Stress evolution and associated microstructure during transient creep of olivine at 1000–1200 °C

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ABSTRACT
We study the mechanical response and correlated microstructure of axial deformed fine-grained olivine aggregates as a function of incremental finite strains. Deformation experiments were conducted in uniaxial compression in an internally heated gas-medium deformation apparatus at temperatures of 1000 and 1200 °C, at strain rates of 10⁻⁶ s⁻¹ to 10⁻⁵ s⁻¹ and at a confining pressure of 300 MPa. Sample volumes are around 1.2 cm³.

Finite strains range from 0.1 to 8.6% and corresponding maximal (final) differential stresses range from 80 to 1073 MPa for deformation at 1000 °C and from 71 to 322 MPa for deformation at 1200 °C. At 1200 °C, samples approach steady state deformation after about 8% of strain. At 1000 °C, significant strain hardening leads to stresses exceeding the confining pressure by a factor of 3.5 with brittle deformation after 3% of strain. Deformed samples were characterized by electron backscatter diffraction (EBSD) and transmission electron microscopy (TEM). EBSD maps with step sizes as low as 50 nm were acquired without introducing analytical artifacts for the first time. The grain size of deformed samples ranges from 2.1 to 2.6 µm. Despite clear strain hardening, texture or microstructure do not change as a function of stress or finite strain. This observation is supported by a constant texture strength (J-index and symmetry (BA-index), constant grain shape and aspect ratio, constant density of geometrically necessary dislocations, grain orientation spread, and constant subgrain boundary spacing and misorientation in between samples. TEM shows that all samples exhibit unambiguous dislocation activity but with a highly heterogeneous dislocation distribution. Olivine grains display evidence of [1 0 0] and [0 0 1] slip activity, but there is no evidence of interaction between the dislocations from the different slip systems. Several observations of grain boundaries acting as dislocation sources have been found. We find no confirmation of increasing dislocation densities as the cause for strain hardening during transient creep. This suggests other, yet not fully understood mechanisms affecting the strength of deformed olivine. These mechanisms could possibly involve grain boundaries. Such mechanisms are relevant for the deformation of uppermost mantle rocks, where the Si diffusion rate is too slow and dislocation glide must be accommodated in another way to fulfill the von Mises criterion.

1. Introduction
Olivine is the most abundant (> 60%vol) and the weakest mineral phase of Earth’s upper mantle. As such, it largely controls the rheology of the upper mantle and in turn the contribution of mantle convection to the formation and motion of tectonic plates. Since the upper mantle encompasses rocks within a wide range of temperatures, its rheology has to be studied in an equally broad thermal range.

Early flow laws quantifying the rheology of olivine were mostly obtained from experiments at temperatures above 1200 °C (e.g. Chopra & Paterson, 1984; Karato et al., 1986; Bai et al., 1991; Hirth & Kohlstedt, 1995; Mei & Kohlstedt, 2000a,b; Faul et al. 2011; Keefner et al., 2011; Tielke et al., 2017) to avoid to reach the brittle field. Experiments performed at temperatures relevant to the uppermost lithospheric mantle (≤1000 °C, e.g. Evans & Goetze, 1979; Raterron et al., 2004; Demouchy et al. 2009, 2013, 2014; Mei et al., 2010; Idrissi et al., 2016; Tielke et al., 2016) report a lower strength of olivine than was expected from extrapolation of high-temperature flow laws. The low temperature plasticity is a key to understanding deformation of the shallow upper mantle (Hirth & Kohlstedt, 1995) but also in subducting slabs (Goetze & Evans, 1979; Kirby, 1980; Karato et al., 2001), where it governs the stress storage capacity and therefore the maximal stress release of deep earthquakes (e.g., Proietti et al., 2016).

Until recently, the deformation and microstructure of olivine were
characterized under a so-called mechanical steady state, when stress becomes constant and independent of plastic strain. Therefore, the mechanical response and evolution of microstructure before reaching steady state (i.e., transient creep) is largely not understood. The initial rapid increase in stress is traditionally attributed to an increase in the dislocation density, which is reported to become constant in single crystals after about 2% of strain (e.g., Durham et al., 1977). Even under constant dislocation densities, the dislocation structures continue to evolve and reaching a constant dislocation structure requires a finite amount of strain. Transient creep at temperatures and pressures relevant for the lithospheric mantle might prove crucial to understanding the initiation and localization of deformation processes that impact lithospheric plate generation and motion.

The main deformation mechanisms of olivine are 1) glide and climb of dislocations in the crystal lattice, typically called dislocation creep 2) diffusion of atoms and vacancies at temperatures above approximately 1200 °C, which is grain size sensitive (i.e., diffusion creep) and 3) grain boundary controlled mechanisms, such as grain boundary migration or grain boundary sliding. The dominant deformation mechanism depends on the stress, temperature, grain size, strain rate, the crystal chemistry, the oxygen fugacity, partial melt content and even the trace amount of hydrogen (e.g. Carter and Avé Lallemant, 1970; Chopra & Paterson, 1981, 1984; Mackwell et al., 1985; Cooper & Kohlstedt, 1984, 1986; Bai et al., 1991; Bai & Kohlstedt, 1992; Mei & Kohlstedt, 2000a,b; Hirh & Kohlstedt, 1995, 2003; Warren & Hirh, 2006; Mackwell, 2008; Paul et al., 2011, 2016; Keefner et al., 2011; Demouchy et al., 2012; Tielke et al., 2015, 2017). Dislocation glide is thought to be the dominant deformation mechanism at low temperatures (< 1200 °C in Fo80) and for grain sizes > 1 mm prevalent in mantle olivine, since Si is the slowest diffusing species in silicates and its diffusivity is highly dependent on temperature (Chakraborty, 2010). Yet, dislocations in olivine can only glide in three independent slip systems ([1 0 0](0 1 0), [0 0 1](0 1 0) or [1 0 0](0 1 0), [0 0 1](0 1 0), [0 0 1] for steatite can only glide in three independent slip systems ([1 0 0](0 1 0), [0 0 1](0 1 0), [1 0 0](0 1 0), [0 0 1](0 1 0), [0 0 1](0 1 0)). In the case of inhomogeneous flow (i.e., flow in polycrystals), at least four independent slip systems must operate to satisfy the von Mises criterion (Mises, 1928; Hutchinson, 1977). Therefore, the currently known dislocation slip systems in olivine are not enough, and additional degrees of freedom must be provided by e.g. dislocation climb, grain boundary sliding or grain boundary migration triggered by disclinations. Disclinations (rotational topological defects) were known to exist in liquid crystals (Friedel, 1922; Frank, 1958) but were thought to be too energetically costly in crystalline solids. Nevertheless, they were recently described along the grain boundaries of copper and olivine (Beausir & Fressengeas, 2013; Cordier et al. 2014).

The slip systems in which dislocations glide in olivine are well documented, and Turner (1942) already anticipated the principal glide system in naturally occurring olivine to be [1 0 0](0 1 0). Most of the early deformation experiments at high temperature targeting slip systems were conducted by Raleigh (1965, 1967, 1968), who concluded that the dominant slip system is dependent on the temperature and the strain rate: at 10−5 s−1, glide on [0 0 1](1 0 0) dominates at < 300 °C, [0 0 1](1 1 0) between 300 and 1000 °C and finally [1 0 0](0 1 0) at temperatures of > 1000 °C (see also Young, 1969). A comparison of microstructures in naturally and experimentally deformed peridotites by Nicolas (1986) confirmed that the same slip systems are active. Such a comparison holds true for mantle rock textures (e.g., Nicolas & Poirier, 1976; Gueguen & Nicolas, 1980; Tommasi et al., 1999, 2016).

First olivine flow laws were provided by Carter and Avé Lallemant (1970) for wet and dry dunites and lherzolites, also observing dislocation climb controlled processes (polyorientation) and recrystallization at temperatures of 1000 °C and 1050 °C respectively. Carter and Avé-Lallemant attributed a weakening effect to the presence of hydrogen in the structure of olivine, expressed by a reduced activation energy of dislocation creep (335 instead of 502 kJ·mol−1). Hirh & Kohlstedt (1996), Mei & Kohlstedt (2000a,b), Demouchy et al. (2012, 2014), Fei et al. (2013), Girard et al. (2013) and Tielke et al. (2017) confirm a weakening effect by water, although the most recent results highlight that it might induce a relatively limited effect (i.e., viscosity diminution by only a factor of 10 for hydrous olivine viscosity at mantle strain rates, Tielke et al., 2017) for hydrogen concentrations reported in spinel-bearing peridotite typical of the lithospheric mantle (Demouchy and Bolfán-Casanova, 2016).

Based on high pressure deformation experiments, Karato and co-workers have proposed a classification in five types of olivine fabrics developing as a function of the applied stress and water content (Jung and Karato, 2001), but until now, the physical mechanism which could explain the crystallographic preferred orientations (CPO) variations is still lacking. Other studies recognized an effect of grain size on the strength of olivine (Goetze, 1978; Karato et al., 1986; Karato and Wu, 1993; Katayama et al., 2004). At smaller grain sizes (sub μm), dislocations are increasingly captured by grain boundaries, resulting in only few organized sub grain boundaries. In such small grains, the rate-limiting factor is dislocation glide or ionic diffusion rates, as opposed to dislocation climb in larger grains and for higher finite strains (e.g., Hiraga et al., 2011; Miyazaki et al., 2015).

In this study, we perform a series of deformation experiments at high pressure and high temperature on fine-grained polycrystalline olivine aggregates at temperatures of 1000 °C and 1200 °C. Experiments are stopped at increasing higher finite strains (0–8.6%, representing all stages of the stress-strain curve. The starting material and deformed samples are characterized by scanning electron microscopy (SEM), electron backscattered diffraction (EBSD) and transmission electron microscopy (TEM).

2. Methods

2.1. Starting material and experimental method

Olivine crystals from San Carlos (Arizona) with the following composition (Mg0.91Fe0.09Ni0.003)2SiO4 (e.g., Buening & Buseck, 1973; Frey & Prinz, 1978) are used for the deformation experiments. The fine powder was obtained by grinding olivine crystals in a fluid energy mill (Gribb & Cooper, 1998). San Carlos olivine naturally has extremely low water content of less than 1 ppm wt H2O (Mackwell et al., 1985; Mei & Kohlstedt, 2000a; Demouchy, 2010). Fine-grained powder of San Carlos olivine was cold-pressed and subsequently hot-pressed in Ni sleeves at 1200 °C and 300 MPa for 3 h, producing dense polycrystalline samples ∼ 8 mm in diameter and ∼ 16 mm long. The Ni sleeves were closed prior to hot-pressing with fitted Ni disks (200 μm thick), held in place by ∼ 2 μl of externally applied superglue. The glue is dried out before insertion into the apparatus, volatilized during hot-pressing and does not induce hydrogen incorporation into the sample. Hot-pressing, as well as axial deformation was performed in a high-pressure high-temperature internally heated, in situ deformation apparatus (also called the Paterson Press; see Paterson, 1990) using Argon as pressure medium. This deformation apparatus allows a high resolution of the applied stress (± 10 MPa) and minimal thermal gradient (± 2 °C) along a hot zone of approximately 20 mm length (Paterson, 1990). Sample volumes of ∼ 1 cm³ and diameter to length ratio of 1:2 reduce boundary effects on the texture evolution and stress distribution. Samples were deformed in axial compression at 300 ± 2 MPa confining pressure, for constant displacement rates (1.25–1.70 × 10−3 mm·s−1). Part of the displacement is taken up by elastic deformation of the apparatus. Consequently, the instantaneous strain rate of the sample changes as stress evolves during the experiment, reaching 10−8·s−1 at mechanical steady state conditions. Temperatures were kept constant during deformation. The Ni capsules containing the olivine samples were mounted with alumina and zirconia pistons inside an iron jacket (e.g., Paterson, 1990; Mei & Kohlstedt, 2000a). The oxygen fugacity was theoretically controlled by the nickel capsule. However, in a recent experimental study, the fO2 in large sample volumes has been shown to decrease towards the sample center, resulting in a linear Log(fO2) ≈ −10, at a distance > 0.5 mm.
from the capsule interface (Fauly et al., 2017, their Fig. 5), which remains in line with common mantle rocks (Frost & McCammon, 2008). Temperature was manually increased at a rate of approximately 10 °C/min before using the automatic temperature controller. Quenching of the samples is achieved by a manually controlled decrease of furnace power, reducing temperature by 100 °C/min. Faster cooling might induce severe damage to the furnace. A total of ten samples were deformed at temperatures of 1000 °C and 1180–1200 °C. Experimental conditions are listed in Table 1.

To account for the load supported by the iron jacket and nickel sleeve, the flow laws reported by Frost and Ashby (1982) (Deformation mechanism maps: the plasticity and creep of metals and ceramics, 1982) were used, following a well-established procedure (Mei & Kohlstedt, 2000a; Chen et al., 2006; Demouchy et al., 2009; Hansen et al., 2011, 2012). The calculated stress was corrected for (1) the instantaneous change in cross-sectional area due to sample shortening, assuming that samples kept their cylindrical shape and constant volume and (2) apparatus stiffness. The stiffness of the deformation apparatus is determined in a separate experiment to be 82.5 kN/mm.

Deformed olivine samples were recovered by cutting the assembly with a low speed saw. Thin sections of the deformed olivine and the hot-pressed samples were cut parallel to the deformation axis (long axis of the cylinder) after removal of 2 mm from the lower- and topmost parts, where stresses are expected to be heterogeneously distributed (Griggs et al., 1960). The heterogeneous stress distribution at the sample ends is caused by friction at the alumina spacer-sample interface, leading to a lower increase in sample radius during shortening compared to the center of the sample (i.e., barreling effect, see Moosbrugger, 2002). Thus the lower- and topmost parts were not mapped by EBSD. For some of the cold-pressed and hot-pressed samples (CMT16-1, CMT16-5, CMT16-7), sections were cut perpendicular to the long axis from the ends of the sample cylinders, to preserve parts of the sample for later deformation.

2.2. Scanning electron microscope based electron backscatter diffraction

Table 1

<table>
<thead>
<tr>
<th>Deformation at 1000 °C</th>
<th>Deformation at 1200 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMT16-6</td>
<td>CMT16-2</td>
</tr>
<tr>
<td>Temperature [°C]</td>
<td>1000</td>
</tr>
<tr>
<td>Duration [min]</td>
<td>11</td>
</tr>
<tr>
<td>Initial length L0 [mm]</td>
<td>13.50</td>
</tr>
<tr>
<td>Strain rate ϵ [s−1]</td>
<td>1.9 × 10−6</td>
</tr>
<tr>
<td>Finite strain ε</td>
<td>0.11</td>
</tr>
<tr>
<td>Diff stress σmax [MPa]</td>
<td>80</td>
</tr>
</tbody>
</table>

2.3. Transmission electron microscopy (TEM)

Transmission electron microscopy (TEM) was used to characterize the microstructures after the deformation experiments. Doubly polished thin sections (30 µm thick) of all deformed samples and hot-pressed sample CMT16-7 were prepared. The sample sections were cut parallel to the deformation axis. They were glued on a grid (Cu or Mo) and ion milled at 5 kV under a low beam angle of 15° until electron transparency was reached. The foils were subsequently covered with a thin layer of carbon. TEM observations were carried out at the University of Lille (France) using a Philips CM30 microscope operating at 300 kV and a FEI® Tecnai G2-20 twin microscope operating at 200 kV.

3. Results

3.1. Mechanical results

The results of the deformation experiments are reported in Table 1 and all stress-strain curves are shown in Fig. 1. Optical microscopy of sample thin sections shows cracks that are sub-parallel to the long axis of the sample cylinder (the compression axis). There is no relative displacement along the cracks or shearing visible on the iron-jacket.
Table 2
Statistical parameters obtained from EBSD maps using MTEX. All values derived from data treatment are given as means, except for max aspect ratio and max grain size. No. EBSD maps gives the number of large scale (global) maps free of artifacts used for the parameter calculation.

<table>
<thead>
<tr>
<th>Powder</th>
<th>Cold-press</th>
<th>Hot-press</th>
<th>Deformation at 1000 °C</th>
<th>Deformation at 1200 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMT16-1</td>
<td>CMT16-5</td>
<td>CMT16-7</td>
<td>CMT17-3</td>
<td>CMT16-6</td>
</tr>
<tr>
<td>No. of EBSD maps</td>
<td>3</td>
<td>3</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Step size [µm]</td>
<td>1</td>
<td>1</td>
<td>0.2</td>
<td>0.2</td>
</tr>
<tr>
<td>No. grains</td>
<td>13,240</td>
<td>45,918</td>
<td>2483</td>
<td>5431</td>
</tr>
<tr>
<td>Indexation rate [%]</td>
<td>28</td>
<td>41.3</td>
<td>97.6</td>
<td>93.8</td>
</tr>
<tr>
<td>Angular deviation [°]</td>
<td>0.85</td>
<td>0.71</td>
<td>0.42</td>
<td>0.46</td>
</tr>
<tr>
<td>Angular deviation (1σ)</td>
<td>(0.38)</td>
<td>(0.34)</td>
<td>(0.16)</td>
<td>(0.19)</td>
</tr>
<tr>
<td>Grain size [µm]</td>
<td>2.1</td>
<td>2.3</td>
<td>1.8</td>
<td>1.4</td>
</tr>
<tr>
<td>Grain size (1σ)</td>
<td>(0.38)</td>
<td>(0.34)</td>
<td>(0.16)</td>
<td>(0.19)</td>
</tr>
<tr>
<td>Max grain size</td>
<td>21.9</td>
<td>23.3</td>
<td>19.8</td>
<td>19</td>
</tr>
<tr>
<td>Area norm. grain size</td>
<td>7.4</td>
<td>7.6</td>
<td>7.8</td>
<td>7</td>
</tr>
<tr>
<td>Kernel av. Mis. [°]</td>
<td>0.81</td>
<td>0.62</td>
<td>0.27</td>
<td>0.37</td>
</tr>
<tr>
<td>Kernel av. Mis. (1σ)</td>
<td>(2.3)</td>
<td>(2.3)</td>
<td>(2.2)</td>
<td>(2.1)</td>
</tr>
<tr>
<td>GOS [°]</td>
<td>0.69</td>
<td>0.65</td>
<td>0.49</td>
<td>0.76</td>
</tr>
<tr>
<td>GOS (1σ)</td>
<td>(3.3)</td>
<td>(2.6)</td>
<td>(3.5)</td>
<td>(3.0)</td>
</tr>
<tr>
<td>PARSi [°]</td>
<td>19</td>
<td>21</td>
<td>42</td>
<td>35</td>
</tr>
<tr>
<td>Sub GB min. [°]</td>
<td>2.4</td>
<td>2.4</td>
<td>3.2</td>
<td>3</td>
</tr>
<tr>
<td>Sub GB min. (1σ)</td>
<td>(1.2)</td>
<td>(1.2)</td>
<td>(1.5)</td>
<td>(1.4)</td>
</tr>
<tr>
<td>J-index 1ppg</td>
<td>1.6</td>
<td>1.6</td>
<td>1.8</td>
<td>1.6</td>
</tr>
<tr>
<td>J-index 1ppg (1σ)</td>
<td>(3.2)</td>
<td>(2.7)</td>
<td>(2.7)</td>
<td>(2.5)</td>
</tr>
<tr>
<td>BAn-index 1ppg</td>
<td>0.2</td>
<td>0.2</td>
<td>0.1</td>
<td>0.5</td>
</tr>
<tr>
<td>Mis2Mean [°]</td>
<td>1</td>
<td>1</td>
<td>3.2</td>
<td>2.7</td>
</tr>
<tr>
<td>Mis2Mean (1σ)</td>
<td>(0.9)</td>
<td>(1.2)</td>
<td>(3.1)</td>
<td>(2.6)</td>
</tr>
<tr>
<td>Aspect ratio</td>
<td>1.4</td>
<td>1.4</td>
<td>1.5</td>
<td>1.5</td>
</tr>
<tr>
<td>Aspect ratio (1σ)</td>
<td>(0.4)</td>
<td>(0.3)</td>
<td>(0.3)</td>
<td>(0.4)</td>
</tr>
<tr>
<td>Max. aspect ratio</td>
<td>4.2</td>
<td>7.7</td>
<td>5.3</td>
<td>5.9</td>
</tr>
</tbody>
</table>

*a* The grain size reported is the geometric mean of the log-normal distribution of the grain equivalent diameter (diameter of a sphere with equivalent area of the respective grain).

*b* Geometric mean of the log-normal distribution of the Kernel Average Misorientation (proxy for density of Geometrically Necessary Dislocations), 2nd order, threshold of 5°.

*c* The grain orientation spread (GOS) is the average deviation in orientation between each point in a grain and the average orientation of the respective grain.

*d* Percentile average relative indented surface boundary, measure of the curvature of grain boundaries. A high PARSi indicates highly curved or indented boundaries.

*e* Subgrain boundaries are defined as 2°–10° misorientation. Subgrain boundary angles increase with polygonization and decrease with subgrain growth.

*f* One point per pixel (1ppp) uses every map pixel with an olivine orientation for the calculation of the texture j-index or BA-index. One point per grain (1ppg) uses the mean orientation of each grain.

*g* Mis2mean gives the misorientation angles from the mean within each respective grain. The given value is the mean over the whole EBSD map.
The permanent finite strains range from 0.1 to 8.6% and the strain rates range from $1.8 \times 10^{-6}\text{ s}^{-1}$ for low finite strain (< 0.5% of strain) to $1 \times 10^{-5}\text{ s}^{-1}$ at high finite strain (> 4% of strain). At 1000 °C, the stress–strain curve departs systematically from linear (elastic) behavior, even for strains below 0.5%. The stress reaches a maximum of 1127 MPa at 3% of strain. From the mechanical data, there is no evidence of plastic deformation and hence no sign of stick-and-slip or brittle failure of the samples.

At the same temperature and very similar displacement rates, the stress–strain curves differ slightly. As an example, samples CMT17-2 and CMT17-4, both deformed at 1000 °C, reach 954 and 787 MPa of stress respectively at 1.6% of strain. Samples CMT16-9 and CMT16-13, both deformed at 1180 °C, reach 265 and 313 MPa of stress respectively at 1.6% of strain. No perfect steady state is reached; stress continues to increase at a very slow rate of 4 MPa per percent of strain. From the mechanical data, there is no evidence of stick-and-slip or brittle failure of the samples.

The Young’s modulus $E$ of San Carlos olivine was calculated assuming the isotropic relation

$$E = 2G(1 + \nu)$$

where $G$ is the shear modulus and $\nu$ is Poisson’s ratio. The shear modulus $G$ was calculated for the respective temperatures and pressures using an initial shear modulus $G_0$ of 77.4 GPa, a pressure dependence $G'/G_0$ of 1.61 and a temperature dependence $\delta G'/\delta T$ of $-0.0130\text{ GPa/K}$ (Liu et al., 2005). By comparison with the deformation curves, neither at 1000 °C nor at 1200 °C do the curves exhibit a linear segment indicating pure elastic deformation. To determine if the deformation is mostly the result of anelasticity (stress and time dependence of elastic strain), we have subjected an already deformed sample (CMT17-4, 1000 °C, 1.6% strain) to axial deformation. The pre-deformed sample reaches significantly higher stresses for the same values of finite strain, as displayed in Supplementary Fig. S2. This suggests, that even at low values of finite strain, there is significant plastic deformation and hence hardening.

### 3.2. Microstructure results

Forward scattered electron images of the starting material and representative deformed samples are shown in Fig. 2 and representative maps of the misorientation to a reference point and to the mean of each respective grain are shown in Fig. 3. Data for samples not shown can be given upon request. Olivine comprises > 99.8% area of all samples, with diopside, enstatite and chromite as secondary phases (< 0.1% area). Indexation rates range from 93.3% to 98.7% in the hot-pressed or deformed samples and 27.9% to 41.3% in the cold-pressed and powder samples. Grains in all samples are tabular, shorter in the compression direction, with a mean aspect ratio of 1.4–1.5 ($d_{\text{max}}/d_{\text{min}}$) and a maximum aspect ratio of 3.0 (Table 1, Fig. 4a). The long axis of the grain is preferentially oriented at 180° to the compression axis (or vertical axis in case of the olivine powder) in all samples, as shown in Fig. 5. Grain boundaries, and part of the sub-grain boundaries, are curved or serrated. Grain internal misorientation occurs in all stages from gradual misorientation to well developed polygonization. Triple junctions are dominant, but apparent 4-grain junctions can be seen in several EBSD maps as well. BSE images reveal occasional cracks inside the grains that are oriented at 60° to each other. These rare cracks are already present in the olivine powder and there is no sign of displacement or sliding along them. Sample CMT17-2, deformed at 1000 °C to a finite strain of 7.3% exhibits a higher density of cracks crossing several grains or on grain boundaries, indicating that CMT17-2 deformed by micro-cracking after reaching a peak stress of 1073 MPa. See Supplementary Fig. S3 for SEM images of sample CMT17-2 showing grain-interior cracks and cracks at an angle of ~45° to the compression axis. Porosity from plucking during sample polishing is higher in samples with shorter hot-press durations or lower hot-press temperatures (e.g. CMT16-5 and corresponding deformed sample CMT16-6). Excluding plug outs, porosity makes up less than 3% area as observed for similar experiments (Beeman & Kohlstedt, 1993; Hirth & Kohlstedt, 1995; Hansen et al., 2012).

Representative grain size distributions obtained using MTEx as the diameter of a circle with an area equivalent to the respective grain are shown in Fig. 6. The grain size distribution is similar for all samples, there is no significant change as a function of finite strain or temperature as shown by Fig. 4b. The mean equivalent diameter calculated from the log-normal distribution ranges from 1.4 ± 0.5 μm to 1.7 ± 2.0 μm (geometric mean, 1.0) for the deformed samples and their respective hot-presses. The high geometric standard deviation reflects the broad distribution of grain sizes. The microstructure of the hot-pressed and deformed samples are identical to the one of the powder and cold-press samples, taking into account measurement errors and higher EBSD step sizes in the powder and cold-press. It is likely inherited from the original powder and shape preferred orientation due to sedimentation effects while loading the capsule. The grain shape is likely related to perfect cleavage of olivine on (0 1 0) (Deer et al., 1997; Hawkes, 1946; Velinskiy & Pinus, 1969; Kuroda & Shimoda, 1967; Kuijper, 1979) during pulverization.
Accounting for sectioning bias (factor of 1.5 for nonspherical olivine grains; Underwood, 1970), this equals a true mean grain diameter of 2.1–2.6 µm. Normalized by area, the distribution broadens with grains up to 27 µm in equivalent diameter taking up a sizeable fraction of the sample area. Due to the calculation method (i.e., only considering grains consisting of more than 5 pixels), the smallest grains have a equivalent diameter of 0.6 µm. For the cold-press and powder samples, a higher EBSD step size of 1 µm was used. Consequently, the average grain size appears to be higher, with 2.1 ± 1.9 µm for the olivine powder and 2.3 ± 1.9 µm for the cold-press (Table 1).

All samples, including the undeformed olivine powder, show an axial [010] pattern of the texture, as shown in Fig. 7. Crystal [100] and [001] axes are aligned in a girdle perpendicular to the compression axis, and [010] axes are parallel to the compression axis. The J-index (area weighted) ranges from 1.2 to 1.8, and is not changing significantly in between samples. Calculated for one mean orientation per grain, the J-index is slightly lower, ranging from 1.1 to 1.2.

The frequency of grain boundaries as a function of their misorientation angle is displayed in Fig. 8. The given histogram (CMT16-9) is representative of all samples. We show the correlated misorientation angles (i.e., from neighboring pixels). In MTEX, the uncorrelated misorientation angle distribution is calculated using the orientation distribution function (ODF) harmonic coefficients and only reflects the influence of the CPO and nothing else. The uniform misorientation angle distribution is calculated using the analytical method of Morawiec (2004). Note that a deviation of the uncorrelated angles from the uniform distribution therefore shows the influence of texture, while the distribution of the correlated angles depends on the occurrence of deformation-related (sub)grain boundaries. The most commonly occurring low angle boundaries have misorientations of up to 20°, with a strong maximum at 10° (10% frequency). High angle boundaries increase in frequency towards a maximum at 90° (8% frequency), resulting in a negatively skewed distribution. The uncorrelated misorientation angles match the uniform distribution, with only a higher frequency of angles below ~70° than in the random distribution. The misorientation from the mean within each grain is log-normal distributed with angles up to 15° as plotted in Fig. 9. The misorientation axis of these low-angle boundaries of up to 15° can be plotted in crystallographic coordinates. The resulting inverse pole figure,
representative for the hot-pressed and deformed samples, is displayed in Fig. 10. Most misorientation axes are close to the crystallographic [0 0 1] direction with a strong maximum of 8× uniform. There is a weak secondary concentration around the [0 1 0] direction. The distribution of the misorientation axes does not change significantly between samples deformed at 1000 °C and 1200 °C or as a function of finite strain.

3.3. Transmission electron microscopy

Typical weak-beam dark-field TEM images of the dislocation microstructures are given in Fig. 11. All samples exhibit unambiguous dislocation activity. The most striking feature is the rather strong heterogeneity of the microstructures in all samples which does not allow a simple trend to be found among them. Even at the lowest temperatures considered (1000–1030 °C) both [1 0 0] and [0 0 1] dislocations activity is found (Fig. 11a and b) from quite small finite strains (1.41% in case of CMT16-2). Dislocation line shapes suggest high lattice friction, but dislocations already organize into subgrain boundaries. Even at the highest temperature considered (1200 °C) deformation is very heterogenous with some grains almost or completely pristine (Fig. 11c) and some other exhibiting dislocation densities of the order of 10^{14} m^{-2}. Several observations of grain boundaries acting as sources for dislocations have been found. The samples contain residual pores which do not appear to be sites where plasticity concentrates. In grains displaying evidence of [1 0 0] and [0 0 1] slip activity, there is no evidence of interaction between the dislocations from the different slip systems.

4. Discussion

4.1. Mechanical data

The maximum differential stresses of our experiments at 1000 °C exceed the confining pressure by a factor of up to 3.5. The confining pressure (P) acts by opposing volume expansion (e.g. Edmond & Paterson, 1972) and against crack nucleation and propagation. The Goetze criterion, first proposed by Briegel and Goetze (1978), defines a critical stress needed to initiate crack nucleation as \( \sigma_1 - \sigma_3 = P \) (see also Kirby, 1980; Evans & Kohlstedt, 1995). With the exception of CMT17-2, there is no evidence for displacement along cracks in SEM images of our samples. On the short time scales of our experiments, the Goetze criterion does not hold true.

For deformation experiments at 1000 °C, the present stress–strain curves are consistent when compared to previous mechanical results for polycrystalline olivine (900 °C, 300 MPa, 1 × 10^{-5}s^{-1}–3 × 10^{-5}s^{-1}, axial compression, with correction for the apparatus stiffness) from Demouchy et al. (2014). The sample CMT17-2 of this study reaches brittle failure at a higher stress of 1127 MPa, while sample PoEM22 of Demouchy et al. (2014) fails at 930 MPa at 900 °C. Note that both CMT17-2 of this study and PoEM22 of Demouchy et al. (2014) did not reach steady state before brittle failure and thus give only a lower bound of strength. For a given strain, samples in this study reach a lower stress, consistent with deformation at higher temperatures. The maximum stresses from our experiments at 1000 °C and existing low temperature (exponential) flow laws are shown in Fig. 12c. The maximum stress for 1.1% of finite strain at 1000 °C (CMT16-8, no steady state or brittle failure) surpasses recent calculations based on 2.5-D dislocation dynamics models for single crystals of olivine oriented for Fig. 4.
easy slip (Boioli et al. 2015; their Fig. 3). In this study, they model a single glide direction, [1 0 0](0 0 1), in a olivine single crystal, therefore giving only a lower bound of the strength. The experiment CMT17-2 reached brittle failure around 3% of finite strain, but the maximum stress before brittle failure of 1127 MPa can be used. The stress surpasses steady state estimations from the single crystal flow laws from Evans and Goetze (1979) and Demouchy et al. (2013), but is in good agreement with the flow law based only on [0 0 1](1 0 0) glide from deformation experiments in shear from Tielke et al. (2016), as seen in Fig. 12. The higher strength of our samples can at least partly be attributed to the lack of sufficient independent slip systems in polycrystalline olivine samples (the so called olivine paradox) and the difference between single crystal and polycrystal rheology (e.g., Evans and Goetze, 1979, their Fig. 14).

The deformation curves of samples CMT16-13 and CMT16-9 obtained at 1200 °C (Fig. 12b) show a quasi-steady-state mechanical behavior. Their respective maximum stresses fit well with the dislocation creep flow law for Åheim dunite from Keefner et al. (2011, grain size of 0.9 mm). The dislocation power law of Hirth & Kohlstedt (2003, using data from Mei and Kohlstedt, 2000b for dry conditions) and the dislocation creep flow law for dry dunites from Chopra and Paterson (1984) surpass the stresses reached in our samples deformed at high temperature by 100 MPa and 200 MPa respectively.

Fig. 6. Representative grain size distribution of (a) a hot-pressed sample (CMT16-5), and (b) a sample deformed at 1200 °C to 8.6% of strain. Grain sizes are calculated as the diameter of a circle with an equivalent area to the grain. N gives the number of grains used for the grain size calculation. The mean is calculated as the geometric mean.

Fig. 7. Lower hemisphere pole figures of the crystal preferred orientation for (a) the olivine powder; (b) a representative hot-pressed olivine sample; (c) the olivine sample deformed at 1000 °C (CMT16-8) and (d) the olivine sample with the highest finite strain deformed at 1200 °C (CMT16-9). The pole figures were calculated using one point per pixel.

Fig. 8. Representative histogram of the correlated, uncorrelated and uniform misorientation angles using a lower bound of 2°. Grey bars show boundaries in CMT 16-9. The black line is calculated for a uniform, randomly oriented sample. The grey line is calculated from uncorrelated measurement pixels (spaced apart) and can be compared to the uniform sample to estimate the texture influence.
While deformation is obviously plastic, one sample (CMT17-2) exhibits a higher density of cracks. Cracks generally exist in a subset of the grains in all samples, including the olivine powder, where they are more common and broader. They are therefore likely to be related to the powder preparation method of the olivine starting material (i.e., fluid energy pulverization). The pulverization might also explain the aspect ratio of 1.4–1.5 seen in all samples, since (010) planes are distinct cleavage planes in iron-rich olivine (Deer et al., 1997; Hawkes, 1946; Velinskiy & Pinus, 1969; Kuroda & Shimoda, 1967; Kuijper, 1979). During hot-pressing these cracks mostly healed. Sample CMT17-2 exhibits a higher density of cracks, which are paired with their acute angle (~60°) orientated nearly parallel to the compression axis; opening grain boundaries, as well as crossing several grains. We conclude that the sample CMT17-2 deformed in the brittle field by micro fracturing for a strain > 3% (at 1127 MPa, where the curve flattens abruptly), while the other samples underwent no brittle failure. Rigid body grain rotation cannot be observed, since the orientation of the long axis of the grains stays constant.

The main deformation mechanism can be identified from comparison of the mechanical data to published flow laws and the alignment of low-angle boundaries. The maximal stresses of our experiments match existing dislocation creep flow laws for temperatures > 1100 °C. Combined with the constant grain size, aspect ratio and TEM images, this suggests that dislocation creep was the dominant deformation mechanism, with a limited contribution of diffusion creep. The alignment of low-angle boundary misorientation axes parallel to the crystallographic [001] axis, shown in Fig. 10, is common for tilt walls composed of [100]hk0 dislocations, while the weak concentration parallel to the [010]axis is common for tilt walls built from [001] (hk0) dislocations (Nicolais et al. 1973; Wenk et al. 1991; Tommassi et al. 2008; Farla et al. 2011; Demouchy et al. 2013, 2014). However, the grain boundary misorientation axes obtained from EBSD maps only represent preserved geometrically necessary dislocations (GND’s, i.e., dislocations contributing to the lattice curvature). Therefore, the slip systems active during deformation might differ slightly (Raleigh, 1965, 1967, 1968; Carter and Avé Lallemant, 1970; Durham, 1975; Poirier, 1975; Tommassi et al., 2008; Soustelle et al., 2010; Demouchy et al. 2014). The distribution of grain boundary misorientation angles (Fig. 8) showing a peak at 90° is typical for a uniform distribution for orthorhombic crystal symmetry (Morawiec, 2004). The secondary peak at 10° matches well with observations from Marquardt et al. (2015), who investigated the grain boundary character distribution of undeformed synthetic and natural polycrystalline olivine.

The microstructures of the hot-pressed and deformed samples are identical. In particular, the distribution of deformation axes does not change in between hot-presses and experiments performed at 1000 °C and 1200 °C and as said above (4.2), it is due to perfect cleavage of olivine on (010) during powder preparation. The {100}(001) slip system therefore seems not to be solely or preferentially activated in samples deformed at 1000 °C, compared to deformation at 1200 °C, which is consistent with experiments of Raleigh (1968) and Phakey et al. (1972). Likewise, the crystal preferred orientation (Fig. 7) does not change or increase in intensity, which is expected for low values of finite strains (Nicholas et al., 1973; Zhang & Karato, 1995; Skemer et al., 2012), and is confirmed here, even if high differential stresses are reached. We tried to hot-press an olivine aggregate for 20 h at 1270 °C (CMT17-3) and it produced only a limited grain growth to 2.8 μm and no significant reduction of dislocation density (as evidenced by the grain orientation spread and kernel average misorientation) or a change in texture (see J- and BA-index in Table 2). To significantly alter the starting microstructure, much longer hot-press durations (>100 h), a higher hot-press temperature (>1300 °C) or an addition of water (drastic change of thermodynamic conditions) would be needed and these grain growth conditions are beyond the scope of this study.

### 4.2. Deformation mechanisms & microstructure

The microstructure of the deformed samples is characterized by a high density of sub-grain boundaries. The misorientation axes of these boundaries are distributed in within the range of 2° to 15°, showing a peak at 9°. This distribution is representative of all samples. The misorientation is log-normal distributed with angles up to 16°.

![Fig. 9. Histogram of the misorientation to the mean within each respective olivine grain (CMT16-9). This distribution is representative of all samples. The misorientation is log-normal distributed with angles up to 16°.](image9)

![Fig. 10. Representative inverse pole figure of the misorientation axes of olivine grain boundaries in crystal coordinates for misorientation angles between 2 and 15°.](image10)

### 4.3. Strain hardening during transient creep

We have two opposing observations, which question the current understanding of strain hardening in olivine. The stress during deformation rises nonlinearly as a function of finite strain before reaching steady state or brittle failure. This hardening is dependent on the deformation temperature and must therefore be based on a thermally activated process. Based on observations on metals, hardening is explained by competing effect of reduction of the grain size, and an increasing dislocation density in the deformed sub-grains acting as obstacles and yielding an increasing sub-grain boundary misorientation (Estrin et al., 1998; Nes, 1997). Yet, the microstructure of our samples does not change with increasing stress or finite strain. Since none of these mechanisms seem to hold true during transient creep of our samples, the question of the mechanism of strain hardening in our sample remains. Thus, we discuss here several possibilities:

(i) **Dislocation interactions**: In metals, plastic strain hardening is commonly interpreted as the consequence of gliding dislocations interacting with the so-called forest dislocations (i.e. dislocations lying in different planes). This effect is reinforced by the possibility in most cubic metals that dislocation interaction can lead to stable...
junctions. In olivine, the two Burgers vectors [1 0 0] and [0 0 1] are perpendicular, which allows only rare and weak junctions (Mussi et al. 2017). Indeed, Fig. 11a and b shows no strong interactions between [1 0 0] and [0 0 1] glide. Other hardening mechanisms resulting from dislocation dynamics are collinear interaction and double cross slip. Our observations although preliminary, do not show more of these features in the low temperature samples.

(ii) Dislocation Mobility: In this temperature range, olivine is expected to involve significant lattice friction and this is supported by observation of the shapes of dislocation lines. However, here again, qualitative indication of lattice friction can be found in all samples (e.g. Fig. 11b and d). It is to be remembered that in the thermally activated regime, temperature enters an exponential and hence that a difference of 200 °C can lead to significant enhancement of dislocation mobility which could not be detected from post-mortem observations. This contribution can therefore not be excluded.

(iii) Grain boundaries acting as obstacles: In low-temperature plasticity, grain boundaries play mostly the role of obstacles leading to the well-known Hall-Petch effect (Hansen, 2004; Louchet et al., 2006). With grain sizes of around 2 μm, the ratio of boundary area to grain volume is high. Therefore, the interaction of dislocations with grain boundaries might play a role by preventing easy transfer of plastic activity from grain to grain. The formation of pile-ups might cause sufficient back stresses to be a source of hardening. However, we did not observe more evidence for pile-ups in samples deformed at the lowest temperatures than at the highest temperature.

(iv) Grain boundaries acting as dislocation sources. For plastic deformation to proceed, dislocation multiplication mechanisms are necessary. Inside grains, multiplication is usually attributed to the operation of Frank-Read sources (Frank & Read, 1950), although they are rarely clearly observed. Our study follows this trend since we did not observe such sources. Instead, we observed several occurrences of dislocations being emitted from grain boundaries. We have no information about the ability of grain boundaries in olivine to act as dislocation sources and how temperature influences this mechanism. We note that the sources observed only produce a few loops. The hypothesis that dislocation nucleation represents a limiting factor in this temperature range cannot be further established or discarded.

(v) Grain boundaries acting as agent of deformation. Dislocation activity might not be the only deformation mechanism acting and in the recent years, a growing body of evidence has lead to establish grain boundary processes as a contribution to creep of olivine aggregates. This is most often attributed to the so-called grain boundary sliding mechanism (i.e., relative tangential motion of adjacent grains parallel to their common boundary; e.g. Hirth & Kohlstedt, 1995; Langdon, 2006; Hansen et al., 2011) although other mechanisms described as shear-induced boundary migration, possibly involving disclinations, have also been proposed (Cordier et al., 2014). Grain boundary sliding was proposed to be active under high stresses and moderate to low homologous temperatures in olivine (T_m of Fo90 = 1765 °C, Bowen and Shairer, 1935; Wang, 2016) but also at temperatures very close to melting temperature and for a restricted type of tilt boundaries (e.g., in gold, Cahn et al., 2006). Without suitable markers (i.e., grid or dot-array of
deposited metal or electron engraving, Quintanilla-Terminel et al., 2017), we cannot determine if these mechanisms were more active at 1200 °C than at lower temperatures.

4.4. Implications for the uppermost mantle

Temperature conditions in our experiments (i.e., 1000 °C) correspond to the uppermost lithospheric mantle, just below the Mohorovičić-Discontinuity. This layer acts as an imperfect mechanical boundary between the convecting mantle and the lithospheric plates. The maximum stresses obtained by our samples at 1000 °C surpass predictions from existing flow laws for this temperature domain (exponential law), but are indeed significantly lower than predictions deduced from high temperature flow laws (power law), without the intervention of additional weakening mechanism such as water weakening (e.g., Mackwell et al., 1985; Demouchy et al., 2012) or partial melting (e.g., Kohlstedt and Zimmermann, 1996). In consequence, the strength of olivine-rich rocks deformed at a temperature of 1000 °C and below, are in the order of 0.5–1 GPa when extrapolated at the strain rate of the mantle (e.g. \(10^{-14} \text{s}^{-1}\)) which matches the requirement of a not so soft lithospheric mantle, able to upkeep large orogens for millions of years (e.g., Mouthereau et al., 2013).

5. Conclusions

- Deformation at 1000 °C surpasses predictions from most previous low-temperature flow laws, but is in good agreement with Tielke et al. (2016, shear on [0 0 1](1 0 0)).
- Deformation at 1200 °C is in good agreement with Keefner et al. (2011), Tielke et al. (2016, shear on [1 0 0](0 0 1)) and Hirth and Kohlstedt (2003).
- The non-linear dependence of stress on finite strain (hardening) during early stage deformation is not correlated to a change in statistical microstructure measured by several parameters or texture strength and symmetry using EBSD maps.
- The samples with constant statistical microstructure however have a strength that is sensitive to finite strain, temperature and strain rate in a similar way to previously reported flow laws.
- The achieved stresses, constant grain sizes and absence of grain shape change fit well to dislocation creep as the main deformation mechanism. Grain boundary mechanisms would suggest a lower strength than observed.
- Glide on [1 0 0](0 k l) is dominant, as evidenced by the alignment of low-angle boundaries parallel to the crystallographic [0 0 1] direction.
- Even at early steady state (at 1200 °C) and stresses of up to...
1073 MPa, a sizeable subset of grains remains free of misorientation, which could be related to the small grain size.

- We further confirm that flow laws obtained at temperatures > 1200 °C cannot predict the strength of olivine deformed at 1000 °C. The observed microstructures cannot explain the change in viscosity.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found in the online version, at http://dx.doi.org/10.1016/j.pepi.2018.03.002.

References


