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Key Points:

- We combine automatic phase mapping (APM) by SEM/EDS with laser ablation mass spectrometry (LAICPMS)
- The new workflow produces paired mineralogical and isotopic data sets for hand specimens and thin sections
- APM + LAICPMS increases throughput of igneous, metamorphic, and sedimentary studies

Supporting Information:

- Supporting Information S1
- Figure S1
- Data Set S1

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High Throughput Petrochronology and Sedimentary Provenance Analysis by Automated Phase Mapping and LAICPMS

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Abstract The first step in most geochronological studies is to extract dateable minerals from the host rock, which is time consuming, removes textural context, and increases the chance for sample cross contamination. We here present a new method to rapidly perform in situ analyses by coupling a fast scanning electron microscope (SEM) with Energy Dispersive X-ray Spectrometer (EDS) to a Laser Ablation Inductively Coupled Plasma Mass Spectrometer (LAICPMS) instrument. Given a polished hand specimen, a petrographic thin section, or a grain mount, Automated Phase Mapping (APM) by SEM/EDS produces chemical and mineralogical maps from which the X-Y coordinates of the datable minerals are extracted. These coordinates are subsequently passed on to the laser ablation system for isotopic analysis. We apply the APM + LAICPMS method to three igneous, metamorphic, and sedimentary case studies. In the first case study, a polished slab of granite from Guernsey was scanned for zircon, producing a 609 \pm 8 Ma weighted mean age. The second case study investigates a paragneiss from an ultra high pressure terrane in the north Qaidam terrane (Qinghai, China). One hundred seven small (25 μ m) metamorphic zircons were analyzed by LAICPMS to confirm a 419 \pm 4 Ma age of peak metamorphism. The third and final case study uses APM + LAICPMS to generate a large provenance data set and trace the provenance of 25 modern sediments from Angola, documenting longshore drift of Orange River sediments over a distance of 1,500 km. These examples demonstrate that APM + LAICPMS is an efficient and cost effective way to improve the quantity and quality of geochronological data.

Plain Language Summary Much progress has been made in improving the accuracy and precision of geochronological data. However, many Earth Science applications do not so much require better, but simply more data. We have developed a new analytical workflow that is more than twice as efficient than the current state of the art. This new workflow combines a fast scanning electron microscope with a mass spectrometer to simultaneously characterise the chemical, mineralogical and isotopic composition of rocks and minerals. We have applied the new instrument suite to three distinct geological settings to generate high quality data in a fraction of the time that it would have taken conventional studies of the same size and scope.

1. Introduction

Geology, like other fields of Science, greatly benefits from the ever increasing pace of technological progress. Continuous improvements in the miniaturization, automation, and affordability of mass spectrometers have provided geologists with unprecedented access to precise chronological data, and have opened up new research fields such as detrital geochronology. Nowadays, it is common for studies to comprise thousands of U-Pb ages in dozens of samples, something that would have been prohibitively expensive just a decade ago. In a parallel trend, geochronological/isotopic data are increasingly combined with compositional measurements (Engi et al., 2017). Similarly, sedimentary provenance studies increasingly employ a combination of detrital geochronology, bulk geochemistry, and heavy mineral analysis (Vermeesch & Garzanti, 2015). Mineral separation is a bottleneck that prevents these exciting new developments from being more widely adopted.

In stark contrast with the spectacular advances in mass spectrometry and computing, the procedures used to separate datable minerals such as zircon or apatite from the host rock have changed little over the past

half century. These procedures typically involve physical disaggregation of the rock with hammers, jaw crushers, and disk mills; submersion in one or two dense liquids such as bromoform, di-iodomethane, or sodium-metatungstate; and magnetic separation. Not only is mineral separation an unpleasant and time consuming enterprise, it also destroys all textural information within a rock sample, increases the chance for sample cross contamination, and may introduce bias (Sircombe & Stern, 2002). For example, small zircon inclusions in other minerals such as biotite or garnet, which may contain key petrological information, will generally be removed by magnetic and density separation techniques. We here present an integrated suite of software and hardware designed to overcome all these problems. The new workflow combines an SEM/ EDS Automated Phase Mapping (APM) system with an LAICPMS system in order to extract both chemical and isotopic information from rocks while eliminating or reducing the need for manual mineral separation. The APM + LAICPMS workflow comprises of three steps (Figure 1).

- 1. APM is applied to a polished and carbon-coated sample surface to find all occurrences of the mineral of interest in it.
- 2. The center point locations of a random subset of these datable grains are extracted and their coordinates are converted into a format that can be read by the laser ablation system.
- 3. The selected grains are analyzed by LAICPMS to determine their isotopic composition and, hence, age.

Neither the combination of SEM/EDS with in situ geochronology nor the application of in situ U-Pb dating to thin sections is new. Previous studies have developed similar approaches to those described in this paper in order to obtain contextual information that would be destroyed by mineral separation, or to overcome severe sample size limitations (e.g., Compston et al., 1984; Simonetti et al., 2006). The APM + LAICPMS approach achieves those same goals, but is also motivated by other factors, namely speed and convenience. Both the QEMSCAN© SEM/EDS system and the quadrupole LAICPMS instrument used in this study are optimized for speed. They produce chemical maps and isotopic compositions faster than any other common technology available today. When used in tandem, they open the door to a host of new and exciting applications. Although ongoing improvements in mass spectrometry and computing technology have resulted in amazing gains in precision and accuracy of geochronological data (e.g., Kuiper et al., 2008; Schmitz & Kuiper, 2013), it may be argued that for some Earth Sciences applications, the current precision and accuracy of geochronological methods are already fit for purpose. In many studies, what is needed is not so much better data, but just more data. The new analytical workflow removes most manual steps in the data generation process while providing valuable compositional and textural information. This will be illustrated with three simple case studies in igneous (section 3), metamorphic (section 4), and sedimentary (section 5) geology.

2. Methods

2.1. Sample Preparation

The analyzed materials may consist of a standard grain mount, a polished thin section, or a hand specimen (Figures 2a, 3a, and 5a). Further preparation details about these are provided in sections 3–5. The only limitations are the size of the SEM and LAICPMS sample trays. For very homogeneous samples, it is useful for three copper tape markers to be attached to the corners of the samples, to serve as reference points for recoordination. Alternatively, marker points can also be created by laser ablation, or one can simply use any easily recognizable sample feature. Carbon coating is applied prior to SEM/EDS analysis, and rubbed off prior to LAICPMS analysis.

2.2. SEM-EDS Automated Phase Mapping

The APM in this study was performed on a QEMSCAN[©] ("Quantitative Evaluation of Minerals by SCANning electron microscopy") "Wellsite" instrument. This is a high throughput SEM developed by CSIRO and marketed by FEI[®] Inc. (Allen et al., 2012; Khosa et al., 2003; Sutherland & Gottlieb, 1991). The Back-Scattered Electron (BSE) signals and Energy-Dispersive X-ray Spectra (EDS) collected by a QEMSCAN instrument are compared with a database (Species Identification Protocol [SIP]) of known materials to determine the chemical and mineralogical composition of samples at high speed and precision. Typical processing speeds are $0.5-2 \text{ cm}^2 \text{h}^{-1}$ at 10 µm spatial resolution (which can go down to 1 µm), depending on whether crystalline rock or grain mounts are analyzed. This allows 3–15 samples to be processed per day, depending on the

application. Postprocessing of archival data sets produces a wealth of additional information that is difficult to obtain using conventional techniques. The software can assess mineral grain sizes and shapes. For whole-rock samples, textural information and grain orientations can be extracted. Mineral associations, inclusions and their host minerals, and types of lithic clasts are valuable additional information that is difficult and time consuming to obtain by conventional methods.

The present study was performed with a QEMSCAN instrument, but many other APM instruments can be combined with LAICPMS as well. Examples of this include the TIMA (TESCAN), MinSCAN (Zeiss), INCAMineral (Oxford Instruments), RoqScan (CGG), and the free iSpectra software, which can be installed on a wide range of "ordinary" SEMs equipped with an EDS detector (Liebske, 2015). Raman spectroscopy is a different option, based on crystallographic structure rather than chemical composition, that holds great promise for the identification of heavy minerals in sediments (Andò & Garzanti, 2014). While SEM-based methods such as QEMSCAN can easily distinguish different chemical compositions in similar minerals and even amorphous matter, but do not recognize crystal structure, Raman is able to distinguish chemically identical polymorphs. But like SEM/EDS, the accuracy of Raman classification is only as good as that of the underlying mineral database. Different instrumental implementations of QEMSCAN and some of the previously mentioned alternatives offer the option to combine EDS and Raman spectroscopy, potentially combining the strengths of both approaches.

2.3. LAICPMS

Zircon age measurements were performed using an NWR193 laser ablation system coupled to an Agilent 7700X quadrupole ICPMS. Stage coordinates were converted from the QEMSCAN output format to the NWR193 laser stage input format by an in-house computer code called CoCo, which stands for Coordinate Converter and can be downloaded from https://github.com/thegeologician/CoCo. The conversion requires the identification of three marker points, which can either be copper markers or any other recognizable sample feature. Vertical focus was verified by manually going through the selected grains and adjusting the z-position if necessary. Ablation was done using a 25 μm spot size, 11 Hz repetition rate, and an energy fluence of \sim 2.5 J/cm². Mass spectrometer measurements included a 15 s blank measurement followed by 25 s of ablation and 22 s of washout time. The first 2 s worth of ablation signal were discarded to remove any surface-adsorbed common Pb. Integration times were set to 0.05 ms for masses 29 and 91, 50 ms for mass 206, 120 ms for masses 207 and 208, and 20 ms for masses 232 and 238. The entire analytical process is highly automated, allowing us to run the ICPMS overnight and date up to 1,000 zircon grains per analytical session at a rate of \sim 100 dates per hour. In our experience, long analytical sessions often yield better results than short ones, because they ensure that the ICPMS is operated under stable conditions. These long sessions require extraordinary spatial repeatability ($<5 \ \mu m$) of the laser ablation stage. This stability is achieved using the ImageLock image recognition function that has recently been added to the NWR193 software. Age calculation followed a standard-sample bracketing approach in which two Plešovice zircon analyses (TIMS reference age 337.13 \pm 0.37 Ma; Sláma et al., 2008) were interspersed between every 50 measurements (Jackson et al., 2004). Additionally, NIST SRM612 silicate glass (Jochum et al., 2011) was analyzed to monitor sensitivity and quantify the U and Th concentrations of the samples. The stage coordinates of the reference materials were set manually. Instrumental mass bias and depth-dependent elemental fractionation of Pb, Th, and U were corrected using Glitter 4.4 (Griffin et al., 2008). The calculated ²⁰⁶Pb/²³⁸U age was used for grains younger than 1,100 Ma, and the ²⁰⁷Pb/²⁰⁶Pb age for older grains. Grains with a complex growth history or disturbed isotopic ratios, with greater than +5/-15% discordance, were rejected. A common lead correction was applied for the igneous and metamorphic samples, using ²⁰⁷Pb instead of ²⁰⁴Pb due to the unresolvable 204 Hg interference on the latter isotope. The radiogenic 207 Pb/ 206 Pb- and ²⁰⁷Pb/²³⁸U-ratios were obtained by projecting the measured ratios onto the Tera-Wasserburg concordia line along a line going through the common ²⁰⁷Pb/²⁰⁶Pb-ratio. The latter was obtained by isochron regression through multiple aliquots (igneous case study, section 3), or from the Stacey and Kramers (1975) crustal evolution model (metamorphic case study, section 4).

3. Application to Igneous Petrochronology

Igneous rocks are, in many ways, ideally suited for radiometric geochronology. First, they often contain abundant datable minerals. In the case of acidic to intermediate lithologies, this is typically zircon, the

mineral of interest in the present study. Second, the interpretation of igneous ages is generally straightforward, as isotopic closure of igneous minerals tends to coincide with their formation. Third, unlike sedimentary rocks, isotopic data from multiple analyses of igneous minerals can be combined into an isochron or weighted mean, resulting in internal quality checks and higher precision. Sample size requirements for igneous rocks are often an order of magnitude lower than for sedimentary rocks. This makes the overhead of mineral separation all the more unreasonable. It seems wasteful to spend a full day to separate 10,000 zircons from an igneous rock sample, only to use a dozen or so for mass spectrometry. The APM + LAICPMS workflow alleviates this concern and has allowed the London Geochronology Centre (LGC) to pursue studies that would previously have been deemed too costly. One illustrative example of such a study was conceived by an amateur geologist (Mr. D. Smith) from Guernsey who approached the LGC with a request to date some regionally important igneous rocks. With no access to mineral separation facilities and no funds to pay for the analyses, this request would normally have been turned down. However, the efficient workflow tipped the balance in favor of scientific curiosity, and the results of this small study are discussed in this section.

3.1. L'Erée Granite (Guernsey)

Guernsey is one of the Channel Islands, located off the French coast between Normandy and Brittany. The geology of this small island is very diverse. Its southern half is dominated by high-grade metamorphic basement rocks (Icart orthogneiss) of Palaeoproterozoic age, intruded by the Perelle diorite, which was foliated during the Cadomian (late Neoproterozoic) Orogeny. Igneous rocks found further north cover the entire range from gabbro to granite (Topley et al., 1990). The British Geological Survey last visited Guernsey 40 years ago and relatively little geological research has been carried out on the island since then. The geological and tectonic history of Guernsey is commonly assumed to match that of the French Northern Armorican Massif (FNAM; D'Lemos et al., 1990). However, tectonic reconstructions by Linnemann et al. (2014) suggest that the Cadomian suture zone may separate Guernsey from the FNAM, implying completely different pre-Cadomian geologic histories between both terranes. The few geochronological constraints on Guernsey include a single zircon U-Pb age of 2,061 \pm 2 Ma for the lcart orthogneiss (Samson & D'Lemos, 1998), a 611 Ma ± 0.5 Ma zircon U-Pb age for the Perelle diorite (Samson & D'Lemos, 1999), a handful of biotite and hornblende ⁴⁰Ar/³⁹Ar ages (565–605 Ma, Dallmeyer et al., 1992), and a single whole-rock Rb/Sr isochron (469 \pm 13 Ma, D'Lemos, 1987) for gabbros and diorites of the Northern Igneous Complex. For this study, we analyzed a previously undated granite sample from the L'Erée granite (49.45°N, 2.65°W). Sample preparation was done by a stone mason on Guernsey and consisted of cutting and polishing a 50 \times 50 \times 6 mm slab of rock. This small tile was shipped to London and analyzed by APM + LAICPMS.

3.2. Results

SEM/EDS phase mapping of the polished tile confirms the granitic lithology of the L'Erée intrusion. Based on a 3 imes 3 cm area scanned at 10 μ m resolution for a duration of 9 h, the sample contains 32.5% (by area) quartz, 27.2% plagioclase feldspar, 9.0% alkali feldspar, 7.8% micas and chlorite, and 8.7% clay minerals (Figure 2a). Note that these modal abundances are measured over a comparatively small area and may not be representative of the bulk rock due to the coarse grain size of this sample (Figure 2a). Accessory minerals include 0.26% apatite and 0.01% zircon, with grain sizes of up to 4,900 μ m² (corresponding to an equivalent circular diameter of \sim 80 μ m). The X-Y coordinates of 40 of these grains (>400 μ m²) were passed on to the LAICPMS system and dated using the U-Pb method. Thirty three of these zircons yielded U-Pb isotopic data. This was further narrowed down to nine dates by visual inspection of the time-resolved ICPMS signals for evidence for overpolishing, which causes the laser beam to drill through grains during the ablation process. Plotting the concordant data on a Tera-Wasserburg diagram yields a common-Pb corrected weighted mean age of 609.3 ± 8.3 Ma (Figure 2b). This value confirms that the L'Erée granite was emplaced during the late Proterozoic Cadomian orogeny, roughly coeval with the Perelle granodiorite. More information could be gleaned from the sample by acquiring additional data (e.g., CL-imagery), or using multicollector mass spectrometry (e.g., including a ²⁰⁴Pb-based common Pb correction). But the key point made by this case study is that the L'Erée granite would have remained undated if it were not for the low cost (in terms of staff time and consumables) of the APM + LAICPMS workflow.

4. Application to Metamorphic Petrochronology

Metamorphic minerals and textures provide valuable information about geological processes occurring in the crust and upper mantle. By tying together the exerimentally determined stability field of metamorphic index minerals with geochronological information obtained from minerals such as zircon, monazite, or K-bearing micas, it is possible to reconstruct P-T-t (pressure-temperature-time) paths and thereby track vertical motions through the lithosphere. This in turn provides valuable constraints on tectonic rates and orogenesis. It is worth emphasizing that textural and compositional information is crucial in these studies. Using the principle of superposition, the timing of (prograde, peak, and retrograde) regional metamorphism can be bracketed by U-Pb dating of inclusion-bearing zircons and zircon inclusions in metamorphic minerals (e.g., Katayama et al., 2001; Liu & Liou, 2011). In this context, the combination of electron microscopy and laser- or ion-microprobe geochronology has proved invaluable (Schaltegger et al., 1999). The new APM + LAICPMS instrument suite makes this powerful combination of methods both faster and more convenient.

4.1. UHP Paragneiss From Qaidam (China)

The discovery of coesite in metamorphic rocks from the Western Alps and Norway by Chopin (1984) and Smith (1984) marked the beginning of Ultra High Pressure (UHP) metamorphism research. Coesite is only stable at pressures of greater than 30 kbar, indicating that these rocks had been buried to depths of greater than 90 km (assuming lithostatic conditions). This is far greater than geologists previously thought possible. The discovery of metamorphic diamonds in Kazakhstan by Sobolev and Shatsky (1990) pushed the limits of orogenic metamorphism further to 40 kbar, or more than 120 km depth (again assuming lithostatic pressure). This means that continental subduction can bury crustal rocks well into the mantle, and these can resurface rapidly enough to retain their characteristics, thus placing important constraints on tectonic processes. The discovery of metamorphic coesite and diamond sparked a global search for these minerals. The north Qaidam terrane is one area in which UHP metamorphism has been confirmed (Mattinson et al., 2007; Song et al., 2003). UHP minerals are often found in relatively small eclogite and peridotite "rafts" contained within volumetrically dominant ortho- and paragneisses. These gneisses may have provided the buoyancy required to lift the dense ultramafic UHP lithologies out of the mantle (Hacker et al., 1995). Although the gneisses do not always contain UHP minerals themselves due to retrograde metamorphism, they do tend to be much richer in datable minerals (notably zircon) than the eclogites and peridotites. These minerals put key constraints on the timing of peak UHP and retrograde metamorphism (Mattinson et al., 2009).

We have reanalyzed a paragneiss sample (D21G-2) from the north Qaidam terrane that was previously studied by Mattinson et al. (2009). Zircons from this sample were originally analyzed by electron microprobe, cathodo-luminescence imaging, and SIMS U-Pb dating. This revealed two distinct zircon age components. The first component contains 1.8–2.5 Ga detrital ages inherited from the sedimentary protolith of the paragneiss. This component is predominantly found in the cores of the zircons. The second component is found in some rims and is much younger, with dates ranging from 390–450 Ma and a concordia age of 426 ± 4 Ma (Mattinson et al., 2009). This date marks a phase of syn-metamorphic overgrowth during peak metamorphism. Unlike the igneous rock sample in the previous section, the metamorphic rock sample was not introduced into the QEMSCAN as a hand specimen but as a polished thin section (Figure 3a).

4.2. Results

SEM/EDS phase mapping shows that sample D21G-2 contains (in decreasing order of abundance) quartz, feldspars, muscovite, biotite, chlorite, garnet, tourmaline, alumino-silicates, apatite, rutile, and zircon (Table 1). One hundred seven of these zircon grains were sufficiently large ($400-12,000 \ \mu m^2$) for U-Pb dating. Forty of these grains were enclosed by a single mineral in cross section. Garnet only makes up 2.4% of the whole rock, but hosts 20% of the completely enclosed zircon grains. Similarly, 55% of the zircons are hosted by mica, which only makes up 24% of the whole rock. Thus, a disproportionately large number of zircon grains are found in garnet and mica, with quartz and feldspar containing far fewer zircons (Table 1). The U-Pb data reveal two distinct zircon populations. 12% of the grains (13/107 zircons) yield Proterozoic (1–2.5 Ga) U-Pb-ages (Figure 3b). These are inherited grains from the sedimentary protolith to this paragneiss. The remaining 88% of the zircon grains cluster between 390 and 500 Ma. This population represents the syn-metamorphic component. Plotting the common-Pb corrected ²⁰⁶Pb/²³⁸U-dates (using the two-stage crustal evolution model of Stacey & Kramers, 1975) as a Kernel Density Estimate (KDE, Vermeesch, 2012) constrains the timing

Table 1

Side-By-Side Comparison of the Relative Abundances (in Area Percent) of the Minerals Found by SEM/EDS Phase Mapping in Metamorphic Sample D21G-2

Mineral	Whole rock	Zircon
Feldspar	14.1	2.5
Garnet	2.4	20
Nica	24	55
Quartz	55	12
Rutile	0.17	2.5

Note. The middle column shows the composition of the entire scanned area. The right column tabulates the mineralogy surrounding the 40 out of 107 dated zircons that were completely surrounded by one of the minerals in the left column. Garnet and mica grains are most likely to host zircon, whereas quartz and feldspars are least likely to do so. The sample also contained 0.90% tourmaline, 0.37% alumino-silicates, and 0.18% apatite but these did not host any dateable zircon.

of syn-metamorphic zircon growth to 400–440 Ma with an errorweighted mean of 419 ± 4 Ma, in good agreement with the results obtained by Mattinson et al. (2009) (Figure 3c). The mineral affinities produced by the SEM/EDS phase mapping allow further inspection of the U-Pb results. Figure 3d simultaneously plots zircon U-Pb age against grain size and mineralogy. This is a potentially very useful way to constrain crystallization histories, albeit one that does not yield any noticeable patterns in sample D21G-2, as there is no discernable difference in the zircon U-Pb ages for the different host minerals. Using the principle of superposition, this observation supports the notion that zircon growth preceded or accompanied the formation of the other minerals in the sample.

5. Application to Sedimentary Provenance Analysis

Sediments and sedimentary rocks, though globally insignificant from a volumetric point of view, nevertheless are the subject of much geological research, because:

- 1. They cover an estimated 66% of the Earth's surface including nearly all agricultural and urban areas (Blatt & Jones, 1975).
- They represent a valuable archive of geological and environmental change, which is key to reconstructing the geologic history of our planet.
- 3. Sedimentary basins form the source and reservoir of most hydrocarbon resources.

Siliciclastic sediments are, by definition, associated with mass transport of Earth materials. The issue of provenance is therefore central to understanding sedimentary environments. Provenance may be traced by "fingerprinting" sediments through chemical, mineralogical, or isotopic means. Conventionally, each of these provenance proxies are characterized on separate aliquots of the same samples. For example, the chemical composition of the bulk sample may be analyzed by X-ray fluorescence (XRF) on one aliquot, framework petrography on another, heavy mineral analysis on a density separate of a third split, and zircon U-Pb dating on a further density separate of the heavy mineral fraction. The labor intensity of this procedure holds back the widespread application of multimethod provenance studies (Vermeesch & Garzanti, 2015). The new APM + LAICPMS instrument suite aims to solve this problem by potentially generating all four aforementioned provenance data sets as part of the same workflow. When APM is performed to locate zircons in a sediment grain mount, it produces compositional and mineralogical data as byproducts (Figure 1). We will illustrate this procedure with a large-scale provenance study of modern sands from Angola.

5.1. Modern Sand From Angola

The Angolan coast stretches from the Cunene River in the south to the Congo River in the north. Spanning over 11° latitude, it is characterized by hyperarid conditions near the border with Namibia (17°S) to humid conditions near the border with the Democratic Republic of Congo (6°S). The geology consists of Palaeoproterozoic basement rocks in the southern and central parts, and metamorphic terranes of Neoproterozoic age in the north. The topography consists of a coastal plain that widens from 100 km in the south to 200 km in the north, and a steep escarpment to the east of this that rises to a ~1,500 m high plateau. Most Angolan rivers flow toward the Indian Ocean, with only the Cunene (11×10^4 km² catchment area) and the Cuanza (15×10^4 km² catchment area) carrying significant amounts of water to the Atlantic Ocean. Atmospheric circulation along the southwest African margin is dominated by the southeasterly trade winds. These winds drive powerful longshore drift currents that carry sediments from the Orange River in South Africa to the Namib and Skeleton Coast sand seas in Namibia, and into the Moçâmedes Desert of southern Angola (Garzanti et al., 2012, 2014, 2017; Vermeesch et al., 2010). Further offshore, the Benguela Current carries cold Antarctic waters northward until 14°–16° latitude, where the Benguela meets the warm Angola Current and veers offshore (Meeuwis & Lutjeharms, 1990).

The purpose of the third case study was to determine how far north Orange River sediment is carried by longshore currents. To this end, 25 samples were collected along the Angolan coast during a single field

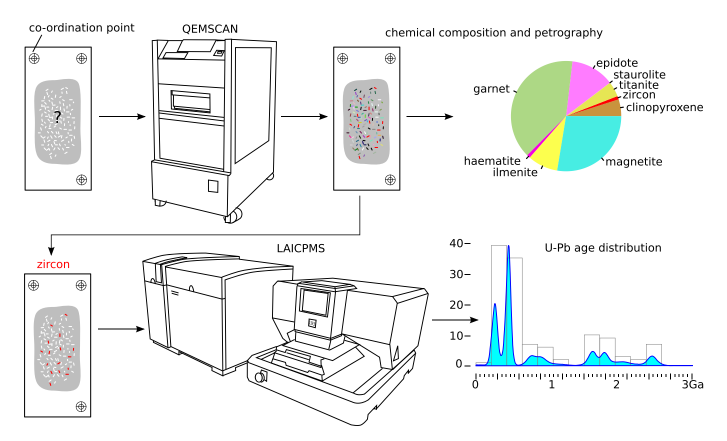


Figure 1. Combining SEM-EDS phase mapping with LAICPMS spot analyses produces paired petrographic and geochronological data sets removing most manual steps required by conventional sample processing routines.

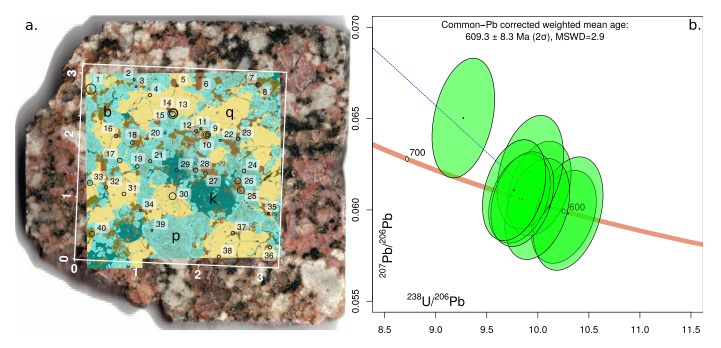


Figure 2. (a) SEM/EDS phase map of a polished hand specimen of l'Erée Granite from Guernsey. A 3×3 cm area of this sample yielded 40 dateable zircon crystals, which are numbered and marked by the black circles whose sizes are proportional to those of the zircons (b, biotite; k, orthoclase; q, quartz; p, plagioclase); (b) Tera-Wasserburg concordia diagram showing the uncorrected U-Pb ratios of the nine most radiogenic zircons, indicating a Cadomian age for this intrusive rock. The dashed blue line marks the mixing line with inherited Pb. Error ellipses are drawn at 95% confidence.

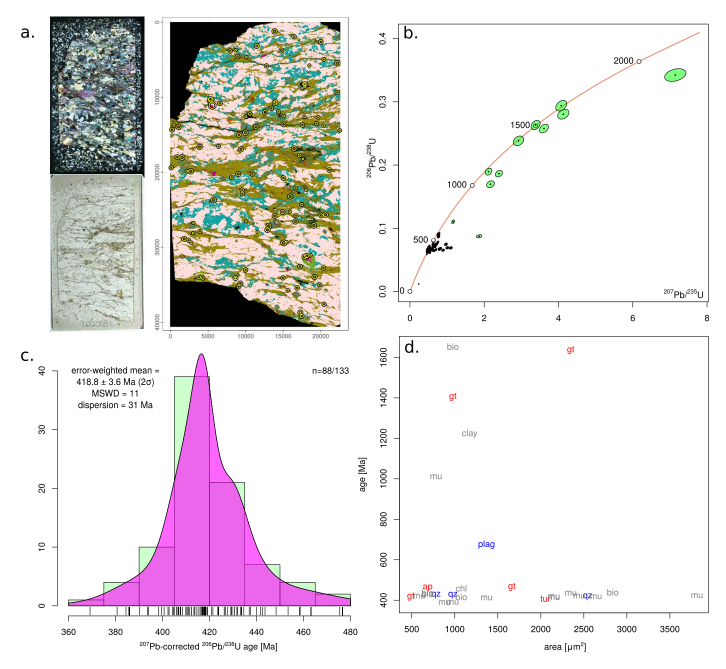


Figure 3. (a) A polished thin section of sample D21G-2 of Mattinson et al. (2009) in XPL (top left) and PPL (bottom left) PPL. A 2×4 cm area of this sample yielded 107 dateable zircon grains shown as yellow circles in the QEMSCAN image (right); (b) (uncorrected) Wetherill concordia diagram of metamorphic sample D21G-2, which contains a mixture of inherited zircons of Proterozoic age and syn-metamorphic zircons of Devonian age; (c) Kernel Density Estimate (KDE; Vermeesch, 2012) of the (common-Pb corrected) syn-metamorphic ages; (d) The zircon U-Pb ages of sample D21G-2 plotted against their apparent area, and labeled with the name of the host mineral (ap, apatite; gt, garnet; qz, quartz; fsp, feldspar).

expedition in June 2015. The sample set comprises of 2 Cunene River samples (labeled 01 and 02 in Figure 4), 10 samples from dunes and beaches in the Moçâmedes Desert (03–12), and 13 beach and river samples along the rest of the Angolan coast, up to the Congo River mouth. From the 32–500 μ m fraction of each sample, heavy minerals were separated by centrifuging in sodium metatungstate (density ~2.90 g/cm³), and recovered by partial freezing with liquid nitrogen. One split of these separates was analyzed by conventional heavy mineral analysis, using methods described by Garzanti et al. (2012). A second split was mounted in epoxy resin on glass slides, polished, carbon coated and analyzed by SEM/EDS. After scanning, the CoCo software recoordinated all the zircon grains in each sample. If more than 130 zircon grains were visible in the

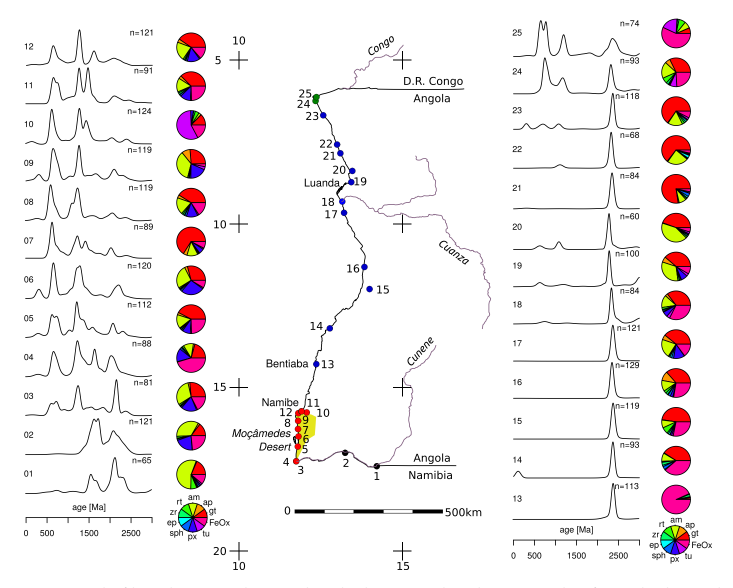


Figure 4. Summary plot of the Angolan provenance data set (samples numbered 1–25), generated using the provenance package of Vermeesch et al. (2016). Colors refer to the Cunene (black), Moçâmedes (red), central (blue) and Congo (green) segments of the Angolan coast. Zircon U-Pb age spectra are shown as Kernel Density Estimates (KDEs; Vermeesch, 2012), heavy mineral data as pie charts with: zr, zircon; rt, rutile; am, amphibole; ap, apatite; gt, garnet; FeOx, iron oxides; tu, tourmaline; px, pyroxene; sph, sphene; ep, epidote.

grain mount, then a random subset of these were selected by the CoCo software (Figure 5a). For samples containing fewer than 130 zircons, all datable grains were analyzed.

5.2. Results

The most abundant heavy minerals are Fe-Ti-Cr oxides, amphibole, epidote, pyroxene, garnet, and zircon. The relative abundances of these minerals vary greatly between samples as a result of the combined effects of provenance and hydraulic sorting (Garzanti et al., 2017; Vermeesch & Garzanti, 2015). Sample 13 in particular is a placer deposit whose heavy mineral composition consists of nearly pure Fe-Ti-Cr oxides (Garzanti et al., 2017). The transparent heavy mineral compositions exhibit similar degrees of variability. For example, sample 01 is dominated by amphibole and contains little pyroxene, whereas sample 10 contains a lot of pyroxene but virtually no amphibole. This variability provides an opportunity to compare and contrast the SEM/EDS results with a conventional heavy mineral analysis. To this end, >200 transparent heavy-mineral grains were either counted under the microscope by the area method or point-counted at suitable regular spacing to obtain real volume percentages (Chayes, 1956a; Galehouse, 1971). Figure 5b compares the

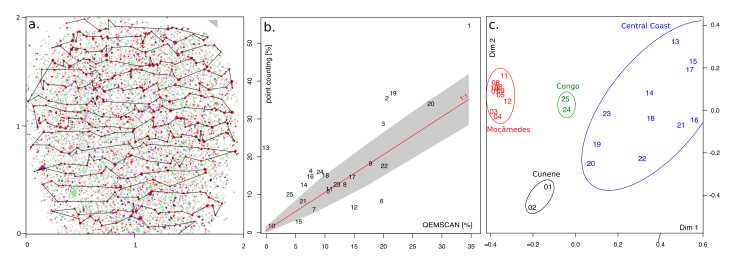


Figure 5. (a) A polished grain mount of a heavy mineral separate from Angola, yielding hundreds of dateable zircons, a random selection of which was made for detrital zircon U-Pb dating (black line). The scale is in centimeter; (b) comparison of the mineral abundances obtained by SEM/EDS phase mapping with conventional heavy mineral point counting for amphibole, which is one of the most abundant transparent heavy minerals in the Angolan data set. The grey-shaded area marks the 95% range around the 1:1 line expected from Binomial counting statistics of 200 conventional heavy mineral counts; (c) multidimensional scaling (MDS) analysis of the detrital zircon U-Pb data, using the Kolmogorov-Smirnov statistic as a dissimilarity measure (Vermeesch, 2013). The age spectra naturally fall into four groups corresponding to distinct geographical provinces, colored as in Figure 4.

resulting modal estimates of the most abundant transparent heavy mineral (amphibole) with those obtained by the SEM/EDS phase mapping. The agreement can only be described as fair (correlation coefficient of 0.76). This likely results from a combination of geometrical and compositional factors that will be discussed in more detail in section 6.

All samples yielded at least 60 concordant U-Pb ages (using a 15% discordance filter), ensuring with 95% confidence that we have sampled all fractions exceeding 8.5% of the total population (Vermeesch, 2004). Wetherill concordia plots of all samples are presented in the supporting information. The detrital zircon U-Pb age spectra fall into four geographical classes: Cunene, Moçâmedes, Central Angola, and Congo. The Cunene River samples (01 and 02) contain two components at 1,300 and 1,700 Ma, respectively. Moçâmedes samples (03–12) exhibit a broader and more complex spectrum with dates ranging from 500 to 2,000 Ma. These spectra look similar to those obtained from modern sand in the Namib and Skeleton Coast deserts, and the Orange River as reported by Vermeesch et al. (2010) and Garzanti et al. (2012, 2014). The Orange signal continues until the town of Namibe at the northern edge of the Moçâmedes desert. The next sample, collected 100 km further north in Bentiaba, has a completely different U-Pb age spectrum that consists of a single, Eburnean age component at ~2,100 Ma. Our data demonstrate that longshore drift of Orange River sediment terminates some 1,500 km north of the Orange River mouth. Thus, the offshore extension of southern Africa's largest river is nearly as long as its onshore course.

The Eburnian age spectrum is found all along the rest of the Angolan coast (samples 03–22) over a distance of more than 1,000 km. It is only a few tens of kilometers south of the Congo River mouth (samples 24 and 25) that the spectrum changes again, to a mixture of four components at 700, 1,000, 2,100, and 2,800 Ma. Joint analysis of all the U-Pb data by Multidimensional Scaling (MDS) confirms the segmentation of the Angolan coast into four zones. MDS is a statistical technique that produces "maps" in which similar samples plot close together, and dissimilar samples plot far apart (Vermeesch, 2013). Using the Kolmogorov-Smirnov statistic as a measure of dissimilarity, the MDS configuration of Figure 5c groups the samples into four clusters that have a one-to-one correspondence with the sampling locations (Cunene, Moçâmedes, Central Coast, and Congo). The comparatively large spread of the Central Coast samples (shown in blue) reflects the high sensitivity of the Kolmogorov-Smirnov statistic to unimodal age distributions. For such distributions, even the slightest shift in the median age will have a large effect on the KS statistic. In contrast, the KS-statistic is relatively insensitive to differences in the tails of complex age distributions and does therefore not easily differentiate between the different samples within the Moçâmedes cluster.

6. Discussion

Extracting or locating dateable minerals is the biggest bottleneck in many geochronological studies. By reducing or entirely removing this bottleneck, the APM + LAICPMS workflow has more than tripled the throughput of the U-Pb operations at University College London, allowing the lab to process up to five samples and 1000 zircon dates per day. Detrital samples generally do not contain sufficient zircon in thin section, and generally still do require mineral separation. However, this is only an abbreviated version of the mineral separation normally employed for zircon. So even though this study uses mineral separates, the process is faster and less toxic than a full separation to concentrate zircon to good purity as would conventionally be done. Furthermore, because zircon does not form a solid solution series, EDS phase mapping is nearly 100% efficient at finding zircon in grain mounts. Hence, small and/or crude mineral separates do not pose a problem. The high throughput approach to mineral separation described in section 5 enables significant time savings compared to conventional mineral separation. The increased throughput allows more samples to be analyzed for any given provenance study. The resulting "big" data sets offer much greater power to resolve the sometimes subtle differences that can exist between age distributions (e.g., Rittner et al., 2016).

In igneous petrology, the APM + LAICPMS workflow avoids even more manual work. In this case, sample preparation is limited to simply polishing a hand specimen, and LAICPMS analysis requires only a dozen or so datable crystals. The resulting gain in throughput would be very advantageous in the case of reconnaissance studies on the one hand, and studies where large numbers of different samples need to be analyzed on the other. It is now widely recognized that many plutons have a complex architecture (Krauskopf, 1968) and are assembled over measurable time intervals (Matzel et al., 2006). The new approach opens the door to high (map-scale) resolution "age mapping" of igneous bodies. For example, working on some Cascades batholiths, Matzel et al. (2006) found 3–5 Myr assembly time scales, interspersed with shorter magmatic pulses, based on detailed TIMS work on 8–12 samples per pluton. Assuming sufficient precision of the LAICPMS measurements, the new method would make practical analysis of many dozens of samples distributed across the batholith, and the SEM/EDS phase mapping would also generate compositional/mineral assemblage information as well.

In some cases, the APM + LAICPMS approach does not just produce more data, but also improves their fitness for purpose. The use of software such as CoCo to automatically pick a random selection of (detrital) zircon grains largely removes the selection bias that human operators inevitably carry. The Angolan samples yielded many dark anhedral grains that would never have been analyzed by a human operator, but were marked as zircons on the SEM/EDS phase maps. These grains always did turn out to be zircons, many of which yielded concordant U-Pb dates. In igneous studies, significant gains in precision can be made by averaging multiple grains. This may be used to offset any loss of age precision resulting from small laser spot sizes, or from analyzing a full suite of trace elements during ICPMS analysis. Thus, the increased throughput of the new workflow may significantly increase the scope of what can be analyzed. Data quality would be further improved by integrating CL imaging in the analytical workflow. Unfortunately this was not possible using the "Wellsite" version of the QEMSCAN instrument used for this study. Other automated SEM-EDS phase imaging systems do allow CL imaging of samples, which yields additional textural information for zircon U-Pb analysis. This is essential for complex igneous and metamorphic samples that contain multiple phases of zircon growth. However, in some sedimentary provenance studies, the actual ages do not matter so much as the shape of the age spectra, which can be used as a "fingerprint" to trace the flow of sand through a sediment routing system (e.g., Garzanti et al., 2012, 2014; Rittner et al., 2016; Vermeesch, 2013; Vermeesch & Garzanti, 2015; Vermeesch et al., 2010). In such cases, it is arguably better to place the laser spot "blindly" so as to avoid selection bias between cores and rims. CL imaging is time consuming but could be integrated in an automated workflow, by targeting a subset of the imaged mineral phases after APM and before LAICPMS analysis. Metamorphic and (particularly) igneous studies generally require far fewer zircon U-Pb dates than sedimentary provenance studies do, so the additional time for CL imaging would be modest and would be counterbalanced by the reduction in the number of mixed analyses that need to be rejected during data processing.

Although this paper has focused on zircon, SEM-EDS phase mapping can identify other accessory minerals amenable to U-Pb geochronology, such as apatite, titanite, rutile, monazite, etc. This opens up the possibility of acquiring detailed P-T-t paths in metamorphic petrology or multiproxy detrital data sets without mineral picking (e.g., O'Sullivan et al., 2016). Applying U-Pb geochronology in thin section and polished hand specimen can also be used to constrain the Pb initial ratio in common Pb-bearing phases such as apatite, rutile, or titanite using the Pb-composition of adjacent feldspars (Chew et al., 2011). Finally, the ability to map accessory mineral occurrence inside rock-forming minerals could be used to constrain the crystallization and cooling history of individual clasts in a conglomerate generating rich provenance data sets.

Despite all these strengths, it is important to also be aware of the limitations of the APM + LAICPMS workflow. First of all, the accuracy of the compositional maps produced by the SEM/EDS instrument is only as good as the compositional database behind the software. Out of the box, SEM/EDS may struggle to correctly identify solid solution series minerals such as amphibole, pyroxene or garnet, and chemically variable minerals such as tourmaline (Nie et al., 2013). This may partly be the reason for the relatively poor agreement between the SEM/EDS and conventional heavy mineral abundances shown in Figure 5b. The other likely reason for the observed scatter is the fact that conventional heavy mineral analysis is based on observations made by looking through a grain mount in transmitted light. Thus, the resulting mineral abundances are effectively done based on the 2D projection of the entire grain on to the focal plane of the petrographic microscope (Chayes, 1956b). In contrast, SEM/EDS uses carbon-coated polished sections which are, essentially, 2-D cross sections through a 3-D lattice of grains (in case of hand specimens or thin sections), or through loose sediment in an epoxy puck or grain mount. In the latter case, the mineralogical composition may differ depending on grain size and the depth of the polish. This is due to the hydraulic equivalence between fine grained dense minerals and coarser grained light minerals (Garzanti et al., 2008), which makes the latter more (or, for epoxy pucks, less) likely to be overpolished than the former. Neither grain size nor polishing depth was controlled in the Angolan case study. Further work needs to be done to understand these issues and demonstrate the reliability of heavy mineral counts produced by SEM/EDS.

7. Conclusions

We here presented a novel workflow for high throughput petrochronology, combining automated phase mapping by high-speed SEM/EDS with isotopic measurements by quadrupole LAICPMS. We successfully applied the new workflow to igneous, metamorphic and sedimentary samples. The igneous study took 12 h to complete, including 30 min to cut and polish the hand specimen, 9 h for the (overnight) APM analysis, 30 min for recoordination, and 30 min of LAICPMS time. The main limitation of the igneous study was the low yield of highquality U-Pb ages, which was caused by the tendency of the laser beam to drill through, or partially fall outside small or overpolished zircon crystals. The metamorphic case study was performed on a polished thin section. Since thin sections are needed for petrographic purposes anyway, the only additional sample preparation step was the application and removal of the carbon coating before and after APM analysis, respectively. One potentially very useful application of the new analytical approach is investigating the association of age components within the mineralogical context. The most significant limitations of our metamorphic study were the absence CL-images to guide the placement of the laser spot, and the use of a single collector guadrupole LAICPMS, which precluded the use of a ²⁰⁴Pb-based common Pb correction. However, adding those two features would slow down the workflow and partially defeat the purpose of the high throughput instrument suite. For these reasons, APM + LAICPMS is best thought of as a complementary approach to igneous and metamorphic petrochronology that is particularly useful for reconnaissance studies. In contrast, the step change in sample throughput achieved by APM + LAICPMS is a game changer for sedimentary provenance studies. Allowing 30 min for the crude mineral separation routine described in section 5.1, 30 min for preparing a polished grain mount, 2 h for the APM analysis, 30 min for the coordinate conversion, and 1 h for LAICPMS measurements, a streamlined workflow produces paired mineralogical and geochronological data sets at a rate of one sample per 5 h. Thus, APM + LAICPMS has the potential to bring the fields of petrochronology and sedimentary provenance analysis into the Era of "big data" science (Vermeesch & Garzanti, 2015).

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