Ultramafic rock carbonation: Constraints from listvenite core BT1B, Oman Drilling Project

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Key Points:

- Large-scale carbonation resulted in pervasive replacement of serpentinite by the magnesite-quartz-fuchsite (listvenite) assemblage.
- Clumped isotope thermometry indicates carbonation temperatures between 50°C and 250°C.
- Carbonation textures indicate a transition from non-equilibrium spheroidal growth to near-equilibrium euhedral growth.
Abstract

The occurrence of the quartz–carbonate alteration assemblage (listvenite) in ophiolites indicates that ultramafic rock represents an effective sink for dissolved CO$_2$. However, the majority of earlier studies of ultramafic rock carbonation had to rely on the surface exposure of reaction textures and field relationships. Here we present the first observations on ultramafic rock alteration obtained from the 300 m deep BT1B drill hole, ICDP Oman Drilling Project, allowing for a continuous and high-resolution investigation. Hole BT1B recovered continuous drill core intersecting surface alluvium, 200 meters of altered ultramafic rock comprising mainly listvenite and minor serpentine bands at 90 m and 180 m depth, and 100 meters of the underlying metamorphic sole. Textural evidence suggests that the carbonation of fully serpentinized harzburgite commenced by non-equilibrium growth of spheroidal carbonate characterized by sectorial zoning resulting from radially oriented low-angle boundaries. In the serpentinite, carbonate spheroids are composed of alternating magnesite cores and dolomite rims, whereas texturally similar carbonate in the listvenite is composed of Fe-rich magnesite cores and Ca-Fe-rich magnesite rims. The distinct compositions and mineral inclusions indicate that the carbonation extent was controlled by fluid accessibility resulting in the simultaneous formation of limited carbonate in the serpentinite bands and complete carbonation in the listvenite parts of BT1B. The presence of euhedral magnesite overgrowing spheroidal carbonate in the listvenite suggests near-equilibrium conditions during the final stage of carbonation. The carbonate clumped isotope thermometry constrains carbonate crystallization between 50°C and 250°C, implying repeated infiltration of reactive fluids during ophiolite uplift and cooling.

1 Introduction

Alteration of the oceanic lithosphere by hydrothermal fluids is the primary driver of the hydrosphere–lithosphere volatile exchange and has significant consequences for rock physical properties and planetary-scale element fluxes. Furthermore, natural carbon uptake in thermodynamically stable carbonate in the oceanic lithosphere represents a means for permanently removing CO$_2$ from the atmosphere (Kelemen & Matter, 2008). The carbon uptake potential is particularly strong in the ultramafic part of the oceanic lithosphere that has high concentrations of divalent metal ions, required for carbonate mineral formation (Kelemen et al., 2011). Peridotite alteration by seawater typically results in the formation of Ca carbonate (calcite and aragonite) (e.g. Bonatti et al., 1980; Grozeva et al., 2017), while the alteration by metamorphic fluids frequently results in Mg-rich carbonate (dolomite and magnesite), accompanied by secondary silicate phases including talc, quartz, and fuchsite (Cr-muscovite) and additional Fe-oxide, -hydroxide and/or sulfide phases. Altered ultramafic rocks consisting of the magnesite–quartz assemblage are termed listvenite and represent the end-product of CO$_2$-fluid driven mafic and ultramafic rock alteration as almost the entire inventory of Mg and Ca cations is incorporated into carbonate minerals (Halls & Zhao, 1995). Natural carbon uptake in ultramafic formations is significant as evidenced by abundantly exposed carbonate-bearing ophiolite complexes worldwide (Akbulut et al., 2006; Azer et al., 2019; Barnes & O’Neil, 1969; Barnes et al., 1978; Beinlich et al., 2010, 2012, 2018, 2020a; Belogub et al., 2017; Emam & Zoheir, 2013; Gahlan Hisham et al., 2018; Garcia del Real et al., 2016; Halls & Zhao, 1995; Hansen et al., 2005; Hinsken et al., 2017; Launay & Fontes, 1985; Menzel et al., 2018; Quesnel et al., 2013; Tominaga et al., 2017; Tsikouras et al., 2006). One of the largest and most well-explored ophiolite complexes is the Samail ophiolite in the Sultanate of Oman and the United Arab Emirates (Chavagnac et al., 2013a, b; Clark & Fontes, 1990; Falk et al., 2016; Falk & Kelemen, 2015; Kelemen & Matter, 2008; Lacinska & Styles, 2012; Lacinska et al., 2014; Mervine et al., 2014; Nasir et al., 2007; Neal & Stanger, 1983, 1985; Paukert et al., 2012; Stanger, 1985; Streit et al., 2012).
the Samail ophiolite, listvenite is present in the basal section above the underlying metamorphic sole. Earlier work has recognized the occurrence of calcite-, dolomite- and quartz-dominated listvenite containing additional fuchsite ± chlorite ± spinel in the form of discontinuous planar lenses and fault-controlled vein systems (Nasir et al., 2007; Stanger, 1985; Wilde et al., 2002). More recently investigated listvenite from Wadi Mansah is characterized by magnesite as the dominant carbonate mineral consistent with peridotite alteration predominately driven by influx of CO$_2$ (Falk & Kelemen, 2015). In addition to the carbonated ultramafic basal section, the Samail ophiolite comprises a complete sequence of oceanic lithosphere from the upper mantle through to the continental crust all of which have recently been subject to comprehensive drilling and subsequent analysis of rock core, logging, hydrological and microbial sampling in the course of the Oman Drilling Project (OmanDP, ICDP Exp. 5057) (Kelemen et al., 2020; Kelemen et al., 2014). To better constrain the conditions and reaction pathways driving complete peridotite carbonation at large scale, here we present the first results of microstructural investigation and carbonate clumped isotope thermometry on samples recovered from Hole BT1B of the OmanDP. BT1B recovered a 300 m long continuous core across the basal thrust of the Samail ophiolite through 200 m of listvenite and intercalated serpentinite into an additional 100 m of the underlying metamorphic sole. Reaction textures are assessed across scales ranging from hand specimen to the nanoscale constraining critical reaction parameters and chemical equilibration during natural, large-scale ultramafic rock carbonation.

2 Geological Setting

The Samail Ophiolite, located in the Sultanate of Oman and the United Arab Emirates, is the most extensive and best-exposed cross-section through the oceanic lithosphere (Fig. 1a). The ophiolite comprises an intact stratigraphy of oceanic crust composed of pelagic allochthonous sediments, overlying pillow basalts and the sheeted dike complex through to gabbro and upper mantle peridotite (Coleman, 1981; Searle & Malpas, 1980). The basal thrust beneath the Samail nappe separates the ophiolite sequence from the underlying metamorphic sole consisting of metasedimentary phyllite, greenschist facies...
metabasalt and minor amphibolite of the Haybi Complex (Searle & Malpas, 1980). The metamorphic sole is underlain by Permian to Late Cretaceous distal to deep-sea sediments, metavolcanic rocks, and shelf-facies limestone of the Hawasina Group (Coleman, 1981; Nasir et al., 2007; Searle et al., 2015; Stanger, 1985). Syn-to post-emplacement tectonic tilting of the stratigraphy has been followed by unconformable deposition of Late Cretaceous to Early Miocene transgressive shallow marine sediments before Late Miocene to Quaternary uplift and erosion (Coleman, 1981; Searle & Malpas, 1980; Searle et al., 2015).

U–Pb dating of magmatic zircon and ⁴⁰Ar/³⁹Ar ages of mica and hornblende from the metamorphic sole constrain the timing of thrusting of the Samail ophiolite onto the autochthonous Proterozoic to Mesozoic sedimentary Arabian continental margin between 95 Ma and 80 Ma (Cowan et al., 2014; Gnos, 1998; Hacker, 1994; Hacker & Gnos, 1997; Hacker & Mosenfelder, 1996; Hacker et al., 1996; Lanphere, 1981; Rioux et al., 2012, 2013, 2016; Tilton et al., 1981; Warren et al., 2005). The recently obtained 97±29 Ma Rb–Sr isochron age of listvenite fuchsite, and the fact that listvenites in the Samail ophiolite are only found along the basal thrust and associated mélangé zones, are consistent with syn-emplacement alteration of the ophiolite (Falk & Kelemen, 2015).

3 Materials and Methods

3.1 Macroscopic sample description and context in drill core

Samples used for this study were collected from core BT1B in the course of shipboard core logging aboard D/V Chikyu in September 2017 (Kelemen et al., 2020). BT1B was drilled in March 2017 in Wadi Mansah (23°21.861’ N, 58°20.149’ E), has a total length of 300.1 m, and comprises upper alluvium (2%), serpentinite (8%), listvenite (56%; quartz + carbonate), and the underlying metamorphic sole (34%) (Fig. 1b).

Serpentinite is present in two bands, one between 80 m and 100 m depth (Upper Serpentinite) and another between 181 m and 186 m (Lower Serpentinite) (Fig. 1b). Serpentinite displays a dark green to black color in hand specimen, has a cryptocrystalline grain size, and is typically massive but intersected by abundant veins of light green serpentine and carbonate (Fig. 1c). Carbonate-rich zones in the Upper Serpentinite occur between 80.54–80.73 m, 81.47–83.63 m and 86.82–88.07 m, and in gradational contacts between serpentinite and listvenite characterized by a distinct increase in carbonate vein abundance and color change to light green. The thickness of these zones ranges from 0.26 m to 3.2 m (Kelemen et al., 2020).

Listvenite is the dominant rock type in the ultramafic part of BT1B and present as three massive bands between the alluvium and the Upper Serpentinite (6.02 m–80.82 m), between the Upper and Lower Serpentinite (100.2 m–181.3 m) and between the Lower Serpentinite and the basal thrust (185.5 m–196.6 m) (Fig. 1b). Listvenite is fine-grained to cryptocrystalline and macroscopically heterogeneous with frequent color transitions on the centimeter to meter scale between dark and light red, orange, green, tan and pale grey (Fig. 1c). Red and orange listvenite is strongly brecciated in the drill core, and individual clasts show evidence for repeated brecciation and re-cementation such as truncation of clast-hosted veins at clast–matrix boundaries. Pale grey, tan and green listvenite are characterized by the presence of cm-sized quartz–fuchsite intergrowths situated in a pale to dark grey matrix. Tan, pale grey and green listvenite are mainly present between the Upper and Lower Serpentinite, are massive, and locally show foliation resembling ductile deformation.

The basal thrust (196.6 m–197.1 m) is developed as strongly foliated lower part of the lower listvenite unit and upper part of the underlying metamorphic sole separated by 0.42 m of brown to grey–green fault gouge. The metamorphic sole is composed of intercalated, foliated grey–green metabasalt and mafic metasediments. Hole BT1B did not reach the base of the metamorphic sole, but a recent review (Soret et al., 2017) reports that its thickness throughout the ophiolite “is commonly 10–100 m and does not exceed 500 m”. Falk & Kelemen
(2015) report bulk compositions and Sr isotope ratios for phyllites and carbonates from the underlying Hawasina formation in outcrop within a few kilometers of Hole BT1B.

The study presented here is based on six serpentinite samples from the Upper Serpentinite and nine listvenite samples consisting of light red, dark red and green listvenite from the Upper and Lower Listvenite (Fig. 1).

3.2 Carbonate micro-sampling for stable isotope analyses

A subset of samples was micro-sampled for carbonate stable isotope analysis (δ13C, δ18O) and clumped isotope thermometry. This sample subset includes matrix and vein magnesite, vein dolomite from the Upper Serpentinite (BT1B 43–02 and 44–03) and listvenite (Fig. 1). The majority of listvenite carbonate samples are from the Lower Listvenite, comprising green, light red and dark red listvenite, plus one additional light red listvenite from the Upper Listvenite. Listvenite matrix carbonate is mainly magnesite except for matrix dolomite from core sections BT1B 72–04 and BT1B 77–03. Listvenite vein dolomite was sampled from the Upper Listvenite (BT1B 32–02) and Lower Listvenite (BT1B 67–04) (see also Fig. S1).

Carbonate was sampled by using a micro drill with 3.8 mm inner diameter and a handheld Dremel tool from thin-section billets at sites selected based on the petrography of the respective thin sections. The relatively large diameter of the drill compared to the typical grain size and vein diameter, required sampling of mostly the homogeneous matrix areas and veins that were macroscopically devoid of crosscutting relationships. Carbonate samples were further crushed to a fine powder using an agate mortar and pestle prior to the analysis.

3.3 Analytical methods

3.3.1 Electron Microscopy

Electron microscopy was conducted on standard petrographic thin sections using an FEI Verios XHR FE-SEM at the Centre for Microscopy, Characterization and Analysis (CMCA), The University of Western Australia. A working distance of 4 mm and an acceleration voltage of 15 keV was used for BSE imaging. Additional TEM foils were prepared from spheroidal carbonate from serpentinite (BT1B 44–01) and listvenite (BT1B 60–03) by focused ion beam (FIB) milling at Utrecht University and CMCA using a FEI Helios Nanolab G3 CX DualBeam FIB-SEM at CMCA. Subsequent scanning transmission electron microscopy (STEM) imaging and quantitative electron dispersive X-ray spectroscopy (EDS) element mapping was carried out using a FEI Titan G2 80-200 at CMCA and a FEI Talos F200X at Utrecht University. Quantitative mineral compositions and element distribution maps were acquired using the JEOL 8530F FE electron microprobe at CMCA using an acceleration voltage of 15 keV and a focused and 5 µm defocused beam for silicate and carbonate minerals, respectively. The analytical procedure and application of reference materials follow the method described in Beinlich et al. (2018). Thin-section phase distribution mapping was conducted using a TESCAN Integrated Mineral Analyser (TIMA) high-resolution Field Emission Scanning Electron Microscope (FESEM) in full liberation mode at the John De Laeter Centre, Curtin University. Operating conditions were 15 keV, 0.8 nA and a working distance of 10 mm. Acquisition time per field was 227 seconds, and the map consists of 464 fields. Mineral abbreviations in the figures and tables follow the suggestions by (Whitney & Evans, 2010).

Crystallographic orientations of dolomite and magnesite in two samples BT1B 44–01 and BT1B 70–01 were measured by indexing Electron Backscatter Diffraction (EBSD) patterns acquired using a Hitachi SU3500 SEM equipped with an Oxford Instruments HKL EBSD Nordlys-Nano detector at the Geological Survey of Japan (GSJ). Before EBSD analysis, the thin sections were polished for several minutes with SYTON fluid using a Buehler VibroMet 2 to remove surface damage. Diffraction pattern acquisition was performed in low-vacuum mode using an accelerating voltage of 15 keV, a working distance of 18 mm, specimen tilt of 70° and step size of 0.5 µm. The orientation data were collected using the AZtec software and processed using the MTEX.
toolbox in Matlab™. All index data in this study represent points with a mean angular deviation (MAD) of <1°. Post-acquisition processing was done using a 10° segmentation angle for grain modeling, a cut-off grain size of 10 pixel, and the nearest-neighbor method to fill non-indexed areas.

3.3.2 X-ray micro-computed tomography

Visualization of the carbonate spherule distribution in serpentinite sample BT1B 44–01 in three dimensions was done using a Zeiss Versa 520 X-ray microscope at CMCA. The X-ray source operated at 80 keV and 7 W, and a Zeiss high-contrast, low-resolution 4X detector was used, achieving a spatial resolution of ~1.5 µm. The sample used was a hand-drilled 6 mm long and 3.8 mm diameter core from the thin-section billet used for petrography. Post-processing of the acquired 3D imaging data was done for a sub-volume of the sample core using the Avizo software, version 9.4.

3.3.3 Stable isotope analysis

Stable isotope analyses were performed at the GeoLab, Utrecht University, The Netherlands. Before isotope analysis, the mineralogy of each sample powder was constrained by XRD (Table S1; Beinlich et al., 2020d). The duration of acid digestion was adjusted to either dolomite or magnesite according to the dominant carbonate species in the sample powder. The carbonate δ13C and δ18O values are reported relative to Vienna Pee Dee Belemnite (VPDB).

Dolomite samples were digested in 103% phosphoric acid at 70°C for 20 minutes and the released CO2 was continuously collected in a liquid nitrogen trap using a Kiel IV carbonate device, coupled to a 253 Plus isotope ratio mass spectrometer (both instruments from Thermo Scientific) and analyzed in Long-Integration Dual-Inlet mode (Müller et al., 2017a; with 600 seconds integration time per aliquot). The weight of individual aliquots of reference materials and unknown samples ranged between 75–95 µg. The analyses were carried out in sequences consisting of 38 to 46 individual aliquots, of which 20 were carbonate reference materials. The reference materials used are calcite ETH-1 (δ13C = 2.20‰, δ18O = 6.70‰, Δ47 = 0.258), ETH-2 (δ13C = -10.0‰, δ18O = -10.0‰, Δ47 = 0.256) and ETH-3 (δ13C = 1.95‰, δ18O = 7.18‰, Δ47 = 0.691) (Bernasconi et al., 2018). Each sample and standard gas was measured against a CO2 reference gas (δ13C = -7.25‰, δ18O = 1.65‰), and each unknown sample was analyzed 10 to 14 times, spread over several measurement days.

Magnesite samples were digested offline, using 10–20 mg solid powder and 1–2 ml 103% phosphoric acid at 100°C for 15–16 hours in individual, sealed vials using a custom-built vacuum line containing a cold trap with liquid nitrogen acetone slush (-96°C) to remove H2O trace quantities from the CO2 gas. Each measurement sequence consisted of 6 samples, of which 3 were identically treated carbonate reference materials (ETH-1, ETH-2, ETH-3). Each sample analysis, including reference materials, was replicated 4 to 10 times. In contrast to the dolomite analyses, the magnesite analyses were conducted using the Dual Inlet of a Thermo Fisher Scientific MAT 253 in the traditional way by 8 alternating reference gas-sample gas cycles (208 seconds sample gas integration time per measurement).

Clumped isotope analyses of dolomite are reported as Δ47 in the absolute reference frame for 70°C reaction temperature without projecting the values down to 25°C acid digestion temperature due to the uncertainties in the dolomite specific temperature-dependent acid fractionation correction (see Müller et al., 2017b, 2019; Murray et al., 2016). The clumped isotope composition Δ47 is the measured abundance of the 13C18O16O isotopologue (m/z 47) in the carbonate acid digestion-released CO2 relative to an abundance that would be expected for a stochastic distribution of all isotopes. The deviation from the stochastic distribution is thermodynamically controlled (Eiler, 2007). The raw signal intensities were corrected for negative pressure baselines derived from the m/z 44 beam in the mass spectrometer following the recipe described in Meckler et al. (2014) and Müller et al. (2017b). We then constructed from the measured raw Δ47 values of the three ETH calcite standards and their accepted values an empirical transfer function to convert the raw Δ47 values of the unknown samples into the
absolute reference frame as described in Bernasconi et al. (2018). For the $\Delta_{47}$ temperature estimates of dolomite, we used the mineral specific $\Delta_{47}$-temperature calibration of Müller et al. (2019) for 70°C acid digestion temperature. For the dolomite $\delta^{18}$O values we applied the mineral-specific acid fractionation factor of 1.009926 for acid digestion at 70°C (Rosenbaum & Sheppard, 1986).

In contrast to dolomite, magnesite samples were processed identically to the calcite samples since earlier magnesite clumped isotope studies did not observe any differences in acid fractionations of magnesite and calcite (García del Real et al., 2016) and more elaborate studies on the magnesite $\Delta_{47}$ acid fractionation do not yet exist. The only published value for the magnesite $\Delta_{47}$ acid fractionation (Müller et al., 2017b) may be erroneous due to incomplete isotopic re-equilibration during heating of the magnesite in this study. Thus, we use the calcite $\Delta_{47}$-temperature calibration of Kele et al. (2015) recalculated with the new IUPAC (International Union of Pure and Applied Chemistry) parameters in Bernasconi et al. (2018) to derive our estimates of the carbonate formation temperatures. To transfer the raw $\Delta_{47}$ values of the magnesites into the absolute reference frame, we constructed an empirical transfer function from the measured raw

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**Figure 2** Representative back-scattered electron (BSE) images and quantitative element distribution maps of carbonate in serpentinite and listvenite. **a** Spheroidal magnesite–dolomite composite grains in the least carbonated serpentinite section 44–01. Quantitative element maps showing the chemical zonation of the carbonate. Ca-rich carbonate is identified as dolomite by Raman spectroscopy (not shown), whereas grain core regions are magnesite. FeO enrichment is confined to the magnesite. **b** Spheroidal carbonate in green listvenite core section 60–03. The carbonate growth morphology and chemical zonation is similar to that in serpentinite but Ca-rich carbonate is identified as Ca-magnesite that is also enriched in FeO. **c** Euhedral magnesite in green listvenite section 60–03. The core region of the euhedral magnesite grains resembles the morphology and composition of spheroidal magnesite shown in Fig. 2b.
$\Delta_{47}$ values of all analyzed ETH-1,-2,-3 standards (runs DI-3 to DI-12; Beinlich et al., 2019) against their accepted values at 70°C acid digestion temperature and applied the recalculated calcite $\Delta_{47}$-temperature calibration of Kele et al. (2015) for 70°C acid digestion temperature (by subtracting an acid fractionation correction factor of 0.062‰ from the intercept of the calibration equation). For the magnesite $\delta^{18}$O values we applied the acid fractionation factor 1.009178 for acid digestion at 100°C (Das Sharma et al., 2002).

4 Results

4.1 Microtexture and phase composition

4.1.1 Serpentinite

Macroscopic description during core logging aboard D/V Chikyu distinguished “serpentinite” from “ophicarbonate” (Kelemen et al., 2020). However, subsequent microtextural analysis shows that also those samples that were macroscopically described as serpentinite are carbonate-bearing. Thus, samples previously classified as either serpentinite or ophicarbonate samples are grouped in this section. The increase in carbonate content, from the least-carbonated serpentinite to listvenite, is gradational. The distinction between carbonate-bearing serpentinite from listvenite is based on the absence of quartz in serpentinite, and the lack of serpentinite in listvenite.

The serpentinite with the smallest proportion of carbonate is dominated by lizardite and chrysotile (see Figs S2a and b, S3, S4) together with chromite, magnetite, bastite intergrowths after pyroxene, minor carbonate and accessory µm-sized awaruite and is free of hematite and goethite. Primary olivine and pyroxene were not found in the investigated samples, and the serpentinite shows the typical mesh (hourglass) texture (Fig. S2a and b). Carbonate occurs in veins and as spheroidal grains within the serpentine matrix. Crosscutting relationships indicate that carbonate vein formation was preceded by at least two generations of serpentine veining. Carbonate veins frequently follow or deflect into earlier serpentine veins and sometimes occur in sub-parallel sets with regular vein spacing of ~130–200 µm (e.g. BT1B 44–03). The veins are composed of dolomite or magnesite or a combination of both. Magnesite veins show in most cases a distinct chemical zonation comprising a Fe-rich central part and Fe-poor outer part at the contact with serpentine (Table S2; Beinlich et al., 2020c). The composition of serpentine after primary olivine is identical within the analytical uncertainty between samples BT1B 44–01 and BT1B 44–03. Serpentine formed after orthopyroxene (i.e. bastite) shows slightly lower MgO and higher FeO concentrations (Table S3; Beinlich et al., 2020b).

Abundant spheroidal carbonate is observed in serpentinite and ophicarbonate samples BT1B 44–01 and BT1B 43–02, respectively (Fig. 2). These grains are characterized by their spherical shape and consistent grain diameters between 60 µm and 80 µm. BSE imaging and element distribution maps show a distinct chemical zonation consisting of relatively large Mg-rich cores, surrounded by a zone of increased FeO, and thin Ca-rich rims (Fig. 2a). Raman spectroscopy (Fig. S4) confirms that the Ca-rich zone of the spheroidal carbonate in serpentinite samples is dolomite. The chemical zonation is in some grains repetitive and defined by a second spherical dolomite zone (Fig. 2a). The outer rim of the carbonate spheroids is in most cases composed of dolomite and less frequently by a thin rim of magnesite (Fig. 3). Magnesite cores and dolomite rims are near their respective endmember compositions, while the outer magnesite core zones are characterized by FeO concentrations of up to 10 wt% (Fig. 2a). Larger matrix carbonate clusters, composed of amalgamated carbonate spheroids, preserve the described chemical and mineralogical zonation (Fig. 2a). Carbonate spheroids in the serpentinite frequently contain sub-µm sized serpentine inclusions (Fig. 3).

4.1.2 Listvenite

Listvenite samples are dominantly composed of carbonate, quartz, and relic chromite and generally lack precursor serpentine and primary magmatic silicate phases. Red listvenite contains additional fine-grained hematite and potentially goethite intergrown with carbonate and quartz. Matrix carbonate is dominated by magnesite, while carbonate veins are either magnesite or
dolomite. Occasionally, the distribution of oxide strings in listvenite resembles that typically associated with a serpentine mesh (or hourglass) texture (Fig. S2c and d). Based on their crosscutting relationships, thin, sometimes Fe-oxide-bearing magnesite veins (10–50 µm) represent in most cases the oldest vein generation. They are crosscut by thicker magnesite veins (< 1 mm), followed by quartz and/or quartz–carbonate veins (50–300 µm), and then by dolomite veins (<500 µm). The thicker magnesite veins occasionally contain euhedral magnesite showing an oscillatory variation in its Ca content. Additionally, coarse-grained magnesite veins frequently show a vein-internal chemical zonation, composed of near-endmember magnesite at the vein-matrix interface as well as high Fe and Ca concentrations in the vein center. Matrix quartz in the listvenite is in most cases fine-grained to cryptocrystalline and shows an interlocked texture. Less frequently, the quartz matrix appears to consist of clusters of sub-µm sized spheroidal quartz, creating an interstitial porosity (BSE images in Fig. 2b and c).

Green listvenite samples (e.g. BT1B 52–01, BT1B 60–03) contain additional fuchsite (Cr-muscovite, Table S3; Beinlich et al., 2020b) as intergrowths with quartz and associated with relic chromite. These samples also contain spheroidal carbonate (~80 µm grain diameter) that texturally resembles the matrix carbonate in serpentinite samples. Spheroidal carbonate in the listvenite occurs in the fine-grained quartz matrix as isolated grains and amalgamated clusters of individual spheroids. Listvenite carbonate spheroids also show a systematic compositional zonation with high Mg concentrations in the core and elevated Ca and Fe concentrations in the rim (Fig. 2b). However, the rims are composed of Ca- and Fe-rich magnesite instead of dolomite. Spheroidal magnesite is frequently overgrown by compositionally distinct euhedral magnesite (Fig. 2c). Spheroidal carbonate in the listvenite contains abundant, fine-grained (≤100 nm) spherical silica inclusions, which may be quartz or amorphous silica (Fig. 4). In addition, magnesite in the listvenite occasionally displays dendritic growth textures with initial Fe-richer branches and later Fe-poorer overgrowths (Fig. S5).
4.2 Carbonate stable isotope composition and clumped isotope thermometry

Dolomite and magnesite extracted from veins in serpentinite and listvenite, and the listvenite matrix were analyzed for their δ¹³C (VPDB), δ¹⁸O (VPDB), and Δ₄⁷ composition. Δ₄⁷ values range between 0.275‰ and 0.548‰. The corresponding spread in formation temperatures for all samples is 45±5 °C (1σ) to 247±52°C (Fig. 5a, Table S1; Beinlich et al., 2020d). Listvenite matrix magnesite formation temperatures range between 52±8°C (BT1B 52–01) and 190±31°C (BT1B 68–02) and listvenite matrix dolomite temperatures between 98±15°C (BT1B 72–04) and 107±19°C (BT1B 77–03) (see also Fig. S1). The dolomite and magnesite veins in the upper serpentinite yielded 227±52°C (BT1B 43–02) and 45±5°C (BT1B 44–03), respectively. In the listvenite, magnesite vein–matrix pairs indicate a higher temperature for the vein (T_vein-T_matrix = 11°C and 56°C), whereas the matrix magnesite in BT1B 67–04 records a temperature 46°C higher than the vein dolomite in the same sample.

Dolomite δ¹³C values range from -2.6‰ to 0.70‰ and magnesite δ¹³C values range from -3.8‰ to 0.60‰. Dolomite δ¹⁸O values range from -13.8‰ to -6.2‰ and magnesite δ¹⁸O values range from -9.3‰ to -6.9‰ (Fig. 5b and c, Table S1; Beinlich et al., 2020d). In the δ¹³C vs. δ¹⁸O space (Fig. 5c), carbonate stable isotope compositions cluster in three groups characterized by equal δ¹³C but distinct δ¹⁸O, and intermediate δ¹⁸O and low δ¹³C. Listvenite vein–matrix magnesite pairs (BT1B 60–03 and BT1B 68–02) show higher matrix δ¹⁸O of 0.23‰ and 0.40‰, while δ¹³C values are 0.26‰ higher and -0.17‰ lower in the matrix compared to the vein. Vein dolomite from BT1B 67–04 has a lower δ¹⁸O compared to the matrix magnesite (Δ¹⁸O = -5.9‰) but slightly higher δ¹³C (Δ¹³C = 0.27‰) (Fig. 5b and c, Table S1; Beinlich et al., 2020d).

5 Discussion

5.1 Incipient carbonation, carbonate nucleation and growth

Conceptually, the per-sample amount of carbonate can be related to reaction progress from incipiently altered serpentinite, containing only some carbonate, to fully carbonated listvenite. Along this conceptual reaction sequence, the carbonate growth texture evolves from isolated spheroids, through clusters of amalgamated

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Figure 5 Stable O- and C-isotope systematics. a Down-hole plot of clumped isotope carbonate formation temperatures for listvenite and serpentinite samples. Colors and symbols refer to the legend shown in Fig. 5c and differentiate between matrix and vein magnesite and dolomite. The downhole stratigraphy is identical to that shown in Fig. 1. b Down-hole plot of magnesite and dolomite δ¹⁸O (VPDB). c Carbonate δ¹⁸O (VPDB) vs δ¹³C (VPDB) showing clustering at high oxygen and carbon isotope ratios, lower δ¹⁸O for some of the listvenite matrix and vein dolomite and lower δ¹³C for vein dolomite and magnesite from serpentinite sample BT1B 44–01.
carbonate spheroids, to spheroids overgrown by compositionally distinct and euhedral carbonate. The spheroidal carbonate texture in serpentinite (e.g. BT1B 44–03; Fig. 2a) thus represents a snapshot of the incipient carbonation stage. The similar spheroid diameters imply nearly constant rates and durations of nucleation and growth. The presence of abundant carbonate veins, even in the serpentinite, suggests that the initial fluid infiltration was channelized and facilitated by tectonic deformation of the serpentinite host rock. This is further supported by 3D textural analysis showing the existence of smaller veinlets connecting individual carbonate spheroids with larger veins (Fig. 6). However, the 3D textural analysis also shows that a relatively large number of carbonate spheroids occur isolated from carbonate veins. Consequently, further migration of reactive fluid must have occurred by diffusion through interconnected, fluid-filled pore space.

Analyses of TEM dark-field images suggest a porosity of 2.68±1.02% for the serpentinite. Most individual pore diameters are less than 100 nm (Fig. 7). At this scale, nanofluidic transport phenomena can be significant and effectively govern metamorphic fluid-flow, mass transport and hence carbon mineralization (Plümper et al., 2017a; Zheng et al., 2018). These transport phenomena are consistent with the relatively even distribution of carbonate spheroids on the thin-section scale (Fig. 8a and b), suggesting that the spatial distribution of carbonate nucleation sites and by implication also the pore fluid saturation state was homogeneous during nucleation. Indeed, the carbonate morphology is in agreement with a strong driving force causing spherical growth instead of sub- to euhedral Mg-carbonate that is frequently observed in similar alteration systems elsewhere (Beinlich et al., 2012; Hansen et al., 2005; Sunagawa, 2005).

Based on the premise that spherulitic growth may result in morphologically similar carbonate and would indicate fast precipitation under far-from-equilibrium conditions (Gránásy et al., 2005; Shtukenberg et al., 2012), we further apply EBSD analysis to representative carbonate spheroids (Fig. 9). This crystallographic examination indicates that spheroidal magnesite is composed of a single crystal and contains abundant, predominantly radial low-angle boundaries (2°–10°) that originate from small dolomite inclusions in the grain centers. High-angle boundaries are exclusively present in the dolomite (Fig. 9a and b). Despite the resemblance with spherulitic carbonate, the fact that carbonate spheroids from BT1B are primarily single crystals is inconsistent with polycrystalline
spherulitic growth. Instead, the shape and orientation of such magnesite grains are similar to sector zoning occasionally found in, e.g., garnet and fluorite (e.g., Bosze & Rakovan, 2002; Stowell et al., 2011). Sectorial zoning is commonly related to compositionally distinct zones and ascribed to orientation-dependent differences in crystal growth rates. However, growth competition may also result in crystallographic misfit between adjacent sectors of identical composition, thus forming low-angle boundaries in chemically homogeneous single crystals that are unrelated to external deformation (see also Pearce et al., 2013).

5.2 Pervasive carbonation and listvenite formation

Listvenite contains spheroidal and euhedral carbonate, both showing a systematic chemical zonation with Mg-rich and Ca-poor grain cores, and Fe–Ca-rich rims. Spheroidal magnesite in the listvenite shows a gradual increase in MgO toward the core and a sharp transition to Ca- and Fe-rich magnesite in the rim. Theoretically, the textural similarity of carbonate in the listvenite and serpentinite may be interpreted as to indicate that spheroidal carbonate in the listvenite is inherited from the initial carbonation stage represented by spheroidal carbonate in the serpentinite. This scenario would involve dissolution of the dolomite rims and re-precipitation of Ca-bearing magnesite in a CO₂-rich reactive fluid. However, the core region of spheroidal carbonate in the listvenite has 4–5 wt.% FeO, whereas the core region of spheroidal carbonate in the serpentinite has less than 2 wt.% FeO. Furthermore, listvenite carbonate spheroids contain abundant sub-μm sized silica inclusions (Fig. 4), whereas serpentinite magnesite contains serpentine inclusions (Fig. 3). These characteristics suggest that the listvenite carbonate did not evolve from spheroidal carbonate in the serpentinite, but instead formed via direct replacement of serpentine, accompanied by the formation of quartz in the matrix. EBSD phase- and misorientation-maps show that spheroidal carbonates in the listvenite also consist of magnesite single crystals with radially orientated low-angle boundaries that originate from a central dolomite inclusion (Fig. 9c and d). Thus, we infer that the listvenite carbonate spheroids also precipitated at relatively high supersaturation, resulting in growth competition between adjacent grain sectors and the spheroidal crystal habit as well as the less frequently observed dendritic texture of the magnesite (Fig. S5) (Haque et al., 2016; Sunagawa, 2005).

Assuming that Ca is primarily sourced from the dissolution of clinopyroxene, the low CaO concentration in magnesite rims in the listvenite compared to the serpentinite is likely related to the larger total carbonate amount in the sample (dilution effect). High concentrations of FeO in rims of spheroidal magnesite and the euhedral magnesite overgrowths (Fig. 2b and c) indicate
increased availability of Fe during a relatively late stage of listvenite formation, i.e. at the time when the carbonate rims and overgrowths formed relative to carbonate core zones. This availability of FeO is attributed to the breakdown of oxide phases (chromite, magnetite) in CO$_2$-rich alteration fluid during listvenite formation, as previously documented for sites in Canada and Norway by Hansen et al. (2005) and Tominaga et al. (2017). Perhaps, this occurred at the same time as crystallization of fuchsite, which formed from reactions involving spinel, silicates and fluid. The transition from spheroidal to euhedral carbonate morphology reflects a driving force reduction during continued crystal growth (Sunagawa, 2005) and may be related to cooling and/or decrease in carbonate saturation.

5.3 Carbonation temperature and depth

The clumped isotope distribution in carbonate is thought to be independent of the isotopic composition of the fluid from which the carbonate precipitates (e.g., Eiler, 2007; Ghosh et al., 2006; Huntingon et al., 2009), though this is a theoretical prediction that has not been rigorously tested over a broad range in fluid composition and temperature, particularly for dolomite and magnesite. Carbonate clumped isotope signatures can experience solid-state reordering during cooling, depending on the carbonate mineralogy and the cooling rate (Passey & Henkes, 2012). However, dolomite is relatively resistant to solid-state reordering at temperatures below 300°C, thus the temperatures in our study likely reflect the dolomite formation temperatures (e.g. Bonifacie et al., 2017; Ryb et al., 2017). In the case of magnesite, solid-state reordering is less well studied. Nevertheless, its sluggish reaction kinetics with phosphoric acid and evidence for an equilibrium blocking temperature of approximately 490°C (García del Real et al., 2016) suggest that also the magnesite clumped isotope composition is resistant to isotopic reordering. Hence, we assume that the measured clumped isotope compositions of our carbonate samples reflect their formation temperatures.

The clumped isotope temperatures complement earlier thermometry work in the Samail ophiolite that described relatively low formation temperatures (clumped isotope temperatures between 23–43°C) for young carbonate veins in peridotite forming during present-day weathering (Streit et al., 2012), slightly higher temperatures for calcite veins in peridotite beneath a Cretaceous unconformity (25–60°C, de Obeso & Kelemen, 2018), and yet higher temperatures for carbonate from outcrop samples of listvenite and the metamorphic sole within a few kilometers of Hole BT1B (listvenite 61–114°C, n=15; metamorphic sole 64–171°C, n=4) (Falk & Kelemen, 2015). Also, Falk & Kelemen (2015) documented the presence of intergrown quartz and antigorite, with and without talc and magnesite, in several samples from a gradational contact between listvenite and serpentinite about 2 km north of Hole BT1B. These mineral assemblages equilibrated with a relatively H$_2$O-
For listvenite carbonate from BT1B analyzed in this study, the clumped isotope data yield a relatively wide range of formation temperatures, 52–247°C. In detail, matrix magnesite from the Lower Listvenite (118.5 m to 165 m depth) formed between 52–190°C, and listvenite matrix dolomite sampled near the basal thrust (178.7 m and 193.2 m depth) formed at temperatures of 98–107°C. Vein dolomite from the red Upper Listvenite formed at ~140°C (67.4 m), whereas vein dolomite and magnesite from the relatively narrow Upper Serpentinite yielded one of the highest (227°C) and lowest (45°C) formation temperatures, respectively (Fig. 5, Table S1; Beinlich et al., 2020d). Magnesite formation temperatures appear to increase with depth, whereas dolomite formation temperatures are relatively even within the uncertainty except for the high-temperature dolomite vein in the Upper Serpentinite (227°C; BT1B 43–02). Thus, our data show that the overall range in carbonate formation temperatures is wider than previously constrained by carbonate sampled in surface outcrops (Falk & Kelemen, 2015). Furthermore, the relatively erratic temperature distribution throughout core BT1B may reflect that...
the depth of carbonate formation varied non-

systematically with time, consistent with the

repeated formation of deformation-induced high-

permeability pathways allowing reactive fluid

infiltration into different structural levels upon

cooling of the system.

Assuming that cooling is related to ophiolite

obduction, the obtained temperatures can be

translated to carbonate formation depth (y), using a

simple equation for conductive heat transfer:

\[ y = \frac{q_0 - \sqrt{2T_0 H} k \rho - 2H k \rho T + q_0^2}{H \rho} \]

where \( q_0 \) is the surface heat flow (0.06±0.01 W∙m\(^{-2}\)

in forearc settings; Blackwell et al., 1990; Lewis et al.,

1988), \( T_0 \) is the temperature at the ocean floor

(277 K), \( H \) is the current mean radioactive heat

generation in the mantle (7.42×10\(^{-12}\) W∙kg\(^{-1}\))

(Turcotte & Schubert, 2014), \( k \) is the thermal rock

conductivity (2.9 W∙m\(^{-1}\)K\(^{-1}\)), \( \rho \) is the density of

partially serpentinized peridotite (2900 kg∙m\(^{-3}\)),

and \( T \) is the carbonate formation temperature (see

also Plümper et al., 2017b).

The calculated depth is mainly dependent on

chosen surface heat flow and thermal conductivity

values. The 2.9 W∙m\(^{-1}\)K\(^{-1}\) thermal rock

conductivity is based on measurements of

serpentinized peridotite from the Mid-Atlantic

Ridge (Kelemen et al., 2004), in agreement with the

thermal properties of olivine and antigorite

(Hofmeister, 1999; Osako et al., 2010). The paleo-

surface heat flow is unknown but can be

constrained based on pressure and temperature

conditions of ~850°C and 1 GPa recorded in the

metamorphic sole underlying the ophiolite (Agard

et al., 2016; Soret et al., 2017). The resulting

surface heat flow (0.06±0.01 W∙m\(^{-2}\)) is consistent

with these criteria and also with the inferred

Permian age of the Samail mantle section and the

~92 Ma mica Ar–Ar ages from the metamorphic

sole (Hacker et al., 1996; Jaupart et al., 2015; van

Hinsbergen et al., 2019). The resulting geothermal

gradient is 25±0.6°C km\(^{-1}\). If our assumptions are

correct, then the clumped isotope temperatures

place the carbonation depth between ~10.1 km at the

highest temperature and 1.7 km at the lowest

temperature, thus recording ophiolite uplift of ~8.4

km (Fig. 10; Fig. S6).

5.4 Implications for reactive fluid composition

Ultramafic rock carbonation to listvenite is

driven by the infiltration of CO\(_2\)-bearing reactive

fluid into the ophiolite causing the dissolution of

serpentine and precipitation of secondary Ca–Mg

and Mg-carbonate in the serpentine and

additional quartz in the listvenite. Primary olivine

was not found in core BT1B suggesting that

serpentinization went to completion before CO\(_2\)-

fluid infiltration, consistent with the mesh-textured

distribution of oxides in listvenite (Fig. S2). Hence,

the carbonation reaction can be conceptualized, in Fe- and Ca-free space, as:

\[
\text{Mg}_5\text{Si}_2\text{O}_8(\text{OH})_4 + 3 \text{CO}_2,\text{aq} \rightarrow \\
3 \text{MgCO}_3 + 2 \text{SiO}_2(\text{aq}) + 2 \text{H}_2\text{O}
\]

In the serpentinite, the reaction neither went to

completion nor resulted in quartz/silica saturation

of the alteration fluid and SiO\(_2\),aq was likely

removed by the alteration fluid. In contrast,

quartz/silica saturation was reached in the

pervasively carbonated parts of BT1B that are now

present as listvenite. The breakdown of relict

clinopyroxene may have provided Ca for dolomite

growth during limited carbonation of the

serpentinite (e.g., in BT1B 44–01), whereas the

significantly larger total amount of carbonate in

the listvenite may have prevented dolomite

precipitation instead of Ca-bearing magnesite.

Additional replacement of primary Cr-spinel

results in the formation of fuchsite and implies the

addition of potassium by the alteration fluid.

Similarly altered ultramafic sequences have been

described from several other ophiolite occurrences

(e.g. Beinlich et al., 2012; Halls & Zhao, 1995;

Hansen et al., 2005; Hinsken et al., 2017) and have

been the subject of thermodynamic models linking

mineral assemblages to fluid composition at a

given alteration pressure and temperature

(Beinlich et al., 2020a; Klein & Garrido, 2011;

Klein & McColloM, 2013). However, the

spheroidal carbonate texture in both the

serpentinite and listvenite and the dendritic

carbonate in the listvenite (Figs 2, 6, 9, and S5) are

indicative of textural disequilibrium complicating

constraints on the alteration fluid carbon
fractionation between 31‰ and 9.5‰ over the same temperature range. For dolomite, the predicted oxygen isotope fractionation varies between 22‰ and 7.2‰ for the temperature range 98–227°C based on the most recent temperature relationship by Horita (2014). In the absence of experimental data on the temperature dependence of magnesite-fluid oxygen isotope fractionation, we here adopt the calibration by Aharon (1988) that is consistent with that by Zheng (1999) and applicable over the investigated temperature range, whereas the magnesite-H₂O equation by Chacko & Deines (2008) is only applicable below 130°C. Furthermore, for oxygen isotope fractionation between dolomite and fluid, we use the calibration by Horita (2014), calibrated over the temperature range 80–350°C. By applying these fractionation factors to the measured oxygen isotope composition of dolomite and magnesite samples that were used for clumped isotope thermometry defines an apparent trend from relatively elevated δ¹⁸Ofluid at a higher temperature (e.g. 12.2‰ at 247°C) to relatively lower δ¹⁸Ofluid at a lower temperature, e.g. -9.9‰ at 45°C. The different calibrations of the oxygen isotope fractionation factors result in a relative variation of δ¹⁸Ofluid of less than 3.7‰ at given temperature (Fig. S6; see also Garcia del Real et al., 2016).

As previously observed for the area around Hole BT1B (Falk & Kelemen, 2015; their Supplementary Figure 8 and related text), and low-temperature carbonates in altered peridotite worldwide (Garcia del Real et al., 2016), the resulting calculated δ¹⁸Ofluid displays a positive correlation with the corresponding carbonate formation temperature (Fig. 10). This correlation may indicate carbonate precipitation from fluids that changed their oxygen isotope ratios and/or were derived from compositionally distinct sources. Low-temperature and low δ¹⁸O fluids may have been sourced from sediment compaction containing seawater-filled pore space during tectonic underthrusting (Falk & Kelemen, 2015; Kelemen et al., 2011), whereas high-temperature fluids with elevated δ¹⁸O may be released by subduction zone devolatilization reactions at the greenschist to amphibolite transition (Menzies et al., 2018). These different fluid sources can be reconciled with the measured carbonate compositions if the seawater δ¹⁸O remained fluid-

Figure 10 Calculated equilibrium fluid δ¹⁸O (VSMOW) (Aharon, 1988; Horita, 2014) plotted against carbonate clumped isotope temperatures. The secondary abscissa shows depth inferred from the carbonate formation temperatures and simple heat conductivity calculation. The gray shaded area and transparent data points outline the range in fluid δ¹⁸O and temperature based on Falk & Kelemen (2015). Error bars reflect the propagated analytical uncertainty, standard deviation of the carbonate formation temperature, and the uncertainty of the mineral–fluid isotope fractionation calibration. Equilibrium fluid δ¹⁸O values calculated using various mineral–fluid oxygen isotope fractionation factors are shown in Fig. S6.

Assuming that the system had sufficient time to attain isotopic equilibrium between dissolved carbon species, the aqueous fluid, and the precipitating carbonate at the temperatures observed in this study, we can evaluate the fluid δ¹⁸O (δ¹⁸Ofluid) based on the measured carbonate composition and available temperature-dependent fractionation factors (Aharon, 1988; Chacko & Deines, 2008; Horita, 2014; Matthews & Katz, 1977; Northrop & Clayton, 1966; Schauble et al., 2006; Schmidt et al., 2005; Zheng, 1999). The predicted oxygen isotope fractionation between magnesite and fluid (10⁷lnαMgsfluid) varies between 34‰ and 7.8‰ over the temperature range 45–247°C. The strongest fractionation is predicted by Chacko & Deines (2008; magnesite-H₂O) at low temperature (34.9‰), which is distinctly stronger than the predicted fractionation by Zheng (1999) of ~27.3‰. The magnesite-fluid calibration by Aharon (1988; using magnesite 2) yields
buffered while trapped in the sedimentary pore space. Also, recent thermodynamic modelling has shown that magnesite–quartz assemblage requires fluid CO$_2$ concentrations of ~2500 ppm (at 100 °C and 5 kbar), i.e. higher than that of low-pressure fluid saturated with calcite, suggesting that CO$_2$ was primarily derived from subduction zone devolatilization (de Obeso et al., 2017; Kelemen et al., 2017).

Alternatively, the correlation between carbonate formation temperatures and calculated $\delta^{18}$O$_{\text{fluid}}$ may be unrelated to the fluid composition and instead reflect a tendency toward departure from equilibrium of fluid–carbonate oxygen isotope exchange with decreasing temperature. Fluid–mineral oxygen isotope exchange may operate independently from the ordering of clumped isotopes in the carbonate mineral. Further experimental work on the relative rates of isotope exchange and ordering is required to fully investigate the apparent correlation of calculated $\delta^{18}$O$_{\text{fluid}}$ and measured clumped isotope temperatures. However, independently determined fractionation factors are mutually consistent despite relative offsets (Fig. S6), and a measurement artifact would be expected to cause a correlation between the measured carbonate clumped isotope composition and $\delta^{18}$O$_{\text{carbonate}}$.

5 Conclusions

Oman DP Hole BT1B intersects 200 m of listvenite containing two relic bands of the precursor serpentinite. The alteration to listvenite was pervasive, forming an assemblage consisting of Mg- and Ca–Mg-carbonate, quartz, minor fuchsite and relic chromite. The presence of hematite and/or goethite in some parts of the core cannot be unambiguously linked to listvenite formation and may also result from a later overprint of already formed listvenite under oxidizing conditions. The least altered parts of BT1B are composed of serpentinite containing minor carbonate, while precursor olivine and pyroxene minerals could not be found during this study and earlier shipboard core logging. While we cannot rule out with certainty that carbonation was accompanied by serpentinization, the presence of oxides strings in carbonate resembling those typically found in serpentinized peridotite suggests that serpentinization went to completion prior to the carbonation.

The entire ultramafic part of the core is pervasively veined. Crosscutting relationships in the serpentinite indicate that carbonation was predated by at least two generations of serpentine veins. In the listvenite, crosscutting relationships indicate a general trend of early magnesite and minor dolomite veins being followed by the formation of a second set of broader magnesite veins, that is subsequently crosscut by quartz and quartz–carbonate veins, and then by dolomite veins. Whether the crosscutting relationships in the listvenite represent a network of fluid pathways during one alteration event or provide snapshots of consecutive fluid infiltration requires continued investigation beyond this contribution.

Carbonation of the serpentinite matrix resulted in the formation of abundant spheroidal magnesite–dolomite composite grains that show an even spatial and grain size distribution at the thin-section scale and appear to have nucleated on µm-sized, early dolomite. The textural analysis indicates that also listvenite formation commenced with the formation of spheroidal carbonate. However, the carbonate spheroids in the listvenite are chemically and mineralogically distinct from those in the serpentinite. This suggests that the carbonate-bearing serpentinite and listvenite are unlikely to represent consecutive steps within the same reaction sequence but are the result of concomitant alteration, controlled by the accessibility of the alteration fluid, consistent with the preservation of serpentinite in BT1B.

The spheroidal and less commonly observed dendritic carbonate growth textures indicate fast precipitation from over-saturated fluid implying non-equilibrium conditions during serpentinite carbonation and the initial stage of listvenite formation. This may explain the pervasive nature of the alteration in those parts of BT1B that were accessible by the alteration fluid and also the lack of discernable talc-bearing transitional assemblages between the serpentinite and the listvenite that would be expected to form at chemical equilibrium in the temperature range defined by the clumped isotope thermometry. Spheroidal magnesite in the listvenite is frequently overgrown by euhedral magnesite and both texturally distinct carbonate generations crystallized together with quartz. The formation of
euhedral carbonate overgrowths in the listvenite reflects a driving force reduction and may be related to cooling and/or decrease in fluid saturation during the final stage of listvenite formation.

Clumped isotope thermometry of selected matrix and vein magnesite and dolomite indicates an overall carbonation temperature range of 45 to 247°C. Carbonate formation temperatures show a non-systematic distribution with sample depth in BT1B, indicating that reactive fluid could infiltrate the precursor ultramafic rock at different structural levels and times. Furthermore, the calculated oxygen isotope composition of the carbonate forming fluid varies between ~9.9‰ and ~12.1‰ and shows an apparent correlation with temperature. This correlation is consistent with previous observations in Oman (Falk & Kelemen, 2015) and suggests that carbonation took place as a multi-stage process driven by compositionally evolving reactive fluids that infiltrated the ophiolite during cooling.

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Data availability
All data are available online. These include the processed carbonate stable isotope data and textural description of clumped isotope sampling spots, the major element composition of carbonate phases, the major element composition of silicate phases, and the complete set of stable isotope data of all analyzed reference materials and unknown samples.
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Supporting Information:

- Figures S1 to S6
- All supporting tables are available at www.pangaea.de and can be found under the DOIs provided in the reference list.
- An animated version of the 3D reconstruction shown in Fig. 6 can be found together with the journal version of this article at DOI: 10.1029/2019JB019060
Figure S1 Sampling locations for clumped isotope analyses in all investigated samples. Images on the left side are full thin-section scans using cross-polarized light. Images on the right side show the location of micro drill spots in the thin section billets.
**Figure S2.** Photomicrographs showing the mesh texture in serpentine (a and b) and listvenite (c and d). a Serpentine sample BT1B 39-04 showing a typical mesh texture composed of lizardite/chrysotile serpentine and magnetite strings. Bastite after primary orthopyroxene is present in the center and bottom (plane-polarized light). b Cross-polarized photomicrograph of the same sample as shown in a. c Listvenite sample BT1B 52-01 showing relics of the serpentinization mesh texture defined by magnetite strings in magnesite and quartz (plane-polarized light). d Cross-polarized photomicrograph of the same sample as shown in c.
Figure S3. Scanning transmission electron microscope (STEM) bright-field images from sample BT1B 44-01 showing that the serpentine matrix is predominantly composed of lizardite and chrysotile (see also Raman spectrum in Fig. S3).
Figure S4. Principal component analysis (PCA) of Raman spectra (left) acquired on a zoned carbonate spheroid in serpentinite sample BT1B 44–01 showing the presence of magnesite in the core surrounded by a thin rim of dolomite in a matrix of chrysotile. Raman spectra (right) are the average spectra for the three zones of the PCA map. Spectra were acquired with a WITec Alpha 300 RA+ at CMCA using the 532 nm laser and a 600 nm/line grating. Raman mapping was carried out at a spatial resolution of 700 nm.
Figure S5. Back-scattered electron (BSE) image of dendritic magnesite in the quartz matrix of listvenite sample BT1B 60-01. The variation in BSE contrast within the magnesite is caused by unquantified variations in FeO concentration.
Figure S6. Calculated equilibrium fluid δ^18O (VSMOW) values plotted against carbonate formation temperatures based on various mineral–fluid oxygen isotope fraction factors (see also Fig. 10). Error bars reflect the propagated analytical uncertainty, standard deviation of the carbonate formation temperature, and the uncertainty of the mineral–fluid isotope fractionation calibration.