Apatite fission-track dating by LA-Q-ICP-MS mapping

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Abstract

Obtaining accurate and precise apatite fission-track (AFT) ages is dependent on producing plentiful high-quality apatite grains from a sample, ideally with high spontaneous fission-track densities (c. >10^5 tracks.cm^-2). Many natural samples, such as bedrock samples from young orogenic belts or low-grade metamorphic samples with low U contents yield low spontaneous fission-track densities. Suchapatites must be counted to avoid biasing the resultant FT age. AFT dating employing LA-Q-ICP-MS spot ablation works very well for grains with high spontaneous fission-track densities which enable potential U-zoning to be detected, while also removing the need for an irradiation step and facilitating simultaneous acquisition of U-Pb and trace element data. The LA-Q-ICP-MS spot ablation thus offers several advantages compared to the External Detector Method (EDM). However, for grains with low spontaneous fission-track densities where U zoning cannot be observed, the LA-Q-ICP-MS spot ablation approach requires the counted area to mimic exactly the site of the laser spot, with the downside that this smaller counted area limits the precision of the resultant AFT age. Here we present an alternative approach to LA-Q-ICP-MS analysis of low fission-tracks density grains by generating a U distribution (^{238}U/^{43}Ca) map of the entire apatite surface by LA-Q-ICP-MS elemental mapping which enables characterization of U zonation. The Monocle plugin for the Iolite LA-ICP-MS data reduction software package is used to display elemental maps and extract mean ^{238}U/^{43}Ca values of the same area counted for the fission tracks. A typical grain-mapping session takes < 5 hours to map 80 grains. The method was employed on the Durango and Fish Canyon Tuff apatite reference materials, and on apatite from six bedrock samples with low fission-track densities (≤ 1.10^5 track.cm^-2). Most apatite samples investigated here were previously dated by the EDM or the LA-Q-ICP-MS ablation spot method. The AFT grain-mapping ages agree with previously published EDM.
or LA-Q-ICP-MS spot ablation ages at the 2σ level. For each apatite sample, we simultaneously acquired U-Pb age and trace element data (Mn, Sr, La, Ce, Sm, Eu, Gd, Lu); here again the data agree with literature constraints (when available) within uncertainties. The mapping approach is therefore a practical solution to low-temperature thermochronology studies employing apatite grains with low spontaneous fission-track densities, while also facilitating investigation of the spatial relationships between thermo- and geochronometric ages and grain chemistry.

Keywords: AFT and U-Pb dating; rare earth elements; Durango; Fish Canyon Tuff; Cyclades; Western Himalaya
1. Introduction

Apatite fission-track (AFT) dating is a well-established thermochronometric method for investigating the thermal history of the crust within a c. 60-120˚C temperature window (Green et al., 1986; Barbarand et al., 2003). The method is widely employed to constrain mountain range development through time, reconstruct the thermal history of sedimentary basins, or determine the provenance of clastic sediments (see Malusà and Fitzgerald, 2019 for a review of AFT applications). However, AFT is a time-consuming dating method since spontaneous fission-tracks, corresponding to the natural fission of \(^{238}\text{U}\) in apatite, must be counted by a human operator. AFT dating also requires determination of the parent U distribution in the dated apatite crystal. This has traditionally been determined using the external detector method (EDM), whereby induced fission-tracks are generated by irradiation of the apatite sample with thermal neutrons in a nuclear reactor. Induced fission of \(^{235}\text{U}\) is registered on a U-free muscovite detector placed in intimate contact with the surface of the apatite crystal; the muscovite detector is then etched and its induced fission track density determined (see Hurford and Green, 1982; Gallagher et al., 1998 and Tagami and O’Sullivan, 2005 for a description of the EDM). The EDM protocol is therefore time consuming as it requires irradiation and counting of both spontaneous and induced fission-tracks. LA-Q-ICP-MS (laser ablation quadrupole inductively-coupled-plasma mass spectrometer) spot analysis has been successfully employed in AFT dating for the \textit{in-situ} measurement of apatite \(^{238}\text{U}\) contents (e.g., Hasebe et al., 2004; 2013; Chew and Donelick, 2012; Cogné et al., 2020). This approach is rapid as it avoids the need for irradiation and counting of induced fission tracks, and can also yield additional information such as apatite Cl contents (Chew et al., 2014a), apatite U-Pb age data (e.g. Chew et al., 2014b) and apatite trace element information which can be exploited to yield host rock-type information (Henrichs et al., 2018; O’Sullivan et al., 2020).

The main advantage of the EDM approach is that identical areas are counted on individual apatite grains and their mirror images in the muscovite detector, and the distribution of induced fission-tracks in the muscovite detector is a reliable proxy map for the U distribution in the apatite grain. This induced fission-track map also records depth-integrated variations in U concentration, as induced fission (like spontaneous fission) generates tracks in the detector up to half a fission-track length.
below the apatite grain surface. The EDM approach can thus be used to assess if U-zonation is present, and as spontaneous and induced fission-tracks are counted on identical areas, within-grain heterogeneity in U concentration can be accommodated by this technique. However, with the LA-Q-ICP-MS spot analysis approach, U zonation can typically only be detected when the spontaneous fission track density is $> \, c. \, 1 \times 10^5 \text{ tracks.cm}^{-2}$. If the dated apatite has a homogeneous fission-track distribution over its entire surface (i.e. no U zonation), the ablation spot can be placed anywhere within the counted area without biasing the accuracy of the resultant AFT age. However, if U zonation is present and the spontaneous fission track density is low (i.e. in low U and/or young samples), it is necessary to match precisely the size and position of the counted area with that of the ablation spot. However, reducing the size of the counted area to that of the ablation spot (typically c. 30 μm in diameter) results in fewer spontaneous fission-track counts and thus less precise single-grain AFT ages (Vermeesch, 2017).

An alternative to laser ablation spot analysis to accommodate potential U zonation in grains with low spontaneous fission track density is to generate a two-dimensional U distribution map over the entire grain surface by LA-Q-ICP-MS mapping. Such an approach would effectively imitate the role of the muscovite detector employed in the EDM, and would mean that the advantages of LA-Q-ICP-MS approach (no need for an irradiation step coupled with simultaneous acquisition of U-Pb and trace element data) could be applied to grains with low spontaneous fission track densities. Here we present such an elemental mapping approach by LA-Q-ICP-MS to AFT dating employing a fast-washout laser cell with an aerosol rapid introduction system (ARIS, van Malderen et al., 2018) which allows for rapid and precise characterisation of elemental distributions on the sample surface (Petrus et al., 2017; Ubide et al., 2015; Chew et al., 2019). Following data reduction with the Iolite software (Paton et al., 2011), mean elemental abundances and mean elemental and isotopic ratios are extracted from user-defined areas on the apatite grain maps (which are similar to the area counted for AFT) using the ‘Monocle’ map interrogation tool for Iolite (Petrus et al., 2017). This grain-mapping approach for AFT dating presented here is easily implemented by any laboratory with an LA-Q-ICP-MS system and we provide details from sample preparation through to the data acquisition protocol, along with recommendations for future research using the approach.
2. Sample information and preparation protocols

2.1 Samples used in this study

We tested the grain-mapping approach for AFT dating on Durango and Fish Canyon Tuff apatites, which are well-characterised in terms of both AFT and U-Pb ages and which we treated as unknowns to assess the reproducibility of the technique (termed “Dur_unk” and “FCT_unk”, respectively). In addition, we also analysed six bedrock samples from Henrichs et al. (2018), Vannay et al. (2004) and Treloar et al. (2000), all of which come from rapidly exhumed terranes and were previously analysed for AFT and/or U-Pb dating. These bedrock samples were selected because of their young AFT ages and their variable U contents (between c. 1 and 300 ppm). Hereafter, AFT ages obtained with the EDM or LA-Q-ICP-MS spot ablation techniques are noted AFT_{EDM} or AFT_{spot}, respectively.

The Durango fluorapatite sample is a crushed fragment of a single large crystal from the iron oxide deposit at Cerro de Mercado, Mexico. The deposit is bracketed by two major ignimbrites from which sanidine-anorthoclase yielded 40Ar/39Ar age of 31.44 ± 0.18 Ma (2σ level; McDowell et al., 2005). Apatite FT_{spot} and U-Pb dating of Durango fluorapatite has yielded ages of 30.6 ± 5.4 Ma (2σ level; Hasebe et al., 2004) and 30.87 ± 0.82 Ma (2σ level; Thompson et al., 2016). Chew et al. (2016) acquired trace element data by solution ICP-MS on aliquots of the same crushed Durango fluorapatite and these data are presented along with our results in Section 4.3.

Fish Canyon Tuff fluorapatite comes from a vast phenocryst-rich dacite with a rhyolitic matrix in the San Juan Volcanic Field of southern Colorado, from which sanidine phenocrysts have yielded an 40Ar/39Ar age of 28.13 ± 0.02 Ma (2σ level; Phillips et al., 2017). Apatite FT_{spot} and U-Pb dating of Fish Canyon Tuff fluorapatite has yielded ages of 29.7 ± 3.8 Ma (2σ level; Hasebe et al., 2004), and 29.1 ± 0.7 Ma (2σ level; Chew et al., 2014b), respectively. Pang et al. (2017) have acquired trace element data (excluding Th, U, Sr) by LA-ICP-MS on Fish Canyon Tuff apatite (see Section 4.3).

RM13 is an upper amphibolite-facies paragneiss sample collected from the central part of the migmatite dome on Paros Island (Greek Cycladic Islands), which yielded a U-Pb age of 11.5 ± 3.8 Ma (95% conf.; model 1 regression; Henrichs et al., 2018). These authors also documented the apatite trace element abundances in sample RM13 (see Section 4.3). There are presently no published AFT ages for sample RM13, but three samples collected from the same unit a few
kilometres to the west yielded AFT$_{EDM}$ central ages ranging from 10.5 ± 2.0 to 12.5 ± 2.8 Ma (2σ level; Brichau et al., 2006).

Apatite samples hb3197 and hb4396 from Vannay et al. (2004) were collected from the Wangtu Gneiss Complex (from the Jutogh Group of the Lesser Himalayan Crystalline Sequence), which yields a U-Pb zircon crystallization age of c. 1.8 Ga (Chambers et al., 2008; Kohn et al., 2010). Although there are no U-Pb age constraints for samples hb3197 and hb4396, monazite from a pelitic schist (Caddick et al., 2007) and uraninite from a leucogranite (Chambers et al., 2008), both from the Jutogh Group, yielded U-Pb ages of c. 10 Ma which dates the latest tectono-thermal event in the Lesser Himalayan Crystalline Sequence (upper amphibolite facies, ca. 640-700˚C; Vannay et al., 2004; Caddick et al., 2007). Samples hb3197 and hb4396 yielded young AFT$_{EDM}$ central ages of 0.7 ± 1.2 Ma, and 1.7 ± 1.0 Ma respectively (2σ level; Vannay et al., 2004). There are no trace element data available for these two apatite samples in the literature.

Apatite samples him610/205, him618/230 and him622/244 from Treloar et al. (2000) were collected along the Astor River in the Shengus Gneiss of the Nanga Parbat massif (Pakistan). Samples him610/205, him618/230 and him622/244 yielded AFT$_{EDM}$ central ages of 1.7 ± 0.2 Ma, 0.4 ± 0.2 Ma, and 0.03 ± 0.04 Ma respectively (2σ level; Treloar et al., 2000). There are presently no U-Pb ages for those three apatite samples, however Treloar et al. (2000) obtained a hornblende Ar/Ar age of 27 ± 1 Ma on sample him610/205 thus documenting a cooling event through 500 °C at 25 ± 5 Ma along the Astor River. Ar-Ar biotite cooling ages as young as 5 Ma are also present in the Nanga Parbat massif (e.g., Zeitler et al., 2001). There are no trace element data available for these three apatite samples in the literature.

2.2 Sample preparation

All samples were pure apatite separates that were processed at the Fission Track Laboratory at the Geology Department, Trinity College Dublin. Each apatite sample was mounted on 15 mm diameter, 2 mm thick epoxy resin discs (Fig. 1a), which was then ground for 20 s and polished for several minutes (using progressively finer diamond-based suspensions) to expose internal apatite surfaces. Apatite mounts were etched in 5.5N HNO$_3$ at 21°C for 20 s to reveal spontaneous fission-tracks (Donelick et al., 1999), and then rinsed thoroughly in deionised water.
Three copper target grids of 3.05 mm diameter (Agar scientific) were affixed to each mount using a water-based glue to enable coordination of the grains (Fig. 1a). Spontaneous tracks were then counted over the entire surface (excluding the outer 10 μm crystal rim; Donelick et al., 2005) of c-axis parallel apatites at x1000 magnification using a Zeiss Axiom Z1m microscope equipped with a camera and the TrackWorks software (Autoscan Systems). The TrackWorks software also permits coordination of the grain centres relative to the target grid positions, delimiting the counted area on each grain and measurement of the etch-pit length (D_{par}). To obtain well constrained ages, we selected from 40 to 70 grains per sample. Fewer grains were counted for the Durango and Fish Canyon Tuff apatite samples (n = 26 and 29, respectively). The grain sizes were typically > 120 μm x 100 μm, and the approach was tested on grains as small as c. 95 μm x 55 μm.

The datafile exported by TrackWorks includes for each grain the sum of the counted spontaneous fission-tracks (N_s), the fission-track density (ρ_s), and the grain X-Y coordinates. The grain coordinates are imported into the laser ablation software (Chromium 2.3, Teledyne CETAC Technologies) to facilitate rapid relocation of the grains for subsequent elemental mapping. Prior to LA-Q-ICP-MS data acquisition, apatite mounts were cleaned in alcohol and deionised water in an ultrasonic bath for 10 minutes to remove any surficial common Pb (Pb_c) contamination to not prejudice U-Pb age measurements.

In this study, we followed the zeta-based approach for AFT dating described by Cogné et al. (2020; see Section 3.4.1). This approach involves (i) FT counting of c-axis parallel, c. 300 μm-long Durango apatite shards on a “zeta” mount (called “Dur_zeta”), (ii) determination of the U/Ca ratios of the Durango shards over one large primary LA-Q-ICP-MS zeta session where grains are ablated three times each, and (iii) calculation of a zeta factor. These Durango shards are then reanalysed during all LA-Q-ICP-MS sessions. While typically up to 20 spot ablations can be placed on each Durango zeta shard, the mapping approach to AFT dating employs much larger areas and hence uses up the pool of FT-counted zeta shards more quickly. It is feasible to reanalyse over a previously ablated surface during subsequent grain mapping sessions (which was undertaken twice during this study), or to repolish the “Dur_zeta” mount between the sessions (which was undertaken four times in total in this study removing a total depth of c. 12 μm). Hence, for the “Dur_zeta” mount the copper target
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grids were replaced by three distinct reference marks each comprising two large ablation patterns for location purposes adjacent to a smaller pattern of the same shape and which was used for the fine-scale referencing (Fig. 1a); these ablation patterns are sufficiently deep (> 50 μm) to survive multiple re-polishing steps.

3. Method

3.1 Data Acquisition

We performed all LA-Q-ICP-MS mapping sessions at the Centre for Microscopy and Analysis at Trinity College Dublin using a Teledyne Photon Machines Analyte Excite 193 nm ArF excimer laser system with a two-volume ablation cell (Müller et al., 2008; van Malderen et al., 2016) coupled to an Agilent 7900 Q-ICP-MS. The aerosol was transported from the laser cell to the mass spectrometer using an aerosol rapid introduction system (ARIS; van Malderen et al., 2018) with short polyetheretherketone (PEEK) tubing and subsequently mixed in a volume-variable smoothing device with Ar carrier gas and N₂ to enhance signal sensitivity and reduce oxide formation.

Laser and Q-ICP-MS operating conditions are summarised in Table 1 and were optimised to satisfy the following criteria: producing a sufficient volume of aerosol during the ablation process (with the main determining factor being optimal acquisition of the full isotope suite required for U-Pb analysis), allocating sufficient mass sweep times per ablation line to produce high-resolution maps that resolve elemental zonation on a typical 100 μm-wide apatite grain, and rapidly acquiring data with limited carry over from zones of varying U concentration or from one ablation line scan (raster) line to another. Operating conditions were optimised daily by LA-Q-ICP-MS tuning on NIST 612 SRM silicate glass to yield (i) maximum sensitivity for ²³⁸U while maintaining a Th/U ratio close to unity and (ii) low production rates of oxides and doubly charged ions which were monitored by analysis of ThO⁺/Th⁺ and Ca²⁺/Ca⁺ respectively (Table 1). Once optimised, the same LA-Q-ICP-MS parameters were applied to both standards and unknowns and across all experiments. The ablation pit depth on the grain surface during a raster line was determined using a Filmetrics white light interferometer and is c. 3 μm-deep (Fig. 1b).

All data were acquired as time-resolved signals derived from a series of horizontal and adjacent parallel raster lines, which populate a user-defined rectangular area defined within the Chromium
2.3 software. For the unknowns, the user-defined area systematically covers the entire grain surface and partly the surrounding epoxy resin (Fig. 1a). Data on primary standards (see the Data Reduction Section 3.2) were acquired from three to four parallel raster lines without overlapping the epoxy resin and for a minimum duration per raster of c. 25 s. We allowed a 10 s background acquisition (washout) between each raster line for both standards and unknowns (Fig. 1a).

A sample-standard bracketing approach is systematically employed to correct for ICP-MS sensitivity drift, which involves intercalating c. 5 to 10 unknowns (corresponding to between c. 45 to 75 raster lines) in between a series of reference materials. These include NIST 612 SRM silicate glass, Durango apatite (fifteen shards of the “Dur_zeta” mount) and the U-Pb standards Madagascar apatite and McClure Mountain apatite (Chew et al., 2014b).

For all mapping sessions (six sessions in total for eight analysed samples), the ICP-MS monitored fourteen masses, which were selected for AFT dating (^{238}U, ^{43}Ca), U-Pb ages (^{238}U, ^{232}Th, ^{206}, ^{207}, ^{208}Pb), and key trace and rare earth element abundances which have petrogenetic significance in apatite (^{55}Mn, ^{88}Sr, ^{139}La, ^{140}Ce, ^{147}Sm, ^{153}Eu, ^{157}Gd, ^{175}Lu; e.g., O’Sullivan et al., 2020). We allocated a dwell time of 10 ms for ^{43}Ca, 2.5 ms for ^{55}Mn, ^{88}Sr, ^{139}La, ^{140}Ce, ^{147}Sm, ^{153}Eu, ^{157}Gd, ^{175}Lu, ^{208}Pb and ^{232}Th, 5 ms for ^{175}Lu, and a longer dwell time of 25 ms for ^{206}Pb, ^{207}Pb, and ^{238}U. Although Cl can exhibit a strong control apatite fission track annealing kinetics (Green et al., 1986; Barbarand et al., 2003) and can be analysed by LA-Q-ICP-MS (Chew et al., 2014a), its elevated background and high first ionisation energy result in low signal/background ratios which makes it challenging to measure with the short dwell times employed in the LA-Q-ICP-MS grain-mapping approach. Chlorine was therefore not monitored in this study.

3.2 Data Reduction

We imported and reduced the data in lolite v2.5 (Paton et al., 2010; 2011). lolite synchronises the ICP-MS data files and laser log files to define individual standard and unknown analyses (termed “integrations” within lolite). The baseline integration (background signal) is defined using the latter portion of the 10 s washout intervals; a smooth spline line is subsequently fitted to these baseline data which are then subtracted from the standard and unknown signals.
Fission-track data (i.e. U/Ca ratios) are reduced using a “Trace_Elements_FTD” data reduction scheme (DRS) and employing semi-quantitative standardisation following Chew and Donelick (2012) and Cogné et al. (2020). NIST 612 SRM silicate glass was used as the primary LA-ICP-MS reference material and previously counted Durango apatite (the “Dur_zeta” mount) as the zeta reference material (see supplementary data table SD1). Uranium concentrations are subsequently normalized relative to \(^{43}\text{Ca}\) to correct for variations in ablation yield. We reduced the trace and rare-earth element data using the Iolite “Trace_Elements” DRS with NIST 612 as the primary reference material and with \(^{43}\text{Ca}\) as an internal elemental standard (the Dur_zeta grains are used as the quality control material for these data, see supplementary data table SD2).

U-Pb data were reduced using the “VisualAge_UcomPbine” DRS (Chew et al., 2014b), which is a modified version of the “Vizual Age” DRS of Petrus and Kamber (2012) that can correct for the presence of Pb\(_c\) in the primary standard. Here, the DRS was run with a \(^{207}\text{Pb}\)-based correction applied to the primary standard and with a linear downhole fractionation correction of zero slope which is appropriate for a shallow rastering approach as time-resolved U/Pb fractionation is absent in line scans (Košler and Sylvester, 2003). Madagascar apatite (U-Pb dated at 473.5 ± 0.7 Ma; Cochrane et al., 2014) is used as the primary matrix-matched standard for reducing the apatite U-Pb data. The quality control materials are Durango apatite and McClure Mountain apatite (523.51 ± 1.47 Ma \(^{207}\text{Pb}/^{235}\text{U}\) ID TIMS age; Schoene and Bowring, 2006). The secondary U-Pb reference material data are reported in the U-Pb supplementary data table SD3.

### 3.3 Extracting data with Monocle

Iolite generates X-Y maps by converting each time-resolved data point of a channel computed by a DRS (e.g., \(^{238}\text{U}/^{43}\text{Ca}\)) into a pixel using Igor Pro’s “Gizmo” Open GL visualisation tool (Paton et al., 2011; Fig. 2). In this study, as all maps were acquired by horizontal rasters with a 18 μm laser beam, the pixels are 18 μm high. The pixels are 3.5 μm wide, which corresponds to the ICP-MS total sweep time (140.5 ms) times the laser scan speed (25 μm.s\(^{-1}\), Table 1). The recently developed Iolite add-on Monocle (Petrus et al., 2017) facilitates display and interrogation of Iolite maps using a flexible “data-extractor” tool (the data are not cropped prior to using Monocle). In this study we extracted data from a user-defined polygon, which closely mimics the area counted for fission tracks. To match the polygon with the counted area, the polygon is defined using the \(^{238}\text{U}/^{43}\text{Ca}\) map and aligned with
the picture of the counted grain (Fig. 2a). The $^{238}\text{U}/^{43}\text{Ca}$ map also allows for identification of any zircon inclusions (which would yield high U but no Ca) which are then excluded from the polygon. While Monocle displays one channel (an elemental abundance or an isotopic ratio) on a map at a time, the “data-extractor” tool retrieves average values for all pixels within the user-defined area for the full suite of channels computed by each different DRS. The Monocle plug-in automatically compiles all extracted average values in an exportable table with their associated internal standard errors.

### 3.4. Age calculation

#### 3.4.1 Apatite fission-track ages

Apatite FT age calculation is performed offline using the information from the Monocle and TrackWorks exported datafiles and employs the Windows Excel spreadsheet provided by Cogné et al. (2020). We refer the reader to the study of Cogné et al. (2020) for further details about the zeta-based calibration and AFT age calculation. In this study, the zeta-factor was obtained using 70 Durango apatite shards analysed during a “primary zeta” session and is $0.70 \pm 0.03$ (zeta-factor of C.A.).

During each of the six grain-mapping analytical sessions undertaken in this study, a pool of 15 of those 70 Durango apatite shards were revisited and analysed. Their respective mean $^{238}\text{U}/^{43}\text{Ca}$ values extracted with Monocle, and the $N_s$ and $\rho_s$ values exported from TrackWorks, are imported into the Cogné et al. (2020) spreadsheet. This spreadsheet calculates (i) a session-specific fractionation factor ($R_i$ ratio) which accounts for systematic variations in the $^{238}\text{U}/^{43}\text{Ca}$ values of Durango apatite shards between the “primary zeta” session and the analytical sessions where FT unknowns are measured and which is related to variations in the ICP-MS tuning conditions, (ii) single and pooled AFT ages of the shards and (iii) single and pooled AFT ages of the unknowns (see supplementary data). As the mapping approach employs a shallow ablation down to c. 3 μm, the $^{238}\text{U}/^{43}\text{Ca}$ ratio is not depth-weighted, which is the only modification in this study made to the approach of Cogné et al. (2020).

We used IsoplotR (Vermeesch, 2008) to display all the single-apatite FT ages of each sample on radial plots (Fig. 3, left panels). The central AFT age of each sample is calculated by IsoplotR and is
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reported in Figure 3 and Table 2; the pooled AFT age is reported when sample passes the $\chi^2$ test ($P(\chi^2) > 0.05$), which suggests one main age population (e.g., Donelick et al., 2005). The uncertainties are given at the 2σ level.

### 3.4.2 Apatite U-Pb ages

U-Pb data extracted with Monocle are imported into the Isoplot 4.15 add-in for Excel (Ludwig, 2012) to propagate uncertainties (following Horstwood et al., 2016 and Drost et al., 2018) and carry out age calculations and plotting. The $^{207}\text{Pb}/^{206}\text{Pb}$ vs $^{238}\text{U}/^{206}\text{Pb}$ data are plotted on a Tera-Wasserburg Concordia plot through which a linear regression is fitted to obtain a lower intercept $^{238}\text{U}/^{206}\text{Pb}$ age. The upper intercept is anchored either with a known initial $^{207}\text{Pb}/^{206}\text{Pb}$ as for the Fish Canyon Tuff apatite ($^{207}\text{Pb}/^{206}\text{Pb}: 0.8444 \pm 0.0006$; Hemming and Rasbury, 2000), or with a $^{207}\text{Pb}/^{206}\text{Pb}$ initial value derived from the Stacey and Kramers (1975) terrestrial Pb evolution model, as for Durango apatite ($^{207}\text{Pb}/^{206}\text{Pb}: 0.84 \pm 0.01$). Anchoring Tera-Wasserburg regressions using the Stacey and Kramers (1975) terrestrial Pb evolution model may not always be appropriate in Cenozoic samples as the amount of radiogenic Pb in-growth in such young samples is typically not significant and the regression will likely be heavily dependent on the choice of the initial $^{207}\text{Pb}/^{206}\text{Pb}$ ratio. We therefore calculated the lower intercept U-Pb ages of the unknowns with an unanchored Model 1 fit to the array. Lower intercept dates are reported with a 95% confidence level (see Table 3 which also includes information on the uncertainty propagation).

### 4. Results

#### 4.1 Apatite fission-track ages

Except for the Durango and Fish Canyon Tuff apatites, most of the bedrock apatite samples analysed in this study have low fission-track densities ($< 1.10^5$ tracks.cm$^{-2}$; Table 2). Detecting U zonation in these apatites was thus impossible when inspecting their spontaneous fission-track distributions. In sample RM13 from Paros in particular, the grain-mapping method reveals complex U/Ca zonation (and thus U-zonation as Ca is assumed stoichiometric) undetected under the microscope, which shows that the FT grain-mapping protocol employed in this study closely emulates the role of the muscovite detector employed in the EDM approach (Fig. 2b).
All apatite samples yield similar pooled and central FT ages at the 2σ level and passed the P(χ²) test as is commonly expected for bedrock (i.e. non-detratal) apatite samples (Table 2). When compared to their literature constraints, the AFT ages obtained from the grain mapping approach are indistinguishable from their accepted ages within 2σ uncertainties; the Durango and Fish Canyon Tuff apatite samples yield pooled FT ages of 29.1 ± 1.6 Ma and 26.6 ± 2.0 Ma, respectively (Fig. 3, left panels). The sole exception arises for samples him610/205 and him622/244, for which an age discrepancy occurs between the central AFT ages obtained in this study (1.1 ± 0.2 Ma and 1.1 ± 0.6 Ma, respectively) and those published in Treloar et al (2000; 1.7 ± 0.2 Ma and 0.03 ± 0.04 Ma, respectively; Table 2). We believe this age discrepancy is due to the larger number of grains analysed for these two samples in this study (n= 40 and 70, respectively) compared to that in the original study (n≤ 20); this point is further addressed in the discussion. Finally, no AFT age was reported for the RM13 sample, but the central AFT age obtained in this study (9.9 ± 1.0 Ma; n=78) agrees with central AFT_{EDM} ages from Paros that range between 12.5 ± 2.8 and 10.5 ± 2.0 Ma (Brichau et al., 2006).

### 4.2 Apatite U-Pb ages

The data extracted from Monocle yield lower intercept $^{238}\text{U}/^{206}\text{Pb}$ ages of 29.2 ± 1.7 / 2.7 Ma (MSWD= 2.1), 30.9 ± 2.7 / 3.5 Ma (MSWD= 2.0) and 10.9 ± 2.0 / 2.1 Ma (MSWD= 1.4) for the Durango, Fish Canyon Tuff and RM13 apatite samples, respectively (Fig. 3, middle panels), which are in good agreement with their literature constraints (Table 3).

The U-Pb age of sample hb3197 is hampered by low $^{238}\text{U}/^{206}\text{Pb}$ ratios with limited spread, and thus yields an uncertainty >100 % (95% confidence, 9 ± 13 / 13 Ma, MSWD= 3.1; Table 3). The U-Pb age of sample hb4396 (1795 ± 42 / 117 Ma; MSWD= 11) corresponds to the age of the Wangtu Gneiss Complex (Chambers et al., 2008; Kohn et al., 2010) implying that regional Miocene metamorphism of the Lesser Himalayan Crystalline Sequence did not reset the Paleoproterozoic U-Pb age of the hb4396 apatite. We note that for the hb4396 sample the MSWD is high, thus highlighting age dispersion.

The lower intercept $^{238}\text{U}/^{206}\text{Pb}$ ages of apatite samples him610/205, him618/230 and him622/244 are 21.3 ± 4.1 / 4.4 Ma (MSWD= 16), 5.2 ± 8.4 / 8.4 Ma (MSWD= 5.8) and 6.7 ± 4.6 / 4.6 Ma (MSWD= 17).
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3.8), respectively (Table 3). The U-Pb age of him610/205 apatites is coherent with the 27 ± 1 Ma hornblende Ar/Ar age obtained on that sample by Treloar et al. (2000). The 95% confidence level uncertainties associated with sample him618/230 and him622/244 are large (>100% and 69%, respectively) due to the limited spread in $^{238}\text{U}/^{206}\text{Pb}$ ratios rendering poor U-Pb age precision, although they are coherent with the late Miocene Ar-Ar cooling ages reported in the Nanga Parbat massif in Zeitler et al. (2001, see Section 2.1). The him-apatite samples also display large MSWDs.

4.3 Trace and rare-earth element contents

Trace and rare-earth element data from Monocle were normalised against chondrite values from McDonough and Sun (1995) and plotted in multi-element spectra diagrams (Fig. 3, right panels). The 26 spectra obtained from Durango apatite show no dispersion and are in excellent agreement with the solution ICP-MS values of Chew et al. (2016) from the same crushed Durango crystal aliquots. The 29 spectra obtained on the Fish Canyon Tuff apatite have limited dispersion and the chondrite-normalised La, Ce, Sm, Eu, Gd and Lu values agree with that obtained by LA-ICP-MS analysis by Pang et al. (2017). The 78 spectra obtained from RM13 Paros apatite agree with the mean spectrum obtained by Henrichs et al. (2018), and we obtain the same range of chondrite-normalised Th and U values with those reported in that study (Table 4). These results show that the grain-mapping approach achieves accurate and precise measurement of trace and rare earth elements.

We note that several of the hb- and him-apatite samples have scattered trace element spectra implying different grain populations (Fig. 3, right panels), although there are no trace element spectra for these samples in the literature to compare our results with. Nonetheless, we explore the trace element chemistry of these samples to extract host rock-type information (e.g. igneous vs metamorphic apatite, which can aid U-Pb data interpretation) using the apatite trace element database compiled from a suite of distinct bedrock lithologies by O’Sullivan et al. (2020). A subset of this database is investigated using principal component analysis (PCA) with the following input variables: Sr, La, Sm, Lu and Eu/Eu* ($\text{Eu/Eu}^* = \text{Eu}_n/(\text{Sm}_n\text{Gd}_n)^{0.5}$; where N = chondrite-normalised). The PCA plot (Fig. 4) shows that most of the hb- and him-samples (except hb4396) plot in the high-grade metamorphic and S-type granite fields, with samples hb3197 and him622/244 trending
towards the low-grade metamorphic field. Sample hb4396 plots in the igneous apatite (I-type and mafic igneous) category (Fig. 4).

5. Discussion

5.1 Apatite fission-track data

The AFT ages obtained with the LA-Q-ICP-MS mapping technique reproduce well with the literature constraints within 2σ uncertainty. More than twenty grains were counted per sample; the AFT pooled age precision is <8 % for the Durango, Fish Canyon Tuff and RM13 Paros samples, and range from 9 to 50% for the youngest (<2 Ma) samples (Table 2).

Samples him610/205 and him622/244 are two samples with an AFT age discrepancy between the central ages obtained and their published age constraints (Table 2). Here we tested whether it is possible to reproduce the EDM central ages of him610/205 and him622/244 samples within 2σ uncertainty using twenty grains (i.e., the number of grains counted in Treloar et al., 2000). We produced an in-house R-script which picks a set of 20 grains at random from our dataset and calculates a central age for this grain subset using IsoplotR. This R-script runs for 1000 iterations and then calculates the percentage of runs whose central ages overlap within 2σ uncertainty with the published age constraints for samples him610/205 and him622/244 listed in Table 2. Using this R-script, 10 % and 50 % of runs overlap within 2σ age uncertainty for him610/205 and him622/244, respectively. Hence, we attribute this age discrepancy to the larger number of counted grains in this study (n= 40 and 70, respectively), which has improved the AFT age precision of both samples.

The samples dated in this study show no AFT age dispersion except for sample him622/244 (Fig. 3). This sample is thus the only sample suitable for discussing the potential relationship between single-apatite FT age and trace element composition. Interestingly, within this sample the five grains with the lowest chondrite-normalised trace element values are those with the lowest $^{238}\text{U}/^{206}\text{Pb}$ ratios, and these five grains also yield the oldest AFT ages (grains coloured in blue on Fig. 3). Although based on just one sample in this dataset, the relationship between single-apatite trace element and FT age should be more routinely checked. This is particularly the case for samples with complex thermal histories as shown by McDannell et al (2019); LA-Q-ICP-MS therefore has great potential in determining the key trace elements for further investigations of FT retentivity.
5.2 Apatite U-Pb and trace element data

The U-Pb dates obtained with the grain-mapping approach on the Durango and Fish Canyon Tuff reference apatites reproduce within 95% confidence uncertainty with their accepted U-Pb ages from LA-Q-ICP-MS spot analyses (with “session-wide” uncertainties < 10 %; Table 3). Therefore, for igneous apatites the accuracy and precision of U-Pb data is not compromised even when mapping with an 18 μm spot size. The Paros (sample RM13) and Himalayan (hb- and him- sample suite) apatites yield greater uncertainties at the 95% confidence level (all >10 % except hb4396; Table 3). These samples are relatively young metamorphic apatites (Fig. 4) characterised by low $^{238}\text{U}/^{206}\text{Pb}$ ratios (high Pb to radiogenic Pb ratios) with a limited spread on Tera-Wasserburg concordia (Fig. 3), which results in poorer precision on the resultant U-Pb age (Henrichs et al., 2018).

Most of the Himalayan apatite sample U-Pb ages are dispersed with large MSWDs (Fig. 3), which based on their scattered trace element spectra could be related to distinct grain populations (e.g. neocrystalline metamorphic apatite and relict higher-grade or magmatic porphyroclasts). Isolating apatite grain populations based on trace-element composition can reduce the U-Pb data dispersion within a sample (Henrichs et al., 2018). This was not undertaken as it was not the primary goal of this study, and only one grain (#31, sample him610/205) was excluded from a Tera-Wasserburg concordia plot due to its highly distinctive trace element spectrum compared to other analyses (grain coloured in red on Fig. 3).

5.3 Advantages and limitations

The elemental mapping method presented herein provides a new approach for laboratories undertaking AFT dating by LA-Q-ICP-MS to deal with low fission-track density apatite grains, while maintaining good age precision. As a result, the entire spectrum of low through to high spontaneous fission-track density grains encountered in natural apatite samples can now be analysed by the same LA-ICPMS instrument without resorting to the EDM with its associated time-consuming irradiation step. Additionally, our approach not only yields accurate and precise AFT dates of low fission-track density apatite grains, but also facilitates simultaneous and accurate U-Pb dating and trace element determinations on the same samples (Fig. 3). While the elemental mapping approach is admittedly slower than single spot ablations, it is still faster than the EDM as it removes the need for sample
irradiation and cooling, mica etching and induced fission-track counting on the mica and dosimeter glasses. Compared to spot ablations, as the whole grain is used to count spontaneous fission-tracks and extract ICPMS data, the siting of the counting area is no longer an issue.

5.4 Recommendations for future work

Below we list a series of recommendations for apatite LA-ICP-MS fission track mapping studies.

1. Rectangular ablation areas (with edges aligned “N-S” and “E-W”) are the simplest to define in most laser ablation software packages, and greatly simplify subsequent data reduction. Apatite grains should therefore be mounted parallel to each other on the grain mounts and their c-axes aligned “N-S” or “E-W” within the sample holder, as this minimizes the amount of epoxy analysed within the rectangular area of ablation.

2. Orientating the line scans parallel to the grain c-axis results in longer rasters and thus best resolves the spatial elemental distribution. Orientating the line scans perpendicular to the c-axis would result in more line scans, which increases the analysis duration as each line scan is followed by a fixed washout interval.

3. Improving the washout in the laser ablation cell (e.g. by using an aerosol rapid introduction system such as in this study) shortens the analysis duration. This is because faster laser stage translation is possible as smearing is reduced, while the washout interval after every line scan can be shortened.

4. Our AFT LA-ICP-MS mapping approach is tailored for samples with low spontaneous fission-track densities. For such samples, similar to McDannell et al. (2019), we advocate analysing significantly more than twenty grains (e.g. n ≥40) per sample to improve age accuracy and precision.

5. In this study, mean values (e.g. U/Ca and U/Pb ratios and other trace elements) were obtained over the entire grain surface. While we did not isolate and pool pixels from the grain maps, it is possible to isolate homogeneous chemical domains on age maps using Monocle (Petrus et al., 2017; Drost et al., 2018) to link apatite U-Pb dates with texturally controlled petrographic elemental information.
6. Conclusions

This work presents an LA-Q-ICP-MS analytical protocol to produce micron-scale elemental ratio maps for fission-track dating of apatite. The protocol is specifically designed for samples with low fission-track density ($\leq 1 \times 10^5$ tracks.cm$^{-2}$), which can be problematic for LA-Q-ICP-MS ablation spot analysis as potential U zoning cannot be detected. Our approach is valid as it produces accurate fission-track dates for Durango and Fish Canyon Tuff and a suite of previously dated samples (six igneous and metamorphic bedrock samples with young AFT ages). The method effectively imitates the role of the muscovite detector employed in the EDM, and therefore provides an alternative approach to fission-track dating of apatite with low U contents and/or young fission-track ages by LA-Q-ICPMS without resorting to the time-consuming EDM. Additionally, the large numbers of grains analysed ($n \geq 40$) for the samples with young ($< 2$ Ma) AFT ages has improved their precision (compared to their literature constraints that used $\leq 20$ grains). Additionally, the method produces U-Pb dates and trace element abundances (Mn, Sr, La, Ce, Sm, Eu, Gd, Lu), which for all samples reproduce with literature constraints (when available) within 95% confidence level.

Finally, the high-resolution imaging protocol integrated with Monocle offers the possibility to isolate pixels of homogeneous chemical domains over large crystals (e.g., Drost et al., 2018). This approach has already been applied to metamorphic apatite petrogenesis study (Henrichs et al., 2019), and has further potential for bioapatite (e.g., bones and teeth) chemical mapping applications, where it would be particularly suited to identifying and isolating zones affected by diagenesis to improve U-Pb dating of fossil bioapatite materials.

Acknowledgements

C.A. thanks Isadora Henrichs and Andy Carter for sharing the Paros and western Himalayan apatite separates respectively, Chris Mark for helpful discussion on apatite U-Pb dating, and Leona O'Connor for the white light interferometer analyses. This research is supported by a research grant from Science Foundation Ireland under Grant Number 13/RC/2092, which is co-funded under the European Regional Development Fund and by PIPCO RSG and its member companies.
References


Chew D.M., Babechuk, M.G., Cogné, N., Mark, C., O'Sullivan, G., Henrichs, I.A., Doepke, D., McKenna, C., 2016. (LA, Q)-ICPMS trace-element analyses of Durango and McClure Mountain


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Figures and Tables

Figure 1. a) Schematic diagram of an apatite mount illustrating the LA-Q-ICP-MS mapping process (features are not to scale). The “ablation pattern” markers serve as location reference points which can survive multiple repolishing events which enables reuse of the Durango “zeta” mounts. They were undertaken with the following laser conditions: 2.5 J.cm\(^{-2}\) fluence, 100 Hz repetition rate and 1000 shot counts. In this protocol the laser stage always scans from left to right and from top to bottom. b) Ablation pit depth measurements from white light interferometry, Y-axis plotted with x 6.5 vertical exaggeration. c) Example of elemental ratio \((^{238}\text{U}/^{43}\text{Ca})\) map from Monocle.

Figure 2. a) Left panel: Optical image of him610/205 grain4 with ablation lines. Middle panel: Monocle image of grain4 showing the \(^{206}\text{Pb}/^{238}\text{U}\) channel (from the Visual_UcomPbine DRS). Right panel: \(^{238}\text{U}/^{43}\text{Ca}\) map (from the Trace_Element_FTD DRS), which is used to define a region of interest to extract AFT, U-Pb and trace element data. b) Example of complex U/Ca zoning in grains from the RM13 Paros sample.

Figure 3. Results from the LA-Q-ICP-MS mapping. From left to right: radial plots of single-apatite FT ages (from isoplotR, Vermeesch 2008), Tera-Wasserburg Concordia diagram (from Isoplot 4.15, Ludwig 2012) and chondrite-normalised multi-element spectra of the analysed samples. The AFT age to be interpreted depending on \(P(\chi^2)\) value is marked by a star.

Figure 4. Principal component analysis of log-normalised Sr, La, Sm, Lu ppm values and Eu/Eu* ratios of the analysed apatite samples plotted against the bedrock apatite database compiled by O’Sullivan et al. (2020). Each 95% confidence ellipse is a lithology group (the data points forming each ellipse have been removed for clarity). ALK: alkali-rich igneous rocks; HM: partial-melts/leucosomes/high-grade metamorphic rocks; IM: mafic I-type granitoids/mafic igneous rocks; LM: low- and medium-grade metamorphic rocks; UM: ultramafic rocks; S: S-type and felsic I-type granitoids.
Table 1. LA-Q-ICP-MS Operating Conditions

<table>
<thead>
<tr>
<th>Laser</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Instrument</td>
<td>Teledyne Photon Machines Analyte Excite ArF 193nm Excimer (HelEx II Active 2-volume Cell)</td>
</tr>
<tr>
<td>Software</td>
<td>Chromium 2.3</td>
</tr>
</tbody>
</table>
| Laser carrier gas | He cell: 0.25 to 0.3 L/min  
He cup: 0.15 to 0.1 L/min  
N₂: 7 to 11 mL/min |
| Washout and background | 10 s (including 3 s laser warm up) |
| Energy density | 2.5 J/cm² |
| Spot size | 18-μm circle (corresponding to the y-axis map resolution) |
| Rastering process | User-defined rectangle automatically populated by raster lines (without overlap). |
| Repetition rate | 53 Hz |
| Scan speed | 25 μm/s |
| Ablation pit depth | c. 3 μm |

<table>
<thead>
<tr>
<th>Inductively coupled plasma mass spectrometer</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Instrument</td>
<td>Agilent 7900 Quadrupole</td>
</tr>
<tr>
<td>Software</td>
<td>MassHunter 4.3</td>
</tr>
<tr>
<td>Plasma ratio frequency power</td>
<td>1550 W</td>
</tr>
<tr>
<td>Sample gas flow</td>
<td>0.60 to 0.70 L/min</td>
</tr>
<tr>
<td>Operating mode</td>
<td>Time-resolved analysis</td>
</tr>
<tr>
<td>Effective mass sweep time</td>
<td>140.5 ms</td>
</tr>
<tr>
<td>Total dwell time</td>
<td>112.5 ms (⁴²Ca: 10 ms; ⁵⁵Mn, ⁸⁸Sr, ¹³⁹La, ¹⁴⁰Ce, ¹⁴⁷Sm, ¹⁵³Eu, ¹⁵⁷Gd, ²⁰⁸Pb, ²³²Th: 2.5 ms; ¹⁷⁵Lu: 5 ms; ²⁰⁶Pb, ²⁰⁷Pb, ²³⁸U: 25 ms) (all sessions)</td>
</tr>
<tr>
<td>Tuning conditions</td>
<td>ThO⁺/Th⁺: 0.2%; ⁴⁴Ca²⁺/⁴⁴Ca⁺: 0.3% (on NIST 612)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Data reduction</th>
<th></th>
</tr>
</thead>
</table>
| Primary standard | For AFT and trace element data: NIST 612 silicate glass  
For U-Pb data: Madagascar apatite |
| Quality control material | Durango apatite (see supplementary data tables) |
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Data Reduction Scheme
- AFT data: “Trace_Elements_FTD”
- Trace element data: “Trace_Elements”
- U-Pb data: “VizualAge_UcomPbine”

Table 2. Apatite fission-track data obtained with the LA-Q-ICP-MS grain mapping approach

<table>
<thead>
<tr>
<th>Sample</th>
<th>Age ± 2σ (Ma)</th>
<th>Published in</th>
<th>n</th>
<th>Ns</th>
<th>Σ Area (cm²)</th>
<th>p</th>
<th>U/Ca</th>
<th>U (ppm)</th>
<th>P(χ²)</th>
<th>Central age ± 2σ (Ma)</th>
<th>Pooled age ± 2σ (Ma)</th>
<th>Dpar (μm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Durango</td>
<td>30.6 ± 5.4</td>
<td>Hasebe et al., 2004*</td>
<td>26</td>
<td>2958</td>
<td>1.2E-02</td>
<td>2.5E+05</td>
<td>5.9E-03</td>
<td>18</td>
<td>1.0</td>
<td>29.3 ± 2.4</td>
<td>29.1 ± 1.6</td>
<td>1.7</td>
</tr>
<tr>
<td>Fish C. Tuff</td>
<td>29.7 ± 3.8</td>
<td>Hasebe et al., 2004*</td>
<td>29</td>
<td>1088</td>
<td>6.5E-03</td>
<td>1.7E+05</td>
<td>4.4E-03</td>
<td>13</td>
<td>1.0</td>
<td>27.1 ± 3.4</td>
<td>26.6 ± 2.0</td>
<td>2.4</td>
</tr>
<tr>
<td>RM13 Paros</td>
<td>10.5 ± 2.0</td>
<td>Brichau et al., 2006+</td>
<td>78</td>
<td>1575</td>
<td>1.6E-02</td>
<td>1.0E+05</td>
<td>7.4E-03</td>
<td>22</td>
<td>1.0</td>
<td>9.9 ± 1.0</td>
<td>9.5 ± 0.6</td>
<td>1.4</td>
</tr>
<tr>
<td>HB3197 (F8)</td>
<td>0.7 ± 1.2</td>
<td>Vannay et al., 2004°</td>
<td>70</td>
<td>22</td>
<td>6.4E-03</td>
<td>3.4E+03</td>
<td>4.0E-03</td>
<td>1</td>
<td>1.0</td>
<td>2.0 ± 1.1</td>
<td>0.6 ± 0.3</td>
<td>2.2</td>
</tr>
<tr>
<td>HB4396 (F10)</td>
<td>1.7 ± 1.0</td>
<td>Vannay et al., 2004°</td>
<td>69</td>
<td>279</td>
<td>7.2E-03</td>
<td>3.9E+04</td>
<td>1.4E-02</td>
<td>4.0</td>
<td>1.0</td>
<td>2.2 ± 0.5</td>
<td>1.9 ± 0.2</td>
<td>1.5</td>
</tr>
<tr>
<td>HIM610/205</td>
<td>1.7 ± 0.2</td>
<td>Treloar et al., 2000°</td>
<td>40</td>
<td>639</td>
<td>4.4E-03</td>
<td>1.4E+05</td>
<td>9.5E-02</td>
<td>287</td>
<td>1.0</td>
<td>1.1 ± 0.2</td>
<td>1.1 ± 0.1</td>
<td>2.5</td>
</tr>
<tr>
<td>HIM618/230</td>
<td>0.4 ± 0.2</td>
<td>Treloar et al., 2000°</td>
<td>41</td>
<td>170</td>
<td>7.0E-03</td>
<td>2.4E+04</td>
<td>5.9E-02</td>
<td>160</td>
<td>1.0</td>
<td>0.3 ± 0.1</td>
<td>0.29 ± 0.05</td>
<td>1.6</td>
</tr>
<tr>
<td>HIM622/244</td>
<td>0.03 ± 0.04</td>
<td>Treloar et al., 2000°</td>
<td>70</td>
<td>60</td>
<td>1.3E-02</td>
<td>4.7E+03</td>
<td>7.0E-03</td>
<td>21</td>
<td>0.6</td>
<td>1.1 ± 0.6</td>
<td>0.5 ± 0.1</td>
<td>1.3</td>
</tr>
</tbody>
</table>

*: LA-ICP-MS spot ablation study with < 20 grains dated
+: The closest study to the RM13 Paros sample (AFTEDM analysis; < 20 grains dated)
°: EDM analysis with ≤ 20 grains dated
n: number of grains analysed in this study
#: shards
Ns: sum of spontaneous fission-tracks
U (ppm): mean U content value (from internal elemental standardisation; Table 4)
P(χ²): Probability to obtain χ² for ν (nb of crystals -1) degree of freedom. If P(χ²) > 0.05 then one population is present and the pooled AFT age can be used.

Central age is calculated with isoplotR. Pooled age is calculated with the spreadsheet of Cogné et al. (2020; see Table SD1).

ζ (zeta-factor) = 0.70 ± 0.03
Table 3. Apatite U-Pb data obtained with the LA-Q-ICP-MS grain-mapping approach

<table>
<thead>
<tr>
<th>Sample</th>
<th>Age ± 2σ (Ma)</th>
<th>MSWD</th>
<th>Published in</th>
<th>n</th>
<th>Age ± 95% conf. (Ma)*</th>
<th>MSWD</th>
<th>initial (^{207})Pb/(^{206})Pb</th>
</tr>
</thead>
<tbody>
<tr>
<td>Durango</td>
<td>30.87 ± 0.82</td>
<td>0.99</td>
<td>Thompson et al., 2016*</td>
<td>26</td>
<td>29.2 ± 1.7 / 2.7</td>
<td>2.1</td>
<td>0.84±0.01 [1]</td>
</tr>
<tr>
<td>Fish C. Tuff</td>
<td>29.1 ± 0.7</td>
<td>1.7</td>
<td>Chew et al., 2014b*</td>
<td>29</td>
<td>30.9 ± 2.7 / 3.5</td>
<td>2.0</td>
<td>0.8444±0.0006 [2]</td>
</tr>
<tr>
<td>RM13 Paros</td>
<td>11.5 ± 3.8°</td>
<td>1.9</td>
<td>Henrichs et al., 2018*</td>
<td>78</td>
<td>10.9 ± 2.0 / 2.1</td>
<td>1.4</td>
<td>0.826±0.005</td>
</tr>
<tr>
<td>HB3197(F8)</td>
<td>NA</td>
<td>NA</td>
<td>70</td>
<td>9.0 ± 13 / 13</td>
<td>3.2</td>
<td>0.691±0.007</td>
<td></td>
</tr>
<tr>
<td>HB4396(F10)</td>
<td>NA</td>
<td>NA</td>
<td>69</td>
<td>1795±42 / 122</td>
<td>11</td>
<td>0.786±0.081</td>
<td></td>
</tr>
<tr>
<td>HIM610/205</td>
<td>NA</td>
<td>NA</td>
<td>40</td>
<td>21.3±4.1 / 4.4</td>
<td>16</td>
<td>0.767±0.063</td>
<td></td>
</tr>
<tr>
<td>HIM618/230</td>
<td>NA</td>
<td>NA</td>
<td>41</td>
<td>5.2±8.4 / 8.4</td>
<td>5.8</td>
<td>0.653±0.032</td>
<td></td>
</tr>
<tr>
<td>HIM622/244</td>
<td>NA</td>
<td>NA</td>
<td>70</td>
<td>6.7±4.6 / 4.6</td>
<td>3.8</td>
<td>0.826±0.009</td>
<td></td>
</tr>
</tbody>
</table>

°: U-Pb age (± 95% conf. level) calculated using a Model 1 regression from Henrich et al. (2018)'s data (see Table SD3).
+*: LA-Q-ICP-MS spot analyses
n: number of grains analysed in this study
#: shards
*: First uncertainty: session-wide estimate (quadratic addition of internal uncertainties and overdispersion of NIST612 data). Second uncertainty: overall propagated systematic uncertainty including a 7% uncertainty derived from the U-Pb results of the Durango zeta apatite shards analysed in all six sessions.
MSWD: mean square of weighted deviates
[1]: anchored using Stacey and Kramers (1975) lead evolution model
[2]: anchored using Hemming and Rasbury (2000)
Table 4. Apatite trace and rare-earth element contents obtained with the LA-Q-ICP-MS grain-mapping approach and used in the PCA.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Th</th>
<th>U</th>
<th>La</th>
<th>Ce</th>
<th>Sr</th>
<th>Sm</th>
<th>Eu</th>
<th>Gd</th>
<th>Lu</th>
</tr>
</thead>
<tbody>
<tr>
<td>Durango</td>
<td>291-491 (363)</td>
<td>15-23 (18)</td>
<td>3508-4704 (4274)</td>
<td>5008-6274 (5664)</td>
<td>491-522 (505)</td>
<td>187-283 (242)</td>
<td>17-21 (19)</td>
<td>164-249 (210)</td>
<td>6-8 (7)</td>
</tr>
<tr>
<td>Fish Canyon Tuff</td>
<td>25-79 (53)</td>
<td>7-19 (13)</td>
<td>1606-2755 (2160)</td>
<td>2876-5152 (3977)</td>
<td>523-612 (560)</td>
<td>132-257 (189)</td>
<td>20-37 (28)</td>
<td>95-187 (138)</td>
<td>4-8 (6)</td>
</tr>
<tr>
<td>RM13 Paros</td>
<td>0-17 (2)</td>
<td>5-81 (22)</td>
<td>83-300 (147)</td>
<td>358-1019 (578)</td>
<td>108-139 (114)</td>
<td>222-449 (291)</td>
<td>14-26 (18)</td>
<td>295-578 (371)</td>
<td>11-32 (17)</td>
</tr>
<tr>
<td>HB3197 (F8)</td>
<td>5-116 (36)</td>
<td>2-27 (12)</td>
<td>46-551 (131)</td>
<td>135-1632 (451)</td>
<td>61-76 (68)</td>
<td>67-739 (215)</td>
<td>3-16 (7)</td>
<td>110-1048 (367)</td>
<td>37-227 (120)</td>
</tr>
<tr>
<td>HB4396 (F10)</td>
<td>21-641 (119)</td>
<td>9-93 (44)</td>
<td>347-2382 (1284)</td>
<td>1058-5931 (3533)</td>
<td>231-459 (330)</td>
<td>132-601 (443)</td>
<td>8-44 (30)</td>
<td>91-405 (299)</td>
<td>3-12 (6)</td>
</tr>
<tr>
<td>HIM610/205</td>
<td>94-541 (358)</td>
<td>66-444 (286)</td>
<td>2-451 (314)</td>
<td>5-897 (602)</td>
<td>297-335 (318)</td>
<td>2-75 (45)</td>
<td>0-9 (7)</td>
<td>4-82 (46)</td>
<td>4-13 (8)</td>
</tr>
<tr>
<td>HIM618/230</td>
<td>2-9 (4)</td>
<td>123-200 (160)</td>
<td>106-185 (134)</td>
<td>467-801 (577)</td>
<td>106-125 (115)</td>
<td>222-320 (259)</td>
<td>5-7 (7)</td>
<td>243-369 (296)</td>
<td>14-39</td>
</tr>
<tr>
<td>HIM622/244</td>
<td>0-30 (2)</td>
<td>1-82 (21)</td>
<td>13-129 (32)</td>
<td>42-478 (117)</td>
<td>81-151 (102)</td>
<td>23-220 (65)</td>
<td>1-5 (3)</td>
<td>40-294 (106)</td>
<td>12-61</td>
</tr>
</tbody>
</table>

Data are given as min-max (mean). Mn concentration and quality control data are provided in Table SD2.
1 raster line of 200 μm-long: 8 s-worth signal

laser displacement (25 μm/s)

position (+10 s)

18 μm spot

copper target grid

user-defined mapping area

for 9 raster lines of 120 μm-wide: 123 s-long acquisition (including washout)

Ablation pit depth observed with white light interferometry

Elemental ratio map from Monocle
a) Elemental ratio map from Monocle (sample Him610/205-Grain4)

b) Example of complex $^{238}\text{U}/^{43}\text{Ca}$ zonation in the RM13 Paros sample
**U/Pb age**

**Central age = 29.3 ± 2.4 Ma**

**Dispersion: 0%, p(χ²) = 1**

**U/Pb age**

**Central age = 9.9 ± 1.0 Ma**

**Dispersion: 0%, p(χ²) = 1**

**Pooled age = 9.5 ± 0.6 Ma**

**Pooled age = 29.1 ± 1.6 Ma**

**Durango (n=26)**

- **AFT age**
  - Pooled age = 26.6 ± 2.0 Ma
  - Central age = 27.1 ± 3.4 Ma
  - Dispersion: 0%, p(χ²) = 1

- **Trace element pattern**
  - Apatite/Chondrite

**Fish Canyon Tuff (n=29)**

- **Central age = 27.1 ± 3.4 Ma**
  - Dispersion: 0%, p(χ²) = 1

- **Pooled age = 26.6 ± 2.0 Ma**
  - Dispersion: 0%, p(χ²) = 1

- **Trace element pattern**
  - Apatite/Chondrite

**RM13 Paros (n=78)**

- **Central age = 9.9 ± 1.0 Ma**
  - Dispersion: 0%, p(χ²) = 1

- **Lower intercept = 9.0 ± 1.3 / 13 Ma**
  - Y-intercept: 0.691 ± 0.007
  - (MSWD = 3.2)

- **Trace element pattern**
  - Apatite/Chondrite

**hb3197 (n=70)**

- **Central age = 2.0 ± 1.1 Ma**
  - Dispersion: 0%, p(χ²) = 1

- **Lower intercept = 2.0 ± 0.6 Ma**
  - Y-intercept: 0.826 ± 0.005
  - (MSWD = 1.4)

- **Trace element pattern**
  - Apatite/Chondrite
**hb4396 (n=69)**

- **pooled age** = 1.9 ± 0.2 Ma
- **central age** = 2.2 ± 0.5 Ma
- Dispersion: 0%, $p(\chi^2) = 1$

**U/Pb age**

- Lower intercept: 1795 ± 42 / 117 Ma
- Y-intercept: 0.786 ± 0.081
- (MSWD= 11)

**Trace element pattern**


g31

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**him618/230 (n=41)**

- **pooled age** = 0.3 ± 0.1 Ma
- **central age** = 0.3 ± 0.1 Ma
- Dispersion: 0%, $p(\chi^2) = 1$

**U/Pb age**

- Lower intercept: 21.3 ± 4.1 / 4.4 Ma
- Y-intercept: 0.767 ± 0.063
- (MSWD= 16)

**Trace element pattern**


g31

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**him622/244 (n=70)**

- **pooled age** = 1.1 ± 0.6 Ma
- **central age** = 1.1 ± 0.1 Ma
- Dispersion: 104%, $p(\chi^2) = 0.6$

**U/Pb age**

- Lower intercept: 6.7 ± 4.6 / 4.6 Ma
- Y-intercept: 0.826 ± 0.009
- (MSWD= 3.8)