Clustering and interfacial segregation of radiogenic Pb in a mineral host – inclusion system

Tracing two-stage Pb and trace element mobility in monazite inclusions in rutile

R. Verberne\textsuperscript{1,2,3}\textsuperscript{*} (0000-0002-5529-1250), Steven M. Reddy\textsuperscript{2,3} (0000-0002-4726-5714), Denis Fougerouse\textsuperscript{2,3} (0000-0003-3346-1121), Anne-Magali Seydoux-Guillaume\textsuperscript{5} (0000-0002-9921-4695), David W. Saxey\textsuperscript{2,3} (0000-0001-7433-946X), William D.A. Rickard\textsuperscript{2} (0000-0002-8118-730X), Zakaria Quadir\textsuperscript{4} (0000-0003-1626-3748), Chris Clark\textsuperscript{3} (0000-0001-9982-7849)

\textsuperscript{1}Centre for Star and Planet Formation, Globe Institute, University of Copenhagen, Øster Voldgade 5–7, DK-1350 Copenhagen, Denmark;

\textsuperscript{2}Geoscience Atom Probe, John de Laeter Centre, Curtin University, Perth, WA 6845, Australia.

\textsuperscript{3}School of Earth and Planetary Sciences, Curtin University, Perth, WA 6845, Australia

\textsuperscript{4}Microscopy and Microanalysis Facility, John de Laeter Centre, Curtin University, GPO Box U1987, Perth, WA 6845, Australia

\textsuperscript{5}UJM-Saint-Étienne, LGL-TPE UMR5276 CNRS, 42023 Saint-Étienne, France

\textsuperscript{*}Correspondence: E-mail: rick.verberne@sund.ku.dk; rick.verberne89@gmail.com
Abstract

Accessory minerals like zircon, rutile and monazite are routinely studied to inform about the timing and nature of geological processes. These studies are underpinned by our understanding of the transfer processes of trace elements and the assumption that the isotopic systems remain undisturbed. However, the presence of microstructures or Pb-bearing phases in minerals can lead to the alteration of the Pb isotopic composition. To gain insight into the relationship between Pb isotopic alterations from inclusions and microstructures, this study focused on inclusions from an ultra-high temperature metamorphic rutile. The studied inclusions are submicron monazites, a common mineral rich in Pb but normally not present in rutile. The sample is sourced from Mt. Hardy, Napier Complex, East Antarctica, an ultra-high temperature (UHT) metamorphic terrane. By applying correlative analytical techniques including electron backscatter diffraction mapping, transmission electron microscopy (TEM), and atom probe tomography, it is shown monazite inclusions are often in contact with low-angle boundaries and yield no preferred orientation. TEM shows the monazite core has a mottled texture due to the presence of radiation damage and nanoclusters associated with the radiation damage defects that are rich in U, Pb, and Ca. Some monazites exhibit a core-rim structure. The rim yields clusters composed of Ca- and Li-phosphate that enclose Pb nanoclusters that are only present in small amounts compared to the core, with Pb likely diffused into the rutile-monazite interface. These textures are the result of two-stages of Pb mobility. Initial Pb segregation was driven by volume diffusion during UHT metamorphism (2500 Ma). The second stage is a stress-induced recrystallization during exhumation, leading to recrystallization of the monazite rim and trace element transport. The isotopic signature of Pb trapped within the rutile-monazite interface constrains the timing of Pb mobility to c. 550 Ma.

Keywords: Rutile, Monazite, Pb mobility, Atom Probe Tomography
Introduction

Rutile (TiO$_2$) is a common accessory mineral that is stable at mid- to lower-crustal metamorphic conditions (Clark et al., 2019; Dachille et al., 1968; Harley, 2016). Zr-in-rutile geothermometry and rutile U-Pb geochronology are increasingly used to provide temperature and time constraints on the evolution of metamorphic terranes (Clark et al., 2018; Kooijman et al., 2010; Meinhold, 2010; Smye and Stockli, 2014; Watson et al., 2006; Zack et al., 2004a). Mineral inclusions in rutile are geologically significant with inclusions of corundum in rutile providing constraints on growth mechanisms (Daneu et al., 2014), whilst the molar fraction of Zr-bearing mineral inclusions can be used to re-integrate Zr concentrations for the estimation of peak metamorphic temperatures (Mitchell and Harley, 2017; Pape et al., 2016). The ability of rutile to trap and isolate inclusions from retrograde metamorphic overprints and fluid alteration events may allow inclusion assemblages in rutile to be used to characterize metamorphic conditions in ultra-high pressure (UHP) and ultra-high temperature (UHT) metamorphic terranes (Hart et al., 2016; Hart et al., 2018).

Despite the uses of inclusion analyses, the presence of small (<1 µm diameter) inclusions in rutile has the potential to compromise compositional and isotopic analysis of the host mineral by techniques, such as Laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS), which analyses significantly larger volumes than the inclusion volume. Little is currently known about the geochemistry of sub-micrometer mineral inclusions or how their presence and composition may affect the analysis of the host. In part, this reflects the difficulty in measuring the trace element and isotopic compositions of such small inclusions. Furthermore, there is little information regarding how such inclusions may evolve as the pressure-temperature conditions of the host rock change over time, or if such inclusions are truly isolated from the matrix of the rock in which the host mineral resides. In this contribution, the structural setting and compositional character of micrometer-scale (200-1100 nm) monazite inclusions within ultra-high temperature metamorphic rutile are studied to provide some constraints on host-inclusion relationships and determine the mechanisms by which compositional heterogeneity in the inclusion may be developed.
The mobility of Pb in monazite has received significant attention in nanoscale studies using transmitted electron microscopy (TEM) and atom probe tomography (APT) (Bingen and Van Breemen, 1998; Fougerouse et al., 2021a; Fougerouse et al., 2018; Grand’Homme et al., 2016; Seydoux-Guillaume et al., 2003; 2012; 2019; Turuani et al., 2022, 2023). However, such studies have typically analyzed the internal parts of large monazite crystals, ignoring the potential complications associated with Pb mobility adjacent to mineral-mineral interfaces that characterize small inclusions.

The complementary use of APT, TEM and electron backscattered diffraction (EBSD) provides unique insight into Pb migration within the inclusions (Reddy et al., 2020). The results obtained show evidence for multiple mechanisms of Pb and trace element mobility, provide important insights into the segregation of Pb to mineralogical grain boundaries and yield important new observations that influence the interpretation of isotopic data from sub-micrometer monazite grains in a mineral – host system.

Geological setting and sample description

The sample used for this study was collected from Mt Hardy in the Napier Complex of East Antarctica (Fig. 1a) and is from an Archean Mg-Al rich metapelitic layer. The Napier Complex consists of a suite of Archean (2990 – 2800 Ma, Black, Williams, & Compston, 1986 Harley & Black, 1987. ortho- and paragneisses that underwent a complex deformational and metamorphic history including two ultrahigh temperature (UHT) metamorphic events at c. 2850 and c. 2580 Ma with their timing constraint by zircon dating. (Clark et al., 2018; Harley, 2016; James and Black, 1981; Sheraton et al., 1987). The c. 2500 Ma UHT event is widely believed to have resulted in the formation of the highest regional-scale temperatures for exposed crustal rocks on Earth, with temperature estimates, coming from the region where Mt Hardy is located, of between 1050–1120 °C (Harley and Motoyoshi, 2000; Hokada, 2001). These high temperatures are thought to have persisted for in excess of 100 Myr (Clark et al., 2018; Harley, 2016). Two younger events, mostly in the southern region of the
Napier Complex have been described by Black et al. (1983); Black et al. (1984). Zircon and monazite U-Pb dating constrain an amphibolite-facies metamorphic event to take place between 1073-1094 Ma. Rb-Sr ages record new magmatic and hydrothermal activity to take place around 522 Ma (Black et al., 1983). A study on galena inclusions in monazite (Turuani et al., 2022) shows the latter event (522 Ma) was more localized while the event at 1073 Ma was more pervasive within the Napier Complex.

The sample is comprised of dry residual layers of porphyroblastic garnet–cordierite–spinel–sapphirine–osumilite (now replaced by symplectites of K-feldspar–cordierite–dendritic orthopyroxene) and quartz–mesoperthite leucosomes (Fig. 1b). Rutile occurs as inclusions in garnet and within both the residual layers and leucosomes, rutile is inferred to form part of the peak UHT assemblage. In this sample, a population of sector zoned “soccer-ball” type zircons interpreted to be metamorphic in origin yielded a concordia age of 2492 ± 7 Ma and a Ti-in-zircon temperature of 915 ± 30°C (Clark et al., 2018). The monazite inclusions reside in two rutile grains. The monazites studied were extracted from rutile in direct contact with garnet, mesoperthite, and traces of biotite. Submicron monazite inclusions are also present in the biotite present in the direct vicinity of the rutile grain of interest. No large monazite grains are present within this sample. However, the earliest monazite in the Napier complex likely crystallized around 3070 Ma (Black et al., 1984).

Methods

Electron backscatter diffraction & electron dispersive X-ray spectroscopy

The thin section has previously been analyzed by LA-ICP-MS. Following LA-ICP-MS, the thin section was re-polished and carbon-coated and analyzed by SEM. SEM characterization was undertaken on a TESCAN Mira3 field emission scanning electron microscope housed at the John de Laeter Centre (JdLC), Curtin University, Perth, Australia and included electron backscatter diffraction (EBSD) and energy-dispersive x-ray spectroscopy (EDS) mapping. Analyses were undertaken with an accelerating voltage of 20 kV and an instrument specific ‘beam intensity’ of 17 using a working distance of 20 mm and 70°
specimen tilt. For mapping of rutile, a 500 nm step size was used and the match units for rutile were obtained from the American Mineralogist crystallography database (801 448-45x) (Swope et al., 1995). Match units for monazite were derived from the crystallographic data of (Ni et al., 1995), following (Erickson et al., 2015). EBSD mapping of the monazite inclusions was done using a 50 nm step size. Post-processing of EBSD data was performed in Matlab®, version R2022b, with the free toolbox MTEX Version 5.8.2 (Bachmann et al., 2010). Images are constructed using local misorientation and the mean misorientation of the rutile grain. The maps were plotted using the ‘roma’ scientific color map (Crameri, 2018). Post-processing involved a noise-reduction by removal of groups of <5 adjacent pixels (‘Wild’ spikes/shards) with crystal orientations within 10° of those of their neighbors. The procedure was followed by applying a 5 × 5 pixel median filter. The filter smooths the orientations in the EBSD data while preserving subgrain boundaries. Slip system characterization was undertaken following established methodologies (Lloyd et al., 1997; Prior et al., 2002; Reddy et al., 2007).

Focused-ion-beam SEM and TEM

Analysis by transmission electron microscope (TEM) and atom probe tomography (APT) requires samples to be prepared in the form of electron-transparent thin foils (TEM) or needle-shaped specimens (APT). These samples were prepared at two different institutes.

Curtin University. TEM and APT specimens analyzed at Curtin University were prepared on a Tescan Lyra3 Ga Focused-Ion Beam Scanning Electron Microscope (FIB-SEM), housed in the John de Laeter Centre, Curtin University. The Tescan Lyra3 comes equipped with an electronically operated nanomanipulator and Pt gas injection system. The FIB was operated with an accelerating voltage of 30 kV for the monoisotopic (mass 69 Da) Ga⁺-source. For backscatter electron (BSE) imaging in SEM-mode, an accelerating voltage of the electron beam was varied between 5, 10 & 20 kV to provide a depth control on the size of the monazite inclusions below the surface. All, TEM and APT, specimens were extracted from one, out of the two, grains that contained monazite inclusions.
The TEM foil was prepared by cutting a vertical slice using the FIB and lifted out using the nanomanipulator. The TEM foil was mounted on a copper half-grid and thinned until electron transparent (< 100 nm) followed by a 2 kV clean-up routine to remove damage and Ga implantation caused by the 30 kV ion beam. After initial imaging by TEM, the sample was further thinned to 30-50 nm to reduce the curvature effect of the inclusion during Scanning TEM (S/TEM) EDS.

Jean Monnet University (UJM). One TEM foil was prepared using a Thermo Fisher Scientific FEI 125 Helios Nanolab 600i focused-ion beam scanning electron microscope (FIB/SEM) hosted by the MANUTECH USD platform in Saint-Etienne (France). Before extraction of the sample, the area of interest is protected with a 1 µm carbon coat. A 30 kV 10 nA Ga-ion beam is used to cut out the foil (15x10x1 µm). The foil is then lifted out and deposited on a Cu half-grid, thinned with an ion beam until ~ 100 nm thickness, and then “cleaned up” at low voltage (2 kV) to remove the amorphous part, resulting in a sample ~60 nm thick.

Transmission Electron Microscopy
TEM analysis at Curtin University was performed on the FEI Talos FS200X Field Emission Gun TEM equipped with a Super-X EDS detector housed in the John de Laeter Centre. The TEM was operated at 200 kV. TEM imaging was conducted in both bright and dark field (BF and DF) modes. The TEM diffraction investigation was conducted with the assistance of Kikuchi patterns that were generated with the convergent beam electron diffraction (CBED). A small spot was used to minimize the beam related damages to the sample, and the sample was tilted to the right diffraction zone axis or two-beam condition to acquire selected area diffraction (SAD) patterns. Both BF and DF TEM imaging were undertaken with the objective apertures after tilting the sample to a diffraction condition within 20°. TEM, STEM, and EDS data acquisition were conducted with the Velox software.

Chemical analysis was done by the attached two pairs of super X detectors. The TEM is fitted with four scanning transmission electron microscopy. (STEM) detection systems: High Angle Annular Dark Field (HAADF), upper Dark Field (DF4), lower Dark Field (DF2), and BF. The contrasts in HAADF are
predominantly made from chemical/phase differences and the contrast in BF is predominantly made from orientation differences. The DF4 and DF2 detectors reveal both chemical and orientation contrasts at different levels.

EDS data was acquired with a beam current of 0.6 nA and collected over 45-90 min for each presented map.

HAADF TEM images provide information based on atomic number (Z) contrast with high z values being brighter.

TEM characterization in UJM Saint-Etienne was performed with a Cs-corrected TEM (NeoARM200F Cold FEG) operated at 200 kV, operated by the CLYM (Consortium Lyon Saint-Etienne de Microscopie) and hosted within the Hubert Curien Laboratory. The instrument is equipped with a wide-angle energy dispersive X-ray (EDX) spectrometer SDD CENTURIO-X from JEOL, two STEM detectors (annular dark field and annular bright field from JEOL and Gatan) with a CMOS camera Gatan Rio (4kx4k), a CCD camera Ultrascan from Gatan (2kx2k), and an electron energy-loss spectrometer (EELS) (Gatan GIF Quantum ER). The resolution in STEM at 200 kV is 0.78 Å due to the presence of a Cs corrector CEOS ASCOR.

Sample preparation for atom probe tomography

For the preparation of APT specimens, monazite was marked by depositing Pt buttons in the middle of the targeted inclusion prior to the deposition of a protective layer (Rickard et al., 2020). APT specimen preparation followed lift-out and sharpening procedures described elsewhere (Rickard et al., 2020; Thompson et al., 2007). A wedge-shaped sample was cut using the ion beam at an angled surface and lifted out using a nanomanipulator. The wedge was brought to the APT specimen holder consisting of a coupon with 22 silicon posts to hold the specimens. The wedge was aligned over the top of a silicon post and a segment was attached using ion beam deposition with a Pt precursor. The wedge above the post was cut leaving ±2 µm of sample on the silicon post. This process is repeated until all material in the wedge has been placed on posts. Following the mounting of specimens on the silicon posts, the specimens were...
then sharpened using an annular milling technique until needle-shaped specimens approximately 2.5 µm in length and less than <100 nm wide at the tip were formed. This was followed by a 2 kV clean-up routine to remove the damaged layer. All specimens were imaged by secondary electron (SE) and backscatter electron imaging (BSE) imaging in SEM-mode only at a working distance of 6 mm and an accelerating voltage of 10 kV.

**Atom probe tomography**

APT analyses were conducted at the Geoscience Atom Probe Facility, Curtin University. This system is a Cameca Local Electrode Atom Probe (LEAP) 4000X HR that allows laser-assisted field evaporation and high mass-resolving power. Data were collected using a 125 kHz pulse rate, a laser pulse energy of 100 pJ, a base temperature of 70° K and an ion detection rate of 0.3% based on (Fougerouse et al., 2018; Verberne et al., 2019). Post-processing was done using Cameca’s Integrated Visualisation and Analysis Software (IVAS) 3.8.0. Reconstructions are based on the density of rutile and using the shank-angle reconstruction mode because of the poly-phased nature of our region of interest (Fougerouse et al., 2021c).

Atoms are evaporated as both single ions and molecular species with a range of different ionization states. Peaks in the mass spectra were labelled per individual isotope for specific ionization states and ranged with a constant width of 0.2 Da. For visualization and compositional analyses, isoconcentration surfaces (Hellman et al., 2000) were computed in IVAS using 1-2 at% for the respective element of interest. These isoconcentration surfaces can be used to isolate each of the individual phases in the datasets. System details, reconstruction parameters and other experimental conditions are summarized in (DR-1) following the recommendation of (Blum et al., 2018a). $^{207}\text{Pb}/^{206}\text{Pb}$ ratios are calculated with corrections for background levels and thermal tails. Pb in these samples can be detected at two different ionization states, $^{207}\text{Pb}^+$ and $^{207}\text{Pb}^{++}$, and corrections are needed to account for the $^{206}\text{Pb}^{++}$ (103 Da) peak tail on the $^{207}\text{Pb}^{++}$ (103.5 Da) peak, but also $^{206}\text{Pb}^+$ (206 Da) and $^{206}\text{Pb}^{++}$ (206 Da) on $^{207}\text{Pb}^+$ (207 Da).
Measurement of isotopic ratios was undertaken using a range of different peak widths and bin sizes following (Blum et al., 2018b).

From the prepared submicron monazites specimens (n=11), APT analysis resulted in three successful runs, with a respective yield of 70, 45, and 9 million detected ions. The analysis of interfaces provides a challenge as the interface is a weak spot in the material. The 3D reconstructions show that in all specimens the monazite is encapsulated within rutile indicating that only the rim of the monazite inclusions was analyzed. Monazites not encapsulated by rutile in the APT specimen likely delaminated during the analysis. The datasets were evaluated by generating isoconcentration surfaces that subdivide the specimens into three components - rutile, monazite and the interface. This allowed for the separate interpretation of the mass spectra of each component.

Results

Petrography and analysis of the rutile host

The studied rutile grain occurs in a rock consisting of residual layers and leucosomes. Monazite inclusions are found in two rutile grains within this thin section, as well as in the directly surrounding biotite (Fig. 2b, c). No monazite inclusions were found in the garnet or mesoperthite. The rutile grain studied is approximately 250 x 200 µm in size, with at least 46 sub-micron monazite inclusions visible at the surface. EBSD mapping of individual monazite grains shows no evidence for a crystal-preferred orientation of the inclusions in either the rutile or the surrounding biotite (fig 2d).

Several different microstructures are revealed by EBSD mapping. Twin boundaries are observed in two orientations corresponding to a 65° rotation and belong to the \{101\} <100> twin relationship. In addition, a network of low-angle boundaries (Fig. 2d, e) is visible. The low-angle boundaries have a misorientation axis of (010), with the dispersion of 1-2°, corresponding to the \{101\}<201> slip system (Fig 2e). The LAB are straight (LAB 1) or bent around the monazite inclusions (LAB 2).

Atom probe observations of the rutile host. Fe and Zr, which are common trace elements in rutile, are detected within the mass spectra. U and Pb are not observed above the background (Fig 6). In
the 3D reconstruction, no heterogeneities can be observed within the rutile matrix this includes ranging
the theoretical U and Pb peaks. The rutile mass spectra are comparable to those obtained during previous
APT studies of rutile e.g., (Verberne et al., 2019; Verberne et al., 2023).

Structural and chemical analysis of the monazite inclusions

The examination of two TEM thin-foils, both incorporating a single monazite inclusion, indicates
variations in the texture and chemical associations of the outer rim of monazite in the two inclusions. In
one monazite, there is a clear core-rim structure (Fig 3). In contrast, the second monazite inclusion only
records the ‘core’ structure throughout the entire inclusion (Fig 4). In both monazites, the core has a high
density of clusters, while the rim structure has a lower density of clusters but this records a discrete
compositional variation (Fig 4, 5).

Trace elements observed within the monazite APT mass spectra (Fig. 6) include Ca$^{++}$ (20 Da), Mg$^+$ (24
Da) and Li$^+$ (6 Da). $^{238}$U was detected as $^{238}$UO$_2^{++}$ at 135 Da and no peak was present for $^{235}$UO$_2^{++}$. Pb in
monazite is observed with two different ionization states: $^{206}$Pb$^+$ and $^{207}$Pb$^+$ (206 and 207 Da) as well as
$^{206}$Pb$^{++}$ and $^{207}$Pb$^{++}$ (103 and 103.5 Da). $^{208}$Pb$^+$ and $^{208}$Pb$^{++}$ were not observed above the background.

Furthermore, no Th-bearing species were identified within the mass spectrum.

Chemical heterogeneities in the monazite core. The inner region of the monazite exhibits a
mottled texture identifiable in BF mode (Fig. 4). In one monazite, this texture is only present in the core-
region, whereas this texture is present throughout the second monazite grain imaged by TEM. The
mottled texture is due to the presence of dark (in BF mode) and bright (in DF mode) spots. These reflect
radiation damage and crystal defects as well as nanoclusters ± 5 nm in size. STEM-EDS revealed some of
these nanoclusters are composed of trace elements (U, Pb, and Ca) (Fig. 7a, b).

Trace element distribution in the monazite rim. This rim is only visible in one of the two
imaged monazites (Fig 4b). The monazite rim is ~100 nm wide. Here, the matrix of the monazite appears
pristine while hosting several nanoclusters 20-30 nm in diameter (Fig 5). Lattice imaging of the clusters
shows a continuous lattice consistent with monazite (Fig 5d-f). Here, we differentiate between a cluster
and inclusion based on similar properties as mentioned in Turuani et al., 2023. The presence of a grain boundary is indicative of an inclusion while a continuous lattice/absence is an indication of a cluster. The clusters in the monazite consist of two types of darker clusters as seen in the HAADF images (Fig 4b, 7c, d) and sometimes encompass one or two bright spots. One type of darker cluster was shown to be rich in Ca (Fig 7c). However, the second type of nanocluster did not yield an EDS response. This indicates the major constituents of this cluster may have an atomic number that is too low for S/TEM-EDS (< Z = 6). Alternatively, this could point to a void and represent porosity within the cluster. The bright spots within both these clusters yielded an increased concentration of Pb (Fig. 7c).

The 3D reconstruction of the monazite APT reveals two types of clusters in the monazite rim (Fig. 8). The first type is dominated by Ca. These clusters have a diameter of 20-30 nm containing up to 10 at. % Ca (Fig. 9). Ca-rich inclusions are often accompanied by an increase in Pb of up to 2-3 at. % as well as Y, up to 1.2 at. %. However, not all Ca-rich inclusions show an increase in the Pb signal. The increase in Ca and other trace elements is compensated by a depletion in REE and O with P remaining relatively constant. Pb is spread over four peaks within the mass spectrum, resulting in insufficient counts for meaningful determination of $^{207}\text{Pb}/^{206}\text{Pb}$ ratios after correcting for the static background and the tails of the preceding peaks. Therefore, despite the presence of Pb within the inclusion, no isotopic composition could be obtained.

The second inclusion type is enriched in Li\(^{+}\), with values up to 30 at. % Li (Fig. 9). These inclusions are ~20 nm in diameter. Li-rich inclusions have Pb concentrations up to 0.6 at. % and therefore incorporate less Pb than the Ca-rich inclusion, yet have elevated levels compared to the monazite matrix. Like the Ca-rich inclusions, the increase of Li and traces is compensated by a depletion in REE and O with P remaining at roughly 10 at. %. (Fig. 9). Pb isotopic compositions could not be obtained from the Li-inclusions for similar reasons as with the Ca-inclusions.

The rutile-monazite interface
The interface between rutile and monazite inclusion is sharp and coherent (Fig 4a). Chemical mapping of the interface by TEM shows the enriched zone in the interface between the rutile and monazite is approximately 2 nm wide at the narrowest part of this boundary.

STEM-EDS analysis was applied to further characterize the different features present within the monazite. The interface between rutile and monazite with core-rim texture yields 2-4 at. % Pb based on a line profile across the interface (Fig. 7d) and with no signals above the detection limit for other trace elements.

The mass spectra obtained from the interface show components of both the monazite and rutile mass spectrum. The boundary is enriched in trace elements including U, Pb, Ca, Li, Mg, Si and Al. Ca and Pb concentrations reach 2-3 at. % in all three specimens. The trace element distribution at the interface of specimen 3 is heterogeneous, most clear for trace elements with a lower overall concentration, e.g. Si and Al (Fig. 10), but similar distributions can be observed for Pb (Fig. 10).

The heterogeneous distribution of trace elements in the interface of specimen 3 is visualized in the reconstruction and measured along a 1d profile (fig 10). The combined concentrations of Al + Si are shown and a contrast in concentration is visible from the apex of the tip to the lower part of the reconstructed volume. The transition from low to higher concentration is accompanied by linear features enriched (labelled A, B, and C) in trace elements. Additionally, clusters of Mg are present at this interface. These clusters contain up to 20 at. % Mg and have a Ca concentration of ~4 at% (Fig. 9), which is higher than the average concentration within the interface. The width of the interface was estimated to be ~8 nm based on the Pb concentration profile.

**Pb isotopic compositions of interfacially segregated Pb**

The mass spectra of the interface between rutile and monazite show $^{206}$Pb and $^{207}$Pb are present as both Pb$^+$ and Pb$^{++}$. The 3D reconstruction shows the Pb is homogeneously distributed along the interface of specimens 1 and 2 and heterogeneously in specimen 3 (fig 10). Pb isotopic signals were retrieved from
the interfaces of all three specimens. The background and thermal peak tails corrected $^{207}$Pb/$^{206}$Pb ratios are: 0.0900 ± 0.0198 (2σ) for specimen 1, 0.1016 ± 0.0224 (2σ) for specimen 2 and 0.1733 ± 0.0156 (2σ) in specimen 3. Hence, the Pb isotopic signals and calculated $^{207}$Pb/$^{206}$Pb ratios for specimens 1 and 2 are identical within error. Specimen 3, however, yields a higher ratio. Calculated model ages for respectively specimens 1, 2 and 3 are 1370 ± 420 Ma (2σ), 1590 ± 420 Ma (2σ) and 2580 ± 150 (2σ). U in the interface was detected as $^{238}$UO$_2^{++}$ (135 Da). The concentration of U compared to Pb is very low with a $^{238}$U/$^{206}$Pb ratio of less than 0.04 in all three specimens. Therefore, it is expected the influence on the $^{207}$Pb/$^{206}$Pb ratio is negligible.

Discussion

Monazite inclusions in rutile

Inclusions typically occur either as a product of exsolution or incorporation during growth. In the rutile studied here, exsolution is unlikely since monazite is present in different host phases (rutile and biotite). The monazite lacks a crystal-preferred orientation that is expected when monazite forms via an exsolution mechanism. Furthermore, P and REE concentrations in rutile are typically very low, with any significant amounts of REE only reported in extra-terrestrial Niobian-rutile (>6 wt. % Nb$_2$O$_5$) (Hlava et al., 1972). Therefore, the monazites are interpreted to be primary inclusions trapped within the rutile.

The encapsulation of monazite within rutile potentially shields the monazite from external processes (Daneu et al., 2014; Hart et al., 2016; Hart et al., 2018). However, the measured Pb isotopic compositions of monazite at inclusion interfaces are different. The rutile host is characterized by the presence of several twin boundaries and LABs, and these could aid in the diffusion of elements in rutile (Verberne et al., 2022). As shown, some monazite inclusions are in direct contact with these LAB’s (Fig 2). Potentially, these inclusions would no longer be fully shielded with trace elements able to migrate out of the rutile-monazite interface. Depending on the timing, inclusions within a single rutile grain could record different isotopic signatures.

Formation of trace-element enriched nanoscale clusters

14
Nanoclusters in the monazite core. The two types of monazite inclusions are: one with a mottled texture due to radiation damage and one with a distinct core-rim structure (Fig. 4, 5). S/TEM EDS shows that some of the dark and bright spots that lead to the mottled texture are enrichment in Ca, U, and Pb (Fig. 7). The texture and formation of trace element enriched clusters are explained via entrapment in defects generated via radiation damage (Seydoux-Guillaume et al., 2018; Seydoux-Guillaume, 2019). The damaged regions stand out due to structural difference with the host monazite while the bright spot (HAADF) stands out due to being enriched in trace elements, creating this mottled texture as previously observed by others (Seydoux-Guillaume et al., 2019; Turuani et al., 2022; 2023).

Clusters within a matrix typically represent a low-energy configuration and are preferable sites for trace elements compared to the crystal lattice of the host. With the rutile present during UHT metamorphism at 2580 Ma (Clark et al., 2019), the temperature conditions are met for Pb diffusion into clusters (as observed in Seydoux-Guillaume et al., 2019).

The mottled texture therefore is the result of an interplay between the formation and healing of radiation damage sites, and the capture of trace elements in these sites during high-temperature events, which allow for more rapid diffusion of elements.

Nanoscale clusters in the monazite rim. The rim hosts Ca-rich and Li-rich nanoscale clusters. Given that the lattice of the Ca-rich cluster is the same as the host monazite (Fig 5) with relatively high REE concentrations these clusters are interpreted to be a solid-solution within monazite rather than represent a separate phase.

The Li-rich nanoscale cluster has low REE concentrations (<4 at. %) with Li concentrations between 24 to 30 at% and P concentrations of ± 10 at. % (fig 9). In the literature, we found no reference for the formation of Li-phosphate in monazite. Nevertheless, the composition of the Li-rich cluster may indicate the formation of a separate Li-phosphate phase.

The Mg-rich clusters are only present within the interface and consist of 20 at. % Mg, 10 at% P and, 4 at% Ca while showing a depletion in REE concentrations compared to the monazite host. As is the case
for the Li-phosphate cluster, the high concentration of Mg suggests the formation of a new phase, via a
similar mechanism to the formation of the Ca- and Li-phosphate clusters.

**Pb-rich nanoclusters.** Bright spots in HAADF mode associated with the Ca- and Li-rich clusters are rich in Pb based on STEM-EDS (Fig 7). Furthermore, APT based proximity histogram of the Ca- and Li-rich cluster also shows elevated values of Pb. The lack of a $^{204}$Pb and $^{208}$Pb signal indicated there is no to a negligible $^{232}$Th or common Pb component. Potentially, these Pb-rich nanoclusters reflect the same clusters observed within the monazite core. The Pb nanocrystals within the Ca- and Li-phosphate cluster have the same general appearance as Pb present in Si-rich cluster recently described in zircon by (Kusiak., 2015, Kusiak et al., 2019; Lyon et al., 2019), called nanospheres and monazite (Turuani et al., 2022; Turuani et al., 2023) referred to as nanocrystals. Considering we observe a similar presence of Pb in the core that is related associated with U decay, it is unlikely these are primary Pb phases. Furthermore, since we lack data on the structure or valence state of the Pb and associate the Pb with U decay these Pb-rich particles do not fall within the definition of metallic Pb nanospheres (Kusiak 2015, Kusiak et al., 2023).

These Pb-rich nanocrystals are intimately linked to the Ca, and Li-rich phosphate clusters, which raises questions about the timing of their formation. One possibility is concurrent formation, with Ca, Li, and Pb migrating at the same time. However, this seems unlikely, as the Pb-rich nanocrystals in the rim are not observed outside of the Ca and Li-rich clusters. If the Pb-rich nanocrystals are indeed the same as those in the core this indicates that during the formation of the rim and the Ca- and Li-rich clusters, the Pb nanocrystals become trapped, preventing small volumes of Pb migrating towards the interface.

**A Pb mobility model**

The texture and compositional variations of the submicron monazite inclusions in rutile provide insight into the tectonothermal history of the studied sample. First, the mottled texture of the monazite (core) is ascribed to both a combination of the presence of radiation damage and the segregation of trace elements into those radiation defects leading to the formation of nanoscale clusters (Seydoux-Guillaume et al., 2019; Seydoux-Guillaume et al., 2002; 2004; 2018; Turuani et al., 2023). Depending on the time of
monazite growth and the diffusion of trace elements into defects the clusters rich in Pb can be dated if
enough time has passed to accumulate radiogenic Pb. This diffusion process is the first stage of trace
element mobility.

Second, in some of the monazite crystals, the rim yields a different distribution of clusters. The texture
indicates partial recrystallization of the monazite rim and encompasses Ca- and Li-phosphate clusters as
well as Pb-rich nanocrystals. The presence of a recrystallized rim in only some of the monazite crystals
shows that the reaction is localized.

Rutile and other repository minerals preserve inclusions due to their low compressibility (Mosenfelder et
al., 2005). When exhumed, rutile exerts an overpressure on inclusions (Hart et al., 2016). Differences in
compressibility between the rutile host and the inclusions will result in a stress acting upon the monazite
(Huang et al., 2010; Zaffiro et al., 2019). Therefore, one explanation could be that the formation of the
rim in the monazite is the result of localized stress-induced recrystallization. Alternatively, the presence
of Li and to a lesser extent Mg could indicate infiltration of a fluid. This too could cause recrystallization
of the monazite rim. In general, rutile is resistant to the infiltration of fluids (Triebold et al., 2007; Zack et
al., 2004b). As argued in recent studies on the nanoscale and microstructures in rutile (Verberne et al.,
2022; Verberne et al., 2023) the presence of twin boundaries and low-angle boundaries could allow for
the transport of trace elements. The same structures potentially facilitate fluid infiltration as well.

However, we observe Li primarily in clusters and only minor concentration in the interface itself.
Therefore, a localized stress-induced recrystallization appears to fit better with these observations.

**Low Th monazite.** The absence or very low concentration of Th in monazite is an odd
observation as ThO$_2$ concentration in monazite are mostly reported to be between 1 – 12 wt. % (e.g.,
Janots et al., 2012, Seydoux-Guillaume et al., 2018). However, a few studies report on monazite with less
than 1 wt. % (e.g., Demartin et al., 1991, and Mannucci et al., 1986). Yet, these studies still show
measureable amounts of Th. Closest to our observations are a study by Janots et al., 2008. X-ray maps of
monazite show a monazite with high Th in the core and a recrystallized rim with low, but not absent, Th
concentrations. Although the analyzed rim is 10 mm wide compared to our 100 nm wide rim, and thus
five orders of magnitude larger. The same underlying principle, recrystallization, would offer an explanation. The recrystallization would have occurred at a low-metamorphic grade Poitrasson et al., 1996, Janots et al. 2008 with Th concentration being positively related to the metamorphic grade (Seydoux 2002). This supports our hypothesis of localized stress-induced recrystallization.

Trace element systematics at the rutile-monazite interface

The distribution of trace elements at the interface in specimen 3 is heterogeneous (Fig. 10). In Figure 10 lower concentrations of trace elements are observed in the upper part of the boundary compared to the lower part of the boundary. The transition occurs across a domain with linear features rich in trace-elements. These linear features are an expression of dislocations at the interface (Dubosq et al., 2019; Fougerouse et al., 2021b; Verberne et al., 2022; Verberne et al., 2023). Heterogeneities at phase boundaries are commonly observed and well-studied within the material sciences (Larere et al., 1988; Sutton et al., 1995; Suzuki et al., 1991; Suzuki et al., 1981). There is extensive evidence that the degree of trace element segregation has an inverse relationship with the interplanar spacing (Sutton et al., 1995). In iron-alloys, it was found that P migrates to boundaries when this spacing decreases (Suzuki et al., 1981). Also, P and S have different degrees of segregation towards boundaries with different orientations in Ni (Larere et al., 1988) and it is apparent that trace elements segregate in different amounts in function to disorientation in olivine (Tacchetto et al., 2021). The segregation into boundaries for ionic systems is more complex since charge compensation needs to be considered (Kliewer, 1965; Sutton et al., 1995). In TiO$_2$ Ikeda and Chiang (1993); Ikeda et al. (1993) demonstrated differences in segregation based on the valence states of trace elements e.g., Al$^{3+}$ and Nb$^{5+}$, boundary type, and the temperature. Here, the curvature of the interface results in a continuous change of opposing crystal faces. Consequently, the spacing between the two phases will vary. This change is reflected in the segregation of trace elements, causing differences in localized trace element concentrations.
Implications for geochronology: a geological history from submicron monazite inclusions

The presence of inclusions with a radiogenic component different from the host mineral can result in a mixed chemical and isotopic system (Fougerouse et al., 2018; Turuani et al., 2022). In this study, Pb is found to have segregated in multiple ways that might have occurred at different moments in time. The clusters still preserved within the monazite core are the result of the interplay between the formation of radiation damage, alpha-healing (Seydoux-Guillaume et al., 2018) and volume diffusion (Seydoux-Guillaume et al., 2019). While recrystallization led to the segregation of Pb from the monazite rim into the rutile-monazite interface. We observe that the recrystallization of the monazite rim is local and not pervasive throughout the rutile grain. Also, we see Pb-enriched nanocrystals trapped within clusters in the recrystallized rim. This localized recrystallization combined with trapping of Pb-enriched nanocrystals would suggest the Pb-enriched features formation and recrystallization occurred in two stages, though without Pb isotopic information from these nanocrystals and the clusters in the core the timing of these stages remains elusive.

Regardless, the mobility results in several Pb reservoirs with, potentially unique, Pb isotopic signatures that record the timing of their formation and the decoupling of the U-Pb system. In ideal situations, this Pb signature can be independently studied (Fougerouse et al., 2018; Peterman et al., 2016; Seydoux-Guillaume et al., 2019; Turuani et al., 2022; Valley et al., 2014; Verberne et al., 2020).

In this study, the only reservoir with a significant amount of Pb, measurable by APT, is the interface between rutile and monazite. Age determination in APT data is based on analyzing the \(^{207}\text{Pb}/^{206}\text{Pb}\) ratios to derive the timing of Pb segregation in contrast to a direct age. The assumption is that this ratio is preserved after segregation. This also means that the presence of U or Th compared to total Pb needs to be minimal. Important to note is that any additional influx of radiogenic Pb derived from the same source material will obscure the determination of the first event. However, such event could possibly also record the latter, preserving both (Taylor et al., 2023). In our results, the absence of \(^{204}\text{Pb}\) and \(^{208}\text{Pb}\) supports the assumption that no common Pb is present and thus all Pb is of radiogenic origin (Peterman et al., 2016; Valley et al., 2014; Verberne et al., 2020). The U/Pb ratios in the boundary are <0.05. This indicates large
amounts of Pb are unsupported and that there is negligible production of radiogenic Pb via continued U
decay after the Pb migration into the rim.

The $^{207}\text{Pb}/^{206}\text{Pb}$ ratio varied between the interfaces in the three APT specimens. The obtained ratios are

$0.0900 \pm 0.0198$ (2σ) for specimen 1, $0.1016 \pm 0.0224$ (2σ) for specimen 2 and $0.1733 \pm 0.0156$ (2σ) in

specimen 3. Specimens 1 and 2 have $^{207}\text{Pb}/^{206}\text{Pb}$ ratios with large uncertainties, however, specimen 3
yields a more precise ratio which can be evaluated. Let us reiterate that the monazite is enclosed within
rutile. Radiogenic is Pb produced by radioactive decay of U and the Pb will be retained in monazite. Later
metamorphic and recrystallization events will mobilize the Pb, isolating it from U. Thus the $^{207}\text{Pb}/^{206}\text{Pb}$
ratio at the time of Pb segregation is preserved. The rocks in the Napier Complex have a complex history.
The oldest potential time of formation is suggested by Black et al., 1984, with monazite crystallization
around 3070 Ma. If we assume an initial crystallization age of 3070 Ma, the current $^{207}\text{Pb}/^{206}\text{Pb}$ of $0.1733$
$\pm 0.0156$ (2σ) indicates Pb production for a duration of 2580 Myr before segregation. This would
correspond with the decoupling of U-Pb occurring around 550 Ma, which would match the late
hydrothermal event in the area (Black, 1983 #18). This shows that analyses of grain boundaries, which
are often avoided during geochronological studies can provide additional information about the geological
history.

CRediT: All authors contributed to the study. Concept: Rick Verberne. Design: Rick Verberne, Steven
Reddy. Sample: Chris Clark. Material preparation: Rick Verberne, Denis Fougerouse, William Rickard,
Anne-Magali Seydoux-Guillaume. Data collection: Rick Verberne, David Saxey, Steve Reddy, Zakaria
Quadir, Anne-Magali Seydoux-Guillaume. Data analysis: Rick Verberne, Steve Reddy, Anne-Magali
Seydoux-Guillaume. The first draft of the manuscript was written by Rick Verberne, with review and
ing editing by Steven Reddy, Denis Fougerouse. All other authors commented on previous versions of the
manuscript. All authors read and approved the final manuscript.

ACKNOWLEDGEMENTS
The work was conducted within the Geoscience Atom Probe Facility at Curtin University. SMR And DWS acknowledge ARC support. The CLyM (Consortium Lyon St-Etienne de Microscopie) is thanked for access to the TEM NeoARM in Saint-Etienne (France). Stéphanie Reynaud (Université de Saint-Etienne, France) is thanked for her help with FIB sample preparation of one TEM foil. A-M Seydoux-Guillaume thanks the Centre National de la Recherche Scientifique, L’institut National des Sciences de l’Univers (CNRS INSU) for financial support. We would also like to thank Dr. Leanne Staddon and an anonymous reviewer for their constructive feedback.

Conflict of Interest Declarations

The authors declare there are no conflicts of interest.

References


Black, L., et al. (1986). "Four zircon ages from one rock: the history of a 3930 Ma-old granulite from

Blum, T. B., Darling, J. R., Kelly, T. F., Larson, D. J., Moser, D. E., Perez-Huerta, A., Prosa, T. J.,
analysis of geological materials: Microstructural Geochronology: Planetary Records Down to

Blum, T. B., Reinhard, D. A., Chen, Y., Prosa, T. J., Larson, D. J., and Valley, J. W., 2018b, Uncertainty
and Sensitivity Analysis for Spatial and Spectral Processing of Pb Isotopes in Zircon by Atom
Probe Tomography: Microstructural Geochronology: Planetary Records Down to Atom Scale, p.
327-350.

Antarctica: implications for the timing of ultra-high temperature metamorphism of the Napier

Clark, C., Taylor, R. J., Johnson, T. E., Harley, S. L., Fitzsimons, I. C., and Oliver, L., 2019, Testing the
fidelity of thermometers at ultrahigh temperatures: Journal of Metamorphic Geology.

Clark, C., Taylor, R. J., Kylander-Clark, A. R., and Hacker, B. R., 2018, Prolonged (> 100 Ma) ultrahigh
temperature metamorphism in the Napier Complex, East Antarctica: A petrochronological
investigation of Earth's hottest crust: Journal of Metamorphic Geology, v. 36, no. 9, p. 1117-1139.


Dachille, F., Simons, P., and Roy, R., 1968, Pressure-temperature studies of anatase, brookite, rutile and

Daneu, N., Recnik, A., and Mader, W., 2014, Atomic structure and formation mechanism of (101) rutile
twins from Diamantina (Brazil): American Mineralogist, v. 99, no. 4, p. 612-624.


James, P., and Black, L., 1981, Archaean Geology.


Sheraton, J. W., Thomson, J. W., and Collerson, K. D., Mafic dyke swarms of Antarctica1987,

Geological Association of Canada.


Langone, A., Saxey, D. W., and Rickard, W. D., 2023, Trace-element heterogeneity in rutile
linked to dislocation structures: Implications for Zr-in-rutile geothermometry: Journal of
Metamorphic Geology, v. 41, no. 1, p. 3-24.

Watson, E. B., Wark, D. A., and Thomas, J. B., 2006, Crystallization thermometers for zircon and rutile:
Contributions to Mineralogy and Petrology, v. 151, no. 4, p. 413-433.


Zaffiro, G., Angel, R. J., and Alvaro, M., 2019, Constraints on the Equations of State of stiff anisotropic
minerals: rutile, and the implications for rutile elastic barometry: Mineralogical Magazine, v. 83,
no. 3, p. 339-347.
Figure captions

**Fig. 1** a) Simplified geological map of the Napier complex after (Carson et al., 2002). b) Photomicrography showing the location of the analyzed rutile grain within the thin section. The rutile is in contact with a coarse garnet that hosts rutile, spinel and sapphirine inclusions and resides in a matrix of mesoperthite, quartz and minor biotite. C and d) Field images of the outcrop at Mt. Hardy. The photos show the large scale foliation present at the sample locality with close up photo of a leucosome rich in coarse-grained garnet and quartz from which the sample was prepared. Sample coordinates 66°49'04.2"S 50°42'60.0"E. Fsp = feldspar, Grt = garnet, Sph = sapphirine, Sp = spinel, Rt = rutile, Msp = mesoperthite. Red arrow indicates studied rutile.

**Fig. 2**: (a): BSE images of analyzed rutile (Rt). Rutile is surrounded garnet (Grt), mesoperthite (Msp), k-feldspar (Kfs) and biotite (Bt). The lift-out locations for TEM and APT specimens are indicated on the image. TEM-1 corresponds to the lift-out taken at Curtin University and this monazite has a recrystallized rim. TEM-2 corresponds with the monazite that does not show the recrystallized rim (b): BSE image showing the presence of the submicron monazites (Mnz) in rutile, as well as corundum (Crn) and Fe-oxide exsolutions. (c): BSE image of biotite directly in contact with rutile, also hosting submicron monazite. (d) Mean-misorientation map showing the subtle differences in orientations that are linked to low-angle boundaries. The black boundaries show the location of twin boundaries (cf. (Verberne et al., 2022)). (e) Mean-misorientation and local misorientation maps for two low-angle boundaries. In both maps a monazite is present, with LAB 1 being in close vicinity and LAB 2 being pinned on the monazite inclusion. The corresponding slip system for the low-angle boundaries is {101}<201>.

**Fig 3.** TEM BF cross-section to illustrate the structure of the monazite inclusion. From outward inward: the rutile host, the interface between rutile and the monazite, the recrystallized rim including 5-20nm clusters and nano-inclusions of Li, Ca and Pb, and the core of the monazite exhibiting a mottled texture.
**Fig. 4:** TEM/STEM characterization of the TEM foils. a) From the left to the right: TEM-Bright Field (BF) image of one entire monazite inclusion in rutile, Selected Area Electron Diffraction (SAED) pattern from the monazite inclusion seen in the [12-1] zone axis and associated with a crystallographic model from CePO₄-monazite in the same orientation, and high-resolution STEM-Annular Dark Field (ADF) image of the monazite-rutile interface. b-Top) Left STEM-ADF and right STEM-ABF images from part of the Mnz inclusion seen in A. b-below) STEM BF and HAADF images from a Mnz inclusion with an alteration rim.

**Fig 5 STEM-ADF images** of clusters in the core and recrystallization rim. a-c) within the core there is a high density of clusters (bright spots) which are trapped within zones of radiation damage (black spots). d-f) The recrystallized rim yields far fewer bright spots, however different inclusions are visible. Note that despite the presence of clusters and inclusions the lattice is almost not affected.

**Fig 6:** Mass spectrum of rutile, monazite and the interface with highlighted trace elements that are enriched in the interface. Li, Ca and U are observed in one location in the mass spectrum. Pb is observed at both Pb⁺ and Pb++. Pb²⁰⁶ is visible at 103 Da and 206 Da, and Pb²⁰⁷ is observed at 103.5 Da and 207 Da, Pb²⁰⁸ is not present above background.

**Fig 7:** TEM EDS spectra and maps

a) TEM EDS analyses conducted on a monazite without recrystallized rim. EDS analyses was performed on the matrix and a cluster to show the difference in composition and the presence of Ca, Pb and U in the cluster. b) TEM EDS maps of the core of the monazite with a recrystallized rim showing the clusters are rich in Ca, Pb, and U. c) TEM EDS maps of clusters present in the recrystallized rim. Ce is shown as an indication for decreased concentrations of major elements. d) TEM EDS interface analysis, showing an EDS map to highlight that Pb is enriched at the interface with a Pb concentration profile (white arrow).

**Fig. 8 (a, b):** Ion maps of specimen 1 and 2 showing monazite (green) is encapsulated within rutile (grey). The interface is enriched in Ca and Pb in both specimens. Also, the curvature over the monazite interface
is clearly visible in the ion maps of specimen 2. (c): Ion maps of specimen 3 not only yields a Pb and Ca enriched interface but also reveal the presence of Ca (d) and Li-rich (e) inclusions. Within the Li-rich inclusion an isoconcentration surface for Pb is shown to demonstrate the presence and equivalence of the inclusions in Fig 4b.

**Fig 9** a) Proximity histograms for the three inclusions observed within the APT data. All inclusions show a slight increase in Pb concentration and a strong increase in one of three cations (Ca, Li, Mg) indicating the formation of a new phase or solid-solution. Ce is shown as a proxy for the REE, which show similar trends at lower concentrations.

**Fig. 10** Ion map showing the heterogeneous distribution of Si + Al within the interface of specimen 3. The 1D concentration profile was taken along the interface as indicated in the top panel. The enrichment of Al, Si and Pb clearly various within each section. The linear features near the middle are interpreted as dislocation present at the interface.
Figure 1
Figure 2
Figure 3

Recrystallized rim  Mottled core  Recrystallized rim

Monazite

TEM-BF

50 nm

interface

interface

interface
Figure 4
Figure 5

Cluster analysis

High density

Low density
Figure 6
Figure 7

EDS spectra for monazite core without recrystallized rim

EDS maps of monazite core with recrystallized rim

Recrystallized Rim

Interface

Pb concentration
Figure 8

(a) Specimen 1

100 nm

(b) Specimen 2

100 nm

Ce  Pb  Li  Ti  Ca

(c) Specimen 3

100 nm  10 nm

fig 9b

d
e

Always consult and cite the final, published document. See http://www.minsocam.org or GeoscienceWorld
Figure 10