TEM investigation of dislocation microstructure of experimentally deformed silicate garnet

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Abstract

Deformation experiments have been carried out on single crystals of garnet (Py25 Al67 Sp2 Gr6) under high confining pressure (P = 6.5 GPa) and temperature (T = 1440°C) in a multi-anvil apparatus. The high pressure sample assembly was designed so as to generate high differential stress. In one experiment, the differential stress was limited by a single crystal of San Carlos olivine (oriented along [010]) added on top of the specimen. Garnet crystals have been plastically deformed which shows that under our experimental conditions, garnet is ductile. The dislocation microstructure, analysed by transmission electron microscopy (TEM), suggests that both ½(111){110} and (100){110} glide systems have been activated. Some dislocations appear to be dissociated. The observations of dislocation junctions and subgrain structures indicate that dislocation climb is enhanced under our experimental conditions.

1. Introduction

Knowledge of rheological properties of major constituents of the mantle is crucial in understanding the dynamics of our planet. Although significant efforts have been devoted to constrain rheology of olivine and pyroxenes, which are the major phases of the Earth’s upper mantle (see for instance Bai et al., 1991 and Raterron et al., 1994 and references therein), experimental data on minerals stable in the deep mantle have been lacking due to the technical difficulties in conducting deformation experiments under very high pressures. Garnets are among the most abundant minerals in the transition zone, between 410 and 660 km depths (e.g. Ringwood, 1991), and in subducted oceanic lithosphere (Irifune and Ringwood, 1987) at depths within the transition zone.

Garnets are often regarded as very strong minerals. Indeed, recent experimental studies suggest that garnets have a higher strength than major co-existing minerals (Ingrin and Madon, 1995; Karato et al., 1995). However, microstructural observations of natural garnets from various origins (Ando et al., 1993; Ji and Martignole, 1994) including very deep samples (Doukhan et al., 1994) show that under some circumstances, garnets have been really plastically deformed. The question is now whether garnet dominates or not the rheology of the garnet-rich assemblages in the transition zone. Studying the rheology
of garnets is thus of primary importance in understanding the dynamics of this region of the Earth.

The present study describes deformation experiments performed on silicate garnets with a multi-anvil high-pressure apparatus. A modified sample assembly designed to induce significant differential stress has been used. This experimental set-up does not allow quantitative strain–stress relations to be obtained. The emphasis will be put on dislocation microstructure characterization by transmission electron microscopy (TEM) and the understanding of physical processes of deformation under mantle pressure and temperature conditions.

2. Experimental procedure

2.1. Description of the samples

A natural, gem-quality, single crystal of garnet was used in the present study. Its structural formula determined by micro-analysis in the TEM (see below) is: Py25 Al67 Sp2 Gr6. A low density of small (less than 0.1 mm in dimension) fluid and solid inclusions was observed within the crystal with the optical microscope prior to deformation. Samples for both infrared measurements and deformation experiments were cut from the inclusion-free part of the crystal. The hydroxyl content of our crystal was determined, by infrared spectroscopy (in the range 3000–4000 cm⁻¹), to be 0.015 wt% H₂O. The integrated molar absorptivity used for this determination corresponds to the pyrope–almandine compositions: εᵢ = 300 l mol⁻¹ cm⁻² (Aines and Rossman, 1984).

The starting garnet crystal was oriented by using the X-ray Laue technique and then cut into 2 or 4-mm-thick slices, the plane of the cut having a normal close to one of the (110) axes. Two cylinders (2.2 mm in diameter) were cored from the slices. A single crystal of natural San Carlos olivine (Arizona, USA) was also oriented (within ±2°) by universal stage and X-ray Laue techniques. A cylinder was then cored along the [010] direction. Olivine and garnet cylinders were slightly shortened and polished (with 3-μm alumina powder) to eliminate chipped faces and provide resistance against fracturing.

Two deformation experiments were performed: one on a 4-mm-length cylinder of garnet single crystal, and the other on a composite crystal consisting of two 2-mm-length cylinders of garnet and olivine joined end to end.

2.2. Deformation experiments

The deformation experiments were performed in a 2000-t Uniaxial Split-Sphere Apparatus (USSA-2000) which has been described previously (e.g. Remsberg et al., 1988). This apparatus can achieve pressures (P) in excess of 9 GPa and temperatures (T) up to 1600°C (for several hours) in the 18-mm

![Fig. 1. Cross section of the 18/12 cell assembly used for the deformation experiments. This assembly utilizes an octahedron of edge length 18 mm which allows large volume sample experiments. The octahedron made of pyrophyllite, a good thermal insulator, is used for long run duration at high temperature. See text for further explanations.](image-url)
cell assembly, for sample volumes of a few tens of cubic millimetres. Such apparatuses are mainly used for hydrostatic pressure experiments with cell assemblies designed to minimize differential stresses, which usually do not exceed 0.4 GPa (Wang et al., 1988; Ingrin and Liebermann, 1989). However, both Liebermann and Wang (1992) and Bussod et al. (1993) showed that a multi-anvil press can also be used as a high P–T deformation apparatus. With this goal, we used the 18/12 pyrophyllite octahedron previously calibrated under pressure at room and high temperature; in order to strongly increase the differential stress within the cell assembly, we placed two non-machinable alumina rods, acting as pistons, on both ends of the sample.

Fig. 1 shows a cross-section of the high-pressure cell used (after modification): the cylindrical sample (garnet single crystal, or garnet–olivine composite crystal) is fitted into a soft iron capsule which mechanically seals during pressure increase at room temperature. The role of the Fe-capsule is to reduce the temperature gradient within the sample, and to chemically isolate the sample from any contamination. Iron is also a relatively soft material with a much lower mechanical resistance compared with alumina, thus preventing fracturing of the sample during cold compression. We also followed the recommendation of Bertran-Alvarez et al. (1992) who showed that the use of Fe-capsules for high T–P experiments is an efficient method to maintain ferrous iron-bearing minerals within their redox stability fields. The Fe-capsule is lined by MgO which isolates the graphite heater from Fe metal and provides good thermal conduction. The external sleeve of the cell is made of zirconia which provides good thermal isolation from the outside, as well as the pyrophyllite octahedron surrounding the cell.

The P–T conditions of the two experiments performed are given in Table 1. A typical P–T path followed during the runs is illustrated in Fig. 2. Despite P–T conditions near the graphite–diamond transition (see for instance Liu and Bassett, 1986), no characteristic decrease of the heater resistance was noticeable, even after 2 h at the peak temperature. No thermocouple was used during the run, as introduction of thermocouples increases the risk of producing fracturing in the sample during compression. The temperature was deduced from the measurement of the power (W) supplied to the heater and by using the calibration curves \( T = f(W) \) previously established at 6 GPa for this cell assembly. The uncertainty reflects reproducibility from run to run (J. Zhang and P. Raterron, personal communication, 1994). Zhang and Raterron also measured the thermal gradient along the assembly axis; from their measurement, we estimate that the temperature at the extremities of the sample at annealing conditions was about 140°C lower than the temperature at the centre of the sample. Taking into account both the temperature uncertainty and gradient we estimate that the temperature of any part of the sample at annealing conditions ranges from 1250 to 1490°C. At the end of the runs, the sample was quenched to room temperature, and the pressure slowly (for about 17 h) decreased to 1 atm in order to prevent fracturing of the sample during depressurization.

The differential stress and strain rate during the annealing are difficult to estimate in a multi-anvil

![Fig. 2. Experimental procedure for a multi-anvil deformation experiment. Pressure versus time (solid line) and temperature versus time (broken line). See text for details.](image-url)
apparatus. Bussod et al. (1993) monitored the relative displacement of the apparatus guide blocks in order to deduce strain rate for the sample, and estimated the applied stress using grain size piezometry on the deformed samples. The purpose of our experiments was not to deduce any rheological law, but to characterize by TEM the active slip systems of silicate garnet; the applied stress and strain rate during the runs were not critical parameters. Run #2083 was performed on garnet single crystal alone and, as reported below, we observed that the sample #2083 deformed plastically during this experiment. Run #2087 was thus performed on a garnet–olivine composite crystal, in order to limit the deviatoric stress in the assembly to the flow stress of olivine (stressed along one of its hard orientations: [010]). When recovered, both the garnet and the olivine crystals were plastically deformed.

2.3. Transmission electron microscopy

Thin sections of the samples (~20 μm) were prepared by mechanical grinding and then optically polished on both faces with cerium oxide. Electron transparent foils were finally obtained by ion milling at 5 kV under a low beam angle of 15°. TEM observations were carried out at 300 kV with a
diffraction conditions difficult to obtain. It is thus
delicate to use the conventional method of character-
isation based on the \( g \cdot b = 0 \) and \( g \cdot b \times u = 0 \) invis-
bility criteria. Indeed, Ando et al. (1993) chose
another method based on the Burgers circuit. Lattice
fringes were counted on high resolution micrographs
along a closed circuit enclosing the dislocation core.
This method based on high resolution micrographs
suffers, however, some limitation as it applies only
to the special case where dislocations are viewed
edge-on. Moreover, the Burgers vectors must be in
the plane of the micrographs, otherwise the full
Burgers vector cannot be determined. In the present
study we have analysed the dislocations in garnet by
large angle convergent beam electron diffraction:
LACBED (Cherns and Preston, 1986; Tanaka et al.,
1988; Cherns and Morniroli, 1994). LACBED has
been applied recently in mineralogy to analyse dislo-
cations in quartz (Cordier et al., 1995). In LACBED
mode, when a dislocation intersects a Bragg line (i.e.
the line where the exact Bragg condition is satisfied)
with the diffraction vector \( g \), the line splits into \( n \)
nodes if \( g \cdot b = n \) (Cherns and Preston rule). If the
dislocation line is long enough, then it is possible to
place it with respect to the Bragg lines so that it
crosses at least three Bragg lines. Three linear equa-

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Fig. 6. Run #2083. Relaxed subgrain boundaries. (a) One disloca-
tion family, pure tilt boundaries. (b) Two dislocation families,
pure twist boundary. Weak-beam dark field micrographs. \( g \): 400.

Philips CM30 microscope. Microanalyses in the TEM
were performed with a Tracor energy dispersive
X-ray attachment equipped with a Ge detector and
an ultra-thin window allowing light elements such as
oxygen to be detected.

As already noticed by several authors, characteri-
zation of dislocation Burgers vectors is difficult in
the garnet structure (Rabier et al., 1976; Ando et al.,
1993; Doukhan et al., 1994). The reason is the large
unit cell which renders satisfactory two-beam

Fig. 7. Run #2083. Dissociated dislocation. Weak-beam dark
field micrograph. \( g \): 400.
Fig. 8. (a) Experimental LACBED pattern close to the [103] zone axis showing the Bragg lines used for dislocation analysis. (b) Corresponding theoretical pattern calculated with a [278 31 820] orientation.
tions of the type \( g_i, b = n_i \) are obtained whose solution uniquely gives the Burgers vector. If the dislocation is too short, then it is successively placed on three Bragg lines and the Burgers vector is deduced as above. This method requires knowledge of:

- \( n \), the number of intensity minima in a dark field pattern,
- the direction of the dislocation line characterized by a unit vector \( u \),
- the \( hkl \) indices of the \( g \) Bragg line,
- the direction of the deviation parameter \( s \).

These two last features are identified by comparisons with theoretical patterns drawn by means of a computer program, based on the kinematical theory, which was especially developed for this application.

3. Microscopic observations

3.1. Garnet samples

The dislocation microstructures are very similar in samples from both \#2083 and \#2087 experiments. They will thus be described together. The dislocations distribution is rather heterogeneous in both samples. Some areas have not been deformed and are completely pristine. Some large areas, on the contrary, exhibit numerous dislocations. Dislocation densities between \( 10^{12} \) and \( 10^{13} \) \( \text{m}^{-2} \) are routinely observed. Careful examination of the microstructure shows that different dislocation patterns can be distinguished. Some dislocations exhibit straight segments aligned along specific crystallographic orientations (Fig. 3): they are clearly confined in their glide plane. However, in most cases, the dislocation lines are curved and do not seem to belong to any specific plane. The dislocation lines are entangled and they often form junctions (Fig. 4). At some places, one can see dislocations which interact with each other and arrange themselves in subgrain boundaries to minimize total elastic energy of the crystal (Fig. 5). Fig. 6 shows a more advanced stage of recovery with relaxed subgrain boundaries. Such well-organized subgrain boundaries are common in our specimens. Careful observation in weak-beam dark field (WBDF) shows that some free dislocations are dissociated (Fig. 7, also see Discussion).

Dislocation Burgers vectors have been analysed by LACBED. Diffraction experiments have been carried out close to the \([103]\) zone axis (Fig. 8). More than 40 dislocations have been fully characterized. Two-thirds of them are of the \( \frac{1}{2} \langle 111 \rangle \) type (Fig. 9(a)) and the rest are of the \( \langle 100 \rangle \) type (Fig. 9(b)). More precisely, we have found \( \frac{1}{2}[111], \frac{1}{2}[111], \)
The microstructure is very heterogeneous in olivine too. Some regions are free of dislocations. Most of the crystal exhibits numerous free dislocations with densities ranging from $10^{10}$ to $5.10^{13}$ m$^{-2}$. Dislocation Burgers vectors have been analysed by means of the usual invisibility criteria. One finds $b = [100]$ and $[001]$. Straight screw $[100]$ dislocation segments appear to be confined in the (010) plane. $[001]$ dislocations have less preferential orientations. In dislocation-rich areas, dislocations form numerous subgrain boundaries. In some highly deformed zones, polygonization and recrystallization are well developed.

4. Discussion

The Burgers vectors of the stable dislocations in bcc structures are $\frac{1}{2} \langle 111 \rangle$ and $\langle 100 \rangle$. These two types of dislocations have already been found in deformed garnets (Rabier et al., 1976; Garem et al., 1982; Ando et al., 1993; Doukhan et al., 1994). However, $\langle 100 \rangle$ dislocations may result from the attractive reaction of $\frac{1}{2} \langle 111 \rangle$ dislocations following the reaction:

$$\frac{1}{2} [111] + \frac{1}{2} [\overline{1} \overline{1} \overline{1}] \rightarrow [001].$$

Indeed, $\langle 100 \rangle$ dislocations identified so far occur as short segments in junctions (Rabier et al., 1976; Garem et al., 1982; Doukhan et al., 1994) suggesting that $\langle 100 \rangle$ slip may not be significantly activated. In contrast, Ando et al. (1993) found that $\langle 100 \rangle$ slip is a dominant deformation mode in natural garnets from peridotites.

A significant part of the dislocations analysed in the present study is of the $\langle 100 \rangle$ type. Does this mean that $\langle 100 \rangle$ slip has been activated? In LACBED mode, dislocations are mainly detected through their effects on the Bragg lines. In principle, a shadow image is superimposed on the LACBED pattern, due to the defocus in the LACBED mode. In our diffraction experiments, the shadow contrast of dislocations was extremely faint and did not allow an easy control of the microstructure during characterization of dislocation Burgers vectors. It is thus difficult to guarantee that $\langle 100 \rangle$ dislocations characterized are not junction products. However, characterization of short dislocation segments in junctions is delicate work and has not been performed and is very unlikely to occur by chance. Moreover, it appeared during LACBED analysis that some regions of the crystal was dominated by $\langle 100 \rangle$ dislocations while some others contained mostly $\frac{1}{2} \langle 111 \rangle$ type. Altogether, these elements suggest that $\langle 100 \rangle$ slip has been activated in our experiments.

The lattice parameter of garnets is rather large resulting in large Burgers vectors. In the present case, $|b|_{\langle 1/2 \langle 111 \rangle \rangle} = 0.91$ nm and $|b|_{\langle 100 \rangle} = 1.05$ nm. The dislocations energy which is proportional to $\mu b^2$ ($\mu$ is the shear modulus) is then large compared with most other minerals and one expects dislocation dissociation to occur in garnets, reducing the core energy. WBDF imaging is a suitable technique to investigate dissociation of dislocations because it produces much sharper images than conventional
bright- or dark-field imaging (which corresponds to a strongly diffracted beam with a small Bragg error). Under optimum conditions, WBDF can reach resolution levels in the 2–5 nm range (Sarikaya and Howe, 1992). In the case of garnets, these levels are difficult to reach. The diffraction vectors which are suitable for dislocation imaging are 400, 420 or 640 (their extinction distances, in almandine, are 135, 160 and 201 nm, respectively). The use of such diffraction vectors corresponds frequently to high $g \cdot b$ products (3 or 4). In such cases, the dislocation image is broad, the intensity diffracted being approximately proportional to $(g \cdot b)^2$. Intensity profiles calculated using the kinematical theory show that for $g \cdot b = 3$ and 4, a double image is produced (Hirsch et al., 1977) which is usually seen on bright- or dark-field images (close to the Bragg conditions). However, both images have markedly different intensities. The dislocation presented in Fig. 7 has been imaged by WBDF (with a large Bragg error); it exhibits two fine bright lines with similar intensities. This contrast cannot be attributed to a double image and those lines would thus correspond to two partial dislocations. The dissociation widths observed are in the range 10–15 nm. The imaging conditions of Fig. 7 correspond to a very low intensity which makes direct observation of these dislocations with the microscope very difficult. As a consequence, it has not been possible yet to perform LACBED determinations and WBDF observations on the same dissociated dislocation. Up until now, we do not know which dislocations ($\frac{1}{2}[111]$ or $[100]$) are dissociated. The only dissociated dislocations that have been reported so far are not mobile dislocations. Doukhan et al. (1994) found dissociated $[110]$ dislocations in subgrain boundaries. According to these authors, these dislocations were the product of junctions and rearrangements within the subgrain boundary. Allen et al. (1987) observed the dissociation of a $\frac{1}{2}[111]$ dislocation into two $\frac{1}{2}[111]$ partials which was interpreted as a grown-in defect. Moreover, no dissociation was reported by Ando et al. (1993) from their high resolution TEM analysis of mobile dislocations. More work is definitely needed to assess the fine core structure of mobile dislocations in garnets.

From a mechanical point of view, the control of the deformation parameters is still very crude in our experiments. During pressurization, the specimen is subjected to high non-hydrostatic stresses. While the specimen is heated, plastic deformation occurs to relax these stresses in the specimen as well as in the surrounding materials of the assembly. Our deformation experiments are thus stress relaxation tests with a differential stress and strain-rate which decrease with time. Such experiments are not suitable for large deformations and it is not surprising that the microstructure remains heterogeneous in both experiments which were at the peak temperature for 1 and 2 h, respectively, as the major contribution to dislocation multiplication must arise from the beginning of the test. Run #2087 was designed to limit the differential stress on the garnet at the level of the flow stress of olivine oriented along $[010]$. The slip directions in olivine are $[100]$, $[001]$ and, to a lesser extent, $[010]$. When olivine is compressed along $[010]$, the Schmid factor is zero for all slip systems. Dislocation glide is then theoretically precluded in such experiments. Indeed, deformation experiments performed under such conditions (Durham and Goetze, 1977) have shown than the creep rate is three orders of magnitude slower than those observed when easy glide can be activated. In principle, the only possible deformation mode for olivine in our experiments is pure climb of edge dislocation segments. Evidence of a glide in $(010)$ has, however, been observed in our olivine specimen. This suggests that the stress field may be more complicated than expected and that stress analysis based on the Schmid law in terms of a simplified stress geometry may not be relevant. This is to be compared with the microstructure in garnets for which dislocations with Burgers vectors $\frac{1}{2}[111]$ and $\frac{1}{2}[111]$ have been indexed (Fig. 10). These dislocations correspond to slip systems with no or little resolved shear stress. (The Schmid factors of these slip systems are not exactly zero as suggested by the simplified representation of Fig. 10, because the compression axis is slightly off the $[110]$ direction - approximately $10^\circ$.)

Although some dislocations appear to be confined in their glide planes, one observes pervasive evidence of dislocation climb: dislocation junctions, well-organized subgrain boundaries. This suggests that diffusion is efficient enough at 1440°C to enhance climb motion of dislocations. Dislocation climb is usually a slow and difficult process because it involves diffusion of point defects (multicomponent
diffusion in the case of garnets). Climb usually offers little contribution to the total deformation. However, this deformation mode is likely to control the deformation rate because it helps dislocations to overcome local obstacles. Climb also largely affects the restoration process. Rabier et al. (1981) suggested, however, that high temperature creep of YIG could involve pure climb motion of dislocations associated to Bardeen–Herring sources. Such a deformation mode could be an alternative explanation for the occurrence of \( \frac{1}{2}[111] \) and \( \frac{1}{2}[\overline{1}11] \) dislocations.

5. Conclusion

In this study, we have shown that high-pressure high-temperature deformation experiments can be performed on silicate garnets in a multi-anvil apparatus. We have shown also that garnet is ductile under our experimental conditions: \( P = 6 \) GPa, \( T = 1440^\circ C \) and high differential stress. TEM analysis of the dislocation microstructure suggests that both \( \frac{1}{2}
\langle 111 \rangle \) and \( \langle 100 \rangle \) slip have been activated. Occurrence of dissociation of mobile dislocations is reported. Dislocation climb appears to contribute significantly to plastic deformation at \( 1440^\circ C \).

References


