Deformation and hydration of the lithospheric mantle beneath the Kaapvaal craton, South Africa

Virginie Baptiste⁎, Andréa Tommasi, Sylvie Demouchy

Géosciences Montpellier, Université Montpellier 2 & CNRS, CC 60, Place E. Bataillon, 34095 Montpellier cedex 5, France

A R T I C L E   I N F O

Article history:
Received 21 October 2011
Accepted 3 May 2012
Available online xxxx

Keywords:
Craton
Kaapvaal
Peridotite
Deformation
Olivine crystal preferred orientations
Water contents

A B S T R A C T

To constrain the relations between deformation and metasomatism in the subcratonic lithospheric mantle, we have analyzed the microstructures and crystal preferred orientations in 50 mantle xenoliths from the Kaapvaal craton. Water contents in olivine and pyroxenes were measured in 14 samples equilibrated at different depths. Coarse-granular microstructures recording deformation by dislocation creep followed by annealing predominate. Mylonitic (sheared) peridotites with partially or totally recrystallized microstructures are however common below 140 km. Refractory compositions predominate, but multiple metasomatic events resulted in orthopyroxene enrichment or secondary crystallization of clinopyroxene and phlogopite. Coherent orthopyroxene and olivine CPO in most coarse-grained peridotites implies in pre- to synkinematic orthopyroxene enrichment or epitaxial growth on primary orthopyroxene. Undeformed, interstitial orthopyroxene, clinopyroxene, and phlogopite with random orientations in coarse-grained peridotites record post-kinematic modal metasomatic events. Deformation of these phases in the sheared peridotites implies that mylonitization results from a later event, which affected locally the deep cratonic lithosphere. Olivine CPO recording dominant [100] glide predominate at all depths. Only two samples, equilibrated at ~3.3 GPa show olivine [001] and orthopyroxene [001] axes subparallel, suggesting dominant [001] glide. Water contents in olivine are maximum (150 wt.ppm H2O) in peridotites equilibrated at ~160 km depth. Peridotites equilibrated below 180 km depth are, in contrast, almost dry. Lack of correlation between olivine mg# and water content indicates that the high water contents in olivine record re-hydration after the extensive partial melting, which produced the cratonic root. The vertical variation in water contents in olivine observed in the Kaapvaal peridotites may result from hydrogen addition or loss during extraction by the kimberlites. Comparison with magnetotelluric electrical conductivity data suggests, however, that the observed vertical variation of water contents in olivine may be representative of the present-day state of the Kaapvaal mantle, implying that extensive metasomatism resulted in hydration of the cratonic mantle at intermediate depths. The annealed microstructures of Kaapvaal peridotites indicate however that this metasomatism was not followed by remobilization of the cratonic root.

⁎ Corresponding author. Tel.: +33 467144912; fax: +33 467143603.
E-mail address: virginie.baptiste@gm.univ-montp2.fr (V. Baptiste).

1. Introduction

Cratons are domains of thick lithosphere with cold geotherms (Boyd et al., 1985; Chevrot and Zhao, 2007; Evans et al., 2011; Jaupart and Mareschal, 1999), which have remained stable for long geologic periods since their formation in the Archean (Pearson et al., 1995). However, while the mantle roots of some cratons, such as the Kaapvaal in South Africa, survived from reworking by plate tectonics and mantle convection since the Archean (Lenardic and Moresi, 1999). These observations led Jordan (1978) to propose the isopycnic (equal density) hypothesis to explain the stability of craton roots: the more refractory composition of the sub-cratonic mantle compensates the increase in density linked to cooling, implying a neutral buoyancy with respect to the asthenosphere. However, numerical models show that buoyancy alone is unable to explain the stability of cratons over several Ga (Doin et al., 1997; Lenardic and Moresi, 1999). The most common explanation for cratonic root longevity is a high viscosity contrast with the surrounding mantle (Doin et al., 1997; Pollack, 1986). The stability of cratonic roots is also favored by higher yield stresses relative to the mantle roots are poorly constrained, and are a major open question in geodynamics.

Studies of kimberlite-embedded xenoliths reveal that most cratonic peridotites are depleted in CaO and Al2O3 and have olivine with high mg# (mg# = MgO/(MgO+FeO)), implying that they were formed by a high degree of partial melting (e.g., Boyd and Mertzman, 1987). These observations led Jordan (1978) to propose the isopycnic (equal density) hypothesis to explain the stability of craton roots: the more refractory composition of the sub-cratonic mantle compensates the increase in density linked to cooling, implying a neutral buoyancy with respect to the asthenosphere. However, numerical models show that buoyancy alone is unable to explain the stability of cratons over several Ga (Doin et al., 1997; Lenardic and Moresi, 1999). The most common explanation for cratonic root longevity is a high viscosity contrast with the surrounding mantle (Doin et al., 1997; Pollack, 1986). The stability of cratonic roots is also favored by higher yield stresses relative to the
surrounding lithosphere and by neighboring weak material, such as mobile belts, which may localize the deformation (Lenardic et al., 2003).

The stability of cratonic roots appears therefore to depend essentially on its rheological properties. As the dominant phase (>50% in volume), olivine controls these properties. The strength of olivine-rich rocks under upper mantle conditions depends strongly on stress, temperature, grain size, and water fugacity (e.g., Demouchy et al., 2009; Hirth and Kohlstedt, 2003) and, on a lesser extent, on pressure, chemical composition, and oxygen fugacity (e.g., Keefner et al., 2011; Raterron et al., 2007). The relatively cold cratonic geotherm results in a viscosity increase of the cratonic root relatively to the surrounding convective mantle, but it cannot generate the 2–3 orders of magnitude viscosity contrast mantle required to ensure craton stability over Gyr time spans (Doin et al., 1997; Pollack, 1986). However, water contents in olivine from the Kaapvaal mantle measured by Fourier transform infrared spectroscopy (FTIR) are often high (up to 100 wt.ppm H₂O), except for the deepest samples (below 180 km), which have <10 wt.ppm H₂O (Granton et al., 2007; Kurosawa et al., 1997; Peslier et al., 2010).

In the present study, we explore the relations between deformation, melt or fluid percolation, and hydration by jointly analyzing microstructures, crystallographic preferred orientations, and water contents in a new series of peridotite xenoliths of the Kaapvaal craton. Previous petrophysical studies carried out on cratonic xenoliths focused on either the relation between deformation and seismic anisotropy (Bascou et al., 2011; Ben Ismaïl et al., 2001) or on the preferred orientation patterns that differ significantly from those obtained under dry conditions; these results were interpreted as due to changes in the dominant olivine slip systems due to variations in the OH⁻ concentration in olivine (Jung et al., 2006).

2. Geological setting

The Kaapvaal craton (South Africa) is an assemblage of Archean terranes that extends over more than 12,000 km². It is mainly composed by granitoids and gneisses that enclose narrow greenstone belts, but Upper Archean and Lower Proterozoic basins cover most of it (Begg et al., 2009; de Wit et al., 1992). Crustal formation occurred essentially between ~3.7 and ~3.1 Ga, followed by terrane assembly and stabilization of the craton between 3.1 and 2.6 Ga (de Wit et al., 1992; Griffin et al., 2003).

Formation of the Kaapvaal mantle root as early as 3.3–3.5 Ga is indicated by Re–Os whole-rock data on peridotite xenoliths (Pearson et al., 1995) and on sulfides in diamond inclusions (Shirey et al., 2002). These conclusions were confirmed by Re–Os analyses of sulfide phases in peridotite xenoliths, which indicate that the Kaapvaal mantle root formed prior to 3 Ga, that is, previously to or simultaneously to the formation of the crust, implying that each terrane carried its own keel during the craton assembly (Griffin et al., 2004).

Recent electrical conductivity measurements suggest that the lithospheric mantle beneath the Kaapvaal craton is currently defined by a high resistivity layer 200–250 km thick (Evans et al., 2011). High seismic velocities imaged in body and Rayleigh wave tomography also indicate a 250 km thick root beneath most of the craton (Chevrot and Zhao, 2007; James et al., 2001). Body-wave data suggest however that the high-velocity cratonic root may locally attain depths of 300 km (James et al., 2001). A slightly thinner high-velocity layer, 175–250 km thick, was imaged recently using SH-waves (Begg et al., 2009). These lithospheric thicknesses estimated from geophysical data are slightly higher, but still consistent with the 185 to 215 km thick Kaapvaal lithosphere constrained by xenolith thermobarometry (Eaton et al., 2009).

Following its stabilization, the Kaapvaal craton was affected by several magmatic events. The most important is the Bushveld complex, which intruded the Kaapvaal craton at 2.05 Ga (Scoates and Friedman, 2008). Another major magmatic event was the extrusion of the Karoo large igneous province at 182 Ma (Riley et al., 2005), which is associated with the Gondwana breakup. The craton was also affected by numerous kimberlitic eruptions. Kimberlitic pipes were mostly emplaced between the Late Jurassic and the Cretaceous (Kramers and Smith, 1983), but also erupted between 1650 and 1200 Ma (Kramers and Smith, 1983) and between 530 and 255 Ma (Alsopp et al., 1985; Kramers and Smith, 1983; Phillips et al., 1998).

These kimberlitic pipes contain xenoliths of the cratonic mantle that have been extensively studied for their microstructures, petrology, and geochemistry. Kaapvaal mantle xenoliths may be classified in two groups: (1) coarse-grained peridotites, which have dominantly refractory compositions and (2) fine-grained sheared peridotites, which are equilibrated at high temperatures and pressures and have, on average, more fertile compositions (Boullier and Nicolas, 1975; Boyd and Mertzman, 1987; Boyd and Nixon, 1975; Nixon et al., 1981). Petrological and geochemical data on Kaapvaal coarse-grained nodules reveal that they represent a highly refractory lithospheric mantle residue, implying ~40% melt extraction, which was subsequently affected by several metasomatic episodes (e.g., Boyd and Mertzman, 1987; Griffin et al., 2003; Kelemen et al., 1998; Nixon et al., 1981; Simon et al., 2007). The high orthopyroxene/olivine ratio that characterizes many low-temperature Kaapvaal xenoliths, for instance, is usually attributed to Si-enrichment due to interaction with subduction-related fluids or intraplate hydrous melts (Bell et al., 2005; Kelemen et al., 1998; Simon et al., 2007; Wasch et al., 2009). Lu–Hf and Sm–Nd model ages on garnet and orthopyroxene clots in peridotite xenoliths from Kimberley suggest multiple, rather than a single Si-enrichment episode, ranging from 1.3 to 1.1 Ga to Neoproterozoic (Wasch et al., 2009). Kaapvaal xenoliths and xenocrysts also display geochemical evidence for refertilization, which added basaltic components like Fe, Ca and Al to a depleted protolith (Griffin et al., 2003). Crystallization of diopside, hkerzolitic garnet, and phlogopite has been proposed to result from interactions with kimberlitic–carbonatitic fluids (Grégoire et al., 2003; Simon et al., 2007). Recent geochemical data indicate that diamonds and subcalcic garnets also result from interaction with reduced asthenosphere-derived fluids, corroborating the hypothesis that the cratonic mantle was originally essentially composed by highly refractory harzburgites and dunites (Malkovets et al., 2007). Melt-related metasomatism has also been identified in sheared peridotites, which may have acquired their fertile composition by interactions with asthenosphere-derived melts shortly before the kimberlitic eruption (O’Reilly and Griffin, 2010). In the present study, we will refer to all such changes in modal composition due to melt or fluid-rock interactions as modal metasomatism.

For the present study, we selected, from the >400 samples of the Kaapvaal mantle xenolith collection of Geosciences Montpellier (France), 50 peridotite xenoliths with varied microstructures and compositions. Study of samples from 9 pipes with ages ranging from 1.2 Ga to 87 Ma: Kimberley, Jagersfontein, Monastery, Lentseng, De Beers, Finsch, Kamfersdam, Premier, and Mothae (Fig. 1 and Table 1), allowed the sampling of possible lateral and temporal variations of deformation and composition in the Kaapvaal mantle. Analysis of a larger number of samples from Jagersfontein and Kimberley (18 samples for each locality) allowed studying vertical variations at these locations.
3. Methods

3.1. Mineral compositions and pressure–temperature estimates

Chemical compositions of olivine, orthopyroxene, clinopyroxene, garnet and spinel were analyzed using a Cameca SX100 electron microprobe at Microsonde Sud facility, in Montpellier (France). Analysis conditions were a 20 kV accelerating voltage and a 10 nA probe current. Core and rim compositions were analyzed systematically. For each mineral phase, three to four grains were measured. The aim was to determine pressure and temperature equilibrium conditions of the selected peridotites to constrain their depth distribution. Equilibrium temperatures were calculated using the two pyroxenes Fe–Mg exchange geothermometer from Brey and Köhler (1990), which has an uncertainty of ±30 °C. In highly-depleted garnet harzburgites that contained no clinopyroxene, the geothermometer of O’Neill and Wood (1979), based on Fe–Mg partitioning between garnet and olivine, which has uncertainties around ±60 °C, was used. For garnet- and clinopyroxene-free harzburgites, we used the geothermometer Li et al. (1995), based on Fe–Mg exchange between olivine and spinel, which yield temperatures within ±50 °C. Equilibrium pressures were calculated using the orthopyroxene–garnet barometer of Nickel and Green (1985), which has uncertainties of ±0.2 GPa. For the spinel-bearing peridotites, pressures were estimated based on the equilibrium temperatures and the geotherm that best fits the equilibrium pressure and temperature data for the garnet-bearing samples.

3.2. Electron-backscattered diffraction (EBSD)

Crystallographic preferred orientations (CPO) of olivine, pyroxenes, and garnet in all 50 samples were measured at the SEM-EBSD facility in Geosciences Montpellier by indexation of EBSD patterns produced by interaction between an electron beam with the crystals in thin sections tilted at 70° to the horizontal. Measurements were performed in a JEOL JSM 5600 scanning electron microscope using an acceleration voltage of 17 kV and a working distance of 23 mm. Maps covering almost entirely each thin section were performed.
using sampling steps of 100, 40 or 30 μm, depending on grain size. Indexation rates range from 40 to 75% depending on the extent of fracturing and serpentinization in the xenolith. Phlogopite is usually poorly indexed. Indexation is also poor in very fine-grained layers in samples displaying mylonitic or fluidal mosaic textures. Orthopyroxene was rarely misindexed as clinopyroxene. Errors in the measurements were reduced by careful post-acquisition data treatment, controlled by comparison between EBSD maps and microscopic observations. Modal composition, grain sizes, and shape-preferred orientations were also obtained from EBSD maps.

Crystal-preferred orientation data is displayed in pole figures, presented as lower hemisphere stereographic projections. To avoid over-representation of large grains, data were plotted as one point per grain. When the foliation and lineation could be identified, the orientation of the main crystallographic directions: [100], [010] and [001] for olivine and pyroxenes, was plotted relatively to the principal axes of the deformation ellipsoid X, Y, and Z. However, in most coarse-grained samples, the identification of the foliation and lineation was not possible and thin sections were cut in random orientations. To allow easy comparison among different samples, we rotated the CPO of all samples into a common orientation, in which the maximum concentration of orthopyroxene [001] axes and of the olivine [010] axes are parallel to the E–W and the N–S directions of the pole figure, respectively. This choice was made on the observation that [001] is the only known glide direction in orthopyroxene; plastic deformation tends therefore to align this axis in the flow direction (cf. review in Frets et al., 2012). This choice allowed presenting the CPO without making an ad-hoc hypothesis on the dominant glide direction in olivine. When the orthopyroxene CPO was too dispersed, the maximum concentration of olivine [100] or [001] axes, depending on which had the strongest concentration, was placed in the E–W direction of the pole figure.

The strength of the fabric was quantified using the dimensionless J-index, which is the volume-averaged integral of squared orientation densities defined by:

\[ J = \frac{1}{4} \int d^2 \phi \, f(\phi) \]

where \( f(\phi) \) is the orientation distribution function (ODF) and \( d\phi \) = d\( \phi_1 \) d\( \phi_2 \) d\( \phi_3 \) sin\( \phi_3 \) d\( \phi_4 \), where \( \phi_1, \phi_2, \) and \( \phi_3 \) are the Euler angles that define the rotations allowing for coincidence between the crystallographic and external reference frames. Olivine CPO in natural peridotites is characterized by J-indices between 2 and 20, with a peak at 8–10 (Ben Ismail and Mainprice, 1998; Tommasi et al., 2000). The J-index for all samples was calculated based on the mean orientation of each grain using the SuperCtcf program by D. Mainprice (ftp://www.gm.univ-montp2.fr/mainprice/CareWare_Unicel_Programs/) with a 10° Gaussian half-width, 1° cells, and truncation of the orientation distribution function (ODF) at 22°.

3.3. Fourier transform infrared spectroscopy (FTIR)

Fourteen doubly-polished thin sections were prepared for unpolarized FTIR analysis. Prior to analyses, all sections were immersed in pure acetone for at least 12 h to dissolve any intergranular CrystalBond glue. FTIR spectroscopy analyses were performed at the Laboratoire des Colloïdes, Verres, Nanomatériaux in Montpellier using a Bruker IFS66v coupled with a Bruker HYPERION microscope and a liquid nitrogen-cooled mercury–cadmium–telluride (MCT) detector. A Globar light source and a Ge-KBr beam splitter were used to generate unpolarized mid-infrared radiation. Measurements on olivine, orthopyroxene, clinopyroxene, garnet, and phlogopite crystals were performed with square aperture sizes ranging from 40 to 100 μm; for each measurement 200 scans were accumulated with a resolution of 4 cm\(^{-1}\). They were preceded by a background measurement, followed by a baseline correction using the OPUS software, and, finally, normalized to a sample thickness of 1 cm. Fractures and inclusions were strictly avoided. The sample thickness was measured using a micrometer with a tolerance of ±1 μm and was always near 500 μm.

The calibration of Paterson (1982) was used to quantify the OH concentration in each mineral phase:

\[ c_{\text{OH}} = \frac{X_i}{1500} \int (3780 - v_i) k(v) dv \]

where \( c_{\text{OH}} \) is the hydroxyl concentration (in mol H/l), \( \xi \) is an orientation factor (1/3 for unpolarized measurements), and \( k(v) \) is the absorption coefficient in cm\(^{-1}\) for a given wavenumber \( v \). \( X_i \) is a density factor (\( X_i(\text{olivine}) = 2695 \text{ ppm wt. H}_2\text{O}; X_i(\text{orthopyroxene}) = 2812 \text{ ppm wt. H}_2\text{O}; X_i(\text{clinopyroxene}) = 2761 \text{ ppm wt. H}_2\text{O}. \) Estimated errors on the resulting water contents are ~30% (Kohlstedt et al., 1996). To compare the present data with previous studies where measurements were done using polarized infrared radiation and the Bell et al. (2003) calibration, OH concentrations for olivine have to be multiplied by a factor between 2 and 4 (Bell et al., 2003). Following Ferot (2011), a factor of 3 was used in the present study.

4. Microstructures

Kaapvaal xenoliths are generally highly fractured due to fast decompression during kimberlite eruption and have abundant alteration products, such as serpentine along grain boundaries and fractures. Original microstructures can however still be clearly observed. Five distinct microstructures were identified following the original classification of Boullier and Nicolas (1975). Among the 50 studied xenoliths, 78% are coarse-grained, displaying either coarse-granular (protogranular) or coarse-porphroclastic microstructures, and 22% are fine-grained, high-temperature sheared peridotites, showing mylonitic, fluidal mosaic, or mosaic microstructures (Fig. 2).

4.1. Coarse-grained peridotites

Coarse-grained peridotites predominate in our xenolith suite; they are present in every studied pipe except Mothae (Fig. 1). They are characterized by plurimillimetric grain sizes (Fig. 2). Most are coarse-granular (CG) peridotites (32/50, 66%), with rare intracrystalline deformation features, but a few are coarse-porphyroclastic (6/50, 12%), exhibiting a well-defined foliation and lineation and frequent subgrain boundaries in olivine.

4.1.1. Coarse-granular peridotites

Coarse-granular peridotites correspond to the protogranular peridotites in the original Boullier and Nicolas (1975) classification, being characterized by large anhedral olivine and orthopyroxene crystals. The coarsest-grained peridotites from Jagersfontein have cm-wide crystals (Fig. 2a). Most coarse-granular xenoliths have, however, 5–8 mm wide crystals in average (Fig. 2b) and a few samples have slightly smaller grain sizes (1–2 mm in average, Fig. 2c).

In most coarse-granular peridotites, olivine and orthopyroxene crystals have irregular shapes with curvilinear boundaries, evolving locally to polygonal arrangements with 120° triple junctions (Fig. 2a–c). No foliation or lineation can be identified. However, some coarse-granular xenoliths from Kimberley have a weak foliation, highlighted by the elongation of olivine and orthopyroxene crystals (aspect ratios range from 1:2 to 1:4, Fig. 2b).

Olivine crystals are usually free of intracrystalline deformation, but subgrains and undulose extinctions are locally observed (white arrows in Fig. 2a–c). Orthopyroxene crystals sometimes display kinks (black arrows, Fig. 2b and c). Small interstitial orthopyroxenes are also observed; they are most often free of any intracrystalline deformation features, but may also display kinks (Fig. 3a). Clinopyroxene exsolution may or...
not be present. Interpenetrating boundaries between orthopyroxene and olivine grains were identified in more than half of the studied samples (Fig. 2a and b). Rounded olivine crystals are occasionally included in orthopyroxene, but orthopyroxene inclusions in olivine are also observed. Large, often undeformed clinopyroxene crystals are observed in some samples. Most clinopyroxene occurs however as small dark green crystals along vein-like structures or as small interstitial crystals. It is often spatially related to phlogopite (Fig. 3b), spinel, and garnet.

When present, garnet forms large rounded 1 to 5 mm wide crystals (Fig. 2b). Garnet rimming orthopyroxene is however observed in lherzolite FRB1339 from Premier (Fig. 3c). Kelyphite rims with variable thickness (~0.06 to 1 mm) are almost always present. When present, spinel occurs as dark, irregularly shaped and mm-sized crystals, as small crystals in kelyphite rims, or as small interstitial crystals along fractures and grain boundaries. Spinel–pyroxene symplectites are observed in two harzburgites. Garnet- and spinel-bearing harzburgite J41 has

---

Fig. 2. Photomicrographs in cross-polarized light showing typical microstructures for the studied xenoliths. (a) Very coarse-granular gt-bearing J34 lherzolite from Jagersfontein, showing centimetric irregularly shaped grains with straight grain boundaries. (b) Coarse-granular texture in gt- and sp-bearing harzburgite KBBF16 from Kimberley, where elongation of olivine crystals marks a weak foliation (S). (c) Sp-harzburgite KBBF10 from Kimberley showing an equigranular, less coarse-grained microstructure. White arrows in (b) and (c) mark undulose extinctions and subgrain boundaries in olivine crystals. (d) Coarse-porphyroclastic spinel harzburgite KBBF8 exhibiting a foliation (S) marked by elongated olivine crystals with well-developed subgrain boundaries and undulose extinction (white arrows); orthopyroxene crystals have irregular shapes with embayments filled by olivine and display kinks locally (black arrow). (e) Mylonite KBBF11 showing strained orthopyroxene and olivine porphyroclasts surrounded by a fine-grained olivine–orthopyroxene matrix; dashed lines highlight very fine-grained orthopyroxene-rich domains. (f) Fluidal mosaic texture in lherzolite KBB63 characterized by elongated garnet crystals enclosed in a matrix composed of polygonal olivine and very fine-grained orthopyroxene-rich domains, highlighted by dashed lines as in (e). (g) Mosaic texture in garnet–lherzolite J47 characterized by olivine and orthopyroxene porphyroclasts enclosed in an olivine-rich polygonal matrix. S marks the foliation trace deduced from crystallographic orientation data. (h) Pie-chart showing the representativity of the different microstructures among the 50 studied xenoliths.
spinel–clinopyroxene symplectites rimmed by garnet (Fig. 3d), whereas harzburgite KBBF10 has spinel–orthopyroxene symplectites.

Phlogopite crystallizes as reaction products in kelyphite rims, along fractures, or as small interstitial crystals along grain boundaries or triple junctions. Millimetric to plurimillimetric, subhedral phlogopite crystals are observed in two samples (KBJ59 and KBBF14, Table 2). Among these samples, garnet- and spinel-bearing lherzolite KBJ59 from Jagersfontein displays an extremely high phlogopite content (14%).

4.1.2. Coarse-porphyroclastic peridotites

Coarse-porphyroclastic microstructures are rarer, but they are observed in 6 harzburgites and lherzolites from the Kimberley, Finsch and Premier pipes, which display equilibrium temperatures ranging from 850 °C to 1464 °C (Table 2). These peridotites have plurimillimetric, lens-shaped olivine crystals (with aspect ratios up to 4:1) marking the foliation and the lineation (Fig. 2d). Olivine crystals usually exhibit undulose extinction and subgrains (white arrows

Please cite this article as: Baptiste, V., et al., Deformation and hydration of the lithospheric mantle beneath the Kaapvaal craton, South Africa, Lithos (2012), doi:10.1016/j.lithos.2012.05.001
Calculation was not possible because of compositional variability leading to large variations (>200° or >1 GPa) in P, T estimates within the sample.

In contrast, sp-

Table 2

<table>
<thead>
<tr>
<th>Texture</th>
<th>Locality</th>
<th>Sample</th>
<th>Pressure (GPa)</th>
<th>Temperature (°C)</th>
<th>Ol mg#</th>
<th>Reference for mg#, T, and P data</th>
<th>Modal compositions</th>
<th>Olivine</th>
<th>CPO</th>
<th>J-index</th>
<th>J-index</th>
</tr>
</thead>
<tbody>
<tr>
<td>CG</td>
<td>Kimberley</td>
<td>KBBF3a</td>
<td>3.8 (1)</td>
<td>926 (1)</td>
<td>90.8</td>
<td>This study</td>
<td>ol 13 4 9 0 0</td>
<td>Axial [100]</td>
<td>7.6</td>
<td>9.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF6b</td>
<td>3.7 (1)</td>
<td>908 (1)</td>
<td>91.0</td>
<td>This study</td>
<td>ol 8 2 4 6 0</td>
<td>Axial [100]</td>
<td>7.0</td>
<td>3.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF9</td>
<td>4.7 (1)</td>
<td>1102 (1)</td>
<td>89.8</td>
<td>This study</td>
<td>ol 67 16 9 8 0</td>
<td>Axial [100]</td>
<td>3.6</td>
<td>7.5</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF10</td>
<td>–</td>
<td>949 (1)</td>
<td>92.4</td>
<td>This study</td>
<td>ol 72 27 1 0 0</td>
<td>Axial [100]</td>
<td>4.7</td>
<td>2.4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF14</td>
<td>–</td>
<td>996 (1)</td>
<td>93.0</td>
<td>This study</td>
<td>ol 16 1 0 1 0</td>
<td>Axial [100]</td>
<td>4.0</td>
<td>6.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF15</td>
<td>4.3 (1)</td>
<td>1067 (1)</td>
<td>92.6</td>
<td>This study</td>
<td>ol 55 61 0 0 0</td>
<td>Ortho</td>
<td>4.3</td>
<td>5.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF16</td>
<td>4.0 (1)</td>
<td>–</td>
<td>93.3</td>
<td>BI</td>
<td>71 26 0 3 0</td>
<td>Ortho</td>
<td>3.0</td>
<td>4.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF20</td>
<td>–</td>
<td>800 (1)</td>
<td>–</td>
<td>BI</td>
<td>65 35 0 0 0</td>
<td>Ortho</td>
<td>10.2</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRB1404</td>
<td>4.4 (1)</td>
<td>987 (2)</td>
<td>93.5</td>
<td>J</td>
<td>59 76 1 4 0</td>
<td>Ortho</td>
<td>3.5</td>
<td>6.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRRE2122</td>
<td>4.4 (2)</td>
<td>994 (2)</td>
<td>93.5</td>
<td>J</td>
<td>65 31 0 4 0</td>
<td>Ortho</td>
<td>2.1</td>
<td>8.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRRE2133</td>
<td>3.8 (1)</td>
<td>851 (1)</td>
<td>91.8</td>
<td>J</td>
<td>68 18 4 10 0</td>
<td>Axial [100]</td>
<td>5.5</td>
<td>10.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRRE1447</td>
<td>4.2 (1)</td>
<td>968 (2)</td>
<td>93.4</td>
<td>J</td>
<td>73 23 1 3 0</td>
<td>Axial [100]</td>
<td>2.4</td>
<td>7.8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRE348</td>
<td>3.7 (1)</td>
<td>936 (1)</td>
<td>92.3</td>
<td>J</td>
<td>58 28 7 7 0</td>
<td>Ortho</td>
<td>4.7</td>
<td>8.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>KBJ50</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>KBJ52</td>
<td>–</td>
<td>900 (4)</td>
<td>92.7</td>
<td>This study</td>
<td>ol 25 7 0 0 0</td>
<td>Ortho</td>
<td>4.3</td>
<td>5.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>KBJ54</td>
<td>3.8 (1)</td>
<td>907 (4)</td>
<td>92.3</td>
<td>This study</td>
<td>ol 25 7 0 0 0</td>
<td>Ortho</td>
<td>4.3</td>
<td>5.7</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>KBJ56</td>
<td>3.5 (1)</td>
<td>895 (2)</td>
<td>92.5</td>
<td>This study</td>
<td>ol 65 28 0 7 0</td>
<td>Axial [100]</td>
<td>8.7</td>
<td>3.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>KBJ59</td>
<td>–</td>
<td>898 (1)</td>
<td>–</td>
<td>BI</td>
<td>65 36 0 0 0</td>
<td>Ortho</td>
<td>1.5</td>
<td>4.8</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>KBJ62</td>
<td>3.1 (1)</td>
<td>759 (2)</td>
<td>92.6</td>
<td>This study</td>
<td>ol 59 29 5 7 0</td>
<td>Axial [100]</td>
<td>10.6</td>
<td>18.4</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>J42</td>
<td>2.4 (1)</td>
<td>654 (1)</td>
<td>92.4</td>
<td>This study</td>
<td>ol 61 39 0 0 0</td>
<td>Ortho</td>
<td>5.8</td>
<td>7.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>J41</td>
<td>3.7 (1)</td>
<td>668 (4)</td>
<td>92.4</td>
<td>This study</td>
<td>ol 28 19 0 0 0</td>
<td>Ortho</td>
<td>10.5</td>
<td>20.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Jagersfontein</td>
<td>J57</td>
<td>3.1 (1)</td>
<td>765 (2)</td>
<td>92.4</td>
<td>This study</td>
<td>ol 19 3 0 0 0</td>
<td>Ortho</td>
<td>8.5</td>
<td>8.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Monastery</td>
<td>ROM23</td>
<td>3.3 (1)</td>
<td>910 (1)</td>
<td>92.2</td>
<td>MO</td>
<td>66 28 4 1 0</td>
<td>Ortho</td>
<td>7.1</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Monastery</td>
<td>ROM49</td>
<td>2.6 (1)</td>
<td>874 (1)</td>
<td>92.2</td>
<td>MO</td>
<td>76 20 3 1 0</td>
<td>Ortho</td>
<td>7.1</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lentseng</td>
<td>PHN4274</td>
<td>4.3 (1)</td>
<td>1015 (1)</td>
<td>93.1</td>
<td>BI</td>
<td>62 33 2 3 0</td>
<td>Ortho</td>
<td>6.1</td>
<td>7.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kamfersdam</td>
<td>PHN5580</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>De Beers</td>
<td>KBB2</td>
<td>4.0 (1)</td>
<td>917 (2)</td>
<td>93.3</td>
<td>This study</td>
<td>ol 66 33 0 1 0</td>
<td>Ortho</td>
<td>6.5</td>
<td>9.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRRE1330</td>
<td>–</td>
<td>744 (1)</td>
<td>–</td>
<td>BI</td>
<td>80 17 3 0 0</td>
<td>Axial [100]</td>
<td>4.7</td>
<td>2.6</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRRE1336</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>FRRE1339</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>PHN5266</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Finsch</td>
<td>FRF1513</td>
<td>5.2 (1)</td>
<td>1037 (2)</td>
<td>92.6</td>
<td>This study</td>
<td>ol 70 27 1 2 0</td>
<td>Ortho</td>
<td>7.0</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF8</td>
<td>–</td>
<td>–</td>
<td>943 (1)</td>
<td>This study</td>
<td>ol 81 19 0 0 0</td>
<td>Axial [100]</td>
<td>8.5</td>
<td>8.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>KBBF18</td>
<td>–</td>
<td>–</td>
<td>850 (1)</td>
<td>92.2</td>
<td>BI</td>
<td>75 17 6 0 0</td>
<td>Axial [100]</td>
<td>2.1</td>
<td>6.5</td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>PHN4276</td>
<td>4.9 (1)</td>
<td>1053 (1)</td>
<td>93.2</td>
<td>BI</td>
<td>95 2 3 3 0</td>
<td>Ortho</td>
<td>3.0</td>
<td>4.1</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Kimberley</td>
<td>PHN5267</td>
<td>6.5 (1)</td>
<td>1464 (1)</td>
<td>93.1</td>
<td>This study</td>
<td>ol 70 19 6 5 0</td>
<td>Ortho</td>
<td>2.3</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Finsch</td>
<td>FRF1501</td>
<td>4.6 (1)</td>
<td>1030 (2)</td>
<td>93.0</td>
<td>This study</td>
<td>ol 70 22 0 8 0</td>
<td>Ortho</td>
<td>3.6</td>
<td>5.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Finsch</td>
<td>FRF1512</td>
<td>4.7 (1)</td>
<td>1058 (2)</td>
<td>92.0</td>
<td>This study</td>
<td>ol 68 22 0 10 0</td>
<td>Ortho</td>
<td>3.5</td>
<td>6.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Finsch</td>
<td>FRF1501</td>
<td>4.6 (1)</td>
<td>1030 (2)</td>
<td>93.0</td>
<td>This study</td>
<td>ol 70 22 0 8 0</td>
<td>Ortho</td>
<td>3.6</td>
<td>5.0</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Finsch</td>
<td>FRF1512</td>
<td>4.7 (1)</td>
<td>1058 (2)</td>
<td>92.0</td>
<td>This study</td>
<td>ol 68 22 0 10 0</td>
<td>Ortho</td>
<td>3.5</td>
<td>6.9</td>
<td></td>
</tr>
</tbody>
</table>

References for P, T data: BI = Ben Ismaïl (1999); B&N = Boyd and Nixon (1978); G = Grant et al. (2007); J = James et al. (2004); MO = Moore (1986).

Texture: CG = Coarse-grained peridotite; CP = Coarse-porphyroclastic; E = Equigranular; FM = Fluidal mosaic; M = Mosaic; MY = Mylonite. CPO types: ortho = orthorhombic, bimodal, axial [100] and axial [010] crystal-preferred orientations, see main text and Fig. 6 for details.

Please cite this article as: Baptiste, V., et al., Deformation and hydration of the lithospheric mantle beneath the Kaapvaal craton, South Africa, Lithos (2012), doi:10.1016/j.lithos.2012.05.001


Calculation was not possible because of compositional variability leading to large variations (>200° or >1 GPa) in P, T estimates within the sample.

Texture: CG = Coarse-grained peridotite; CP = Coarse-porphyroclastic; E = Equigranular; FM = Fluidal mosaic; M = Mosaic; MY = Mylonite. CPO types: ortho = orthorhombic, bimodal, axial [100] and axial [010] crystal-preferred orientations, see main text and Fig. 6 for details.
serpentinite, but they still preserve the original polygonal shapes. Orthopyroxene porphyroclasts are rare; they are strongly altered and generally show abundant clinopyroxene exsolutions.

4.2. Sheared peridotites

Sheared peridotites are characterized by a bimodal grain size distribution where rare porphyroclasts are enclosed in a fine-grained matrix (grain sizes range from tens to few hundreds of micrometers). They also usually display a well-developed foliation, but the later may be absent (mosaic microstructure). Two other microstructures: mylonitic and fluidal mosaic, may be discriminated based on the average grain size of the recrystallized matrix. In our sample set, mylonitic microstructures are only observed in two xenoliths from Kimberley, fluidal mosaic structures in five xenoliths from Jaggersfontein, and mosaic microstructures in four samples from Jaggersfontein and one from Mothae.

The mylonites display large asymmetric olivine and orthopyroxene porphyroclasts (1–5 mm long), with well-developed undulose extinctions, closely-spaced subgrain boundaries, and recrystallization tails enclosed in a fine-grained matrix composed of recrystallized olivine (30–100 μm wide) and very fine-grained (<25 μm) orthopyroxene-rich bands (Fig. 2e). Phlogopite often displays sigmoidal shapes and strong undulose extinction. Euhedral olivine crystals devoid of any internal deformation (tablets) commonly crystallize on olivine porphyroclasts and sometimes also on matrix crystals (Fig. 3f).

In fluidal mosaic peridotites, orthopyroxene and clinopyroxene porphyroclasts are also enclosed in a fine-grained matrix with a polygonal texture, but olivine grains in the matrix are larger (up to 0.5 mm wide; Fig. 2f). Olivine is usually entirely recrystallized, but rare porphyroclasts are preserved in gt- and sp-lherzolites KBJ30 and KBJ33. Orthopyroxene porphyroclasts are elongated and show asymmetric recrystallization tails composed of very fine-grained orthopyroxene with polygonal boundaries and undulose extinctions (Fig. 3g). These recrystallization tails evolve into mm-scale, orthopyroxene-rich bands within the olivine-rich matrix (Fig. 3g). Olivine crystals in the polygon are polygonal and free of any internal deformation features (Fig. 3g). Garnet forms large porphyroclasts, which may be elongated parallel to the foliation, as in mosaic fluidal peridotites KBJ63 and KBJ14 (Fig. 2f). It can also be rounded, as in samples KBJ30 and KBJ33, in which garnet is more abundant (>15%).

The four studied mosaic peridotites (sp-lherzolite J47, gt- and sp-lherzolites J63, KBJ30, KB60) are characterized by large olivine porphyroclasts with strong undulose extinction and rare subgrain boundaries, and by irregularly-shaped orthopyroxene porphyroclasts enclosed in a polygonal olivine-rich matrix (Fig. 2g). Contrary to the previous textures, these samples show neither a clear foliation nor a compositional banding. Crystallization of olivine tablets on orthopyroxene-rich bands (Fig. 2e). Phlogopite often displays sigmoidal shapes and strong undulose extinction. Euhedral olivine crystals devoid of any internal deformation (tablets) commonly crystallize on olivine porphyroclasts and sometimes also on matrix crystals (Fig. 3f).

4.3. Compositions and equilibrium conditions

Most coarse-grained peridotites are harzburgites or clinopyroxene-poor lherzolites with high modal contents of orthopyroxene (≥20%, Fig. 4). Coarse-grained lherzolites are rarer. A single very coarse-grained peridotite from Jaggersfontein has a wehrlitic composition that is accompanied by enrichment in phlogopite. In contrast, half of the analyzed sheared peridotites from Jaggersfontein has a wehrlitic composition that is accompanied by enrichment in phlogopite. Garnet modal content is highly variable (0–18%, Table 2) and shows no correlation with olivine or clinopyroxene contents. Spinell, when present, is always a minor phase (<1%).

The mineral compositions of the analyzed samples are presented in the online Supplementary Material Table 1. Olivine mg# varies between 87.7 and 94.3 (Table 2). Mg# tends to be higher in coarse-grained peridotites (average mg# = 92.1) and lower in the sheared peridotites (average mg# = 91.1). However, mylonites from Kimberley are harzburgites with refractory olivines (average mg# = 93.4) and one of the lowest mg# (88.9) pertain to olivine crystals from coarse-granular peridotite. Low mg# in olivine are sometimes, but not always, accompanied by modal metasomatism (enrichment in clycophyroxene or phlogopite).

Orthopyroxene is enstatite; it has very low Al₂O₃ contents (<1.10 wt.%), except for five samples (KBBF10, FRB1330, FRB1139, PHN1925, and PHN5266), where Al₂O₃ contents up to 2.76 wt.% are observed. Orthopyroxene has also Ti-poor compositions (<0.30 wt.%). Clinopyroxene is diopside in 11 samples and augite in 14 out of 25 samples. Garnet is always pyrope-rich and is Ca-saturated except in 4 samples (FRB1136, KBJ8, KBJ50 and KBJ56). Spinel is generally Cr-rich (cr# > 50, reaching 94 in harzburgite KBBF14). However, phlogopite-rich harzburgite KBJ59 has Al-rich spinels (cr# = 9).

Coarse-grained peridotites display the widest range of equilibrium temperatures, from 650 °C to 1464 °C, and pressures, from 2.4 to 6.5 GPa (Fig. 5). Equilibrium temperatures and pressures for sheared peridotites are comprised between 1023 and 1306 °C and 4.6–6.5 GPa. Pressure and temperature estimates for the coarse-grained xenoliths are consistent with a geotherm leading to a surface heat flux of 40 mW/m², which intercepts the 1283 °C adiabat at 195 km of depth as illustrated in Fig. 5. Most sheared peridotites plot however slightly above this geotherm, in agreement with previous data for Kaapvaal sheared xenoliths (Finnerty and Boyd, 1987; Kennedy et al., 2002).

5. Crystal-preferred orientations

5.1. Coarse-granular and coarse-porphyrcolastic peridotites

Coarse-granular and coarse-porphyrcolastic peridotites show similar olivine CPO patterns, which range from orthorhombic to axial[100] or axial-[010] (Fig. 6). Orthorhombic olivine CPO predominates (~60% of the coarse-grained peridotites). This CPO is characterized by point concentrations of the three crystallographic axes, with [100]
James et al. (2003) and Stachel et al. (2003), respectively.

dicate data calculated using the O’Neill and Wood (1979), the Brey and Köhler (1990),
(Fig. 6).

display a stronger concentration of [001] axes than of [100] axes

strength and pattern (Fig. 7b) or composition (Table 2). There is,

study were obtained for coarse-grained peridotites with orthorhomb-
(Fig. 7a); the lowest and the highest J-index values in the present

clear correlation between CPO strength and microstructure

are comprised between 1.5 and 10.6, which correspond to very

distribution of [100] and [001] axes. Finally, two coarse-grained peri-

coarse-grained harzburgites have characteristic axial-[010] CPO,

axes exhibiting the strongest point concentration (Fig. 6). Seven

axes are aligned perpendicular to the plane that contain most [001]

and [100] axes. In twelve other samples, [100] axes are normal to

the plane containing the [001] and [010] axes. Two samples (KBBF9
and FRB1501) have, however, an axial-[001] pattern, characterized

by [100] and [010] axes distributed in plane normal to the point

concentration of [001] axes. Six coarse-grained harzburgites (KBBF18,
FRB1404, FRB1422, FRB1423, KB52, J41 and J57) have an almost random

orthopyroxene CPO. Three of these samples (J41, FRB1404, and FRB1422) are particularly enriched in orthopyroxene, with modal

contents greater than 30%. In coarse-grained peridotites PHN2760
and KB50, the number of analyzed orthopyroxene crystals is too

small to identify a CPO pattern.

5.2. Sheared peridotites

Mylonitic, fluidal mosaic, and mosaic peridotites show weak oliv-

cine CPO: J-indexes are comprised between 1.9 and 4.6 (Fig. 7a,
Table 2). Olivine CPO patterns are nevertheless well-defined, with ei-

ther an orthorhombic or an axial-[010] symmetry (Fig. 6). One

mylonite (KBB11) and 3 fluidal mosaic lherzolites (KB8, KB14 and
KB33) have typical orthorhombic olivine CPO with [100] axes aligned

close to the lineation and [010] axes normal to the foliation. Seven

garnet–lherzolites from Jagersfontein, Kimberley and Mothae show,

however, a particular axial-[010] pattern characterized by a bimodal

alignment of [100] and [010] axes in the foliation plane (Fig. 6).

None of the maxima are parallel to the lineation, which is underlined

by the elongation of the olivine crystals and the maximum concentra-
tion of orthopyroxene [001] axes. These bimodal olivine CPO samples

have systematically weak CPO (Fig. 7b). There is also a clear depth
distribution: bimodal olivine CPO are characteristic of sheared peri-
dotites equilibrated at pressures >4.5 GPa (Fig. 7c).

Orthopyroxene CPOs are weak, as indicated by J-indexes between

1.6 and 4.9. CPO patterns are nevertheless clear and correlated to the

olivine CPO, although a small obliquity (10°) between the CPO of the

two minerals is observed for most samples. [001] is always aligned in

the lineation. In six xenoliths (mosaic peridotites KB60 and J47,

mylonite KBFF11, fluidal mosaic peridotites KBJ33, KB14 and KB63),

[100] axes are concentrated normal to the foliation. In four other sam-

ples (mosaic lherzolite KBJ30, mylonite FRB1402, fluidal mosaic

lherzolites KB6 and KB8), [010] axes are aligned normal to the folia-
tion, whereas [100] axes are distributed in the foliation plane and

perpendicular to the lineation. Clinopyroxene CPOs (presented in

the online supplementary material) are weak, but correlated to the

orthopyroxene CPO in all sheared peridotites, except in mylonite

FRB1402.

6. FTIR spectroscopy and water contents

6.1. Olivine

The olivine spectra of xenoliths from Kimberley are heterogeneous

and complex, often displaying more than six O–H absorption bands

(Fig. 8a). This large number of absorption bands suggests H incorp-

oration in a variety of point defects (e.g., Berry et al., 2007; Kovacs et

al., 2010; Miller et al., 1987). A total of nine bands were identified in

the studied xenoliths from Kimberley, but in most samples just a

few of these bands are observed. Absorption bands can be divided

into two groups: group I with absorptions between 3650 and

3450 cm−1 and group II between 3450 and 3200 cm−1 (Bai and

Kohlstedt, 1993). The band at 3571 cm−1 exhibits the maximum O–

H absorbance, in agreement with observations in many natural olivi-

nes (Miller et al., 1987). The intensity of other group I absorption

bands is variable. The 3610, 3590 and 3475 cm−1 bands usually repre-

sent major peaks as observed for samples PHN2760 and KBBF16

spectrum. Group II absorption bands have minor peaks except in sample

KBBF16 where they appear to be the dominant defect sites for H+ in-

corporation (Fig. 8a). O–H absorption bands at 3709, 3685 and

3645 cm−1 are attributed to hydrous minerals as serpentine, while

Please cite this article as: Baptiste, V., et al., Deformation and hydration of the lithospheric mantle beneath the Kaapvaal craton, South Africa,
Lithos (2012), doi:10.1016/j.lithos.2012.05.001
Fig. 6. Representative olivine and orthopyroxene crystal preferred orientations (CPO) of coarse-grained and sheared-peridotites. Arrows indicate the dominant dislocation glide planes in orthopyroxene and samples displaying dominant [001] glide in olivine. Lower hemisphere, equal-area stereographic projections. N is the number of measured grains. Contours at 1 multiple of a uniform distribution. CPOs were not contoured when less than 100 grains could be measured.
the band at 3675 cm\(^{-1}\) is likely to be attributed to talc (Miller et al., 1987; Post and Borer, 2000).

Sample KBJ59 from Jagersfontein, which contains high modal contents of phlogopite (14%), presents very different olivine spectra (Fig. 8a). The major peak is still located at 3571 cm\(^{-1}\), but important O–H absorption bands are also present at 3598, 3571, 3525, and 3484 cm\(^{-1}\). Similar spectra have been described in other Kaapvaal kimberlite xenoliths from Jagersfontein, Lesotho, and Wesselton (Grant et al., 2007; Kurosawa et al., 1997).

6.2. Pyroxenes and garnet

Orthopyroxene spectra are more homogeneous than the olivine ones (Fig. 8b). However, most measured spectra are strongly affected by alteration, exhibiting the O–H absorption bands at 3690 cm\(^{-1}\) and 3650 cm\(^{-1}\) usually attributed to hydrous sheet silicates (Miller et al., 1987; Skogby et al., 1990). The main absorption band associated to O–H bonds in orthopyroxene is observed at 3600 cm\(^{-1}\), followed by two peaks attributed to serpentine.
<table>
<thead>
<tr>
<th>Locality</th>
<th>Sample</th>
<th>Pressure (GPa)</th>
<th>Temperature (°C)</th>
<th>Mg #</th>
<th>Geothermobarometry data reference</th>
<th>Texture</th>
<th>Olivine CPO</th>
<th>Mineral section thickness (μm)</th>
<th>Paterson calibration (OH concentrations (H/10^6 Si))</th>
<th>Olive–Bell calibration (OH concentrations (ppm wt H2O))</th>
<th>Average OH concentrations (ppm wt H2O)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kimberley</td>
<td>FRB348</td>
<td>3.7(1)</td>
<td>938(1)</td>
<td>92.3</td>
<td>J</td>
<td>CG</td>
<td>Ortho</td>
<td>ol</td>
<td>cpx-1 508 S* 146</td>
<td>cpx-2 508 161</td>
<td>cpx-4 508 157</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>cpx-5 513 21</td>
<td>cpx-6 513 44</td>
<td>cpx-8 513 126</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-1 511 644 162</td>
<td>ol-2 511 41</td>
<td>ol-3 511 40</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-4 519 684 119</td>
<td>ol-5 519 134</td>
<td>ol-6 519 89</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-7 519 1786 124</td>
<td>ol-8 519 160</td>
<td>ol-9 149</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 519 1356 122</td>
<td>ol-12 519 160</td>
<td>ol-14 152</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 505 69 23</td>
<td>ol-2 505 67</td>
<td>ol-3 67</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 505 84 31</td>
<td>ol-5 505 84</td>
<td>ol-6 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 505 84 31</td>
<td>ol-8 505 84</td>
<td>ol-9 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 505 84 31</td>
<td>ol-12 505 84</td>
<td>ol-14 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 505 84 31</td>
<td>ol-2 505 84</td>
<td>ol-3 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 505 84 31</td>
<td>ol-5 505 84</td>
<td>ol-6 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 505 84 31</td>
<td>ol-8 505 84</td>
<td>ol-9 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 505 84 31</td>
<td>ol-12 505 84</td>
<td>ol-14 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 505 84 31</td>
<td>ol-2 505 84</td>
<td>ol-3 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 505 84 31</td>
<td>ol-5 505 84</td>
<td>ol-6 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 505 84 31</td>
<td>ol-8 505 84</td>
<td>ol-9 84</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 505 84 31</td>
<td>ol-12 505 84</td>
<td>ol-14 84</td>
</tr>
<tr>
<td>Finsch</td>
<td>FRB1501</td>
<td>4.6(3)</td>
<td>1030(2)</td>
<td>93.0</td>
<td>This study</td>
<td>CP</td>
<td>Ortho</td>
<td>ol</td>
<td>ol-5 500 112 7</td>
<td>ol-6 500 100</td>
<td>ol-7 100</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-8 500 54 3</td>
<td>ol-9 500 54</td>
<td>ol-10 54</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 500 69 4</td>
<td>ol-2 500 63</td>
<td>ol-3 63</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 500 81 5</td>
<td>ol-5 500 81</td>
<td>ol-6 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 500 81 5</td>
<td>ol-8 500 81</td>
<td>ol-9 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 500 81 5</td>
<td>ol-12 500 81</td>
<td>ol-14 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 500 69 4</td>
<td>ol-2 500 63</td>
<td>ol-3 63</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 500 81 5</td>
<td>ol-5 500 81</td>
<td>ol-6 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 500 81 5</td>
<td>ol-8 500 81</td>
<td>ol-9 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 500 81 5</td>
<td>ol-12 500 81</td>
<td>ol-14 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 500 69 4</td>
<td>ol-2 500 63</td>
<td>ol-3 63</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 500 81 5</td>
<td>ol-5 500 81</td>
<td>ol-6 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 500 81 5</td>
<td>ol-8 500 81</td>
<td>ol-9 81</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 500 81 5</td>
<td>ol-12 500 81</td>
<td>ol-14 81</td>
</tr>
<tr>
<td>Premier</td>
<td>FRB1512</td>
<td>4.7(2)</td>
<td>1058(2)</td>
<td>92.0</td>
<td>This study</td>
<td>CP</td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 520 272 17</td>
<td>ol-2 520 67</td>
<td>ol-3 67</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 400 355 22</td>
<td>ol-5 400 65</td>
<td>ol-6 65</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 400 355 22</td>
<td>ol-8 400 65</td>
<td>ol-9 65</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 400 355 22</td>
<td>ol-12 400 65</td>
<td>ol-14 65</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-1 520 272 17</td>
<td>ol-2 520 67</td>
<td>ol-3 67</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-4 400 355 22</td>
<td>ol-5 400 65</td>
<td>ol-6 65</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-7 400 355 22</td>
<td>ol-8 400 65</td>
<td>ol-9 65</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-10 400 355 22</td>
<td>ol-12 400 65</td>
<td>ol-14 65</td>
</tr>
<tr>
<td>Mothae</td>
<td>PHN1925</td>
<td>4.6(3)</td>
<td>1306(1)</td>
<td>90.7</td>
<td>This study</td>
<td>M</td>
<td>Axial [010]</td>
<td>ol</td>
<td>ol-5 390 92 6</td>
<td>ol-6 390 17</td>
<td>ol-7 17</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-8 390 206 13</td>
<td>ol-9 390 38</td>
<td>ol-10 38</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Ortho</td>
<td>ol</td>
<td>ol-12 390 206 13</td>
<td>ol-15 390 38</td>
<td>ol-16 38</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>ol-12 390 206 13</td>
<td>ol-15 390 38</td>
<td>ol-16 38</td>
</tr>
</tbody>
</table>
absorption bands centered at 3511–3520 and 3559–3541 cm$^{-1}$ (Fig. 8b). Broader minor peaks around 3410, 3300, and 3060 cm$^{-1}$ are also observed.

Clinopyroxene spectra are homogeneous (Fig. 8b). They exhibit typical patterns for mantle-derived clinopyroxenes, with major O–H absorption bands at 3640, 3535, and 3454 cm$^{-1}$ (Grant et al., 2007; Li et al., 2008; Skogby et al., 1990). Garnet crystals were usually contaminated by serpentine and t alc, displaying absorption bands 3690, 3683 and 3675 cm$^{-1}$. Spectra obtained from large, clear garnet crystals are however exempt of peaks; their spectra are flat (Fig. 8b), indicating anhydrous gt.

6.3. Phlogopite

Spectra in large undeformed phlogopite crystals from spinel peridotites KBBF18 and KB59 have three major O–H absorption bands at 3712, 3706, and 3544 cm$^{-1}$ (Fig. 8c). Minor bands can be seen at 3650, 3228 and 3126 cm$^{-1}$. Peaks at 3712 and 3650 cm$^{-1}$ are likely to be related to serpentine (Miller et al., 1987). However, an absorption band at ~3706 cm$^{-1}$ was previously observed in phlogopite by Miller et al. (1987).

6.4. OH concentrations

OH concentrations in olivine for each sample were calculated by averaging the OH concentration estimated from spectra from 2 to 7 olivine crystals in the sample (Table 3). OH concentrations in olivine could only be determined precisely for ten samples, because spectra in olivine crystals from samples KKF18, FRB348, FRB1404, and KBBF11 displayed extensive contamination by serpentine, characterized by a broad band centered at 3690 cm$^{-1}$, and covering the whole range of wavenumbers for O–H stretching bands in olivine (3750–3400 cm$^{-1}$). Examples of serpentine-contaminated and good-quality olivine spectra are presented in Fig. 9. Water content in olivine varies between 3 and 50 wt.ppm H$_2$O using the Paterson calibration (9 and 150 wt.ppm H$_2$O after applying a conversion factor of 3 for comparison with data obtained using the Bell et al. (2003) calibration, Table 3). These values are among the highest OH concentrations in olivine measured in Kaapvaal xenoliths (Fig. 10). Still, higher OH concentrations are observed in olivine megacrysts from the Monastery kimberlite (Fig. 10).

Orthopyroxene grains are significantly more affected by alteration than olivine. Most orthopyroxene spectra exhibit some contamination by serpentine and/or amphibole, hindering an accurate determination of their OH concentration. However, in four samples, OH concentrations between 49 and 96 wt.ppm H$_2$O could be measured (Table 3). Sample FRB348 was the only specimen where clinopyroxene spectra were free of any contamination. OH concentrations in those clinopyroxene vary between 146 and 161 wt.ppm H$_2$O (Table 3).

Homogeneity in OH concentrations within a grain was checked by FTIR measurements along profiles with 15 or 25 μm step sizes across randomly oriented olivine (in samples FRB1512 and KB59) and orthopyroxene crystals (in FRB1513 and FRB348). All profiles in olivine indicate homogeneous OH concentrations, except in the vicinity of serpentine-filled cracks and grain boundaries as illustrated in Fig. 9. These observations are in agreement with FTIR profiles in olivine from kimberlite xenoliths by Grant et al. (2007), Peslier et al. (2008), and Kamenetsky et al. (2008), which do not show any evidence of diffusive dehydration or hydration. In orthopyroxene, contamination by serpentine and/or amphibole was too extensive that variations in water contents within a grain could not be quantified.

7. Discussion

The studied xenoliths represent samples of the cratonic mantle at depths ranging from ~50 to 250 km (Fig. 5). Extraction ages vary. Most samples come from kimberlite pipes erupted between 124 and 85 Ma, but the Premier kimberlite sampled an earlier stage of the cratonic lithosphere evolution (2000–1650 Ma, Kramers and Smith, 1983). No significant difference in deformation microstructures, CPO, or composition is observed between xenoliths from Premier and other pipes, but one should keep in mind that only 5 xenoliths from Premier were analyzed in this study. Analysis of the microstructures allows two groups to be discriminated: coarse-grained harzburgites and sheared lherzolites, which also differ by their composition (the latter are more fertile, in average). They also display different depths of origin; sheared peridotites have systematically equilibrium pressures ≥4.6 GPa, whereas coarse-grained microstructures occur over the entire sampled depth range. In the following these two groups will be treated separately. First, the deformation processes and the relations between deformation, annealing, and metasomatism in coarse-grained peridotites will be discussed. Then, we will examine the mechanisms and conditions of deformation that produced the sheared peridotites and discuss their timing and origin. Finally, we discuss the implications of the water content measurements for the hydration state of the Kaapvaal craton root.

7.1. Deformation and annealing in the coarse-grained peridotites

Coarse-granular microstructures, characterized by plurimilimetric anhedral olivine and orthopyroxene crystals with rare intracrystalline deformation features, like subgrains in olivine or kinks in orthopyroxene

Notes to Table 3:
(1)Brey and Köhler thermometry and Barometry; (2)O’Neill and Wood (1979) thermometry; (3)Nickel and Green (1985) barometry.
(4)OH concentrations from this study calculated using the calibration of Paterson (1982) have ±30% error (Kohlstedt et al., 1996); (5)OH concentrations corrected by a factor of 3 according to Bell et al. (2003). S = serpentine contamination prevented OH concentration estimations.


Texture: CG = Coarse-grained peridotite; MY = Mylonite.
CPO types: ortho = orthorhombic, and axial [100] crystal-preferred orientations; see main text and Fig. 6 for details.
peridotites are sampled by Jagersfontein, whereas Kimberley peridotites have finer-grained and least annealed microstructures, which preserve a weak foliation, marked by the elongation of olivine and orthopyroxene crystals (aspect ratios range from 1:2 to 1:4, Fig. 2b). Coarse-porphyroclastic microstructures are also more common in Kimberley (Table 2). This suggests either a stronger annealing or lower stress deformation in the lithospheric mantle beneath Jagersfontein. The remaining sites show, however, intermediate textures with no systematic variation.

In both coarse-granular and coarse-porphyroclastic xenoliths, the well-defined olivine CPO implies that dislocation creep was the dominant deformation process. Three olivine CPO patterns were identified: (1) axial-[100], (2) axial-[010], and (3) orthorhombic. Or-thorhombic olivine CPOs are observed over the whole pressure range, while axial-[100] CPOs tend to concentrate on the top of the column (2.4–3.8 GPa). These olivine CPO patterns, characterized by a stronger concentration of [100] axes relative to [010] or [001] are coherent with simple or pure shear with dominant activation of high-temperature, low pressure, dry (010)][100] or [0K][100] slip systems, respectively (cf. Tommasi et al., 2000).

Axial-[010] olivine CPO, characterized by a point concentration of [010] and girdle distributions of [100] and [001] are however common in xenoliths equilibrated between 3.5 and 4.7 GPa. Such olivine CPO pattern may result from transpressional deformation (Tommasi et al., 1999), from changes in dominant glide system due to variations in physical parameters (differential stress, pressure, or water contents in olivine) during deformation, or from recrystallization (Falus et al., 2011; Tommasi et al., 2008). However, transpressional deformation should also result in dispersion of orthopyroxene [001] axes in the foliation plane, which is not observed in most studied peridotites, which show predominantly point distribution of orthopyroxene [001] axes in xenoliths equilibrated between 3.5 and 4.7 GPa (Raterron et al., 2007). Equilibrium pressures for these CPOs, characterized by a stronger concentration of [100] axes relative to [010] or [001] are coherent with simple or pure shear with dominant activation of high-temperature, low pressure, dry (010)][100] or [0K][100] slip systems, respectively (cf. Tommasi et al., 2000).

Occurrence of axial-[010] olivine CPO patterns in peridotites equilibrated at pressures 3.5–4.7 GPa might suggest that these CPOs may record a change in deformation mechanism associated with an increase in confining pressure or in water contents, as the latter also increase in this depth range (Fig. 10). Experimental studies on a Griggs press (confining pressures ~2 GPa) suggested that high water fugacities and differential stresses may induce a transition from dominant [100] to dominant [001] glide (cf. review in Jung et al., 2006). Yet, according to these experiments, the water contents in olivine measured in the present study (Table 3) are not high enough to induce dominant [001] glide; they rather result in the so-called E-type fabric, characterized by [100] and [001] axes aligned parallel to the lineation and normal to the foliation, respectively. The lack of clear foliation and weak CPO (supplementary online material) in the coarse-granular peridotites for which water contents in olivine could be measured in this study does not allow discrimination of which is the dominant glide plane: (010) or (001).

Transition from dominant [100] to dominant [001] glide was also observed in experimental deformation at high confining pressures (Couvy et al., 2004); the pressure at which this transition occurred varied between experiments, ranging from 3 to 3.5 GPa (Jung et al., 2008) to ~7 GPa (Raterron et al., 2007). Equilibrium pressures for samples with axial-[010] olivine CPO (between 3.5 and 4.7 GPa) are in the same range as the Jung et al. (2008) values. Yet, orthorhombic olivine CPOs are even more common at these depths (Fig. 7b). Soile two samples have olivine [001] axes aligned parallel to the maximum concentration of orthopyroxene [001] axes, suggesting dominant
A alternative explanation for the axial-[010] olivine CPO patterns in the coarse-grained peridotites from the Kaapvaal is dispersion of the [100] and [001] in response to recovery and subgrain rotation recrystallization (Falus et al., 2011). This interpretation is consistent with the observation that low-angle misorientations are dominantlyaccommodated by rotations around the [010] axis. CPO dispersion during static recrystallization may also explain the predominance of weak olivine CPO in the Kaapvaal craton (Fig. 7a). Low olivine J-indexes also predominate in peridotite xenoliths from the Siberian craton (Bascou et al., 2011). In general, cratonic peridotites yield lower J-indexes than peridotites from other tectonic settings (Fig. 11). However, although weakening of olivine CPO associated with modal metasomatism has been described in other environments (Bascou et al., 2008; Morales and Tommasi, 2011; Tommasi et al., 2004, 2008), in the present dataset, there is no clear relation between the olivine J-index, modal contents, or olivine mg# (Table 2).

Orthopyroxene CPOs are consistent with deformation by dislocation creep with activation of [001][100] and [001][010] slip systems. They are always correlated to olivine fabrics, meaning that both minerals underwent the same deformation. Well-defined CPOs are observed even in harzburgites showing high modal orthopyroxene, suggesting that they were deformed during or after the metasomatism leading to Si-enrichment. This interpretation is consistent with the observation of kinked interstitial orthopyroxene (Fig. 3a).

7.2. Evidence for metasomatism and timing relative to deformation

Textual evidence for reactive percolation of Si-rich fluids and/or melts is common in the coarse-granular harzburgites. It encompasses: (1) interpenetrating orthopyroxene and olivine boundaries (Fig. 2a–d), (2) small interstitial orthopyroxene crystals with cusped grain boundaries (Fig. 3a), (3) interstitial garnet in peridotite FRB1339 (Fig. 3d), (4) crystallization of small, interstitial, Cr-rich clinopyroxene, often associated with spinel and phlogopite (Fig. 3d), and (5) phlogopite rimming garnets or as interstitial crystals. Observation (1) may result either from orthopyroxene consumption during the extensive partial melting event that produced the most refractory olivines (Boyd and Mertzman, 1987) or from orthopyroxene crystallization at the expenses of olivine in response to reactions with Si-rich aqueous fluids or hydrous melts (Bell et al., 2005; Kelemen et al., 1998; Wasch et al., 2009). In contrast, observation (2) points to late crystallization of orthopyroxene and is consistent with the excess modal content of orthopyroxene, which characterizes the Kaapvaal harzburgites (Bell et al., 2005). The remaining observations point to partial crystallization or reaction with percolating melts at high pressure (3) and with K-rich compositions (4 and 5).

Analysis of the microstructures may help in constraining the relative timing between deformation and episodes of reactive fluid percolation, even if some observations are ambiguous. Kinks in interstitial orthopyroxene (Fig. 3a) indicate pre- to syn-kinematic silica enrichment. However, undeformed orthopyroxenites with clear interstitial habitus are also observed in some samples, suggesting strongly that Si-enrichment may have continued post-kinematically. As discussed in the previous section, olivine and orthopyroxene CPOs are consistent, suggesting that both minerals recorded the same deformation, even in orthopyroxene-rich harzburgites, implying pre- to syn-kinematic metasomatism. Orthopyroxene CPO is however usually weak (the strongest values in Table 2 probably result from overestimation of the orthopyroxene CPO strength due to measurement of too few crystals), implying that post-kinematic orthopyroxene addition is also possible.

Phlogopite and clinopyroxene enrichment, on the other hand, is essentially post-kinematic, as indicated by the interstitial shapes and undeformed character of these minerals (Fig. 3d). The actual timing and processes responsible for this late metasomatism by K-rich fluids or melts (Bell et al., 2005; Grégoire et al., 2003; Griffin et al., 2003), are however still unclear; it may have happened at any point between the last deformation episode and shortly before extraction by the kimberlitic magmatism. In sample KBBF18, alignment of undeformed phlogopite and clinopyroxene parallel to the foliation (Fig. 3d) suggests nevertheless that the pre-existing deformation structure may have controlled the melt infiltration.

7.3. Sheared peridotites

Sheared peridotites are characterized by high equilibrium temperatures (between 1000 and 1300 °C) and relatively fertile compositions as pointed out by Nixon et al. (1981). They probably represent ancient coarse-granular rocks that underwent extensive dynamic and static recrystallization. Harzburgite PHN2760, with its very unusual texture (Fig. 4d), may indeed be an illustration of the first steps of formation of the sheared peridotites, which was followed by extensive fracturing and low-temperature serpentinitization due to reaction with the kimberlitic melt during extrusion.

The asymmetry of the porphyroclasts in the mylonite implies simple shear deformation. Moreover, the fine recrystallized grain sizes indicate deformation under high deviatoric stresses and deformation rates (Austin and Evans, 2007), in particular considering the high equilibrium temperatures of these peridotites. Both orthopyroxene and olivine underwent dynamic recrystallization but the degree of recrystallization and the recrystallized grain size are larger in olivine than in orthopyroxene, consistent with the higher strength of orthopyroxene relative to olivine. Development of shear bands along fine-grained recrystallized orthopyroxene-rich layers suggests, that more effective grain refinement in the higher strength orthopyroxene-rich domains lead to strain localization. Boullier and Gueguen (1975) proposed that the olivine-rich domains deformed by dislocation creep whereas the fine-grained recrystallized orthopyroxene-rich layers deformed by grain sensitive creep.

Evidence of plastic deformation by dislocation creep, such as undulose extinction, subgrain boundaries, and recrystallization tails, has also been identified in mosaic and fluidal mosaic peridotites. However, the presence of polygonal recovered olivine crystals in the matrix implies that some annealing followed the deformation. The variable grain size and amount of polygonal crystals in the matrix may reflect different temperatures, annealing times, or variable
influence of fluids on the annealing process. Euhedral olivine crystals free of any deformation features (usually called tablets) are commonly observed on both strained porphyroclasts and the matrix of mosaic and fluidal mosaic peridotites. Such euhedral crystals have been often described in sheared xenoliths from South Africa (e.g., Boullier and Nicolas, 1975; Drury and Van Roermund, 1989; Skemer and Karato, 2008). Drury and Van Roermund (1989) proposed that these crystals record fluid-assisted static recrystallization, where a thin fluid film along grain boundaries enhances grain boundary mobility. Fluids may also have played an essential role in the formation of the mosaic texture, which may represent a more developed stage of fluid-assisted static recrystallization. The preservation of the fine-grained texture and deformation features in the orthopyroxene-rich shear bands may be explained by slower grain boundary migration in orthopyroxene in comparison to olivine (Skemer and Karato, 2008). Mosaic and fluidal mosaic peridotites may thus derive from prophyroplastic and mylonitic peridotites to which annealing is superimposed, respectively (Boullier and Nicolas, 1975).

Sheared peridotites display either orthorhombic or bimodal olivine CPO. The orthorhombic CPOs are consistent with deformation by dislocation creep activating high-temperature, low pressure (010)[100] slip system. The axial-[010] with a bimodal distribution of [100] and [001] axes suggest, on the other hand, simultaneous activation of the (010)[100] and (010)[001] slip systems. Similar olivine CPO were described in deep mantle xenoliths from the Tanzanian craton (Vauchez et al., 2005) and have been interpreted as resulting from activation of the two slip directions at high pressure and probably high stresses. This interpretation is consistent with the observation that bimodal olivine CPO are characteristic of sheared peridotites equilibrated at pressures >4.5 GPa (Fig. 7c). Olivine CPO strength is always low (Fig. 7a), probably due to the extensive recrystallization (Faluss et al., 2011). Orthopyroxene CPOs are coherent with deformation by dislocation creep with contribution of both [001] [110] and [001] [010] systems. Clinopyroxene CPO is consistent with deformation by dislocation creep, with dominant activation of the [001] [110] systems (Bascou et al., 2002). Both pyroxenes CPOs are correlated with the olivine CPO (Fig. 6 and online supplementary material), indicating that all major rock-forming minerals underwent the deformation producing the mylonitic microstructure. Indeed, clinopyroxene occurs as strained porphyroclasts and has CPO coherent with the olivine and orthopyroxene, indicating pre-mylonitization reorientation. Finally, alignment of recrystallized phlogopite in the mylonitic foliation in KBBF11 indicates that the mylonitization postdates potassic metamorphism. This observation contrasts with the pre-kinematic character of phlogopite in coarse-grained harzburgites, suggesting that mylonitization corresponds to a later deformation event.

In summary, microstructures and CPO in the sheared peridotites are consistent with post-metasomatism deformation by dislocation creep under high stress and strain rates. Fluidal mosaic and mosaic microstructures may be explained by subsequent fluid-assisted static recrystallization that may be related to the kimberlitic magmatism. The processes leading to this deformation have been widely debated in the literature. The kinked geotherm obtained from thermobarometry in mantle xenoliths from the Kaapvaal made Boyd and Nixon (1975) propose that the sheared peridotites may result from localized shearing at the lithosphere–asthenosphere boundary (LAB). This interpretation was re-proposed by Kennedy et al. (2002) for the sheared peridotites of the Slave craton. However, already in the 70s, Goetze (1975) highlighted that the observed geotherm disturbance is not consistent with shear heating and that the strain rates estimated from the microstructures and equilibrium temperatures of these rocks are too high to be produced by plate tectonics. Similar conclusions were reached by Boullier (1977) and more recently by Skemer and Karato (2008). Our observations are consistent with these conclusions, favoring the model proposed by Green and Gueguen (1974), who viewed the deformation that produced the sheared peridotites as due to ascending kimberlitic diapirs, instead of a quasi-stationary mantle flow at the LAB.

7.4. Hydrostatic state of the Kaapvaal mantle lithosphere

OH concentrations in olivine obtained in this study are highly variable, however they show a general increase with pressure up to 5 GPa and a sharp decrease for pressures >5.9 GPa (Fig. 12, Table 3). Maximum water contents in olivine measured in this study are among the highest ones measured in mantle peridotite xenoliths, reaching values close to those measured in kimberlite-borne megacrysts (Fig. 10). In samples equilibrated at pressures higher than 5.9 GPa, reliable estimates of water contents in olivine could be obtained for two samples, one coarse-grained peridotite and a mylonite. Both show <10 wt. ppm H2O (Table 3), consistently with previous data in four other deep Kaapvaal peridotites by Peslier et al. (2010).

The observed increase in OH concentrations in olivine with depth up to 5 GPa is consistent with the increase of water solubility with increasing $f_{H_2O}$ and, hence, with pressure (Kohlstedt et al., 1996). Measured water contents remain, however, well below the olivine solubility curves of Hirschmann et al. (2005) and Kohlstedt et al. (1998), suggesting that even the most water-rich peridotites in the Kaapvaal are undersaturated in water. However, these water solubility curves in olivine only consider the effect of pressure. They do not include the effects of temperature, iron content in olivine, or the change of water fugacity with increasing solute dissolution at high pressures and temperatures and probably overestimate the solubility in the deepest part of the cratonic root (Bali et al., 2008; Ferot, 2011). Most analyzed peridotites contain phlogopite. Since hydrogen is compatible in sheet silicates, crystallization of phlogopite does not require prior saturation of the nominally anhydrous minerals (NAMs) that compose the peridotite. A recent experimental study shows that water solubility curves of mantle minerals are strongly modified by the presence of pargasite (Green et al., 2010). A similar behavior might be expected for olivine co-existing with phlogopite, but the solubility of water in NAMs in a phlogopite-bearing assemblage has not been quantified yet. In the present dataset, OH concentrations in olivine are uncorrelated with the presence or absence of phlogopite (Tables 2 and 3), although less IR peaks are observed in the phlogopite-rich sample KBJS9 (Fig. 8).

Are the measured values in olivine representative of OH concentrations at different depths in the Kaapvaal mantle? Since H behaves...
as an incompatible element during partial melting (Bolfan-Casanova, 2005; Dixon et al., 2002; Hirschmann et al., 2005), olivines with high forsterite contents are expected to have low OH concentrations. This relation is not observed in the Kaapvaal peridotites; high OH concentrations were measured in olivines with mg# ranging from 90 to 94 (Fig. 13). This implies that the high water contents in olivine from intermediate depths in the cratonic mantle record re-hydration after the main partial melting episode that produced the mantle root. This conclusion brings additional questions. When and how did this water addition occur? What is its spatial extent?

High OH contents in olivine may result from hydration: (A) by solid-state diffusion of hydrogen from a volatile-rich asthenosphere, (B) during transport by the volatiles-rich kimberlitic magma, or (C) during metasomatism of the craton root by water-rich fluids or melts (Fig. 14). Re-hydration of lithospheric mantle in a continuous way by solid-state diffusion of hydrogen from the asthenosphere is not consistent with the observed low OH contents in olivine from peridotites equilibrated at >5.9 GPa. Moreover, even if the fast hydrogen flux along grain boundaries is taken into account (Demouchy, 2010), solid-state diffusion is a process too slow. For an average grain size of 1 μm, which is much smaller than even the smallest grains in the mylonites, and an effective hydrogen diffusion coefficient of 4.48 × 10⁻⁹ m²/s at 1250 °C (Demouchy, 2010) more than 3 Gy would be needed to re-hydrate the entire cratonic root, and if a more representative grain size of 3 mm is considered, the H diffusion front would have progressed by less than 10 km over 3 Gy.

Scenario (B), that is, hydration during xenolith extraction by the kimberlites, is based on the observation that olivine megacrysts from kimberlites have high OH⁻ contents (Fig. 10), suggesting that the kimberlite magma is volatile-rich and, perhaps, water-saturated. It is also consistent with experimental data and with dehydration FTIR profiles in olivines from basalt-borne peridotite xenoliths, which both show extremely fast hydrogen diffusion in olivine at high temperatures (Demouchy and Mackwell, 2006; Demouchy et al., 2006; Peslier and Luhr, 2006). Hydrogen addition to xenolithic minerals may therefore occur during the transport of the xenoliths, even if FTIR profiles do not record these exchanges (Kamenetsky et al., 2008).

In scenario (C), the re-hydration of the cratonic mantle is proposed to be associated with metasomatism by water-rich fluids or melts, for which there is abundant modal and geochemical evidence, like the Si-enrichment (Fig. 3a; Bell et al., 2005; Kelemen et al., 1998; Simon et al., 2007; Wasch et al., 2009) or the crystallization of secondary garnet, clinopyroxene and phlogopite (Fig. 3b and d; Grégoire et al., 2003; Simon et al., 2007). In this case, hydration may be both spatially and temporally heterogeneous, since it may be related to many distinct metasomatic episodes: from (i) early metasomatism by Si- and volatile-rich fluids that produced the high modal content of orthopyroxene in Kaapvaal harzburgites (Bell et al., 2005), which was proposed, based on Lu–Hf and Sm–Nd model ages, to be multiple episodes between 1.3 Ga and 600 Ma (Wasch et al., 2009), to (ii) late metasomatism by K-rich melts, which has caused phlogopite crystallization (Grégoire et al., 2003; Simon et al., 2007). This spatial and temporal variability is coherent with the OH⁻ concentration heterogeneity of the Kaapvaal olivines (Fig. 10).

In scenarios B and C, the observed vertical variation in OH⁻ concentrations may be accounted by considering the relative changes in water and carbon activity with pressure in volatile-rich (C–O–H) fluids, which at pressures > 5 GPa, have high carbon and low water activities (Dixon et al., 1995; Newman and Lowenstern, 2002) and the variation in the mantle redox state with depth, as proposed by Peslier et al. (2010). If metasomatism is associated with percolation of basaltic magmas, change in the melt composition with progressive enrichment in volatiles during reactive percolation (O’Nions and McKenzie, 1988) may also account for the contrast in OH⁻ contents between the deepest and intermediate depth peridotites.

The two last scenarios have different implications for the cratonic root stability. In the first one, the root is essentially dry and the high water contents in olivine measured in the present study and in previous studies result from “contamination” during the ascent in the kimberlitic magma. In the second one, metasomatism may have hydrated and, hence, weakened at least parts of the cratonic root after...
its formation. The very nature of xenolithic sampling by kimberlites prevents the evaluation of the actual volumes of the mantle affected by the metasomatic processes. Comparison between the magnetotelluric electrical conductivity data in a station close to Jagersfontein and the MT signal modeled using a constant water concentration in the cratonic mantle or a variation in water contents with depth similar to the one observed in this study (cf. the Fig. 8 in Fullea et al., 2011) suggests however that the vertical variation of water contents in olivine observed in the present study and in Peslier et al. (2010) may be representative of the present-day hydration state of the Kaapvaal mantle, favoring the last scenario and implying that metasomatism resulted in extensive re-hydration of the cratonic mantle at intermediate depths.

The annealed microstructures of Kaapvaal peridotites indicate however that rehydration was not followed by remobilization of the cratonic root. Several explanations may be proposed: (i) the extensive hydration of the intermediate depths of the cratonic root is recent and its mechanical effects have not been felt yet, (ii) the distribution of the hydrated domains is too heterogeneous, or (iii) the dry layer at its base (Fig. 10) protects the craton root from thermo-mechanical erosion by the convective mantle as proposed by Morency et al. (2002). The latter hypothesis is appealing, but convective models testing the stability of a block with thickened lithosphere and [001] glide on the (010) plane, probably due to high pressure and high equilibrium temperatures imply deformation under high stresses.

Lack of correlation between olivine mg# and water content indicates that the high water contents in olivine record re-hydration after formation of the refractory mantle root. Hydration of olivine may have occurred during either extraction of the peridotites by volatile-rich kimberlites or metasomatism by water-rich fluids/melts. Magnetotelluric electrical conductivity data in the central Kaapvaal is, however, best explained by models with a vertical variation of water contents in olivine similar to the one measured here, suggesting that the observed water contents in olivine may be representative of the present-day hydration state of the Kaapvaal mantle, favoring the metasomatism scenario. The predominance of highly annealed microstructures in the cratonic root indicates nevertheless that this re-hydration did not result in remobilization of the cratonic root.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.lithos.2012.05.001.

Acknowledgments

The authors thank A. Vauchez, J-M. Dautria, and S. Keshav for helpful discussions. C. Nevada and D. Delmas are thanked for providing high-quality polishing of sections for EBSD measurements. Electron microprobe analyses were carried out with the help of C. Merlet at the Service Microscosonde Sud, Université Montpellier 2. FTIR analyses were performed with the assistance of D. Maurin at the Lab. Colloids, Verre et Nanomatériaux, at Université Montpellier 2, France. AT has been funded by CRYSTAL2PLATE, a EU FP7-funded Marie Curie Action under grant agreement PITN-GA-2008-215353. The EBSD-SEM national facility in Montpellier is supported by the Institut National de Sciences de l’Univers (INSU) du Centre National de la Recherche Scientifique (CNRS), France and by the Conseil Régional Languedoc-Roussillon, France.

8. Conclusions

Coarse-granular refractory harzburgites predominate in the Kaapvaal mantle, being observed in all studied pipes at all depths. Their highly recovered microstructures and very coarse grain sizes contrast with the olivine and orthopyroxene CPO, indicating that deformation by dislocation creep was followed by effective annealing. This supports an early deformation episode, possibly associated to the formation of the cratonic root in the Archean, followed by slow cooling and a long quiescence time. Fluid-enhanced grain boundary migration during the multiple metasomatic events that affected the Kaapvaal mantle may also have favored annealing. Coarse-porphyroclastic peridotites are rarer, but present at all depths in Kimberley and in deep levels in Finsch and Premier. They represent domains less annealed or deformed shortly before extraction. Sheared peridotites with mylonitic, fluidal or mosaic microstructures, are common below 140 km depth. Their fine recrystallized grain sizes and high equilibrium temperatures imply deformation under high stress and high strain rates, not consistent with a quasi-stationary mantle flow at the lithosphere–asthenosphere boundary.

Textures and compositions of the studied xenoliths provide evidence for multiple metasomatic events with a spatially heterogeneous distribution. Analysis of the microstructures and comparison between the olivine and secondary mineral CPOs indicate that silica-enrichment episodes, which resulted in high modal orthopyroxene, were pre- to post-kinematic, and that metasomatism by K-rich fluids or melts and refertilization was post-kinematic. Deformation of phlogopite and clinopyroxene in the sheared peridotites implies that mylonitization post-dated all metasomatic events.

Water contents in olivine are strongly variable, but tend to increase with depth up to 150 km, where they are among the highest measured in mantle peridotites (~150 wt. ppm H2O; Bell et al., 2003 calibration). The deepest analyzed samples are, in contrast, almost dry. High water contents in coarse-grained peridotites are not accompanied by dominant activation of [001] glide. The bimodal olivine CPO in sheared peridotites indicates nevertheless activation of both [100] and [001] glide on the (010) plane, probably due to high pressure and stresses.

References


