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Notes

Eocene partial melting recorded in peritectic garnets from kyanite-gneiss, Greater Himalayan Sequence, central Nepal

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Abstract: Anatectic melt inclusions (nanogranites and nanotonalites) have been found in garnet of kyanite-gneiss at the bottom of the Greater Himalayan Sequence (GHS) along the Kali Gandaki valley, central Nepal, c. 1 km structurally above the Main Central Thrust (MCT). *In situ* U–Th–Pb dating of monazite included in garnets, in the same structural positions as melt inclusions, allowed us to constrain partial melting starting at c. 41–36 Ma. Eocene partial melting occurred during prograde metamorphism in the kyanite stability field (Eo-Himalayan event). Sillimanite-bearing mylonitic foliation wraps around garnets showing a top-to-the-SW sense of shear linked to the MCT ductile activity and to the exhumation of the GHS. These findings highlight the occurrence of an older melting event in the GHS during prograde metamorphism in the kyanite stability field before the more diffuse Miocene melting event.

The growth of prograde garnet and kyanite at 41–6 Ma in the MCT zone, affecting the bottom of the GHS, suggests that inverted metamorphism in the MCT zone and folded isograds in the GHS should be carefully proved with the aid of geochronology, because not all Barrovian minerals grew during the same time span and they grew in different tectonic settings.

The Himalayan mountain belt, which developed during the India–Asia collision from c. 55 Ma, is regarded as a classic collisional orogen. It is characterized by the impressive continuity over 2500 km of tectonic units, thrusts and normal faults, as well as large volumes of high-grade metamorphic rocks and granite exposed at the surface. This constitutes an invaluable field laboratory to unravel the tectonic and metamorphic evolution of crystalline units and the mechanisms of exhumation of deep-seated rocks in orogens.

Several first-order tectonic discontinuities have been recognized in the Himalaya. From bottom to top these are the Main Frontal Thrust, the Main Boundary Thrust, the Main Central Thrust (MCT) and the South Tibetan Detachment (STD) System (STDS), respectively (Gansser 1964, 1983; Le Fort 1975; Burchfiel *et al.* 1992; Hodges 2000; Yin 2006). Among these the MCT and STD bound the Greater Himalayan Sequence (GHS) in between,

containing most of the metamorphic rocks of the Himalaya (Burchfiel *et al.* 1992; Hodges 2000; Mukherjee 2013).

In recent years, more attention has been focused on the internal structure of the GHS where increasing evidence of the occurrence of tectono-metamorphic discontinuities highlights their primary role in the exhumation of the GHS (Goscombe *et al.* 2006; Carosi *et al.* 2007, 2010; Imayama *et al.* 2012; Mukherjee *et al.* 2012; Montomoli *et al.* 2013; Larson *et al.* 2013; Mukherjee, this volume, in prep). Although there are many studies on metamorphism, melt generation and deformation concentrated in the same time span as activity on the MCT and STD (c. 23–17 Ma; Godin *et al.* 2006), much less data on deformation, metamorphism, melt generation and geochronology in the GHS is available for the large time span (c. 30 myr) between collision at c. 55 Ma and MCT-STD activities.

U–Th–Pb metamorphic ages between 50 and 25 Ma on monazite and zircon have been reported only sporadically in the GHS (Hodges *et al.* 1996; Coleman 1998; Vance & Harris 1999; Simpson *et al.* 2000; Catlos *et al.* 2001, 2002, 2007; Kohn *et al.* 2004, 2005; Carosi *et al.* 2010; Groppo *et al.* 2010; Corrie & Kohn 2011; Imayama *et al.* 2012; Rubatto *et al.* 2012).

Melt generation and granite emplacement have been related mainly to the exhumation stage of the GHS (Harris & Massey 1994; Patiño-Douce & Harris 1998) during the Early Miocene (Yin 2006; Searle 2013). Streule *et al.* (2010) recognized in the Makalu area two main phases of leucogranite production: (i) the older one from 24 to 21 Ma and (ii) the most recent one, related to the genesis of the cordierite-bearing leucogranite at *c.* 16 Ma.

Very little evidence of partial melting during prograde metamorphism has been reported until now in the southern part of the belt (Godin *et al.* 2001; Imayama *et al.* 2012). The GHS of eastern Nepal and Sikkim (India) shows the occurrence of melting at the bottom of the GHS (Barun gneiss) at nearly 33–27 Ma (Imayama *et al.* 2012; Ferrero *et al.* 2012; Rubatto *et al.* 2012).

In this paper, we report old ages of monazite up to 41–36 Ma in the lower part of the kyanite-gneiss in the GHS in the Kali Gandaki valley (central Nepal) (Fig. 1) as well as evidence of coeval partial melting as attested by melt inclusions in garnets. Compositional X-ray maps of garnets allow us to better constrain monazite ages and melting within the prograde/retrograde path of the rocks. These new data enable refined tectono-metamorphic evolution of the kyanite-gneiss in central Himalaya and constrain current tectonic models widely accepted for the exhumation of the GHS.

Geological outline

The main structure of the Himalaya can be divided into a few principal tectonic zones, separated by prominent tectonic discontinuities that can be followed along the entire length of the belt (Gansser 1964; Le Fort 1975; Upreti 1999; Hodges 2000; Yin 2006).

Along a north–south structural transect perpendicular to the belt in Nepal, these principal tectonic zones occur as follows: the Terai, the Siwalik (Sub-Himalayan), the Lesser Himalayan Sequence (LHS), the GHS and the Tethyan Sedimentary Sequence (TSS).

The Terai unit is the northern edge of the alluvial plain of the Ganges and Indus rivers (Indo-Gangetic Plain), the foreland basin of the Himalaya with the most recent alluvial sediments (Upreti 1999).

The Sub-Himalayan unit (Siwalik Group) represents the foreland basin, made up by a Tertiary

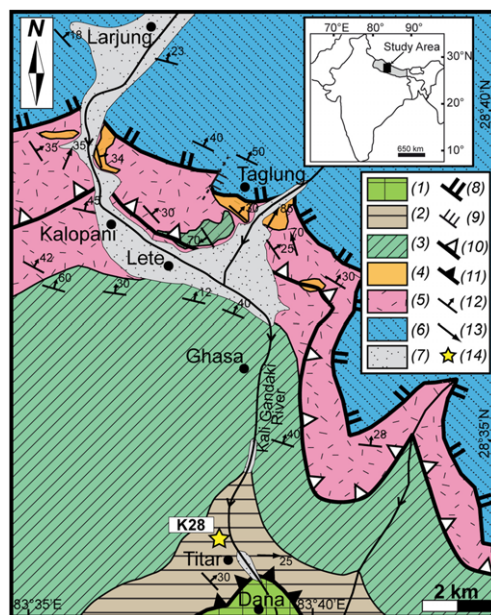


Fig. 1. Geological sketch map of the Kali Gandaki valley (modified after Vannay & Hodges (1996)) with locations of the studied samples. 1, Lesser Himalaya (quartzite); 2, kyanite-gneiss (GHS); 3, calc-silicate and marble (GHS); 4, metapelite (GHS); 5, orthogneiss (GHS); 6, Tethyan Sedimentary Sequence (TSS); 7, alluvial debris; 8, STD; 9, minor normal fault; 10, Kalopani shear zone; 11, MCT; 12, main foliation; 13, object lineation; 14, studied samples.

molasse in a sedimentary sequence that varies from 2 to 10 km in thickness (DeCelles *et al.* 1998; Upreti 1999; White *et al.* 2002; Szulc *et al.* 2006).

The LHS is bound at the base by the Main Boundary Thrust (MBT) and at the top by the MCT, which separates it from the overlying GHS. Both these faults show a top-to-the-SW sense of movement. The LHS mainly consists of lower greenschist to lower amphibolite-facies clastic metasedimentary rocks, organized according to a structurally complex system of fold-and-thrust nappes (DeCelles *et al.* 1998; Hodges 2000) with stratigraphic thickness exceeding 8–10 km, as suggested by palinspastic reconstructions (Schelling 1992). The predominant rock types are impure quartzites and psammitic slates, phyllites and schists, with subordinate impure marbles, metamorphosed mafic rocks and augen orthogneisses (Hodges 2000).

The uppermost tectonic domain to the north is the TSS, tectonically separated from the lower GHS by a system of normal faults and shear zones (STDs) (Burchfiel *et al.* 1992). It comprises a nearly continuous sequence of upper Palaeozoic to Eocene sediments, which were deposited on the

northern passive margin of the Indian plate (Gaetani & Garzanti 1991). The sequence was generally deformed under very low-grade metamorphic conditions, with the highest metamorphic grade corresponding to the greenschist facies, at the base of the sequence in the Cambro-Ordovician rocks affected by the activity of STDS (Crouzet *et al.* 2007; Antolín *et al.* 2011; Dunkl *et al.* 2011).

The metamorphic core of the Himalaya is represented by the GHS, a continuous belt of high-grade metasedimentary and meta-igneous rocks with associated Miocene leucogranites (Carosi *et al.* 1999; Hodges 2000; Visonà & Lombardo 2002; Visonà *et al.* 2012). They range in age from Neoproterozoic to Ordovician, and have been classically divided into three main lithotectonic units (Units 1, 2 and 3) (Searle & Godin 2003).

The base of the GHS (Unit 1) consists of predominantly clastic metasedimentary rocks, represented by biotite-muscovite gneisses, although mica schists and phyllites, calc-schists, quartzites, para-amphibolites and subordinate impure marbles are also present. In the upper part of the unit concordant leucosomes or discrete leucogranitic dykes and sills are found within migmatitic gneisses (Hodges 2000).

The middle part of the GHS (Unit 2) is characterized by middle to upper amphibolite-facies calcareous rocks. The predominant rock type is banded calc-silicate gneiss; other lithologies include marble, calc-schist, quartz-rich psammitic schist, para-amphibolite and orthoquartzite (Hodges 2000).

Unit 3 is constituted by augen orthogneiss and migmatitic gneiss, with a few metasedimentary rocks (Hodges 2000). These metapelites and calc-silicates become more frequent towards the top of the unit, and have been correlated recently with the Cambro-Ordovician sequences of the TSS (Larson *et al.* 2010; Searle 2010). This portion of the sequence is intruded by a dense network of leucogranitic dykes and sills, aged between 22 and 12 Ma, which show an increase in deformation approaching the STDS, at the contact zone with the TSS (Carosi *et al.* 2002, 2007).

The Kali Gandaki valley, in central Nepal, is one of the most studied sections of the chain because it offers an excellent exposure of all the tectonic units (Colchen *et al.* 1986; Vannay & Hodges 1996; Upreti & Yoshida 2005). Here the GHS is c. 10 km thick. It shows a NE-dipping prominent high-grade foliation with SW-verging isoclinal folds (Brown & Nazarchuk 1993; Godin 2003; Kellett & Godin 2009; Searle 2010).

In the Kali Gandaki valley, Unit 1 is represented by kyanite-garnet-bearing gneiss and migmatite that lie in the structurally lowest position and Unit 2 consists of diopside-garnet-amphibole-bearing calc-silicate gneiss, while Unit 3 is characterized by

migmatitic quartzo-feldspathic sillimanite-bearing schists and calc-silicate gneiss which are intruded by variably strained garnet-tourmaline augen orthogneiss sills (Godin *et al.* 2001). The high-grade granulitic and migmatitic metasediments of Unit 1 correlate with the Barun Gneiss in the Everest–Makalu region (Bordet 1961; Schärer 1984; Brunel & Kienast 1986; Lombardo *et al.* 1993; Pognante & Benna 1993; Groppo *et al.* 2012). As a whole the Barun gneiss and its laterally equivalent Kangchenjunga Gneiss and Darjeeling Gneiss (Goscombe *et al.* 2006) form a continuous horizon for at least 200 km along the Himalayan chain, from eastern Nepal to Sikkim and Bhutan. Recent studies documented very similar Grt–Kfs–Ky–Sil rocks at the easternmost margin of the Himalayan belt, 1000 km east of the Barun Gneiss type locality (Guilmette *et al.* 2010), representing a truly significant portion of the metamorphic core of the Himalaya (Groppo *et al.* 2012).

Petrography and microstructures of kyanite-gneiss

Thin sections were obtained from samples K-28 collected in the Kali Gandaki valley close to Titar village (N28°34'13"; E83°38'20", 1787 m above sea level) (Fig. 1). The samples come from the kyanite-gneiss in the lower part of the GHS (Unit 1) involved in the MCT deformation. Samples K-28 (a, b, c) from the kyanite-gneiss were studied in detail. Single garnets used for remelting experiments were picked up from floating garnet grains at the base of the outcrop.

The mineral assemblage of the gneisses is: quartz + plagioclase + biotite + garnet + kyanite + muscovite + chlorite ± sillimanite. Accessory minerals are monazite, xenotime, apatite, zircon, tourmaline, rutile, ilmenite and pyrite. Note that rutile occurs only within garnet and kyanite grains, while ilmenite seems the stable matrix Ti-phase.

Samples K-28 (a, b, c) have a porphyroclastic texture with subhedral to anhedral porphyroclasts of Grt and Ky. These porphyroclasts are surrounded by a spaced and anastomosing foliation marked by recrystallization of biotite and muscovite into small and oriented crystals. The portions between cleavage domains identify areas with less deformation characterized by large crystals of biotite, garnet and kyanite, with different orientations with respect to the main foliation.

Both quartz and plagioclase grains show microstructures testifying to the recovery and recrystallization processes. Quartz grains present lobate grain boundaries, elongate subgrain extinctions and are often squarish (chessboard extinction). Plagioclase feldspar shows lobate grain boundaries,

deformation twins and a lack of internal microfractures. Kyanite is often twinned and sometimes shows undulose extinction. Pinning and windows microstructures (Passchier & Trouw 2005) are observed, indicating Grain Boundary Migration recrystallization at relatively high temperature of deformation. Kinematic indicators, such as kyanite and mica fish (Passchier & Trouw 2005) and shear bands, point to a top-to-the-SW sense of shear related to the ductile shearing of the MCT.

Due to the overprint of deformation after melting, recognition of microstructures linked to the presence of former melt is a difficult task. However, using the criteria recently reviewed by Holness *et al.* (2011) and Vernon (2011), we could detect micro-evidence of former melt: (a) corroded quartz grains rimmed by feldspar in Q-domain; (b) tiny films of feldspars with low dihedral angles, interpreted as pseudomorphs of melt filling pores; (c) 'string of beads' microstructures; and (d) nanogranite inclusion within peritectic garnet. Pseudomorphs after melt films around kyanite (Fig. 2a) and garnet indicate that partial melting involved kyanite as part of the mineral assemblage (see next paragraph).

Needles of fibrolitic sillimanite occur rarely within the fractured kyanite, sometimes at the rims of biotite and muscovite in contact with garnet (Fig. 2a) and at the boundary of plagioclase. This is microstructural evidence of the P – T path of the rock crossing the stability field of sillimanite after the main assemblage Ky – Bt – Grt –melt equilibrated (mineral abbreviation according to Whitney & Evans (2010)).

The mineral assemblage $Qtz + Pl + Bt + Ms \pm Grt \pm Ky$ characterizes the prograde metamorphism

(Vannay & Hodges 1996), whereas the growth of sillimanite occurred later during decompression.

Occurrence of nanogranites in garnet

Melt inclusions trapped in peritectic minerals formed during incongruent melting reactions are a novel tool for the microstructural and microchemical investigation of the genesis of crustal melts (Cesare *et al.* 2011). When slowly cooled, as in migmatites and granulites, these tiny inclusions crystallize in cryptocrystalline aggregates named 'nanogranite' (Cesare *et al.* 2009).

Melt inclusions have been discovered in the kyanite-gneiss (K-28) from this study, as well as in most of the garnets collected from the same outcrop.

Garnet crystals range from 3–14 mm in diameter; those observed in thin sections from sample K-28 are usually smaller than the single garnets collected at the bottom of the rock wall and used for the experiments. Systems of parallel fractures cut garnets at high angle to the main foliation. Biotite overgrowths rim those garnets (Fig. 3a). Most of the analysed garnets contain hundreds of nanogranite inclusions that are grouped in clusters and sometimes show a systematic microstructural distribution. However, garnets free of anatectic melt inclusions have also been observed.

Nanogranite inclusions in garnets are up to 50–60 μm , but most are 10–20 μm . Often the inclusions are crossed by thin cracks, (deformation or decrepitation cracks) (Fig. 3b). The shape of nanogranites from the Kali Gandaki valley is more irregular than those reported (Cesare *et al.* 2011;

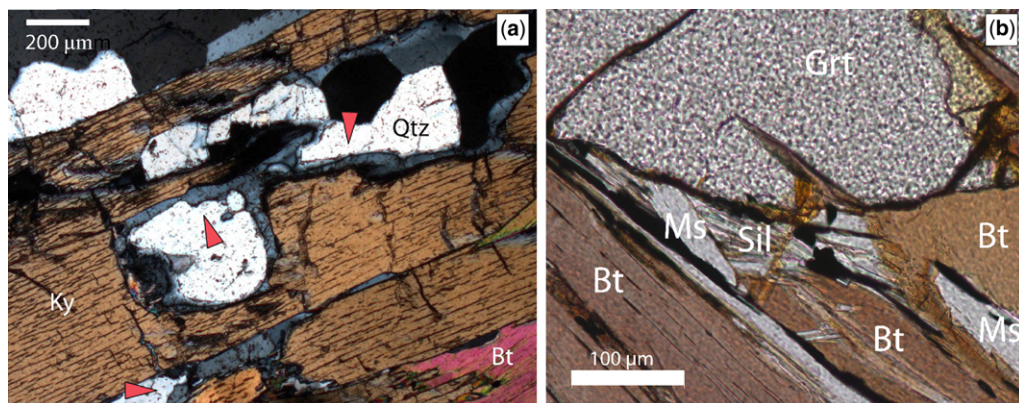


Fig. 2. (a) Thin layers of plagioclase at the boundaries between quartz (Qtz) and kyanite (Ky) constitute pseudomorphs after melt films and indicate the former presence of melt coexisting with kyanite. (b) Needles of acicular sillimanite (Sil; fibrolite) in the Ms–Bt matrix surrounding a garnet (Grt) porphyroblast.

EOCENE PARTIAL MELTING IN THE GHS

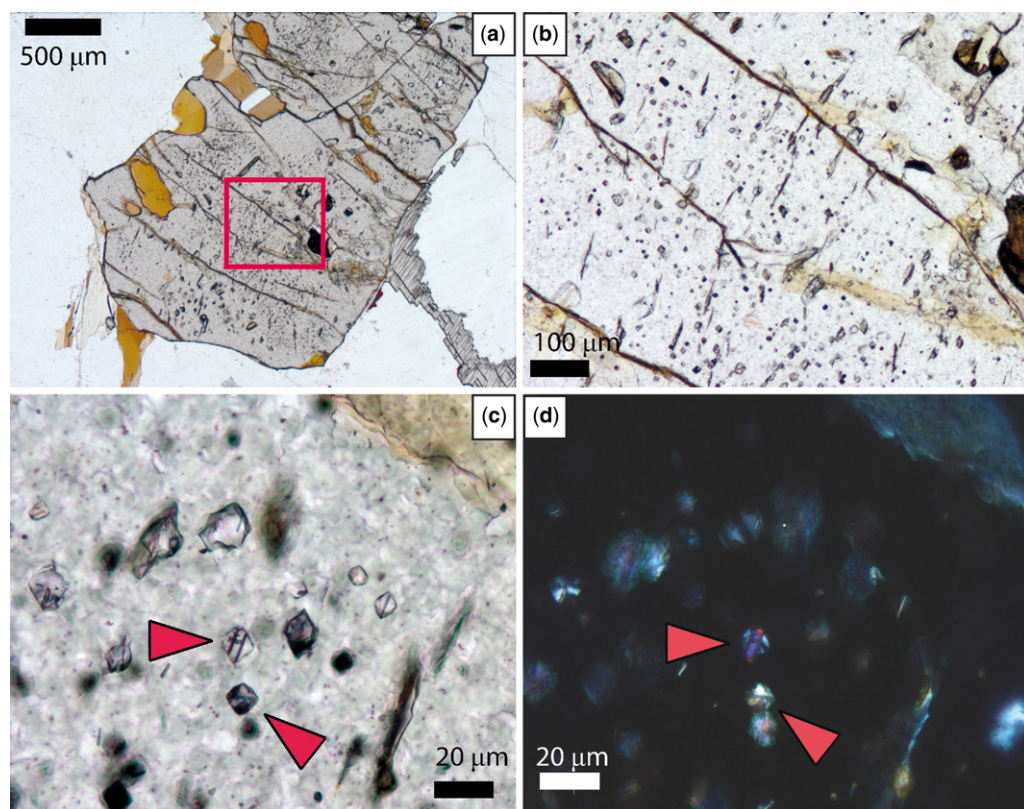


Fig. 3. (a) A porphyroblast of garnet in contact with kyanite, showing abundant inclusions and a set of subparallel, spaced fractures. The outlined box denotes area enlarged in (b). (b) An inclusion-rich area of the garnet. A second set of small fractures, oriented orthogonally to the first, can be observed. (c, d) Close up (plane-polarized light and crossed polars, respectively) of crystallized melt inclusions from the area in (b). Inclusions (arrows) show negative crystal shapes and consist of polycrystalline aggregates.

Ferrero *et al.* 2012; Bartoli *et al.* 2013a, b), and negative crystal shapes are less numerous. Crypto-crystalline aggregates of quartz + plagioclase + muscovite + chlorite commonly constitute nanogranites. Chlorite, instead of biotite, within nanogranites, is unusual if compared with those in the literature (Fig. 3c, d).

Tiny inclusions (1–10 μm) of ilmenite, rutile, apatite, zircon, monazite, biotite, chlorite, quartz and Ca-rich plagioclase can be found within the clusters of nanogranites. The same mineral phases also occur as solid inclusions in nanogranite-free garnets, or even as trapped minerals in nanogranites (Fig. 4). Given the common association of these accessory phases and nanogranite inclusions, we think that they could have played a role as discontinuities on the surface of the growing garnet, thus promoting the trapping of melt drops. Kyanite never occurs as mineral inclusion within garnet.

Garnet compositional maps and chemistry

To obtain the chemical composition of the garnets in the studied samples, X-ray maps and chemical profiles were acquired using a CAMECA SX100 electron microprobe (EMP) equipped with five wavelength-dispersive spectrometers, hosted at the Institut für Mineralogie und Kristallchemie of Stuttgart University. Synthetic and natural crystals and pure oxides were used as standards. Beam current and acceleration voltage were 15 nA and 15 kV, respectively. Garnet analyses were recalculated on the basis of 24 oxygens using the software MINCALC-V5 (Bernhardt 2010). X-ray concentration maps (for Mn, Ca, Mg, Fe) were obtained with a beam current of 50 nA and subsequent computer-aided evaluation. Figure 5a, b presents garnet compositional maps and profiles. Garnet is rich in the almandine component. This mineral is compositionally inhomogeneous and near concentrically

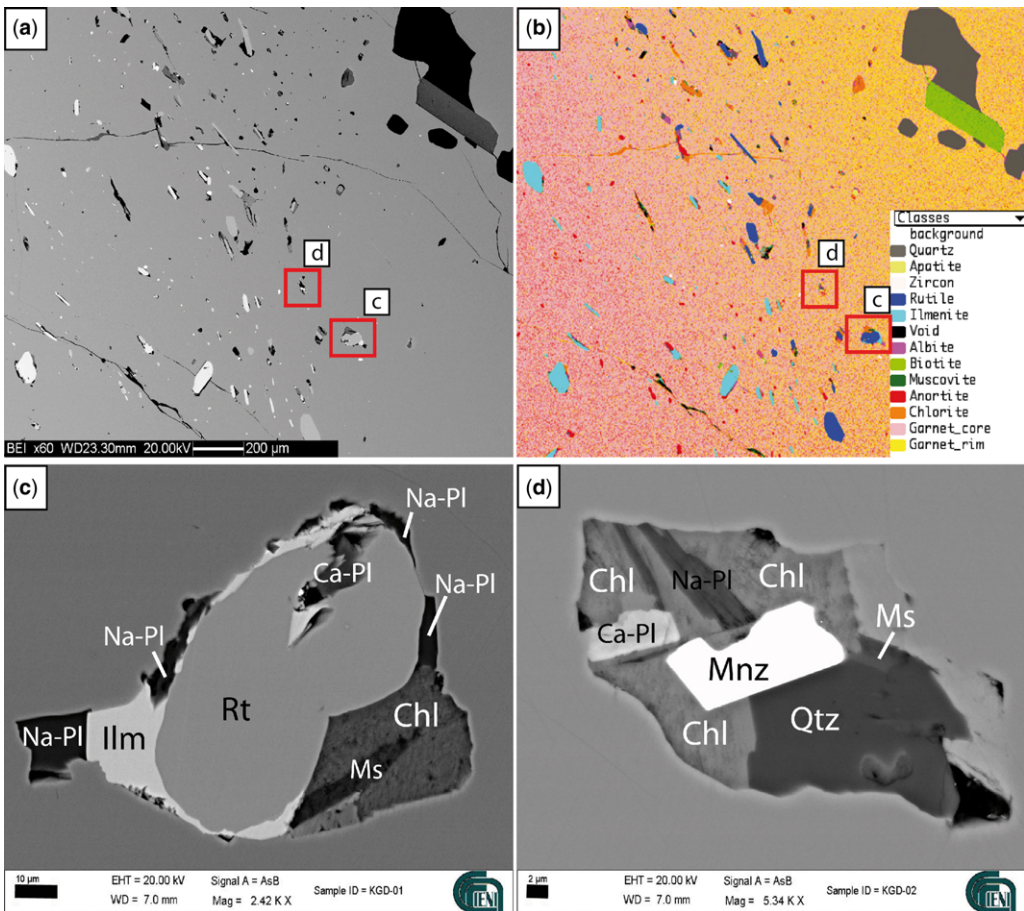


Fig. 4. (a) Backscattered scanning electron microscope (SEM) image of a portion of garnet rich in solid inclusions and nanogranites. The small red boxes indicate the areas enlarged in (c) and (d). (b) The phase reconstruction (based on processing of multispectral image) of the same area as in (a). (c) Crystallized melt inclusion mostly occupied by a trapped rutile partially rimmed by ilmenite. (d) Nanogranite with a trapped crystal of monazite.

zoned (Fig. 5a), with a core where Mn and Ca are higher and Mn and Fe are lower, which is interpreted as indicative of prograde zoning. According to the compositional profile in Figure 5b, the maxima of spessartine and grossular, in the core region, are 9 and 5 mol%, respectively. These values decrease towards the rims, where a spessartine content of 2.5 mol% ($X_{Mn} = 0.025$) and a 3.0 mol% ($X_{Ca} = 0.03$) of grossular are observed. Almandine content increases from core ($X_{Fe} = 0.68$) to rim ($X_{Fe} = 0.76$). The pyrope component increases from core ($X_{Mg} = 0.17$) to near rim position ($X_{Mg} = 0.22$), where it diminishes towards the outer rims (X_{Mg} up to 0.18). The Fe/(Fe + Mg) ratio (hereafter referred as Fe#) decreases from core to near rim (from 0.80 up to 0.77). An upward inflection of Fe# (up to 0.81) and slight increase in X_{Mn} , coupled with a decrease in X_{Mg} (0.18) is

deduced in the outer rims, suggesting partial retrogression/re-equilibration of the garnet (Kohn & Spear 2000).

A similar zoning pattern, but higher spessartine (13 mol%) and grossular (13 mol%) contents, has been observed in both the large garnets used for the remelting experiments. Their equatorial sections show a characteristic sequence of inclusions that is, from the core to the rim: Ca-rich plagioclase + ilmenite → rutile + melt inclusions → quartz + muscovite, biotite (larger inclusions). Chlorite is often found in fractures and is probably a secondary mineral of retrograde origin (Fig. 4b). This sequence matches the above-described concentric chemical zoning of the garnet: the borders of nanogranite-rich areas towards the core correspond to a slight and gradual decrease in Ca and Mn and enrichment in Fe and Mg.

EOCENE PARTIAL MELTING IN THE GHS

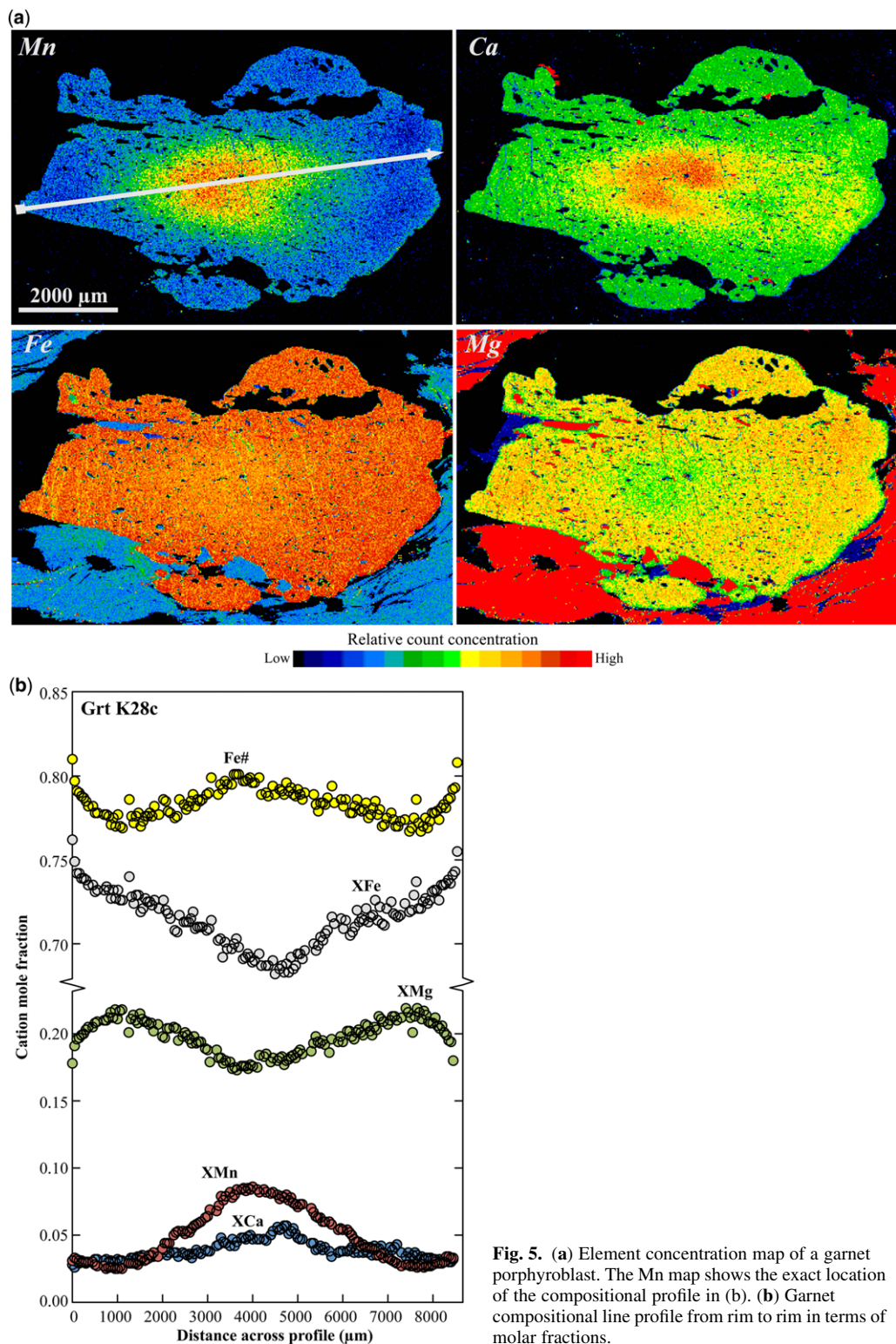


Fig. 5. (a) Element concentration map of a garnet porphyroblast. The Mn map shows the exact location of the compositional profile in (b). (b) Garnet compositional line profile from rim to rim in terms of molar fractions.

Microchemical characterization of melt (remelting experiments)

To obtain the composition of the primary anatectic melts, nanogranites need to be fully re-homogenized, bringing them back to the temperature of trapping or higher. Therefore remelting experiments were performed using a single-stage piston cylinder apparatus at the Laboratory of Experimental Petrology, University of Milan, Italy. Fragments of garnet that contain nanogranites were sealed in a golden capsule with silica powder and then loaded in the apparatus. At the end of the experiments the capsules were mounted in epoxy and uncovered through polishing (Bartoli *et al.* 2013a, b). Successfully remelted inclusions were then analysed using EMP for major elements, with correction for Na and K loss.

We experimented using fragments of garnet crystals coming from the outcrop where sample K-28 was collected. Since the garnets were too large to be loaded into the experimental capsule, they were mounted in epoxy and then cut on the equatorial plane.

In the experimental runs, the main controlling parameter was temperature, which needed to be high enough to cause total rehomogenization, but not chemical interaction with the garnet host. Therefore it is common practice to run more experiments at different temperatures, and to analyse the one that produces total remelting at the lowest temperature. The value of experimental pressure, which does not correspond to the pressure of entrapment, needs only to be high enough to avoid inclusion decrepitation (Bartoli *et al.* 2013a, b), and was increased in the second run to 12 kbar. This value is 2 kbar greater than the pressure estimated by Vannay & Hodges (1996).

For the first experiment (24 h duration at 900 °C and 8 kbar) we cut thick sections from these exposed planes and fragmented them according to nanogranites-rich clusters. Thus we obtained small wafers of garnet *c.* 2.5 mm large and 150–200 µm thick that were loaded into the gold capsule. Up to eight of these garnet wafers, each containing nanogranites, could be loaded in a single capsule. For the second experiment (24 h at 820 °C and 12 kbar) we tried a new approach by drilling small cores of garnet using a diamond hollow-core micro-drill. We obtained four cylinders of garnet of diameter 2.5 mm and up to 4 mm thick. Two were loaded into the capsule.

The experiment at 900 °C produced overheating of the melt inclusions, and partial decrepitation with some interaction with the host garnet (Bartoli *et al.* 2013a, b). The experiment run at 820 °C correctly reproduced the experimental temperature for remelting: here many inclusions appear crystal-free

and fully re-homogenized, with negative crystal shape and without cracks that would indicate overpressure and decrepitation (Fig. 5d). The lack of decrepitation is important to preserve the volatile content of the glass.

Interestingly, despite coming from garnets with very similar composition and zoning, the two experiments provided markedly different compositions for re-homogenized glass (Table 1). The glass produced at 900 °C is granitic (average SiO₂ 71 wt%) and slightly peraluminous (average ASI 1.02). FeO is usually <2.5%, and average wt% oxide contents are Na₂O = 4%, K₂O = 5.2%, CaO = 0.4 and MgO = 0.2. The EMP total is 96–97%, but this value could be an overestimation due to the possible loss of volatiles during inclusion decrepitation (Bartoli *et al.* 2013a, b). Despite the experimental overheating of these nanogranites, the consistent results suggest that the measured granitic composition is reliable.

The inclusions remelted at 820 °C (Fig. 6), which did not decrepitate and therefore preserved their volatile contents, can be defined nanotonalites: they have an average SiO₂ of 66.8 wt% and are strongly peraluminous (average ASI 1.39). Average wt% oxide contents are FeO = 2.78%, Na₂O = 2.4%, K₂O = 1.2% and MgO = 0.45%. A characteristic feature of the chemistry of these melts is an anomalously high CaO content (average 2.6 wt%) with respect to the analysis of other re-homogenized melt inclusions available in the literature. This could be due to a difference in the original chemistry of the protolith, in agreement with experimental evidence (Ferri *et al.* 2009). The composition of nanotonalites resembles the glass produced by García-Casco *et al.* (2003) during the experimental fluid-present melting of a natural pelite. Similar Ca-rich silicic melts in equilibrium with Grt, Ky and Bt were also produced by the fluid-saturated melting at 10 kbar and 740 °C of a Na-free synthetic metapelite (Ferri *et al.* 2009).

Assuming the fluid content of the melts as 100 – EMP total, this value is quite high, around 10 wt% in the inclusions re-homogenized at 820 °C, and supports the fluid-present character of the melting event for this sample.

U–Pb geochronology

Methodology

Monazite, a light rare earth element phosphate (LREEPO₄), is ubiquitous in the Himalayan metapelitic rocks (Tobgay *et al.* 2012; Rubatto *et al.* 2012). This accessory mineral is a powerful geochronometer (Parrish 1990; Rubatto *et al.* 2012) and, due to its ability to undergo dissolution/re-precipitation in response to deformation events

Table 1. EMP composition of melts from the two remelting experiments

	820 °C, 12 kbar						900 °C, 8 kbar				
	KGD-17	KGD-24	KGD-26	KGD-30	KGD-32	KGD-33	HIM 1-1	HIM 1-4	HIM 1-8	HIM 2-2	HIM 2-3
SiO ₂	66.96	66.14	64.51	66.59	66.67	63.81	71.62	72.11	69.88	70.53	66.03
TiO ₂	0.13	0.11	0.00	0.00	0.43	0.17	0.00	0.00	0.05	0.19	0.13
Al ₂ O ₃	15.10	13.45	14.01	13.74	13.88	15.37	13.75	13.36	12.84	12.98	15.39
FeO	2.41	2.15	3.01	2.90	2.70	2.29	2.07	2.19	2.07	2.57	2.26
MnO	0.21	0.08	0.06	0.05	0.02	0.23	0.00	0.00	0.12	0.29	0.17
MgO	0.25	0.48	0.37	0.43	0.84	0.35	0.17	0.17	0.07	0.23	0.16
CaO	2.53	1.40	2.83	2.77	2.01	1.80	0.25	0.25	0.50	0.34	0.55
Na ₂ O	3.01	2.52	1.87	2.11	2.45	4.33	4.37	3.99	3.77	4.02	4.57
K ₂ O	1.52	2.25	1.13	1.61	1.50	1.45	5.66	5.45	5.62	4.34	4.89
P ₂ O ₅	0.43	0.70	0.30	0.00	0.27	0.32	0.00	0.05	0.00	0.05	0.05
Total	92.54	89.28	88.09	90.20	90.77	90.13	97.89	97.57	94.92	95.54	94.20
ASI	1.35	1.47	1.48	1.34	1.49	1.28	1.00	1.03	0.97	1.09	1.11

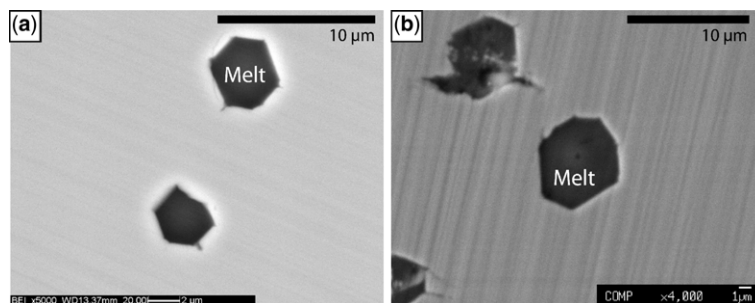


Fig. 6. Example of crystallized inclusions successfully remelted at 820 °C. Inclusions contain a homogeneous melt that has been analysed by electron microprobe.

(Williams & Jercinovic 2002; Dumond *et al.* 2008) and/or changes in P – T condition (e.g. Spear 2010; Spear & Pyle 2002, 2010), offers the possibility to put an absolute time constraint, via *in situ* geochronology (textural geochronology), on the pressure–temperature–deformation history of the rock (Foster *et al.* 2000; Williams & Jercinovic 2002, 2012; Foster & Parrish 2003; Gibson *et al.* 2004).

Sample K-28c was selected for U–Th–Pb geochronology. Prior to isotopic analyses, the thin section was imaged by backscattered electrons to locate, characterize (dimensions, microstructural position and internal structures) and select the most suitable monazite crystals.

Monazite grains were analysed *in situ* by laser ablation–inductively coupled plasma–mass spectrometry (LA–ICP–MS) on 30 µm-thick sections at the CNR–Istituto di Geoscienze e Georisorse U.O.S. of Pavia (Italy) using an Ar–F 193-nm excimer laser (GeolLas 102 from Micro-Las) coupled with a magnetic sector ICP–MS (Element I from Thermo-Finnigan). The full description of the analytical procedure is reported in Paquette & Tiepolo (2007) and Tiepolo *et al.* (2003). Single analyses were performed by one-minute acquisition of the background signal followed by the acquisition of the ablation signal of masses of ^{202}Hg , $^{204}(\text{Hg} + \text{Pb})$, ^{206}Pb , ^{207}Pb , ^{208}Pb , ^{232}Th , ^{238}U , for at least 30 s. ^{202}Hg is acquired in order to correct the isobaric interference of ^{204}Hg on ^{204}Pb and to evaluate the presence of common lead in the sample. The ^{235}U is calculated from ^{238}U on the basis of the mean ratio $^{238}\text{U}/^{235}\text{U} = 137.818$ recently proposed by Hiess *et al.* (2012). Analytical conditions were 12 J cm $^{-2}$ energy density, 3 Hz repetition rate and 10 µm diameter spot size. Technical and data acquisition parameters are summarized in Appendix A. Time-resolved signals were carefully inspected to verify the presence of perturbations related to inclusions, fractures or mixing of different age domains. Laser-induced elemental fractionation and mass bias were corrected using the matrix-matched

external monazite standard (Moacir monazite: Cruz *et al.* (1996) and Seydoux-Guillaume *et al.* (2002a, b)) considering the values, re-calibrated for isotopic disequilibrium, reported by Gasquet *et al.* (2010). Analytical details and results of Moacir standard are reported in Appendix B. External standards and unknowns were integrated over the same time intervals to ensure the efficient correction of fractionation effects. Data reduction was carried out with the software package GLITTER[®] (van Achterbergh *et al.* 2001). In order to have a better estimate of uncertainty affecting the isotopic ratios, the individual uncertainties given by GLITTER[®], were propagated relative to respective reproducibility of the standard following the procedure as reported in Horstwood *et al.* (2003). After this error propagation, each analysis is regarded as accurate within quoted errors. Analytical details and results are reported in Table 2. Data processing and graphic representations were made with Isoplot 3.0 software (Ludwig 2003).

Monazite textural, internal features and U–Th–Pb data

Monazite occurs in different textural positions. Generally, monazite grains were found as inclusions within micas, mainly biotite, and quartz; they show irregular shapes and dimensions up to 180 + 230 µm. Rarely, monazite crystals were also observed as inclusions within garnet and kyanite porphyroblasts. In these cases, monazite shows sub-rounded shapes and fractures and is smaller with respect to the monazite grains included by the other mineral phases. Backscattered electron (BSE) images revealed that monazite grains commonly show bands, often very narrow, or domains with different BSE properties (Fig. 7a, b). Rarely, monazite appears completely homogeneous or shows irregular inner domains with different BSE properties with respect to the surrounding domains (Fig. 7c, d).

Table 2. LA-ICP-MS U–Th–Pb isotope analyses and calculated ages of monazite from the K-28c sample

27 July 2010, IGG-CNR U.O.S. of Pavia					Data for Wetherill plot							Ages							Concordant ages				
Identifier	Monazite number	Textural position	Spot location	Zoning	²⁰⁷ Pb/ ²⁰⁶ Pb	1σ %	²⁰⁷ Pb/ ²³⁵ U	1σ %	²⁰⁶ Pb/ ²³⁸ U	1σ %	Rho	²⁰⁸ Pb/ ²³² Th	1σ %	²⁰⁷ Pb/ ²⁰⁶ Pb	1σ abs	²⁰⁷ Pb/ ²³⁵ U	1σ abs	²⁰⁶ Pb/ ²³⁸ U	1σ abs	²⁰⁸ Pb/ ²³² Th	1σ abs	% U– Pb disc	2σ abs
J127a005	3	Bt	core	hom.	0.05823	0.00101	0.02299	0.00037	0.00286	0.00003	0.8	0.00089	0.00001	538	9.4	23	0.4	18	0.2	18	0.2	20.2	
J127a006	0	Qtz	rim	bright	0.05159	0.00091	0.03695	0.00059	0.00519	0.00007	0.8	0.00162	0.00002	267	4.7	37	0.6	33	0.4	33	0.5	9.4	
J127a007	0	Qtz	rim	dark	0.04679	0.00084	0.04076	0.00068	0.00632	0.00008	0.8	0.00198	0.00003	39	0.7	41	0.7	41	0.5	40	0.7	–0.1	40.6
J127a008	17+	Ky	core	hom.	0.05112	0.00092	0.03893	0.00065	0.00553	0.00007	0.7	0.00160	0.00002	246	4.5	39	0.6	36	0.4	32	0.5	8.3	
J127a009	17+	Ky	core	hom.	0.04893	0.00092	0.04533	0.00079	0.00672	0.00008	0.7	0.00192	0.00003	144	2.7	45	0.8	43	0.5	39	0.7	4.1	
J127a010	18	Ky	rim	dark	0.05532	0.00098	0.03287	0.00054	0.00431	0.00005	0.7	0.00116	0.00002	425	7.6	33	0.5	28	0.3	23	0.4	15.6	
J127a011	18	Ky	core	bright	0.04820	0.00120	0.04614	0.00109	0.00695	0.00009	0.6	0.00189	0.00003	109	2.7	46	1.1	45	0.6	38	0.7	2.5	
J127a012	18	Ky	rim	dark	0.05448	0.00106	0.03156	0.00057	0.00420	0.00005	0.7	0.00115	0.00002	391	7.6	32	0.6	27	0.3	23	0.4	14.4	
J127a013	17	Bt	rim	dark	0.04733	0.00091	0.04407	0.00079	0.00676	0.00008	0.7	0.00199	0.00003	66	1.3	44	0.8	43	0.5	40	0.7	0.8	43.4
J127a014	25	Bt	core	bright	0.04917	0.00095	0.04399	0.00079	0.00649	0.00008	0.7	0.00197	0.00003	156	3.0	44	0.8	42	0.5	40	0.7	4.6	
J127a015	15	Bt	rim	bright	0.05723	0.00109	0.03308	0.00058	0.00419	0.00005	0.7	0.00114	0.00002	500	9.6	33	0.6	27	0.3	23	0.4	18.4	
J127a016	15	Bt	core	dark	0.04780	0.00089	0.04219	0.00073	0.00640	0.00008	0.7	0.00188	0.00003	89	1.7	42	0.7	41	0.5	38	0.7	2.0	41.1
J127a017	15	Bt	rim	dark	0.05765	0.00117	0.02954	0.00055	0.00372	0.00005	0.7	0.00104	0.00001	516	10.4	30	0.6	24	0.3	21	0.2	19.0	
J127a018	36	Bt	core	hom	0.05249	0.00104	0.03499	0.00064	0.00484	0.00006	0.7	0.00147	0.00002	307	6.1	35	0.6	31	0.4	30	0.4	10.9	
J127a019	32+	Qtz	core	bright	0.04772	0.00124	0.03903	0.00096	0.00593	0.00008	0.5	0.00183	0.00003	85	2.2	39	1.0	38	0.5	37	0.7	2.0	38.1
J127a020	32+	Qtz	rim	dark	0.05534	0.00111	0.02477	0.00046	0.00325	0.00004	0.7	0.00090	0.00001	426	8.5	25	0.5	21	0.3	18	0.2	15.8	
J127a021	31	Ms	core	hom.	0.04844	0.00091	0.04208	0.00073	0.00630	0.00008	0.7	0.00192	0.00003	121	2.3	42	0.7	40	0.5	39	0.7	3.3	
J127a022	29	Bt	core	dark	0.05101	0.00097	0.04489	0.00079	0.00638	0.00008	0.7	0.00190	0.00003	241	4.6	45	0.8	41	0.5	38	0.7	8.1	
J127a023	45	Bt	rim	dark	0.04755	0.00087	0.03760	0.00063	0.00573	0.00008	0.8	0.00175	0.00002	77	1.4	37	0.6	37	0.5	35	0.5	1.7	36.8
J127a024	45	Bt	core	bright	0.04868	0.00092	0.04380	0.00076	0.00653	0.00008	0.7	0.00195	0.00003	132	2.5	44	0.8	42	0.5	39	0.7	3.6	
J127a027	50	Bt	core	bright	0.04704	0.00109	0.04129	0.00090	0.00636	0.00008	0.6	0.00188	0.00003	51	1.2	41	0.9	41	0.5	38	0.7	0.5	40.9
J127a028	50	Bt	rim	dark	0.04861	0.00101	0.03656	0.00071	0.00546	0.00007	0.6	0.00162	0.00002	129	2.7	36	0.7	35	0.4	33	0.5	3.7	
J127a029	52	Bt	core	dark	0.04686	0.00127	0.04269	0.00111	0.00660	0.00009	0.5	0.00199	0.00003	42	1.1	42	1.1	42	0.6	40	0.7	0.1	42.4
J127a030	52	Bt	rim	dark	0.05981	0.00130	0.02855	0.00058	0.00346	0.00005	0.6	0.00099	0.00001	597	13.0	29	0.6	22	0.3	20	0.2	22.1	
J127a031	61+	Chl	core	dark	0.04715	0.00090	0.04005	0.00071	0.00616	0.00008	0.7	0.00171	0.00002	57	1.1	40	0.7	40	0.5	35	0.5	0.7	39.6
J127a032	61+2	Grt	core	hom.	0.04674	0.00118	0.03656	0.00087	0.00567	0.00008	0.6	0.00164	0.00002	36	0.9	36	0.9	36	0.5	33	0.5	0.0	36.4
J127a033	61+3	Grt	core	hom.	0.05723	0.00158	0.04136	0.00108	0.00524	0.00007	0.5	0.00144	0.00002	500	13.8	41	1.1	34	0.5	29	0.4	18.1	
J127a034	62	Bt	rim	dark	0.05008	0.00100	0.02278	0.00042	0.00330	0.00004	0.7	0.00093	0.00001	199	4.0	23	0.4	21	0.3	19	0.2	7.1	
J127a035	62	Bt	core	nar. band.	0.04748	0.00110	0.04311	0.00094	0.00659	0.00009	0.6	0.00204	0.00003	73	1.7	43	0.9	42	0.6	41	0.7	1.2	42.3
J127a036	60	Bt	core	bright	0.04828	0.00086	0.04086	0.00067	0.00614	0.00008	0.8	0.00189	0.00003	113	2.0	41	0.7	39	0.5	38	0.7	3.0	
J127a037	56	Bt	rim	bright	0.05050	0.00089	0.04243	0.00068	0.00609	0.00008	0.8	0.00178	0.00002	218	3.8	42	0.7	39	0.5	36	0.5	7.2	
J127a038	75+3	Qtz	core	bright	0.04653	0.00121	0.02093	0.00052	0.00326	0.00004	0.5	0.00093	0.00002	25	0.7	21	0.5	21	0.3	19	0.4	0.2	21.0
J127a039	75+3	Qtz	rim	dark	0.05134	0.00110	0.02176	0.00043	0.00307	0.00004	0.7	0.00080	0.00001	256	5.5	22	0.4	20	0.3	16	0.2	9.6	
J127a040	68+	Qtz	core	dark	0.05442	0.00109	0.04851	0.00090	0.00647	0.00008	0.7	0.00198	0.00003	388	7.8	48	0.9	42	0.5	40	0.7	13.6	
J127a041	68+	Qtz	rim	dark	0.04823	0.00116	0.03534	0.00080	0.00532	0.00007	0.6	0.00172	0.00003	111	2.7	35	0.8	34	0.5	35	0.6	3.0	
J127a042	75	Bt	core	dark	0.04647	0.00128	0.04183	0.00110	0.00654	0.00009	0.5	0.00201	0.00003	22	0.6	42	1.1	42	0.6	41	0.7	–1.0	42.0
J127a043	75	Bt	core	dark	0.04687	0.00182	0.04522	0.00169	0.00699	0.00010	0.4	0.00204	0.00003	43	1.7	45	1.7	45	0.6	41	0.7	0.0	44.9
J127a044	60	Bt	rim	bright	0.04707	0.00087	0.04186	0.00071	0.00645	0.00008	0.7	0.00188	0.00003	53	1.0	42	0.7	41	0.5	38	0.7	0.5	41.4
J127a045	56	Bt	core	dark	0.04758	0.00105	0.04262	0.00088	0.00650	0.00008	0.6	0.00197	0.00003	78	1.7	42	0.9	42	0.5	40	0.7	1.4	41.8
J127a046	56	Bt	rim	dark	0.05383	0.00109	0.04433	0.00084	0.00597	0.00008	0.7	0.00180	0.00003	364	7.4	44	0.8	38	0.5	36	0.7	12.9	

Abbreviations: % U–Pb disc. = % of U–Pb discordance; Prop = propagated error.

Forty *in situ* U–Th–Pb analyses were carried out on monazite grains in different textural positions (Table 2): within quartz (four grains) and muscovite (one grain) along the foliation planes, and included in garnet (two grains), kyanite (two grains), biotite (13 grains) and chlorite (one grain). Fifteen analyses yielded U–Pb concordant data ranging from 45 to 21 Ma, defining two major clusters at about 41 and 36 Ma (Fig. 8).

The U–Pb analyses of monazite included in garnet and kyanite porphyroclasts did not yield concordant data, except for a single value, at about 36 Ma, obtained from a small monazite grain ($18 + 10 \mu\text{m}$) included in a garnet rim (Fig. 8). All these monazite grains are fractured or located near fractures connecting the grains, or portions of them, with the matrix. Fractures and cracks may promote fluid percolation inducing U–Th–Pb resetting (Montel *et al.* 2000; Martin *et al.* 2007; Langone *et al.* 2011). Interestingly, we have observed that garnet and kyanite appear intensely deformed (i.e. fractures, kink bands and/or folds) as a result of stress partitioning around porphyroblasts during deformation (Kenkmann & Dresen 1998).

The older data points, defining the major cluster at about 41 Ma, refer mainly to monazite grains located within the large biotite crystals forming the microlithons (Fig. 8). Integrating the U–Pb concordant data with microstructural observations we interpret these old ages as evidence of the early prograde stage of deformation. Interestingly, two U–Pb concordant data obtained from monazite lying along the foliation planes define the younger cluster at about 36 Ma (Fig. 8) together with the U–Pb concordant value obtained from the small monazite grain included within the garnet rim. These values could be related to the late prograde stage of deformation.

Discussion

Melt inclusions with granitic and tonalitic composition have been found in peritectic garnet belonging to kyanite-gneiss of Unit 1 of the GHS in the Kali Gandaki valley in central Nepal. They testify melt entrapment during garnet growth. Kyanite and garnet were later deformed by a

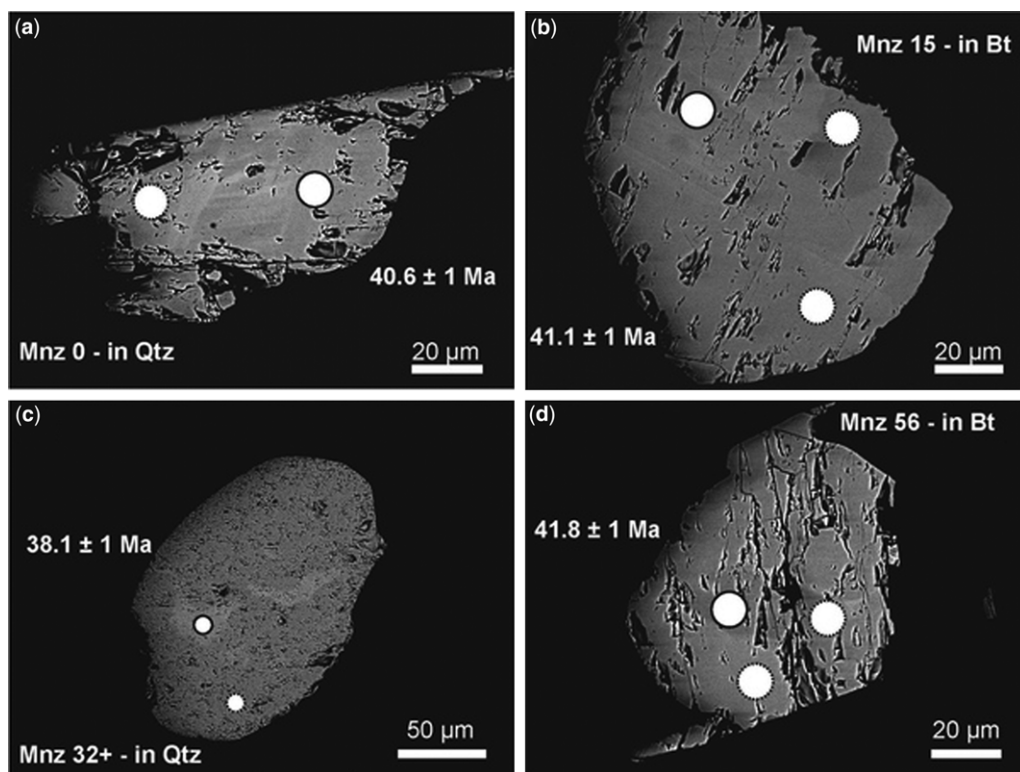


Fig. 7. Backscattered-electron images at high magnification showing monazite shapes and internal features. The white circles indicate locations of the analytical spots providing concordant, continuous contour line and discordant, dashed contour line, U–Pb data.

EOCENE PARTIAL MELTING IN THE GHS

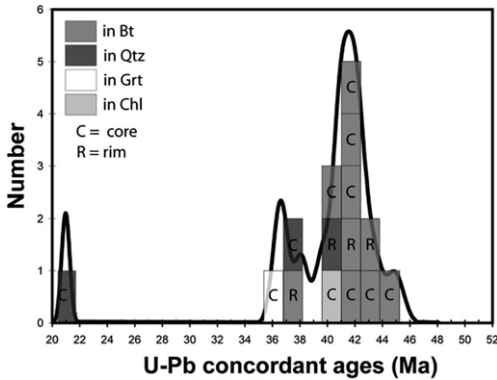


Fig. 8. Probability density plot of monazite concordant data.

pervasive top-to-the-SW shearing linked to Main Central Thrust Zone (MCTZ) activity involving both the upper portion of the LHS and the lower-most part of the GHS. The microstructural position of melt inclusions in the garnets, together with the variation of the chemical composition of garnet, constrained by X-ray maps, allows the constraint of the timing of anatexis during the prograde part of the metamorphism, in the stability field of kyanite.

Remelting experiments were successfully conducted at 820 °C. This temperature is much higher

than those obtained by geothermometric calculations on mineral rims used to obtain pressure (P)–temperature (T) data in the same rocks by Vannay & Hodges (1996), which do not exceed 650–700 °C (Fig. 9). Note, however, that the P – T path of Figure 9 does not enter in the sillimanite stability field, whereas the studied rocks show sillimanite growth after kyanite. They are also higher than the temperatures (675–775 °C) at which García-Casco *et al.* (2003) experimentally obtained peraluminous tonalitic melts. Note, however, that in the latter experiments staurolite is part of the stable assemblage, whereas it is not observed in sample K-28. Although in other studies (Bartoli *et al.* 2013a, b) the experimental temperature of successful remelting of nanogranites agrees well with independently (petrologically or geochemically) constrained temperature conditions for anatexis, in this study we cannot exclude that for kinetic reasons the complete remelting of nanotonalites required temperatures in excess of the trapping values. The quantitative evaluation of P – T conditions of anatexis in sample K-28 is beyond the scope of this research.

In situ U–Th–Pb geochronology on monazite in the same microstructural position as the nanogranites (further confirmed by microchemical zoning in garnet) generates information of the timing of their entrapment at *c.* 41–36 Ma.

Partial melting at 31 Ma in the lower section of the GHS in the Arun valley, eastern Nepal, has

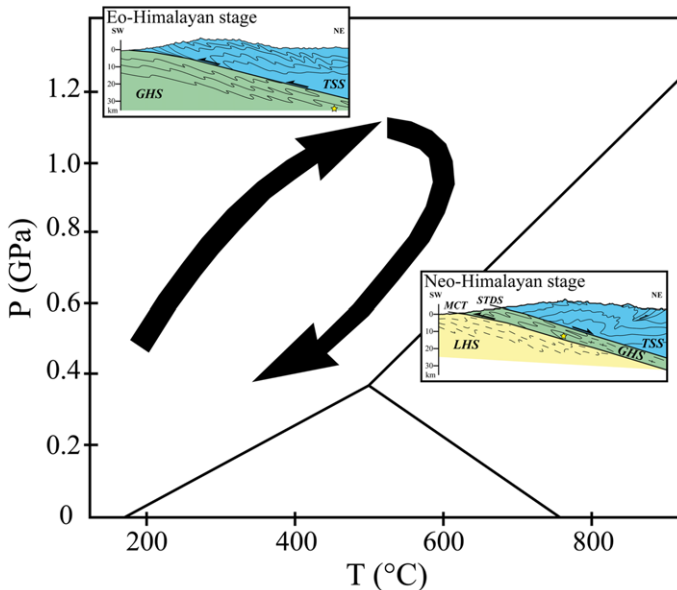


Fig. 9. P – T – t path for the lower GHS–MCT zone as proposed by Vannay & Hodges (1996). Burial of the GHS is attributed to the Eo-Himalayan stage and decompression and exhumation to the Neo-Himalayan stage.

been attributed by Groppo *et al.* (2010) to partial melting before MCTZ activity.

In central and eastern Nepal, partial melting, recorded by melt inclusions included within garnets, occurred during prograde metamorphism in the kyanite stability field in relation to increasing *P* and *T* due to underthrusting of the GHS following continental collision (Fig. 9).

The occurrence of nanotonalite melts with high Ca composition fits well with the Ca-rich melts occurring in eastern Himalaya (Zeng *et al.* 2012) and in southern Tibetan gneiss domes (King *et al.* 2011) developed under fluid-assisted partial melting at higher pressure conditions (>8 kbar) without the contribution of muscovite (Patiño-Douce & Harris 1998).

The well-known and widespread occurrence of partial melting in the GHS, responsible for the generation of Miocene leucogranites, attributed to the exhumation stage during Lower Miocene by activation of MCTZ and STDS, has been put in relation to decompression (Harris & Massey 1994; Hodges 2000; Godin *et al.* 2001; Searle 2013).

This is also in agreement with the finding of several melting stages found by Imayama *et al.* (2012) and Rubatto *et al.* (2012) in eastern Nepal. In this framework relics of Oligocene melts are not limited to southern Tibetan gneiss domes but they have been recorded also within the GHS in the frontal part of the belt.

This earlier melting has been interpreted to have triggered the initiation of the channel flow (Beaumont *et al.* 2001; Mukherjee & Koyi 2010*a, b*; King *et al.* 2011) whereas the Miocene melting testified by the huge amount of high Himalayan granites (Visonà & Lombardo 2002; Visonà *et al.* 2012) has been related to the general exhumation stage of the GHS (Harris & Massey 1994; Hodges 2000).

Monazite ages, related to nanogranites in garnets, at 41–36 Ma (this work) and at 31–28 Ma (Groppo *et al.* 2010; Imayama *et al.* 2012; Rubatto *et al.* 2012) in central and eastern Nepal, respectively, point to a quite larger time span for the occurrence of prograde partial melting in the GHS.

It is worth noting that younger ages of garnets have been reported in the MCTZ in other sections of the belt (i.e. central Nepal: Catlos *et al.* (2001); Bhutan: Tobgay *et al.* (2012); western and central Nepal: Montomoli *et al.* (2013) and Larson *et al.* (2013)). Moreover younger ages have been attributed to a different tectonic setting, that is, retrograde metamorphism during exhumation. The new data show that the garnets in the same level inside the MCTZ do not define a unique retrograde metamorphic isograd because they show different ages and their growth is related to both prograde and retrograde metamorphic processes.

The occurrence of top-to-the-SW shear zones recently recognized in the GHS (Carosi *et al.* 2010; Montomoli *et al.* 2013; Larson *et al.* 2013) influenced the metamorphic history of the tectonic unit and the path of the rocks within it and different slices underwent prograde and retrograde metamorphism at different times. Connecting the metamorphic isograds in the upper part with the ones in the lower part of the GHS could be misleading without an accurate check of the timing of metamorphism. The new data in the Kali Gandaki valley do not confirm the occurrence of simply folded isograds proposed for the whole GHS by Searle & Szluc (2005) and Searle (2010).

The data do not exclude, however, the occurrence of an earlier hot channel active soon after the collisional stage but before MCTZ-STDS activity, even if, as suggested by Prince *et al.* (2001) and King *et al.* (2011), the melt volume started to be enough to lower the viscosity required for crustal flow only after 28 Ma as indicated by the wide occurrence of granites.

Different mechanisms of exhumation have been proposed by Carosi *et al.* (2010, 2013), Imayama *et al.* (2012) and Montomoli *et al.* (2013, 2014) involving the downward and southward progressive migration of ductile shearing within the GHS, allowing the progressive exhumation of crustal slices of the GHS. In this framework the occurrence of folded isograds within the whole GHS, as postulated by the ductile extrusion and channel flow models, is not confirmed.

The interpretation based only on the simple occurrence of minerals in specimens and thin sections at the bottom and at the top of the GHS is not enough to demonstrate the occurrence of folded isograds. The same applies to the mapping of the inverted metamorphism so that mapping isograds in the field and thin sections can result only in an 'apparent' inverted metamorphism.

Conclusion

Melts entrapped in garnets during prograde metamorphism in the kyanite stability field have been documented in the kyanite-gneiss at the base of the GHS in the Kali Gandaki valley. *In situ* U–Th–Pb monazite geochronology constrains their entrapment at 41–36 Ma. Subsequent deformation happened in the sillimanite stability field during decompression and top-to-the-SW shearing linked to the MCT activity.

The composition of Ca-rich melts is in agreement with the occurrence of similar melts found in the Namche Barwa syntaxis and in the southern Tibetan gneiss domes (King *et al.* 2011; Zeng *et al.* 2012) produced by muscovite-absent melting

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under high pressure conditions. This first stage of melting was followed by the widespread occurrence of younger melts producing the well-known high Himalayan Miocene leucogranites during the overall decompression of the GHS.

Monazite ages, related to nanogranites in garnets, at 41–36 Ma in the Kali Gandaki valley and at 31–28 Ma in central and eastern Nepal, point to a quite larger time span for the occurrence of melting during prograde metamorphism in the GHS.

Garnets showing both prograde growth at 41–36 Ma and growth in decompression suggest that the explanation of inverted Barrovian isograds in the MCT zone is not straightforward and should be carefully constrained with the aid of *in situ* geochronology.

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Appendix A. Synthesis of the LA-ICP-MS technical and data acquisition parameters

Inductively coupled-plasma mass spectrometry (ICP-MS)	
Model	Element I, ThermoFinnigan Mat
Type	Single-collector double-focusing magnetic sector field
ICP torch	Capacitive decoupling CD-2
RF power	1200 W
Gas flow:	
cooling	13.5 l min ⁻¹
auxiliary	1.5 l min ⁻¹
Carrier I (Ar)	1.0 l min ⁻¹
Carrier II (He)	1.0 l min ⁻¹
Laser ablation (LA)	
Model	Geolas 200 Q-Microlas
Type	Excimer ArF
Wavelength	193 nm
Repetition rate	3 Hz
Energy density	12 J cm ⁻²
Spot diameter	10 mm
Data acquisition	
Determined isotopes	²⁰² Hg, ²⁰⁴ (Pb + Hg), ²⁰⁶ Pb, ²⁰⁷ Pb, ²⁰⁸ Pb, ²³² Th, ²³⁸ U
Data reduction software	GLITTER (van Achterbergh <i>et al.</i> 2001)

Appendix B. LA-ICP-MS U–Th–Pb isotope analyses and calculated ages of monazite standard

Identifier	Comments	Run position	²⁰⁷ Pb/ ²⁰⁶ Pb			²⁰⁷ Pb/ ²³⁵ U			²⁰⁶ Pb/ ²³⁸ U			²⁰⁶ Pb/ ²³² Th			Ages							
			Ratio	1 σ %	(Prop)	Ratio	1 σ %	(Prop)	Ratio	1 σ %	(Prop)	Ratio	1 σ %	(Prop)	1 σ abor	206Pb/238U	1 σ abor	206Pb/232Th	1 σ abor			
J127a002	MOACIR	Begin	0.05673	0.00052	0.00090	0.64147	0.00473	0.00912	0.08204	0.00062	0.00103	0.02538	0.00033	0.00037	481	8	503	7	508	6	507	7
J127a003	MOACIR	Begin	0.05591	0.00051	0.00088	0.63844	0.00477	0.00950	0.08286	0.00063	0.00124	0.02514	0.00033	0.00037	449	7	501	7	513	8	502	7
J127a004	MOACIR	Begin	0.05728	0.00053	0.00091	0.65189	0.00494	0.00973	0.08257	0.00063	0.00123	0.02537	0.00033	0.00037	502	8	510	8	511	8	506	7
J127a026	MOACIR	Middle	0.05539	0.00057	0.00091	0.64182	0.00554	0.00994	0.08401	0.00063	0.00123	0.02495	0.00033	0.00037	428	7	503	8	520	8	498	7
J127a047	MOACIR	End	0.05745	0.00061	0.00096	0.65477	0.00577	0.01021	0.08262	0.00062	0.00123	0.02543	0.00033	0.00037	509	8	511	8	512	8	508	7
J127a048	MOACIR	End	0.05647	0.00060	0.00094	0.64467	0.00575	0.01009	0.08276	0.00062	0.00123	0.02523	0.00033	0.00037	471	8	505	8	513	8	504	7
J127a049	MOACIR	End	0.05634	0.00060	0.00094	0.63183	0.00561	0.00988	0.08130	0.00061	0.00121	0.02524	0.00033	0.00037	466	8	497	8	504	8	504	7

27 July 2013, IGG-CNR U.O.S. of Pavia

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