The thermal structure of continental crust in active orogens: insight from Miocene eclogite and granulite xenoliths of the Pamir Mountains

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ABSTRACT Rare ultrahigh-temperature–(near)ultrahigh-pressure (UHT–near-UHP) crustal xenoliths erupted at 11 Ma in the Pamir Mountains, southeastern Tajikistan, preserve a compositional and thermal record at mantle depths of crustal material subducted beneath the largest collisional orogen on Earth. A combination of oxygen-isotope thermometry, major-element thermobarometry and pseudosection analysis reveals that, prior to eruption, the xenoliths partially equilibrated at conditions ranging from 815 °C at 19 kbar to 1100 °C at 27 kbar for eclogites and granulites, and 884 °C at 20 kbar to 1012 °C at 33 kbar for garnet–phlogopite websterites. To reach these conditions, the eclogites and granulites must have undergone mica-dehydration melting. The extraction depths exceed the present-day Pamir Moho at ~65 km depth and suggest an average thermal gradient of ~12–13 °C km⁻¹. The relatively cold geotherm implies the introduction of these rocks to mantle depths by subduction or gravitational foundering (transient crustal drip). The xenoliths provide a window into a part of the orogenic history in which crustal material reached UHT–(U)HP conditions, partially melted, and then decompressed, without being overprinted by the later post-thermal relaxation history.

Key words: garnet-omphacite xenoliths; granulite xenoliths; oxygen isotopes; Pamir; partial melting.

INTRODUCTION

The thermal structure of collisional orogenic belts depends on a variety of factors including the rate of subduction, accretion and denudation, the radiogenic heat production, thermal conductivity and the rheology of the underlying lithospheric mantle (e.g. Royden, 1993; Peacock, 1995; Huerta et al., 1998; Pope & Willett, 1998). As most of these factors evolve over the course of orogenesis, the thermal structure changes as the lithosphere thickens, undergoes thermal relaxation and collapses. Therefore, understanding the thermal structure is important for predicting the overall rheological behaviour of collisional orogenic belts (e.g. is partial melting possible? England & Thompson, 1984; Patiño Douce & McCarthy, 1998; Hacker et al., 2000; Erkan & Blackwell, 2008). However, access to rocks from different parts of thickened crust within collisional orogenic belts and to different stages of the thermal evolution (e.g. shortening, thermal relaxation, and collapse) is generally limited.

A suite of Miocene xenoliths erupted in the Pamir Mountains of Tajikistan consists of eclogites and granulites with bulk compositions that suggest that the xenoliths have crustal protoliths (Fig. 1; Dmitriev, 1976; Ducea *et al.*, 2003; Hacker *et al.*, 2005). Some of the eclogite xenoliths equilibrated at 25–28 kbar and 1000–1100 °C (Hacker *et al.*, 2005), just below the coesite-stability field. A group of plagioclase-bearing granulite xenoliths, not studied by Hacker *et al.* (2005), probably equilibrated at lower pressure. All xenoliths were erupted at the same time (11.5 \pm 0.2 Ma, ⁴⁰Ar–³⁹Ar; Hacker *et al.*, 2005) and thus provide insight into the evolving thermal structure of the largest collisional belt in the world, the Pamir Mountains–Tibet system.

To understand the transient thermal structure of orogenic systems, the P-T-t path that the rocks experienced must be determined. Because UHP rocks represent the deepest exposed portions of orogenic belts, numerous studies have focused on determining the peak pressures and temperatures achieved by these rare rocks (e.g. Nakamura & Banno, 1997; Ravna, 2000; Krogh Ravna & Terry, 2004; Nakamura & Hirajima, 2005; Hacker, 2006; Ernst *et al.*, 2007). Most temperature estimates of eclogites have relied on Fe–Mg equilibrium between garnet and clinopyroxene. The accuracy of this thermometer is affected, however, by the Fe³⁺/Fe²⁺ ratios, which are typically not measured (e.g. Krogh Ravna & Paquin, 2003;

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Fig. 1. (a) Digital elevation map of the Pamir Mountains–Tibet–Himalayan system; (b) southeastern Pamir xenolith locality plotted in relation to the Pamir Mountains and Hindu-Kush seismic zones; seismicity from Engdahl *et al.* (1998); (c) simplified geological and structural map of the Pamir Mountains and northern Karakoram, showing Palaeozoic to Tertiary magmatic belts and sutures, Cenozoic gneiss domes, major Cenozoic faults and the location of the Dunkeldik magmatic field (marked by box) (strongly modified from Vlasov *et al.*, 1991; Schwab *et al.*, 2004; Hacker *et al.*, 2005).

Proyer et al., 2004; Stípská & Powell, 2005). Oxygenisotope thermometry represents a complementary method and a check for major-element thermometry. For the Pamir Mountains xenoliths, oxygen diffusion should have been negligible during the rapid eruption and cooling experienced by these xenoliths (Hacker et al., 2005), and therefore, fractionation of oxygen isotopes among the minerals should reflect pre-eruption metamorphic equilibrium (e.g. Sharp et al., 1992, 1993; Rumble & Yui, 1998; Zheng et al., 1998; Valley, 2001; Schulze et al., 2003a). In addition, some complications associated with major-element thermobarometry, such as non-ideal mixing, H₂O activity and pressure dependence, do not affect oxygen-isotope fractionation (Hoering, 1961; Wolfsberg, 1972; Clayton et al., 1975). A combination of conventional thermobarometry and oxygen thermometry techniques can be applied to the Pamir Mountains xenoliths to assess mineral equilibrium and pre-eruption P-T conditions, which were reached during crustal thickening of a large orogen.

In this study, phase equilibria and pre-eruption P-Tconditions are estimated from eclogite and granulite xenoliths by combining oxygen-isotope thermometry, major-element thermobarometry and pseudosections. The samples represent a range of protoliths, from gabbro-granodiorite (for the eclogites) to peraluminous metasedimentary rocks (for the granulites). To further constrain the thermal structure of the region, we also discuss the P-T equilibration of several websterite xenoliths. It is found that at depths of ~ 60 -100 km, the Miocene crust of the Pamir Mountains consisted of eclogite interlayered with garnet-kyanite granulites and websterites. The xenoliths record probable subduction of crustal material in an actively shortening collisional orogen. The granulite xenoliths suggest some cooling and decompression prior to the final eruption at 11 Ma (Hacker et al., 2005).

GEOLOGICAL SETTING

The Pamir Mountains represent the westernmost extent of the Pamir Mountains-Tibet orogen (Fig. 1a). Similar to Tibet, the Pamir Mountains constitute a plateau with a thick crust (~65 km; Beloussov et al., 1980; Mechie et al., 2011), resulting from 1800-2100 km of Cenozoic intracontinental shortening (Johnson, 2002) associated with the India-Eurasia collision. However, in comparison to Tibet, the shortening has been accommodated in half the orogenic width. The Pamir Mountains are still shortening at a rate of 16–20 mm yr^{-1} (Reigber *et al.*, 2001; Mohadjer et al., 2010) and are underlain by intermediatedepth seismicity that may be associated with intracontinental subduction (Fig. 1b; e.g. Pegler & Das, 1998; Negredo et al., 2007). Three belts of mid- to lower-crustal gneiss domes were exhumed chiefly between 25 and 10 Ma (Fig. 1c, Robinson et al., 2007; McGraw et al., 2010; Stearns et al., 2011). A variety of

Cretaceous granitoids to Miocene shoshonitic/calcalkaline igneous bodies intruded the Pamir Mountains crust, including the domes (Schwab *et al.*, 2004).

The volcanic pipes that host the xenoliths studied in this article (first described by Dmitriev, 1976) belong to the Dunkeldik magmatic field in the southeastern Pamir (Fig. 1c). The Dunkeldik magmatic field is the result of the youngest known magmatism in the Pamir Mountains (c. 11 Ma; Hacker et al., 2005) and consists of ultrapotassic (4–7 wt% K_2O ; $K_2O/Na_2O = 4-7$) dykes, pipes and sub-volcanic bodies that range from alkali basalt, to trachyte, syenite and carbonatite (Dmitriev, 1976). The field is exposed between the Late Triassic-Early Jurassic Tanymas suture and the Late Jurassic-Early Cretaceous Rushan-Pshart suture, in an area of active deformation related to the Karakoram fault zone (Fig. 1c; Dmitriev, 1976; Strecker et al., 1995). The Dunkeldik xenolith suite consists of 38% eclogite and garnet-omphacite granulite, 19% phlogopite pyroxenite and glimmerite, 15% biotite-garnet clinopyroxenite, 8% garnet-biotite gneiss, 6% garnetkyanite granulite, 4% garnet gneiss, 3% phlogopitegarnet websterite and 7% other rocks (Lutkov, 2003). The xenoliths are up to 50 cm in diameter and most are foliated. Hacker et al. (2005) studied six of these xenoliths, including sanidine \pm biotite eclogite, felsic garnet-sanidine-kyanite \pm biotite granulites and glimmerite. Major-element thermobarometry revealed equilibration of the eclogites at 1000-1100 °C and 25-28 kbar. The rocks were interpreted to be residues of high-pressure dehydration melting and K-rich metasomatism of granodioritic to gabbroic protoliths. Our study not only focuses on seven garnet-kyanite granulites, but also includes one garnet-omphacite granulite, two eclogites, one garnet-biotite gneiss and five websterites; these rocks equilibrated at a broader range of pressures and temperatures and, thus, provide more constraints on the thermal structure of the Miocene Pamir crust. Analytical methods are outlined in Appendices S1 and S2.

RESULTS

Mineral chemistry and bulk-rock compositions

The studied xenoliths form a continuous compositional series, from quartzofeldspathic garnet-kyanite granulites to eclogite (Table 1). Their protoliths were likely magmatic (i.e. granodioritic-gabbroic) and metasedimentary rocks. Mineral compositions for all samples are reported in Table S1.

Garnet-biotite gneiss

Garnet-biotite gneiss DK83 has a grano-lepidoblastic texture, with 22 vol.% biotite that defines a strong foliation along with quartz and feldspar (Fig. 2a). Grain sizes average 0.5–1.0 mm. The matrix biotite is Ti rich (TiO₂ = 7.1 wt%), whereas biotite inclusions

Sample	DK7	DK13	DK32	DK43	DK63	DK69	DK71	DK74	DK83	DK84	1309
XRF											
SiO ₂	69.57	59.01	50.44	69.01	49.20	63.13	57.60	60.43	62.44	48.74	56.05
TiO ₂	1.22	1.26	1.63	0.77	1.19	0.69	1.13	0.80	0.66	1.16	0.90
Al_2O_3	14.42	19.95	17.40	13.43	24.66	15.08	17.42	19.63	14.72	19.65	19.12
FeO*	5.97	8.38	9.33	5.43	14.21	6.04	8.60	6.86	4.88	9.57	7.15
MnO	0.15	0.18	0.17	0.12	0.82	0.29	0.17	0.15	0.10	0.23	0.12
MgO	1.89	2.83	4.01	1.92	3.57	2.05	3.24	2.77	2.81	4.25	3.72
CaO	0.88	1.43	9.59	0.59	1.68	3.33	2.59	2.10	3.28	7.54	6.70
Na ₂ O	0.62	0.88	2.96	0.93	0.57	1.61	0.82	0.90	2.51	2.42	2.46
K_2O	3.89	4.04	2.03	3.77	3.87	4.75	4.86	3.89	4.21	4.18	3.80
P_2O_5	0.03	0.18	0.25	0.03	0.26	0.12	0.22	0.04	0.32	0.25	0.20
Sum	98.64	98.13	97.79	96.00	100.03	97.08	96.64	97.56	95.93	97.99	100.22

Table 1. Measured bulk compositions of the Pamir Mountains xenoliths.

Oxides in wt%. Analyses accomplished using the XRF at Washington State University (Johnson et al., 1999).



Fig. 2. Representative photomicrographs from the Pamir Mountains xenoliths: (a) garnet-biotite gneiss DK83 with lobate quartz grain boundaries; (b) kinked and tabular kyanite surrounded by feldspar and quartz in granulite DK63; (c) tabular twinned kyanite with undulatory extinction in granulite DK63; (d) garnet encloses elongate kyanite in granulite DK69, with a ribbon of sanidine also parallel to foliation; (e) carbonate and sanidine veins cut clinopyroxene in eclogite DK84; and (f) garnet rimmed mostly by altered omphacite in eclogite DK84. Carb = carbonate; Cpx = clinopyroxene; Grt = garnet; K-spar = alkali feldspar; Ky = kyanite; Qz = quartz; Plag = plagioclase Rt = rutile.

in garnet have less TiO₂ (~4.7 wt%) and higher Mg# (molar Mg/(Mg + Fe); 0.66 v. 0.63 for matrix biotite). The garnet (~12 vol.%) is anhedral and shows weak zoning, with Alm₅₀Prp₀₁Grs₃₈Sps₁₁ cores and Alm₅₃Prp₀₃Grs₃₅Sps₀₉ rims (Fig. 3a); these compositions are significantly different from the garnet in other xenoliths (Fig. 3a; see below). Homogeneous plagioclase (An₂₈₋₂₉Ab₆₁₋₆₂Or₀₉₋₁₀) and alkali feldspar (An₀₀Ab₀₂Or₉₈) are present throughout the matrix (Fig. 3c). Matrix quartz has lobate boundaries; some grains form ribbons or have chessboard extinction. Throughout the sample, carbonate veins parallel the foliation, and carbonate-filled cracks occur in garnet. Carbonate + K-feldspar aggregates replace an unknown mineral. Overall, the garnetbiotite gneiss represents a rare rock type within the xenolith suite; there are few xenoliths that contain hydrous minerals.

Garnet-omphacite granulite

In general, the garnet–omphacite granulite DK32 is similar in texture and composition to the eclogites and garnet–kyanite granulites (described in the next sec-



Fig. 3. (a) Grossular–almandine–pyrope compositions of garnet. Arrows show core to rim zoning; (b) Ca-Tschermak, Mg-number and acmite + jadeite components of clinopyroxene; and (c) albite–anorthite–orthoclase compositions of feldspar with solvus isotherms at 17 kbar (Elkins & Grove, 1990). For all samples, the black fill indicates the core composition, whereas the white fill indicates the rim composition.

tions) but contains plagioclase and omphacite in equilibrium with garnet, kyanite and alkali feldspar.

The grains in xenolith DK32 have an average size of 0.5–1.5 mm. Garnet cores are Alm₄₈Prp₂₆Grs₂₅Sps₀₁, and rims are Alm₄₄Prp₂₁Grs₃₄Sps₀₁ (Fig. 3a). Omphacite shows a rimward decrease in jadeite content from 37 to 33 mol.% and in Mg# from 0.67 to 0.64 in the outermost 50 μ m (Fig. 3b). Micron-scale carbonate veins (Ca_{0.58-0.60}Mg_{0.28-0.30}Fe_{0.11-0.12})CO₃ cut the rock, and 100–500 μ m calcite grains (Ca_{0.96}Mg_{0.02}Fe_{0.02}) CO_3 are spatially associated with plagioclase and omphacite. Coronas around clinopyroxene consist of quartz, oligoclase $(An_{15}Ab_{73}Or_{12})$ and sanidine $(An_{00}Ab_{02-06}Or_{94-98})$ (Fig. 3c). Clinopyroxene in these coronas has a much lower jadeite content of 11 mol.%, and clinopyroxene in fractures that cut garnet has only 4 mol.% jadeite. Homogeneous oligoclase (An₁₆Ab₇₃Or₁₁) throughout the thin section appears to be in textural equilibrium with garnet and omphacite (Fig. 3c). Most

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quartz grains and minor, skeletal kyanite are surrounded by alkali-feldspar coronas $(An_{00-03}Ab_{02-04} Or_{93-98}; Fig. 3c)$.

Garnet-kyanite granulites

In general, the garnet–kyanite granulites only show minor alteration at the tens of micron scale, mostly on the rims of garnet, including symplectite coronas composed of two feldspars $(An_{00-01}Ab_{03-04}Or_{96-99})$ and $An_{01-03}Ab_{23-31}Or_{66-75}) \pm$ quartz \pm spinel. Veins of similar feldspars locally cut garnet and surround garnet inclusions; in the matrix, Or_{96-99} feldspar forms coronae around kyanite. The veins also contain local carbonate ($Ca_{50-59}Mg_{25-36}Fe_{12-19}$). Minor-oriented ilmenite lamellae are found in rutile. Below, each sample is described in more detail.

The plagioclase-bearing garnet-kyanite granulite DK69 contains abundant, up to \sim 1 cm long kyanite.

Garnet, plagioclase, minor alkali feldspar and quartz are also present and have an average grain size of 1-2 mm. Minor graphite is present in the matrix. Garnet occurs both as porphyroblasts and as coronae on kyanite (Fig. 2d); all garnet grains are zoned from Alm₅₀Prp₃₅Grs₁₂Sps₀₃ cores to Alm₄₈Prp₃₂Grs₁₇Sps₀₃ rims, with similar rim compositions at the kyanite interface as well as at the interface with feldspar Homogeneous (Fig. 3a). $An_{29-32}Ab_{58-60}Or_{09-11}$ plagioclase dominates the matrix, whereas plagioclase included in garnet is zoned from cores of An₃₃₋₃₄Ab₅₇₋₅₈ Or_{08-09} to rims of $An_{29-32}Ab_{46-55}Or_{14-25}$ (Fig. 3c). K-feldspar inclusions in garnet have An₀₄Ab₂₆Or₇₀ rims and An₀₄Ab₃₀Or₆₆ cores or are homogeneous Or₉₉. The matrix alkali feldspar $(An_{04-05}Ab_{28-30}Or_{65-67})$ occurs in ribbons parallel to the elongate garnet and kyanite grains, and is associated with calcite ($Ca_{0.95-0.97}$ $Mg_{0.01}Fe_{0.01-0.04}$)CO₃ nodules, which are also present as inclusions in garnet.

Quartz, sanidine and elongate garnet define a strong foliation in DK7. The sample is porphyroblastic, with an average grain size of 1–2 mm. Garnet is up to 1 cm in diameter and homogenous (Alm₅₃Prp₃₇Grs₀₉Sps₀₁; Fig. 3a); it contains inclusions of quartz, sanidine, kyanite and graphite. Matrix quartz displays chessboard extinction. Matrix kyanite is bent around garnet grains and is typically surrounded by a moat of alkali feldspar similar in composition to the coarser feldspar grains (An_{02–03}Ab_{26–29}Or_{68–72}; Fig. 3c). Abundant fluid/melt inclusions in the kyanite are elongate parallel to the fold hinges in the host grain and sub-parallel to the matrix foliation. Rutile crystals are large (>1 mm), homogeneous and xenoblastic.

Granulite DK13 has a strong foliation defined by elongate garnet, kyanite and quartz. The sample consists of ~41 vol.% garnet, ~15% quartz, ~12% kyanite and ~30% alkali feldspar (An₀₁₋₀₃Ab₂₆₋₃₀Or₆₈₋₇₆; Fig. 3c). The grain size averages 1–2 mm. The majority of garnet show only micron-scale rim alteration and is homogeneous Alm₅₅Prp₃₅Grs₀₉Sps₀₁ (Fig. 3a). Elongated kyanite, as well as sanidine, mica, quartz, rutile and apatite, are included in garnet. As in DK7, kyanite contains abundant fluid/melt inclusions, is bent and is rimmed by alkali feldspar of the same composition as in the matrix.

Granulite DK43 has a strong foliation defined by quartz (~63 vol.%), alkali feldspar (~17%), garnet (12%) and kyanite (~8%). The grain-size averages 1–2 mm, with coarser quartz and feldspar (~1 mm) than in other samples. Garnet forms small (2 mm), homogeneous (Alm₅₆Prp₃₉Grs₀₄Sps₀₁), typically elongated xenoblasts (Fig. 3a), with few quartz and kyanite inclusions. The matrix alkali feldspar is homogeneous An₀₃Ab₃₆Or₆₁ (Fig. 3c). Most kyanite grains are surrounded by moats of late feldspar (Or_{97–99}) (Fig. 3c); some have undulatory extinction and fluid/melt(?) inclusion trails.

Granulite DK63 is characterized by \sim 15 vol.% deformed kyanite in up to 0.5 cm elongate crystals that

define the foliation (Fig. 2b). Quartz, sanidine and garnet have an average grain size of 0.5–1.5 mm. Kyanite is commonly twinned and displays undulatory extinction (Fig. 2b,c); kyanite included in garnet is undeformed. Garnet is slightly zoned, with inclusion-rich, altered and fractured cores of Alm₅₉Prp₂₇Grs₁₀ Sps₀₄ and inclusion-free, homogeneous rims of Alm₆₀Prp₂₉Grs₀₈Sps₀₃ (Fig. 3a). Alkali feldspar is homogeneous An₀₁Ab₂₁Or₇₈ (Fig. 3c).

Sample DK71 contains subidioblastic garnet in a matrix of alkali feldspar (~30 vol.%), quartz (~20%) and kyanite (~8%). The feldspar and kyanite define the foliation and have an average grain size of 1–2 mm. The garnet grains are zoned in their outermost ~50 μ m, from Alm₅₇Prp₃₈Grs₀₄Sps₀₁ to Alm₅₃Prp₃₆ Grs₁₀Sps₀₁ rims (Fig. 3a). Alkali feldspar is homogeneous An₀₂Ab₂₆Or₇₂ (Fig. 3c). Kyanite is kinked, has undulatory extinction and is typically surrounded by late An₀₄Ab₂₈Or₆₈ feldspar. Similar coronas between kyanite and garnet contain An₁₅Ab₃₈Or₄₇ feldspar. Quartz forms elongate ribbons that wrap around garnet.

Granulite DK74 is granoblastic, with ~1 mm grains of garnet, quartz, kyanite and alkali feldspar that define a weak foliation. Garnet is weakly zoned, from $Alm_{51}Prp_{35}Grs_{13}Sps_{01}$ cores to $Alm_{51}Prp_{36}Grs_{12}Sps_{01}$ rims (Fig. 3a). They are sub- to idioblastic and have ~10–20 μ m rims of feldspar and carbonate (Ca_{51–56} Mg_{30–34}Fe_{13–16}). Feldspar (An_{00–03}Ab_{00–08}Or_{92–99}) plus carbonate coronas surround quartz and kyanite that are adjacent to garnet. The matrix feldspar is similar in composition to the corona feldspar but locally has more sodic (An₀₂Ab₁₆Or₈₃) rims (Fig. 3c). The alkali feldspar in the matrix and in garnet strain shadows is zoned from An₀₀Ab₁₆Or₈₄ cores to An₀₀Ab₂₉Or₇₁ rims (Fig. 3c). Garnet contains inclusions of Or₉₉ and An₀₅Ab₆₈Or₂₇ alkali feldspar.

Sanidine eclogites

The garnet- and omphacite-dominated samples, including DK84 and 1309, contain kyanite, quartz, sanidine, and minor rutile, apatite, zircon and monazite. The samples are transected by small veins of feldspar or carbonate (see below; Fig. 2e). In addition, garnet and omphacite have $< 10 \ \mu$ m thick quartz + feldspar coronas.

Eclogite DK84 (Fig. 2e,f) has a foliation defined by the shape-preferred orientation of omphacite, quartz and sanidine. The garnet grains are zoned, and different grains have distinct rim compositions. Some have $Alm_{40}Prp_{25}Grs_{34}Sps_{01}$ core, a $Alm_{44}Prp_{31}Grs_{24}Sps_{01}$ mantle and $Alm_{47}Prp_{25}Grs_{27}Sps_{01}$ rim (Fig. 3a); others have rims of $Alm_{42}Prp_{22}Grs_{34}Sps_{02}$. Most garnet has a ~50 μ m corona of intermingled $An_{01}Ab_{12}Or_{87}$ and $An_{04}Ab_{61}Or_{35}$ feldspar (Fig. 3c). The omphacite grains have cores of 32 mol.% Jd and Mg# 0.71 and rims of 42 mol.% Jd and Mg# 0.64 (Fig. 3b). Micron-scale carbonate veins (Ca_{0.52-0.62}Mg_{0.25-0.30}Fe_{0.12-0.18})CO₃ traverse the sample and cut garnet (Fig. 2e). Coronas (>10 μ m thick) of sodic clinopyroxene + plagioclase have replaced <10% of the omphacite. Homogeneous sanidine (An₀₄₋₀₆Ab₄₃₋₅₀Or₄₄₋₅₄) is interspersed and in textural equilibrium with omphacite and garnet (Fig. 3b). Minor quartz (~10%) is typically associated with kyanite, but also occurs as inclusions in omphacite. Quartz–omphacite boundaries are separated by a film of quartz and feldspar (An₀₁Ab₀₃Or₉₆). Kyanite grains are surrounded by coronas of two feldspars, An₀₁Ab₂₅Or₇₄ and An₁₂Ab₈₁Or₀₇.

Eclogite 1309, described in Hacker *et al.* (2005), consists of garnet, omphacite, sanidine, kyanite, quartz and rutile. Garnet and clinopyroxene are homogeneous, whereas the sanidine is zoned outward from a $Or_{73-74}Ab_{25}$ core to a $Or_{56-68}Ab_{31-43}$ rim. Like DK84, veins of carbonate cut the sample.

Garnet-phlogopite websterites

The websterite xenoliths (samples 1170, DK17, DK24, DK57, DK70) are foliated and range from granoblastic to porphyroblastic and grano-lepidoblastic. The foliation is defined by the shape-preferred orientation of phlogopite and/or pyroxene. They consist mainly of orthopyroxene, clinopyroxene, garnet and phlogopite and <1 vol.% apatite and pyrrhotite or rutile. Grain sizes average 1-2 mm, but poikiloblastic garnet reaches 3-5 mm. Unlike the eclogites, the garnet in websterites shows no significant compositional variability in individual samples (Table S1); compositions range from Prp₅₅Alm₃₃Grs₁₁Sps₀₁ to Prp₃₅Alm₅₆ Grs₀₈Sps₀₁. Pyroxene-dominated patches are granoblastic-polygonal, with typical 120° triple junctions. Clinopyroxene ranges from sodic augite to omphacite but is nearly homogeneous in each sample; the overall compositional range is $Di_{44-68}Hed_{12-15}Jd_{5-25}$ $CaTs_{01-06}(En + Fs)_{08-15}$ (Table S1). Some matrix clinopyroxene is surrounded by $< 10 \ \mu m$ symplectite rims of a Na-poor clinopyroxene and ternary feldspar. Orthopyroxene grains are in general homogeneous with compositions falling in the En₇₁₋₇₈Fs₁₆₋₂₃ (Di + Hd)₀₂₋₀₅MgTs₀₁₋₀₂ range; some larger grains display Al zoning (Table S1). Matrix orthopyroxene grains are coated by narrow selvages of unidentified, late submicroscopic minerals (likely phyllosilicate and/or amphibole). Phlogopite is in general homogeneous in composition (Table S1); in some websterites, it appears to be in textural equilibrium with other phases, whereas in other samples, it is secondary after pyroxene, indicating late potassium metasomatism. The websterites show varying degrees of alteration and local disequilibrium, which are mostly confined to the rinds of the xenoliths where late phlogopite formation and clinopyroxene breakdown to symplectite are abundant. Garnet and pyroxene away from the rinds are much less altered. The outermost mineral rims locally display distinct Mg# increases even in the cores of some of the xenoliths.

Oxygen-isotope data

Oxygen isotopes from the eight granulites and two eclogites were first measured using the laser-fluorination technique on bulk separates of kyanite, garnet, quartz and rutile to obtain high-precision measurements at the millimetre- to centimetre-scale (Fig. 4, Table 2). The $\delta^{18}O_{quartz}$ values range from 8.4% to 15.4% (Fig. 4), with the lighter values from eclogites 1309 and DK84, and garnet-omphacite granulite DK32. Garnet δ^{18} O values parallel those in quartz, and range from 6.8% in eclogite 1309 to 14.3% in garnet-kyanite granulite DK7 (Fig. 4a). Garnet in garnet-omphacite granulite DK32 is inhomogeneous, with a $\sim 0.9\%$ difference between the two laser-fluorination analyses. Kyanite δ^{18} O values range from 7.7% to 14.4% (Fig. 4b). Rutile yielded the lightest δ^{18} values 6.6–11.0% in DK32 and DK84, and 9.9–11.9% in the remaining granulites (Fig. 4c).

To determine whether the minerals are homogeneous in δ^{18} O at single grain and sample scales, *in situ* oxygenisotope measurements on quartz, rutile, kyanite, garnet and zircon were performed by ion microprobe (Fig. 5, Tables 3 & S2). The ion-microprobe analyses targeted the extremes in major-element zoning where found in garnet; in other minerals, cores and rims were analysed. Thus, the *in situ* data are not necessarily representative of the average values measured by laser fluorination. For the ion-microprobe analyses on rutile, only the precision within grains can be compared and not from grain to grain. The instrumental mass fractionation for rutile varies according to the orientation of the crystal lattice relative to the primary and secondary beams of the ion microprobe, and it is not possible to accurately convert the raw ion-microprobe data to the VSMOW scale for this mineral. Thus, grain-to-grain rutile comparison is not possible as each grain may have a different orientation. Overall, the results for the different minerals show some zoning in the δ^{18} O of individual grains as well as grain-to-grain variability for the minerals except for rutile.

The kyanite ion-microprobe values of δ^{18} O are similar to those obtained through laser fluorination and overlap with 99% confidence (Fig. 5a). The kyanite grains reveal a spread in values (0.2–0.9‰ 2 σ standard deviation) when including all of the data for the individual samples (Fig. 5a, Table S3). The spread is not caused by zoning within individual grains but rather differences from grain to grain.

The ion-microprobe results on garnet cluster near the laser data, but like kyanite, reveal heterogeneity (Fig. 5b, Table S3). As described above, most garnet is zoned in major cations; however, δ^{18} O does not typically follow the same zoning trend. In general, individual grains are homogeneous and the variability in δ^{18} O measurements reflects grain-tograin differences. The garnet in granulite DK32 tends to have lighter rims (~12.1‰) than cores (~12.5‰).



Fig. 4. Values of δ^{18} O laser fluorination (a) garnet, (b) kyanite and (c) rutile *v*. δ^{18} O(quartz) with isotherms at 200 °C intervals. Analytical uncertainties are smaller than the symbol size and are thus omitted.

Quartz δ^{18} O values show the smallest range of oxygen-isotope ratios, except for sample DK69 (Fig. 5c), for which ion-microprobe analyses reveal two populations, one ~13.7% and a second averaging 10.8%. The latter, lighter values correspond to quartz grains found in a late alkali-feldspar + carbonate ribbon formed parallel to the foliation. Xenoliths

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Sample	R ock type (δ ¹⁸ O Grt ‱ VSMOW)) Average ($^{0}_{\lambda}$	2 SD ^a 00 VSMOW)	δ ¹⁸ Ο Qtz) (‰ VSMOW	V) Average (?,	2 SD ^a ‱ VSMOW) (δ ¹⁸ Ο Ky ‰ VSMOW) .	Average (%	2 SD ^a VSMOW) ((δ ¹⁸ O Rt ‱ VSMOW) .	Average (%	2 SD ^a , VSMOW)	T(Grt-Qtz) (°C)	2 SD (°C) ^b (T Ky-Qtz) (°C)	2 SD (°C) ^b (T Rt-Qtz) (°C)	2 SD (°C) ^b
DK7	Granulite	14.3	14.30	0.17	15.28	15.35	0.17	14.3 14.30	14.35	0.17	11.89	11.86	0.17	- 1351	+ 223/-157	1223	+219/-152	885	+ 42/-38
DK13	Granulite	13.62	13.73	0.17	15.39	15.45	0.17	14.08	14.05	0.17	11.91	11.91	0.17	994	62-/26+	995	+125/-96	878	+41/-37
DK32	Eclogite	12.52	12.97	0.17	14.28 14.05	14.17	0.17	70.11			10.92	10.97	0.17	1246 -	+177/-131			937	+ 48/43
DK43	Granulite	13.37	13.44	0.10	14.17	14.17	0.10	13.86 13.82	13.84	0.10	10.94	10.93	0.10	- 1674	+425/-162	2358	+866/-434	930	+27/-25
DK63	Granulite	10.89	10.82	0.17	12.63	12.52	0.17	11.29	11.23	0.17	9.12	9.12	0.17	1005	+100/-81	1048	+143/-108	901	+ 44/-39
DK 69	Granulite	12.27	12.32	0.10	13.15	13.28	0.10	12.65	12.65	0.10	10.02	10.03	0.10	1425	+216/-111	1609	+252/-180	927	+ 27/-25
DK71	Granulite	12.87	12.89	0.10	13.43	13.44	0.10	13.22	13.27	0.10	9.82 9.94	9.88	0.10	- 1971	+139/-239	3313	+ 4549/-917	875	+ 23/-22
DK74	Granulite	9.42 9.24	9.33	0.17	10.88	10.89	0.17	9.74 9.96	9.85	0.17	8.05	8.05	0.17	1059	+354/-91	1198	+206/-145	1012	+ 58/-51
DK84	Eclogite	7.38	7.51	0.17	9.54	9.54	0.17	8.32	8.41	0.17	6.70 6.58	6.64	0.17	896	+75/-63	1141	+180/-130	1000	+ 56/-50
1309	Eclogite	6.80	6.80	0.20	8.40	8.40	0.20	7.70	7.70	0.20				1028 -	+201/-137	1520	+946/-363		
Temper: ^a 2 SD u ^b 2 SD te	atures were incertainties emperature	calculated u. were calcula uncertainties	sing the tem ted based o	perature cot n the analyt ated by taki	efficients from ical 2 SD unc ng the square	n Valley (2005 certainty on the stroot of the states	3). he standard L sum of the squ	JWG-2. uares of the un	certainties c	of the individ	tual mineral a	nalyses.							



Fig. 5. Values of laser fluorination δ^{18} O ν . ion microprobe δ^{18} O for (a) kyanite, (b) garnet, (c) quartz and (d) rutile. Note the heterogeneity found in the individual measurements. Rutile analyses by ion microprobe vary systematically from grain to grain due to orientation effects and are only accurate for assessing intra-mineral homogeneity.

Table 3.	Average	WiscSIMS	ion-microprobe	oxvgen-isotope	analyses an	d calculated	temperatures.

Sample	Rock type	δ ¹⁸ O Grt (%o VSMOW)	2 SD ^a (% VSMOW)	δ^{18} O Qtz (% VSMOW)	2 SD ^a (% VSMOW)	δ^{18} O Ky (% VSMOW)	2 SD ^a (% VSMOW)	δ ¹⁸ O Zirc (% VSMOW)	2 SD ^a (% VSMOW)	δ^{18} O Rt (% VSMOW)
DK7	Granulite	14.28	0.31	15.38	0.32	14.81	0.31	14.30	0.32	12.62
DK13	Granulite	13.87	0.20	15.77	0.21	14.49	0.21	13.76	0.21	12.08
DK32	Eclogite	12.23	0.25	14.32	0.27					10.74
DK43	Granulite	13.25	0.38	14.69	0.39	13.60	0.39	13.59	0.41	10.17
DK69	Granulite	11.66	0.29			12.60	0.42	12.75	0.32	10.73
DK71	Granulite	12.68	0.45	13.40	0.27	13.15	0.27	12.87	0.22	8.80
DK74	Granulite	9.06	0.23	11.01	0.28	9.56	0.28	9.47	0.45	7.18
DK84	Eclogite	7.31	0.22	9.61	0.34	8.10	0.22	8.14	0.21	6.24

Temperatures are calculated using the temperature coefficients from Valley (2003).

^a2 SD uncertainties are calculated based on the analytical 2 SD uncertainty on the standard UWG-2.

^b2 SD temperature uncertainties are calculated by taking the square root of the sum of the squares of the uncertainties of the individual mineral analyses.

DK 32 and DK 43 each contain one heavier δ^{18} O quartz outlier; these are from quartz inclusions in garnet.

Of all the minerals, rutile reveals the broadest variability in the δ^{18} O raw values measured by ion microprobe (Fig. 5d). The heterogeneous nature of the measurements is mostly due to differences from grain to grain, which as described above is likely biased by crystal-orientation effects.

P–T estimates

Oxygen-isotope thermometry

Oxygen fractionation by a range of different mineral pairs (e.g. Zheng, 1993a,b; Chacko et al., 2001; Valley, 2003) allows the potential of multiple temperatures to be obtained from a single sample. Thus, oxygen isotopes reveal either equilibrium temperatures or information concerning disequilibrium among minerals, which is crucial for interpreting P-T paths. However, the temperature sensitivity of most mineral pairs is poor at ultrahigh temperatures. For instance, at 1000 °C, even if both minerals are homogeneous, an uncertainty of 0.5% in fractionation between the minerals shifts temperature by +128/-97 °C for quartz-rutile, +240/-153 °C for quartz-garnet, +317/-182 °C for guartz-kyanite and +301/-176 °C for quartz-zircon pairs. In the xenoliths, there is local textural disequilibrium in areas affected by K-feldspar + carbonate metasomatism, and in DK32, relict kyanite suggests that at least some of the xenoliths have undergone decompression.

Even though the majority of the minerals appear to be texturally in equilibrium in the studied xenoliths, oxygen-isotope fractionation for the mineral pairs quartz–garnet, quartz–kyanite and quartz–zircon all yield geologically unrealistic temperatures and/or large uncertainties (see Appendix S2). The quartz– rutile pairs, however, yield the most precise temperatures at these high-grade conditions. The $\Delta^{18}O_{quartz-rutile}$ laser-fluorination values range from 2.84‰ in DK74 to 3.56‰ in DK71, indicating a relatively narrow range in temperatures of 1012 + 58/–51 °C to 875 + 23/–22 °C (Fig. 4c; for individual temperature results, see Table 2). A discussion of the other mineral pairs and their uncertainties can be found in Appendix S2.

The rutile oxygen-isotope results obtained by ion microprobe are more heterogeneous than those obtained by laser fluorination, and most likely reflect orientation effects in rutile (see Appendix S2 for a detailed discussion). Based on these observations, only the laser fluorination $T(\varDelta^{18}O_{quartz-rutile})$ are used to evaluate the thermal history. However, the ion-microprobe analyses of rutile can accurately reflect zonation within single crystals. The ion-microprobe analyses indicate that most rutile is homogeneous within a single grain, supporting the use of laser-fluorination data to calculate temperature.

Major-element thermobarometry

To further evaluate the equilibrium conditions of the different xenoliths, major-element thermobarometry was applied to the samples that had an appropriate mineral assemblage. In this respect, it is important to determine which mineral compositions do reflect preeruption equilibrium. Volume diffusion of Fe and Mg during eruption is expected at length scales $< 10 \ \mu m$ (see Hacker et al., 2005 for a more detailed discussion). Mineral zoning in DK32 and DK84 occurs within garnet and omphacite on a large scale ($> 50 \ \mu m$) and therefore represents pre-eruption heterogeneity. The garnet zoning is slightly different in the two samples: DK32 shows a continuous increase in grossular and decrease in both almandine and pyrope from core to rim. The DK84 garnet has outward decreases in grossular and increases in pyrope, but in the outermost \sim 250–300 μ m of the grain, the opposite trend is observed. Garnet zoning similar to that in DK32 was observed in another Dunkeldik eclogite xenolith by Hacker et al. (2005).

The intersection of net-transfer and ion-exchange reactions defines pre-eruption equilibration pressures and temperatures and was used for the thermobarometry. The near-rim compositions of garnet, clinopyroxene and feldspar (see mineral chemistry section) were used to calculate these reactions with THERMO-CALC (Holland & Powell, 1998), assuming no Fe^{3+} in the minerals. If Fe^{3+} is present, the omphacite will take up more Fe^{3+} than the garnet and Fe–Mg exchange thermometry will predict lower temperatures. Conversely, this implies that temperatures derived from Fe-Mg exchange between garnet and clinopy-roxene represent maxima if Fe^{3+} is ignored (Krogh Ravna & Paquin, 2003). Grains used for thermobarometry were carefully selected to avoid any visible signs of disequilibrium. Both the garnet-kyanite and the garnet-omphacite granulites typically contain multiple feldspars. Late, Or-rich (>97 mol.%) feldspar moats around kyanite were not used for thermobarometry. Instead, only coarse matrix feldspar in textural equilibrium with garnet and omphacite was used. Uncertainties in the major-element thermobarometry are a consequence of the assumption of equilibrium among the mineral assemblage, the accuracy and precision of the electron-microprobe analyses, and the calculations using THERMOCALC. The pressure and temperature from garnet-omphacite granulite DK32 have been estimated from Fe-Mg exchange between garnet and clinopyroxene and the nettransfer reaction albite = jadeite + guartz. These two reactions intersect at ~880 °C and 19 kbar (Fig. 6a). Kyanite is present in the sample, but its relict appearance suggests that it is no longer part of the equilibrium assemblage; this it was not used for thermobarometry.

For eclogite DK84, in which various garnet rim compositions have been observed, different garnet-

clinopyroxene pairs yield a wide range of temperatures. The Fe–Mg exchange reaction between garnet and clinopyroxene and the net-transfer reactions albite = jadeite + quartz and diopside + kyanite = grossular + quartz intersect at a pressure of ~18 kbar and 815 °C for one garnet–clinopyroxene pair (Fig. 6b) and ~24 kbar and 1025 °C for a second pair (Fig. 6c). These results further emphasize disequilibrium in DK84.

In comparison with the garnet-omphacite granulite DK32 and eclogite DK84, constraining the preeruption P-T conditions of the clinopyroxene-free granulites is severely limited by the fact that the assemblage garnet-kyanite-sanidine-quartz is stable over a broad P-T range (e.g. DK7; Fig. 7), within which mineral compositions change insignificantly. The only net-transfer reaction applicable to these rocks is 3 anorthite (in sanidine) = grossular + 2 kyanite + quartz (GASP; Ghent, 1976). However, the Ca exchange between garnet and alkali feldspar is negligible in the P-T range of interest (< 1 mol.% variation of anorthite and grossular; see pseudosection description below). The anhydrous nature of the samples suggests that they formed by dehydration melting of metapelites at >900 °C (Patiño Douce & McCarthy, 1998), which yields a loose first-order temperature estimate. Oxygen-isotope equilibria thus provide the more reliable pre-eruption temperature estimates.

The pre-eruption pressure for granulite DK69 can be calculated using the GASP barometer; this reaction yields a pressure of 17 kbar using the $T(\Delta^{18}O_{quartz-rutile})$ of ~930 °C (see below). Feldspar-solvus thermometry can also be applied to DK69. At the ~17 kbar obtained from GASP, the calibration of Elkins & Grove (1990) yields ~925 °C (±3 kbar uncertainty translates to a ±8 °C uncertainty) for matrix plagioclase (An₂₉Ab₆₁Or₁₀) and sanidine (An₀₅Ab₂₉Or₆₆). Using the composition of feldspar rims (An₂₉Ab₄₆Or₂₅ and An₀₄Ab₂₆Or₇₀) included in garnet yields a higher temperature of ~1025 °C at ~17 kbar.

The pre-eruption P-T conditions of five garnetphlogopite websterites were evaluated using equilibria among garnet, orthopyroxene and clinopyroxene. Here, we summarize the results; more details will appear in a separate paper (P. Luffi, unpublished data). Pressures and temperatures were calculated using the Al-in-orthopyroxene barometer of Nickel & Green (1985), based on the distribution of Al between garnet and orthopyroxene, in combination with the orthopyroxene-clinopyroxene solvus thermometer of Tavlor (1998). Nimis & Grütter (2010) demonstrated that this is the most reliable method to assess equilibrium P-T conditions of fertile peridotites and pyroxenites. Only mineral analyses of adjacent grains were used. Results indicate that the websterites equilibrated in the 20-33 kbar/884-1012 °C range, which is similar to our estimates for the eclogite xenoliths. Individual samples yield core and rim P-T values that



Fig. 6. Calculated pressures and temperatures for (a) garnetomphacite granulite DK32, (b) eclogite DK84 grt 1, and (c) eclogite DK84 grt 2. Because DK84 attained only local equilibrium, plots (b) and (c) show reactions calculated using two different sets of adjacent grains of garnet, clinopyroxene and feldspar. All were calculated in the system Na₂O–CaO–K₂O– FeO–MgO–Al₂O₃–SiO₂. Ab = albite; Alm = almandine; Di = diopside; Grs = grossular; Hd = hedenbergite; Jd = jadeite; Ky = kyanite; Qz = quartz; Prp = pyrope.



Fig. 7. Pseudosection for typical garnet–kyanite granulite DK7 calculated in the Na₂O–CaO–K₂O–FeO–MgO–Al₂O₃–SiO₂ system, showing broad stability field of garnet–kyanite–sanidine–quartz assemblage. (1) 2Fsp Grt Cpx Ky Qz; (2) Fsp Grt Sil Qz Spl; (3) Fsp Grt Qz Spl; (4) Fsp Grt Qz Spl Opx. Coe = coesite; Cpx = clinopyroxene; 2Fsp = plagioclase and alkali feldspar; Fsp = feldspar; Grt = garnet; Ky = kyanite; Qz = quartz; Opx = orthopyroxene; Sil = sillimanite; Spl = spinel.

are similar within calibration errors (± 2 kbar, ± 15 –30 °C).

The garnet-biotite gneiss reveals the lowest P-T estimates of the sample suite. Both garnet core compositions + biotite inclusions in garnet and garnet rim compositions + adjacent matrix biotite give 790–850 °C for Fe-Mg exchange, using Holdaway & Lee (1977), Bhattacharya *et al.* (1992) and Perchuk & Lavrent'eva (1983).

Pseudosection calculations

The mineral assemblages dominating several of the studied xenoliths are not appropriate for determining the pre-eruption pressures via major-element thermobarometry. To place some constraints on such pressures, pseudosections were calculated using the bulk composition of the xenoliths. Such pseudosections reveal the P-T region in which the dominant mineral assemblages equilibrated (e.g. for the granulites, including DK7, Fig. 7). In addition, in the case of samples for which thermobarometric results have been calculated, such pseudosections can be used to evaluate the significance of the obtained pressures and temperatures and to show whether these values correspond to bulk or rather small-scale equilibrium. The pseudosections were calculated using Theriak/Domino 01.08.09 (De Capitani & Petrakakis, 2010) with the JUN92 thermodynamic database (based on endmember and solution models of Berman, 1988, 1991) completed with the omphacite solution model of Meyre *et al.* (1997) and the phengite solution model of Keller *et al.* (2005). To evaluate to what extent the topology of the obtained pseudosections depends on the chosen thermodynamic models in the *P*–*T* range of interest, we have also employed Perple_X 7 (Connolly & Petrini, 2002) with the Holland & Powell (1998) thermodynamic database, fitted with the garnet solution model of White *et al.* (2007), the feldspar solution model of Fuhrman & Lindsley (1988) and the omphacite model of Green *et al.* (2007).

The presented results (Figs 7–9 & S1) were calculated in Theriak/Domino; a pseudosection calculated with Perple_X for xenolith DK32 is shown in Fig. S2 for comparison. A comparison of Figs 8 & S2 corresponding to sample DK32 suggests that the pseudosections built with Theriak/Domino and Perple_X are similar in the P-T region of interest, and therefore insensitive to the employed thermodynamic models. Mineral abbreviations in the figures are after Whitney & Evans (2010), and several minor fields in the lowpressure/high-temperature region of the calculated pseudosections are too small to list the compositions and are labelled by numbers.

Whole-rock compositions used in these calculations are shown in Table 1; the models are calculated in the (Na₂O-CaO-K₂O-FeO-MgO-Al₂O₃-**NCKFMAS** SiO₂) system. To preserve consistency with the assumptions in the major-element thermobarometry, the pseudosections were calculated in Fe³⁺-free systems. All pseudosections, except for DK83, were calculated without H₂O due to their anhydrous mineral assemblage. Due to the abundant biotite found in the thin section, it is assumed that DK83 does not represent a restite as interpreted for the other studied xenoliths. To equilibrate for the amount of biotite observed in DK83, 0.85 wt% H₂O was added to its bulk composition. Significantly more water would suppress the alkali feldspar observed in the assemblage; significantly less water would extend the stability of pyroxene against biotite and suppress the garnettwo-feldspar-biotite-quartz assemblage dominating this sample.

Uncertainties in the pseudosections are a consequence of the chosen activity models, the uncertainty in the thermochemical data, the assumption of equilibrium and the assumption that the chosen bulk composition is that with which all minerals equilibrated. In the ideal case of equilibrium, isopleths of the observed mineral rim compositions should intersect in a P-T point in an assemblage field that includes all the minerals in the sample. This is rarely the case in natural rocks, and samples containing zoned minerals, relict phases, evidence for metasomatism and/or two or more generations of a mineral, like the xenoliths studied here, will deviate from this ideal condition. Perhaps, the biggest limitation is that there is no



Fig. 8. Pseudosection and isopleths for almandine, grossular and pyrope for the garnet-omphacite granulite DK32 calculated in the Na₂O-CaO-K₂O-FeO-MgO-Al₂O₃-SiO₂ system; the molar isopleths of the rim garnet intersect at ~960-975 °C and 20-21 kbar. For the pseudosections for this sample and the eclogites, it was assumed that a small fraction of total Fe in omphacite is accommodated as Fe^{3+} and that the jadeite–aegirine binary be approximated as ideal. Because the Meyre et al. (1997) solution model excludes aggirine, the calculated jadeite mol. fraction is greater than analysed in the sample. The blue, red, green and orange bands correspond to $\pm 1 \text{ mol.}\%$ uncertainty around grossular, pyrope, almandine and jadeite values observed in the garnet and clinopyroxene rims. Coe = coesite; Cpx = clinopyroxene; Fsp = feldspar; Grt = garnet;Ky = kyanite; Ol = olivine; Opx = orthopyroxene;Qz = quartz.

internally consistent set of activity models nor a thermodynamic database for silicate melts for the compositions, high pressures and temperatures of interest.

For granulite DK32, the molar isopleths of grossular, almandine and pyrope rim compositions intersect at ~960–975 °C and 20–21 kbar in the equilibrium assemblage of garnet–clinopyroxene–sanidine–plagioclase–quartz–(rutile) (Fig. 8). DK32 also contains kyanite, but the presence of feldspar rims around kyanite and the fact that eclogites from this same xenolith suite record higher pressures and temperatures (Hacker *et al.*, 2005) suggest that the kyanite is relict and that DK32 cooled and decompressed from the sanidine–kyanite–clinopyroxene–garnet–quartz stability field. The core-composition isopleths of the garnet intersect at unrealistically high temperatures above those shown on the pseudosection.

Estimating the pre-eruption P-T conditions for eclogite DK84 is difficult because the garnet has dif-



Fig. 9. Pseudosection for garnet–biotite gneiss DK83, calculated in the Na₂O–CaO–K₂O–FeO–MgO–MnO–Al₂O₃–SiO₂ system. Fields in grey are likely in the supersolidus region and are thus metastable with respect to melt. The solidus is based on a greywacke composition from Vielzeuf & Montel (1994) and Auzanneau *et al.* (2006). (1) Fsp Grt Cpx Phe Qz H₂O; (2) 2Fsp Grt Cpx Phe Qz H₂O; (3) 2Fsp Grt Cpx Bt Phe Qz H₂O; (4) 2Fsp Grt Cpx Opx Bt Qz H₂O; (5) 2Fsp Grt Bt Phe Qz H₂O; (6) 2Fsp Grt Bt Phe Qz; (7) 2Fsp Grt Cpx Dt Qz; (8) 2Fsp Grt Opx Bt Qz; (9) Fsp Grt Cpx Phe Ky Coe; Coe = coesite; Cpx = clinopyroxene; Fsp = feldspar; Grt = garnet; Ky = kyanite; Opx = orthopyroxene; Qz = quartz; Phe = phengite.

ferent rim compositions (Fig. 3a), indicating that only local equilibrium has been achieved. Further complications arise from the presence of secondary alkali feldspar and sanidine not equilibrated with the primary assemblage. These factors explain why the molar isopleths for garnet do not intersect within the calculated stability field for the assemblage observed in the xenolith (Fig. S1).

The pseudosection constructed for garnet-biotite gneiss DK83 is shown in Fig. 9. According to Theriak/Domino, garnet, phengite, omphacite and alkali feldspar are stable in the low-temperature-high-pressure domain of the examined P-T range. There is a P-T field (600-840 °C and 5-14 kbar) in which the assemblage two feldspars-quartz-biotite-garnet \pm H_2O is stable. In the case of DK83, the model is limited by uncertainties in the H₂O contents in the rock (here estimated by the modal biotite) and, thereby, by the approximate constraints on the solidus position. Nevertheless, the temperature range of stability of the observed assemblage estimated from the pseudosection is consistent with conventional thermometry results from this sample. Thus, we conclude that DK83 must have equilibrated somewhere in the 5-14 kbar range.

Metamorphic pressures

To determine metamorphic pressures in samples for which major-element thermobarometry was not possible, we combine the oxygen-isotope quartz-rutile temperatures and the calculated stability field of the mineral assemblages. To ensure that results of such a combination are meaningful, the oxygen-thermometry results are first compared with the major-element thermometry and pseudosections. Individual samples yield consistent temperatures for all of the techniques. For example, for garnet-omphacite granulite DK32, major-element thermobarometry indicates $\sim 890 \pm$ 70 °C/19 \pm 2 kbar, the pseudosection yields ~960-975 \pm 70 °C/20–21 \pm 4 kbar and laser fluorination $T(\Delta^{18}O_{\text{quartz-rutile}}) = 940 \pm 45$ °C. The uncertainties associated with the thermobarometry and pseudosection P-T estimates are similar to those of the quartzrutile thermometry, and the overall results suggest a pre-eruption P-T condition of 900 °C/20 kbar. Exchange thermometry serves as a check for the oxygen-derived temperatures for granulite DK69. Feldspar-solvus thermometry yielded a temperature of 925 ± 50 °C, whereas laser fluorination resulted in $T(\Delta^{18}O_{\text{quartz-rutile}}) = 930 \pm 42 \text{ °C}$. In conclusion, the results from both DK32 and DK69 suggest that the quartz-rutile $T(\Delta^{18}O)$ temperatures can be directly applied to the samples not suitable for conventional thermometry.

For most samples, the mineral assemblage stability field is large (e.g. DK7, Fig. 7). For granulite DK13, $T(\Delta^{18}O_{\text{quartz-rutile}}) = 877 \,^{\circ}\text{C}$ superposed on the stability field of garnet + sanidine + kyanite + quartz indicates a pressure of 12-25 kbar. For DK43, $T(\Delta^{18}O_{\text{quartz-rutile}}) = 930 \text{ °C}$, the same mineral assemblage indicates 13-26 kbar. Xenolith DK71 has a laser-fluorination $T(\Delta^{18}O_{quartz-rutile})$ of 875 °C, corresponding to a pressure of 12-25 kbar. At $T(\Delta^{18}O_{\text{quartz-rutile}}) = 1011 \,^{\circ}\text{C}$, the garnet-sanidinekyanite-quartz-rutile assemblage of DK74 is stable at 15-29 kbar. DK69 is the only clinopyroxene-free garnet-kyanite granulite for which a pre-eruption pressure can be determined. This sample equilibrated at \sim 930 °C, and on the pseudosection, falls in the same pressure range as the other granulites (13–26 kbar); the GASP barometer yields a more precise pressure of 17 kbar. Finally, the 790-850 °C cation-exchange temperature inferred for the garnet-biotite gneiss (DK83) indicates a pressure <14 kbar, based on the pseudosection (Fig. 9). Overall, the equilibrium pressures for the granulites lacking clinopyroxene range from ~ 12 kbar (the stability field of kyanite) up to the clinopyroxene-in boundary (~ 25 kbar at 900 °C).

Depth-profiling U-Pb zircon analyses

Zircon U–Pb SHRIMP depth-profiling analyses were obtained from xenoliths DK32 and DK84 (Table S4).

The depth-profiling technique may allow acquisition of the isotopic signature of the last event to affect zircon and thus may indicate the pre-eruption age of the zircon. The unpolished rims of 15 zircon analysed from the granulite DK32 yielded a lower intercept date of 19.9 \pm 3.3 Ma (Fig. 10a). The zircon dates from eclogite DK84 are more scattered: most of the 20 zircon analysed yielded ²³⁸U/²⁰⁶Pb rim dates from 97.6 \pm 3.7 to 23.6 \pm 1.0 Ma (Table S4), whereas five zircon yielded a lower intercept date of 12.9 \pm 1.0 Ma (Fig. 10b), indistinguishable from the eruption age of 11.5 \pm 0.2 Ma (Hacker *et al.*, 2005), given current uncertainties in intercalibration of the U–Pb and K–Ar decay schemes (Renne *et al.*, 2010).

DISCUSSION

Origin of the Dunkeldik xenoliths

The origin of the Dunkeldik xenoliths is partly obscured by partial melting and infiltration that produced potassium feldspar + carbonate. However, oxygen isotopes measured from the xenoliths provide insight into the fluid–rock interaction history. The δ^{18} O values measured from all the minerals in the studied xenoliths are heavy compared to mantle values, with the weighted-mean values for quartz ranging from 9.5% to 15.4%. These heavy δ^{18} O signatures – including those from garnet cores - mean that even the eclogites DK84 and 1309 must have been altered at shallow crustal depths or have a metasedimentary protolith (most mafic eclogite xenoliths have δ^{18} O values lighter than +9.0%; e.g. Garlick et al., 1971; Deines et al., 1991; Jacob et al., 1994; Schulze et al., 2003b). The oxygenisotope data for the granulites lacking clinopyroxene are consistent with the metasedimentary origin inferred from the bulk chemistry of the samples [molar $Al_2O_3 > (CaO + Na_2O + K_2O)$; Table 1], mineral parageneses, melting history (Hacker et al., 2005) and zircon ages (Ducea et al., 2003). The xenolith suite thus covers a range of crustal rocks that have magmatic and sedimentary protoliths.

Pre-eruption *P*–*T* conditions

Most of this study focuses on the understanding of the last P-T conditions of the various xenoliths prior to their eruption. Textures in the samples argue that the xenoliths were not greatly affected by the host magma during transport to the surface. In addition, ascent rates of the xenoliths are expected to be ~0.1–3 m s⁻¹ (Spera, 1984), and thus, the xenoliths are thought to have cooled rapidly (< < 1 year) to below their closure temperatures for oxygen diffusion. In such conditions, re-equilibration is unlikely during eruption (Hacker *et al.*, 2005) because grain-boundary and volume diffusion in the xenoliths and constituent minerals are expected to be slow even at magmatic temperatures (<1000–1200 °C; e.g. Yund, 1997; Ganguly *et al.*,



Fig. 10. Tera–Wasserburg plots generated using ISOPLOT 3.00 (Ludwig, 2003) of zircon SHRIMP depth-profiling results from (a) garnet–omphacite granulite DK32 and (b) eclogite DK84. Each plot represents a single depth-profiling analysis with its associated error; (c) U–Pb xenolith zircon and monazite ages (this study; Ducea *et al.*, 2003) compared with U–Th–Pb zircon, monazite, xenotime and uraninite ages from the southern Pamir, eastern Hindu Kush and the Karakoram.

1998; Dimanov & Sautter, 2000; Cole & Chakraborty, 2001; Milke *et al.*, 2001; Valley, 2001; Page *et al.*, 2007a,b). For oxygen diffusion, closure temperatures are estimated at > 1500 °C and diffusion distances are

 $< 5 \ \mu m$ for all the analysed phases assuming the fast cooling/eruption rates and using the diffusion characteristics summarized by Valley (2001) and Cole & Chakraborty (2001). Thus, the elemental and isotopic measurements used in this study are considered the representatives of the pre-eruption conditions and not affected by the magma during eruption.

Thermobarometry results from this study and Hacker *et al.* (2005) suggest a broad range of preeruption P-T conditions for the Dunkeldik xenolith suite (Fig. 11). The anhydrous granulites and eclogites yield conditions from ~815 °C at 18 kbar to 1100 °C at 28 kbar and the garnet-biotite gneiss yields 790–850 °C at pressures between 5 and 14 kbar. The websterites appear to have equilibrated at temperatures that overlap those of the eclogites and granulites; however, the pressures recorded in the websterites appear to be 5–10 kbar greater.

The P-T results, combined with the dearth of hydrous phases in most of the xenoliths, corroborate the earlier view (Hacker *et al.*, 2005) that the Dunkeldik xenoliths underwent partial melting and that the granulites with a metasedimentary protolith represent the residue of this melting. The absence of hydrous minerals – combined with the bulk composition – suggests that temperatures were >900 °C for most of the xenolith suite (Patiño Douce & McCarthy, 1998). The single garnet–biotite gneiss is an outlier and represents the coldest end of the suite.

Whereas most of the xenoliths record high-grade pressures and temperatures, several samples reveal evidence of decompression and/or cooling prior to eruption. As described above, xenolith DK32 contains relict kyanite that is not in equilibrium with the twofeldspar-quartz-clinopyroxene-garnet assemblage. This relationship suggests that DK32 left the stability field of sanidine-kyanite-clinopyroxene-garnet-quartz (an assemblage observed in other eclogite xenoliths from Dunkeldik) via decompression and, probably, cooling. Eclogite DK84 also contains evidence for decompression prior to eruption. It yields multiple pressures and temperatures from garnet with different rim compositions, presumably reflecting cooling and decompression from 1025 °C/24 kbar to 815 °C/18 kbar. Finally, we interpret the granulites as melting residues, and in order for these rocks to have undergone phengite and biotite consumption, they must have reached $> \sim 1050$ °C (at > 20 kbar; Patiño Douce & McCarthy, 1998); however, many of the samples record lower temperatures, again suggesting cooling. Granulite DK69 also records multiple temperatures. with garnet inclusions yielding > 1000 °C, whereas the matrix yields a lower temperature near 900 °C.

Miocene Pamir lithosphere

The xenolith pressures of \sim 19–33 kbar indicate that crustal material, represented by Miocene eclogites, granulites and websterites, was taken to mantle depths



Fig. 11. P-T diagram for all of the studied eclogites, granulites and the garnet-biotite gneiss. Lines for samples DK32 and DK84 show the garnet-clinopyroxene thermometer, garnet + quartz = clinopyroxene + kyanite thermometer and the albite-breakdown barometer calculated using THERMOCALC. The black box for DK32 represents the P-T conditions based on the intersection of rim isopleths on the pseudosection computed by Domino, using the thermodynamic models of Berman (1988, 1991). Granulite temperatures are from quartz-rutile oxygen-isotope thermometry and the pressure range is from pseudosections computed using Domino. The DK83 gneiss temperature is derived from garnet-biotite Fe-Mg exchange and pressure is derived from the pseudosection computed by Domino. The darker grey-scale ellipse represents the pressures and temperatures calculated for the websterites. The lighter grey-scale ellipse shows the P-T range represented in the Pamir Mountains gneiss domes (McGraw *et al.*, 2010). The dehydration-melting interval is from experiments by (Vielzeuf & Holloway, 1988; Vielzeuf & Montel, 1994; Montel & Vielzeuf, 1997; Patiño Douce & Harris, 1998; Schmidt *et al.*, 2004; Patiño Douce, 2005; Auzanneau *et al.*, 2006).

(~65–110 km). Late Mesozoic zircon and monazite ages from the xenoliths (Fig. 10c; this study, Ducea *et al.*, 2003) preclude derivation of the xenoliths from Indian crust (e.g. Hodges, 2000) and suggest instead an affinity to the southern Pamir–eastern Hindu Kush–Karakoram–Kohistan–Ladakh continental and oceanic arc rocks of southern Asia (e.g. Parrish & Tirrul, 1989; Hildebrand *et al.*, 1998, 2001; Fraser *et al.*, 2001). In particular, the protracted Tertiary high-grade thermal and magmatic history of the Hindu Kush, Karakoram and southern Pamir appears to be reflected in the xenoliths (Fig. 10c).

Using the six investigated garnet-omphacite granulite and eclogites from this study and from Hacker et al. (2005), the rocks define a geotherm of 12-13 °C km⁻¹ in the 65–110 km depth range (cold geotherm of Fig. 11). The metasedimentary xenoliths are compositionally similar to Barrovian metamorphic rocks exposed in the nearby, southwestern Pamir Shakhdara gneiss dome (Fig. 1c) that reached peak pressures of 8-12 kbar and temperatures of 675-800 °C (Fig. 11; McGraw, 2010). Metamorphism in the Shakhdara dome is Miocene and older (McGraw et al., 2010; Stearns et al., 2011). Similar metamorphic and magmatic ages characterize the basement domes of the eastern Hindu Kush and Karakoram to the south of the Pamir Mountains (Fig. 10c). They reflect a protracted tectonothermal reworking of the upper and middle crust of the southern Asian plate margin, which

is contemporaneous with the UHT–(U)HP metamorphism, melting, and subsequent decompression and cooling of the lower crust, reflected by the xenoliths.

The lower crust of continental cratons can be cold (< 500 °C at 40 km; average thermal gradient of 12.5 °C km⁻¹; Hyndman, 2010), whereas thermal gradients in continent-continent collisions can be as steep as 25–40 °C km⁻¹ (hot geotherm of Fig. 11; e.g. Lee et al., 2004; Root et al., 2005; Hyndman, 2010). The relatively low ratio of heat conduction to advection in Earth's crust means that a typical continent-continent collision is characterized by shallow thermal gradients during thickening stages and by steep thermal gradients during thinning stages (Oxburgh & Turcotte, 1974). For example, during the Miocene, the lower crust of southern Tibet was characterized by a thermal gradient of ~16 °C km⁻¹ (1130–1330 °C/22–26 kbar), as defined by felsic and mafic crustal xenoliths similar to those analysed for this study that were erupted in an ultrapotassic dyke (Chan et al., 2009). In comparison, the lower crust of the central Tibetan Plateau is now hot (>1000 °C at 40 km depth, ~ 25 °C km⁻¹; Hacker et al., 2000), and development of that steep thermal gradient may have required 50 Ma of thermal relaxation assisted by radiogenic heating (LePichon et al., 1997; McKenzie & Priestley, 2008) or magmatic input (e.g. Ding *et al.*, 2003). Over the same time interval, the southern Tibetan Plateau remained refrigerated by subduction (Hetenvi et al., 2007).



Fig. 12. Schematic N–S cross-section of the India–Asia collision zone with speculative deep extrapolation along \sim 73°E. The cross sections represent different times in the Pamir Mountains evolution from the (a) present day, (b) *c*. 20 Ma, (c) *c*. 10 Ma and (d) *c*. 5 Ma. The location of the possible source material for the xenoliths is indicated within the cross-section, and figure (a) shows the seismic data used for interpretations of the deep roots of the Pamir Mountains.

In light of these considerations, the fact that the Pamir Mountains xenoliths define an average thermal gradient of $12-13 \text{ °C km}^{-1}$ (Fig. 11) in a collisional orogen indicates that they had not reached thermal equilibrium corresponding to a mature continentcontinent collision. Instead, the results support the view that the crustal material reached ~1000-1100 °C at >90 km prior to decompressing and cooling to lower pressures and temperatures of $\sim 800 -$ 900 °C/18 kbar and then being entrained in their host magma. This inference is supported by features of the xenoliths described above, such as oxygen-isotope disequilibrium and multiple compositions of garnet within a single thin section.

How the crustal material reached depths in the Miocene greater than the present-day Moho (~65 km depth for the southern Pamir; Mechie *et al.*, 2011) re-

mains an open question. U–Pb zircon data suggest that the material most likely was ablated from the Asian upper plate during the subduction of India (Ducea *et al.*, 2003; Hacker *et al.*, 2005). Alternatively, it may have been introduced to the mantle by intracontinental subduction (Meyer *et al.*, 1998) or could have sunk into the mantle owing to a gravitational instability (Hacker *et al.*, 2005). We exclude DK83, the garnet–biotite gneiss, from consideration in these models because it equilibrated at shallower pressure than the other xenoliths; it was probably located within the mid-crust and collected during eruption.

As described, the P-T history recorded in the eclogitic and granulitic xenoliths implies that the crustal material did not fully equilibrate at peak pressures and temperatures or if the samples did, the record of that equilibration was obliterated by the decompression

and cooling. Thus, the samples must have reached maximum depths and then decompressed/cooled potentially via buoyancy-driven exhumation after melting and melt extraction but prior to the eruption of the xenoliths at *c*. 11.5 Ma. In the gravitational instability model, the denser part of the descending body may have sunk into the mantle, whereas the felsic rocks separated from the dense lithologies and rose buoyantly. In the subduction model, felsic material could have been carried downward with the dense, subducting plate until heating decreased its viscosity sufficiently to allow diapiric rise of the felsic material through the overlying mantle wedge (e.g. Gerya & Yuen, 2003; Keppie *et al.*, 2009; Behn *et al.*, 2011).

Our preferred model (Fig. 12) for the Neogene to recent crustal and lithospheric mantle evolution beneath the southern Pamir calls for early Miocene steep subduction of Indian lithosphere beneath the southern Pamir region, similar to the present-day Hindu Kush slab. This induced local subduction erosion or gravitational foundering of Asian crust, which served as the protolith for the Pamir Mountains xenoliths. The deeply subducted Indian lithosphere slab subsequently broke off. Since the Late Miocene, Indian mantle lithosphere has been underthrusting nearly horizontally, effectively cooling the overlying Asian crust. The leading edge of India is currently outlined by the Pamir Mountains seismic zone (Fig. 1b). This underthrusting Indian mantle lithosphere probably corresponds to the cool upper mantle underneath the inferred from wideangle seismic data Pamir (Mechie et al., 2011).

CONCLUSIONS

Ultrapotassic volcanic rock in the southeast Pamir contains crustal xenoliths scavenged from depths of \sim 40 km (garnet-biotite gneiss) to \sim 90 km (eclogites, granulites and websterites), with most of the granulites. eclogites and websterites intermingled at 60 to 100 km depths. Most of these samples achieved ultrahightemperatures of 1000-1100 °C, leading to mica-dehydration melting. The resulting UHT-near-UHP xenoliths represent the residue of these melts and define an average thermal gradient of 12–13 °C km⁻¹ for depths of 60–100 km. Thermobarometry suggests that the eclogites and granulites cooled and decompressed from >1000 °C to ~900 °C before eruption. The xenoliths may have reached great depth through subduction erosion, intracontinental subduction or a smaller gravitational instability. Regardless, the thermobarometry, oxygen-isotope data and textures indicate that the rocks did not thermally equilibrate at temperatures expected for these depths, but instead cooled to \sim 900 °C. The xenoliths provide a rare window into the fate of crustal material taken to mantle depths and highlight the dehydration melting and subsequent cooling/decompression that occurs when the crustal material reaches P-T conditions at which phengite/biotite are no longer stable.

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SUPPORTING INFORMATION

Additional Supporting Information may be found in the online version of this article:

Figure S1. Pseudosection for eclogite DK84. Excess SiO_2 was added to the bulk composition (Table 1) in order to suppress corundum at high pressure. Corundum is not found in the mineral assemblage. Multiple composition garnet rims were found within a single thin section of DK84; garnet isopleths are drawn for the garnet rim composition $Alm_{47}Prp_{25}Grs_{27}Sps_{01}$. Coe = coesite; Cpx = clinopyroxene; Fsp = feld-spar; Grt = garnet; Ky = kyanite; Ol = olivine; Opx = orthopyroxene; Qz = quartz.

Figure S2. Pseudosection for garnet–omphacite granulite DK32, calculated using Perple_X. Coe = coesite; Cpx = clinopyroxene; Fsp = feldspar; Grt = garnet; Ky = kyanite; Ol = olivine; Opx = orthopyroxene; Qz = quartz.

Table S1. Electron microprobe mineral composi-tions from the Pamir Mountains xenoliths.

Table S2. Individual WiscSIMS ion-microprobe oxygen-isotope spot analyses of unknowns and standards.

Table S3. SHRIMP U–Pb zircon depth-profilingisotopic data.

Appendix S1. Analytical methods used in this study.

Appendix S2. Uncertainties associated with oxygenisotope thermometry using the quartz–garnet, quartz– kyanite and quartz–zircon pairs.

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