast (Figure 1). The ophiolite was formed between 83 21 Ma with the opening of the South Loyalty Basin and its subsequent closing at 34 Ma, timing of its 22 obduction on the Norfolk continental basement (Cluzel et al., 2001; Crawford et al., 2003; Schellart 23 et al., 2006; Whattam et al., 2008; Whattam, 2009; Ulrich et al., 2010; Cluzel et al., 2012). Since its 24 emergence, the uppermost part of the Peridotite Nappe has undergone an intense laterization. This 25 led to the development of a thick laterite bed (up to 60 m thick; Sévin et al., 2012) that owns ~30% 26 of the world nickel resources. The whole ophiolite is formed of upper mantle rocks (mainly 27 harzburgites) with minor cumulates (Prinzofher, 1981). Peridotites are highly serpentinized, 28 particularly at the base of the ophiolite which is made of a thick (up to 400 m thick, Audet, 2010; 29 Ulrich et al., 2010) and silicified serpentinite sole where large amount of magnesite veins have 30 crystallized (Quesnel et al., 2013).

Samples presented in this study were collected in the serpentinite sole of the Koniambo
 massif (Figure 1). Similar outcrops occur on the serpentinite sole from other peridotite massifs in

New Caledonia (Ulrich, 2010). In the field, the whole sole is highly deformed, finely schistose and/ 1 or intensely brecciated, and has recorded multiple serpentinization events (Ulrich, 2010): Massive 2 serpentinization, which is of a typical bottle-green color in the field (Figure 2), is cross-cutted by 3 light-green colored serpentine veins. The latter is associated with black magnetite impregnations 4 and micro-cracks filled by fibrous serpentine (chrysotile). Magnesite occurs as millimeter to multi-5 decimeter stock-work veins with a typical cauliflower-like texture, cross-cutting serpentinites 6 (Figure 2). The magnesite is mainly snow-white colored, but can also appear greenish depending of 7 the amount of intergrown serpentine. On the basis of structural observations, Quesnel et al. (2013) 8 distinguished two types of veins. The first type of veins is observed along and/or within the margins 9 of top to the SW-centimeter shallow dipping shear zones. The second type of veins corresponds to 10 steeper veins occasionally crosscut by the low-dipping shear zones. Magnesite can also develop 11 pervasively by precipitating in large and massive serpentine blocks (Figure 2 a, b, c). In this case, 12 the magnesite seems to develop first at the expense of the light-green colored serpentine to 13 progressively extend to bottle-green serpentine, a relationship that is not clear when magnesite fills 14 the main shear zones, as described by Quesnel et al. (2013). 15

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- 17 2. ANALYTICAL METHODS
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19 1. X-Ray Diffraction

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Analyzes were performed at the Institut des Sciences de la Terre (ISTerre, Grenoble, France) on sample powders obtained after the crushing of separated magnesite veins and serpentinite host rock mineral fractions. X-ray diffraction (XRD) patterns were recorded with a Bruker D5000 powder diffractometer equipped with a Kevex Si(Li) solid state detector using CuK α_{1+2} radiation. Intensities were recorded at $0.02^{\circ} 2\theta$ step intervals from 5 to 80° , with a 6 seconds counting time per step. Size of the divergence slit was 0.298° .

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- 2. Raman Spectroscopy
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Raman spectroscopy measurements were performed at the Ecole Normale Superieure of Lyon and at GéoRessources Nancy, France, in both cases using a Horiba Jobin-Yvon LabRam HR800 spectrometer and a visible ionized argon laser source with a wavelength of 514 nm. Output laser power was 100 mW, and measurements were performed using an Olympus lens of x100 to focus the 1 laser beam onto an area that was 1 μ m in diameter. Analyzes were done on macroscopic samples 2 and on thin sections. Spectra result from the average of 5 acquisitions of 10 s to 20 s to optimize the 3 signal/noise ratio. Two regions of the Raman spectra were investigated: 150-1250 cm⁻¹ for structural 4 bonding characterization and 2800-3900 cm⁻¹ for the characterization of the hydroxyl groups.

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3. Micro-X-ray Fluorescence

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Micro-X-ray Fluorescence (µ-XRF) analyses were performed on a 5 mm-thick rock section 8 (Figure 2d) using EDAX Eagle III spectrometer at ISTerre (Grenoble, France). The X-ray tube 9 consists of a Rh anode operating at 250 µA with an acceleration voltage of 40 kV. Polycapillary 10 lenses were used to focus the X-ray beam down to 40 µm full-width-at-half-maximum at the sample 11 surface. An energy dispersive X-ray detector with resolution of 140 eV was used to measure 12 fluorescence spectra. Chemical maps were recorded with a matrix of 256 x 200 pixels, a 40 µm step 13 interval in both directions, and a dwell time of 1 s per pixel. For each map, the grey-scale 14 corresponds to the intensity of the Ka-lines of the different elements (Si, Mg, Fe, Al, Ni, Ti, Mn, Cr, 15 Ca, K) calculated from the integration of a specified region of interest (ROI) of the energy-range of 16 XRF spectra. Then, ROI maps (see electronic supplements, Figure S4) are used to calculate phase 17 maps thanks to a new routine, specially developed Matlab[©]-based code, following the same 18 approach than that successfully applied first to the computing of mineral-phase maps from 19 hyperspectral µ-XANES mapping (Muñoz et al., 2008). In the case of hyperspectral µ-XRF maps, 20 the phase map calculation consists first of determining pure mineral phases that are expected to be 21 present in the sample, in order to create standard spectra (or "pure" spectra). Then for each pixel of 22 the map, a linear combination of the different standard spectra is performed in order to fit each 23 single spectrum. Results provide quantitative phase maps showing the distribution of minerals 24 previously identified in the sample (*e.g.*, based on XRD and/or Raman analyses). This approach is 25 particularly useful to highlight relationships between minerals, especially for the characterization of 26 finely-divided mineral assemblages, *i.e.* when the beam is larger than grain size (such as here, 27 typically below 1 micron). Concentration maps (in wt.%) are finally recalculated on the basis of 28 phase distribution maps considering the chemical composition of standards (Table 1) 29

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4. RESULTS

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1. Mineralogy and chemical compositions

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a. Bulk mineralogical compositions

Figure 3 shows the X-ray diffraction patterns obtained on separated fractions of serpentinite and magnesite. The serpentinite mainly consists of serpentine with minor amount of magnetite and chromite. Serpentine occurs as lizardite and chrysotile. The occurrence of chrysotile is consistent with the presence of fibrous serpentine in microcracks as described above. Despite a careful separation, small peaks of magnesite and silica are present in the diffraction pattern of the serpentinite, suggesting that both minerals also occur at the micrometer scale within the hosted rock.

XRD pattern of magnesite powder shows that the carbonate exhibits its most characteristic 10 reflections at the following d values (in Å, arranged according to decreasing intensities): 2.74, 2.10, 11 1.70, 2.50, and 1.94 (Figure 3). In addition, the diffraction pattern shows that magnesite 12 systematically integrates small amounts of sepiolite (Mg₄Si₆O₁₅(OH)₂•6(H₂O)), a mineral that is 13 14 frequently described in association with carbonate in ultramafic environments (e.g. Birsoy, 2002; Yaliçin and Bozkaya, 2004; Boschetti and Toscani, 2008). On the basis on this XRD pattern, 15 magnesite is the only carbonate to crystallize. Neither dolomite nor calcite are formed, contrarily to 16 numerous magnesite deposits previously described in the literature (Griffis, 1972; Dabitzias, 1980; 17 Jedrysek and Halas, 1990; Fallick et al., 1991; Abu-Jaber and Kimberley, 1992b; Lugli et al., 2000; 18 Zedef et al., 2000; Ghoneim et al., 2003; Jurković et al., 2012). 19

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b. Optical microscopy

Figure 4 shows the typical mineralogical textures of carbonated serpentinites composing the 22 sole of the New Caledonia ophiolite. In thin section, the serpentinite does not exhibit any relic of 23 primary minerals (*i.e.* olivine and pyroxene). However, the habits of grains and original textures of 24 primary minerals (e.g. cleavage planes of orthopyroxene) have been preserved (typical 25 pseudomorphic "mesh" texture, Figure 4a, b) and indicate that the parent rock was a harzburgite. As 26 highlighted by XRD analyzes, opaque minerals associated with the serpentine are small grains of 27 magnetite and chromite disseminated in the mesh. In these rocks, magnesite occurs as finely 28 disseminate grains (*i.e.* cryptocrystalline texture) that developed both at the rim and in cracks 29 affecting serpentine grains (Figure 4b, d). These grains progressively aggregate to form larger zones 30 with a typical granular texture (Figure 4a-c). The sepiolite, identified by XRD and optical 31 microscopy, occurs as brown fibers interstitially to the magnesite nodules (Figure 4c). Silica also 32 occurs close to magnesite. Microscopic observations show that the nature of the silica is variable: it 33

occurs as an amorphous solid (gel-like) surrounding magnesite aggregates and serpentine grains in 1 area where magnesite dominates (Figure 4a, c, d). Where serpentine is dominant, silica consists of 2 crystalline to fine grains, forming vugs with a typical colloform texture propagating in the 3 serpentine mesh (Figure 4e, f). In this case, the nature of silica ranges from amorphous-like near the 4 rim of the vug to quartz-like at the center (Figure 4c, e, f). 5

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c. Raman spectroscopy

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Raman analyses were performed to further identify minerals that compose the serpentinite. 8 This technique is complementary to XRD, since it is particularly efficient to distinguish among the 9 different varieties of serpentine (e.g. Lemaire, 2000; Auzende et al., 2004) as well as silica 10 polymorphs (e.g. Götze et al., 1998; Pop et al., 2004). Figure 5 shows Raman spectra collected on 11 carbonated serpentinites. Results show that lizardite is the dominant serpentine variety that occurs 12 as individual grains and in the mesh, and corresponds to the bottle-green colored serpentine 13 described in the macroscopic observations. Chrysotile (not shown in Figure 5) has also been 14 detected, as already highlighted by XRD. In addition, light green serpentine was identified as 15 polygonal serpentine. It shows quite similar patterns to those of the lizardite at low wavenumbers, 16 but strongly differs at high wavenumbers (from 3500 to 3800 cm⁻¹, corresponding to OH group), 17 where the polygonal serpentine is characterized by a large peak composed by two bands centered at 18 3689 and 3700 cm⁻¹ (e.g. Lemaire, 2000; Auzende et al., 2004). 19

The Raman spectrum of magnesite is characterized by four distinct bands located at 209, 327, 20 737 and 1094 cm⁻¹ (Figure 5), consistently with the work of Krishnamurti (1956). The lack of bands 21 at 3448 and 3648 cm⁻¹ (*i.e.*, in the OH region; typical of hydromagnesite) shows that magnesite is 22 anhydrous. 23

Silica polymorphs are clearly identified using Raman spectroscopy. Vug rims consist of opal-24 CT (Silica #1, Figure 4e) while brown coronas (Silica #2, Figure 4e) and white fine grains (Silica 25 #3, Figure 4e) inside of the vugs are identified as chalcedony (Figure 5). Although significantly 26 different in microscopic observations (Figure 4e, f), both chalcedonies display very similar Raman 27 spectra, except on the intensity of the band located at 501 cm⁻¹ which is significantly higher in the 28 white chalcedony (electronic supplements, Figure S1). Notice that in all silica polymorphs, the large 29 bands observed between 3100 and 3900 cm⁻¹ indicate the presence of molecular water. 30

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d. XRF mapping 32

The elemental distribution, expressed in weight percent, in a typical serpentinite texture 1 surrounded by magnesite is shown in Figure 6. On the basis of chemical measurements, it was not 2 possible to discriminate lizardite from polygonal serpentine, both having very similar compositions. 3 Therefore, we only refer to serpentine in the followings, focusing on the nature of processes that 4 preferentially affect the serpentine grains or the mesh. The distribution maps of MgO and SiO₂ 5 highlight serpentine grains (s) as well as mesh texture (m). The chemical composition of these 6 grains is consistent with the stoichiometry of serpentine minerals, with about 43 wt.% for both MgO 7 +Fe₂O₃ and SiO₂. However, SiO₂ concentration in the mesh is significantly higher, and can reach up 8 to 65 wt.%. To better understand such differences in chemical compositions, we calculated the 9 phase distribution maps according to the minerals that are expected to be present in this sample 10 (Figure 7 and electronic supplements, Figure S4). Phase distribution reveals that the central zone of 11 the mapped area mainly consists of serpentine minerals. The amount of serpentine is at least 50 % 12 in the mesh and close to 100 % in grains. Silica distribution is relatively homogeneous in the mesh, 13 with about 30 % silica, but shows peculiarities close to the magnesite, where the silica content 14 reaches about 100 %. In contrast, the amount of silica in serpentine grains is well below 5 %. 15 Finally, the calculation of phase distribution maps was obtained with about 20 % of magnesite in 16 the serpentinite texture (*i.e.* the central area). 17

Figure 8 shows density correlation diagrams between serpentine, magnesite and silica based 18 on the phase distribution maps for the two regions delimited by squares in Figure 7. These diagrams 19 statistically illustrate the description made above on the phase maps and reveal two distinct 20 processes. In the region 1, the diagrams show a clear anticorrelation between silica and serpentine, 21 corroborating that a great part of the pixels of this region corresponds to "silicified serpentine". At 22 the opposite, the relationships between magnesite and serpentine on one hand and those between 23 silica and magnesite on the other hand are less straightforward. To better understand these two 24 diagrams, the region 2 (dashed square in Figure 7) was delimited around a heterogeneous serpentine 25 grain that appears partially altered (Figure 9). This alteration occurs around fractures in the grain 26 and is expressed by a lighter color of the serpentine in microscopic view. Considering this grain, a 27 statistical analysis shows that silica and magnesite are correlated to each other, whereas they are 28 both anticorrelated to serpentine (Figure 8, region 2). This point is particularly interesting since the 29 phase maps clearly highlight here a process of replacement of serpentine by an assemblage of 30 magnesite and silica. Even if silica and serpentine are anticorrelated in both regions, carbonation 31 and silicification processes do not follow rigorously the same trend. On the basis of our specific 32 treatment of the XRF maps, we demonstrate here that the serpentinite is affected by two distinct 33

weathering processes: while the mesh texture is mainly affected by a silicification process, the
 serpentine grains are mainly affected by the crystallization of a magnesite + amorphous silica
 assemblage.

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5 5. DISCUSSION

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Serpentinite carbonation in New Caledonia has a supergene origin, following a per descensum 7 model (Glasser, 1904; Trescases, 1973; Ulrich, 2010; Quesnel et al., 2013). The atmospheric carbon 8 dioxide is first dissolved in meteoric water. The CO₂-enriched fluids then circulate through the 9 lateritic cover, dissolve the residual Mg-rich minerals before driving to the precipitation of 10 magnesite in fractures and to a porosity reduction in the serpentinite, within the serpentinite sole. 11 This carbonation process leads to the formation of clusters of magnesite, potentially very large, and 12 usually very localized to the serpentine rich fractures or volumes. Although this process is most 13 easily observed in the field, our results show that the serpentine dissolution leads to the local 14 crystallization of magnesite and silica. This diffuse process is potentially the main mechanism of 15 carbonation of the serpentinite sole. 16

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1. Carbonation after serpentine dissolution

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On the basis of our results of our mineralogical investigations, several features provide 20 evidence that magnesite precipitation occurs through serpentine dissolution. First, the development 21 of magnesite along the edges of adjacent serpentine grains is a characteristic of carbonation by *in* 22 situ replacement of the serpentine (Figures 4b, 4d and 9). When serpentine grains are preserved, 23 carbonation is limited to the rims, the core remaining unaffected. At the opposite, the dissolution 24 and replacement of the serpentine are more efficient in case of fractured grains (Figure 9). This 25 observation is consistent with experimental studies showing that intense grain fracturing is required 26 to ensure complete carbonation (e.g. Haug et al., 2011; Kelemen et al., 2011; Hövelmann et al., 27 2012; van Noort et al., 2013). Another way to illustrate the progressive serpentine carbonation is 28 given by Figure S2 (in electronic supplements). Here, the RGB map (for Red, Green, Blue, see the 29 figure caption for more details) shows that magnesite is ubiquitous, formed by the replacement of 30 the serpentine grains, and that only a few of the latter are preserved from the carbonation process. In 31 addition, considering the stoichiometry and the molar volumes of the mineral species involved in 32 reaction (4), volumes of magnesite and silica produced by the dissolution of 1 mole of serpentine 33

(108 cm³) are respectively 85 cm³ and 58 cm³ ($V_{Mgs}/V_{Silica} \sim 1.45$), leading to a volume increase of 1 about 30 %. Statistics calculated on the serpentine grain presented in Figure 9 (corresponding to the 2 region 2, Figure 7) show that the assemblage consists of about 62 vol.% of serpentine, 21 vol.% of 3 magnesite and 15 vol.% of silica (V_{Mgs}/V_{Silica} ~1.4; Figure 8). This assemblage compares well with 4 the result of dissolution of 38 vol.% of serpentine, which leads to the formation of ~22 vol.% of 5 magnesite and ~15 vol.% of silica. These estimates testify that results given in the phase maps are 6 in good agreement with the stoichiometry of reaction (4). Second, carbonation by serpentine 7 dissolution is also evidenced by the precipitation of an amorphous silica layer at the grain rims 8 (Figure 4d). This observation is consistent with the previous studies that systematically reported the 9 formation of a Si-rich layer after Mg-bearing mineral breakdown (e.g. Luce et al., 1972; Lin and 10 Clemency, 1981; Guthrie et al., 2001; Schulze et al., 2004; Giammar et al., 2005; Bearat et al., 11 2006; Sipilä et al., 2008; Andreani et al., 2009; Daval et al., 2011; King et al., 2011, Hövelmann et 12 al., 2011; 2012). Third, sepiolite, which is described in our samples in association with magnesite 13 (Figures 3 and 4c), also formed subsequently to the dissolution of serpentine (e.g. Jones and Galàn, 14 1988; Birsoy, 2002; Yaliçin and Bozkaya, 2004; Andreani et al., 2009). Similar occurrences of 15 sepiolite have been reported during natural (Yaliçin and Bozkaya, 2004) and experimental 16 (Andreani et al., 2009) carbonation of Mg-bearing minerals (serpentine and olivine). Actually, 17 sepiolite may form in different ways: 18

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$$2Mg_3Si_2O_5(OH)_4 + 2(2H^+ + CO_3^{2-}) + 2H_4SiO_4 \rightarrow 2MgCO_3 + Mg_4Si_6O_{15}(OH)_2.6H_2O + 3H_2O$$
(5)
21 Serpentine + CO_{2(aq)} + Silica_(aq) \rightarrow Magnesite + Sepiolite + Water

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23 $3Mg_3Si_2O_5(OH)_4 + 5(2H^+ + CO_3^{2-}) \rightarrow 5MgCO_3 + Mg_4Si_6O_{15}(OH)_2.6H_2O + 4H_2O$ (6) 24 Serpentine + $CO_{2(aq)} \rightarrow Magnesite + Sepiolite + Water$

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$$\begin{array}{ll} & Mg_{3}Si_{2}O_{5}(OH)_{4}+2.5(2H^{+}+CO_{3}^{2-}) \rightarrow 2.5MgCO_{3}+0.125Mg_{4}Si_{6}O_{15}(OH)_{2.6}H_{2}O+1.25H_{4}SiO_{4}+1.125H_{2}O \\ & \\ & Serpentine+CO_{2(aq)} \rightarrow Magnesite+Sepiolite+Silica_{(aq)}+Water \end{array}$$

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All these reactions contribute to form magnesite, but they differ as reactions (5) and (6) conserve all Mg and Si in solids and are characterized by a volume gain (~60% and ~30%, respectively) while reaction (7) is balanced on volume. In addition, Eq. (5) consumes silica while Eq. (7) produces it. Only on the basis of cm-scale maps, it is difficult to estimate which reaction occured: in our sample, sepiolite represents less than 1 vol.% (see electronic supplements, Figure S4). Among the reactions written above, Eq. (7) is the one that produces the smallest amount of sepiolite, regarding the amount of magnesite that stoichiometrically precipitates. In contrast, Birsoy (2002) demonstrated that sepiolite formation is much more favored in the presence of Si-rich solution. In that case, sepiolite in our sample may derive from Eq. (5).

The fact that some serpentine grains are only partially transformed into magnesite indicates 5 that the carbonation process was not completed. The reason why the reaction does not go to 6 completion may be due to (1) silica precipitation, (2) pH increase and/or (3) porosity decrease. 7 Numerous studies proposed that the formation of a Si-rich layer at the rims of serpentine grain 8 during the first steps of dissolution might inhibit further Mg diffusion, potentially retarding or even 9 stopping the process of carbonation (Gerdemann et al., 2003; Schulze et al., 2004; Alexander et al., 10 2007; Daval et al., 2011). Although Si-rich layer effectively reduces the accessibility of fluids to the 11 reactive surface of minerals, recent investigations demonstrate that some permeability is maintained 12 as carbonation remains active even after its development (Béarat et al., 2006; Andreani et al., 2009; 13 Hövelmann et al., 2012). Thus, it is not obvious that precipitation of Si-rich layer led the serpentine 14 carbonation to stop. Alternatively, following reaction (4), the dissolution of serpentine consumes 2 15 moles of H⁺ for 1 mole of CO₃²⁻, leading to a progressive increase of fluid pH. According to Barnes 16 17 et al. (1978), fluids reacting with serpentinite in New Caledonia have pH ranging from 9 to 11. At 18 pH 9, HCO_3^{-1} dominates over $CO_3^{2^{-1}}$, making the serpentine dissolution possible following reactions 19 (1) and (2). At pH 10.5, CO₃²⁻ species become dominant and at pH 10.8, HCO₃⁻ represent less than 20 20% of the carbonate species in the fluid. In these conditions, magnesite precipitation is favored, 21 but serpentine dissolution is scarce due to the lack of H⁺ ions (e.g. Teir et al., 2007). Thus, intensive 22 exchange between H⁺ ions and Mg²⁺ cations on the serpentine surface leads to a progressive 23 increase of pH that may inhibit further serpentine dissolution and subsequent carbonation. There is 24 no evidence to argue against such process in New Caledonia, but it requires that atmospheric CO₂ is 25 not freely available and thus a roughly closed system (Jurković et al., 2012). A third explanation 26 may account for the inhibition of carbonation. Hövelmann et al. (2012) recently investigated the 27 microstructure and porosity evolution as a function of carbonation reaction progress in natural 28 peridotite. On the basis of their experimental results, they reported that a carbonation extend of ~ 10 29 % leads to a closure of 50 % of the initial porosity. They demonstrated that magnesite precipitation 30 in fracture pore space reduces the permeability and progressively stops the fluid pathway, 31 preventing further reaction between the fluid and the silicate surface and ultimately ends the 32 carbonation process. Contrarily to serpentinization, which is able to propagate through a reaction-33

induced fracturing mechanism (e.g. Plümper et al., 2012), carbonation is self-limited as the reaction 1 will be inhibited due to magnesite growing that ultimately clogs the system. This implies that no 2 volume gain occurs at the rock-scale (Beinlich et al., 2012). Numerous studies highlight the 3 necessity of active fracturing to ensure complete carbonation (e.g., Kelemen and Matter, 2008; 4 Kelemen et al., 2011; Boschi et al., 2012). In our samples, no evidence of carbonation-induced 5 fracturing has been observed so far, even if Quesnel et al. (2013) show the syn-kinematic character 6 of magnesite veins at the outcrop scale. At the millimetric scale, volume gain associated with 7 reaction (4) may have inhibited the complete carbonation of serpentinite by clogging the reacting 8 zone. Such a process may prevent the infiltration of additional fluids, resulting in the partial 9 carbonation of serpentine grains as illustrated by Figure 7 (see also electronic supplements, Figure 10 S2). 11

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2. Redox conditions during magnesite precipitation

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The behavior of iron during carbonation has been poorly investigated. Such information 15 would be indicative of redox conditions during magnesite precipitation. In serpentinite, iron is 16 mainly hosted by iron oxides (magnetite, Fe₃O₄) surrounding serpentine grains or disseminated in 17 the mesh texture (see electronic supplement, Figure S5). Iron content in serpentine is ~ 2 wt.% 18 (expressed as FeO, Table 1). It substitutes Mg²⁺ cations in octahedral sites (O'Hanley, 1996). During 19 serpentine dissolution, iron is released from the mineral structure and may subsequently form Fe-Si-20 rich layer that progressively evolves into magnetite at the interface between serpentine and aqueous 21 fluid (Fallick et al., 1991; Alexander et al., 2007; Andreani et al., 2009, Saldi et al., 2013). 22 Alternatively, it may precipitate as siderite (FeCO₃) under rather high pCO_2 and specific pH range 23 from 5.5 to 7.5 (Ohmoto et al., 2004) or be integrated in magnesite by substituting Mg²⁺ cations 24 (Abu-Jaber and Kimberley, 1992b; Hansen et al., 2005). In each of these minerals, iron oxidation is 25 expected to differ according to oxidizing conditions: it is mostly oxidized in magnetite (i.e. 2/3 of 26 iron is ferric iron) while it is in ferrous state in siderite or magnesite. In our sample, XRF analyzes 27 show that the iron content in magnesite is very low (Table 1), suggesting that its integration into the 28 structure of the magnesite was very scarce. In addition, we never identified siderite in our samples 29 so far. As stated above, siderite precipitates at pH conditions that are significantly lower than those 30 of New Caledonia waters, so it is likely that pH was too high to make siderite precipitate. Magnetite 31 is ubiquitous in our sample. Therefore, discriminating magnetite grains related to prior 32 serpentinization events to those potentially derived from carbonation is not obvious. However, 33

microscopic observations show that numerous magnetite grains have precipitated inside of partially 1 carbonated serpentine grains (Figures 4d and 9) while magnetite related to serpentinization 2 generally forms outside of serpentine grains. Schematic representation in Figure 9 highlights the 3 systematic association of magnetite with cracks in the serpentine grain. According to Andreani et al. 4 (2009), such grain fracturing may be considered as zones of localized fluid flow that favor the 5 precipitation of magnetite during the first step of mineral dissolution. In contrast, their experiments 6 show that carbonation initiates in domains of reduced fluid flow zones (fractures of a smaller size) 7 where chemical gradient are small and thus facilitate local supersaturation, high pH and more 8 reducing conditions. These results indicate that magnetite associated with the carbonation process 9 may be used as a proxy to estimate the fluid flow rate at the serpentine grain scale: Fractures filled 10 by magnetite correspond to high fluid flow zones characterized by more oxidizing conditions. In 11 these regions, dissolution occurs but not magnesite precipitation (Andréani et al., 2009). 12 Magnesium migrates to zones of reduced fluid flow, corresponding to regions dominated by 13 magnesite or partially carbonated serpentine (Figures 9. 10). In these areas magnesite precipitates 14 due to local supersaturation under more reducing conditions. This underlies that redox gradients 15 occur even at grain scale. 16

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18 3. Serpentine mesh silicification

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a. Silica mobilization and precipitation

Intense silicification of the serpentine mesh was revealed during this study based on XRF 21 measurements and Raman spectroscopy (Figures 6-8, electronic supplements, Figures S2 and S3). 22 For instance, the region 1 (Figure 7) is composed of 62 vol.% of serpentine, 24 vol.% of silica and 7 23 vol.% of magnesite in average. The precipitation of 7 vol.% of magnesite consumes 12 vol.% of 24 serpentine and forms 5 vol.% of silica, on the basis of reaction (4). Such excess of silica necessarily 25 involves a contribution of silica from outside of the region 1. Phase maps show that pure silica 26 mainly occurs in gaps between magnesite aggregates (left-side of the map, Figure 7 and electronic 27 supplements, Figure S2). However, when aggregates agglomerate subsequently to magnesite 28 growth (right-side of the map, Figure 7 and electronic supplements, Figure S2), silica is expelled 29 from the magnesite. Numerous experimental studies indicate the progressive migration of silica 30 rather than being rejected from the growing magnesite (Schulze et al., 2004; Hövelmann et al., 31 2011; 2012). As stated above, the first steps of serpentine dissolution release silica that immediately 32 precipitates as a Si-rich layer. As carbonation reaction proceeds, the released silicon may feed the 33

growth of the Si-rich layer, but Hövelmann et al. (2012) reported that such inward growth is 1 limited. This limitation is mainly associated with pH increase of fluid during the carbonation 2 process. At pH>9, magnesite precipitates while silica is solubilized and is therefore able to migrate 3 away from the solid-fluid interface. Occurrence of water within silica is demonstrated by the Raman 4 spectrum measured in the region of high frequencies (Figure 5), which exhibits a typical spectrum 5 of molecular water. As illustrated by XRF mapping, fluids enriched in silica subsequently to 6 serpentine dissolution, propagates in the serpentine meshwork. Similar observations have been 7 reported in serpentinites from the Oman and the Ligurian ophiolites (Stanger, 1985; Boschi et al., 8 2009; Lacinska and Styles, 2012). According to Lacinska and Styles (2012), the well-preserved 9 mesh texture (as observed in our samples) induces a combination of iso-volumetric processes of 10 slow rate dissolution of the mesh serpentine and immediate local precipitation of silica. They also 11 argue that precipitation of silica is favored at near-neutral pH conditions. Such conditions differ 12 from those we assume in our system (high alkali pH). However, Williams and Crerar (1985) 13 attributed the precipitation of amorphous silica phases in nature due to the formation of dense 14 colloids in supersaturate alkaline aqueous solutions. In addition, progressive dissolution of silica 15 causes pH of the solution to drop (Williams and Crerar, 1985; Williams et al., 1985). Following 16 these observations, we infer that such process may favor the dissolution of the serpentine mesh and 17 the subsequent grain-by-grain replacement by amorphous silica phases, as postulated by Lacinska 18 and Styles (2012). Such a process involves the removal of substantial amount of magnesium. As 19 shown above, serpentine grain carbonation occurs stoichiometrically, *i.e.* magnesium released after 20 serpentine grain dissolution, then immediately precipitates as magnesite. Thus, it is likely that the 21 magnesium released during serpentine mesh silicification migrates out of the reaction zone. This 22 assumption is consistent with the study of Boschi et al. (2009) that reported similar magnesium 23 mobilization after serpentine dissolution and silica precipitation. 24

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b. Silica evolution

Raman spectroscopy shows that silica formed in response to serpentine carbonation crystallizes as opal-CT and chalcedony (Figure 5). Such silica polymorphs are commonly described in association with magnesite and are consistent with formation at low temperature conditions (Boydell, 1921; Bodenlos, 1950; Dabitzias, 1980; Pohl, 1990; Abu-Jaber and Kimberley, 1992b; Klein and Garrido, 2011). According to Lacinska and Styles (2012), the formation of opal-CT indicates precipitation from supersaturated fluids. Williams *et al.* (1985) and Williams and Crerar (1985) show that the precipitation of silica polymorphs is driven by multiple steps of dissolution-

precipitation. Systematically, studies made on the silica diagenesis report that saturated silica 1 solutions do not form opal-CT directly, but follow a sequential crystallization with first the 2 precipitation of amorphous opal-A (Graetsch et al., 1985; Williams et al., 1985; Williams and 3 Crerar, 1985; Heaney, 1993; Lacinska and Styles, 2012). Opal-A then evolves by means of 4 dissolution-precipitation with concurrent ordering of the structure and removal of water, forming a 5 pathway as follow: opal-A (amorphous) \rightarrow opal-CT (cristobalite-tridymite assemblage) \rightarrow 6 chalcedony \rightarrow quartz. According to Williams *et al.* (1985), the relationship between solubility and 7 surface area or particle size is sufficient to explain such evolution. In our samples, only opal-CT and 8 chalcedony have been identified so far. One possible explanation to account for the absence of opal-9 A is the complete replacement by opal-CT. The association with magnesite supports this hypothesis, 10 since carbonates are thought to enhance the formation of opal-CT (Williams and Crerar, 1985). 11 Quartz has never been described associated with serpentinite in New Caledonia, so it is likely that 12 the transition chalcedony to quartz does not occur. This reaction is very slow and is more likely to 13 occur in a closed system, from the precipitation of fluids undersaturated in silica with respect to 14 opal-A, opal-CT or chalcedony (Lund, 1960 and references therein; Williams et al., 1985; Williams 15 and Crerar, 1985). In our samples, vugs containing chalcedony are systematically characterized by 16 the presence of a hole on its center (Figure 4). Similar observations were reported by Lund (1960) 17 from silicified corals where both chalcedony and quartz precipitated. This author concluded that the 18 hole served as a conduit for the continuous circulation of dissolved silica, resulting in the 19 precipitation of chalcedony. At the opposite, when no hole was observed, chalcedony was 20 completely replaced by quartz. 21

As an example, the concurrent reordering that occurs during the diagenetic pathway of silica 22 is illustrated by the behavior of chalcedony (Figures 4 and electronic supplement, Figure S1). 23 Chalcedony, as many of the microcrystalline SiO₂ varieties, consists of an intimately intergrowth of 24 α -quartz and moganite. Moganite is a silica polymorph that typically contains up to 3 wt.% of water 25 which is not a constituent of the structure. Using Raman spectroscopy, Pop et al. (2004) showed 26 that during the opal-CT to chalced ony transition, moganite starts growing after α -quartz and 27 preferentially in the most crystallized areas. It can be used as a proxy to evaluate the ordering of the 28 chalcedony during its transition from opal CT to quartz (Götze et al., 1998; Rodgers and Cressey, 29 2001; Pop *et al.*, 2004). In Figure 4c, both silica #2 and silica #3 have been identified as chalcedony 30 but differ in microscopic observation by their colors. In Raman spectroscopy, silica #2 and silica #3 31 almost display the same patterns, except the intensity of the band at 501 cm⁻¹ (moganite) that is 32 higher in the white chalcedony (electronic supplement, Figure S1). The moganite content (in wt.%) 33

of chalcedony can be calculated using the Raman band integral ratios I₍₅₀₁₎/I₍₄₆₅₎ (*i.e.* moganite/ quartz) and applying the calibration curve proposed by Götze *et al.* (1998). We find that both chalcedonies are dominated by moganite, which represents 77 wt.% in brown chalcedony (silica #2, Figure 4c) and 81 wt.% in white chalcedony. The enrichment of moganite traduces the progressive ordering of chalcedony which ultimately transforms in quartz given sufficient time (Williams and Crerar, 1985; Heaney and Post, 1992; Rodgers and Cressey, 2001; Lynne *et al.*, 2007).

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4. Coupled carbonation-silicification in dissolution-precipitation processes: a summary

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10 Based on our results and those from previous experimental studies, we infer that serpentine 11 carbonation occurred due to the circulation of high pH meteoric waters dissolving the serpentine. 12 Serpentine dissolution started at grain boundaries and in large grain fractures that correspond to 13 regions where the fluid flow is the highest (Figure 10a). In these regions, intensive exchanges 14 between H⁺ and Mg²⁺ ions led to the development of a Si-rich layer and the precipitation of 15 magnetite at the mineral/water interface (Figure 10b). In contrast, magnesite precipitation was not 16 favored in these zones of high fluid flow, which are characterized by strong chemical gradients and local oxidizing conditions (Andreani et al., 2009). Released magnesium therefore migrated to 17 18 regions of reduced fluid flow, where magnesite nucleated at the expense of serpentine surfaces by a 19 process of dissolution-precipitation due to local supersaturation and more reducing conditions 20 (Figure 10b-c). Here, the growth of magnesite was mainly fed by magnesium coming from the 21 dissolution of adjacent serpentine grain. Potentially, distal contributions may have occurred (i.e. 22 coming from the laterization, Barnes et al., 1978), since magnesite incorporate calcium while this 23 element is not concentrated in serpentine (Table 1). Silica released during the carbonation process 24 may first have precipitated *in situ* as amorphous silica (opal A/CT), but progressive pH increase 25 during the reaction facilitates the silicon solubilization and subsequent migration away from the fluid-solid interface (Figure 13c; Schulze et al., 2004; Andreani et al., 2009; Hövelmann et al., 26 27 2011; 2012). Silica was then able to propagate in the serpentine mesh, silicifying the latter by precipitating first as an amorphous gel that progressively orders its crystalline structure to 28 ultimately evolve as quartz if given sufficient time (Figure 10c; Williams and Crerar, 1985; Heaney 29 and Post, 1992; Rodgers and Cressey, 2001; Lynne et al., 2007). Magnesium leached during the 30 serpentine mesh carbonation migrated out from its original reacting zone and potentially fed larger 31 32 magnesite deposits in syn-tectonic fractures, as previously proposed by Boschi et al. (2009) in the 33 Ligurian ophiolite. Carbonation process ended when the overall serpentine was converted into an assemblage of magnesite + silica or, alternatively, when magnesite precipitation induced closure of
 the initial porosity (Figure 10d, Hövelmann *et al.*, 2012).

In agreement with recent studies from Andreani et al. (2009), Boschi et al. (2009), or 3 Hövelmann et al. (2011; 2012), our study demonstrates the importance of active fracturing in the 4 idea of in situ CO₂ sequestration. Although the volume extend expected during serpentine 5 carbonation may lead to the system clogging, serpentine mesh silicification involves substantial 6 removal of magnesium. This magnesium may migrate out of the reaction zone and subsequently 7 precipitates as massive magnesite veins along the main structural discontinuities such as those 8 described in New Caledonia by Quesnel et al. (2013). As previously underlined by Boschi et al. 9 (2009), such a process may be considered as an alternative, efficient way for CO_2 sequestration. 10

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1 Figure captions

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Figure 1: Localization of the main ophiolitic occurrences composing the Peridotite Nappe.
Simplified geological map of the Koniambo massif is modified from Maurizot *et al.* (2002).

5

Figure 2: a, b, c) Snow-white colored magnesite veins crystallized in massive serpentinite
blocs, Koniambo massif. Magnesite show a typical cauliflower texture and is closely associated
with silica veins (in brown). d) Typical sample of carbonated serpentinite. Orange dashed line
square localizes the mapped area by μ-XRF presented in Figures 6 to 8 and in Figures S2 and S3.

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Figure 3: Typical XRD patterns of serpentinite and magnesite. Liz: Lizardite; Chr: Chromite;
 Opl: Opal; Mgs: Magnesite; Mgt: Magnetite; Ctl: Chrysotile. Numbers upon the magnesite peaks
 correspond to *d* values, given in Å.

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Figure 4: a) Microphotography under polarized light illustrating the development of 15 magnesite and silica in serpentinite. b) Microphotography under crossed-polarized light showing 16 the development of magnesite along the rims of serpentine grains (indicated by the red arrows). 17 Notice that the serpentinization also affects the orthopyroxene c) Magnesite grain aggregates 18 surrounded by colloform amorphous silica gel. Associated brown and fibrous mineral corresponds 19 20 to sepiolite (observation under polarized light). d) Nucleation of magnesite grain on serpentine 21 surface surrounded by colloform amorphous silica gel. Black dots inside of the serpentine grain correspond to magnetite (observation under crossed-polarized light). e) Microtextures of a silica 22 vein under polarized light (mesh: serpentine mesh; #1: opal-CT, #2 and #3: Chalcedony; 23 identifications made by raman spectroscopy, see Figure 5). e) Same as e) under crossed-polarized 24 light. Mineral abbreviations: Mgs: magnesite; Mgt: Magnetite; Opx: Orthopyroxene; Sep: sepiolite; 25 Srp: serpentine 26

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Figure 5: Typical Raman spectra observed for serpentine (lizardite and polygonal), chalcedony, opal-CT and magnesite composing the serpentinite sole of the New Caledonia ophiolite.

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1 2 **Figure 6:** Quantitative chemical maps of MgO, SiO₂ and Fe₂O₃ (in wt.%) calculated on the basis of μ -XRF measurements (EDAX Eagle III). Complete procedure for the map calculation is detailed in the text.

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Figure 7: Spatial repartition and quantification of mineral phases (in %) calculated on each
 pixel of the mapped area (Figure 2d), based on μ-XRF measurements (EDAX Eagle III). Square delimited regions correspond to areas used to construct correlation diagrams in Figure 8.

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9 Figure 8: Correlation diagrams (represented as density fields) between serpentine, magnesite 10 and silica based on the phase distribution maps for the two square-delimited regions in Figure 8. *n* 11 corresponds to the total number of pixels composing each region. Colorbar values correspond to the 12 number of pixels in one grid cell of a size of 5% by 5%.

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Figure 9: Microphotography under polarized light and schematic representation of thepartially carbonated serpentine grain (corresponding to the Region 2 in Figure 7).

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Figure 10: Schematic sketch of coupled carbonation-silicification in dissolution-precipitation 17 processes. a) High pH meteoric waters (vellow arrows) percolating in serpentinite porosity start to 18 dissolve the serpentine. b) In zones of high fluid flow (see the text for more details), dissolution of 19 the serpentine surface releases Mg²⁺ cations (white arrows) that migrate to zones of reduced fluid 20 flow, leaving behind a Si-rich layer. Iron released during this dissolution step immediately 21 precipitates as magnetite due to local oxidizing conditions favored by the constant renewal of water 22 that characterizes zones of high fluid flow (Andreani et al., 2009). c) Magnesite precipitates at the 23 expense of serpentine surface in zones of reduced fluid flow due to local supersaturation and more 24 reducing conditions. Released silica may precipitate as amorphous silica interstitially to magnesite 25 aggregates, but increasing pH as the carbonation proceeds leads to its solubilization and subsequent 26 migration in the serpentine mesh (orange dashed arrows). d) Silica, in aqueous form, propagates in 27 the serpentine mesh and finally precipitates in amorphous opal A subsequently evolving to opal CT, 28 chalcedony and ultimately quartz. Complete serpentine replacement in a magnesite+silica 29 assemblage may occur. Alternatively, the growing of magnesite may lead to the closure of initial 30 porosity, preventing additional fluid circulations and thus ending the carbonation process before 31 going to completion (Hövelmann et al., 2012). 32

1 Table

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Table 1: *In situ* concentration measurements by μ-XRF for serpentine, magnesite, silica,
sepiolite, magnetite and chromite used for the calculation of quantitative maps shown in Figure 6
(see also electronic supplements, Figure S6).

6

Elements	Serpentine	Magnesite	Silica	Magnetite	Sepiolite	Chromite
MgO	40.25	46.84	-	-	24.72	12.34
Al ₂ O ₃	0.41	-	-	-	0.46	23.22
SiO ₂	43.60	0.39	100.00	-	62.66	0.08
K ₂ O	-	-	-	-	0.01	0.01
CaO	-	0.54	-	-	0.2	0.07
TiO ₂	0.01	-	-	-	0.02	-
Cr ₂ O ₃	0.02	-	-	-	0.02	48.20
MnO	0.03	-	-	-	0.67	0.20
FeO	2.51	0.02	-	100.00	0.55	15.82
NiO	0.17	-	-	-	0.11	0.06
CO ₂	-	52.21	-	-	-	-
H ₂ O	13.00	-	-	-	10.58	-
Total	100.00	100.00	100.00	100.00	100.00	100.00

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19 Supplementary figure captions

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Figure S1: Raman spectra of brown chalcedony (Silica #2, Figure 4) and white chalcedony (Silica #3, Figure 4). Both only differ by the intensity of the moganite band (501) which is significantly higher in the white chalcedony.

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Figure S2: RGB (for Red-Green-Blue) map calculated by the superposition of the three phase maps shown in Figure 7. One color is attributed to each phase (Red: Magnesite; Green: Serpentine; Blue: Silica), in order to highlight the relationship between mineral phases. Pixels characterized by a mix of green and blue show the progressive silicification of the serpentine mesh. Greenish to brownish pixels (mix of red and green) correspond to partially carbonated serpentine grains.

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Figure S3: Progressive silicification of the serpentine mesh illustrated by Raman
 spectroscopy.

2	Figure S4: µ-XRF maps of elemental concentrations based on ROI (region of interest)
3	measurement at element K-edge. These maps correspond to raw data used for the calculation of
4	mineral phase maps (in %) shown in Figure 6 and Figure S5, and quantitative maps shown in Figure
5	6 and Figure S6.
6	
7	Figure S5: Maps of mineral phases calculated calculated on the basis of μ -XRF
8	measurements (EDAX Eagle III). The «Total» map corresponds to the sum of phase maps and is
9	used to verify the consistency of the calculation (each pixel of the map have to be close to 100 %).
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11	Figure S6: Quantitative maps (in wt.%) calculated on the basis of μ -XRF measurements
12	(EDAX Eagle III). The «Total» map corresponds to the sum of quantitative maps. Map of $\mathrm{H_{2}O}$
13	+CO ₂ , elements that cannot be measured by μ -XRF, is calculated by subtracting the Total map to
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SiO₂ (wt.%)





























Intensity (a.u.)







Optical



Magnesite (%)



Magnetite (%)







Serpentine (%)





Sepiolite (%)









