Redox preconditioning deep cratonic lithosphere for kimberlite genesis – evidence from the central Slave Craton

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We present the first oxygen fugacity ($f_{O_2}$) profile through the cratonic lithospheric mantle under the Panda kimberlite (Ekati Diamond Mine) in the Lac de Gras kimberlite field, central Slave Craton, northern Canada. Combining this data with new and existing data from garnet peridotite xenoliths from an almost coeval kimberlite (A154-N) at the nearby Diavik Diamond Mine demonstrates that the oxygen fugacity of the Slave cratonic mantle varies by several orders of magnitude as a function of depth and over short lateral distances. The lower part of the diamond-bearing Slave lithosphere (>120–130 km deep) has been oxidized by up to 4 log units in $f_{O_2}$, and this is clearly linked to metasomatic enrichment. Such coupled enrichment and oxidation was likely caused by infiltrating carbonate-bearing, hydrous, silicate melts in the presence of diamond, a process proposed to be critical for “pre-conditioning” deep lithospheric mantle and rendering it suitable for later generation of kimberlites and other SiO$_2$-undersaturated magmas.

The $f_{O_2}$ of the Earth's deep interior is critically important, influencing diverse processes in the solid and volatile cycles of our planet. It controls the speciation of volatile components (CHONS) in the mantle, including diamond/graphite versus carbonate stability. This profoundly influences melting temperatures and types of partial melts of mantle rocks and the nature of mantle metasomatism. It affects volatile solubilities in magmas and hence outgassing of CHONS-volatiles and so was influential in the formation and nature of the atmosphere. Knowledge of the distribution of the mantle’s $f_{O_2}$ with depth, location and time is critical for understanding large-scale volatile cycles and fluxes between the crust, ocean and atmosphere, and deep Earth (mantle, core)1.

In the absence of partial melts or fluids, the $f_{O_2}$ of the Earth’s upper mantle is controlled internally by redox sensitive reactions involving Fe-bearing mineral components in spinel, garnet and pyroxenes in which Fe has variable oxidation states. Because of the positive molar volume changes of these reactions2–4, $f_{O_2}$ should decrease steadily with increasing pressure (depth) in the upper ≈250 km of the peridotite-dominated mantle. At pressures around 8–9 GPa (≈250–300 km deep) the depth-$f_{O_2}$ trend should intersect the Ni precipitation curve, and an FeNi metallic alloy is expected to form4. The $f_{O_2}$ of peridotite will then remain close to that of the Ni precipitation curve (≈IW to IW-1 log unit)3 down to the transition zone at ≈410 km depth4.

Experimental calibrations of redox controlling reactions in garnet peridotite assemblages2,3, coupled with determinations of the Fe$^{3+}$/Fe of garnet from peridotite xenoliths transported by kimberlite magmas to the surface from depths as great as 220 km (~7 GPa), enable determination of the $f_{O_2}$ of the cratonic mantle lithosphere. There is now data from several different cratons6–13. We have determined the oxygen fugacities recorded by garnet peridotite xenoliths hosted by two nearly coeval and diamondiferous kimberlites from the geographically close Ekati and Diavik mine leases in the central Slave Craton, Canada. In combination with conventional thermobarometry, this allows construction of the first $f_{O_2}$

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profile through the cratonic lithospheric mantle under the Ekati Diamond Mine and demonstrates that the $f_{O_2}$ of the Slave cratonic mantle can vary by several orders of magnitude as a function of depth and over short lateral distances. The implications for kimberlite genesis are then discussed. No redox data has previously been reported from xenoliths from the Panda kimberlite.

**Results and Calculations**

**Sample locations and other details.** Fresh garnet peridotite xenoliths recovered from the Panda kimberlite on the Ekati Diamond Mine lease and from the A154-North (A154-N) pipe on the nearby Diavik lease were investigated. The Panda samples consist of 30 fragments of peridotite approximately 1 cm across, derived from fragmentation of larger xenoliths during processing at the mine. In all cases, they contain relict, fresh primary mineral phases. Major and minor element compositions for all mineral phases were presented by Menzies et al.14. The Diavik suite includes 10 new and very fresh garnet lherzolite xenoliths recovered from the A154-N kimberlite pipe at the Diavik Diamond Mine in Lac de Gras, Northwest Territories, Canada. The data from these new samples is presented in Supplementary Tables 1–5 and is supplemented by earlier data from other garnet peridotite xenoliths from the same pipe8. The two host kimberlites were emplaced at 53.3 ± 0.6 Ma (Panda)15 and 56.0 ± 0.7 Ma (A154-N)16, and are located about 30 km apart near the edge of Lac de Gras in the central Slave Craton (see Fig. 1 of Creighton et al.8 for a location map).

**Major, minor and trace element compositions of mineral phases.** Electron microprobe analyses of the major and minor element abundances and laser-ablation inductively couple plasma mass spectroscopy (LA-ICPMS) analyses of trace element abundances in constituent phases in the new A154-N garnet lherzolite xenoliths are presented in Supplementary Tables 1–4. The data presented are averages of multiple analyses of each phase in each sample. In Supplementary Table 4 $Fe^{3+}/\Sigma Fe$ data for garnets from the A154-N (Diavik) suite, determined by Mössbauer spectroscopy, are presented and the wt% oxide and cation analyses have incorporated..

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**Figure 1.** Garnet (A) Ti, (B) Yb, (C) Y and (D) Ga vs. pressure (GPa) for the Panda and A154-N samples. Data for some A154-N samples are from Creighton et al.8. Depleted samples contain garnet with <200 ppm Ti, following Griffin and Ryan8.
this data. In Supplementary Table 5 Fe\(^{3+}/\sum\)Fe data for garnets from the Panda (Ekati) suite, determined by X-ray Absorption Near-Edge Structure spectroscopy (XANES), are presented. Major and minor element chemistry for garnets from Menzies et al.\(^\text{13}\) are also re-presented, but with Fe\(_2\)O\(_3\) and Fe\(^{3+}\) included in the wt% oxide and cation analyses respectively, calculated using the XANES measurements. All XANES Fe\(^{3+}/\sum\)Fe measurements are presented. New LA-ICPMS garnet trace element data are also presented.

Major element chemistry of mineral phases in the Panda samples and in some of the A154-N samples has been comprehensively described previously\(^\text{8,14}\). The mineral chemistry of the new A154-N samples is typical of other fertile garnet peridotite xenoliths from various locations around the world. Olivine Mg\# (where Mg\# = 100×Mg/\([Mg+Fe]\)) varies from 90.0 to 92.9. All new A154-N samples are lherzolites and clinopyroxene is Cr-diopside-rich with Mg\# ranging from 90.5 to 95.2, Na\(_2\)O from 0.94 to 1.97 wt% and TiO\(_2\) up to 0.23 wt%. Orthopyroxene has Mg\# from 91.2 to 93.6, CaO from 0.22 to 1.09 wt% and Al\(_2\)O\(_3\) from 0.41 to 0.70 wt%. Garnets contain from 1.77 to 9.90 wt% Cr\(_2\)O\(_3\), 4.32 to 6.95 wt% CaO and are classified as lherzolitic G9 types\(^\text{17}\). CaO and Cr\(_2\)O\(_3\) are very well correlated. Fe\(^{3+}/\sum\)Fe was determined using Mössbauer spectroscopy and ranges from 0.022 to 0.105, similar to other suites of garnet peridotites\(^\text{5,10,18}\). Mineral grains from all samples are homogenous within analytical uncertainty on an intra- and intergrain basis.

Trace element abundances in mineral phases present in the samples from the Panda kimberlite and the new samples from the A154-N kimberlite were determined using LA-ICPMS (see Methods) and are presented in Supplementary Tables 2, 4 and 5. Data for the remaining A154-N samples of Creighton et al. has not been presented\(^\text{8}\).

**Thermobarometry of the garnet peridotites.** Based on conventional thermobarometry\(^\text{19,20}\), the Panda samples equilibrated at pressures from 2.3 ± 0.3 to 6.9 ± 0.3 GPa and temperatures from 717 ± 20 to 1271 ± 20°C. The A154-N samples equilibrated at pressures from 2.0 ± 0.3 to 5.8 ± 0.3 GPa and temperatures from 457 ± 20 to 1269 ± 20°C (Supplementary Table 6). The coeval and geographically close nature of the Panda and A154-N kimberlites and the sampling of material over a depth interval that includes most of the vertical lithospheric section, afford an excellent opportunity to assess lateral and vertical heterogeneity in cratonic lithospheric mantle down to nearly 7 GPa (≈200 km depth).

**Trace element mineral chemistry of the garnet peridotites.** Abundances of incompatible trace elements in garnet (and clinopyroxene) are presented in Supplementary Tables 2, 4 and 5. Abundances of trace elements such as Y, Yb, Ga and Ti in garnet have long been recognized as indicators of the relative depleted versus enriched nature of garnet peridotite xenoliths\(^\text{7}\). When abundances of these elements in garnet are plotted against pressure (P), it is clear that the Slave lithosphere under the A154-N pipe is highly depleted at depths shallower than ≈135 km (4.5 GPa) (Fig. 1). At greater depths, the garnets contain a much larger range and mostly higher abundances of these elements, indicating that the material present is variably and often strongly enriched. Under the Panda pipe at the nearby Ekati Diamond Mine, the lithospheric architecture is broadly similar to that under the nearby A154-N pipe, although the proportion of samples containing enriched garnet is lower. Also, at Ekati more depleted material was sampled from P > 4.5 GPa than was the case at Diavik. As for A154-N, Panda garnets derived from P ≤ 4.5 GPa are exclusively depleted in these trace elements relative to most higher-pressure garnets (Fig. 1).

Cratonic lithospheric garnet normalized Rare Earth Element patterns are distinctive – “sinusoidal” patterns are usually found in garnets in more depleted lithologies (harzburgites) but “normal” patterns are usually in more fertile lherzolitic lithologies\(^\text{22}\). This is also observed in the current samples in that the shallower, depleted samples mostly exhibit “sinusoidal” patterns and the deeper, enriched samples have “normal” patterns (Fig. 2). The Slave peridotitic garnet compositions therefore indicate strikingly that the upper part of the lithosphere (P < 4.5 GPa) is depleted, but at P > 4.5 GPa, both depleted and variably enriched material is present. Thus, the central Slave cratonic lithosphere is characterized by two layers, the upper depleted and the lower containing both enriched and depleted components, with a remarkably sharp contact between the two at a depth of ≈140 km. Broadly similar inferences were made by Griffin et al.\(^\text{23}\) based on garnet and Cr-spinel heavy mineral concentrates from Cretaceous-Tertiary central Slave Craton kimberlites.

**Oxygen fugacity determinations.** Oxygen fugacity conditions experienced by the xenoliths prior to entrainment in the host kimberlites were calculated using the garnet peridotite oxygen barometer of Stagno et al.\(^\text{3}\). Relative to the fayalite-magnetite-quartz (FMQ) redox buffer calculated for the pressure-temperature (PT) conditions of equilibration of each mantle xenolith, we obtained ΔfO\(_2\) [FMQ] from −0.47 to −3.89 log units (±0.6 log units) for the Panda suite and 0.18 to −3.36 log units for the new A154-N suite (Supplementary Table 6). Combining the new A154-N data with those from Creighton et al.\(^\text{1}\), recalculated using the Stagno et al.\(^\text{3}\) calibration, defines a range of ΔfO\(_2\) [FMQ] from +0.95 to −4.11 log units over a PT range of 2.01 to 6.85 GPa and 457 to 1346 °C for the full Diavik suite.

**Discussion**

In Fig. 3, the variation in fO\(_2\) as a function of depth for the A154-N and Panda samples is plotted, revealing several features:

1. ΔfO\(_2\) [FMQ] varies by up to ≈4 log units at almost any depth; Lateral heterogeneity in oxygen fugacity of 3 log units has also been observed in the northern part of the Slave Craton using eclogite and pyroxenite xenoliths\(^\text{24}\).

2. The shallow, depleted Panda samples and some depleted A154-N samples define a broad trend of decreasing fO\(_2\) with increasing pressure down to 5.0 GPa.
At P > 4.5 GPa, there is a trend to more oxidized values than expected by deeper extrapolation of the lower pressure fO2-P trend (grey field on Fig. 3). Samples defining this oxidized zone (yellow field on Fig. 3) include nearly all enriched A154-N samples, all enriched Panda samples and 5 of the depleted Panda samples. These 5 depleted Panda samples were classified as depleted on the basis of their low garnet Ti contents (<200 ppm), but they are relatively enriched in Yb, Y and Ga (Fig. 1). Hence, this deep, oxidized domain corresponds quite precisely to the trace element enriched layer in the lower part of the lithosphere and to the diamond stability field defined in P-T-fO2 space, although 8 of the A154-N samples of Creighton et al. lie in the carbonate stability field (Fig. 3).

Figure 2. Primitive Mantle normalized garnet REE patterns from depleted and enriched garnets from the Panda and new A154-N samples. Depleted garnets contain <200 ppm Ti. Nearly all depleted garnets have strongly sinusoidal REE patterns, whereas the majority of enriched garnets have normal patterns. Sample numbers are indicated in the key. In the case of the Diavik samples, the samples numbers quoted in the key correspond to the last three digits of the full sample numbers listed in Supplementary Table 4.
Thus, although the central Slave cratonic lithosphere at \( P < 4.5 \text{ GPa} \) decreases in \( f_{\text{O}_2} \) with increasing pressure as predicted by thermodynamic considerations, at \( P > 4.5 \text{ GPa} \), there are no samples with \( f_{\text{O}_2} \) values on the extrapolation of this trend to higher pressures. Instead almost all samples from these depths are at least 1–4 log units higher in \( f_{\text{O}_2} \). Given that the majority of these deep and oxidized samples are enriched relative to the samples from shallower depths (\( P < 4.5 \text{ GPa} \)), we argue that they have been metasomatised and that the metasomatism was strongly oxidising. This effectively requires addition of oxygen to the deep cratonic lithosphere. Likely candidates for the responsible metasomatic agents are carbonate-bearing, silicate melts\(^3, 25\), possibly derived from the deeper, reduced asthenosphere\(^26\) (at \( \geq 230 \text{ km depth} \), i.e. \( P > 7 \text{ GPa} \)), or saline, hydrous fluids derived from a postulated subducted slab beneath the base of the Slave cratonic lithosphere\(^27\). Such melts may contribute to the commonly observed elevated modes of orthopyroxene in some garnet peridotite xenolith suites\(^28\), in a manner similar to that proposed by Kelemen\(^29\).

At these pressures the mantle may be still under-saturated in metallic FeNi\(^4\). Stagno \textit{et al.}\(^3, 25\) showed that silicate melts with dilute carbonate contents could be stable at the relatively reduced \( f_{\text{O}_2} \) values of the mantle at these depths because of the lower carbonate activity relative to that of low-SiO\(_2\) carbonatite melts (Fig. 3). If these carbonated, silicate melts segregate and percolate upwards through parts of the deep cratonic lithosphere they would encounter progressively cooler conditions along the cratonic geotherm, at some point freezing into the lithospheric mantle as the appropriate peridotite+volatile solidus is locally crossed\(^30\). The oxidized carbon species exsolved from the melt during crystallisation would reduce to C or CH\(_4\) (\( \text{CO}_2 + \text{C} = \text{CO} + \text{CH}_4; \text{CO}_2 + 2\text{H}_2\text{O} = \text{CH}_4 + 2\text{O}_2 \)). Very minor amounts of such C and H might be incorporated into nominally volatile-free, peridotitic silicate minerals (such as olivine, ortho- and clinopyroxene, garnet)\(^31, 32\), or if in excess, may form CH\(_4\) and/or diamond when in contact with reduced deep peridotite. Some of the Fe\(^{2+}\) in the melt and wall rock would oxidize to Fe\(^{3+}\) (\( 2\text{FeO} + 1/2\text{O}_2 = \text{Fe}_2\text{O}_3 \)). This and other components would be incorporated into garnet and pyroxenes, leading to the observed increase in lithospheric \( f_{\text{O}_2} \) associated with metasomatism. The increased activity of H\(_2\text{O}\) may cause a substantial drop in the solidus temperature of peridotite and consequent partial melting, if the quantity of newly formed H\(_2\text{O}\) exceeds the storage capacity of nominally anhydrous minerals in peridotite at given pressure, temperature and \( f_{\text{O}_2} \) (refs 30 and 33), a process known as hydrous redox melting\(^34, 35\).

An alternative scenario for metasomatism of the deep Slave cratonic lithosphere has been proposed by Weiss \textit{et al.}\(^27\), whereby hydrous, carbon-bearing and saline fluids in a subducted slab underlying the cratonic lithosphere...
percolate into the overlying lithosphere, evolving to silicic and carbonatitic melts which are preserved as inclusions in syngenetic fibrous diamonds. Similarly, Miller et al. proposed a model of fibrous and non-fibrous diamond formation from reduction of carbonatitic fluids under Ekati, accompanied by enrichment of the mantle in Ca and REE. Reduction of carbonate components in such fluids to diamond could conceivably be accompanied by oxidation of Fe in residual fluids and silicate phases which might crystallise from them.

The observations and processes described above have important general implications for the petrogenesis of kimberlites, some of which (including those discussed here) are primary hosts of economic diamond deposits. High pressure experimental investigations show that group I kimberlite magmas likely formed from low degree melting of carbonate + H2O-bearing garnet harzburgite at pressures near the base of typical cratonic lithosphere37–40, whereas group II kimberlitic magmas (orangeites) may have formed from phlogopite-bearing peridotite41. The pressure-temperature conditions for the origins of group I kimberlites were proposed to be those at which the primary melt is saturated in magnesite-bearing garnet peridotite, but will vary with the details of volatile contents and species37. Ca-bearing magnesite is most likely a necessary phase in the peridotitic melting assemblage to form primary, carbonate-rich melts which can evolve to kimberlite37.

This introduces a problem, because at depths corresponding to the likely source regions of kimberlites (near the base of cratonic lithosphere at ≈5–7 GPa [160–230 km], or deeper in the asthenosphere) ambient peridotite mantle oxygen fugacity is too low for crystalline carbonate stability. Predictions from thermodynamic arguments and trends from direct measurements of depleted garnet peridotite xenoliths, as presented here, show that the mantle’s fO2 at these depths (≈160–230 km) is approaching the Ni precipitation curve at around the iron-wüstite (IW) buffer42. Hence, the pre-metasomatic, deep cratonic lithosphere may have been too reduced for carbonate to be stable as a crystalline solid solution (i.e. Ca-magnesite). Even deep lithosphere oxidized by metasomatism, as represented by the xenoliths discussed here, is mostly too reduced for crystalline carbonate stability. Rather such pressure-temperature-fO2 conditions favour the stability of diamonds as the solid carbon phase in equilibrium with a metasomatic, carbonate-poor, but water-bearing silicate melt.

Tappe et al. have proposed a model for genesis of the Lac de Gras kimberlites, based in part on radiogenic isotope systematics. Lac de Gras kimberlites exhibit restricted ranges in Sr and Nd isotopic compositions, close to bulk earth and CHUR values, but extreme heterogeneity in Hf isotope compositions (ΔεHf varies from +5 to −5). In addition, the Lac de Gras kimberlites are known to have transported ultra-deep diamonds43 to the surface. The decoupling of different radiogenic isotope systems and evidence of ultra-deep components in the kimberlites has been explained by a model in which ancient, deeply recycled crustal lithologies (both MORB43 and OIB) in the Mantle Transition Zone (MTZ) underwent partial melting. These melts refertilised refractory peridotite wall-rock in the MTZ or deep upper mantle. Vigorous mantle convection transported the refertilised domains upwards, in some cases allowing transport of ultra-deep diamonds (and their inclusions) from the deep upper mantle or MTZ, to shallower levels. Some of the refertilised domains may have reached depths at which the peridotite mantle was no longer metal saturated (≤300 km) and may have undergone redox partial melting to produce CO2-bearing, silicate magmas45. These magmas were then transported into the deep cratonic lithosphere, metasomatically enriching and oxidizing it over time.

Progressive, long term oxidation of conduits or zones in the deep cratonic lithosphere by infiltrating carbonate-bearing silicate melts, in the manner described above, or by the incorporation of subducted carbonate-bearing crustal material, and associated complex redox-melting and freezing reactions46, 47, could lead to pressure-temperature-fO2 conditions under which crystalline carbonate phases (Ca-magnesite) become stable in deep, refractory cratonic peridotite relative to diamond or CH4 fluids. These may ultimately constitute fO2 heterogeneities, which could be oxidized and chemically enriched, localized sources of carbonatites, kimberlites and other highly silica-undersaturated magnas46, 47. This metasomatism may also add accessory phlogopite or K-richertite to the deep lithosphere48, phases implicated in the petrogenesis of some alkali-rich magnas48, 49, 50. Therefore, carbonatites and kimberlites may form at these depths only after long-term redox pre-conditioning of source regions near the base of the cratonic lithosphere (≈5–7 GPa) by metasomatism, where the pre-conditioning involved trace element enrichment and oxidation to levels of carbonate, rather than diamond stability.

Methods
Sample preparation. The Panda samples were mounted in 1-inch diameter round epoxy buttons, sectioned and polished to expose mineral phases. The new A154-N samples were presented as polished thin-sections (thickness 60 μm).

Conventional electronprobe microanalysis. The major and minor element compositions of mineral phases from the new A154-N garnet peridotite samples were determined by wave-length dispersive electronprobe microanalysis (EPMA) using the 4–spectrometer Cameca SX100 instrument at the Australian National University (ANU). Calibration was performed using a range of well-characterised natural minerals standards. Column conditions were 15 kV and 20 nA, with the beam focused to 1 μm diameter. Peak counting times varied from 10–60 seconds, depending on element abundance, and background counting times varied from 5–30 seconds.

Laser ablation-ICPMS. Trace element abundances in garnet and clinopyroxene in the xenoliths were measured by laser ablation ICP-MS, using the instrument at RSES, ANU. This consists of an Excimer 193 nm laser coupled to a purpose built sample cell and an Agilent 7500 ICP-MS.

Analytical conditions used were typically as follows. The laser spot size was varied from 135 to 225 μm depending on the nature of the phase being analysed. The laser pulsed at 5 Hz. For each analysis a gas blank (laser off) was collected for 25 seconds, the laser triggered, and the signal collected for a further 35 seconds. The ablation...
atmosphere was He and the ablated material was delivered to the Ar plasma in an Ar/H₂ mixture. The ionised material was analysed in peak hopping mode by the quadrupole mass spectrometer.

The analytical protocol typically consisted of analyses of the calibrating standard (NIST612 glass) bracketing 6–10 unknowns. BCR2g glass or garnet MU53388 were also analysed as unknowns to provide a check on data quality. In almost all cases, measured trace element abundances in BCR2g and MU53388 were within 5–10% of accepted values.

⁴³Ca was used as the internal reference element, based on CaO determined by EPMA. The values for NIST612 of Pearce et al.[54] were used in the data reduction. A linear drift correction was applied to the background corrected signal for each analysed mass, based on interpolation between bracketing analyses of calibration standards[51].

Mössbauer spectroscopy. Garnet Fe⁴⁺/∑Fe from the new A154-N samples was measured using Mössbauer spectroscopy applied to ≈20 mg samples of clean, inclusion and alteration-free garnet fragments, separated by hand-picking under a binocular microscope from crushed portions of all xenoliths except 60505115. Garnet grains from sample 60505115 exhibited cloudy interiors as a result of the presence of abundant inclusions, and were therefore not considered suitable for bulk analysis using Mössbauer spectroscopy. Separated material from the other samples was washed for 5 minutes in HF acid to remove any surface alteration. The acid-washed grain fragments were then inspected with a binocular microscope to ensure that they were free of inclusions and alteration material. Analysis and data reduction were performed at the University of Frankfurt following well-established procedures of Woodland and Koch[10] and Woodland and Ross[55]. Garnet grains separated from these samples have been used as calibration standards for the XANES method for determining Fe³⁺ in mantle garnet[7, 18, 56, 57].

X-ray Absorption Near-Edge Structure (XANES) spectroscopy. Garnet Fe³⁺/∑Fe was determined on the Panda xenoliths using the Fe K-edge XANES method of Berry et al.[56]. Spectra were collected at the X-ray Fluorescence Microscopy beamline at the Australian Synchrotron[58]. The method was calibrated using homogeneous garnet standards from the new A154-N samples, for which Fe³⁺/∑Fe was previously determined by Mössbauer spectroscopy and the suite of Kaapvaal samples of Woodland and Koch[10], for which garnet Fe³⁺/∑Fe was also determined by Mössbauer spectroscopy. Up to three individual garnet grains per sample were measured.

Following Berry et al.[56] the calibration curve obtained at the beamline related the normalized intensity ratios of the post-edge features in the XANES spectrum at 7138.4 and 7161.7 eV to the Fe³⁺/∑Fe of the standard garnets as determined by Mössbauer spectroscopy (Fig. 4). The precision of the XANES Fe³⁺/∑Fe measurements is estimated at ±0.012, comparable with that of the Mössbauer measurements[10].

Thermobarometry. Equilibration pressures and temperatures were calculated for the new A154-N and Panda xenoliths using conventional thermometers and barometers. The results are presented in Supplementary Table 6. For the A154-N samples, the approach of Nimis and Grütter[20] was used as this has been shown to be a suitable thermometer and barometer combination for garnet lherzolite assemblages[20]. However, because many of the Panda samples do not contain modal clinopyroxene, the garnet-olivine thermometer of O’Neill and Wood[59] combined with the garnet-orthopyroxene barometer of Brey and Köhler[19] was preferred. Fe³⁺ in garnet determined by XANES spectroscopy was incorporated into the calculations.
Calculation of oxygen fugacity. The presence in the new samples of fresh garnet, olivine and orthopyroxene was used as a basis for selection for garnet Fe3+ analysis, to enable calculation of the samples’ O2 fugacity. This information (Supplementary Tables 4 and 5), combined with the results of thermobarometry and the major element compositions of olivine and orthopyroxene measured by electronprobe microanalysisPorcelli et al., enabled calculation of the fO2 conditions which the xenoliths experienced prior to entainment in the host kimberlite (Supplementary Table 6).

All fO2 calculations presented in this paper were calculated (or re-calculated) using the experimental calibration for garnet peridotite assemblages of Stagno et al. and garnet Fe3+/ΣFe determinations by Mössbauer spectroscopy (the new samples from A154-N, following the method of Woodland and Koch(90), or Fe K-edge XANES (the Panda samples, following the method of Berry et al.)). This data was coupled with new mineral compositional data for the new A154-N samples, previously reported data for the Panda suite and the earlier A154-N suite and with the thermobarometric data as reported above. Uncertainties in the fO2 calculations are estimated at ±0.6 log units.

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