

U-Pb Zircon LA-ICP-MS Techniques (University of British Columbia)

Zircons were separated from their host rocks using conventional mineral separation methods and sectioned in an epoxy grain mount along with grains of internationally accepted standard zircon (FC-1, a ~1100-Ma zircon standard), and brought to a very high polish. The grains were examined using a stage-mounted cathodoluminescence imaging setup that makes it possible to detect the presence of altered zones or inherited cores within the zircon. The highest quality portions of each grain, free of alteration, inclusion, or cores, were selected for analysis. The surface of the mount was then washed for ~10 minutes with dilute nitric acid and rinsed in ultraclean water. Analyses were carried out using a New Wave 213nm Nd-YAG laser coupled to a Thermo Finnigan Element2 high resolution ICP-MS (inductively coupled plasma mass spectrometer). Ablation took place within a New Wave "Supercell" ablation chamber, which was designed to achieve very high efficiency entrainment of aerosols into the carrier gas. Helium was used as the carrier gas for all experiments, and gas flow rates, together with other parameters such as torch position, were optimized before beginning a series of analyses. We typically used a 25-micron spot with 60 percent laser power and did line scans rather than spot analyses to avoid within-run elemental fractions. Each analysis consisted of a 7-second background measurement (laser off) followed by a ~28-second data acquisition period with the laser firing. A typical analytical session consisted of four analyses of the standard zircon, followed by four analyses of unknown zircons, two standards, four unknowns, and so forth, and finally four standard analyses. Data were reduced using the GLITTER software package developed by the GEMOC group at Macquarrie University, which subtracts background measurements, propagates analytical errors, and calculates isotopic ratios and ages. This application generated a time-resolved record of each laser shot. For detrital zircon samples, 60-100 grains were analysed.

U-Pb zircon LA-ICP-MS Techniques (Apatite to Zircon, Inc.)

Mineral separates were obtained at the laboratories of Apatite to Zircon, Inc., in Viola, Idaho. Lithium polytungstate and a centrifuge were used in place of the conventional Wilfley table, thus guarding against loss of zircon grains that might inadvertently be washed away, undetected, in the conventional method. Zircons (both standards and unknowns) were then mounted in 1-cm² epoxy wafers and ground down to expose internal grain surfaces before final polishing. Grains, and the locations for laser spots on these grains, were selected for analysis from all sizes and morphologies present using transmitted light with an optical microscope at a magnification of x 2000. This approach is used instead of cathodoluminescence 2-D imaging because it allows the recognition and characterization of features below the surface of individual grains, including the presence of inclusions and the orientation of cracks, which could otherwise result in spurious isotopic counts.

Isotopic analyses were performed with a New Wave UP-213 laser ablation system in conjunction with a ThermoFinnigan Element2 single collector double-focusing magnetic sector inductively coupled plasma-mass spectrometer (LAICP-MS) in the GeoAnalytical Lab at Washington State University. In comparison to a quadrupole ICP-MS, the Element2 has flat-top peaks and higher sensitivity, resulting in larger Pb signals, better counting statistics, and more precise and accurate measurement of isotopic ratios. For all analyses (both standard and unknown), the diameter of the laser beam was set at 20 µm and the laser frequency was set at 5 Hz, yielding ablation pits ~10-15 µm deep. He and Ar gas were used to deliver the ablated material into the plasma source of the mass spectrometer. Each analysis of 250 cycles took approximately 30 seconds to complete and consisted of a 6-second integration on peaks with the laser

turned off (for background measurements) followed by a 25-second integration with the laser firing. A delay of as much as 30 seconds occurred between analyses in order to purge the previous analysis and prepare for the next. The isotopes measured included ^{202}Hg , $^{204}(\text{Hg} + \text{Pb})$, ^{206}Pb , ^{207}Pb , ^{208}Pb , ^{232}Th , ^{235}U , and ^{238}U . The Element2 detector was set at analog mode for ^{232}Th and ^{238}U and at pulse counting mode for all other isotopes. Common Pb correction was made by using the measured ^{204}Pb content and assuming an initial Pb composition from Stacey and Kramers (1975). Interelement fractionation of Pb/U is generally <20 percent, whereas fractionation of Pb isotopes is generally <5 percent. At the beginning of each LA-ICP-MS session, zircon reference materials (Peixe and FC1) were analyzed until fractionation was stable and the variance in the measured $^{206}\text{Pb}/^{238}\text{U}$ and $^{207}\text{Pb}/^{206}\text{Pb}$ ratios was at or near 1 percent. In order to correct for interelement fractionation during the session, these standards were generally reanalyzed after each 15-25 unknowns. Fractionation also increases with depth into the laser pit. The accepted isotopic ratios were accordingly determined by least-squares projection through the measured values back to the initial determination.

U-Pb Zircon LA-ICP-MS, Apatite to Zircon, Inc. and Geosep Services

Mineral separates were obtained at the laboratories of Geosep Services in Moscow, Idaho. Lithium polytungstate and a centrifuge were used in place of the conventional Wilfley table, thus guarding against loss of zircon grains that might inadvertently be washed away, undetected, in the conventional method. Zircons (both standards and unknowns) were then mounted in 1-cm² epoxy wafers and ground down to expose internal grain surfaces before final polishing. Grains, and the locations for laser spots on these grains, were selected for analysis from all sizes and morphologies present using transmitted light with an optical microscope at a magnification of x 2000. This approach is used instead of cathodoluminescence 2-D imaging because it allows the recognition and characterization of features below the surface of individual grains, including the presence of inclusions and the orientation of cracks, which could otherwise result in spurious isotopic counts.

Data were collected for the following isotopic masses: ^{202}Hg , $^{204}\text{Hg}+^{204}\text{Pb}$, ^{206}Pb , ^{207}Pb , ^{208}Pb , ^{232}Th , ^{235}U , and ^{238}U (250 data scans over 30 s) followed by ^{28}Si and ^{91}Zr (5 data scans over 4 s). The instruments used were a New Wave YAG 213 nm laser ablation (LA) system in line with a Finnigan Element2 magnetic sector, inductively coupled plasma, mass spectrometer (ICP-MS) at the Washington State University Geoanalytical Laboratory in Pullman, Washington, U.S.A. (e.g., Chang et al., 2006). All analyses were performed using a 20 μm spot. Following approximately 6 s of background data collection, laser ablation commenced and data were collected for the ablated material. Ablated material was transported to the plasma line using He; Ar was the plasma gas.

Zircon reference material for which independently accepted ages are published were designated as primary, secondary, and tertiary for purposes of U-Pb age calibration depending on the anticipated age of the unknowns (see table below). Two primary and two secondary standard spots were analyzed prior to and following each group of ~25-30 tertiary standards and/or unknown sample spots. Five spots of each tertiary standard were analyzed near the beginning and again near the end of the session.

Standard	Standard	U-Pb age ($\pm 2\sigma$)	Reference
FC	Duluth complex	$1099.0 \pm 0.6 \text{ Ma}$	Paces and Miller, 1993

F5	Duluth complex	1099.0 ± 0.6 Ma (assumed equal to FC-1)	Paces and Miller, 1993
IF	Fish Canyon Tuff	28.201 ± 0.012 Ma	Lanphere et al., 2001; Kuiper et al., 2008
MD	Mount Dromedary	99.12 ± 0.14 Ma	Renne et al., 1998
PX	Peixe	563.5 ± 1.6 Ma	Gehrels et al., 2008
R3	Braintree complex	418.9 ± 0.4 Ma	Black et al., 2004
T2	Temora 2, Middledale gabbroic diorite	416.78 ± 0.33 Ma	Black et al., 2004
TR	Tardree Rhyolite	61.23 ± 0.11 Ma	Dave Chew, personal communication

Data modeling – Previous LA-ICP-MS studies of U-Pb zircon dating deployed the so-called intercept method which assumes that isotopic ratio varies linearly with scan number due solely to linearly varying isotopic fractionation (Chang et al., 2006; Gehrels et al., 2008). For the intercept method, a line is fitted to background-corrected isotopic ratio (e.g., $^{206}\text{Pb}/^{238}\text{U}$) versus data scan number and the intercept of the fitted line (corresponding to data scan number = 0) is used as the isotopic ratio for age calculation and the error on the intercept is used for age error calculation. For this study, individual isotopes were modeled by fitting a sum of 10 Gaussian equations (“Gaussians”) to the raw signal data (not background corrected) using chi-squared minimization. Two fitting passes were performed: after the first pass, all raw signal values greater than two standard deviations away from the sum of fitted Gaussians were designated outliers; the second pass fit the sum of Gaussians to the data excluding the outliers. The advantage of the present approach is that it avoids the assumption of linearly varying isotopic ratio with scan number, an assumption easily violated for zircons that may contain useful information (e.g., a zircon for which the ablation pit variably penetrates two zones having different U-Pb ages).

Measured background values for each isotope at each LA-ICP-MS spot were calculated as follows: a) the final background scan was assigned as the scan closest to the global minima ^{232}Th and ^{238}U values; if no such global minima were found, the analysis was deemed a failure, b) a line was fitted to the background values, outliers identified, and a line again fitted to the data excluding the outliers, c) for a fitted line exhibiting a negative slope (indicative of a decaying background), the value of the line at the last background scan was assigned as the background value; for a fitted line exhibiting a zero or positive slope, the mean value of the data excluding the outliers was assigned as the background value, and d) the error of the background value was set equal to the standard deviation of the all background values (excluding outliers) about their fitted line (negative slope) or mean (zero or positive slope).

Session-wide fitted background values for each isotope were determined using all zircon standards and applied to all spots in the session. These steps were taken for each isotope: a) measured background value versus spot number in the session was fitted to a 3rd-order polynomial, outliers identified, and the

fitting repeated excluding the outliers, and b) fitted background at each session spot was calculated using the 3rd-order polynomial. Session-wide fitted background error was set equal to the standard deviation of the measured background values (excluding outliers) about their respective fitted 3rd-order polynomial. For any spot (standard or unknown) where the measured background value exceeded the session-wide fitted value by more than 2σ , the background error was set equal to 1σ plus one half of the amount by which the measured background value exceeded the session-wide fitted value by 2σ .

The sum of fitted Gaussians was used here primarily to identify outlier data and characterize signal noise. After the second fitting pass, the standard deviation of the non-outlier data about their respective sum of fitted Gaussians was taken as the absolute signal error for each data scan. When N data scans contribute to a single isotopic signal value used for age calculation (only concordant scans when the number of concordant data scans is greater than zero; all data scans for common Pb-correction based on isotopic sums), the error of the single isotopic signal value was set equal to the product of a) $N^{1/2}$ and b) the absolute signal error for each data scan.

Pb/U Fractionation Factor—Fractionation factors were determined for each data scan of each primary standard spot. For a particular isotopic ratio (e.g., $^{206}\text{Pb}/^{238}\text{U}$), the fractionation factor as used here equals the accepted isotopic ratio divided by the measured ratio. A two dimensional grid (spot number, scan number) of fractionation factors for each isotopic ratio was constructed for the session as a whole by fitting a series of 4th-order polynomials (excluding outliers). Under the operating conditions of the LA-ICP-MS sessions in this study, fractionation factors were found to vary strongly with scan number, decreasing with increasing scan number (presumably due to increasing ablation pit depth and the effect this had on fractionation; e.g., Paton et al., 2010). Fractionation factors were calculated using isotopic values based on the sum of fitted Gaussians. Ages, including when the standards were treated as unknowns, were calculated using raw isotopic signal values (excluding outliers) to avoid any bias due to artifacts of the fitting of the sum of Gaussians.

Fractionation Factor Adjustment for Integrated α -damage—Zircon is widely known to accumulate α -radiation damage (e.g., Zhang et al., 2009 and references therein). It is assumed here that increased α -damage in a zircon leads to a decrease in the hardness of the zircon; this in turn leads to a faster rate of laser penetration into the zircon during ablation leading to shift in isotopic fractionation. Ages calculated for the primary, secondary, and tertiary zircon standards, when those standards were treated as unknowns, were used to construct a fractionation factor correction curve (exponential form). Much previous work has attempted to understand the chemical basis for why some standards work better with some zircons. The notion of matrix-matched standard and unknown zircons has been proposed largely on the basis of trace element chemistry (e.g., Black et al., 2004). Here, time and crystallographic damage, parameters invisible to instruments used to characterize trace element chemistry, were introduced and applied in conjunction with U and Th chemistry.

Common Pb Correction—Common Pb was subtracted out using the Stacey and Kramer (1975) common Pb model for Earth. Ages and common Pb ratio were determined iteratively using a pre-set, session-wide minimum common Pb age value (default for each session was the age of the oldest age standard which for both Apatite and Zircon was 1099 Ma FC-1 and/or FC-5z).

Preferred Age—Uranium decay constants and the $^{238}\text{U}/^{235}\text{U}$ isotopic ratio reported in Steiger and Yäger (1977) were used in this study. $^{207}\text{Pb}/^{235}\text{U}_c$ ($^{235}\text{U}_c = 137.88^{238}\text{U}$), $^{206}\text{Pb}/^{238}\text{U}$, and $^{207}\text{Pb}/^{206}\text{Pb}$ ages were calculated for each data scan and checked for concordance; concordance here was defined as overlap of all three ages at the 1σ level (the use of 2σ level was found to skew the results to include scans with significant common Pb). The background-corrected isotopic sums of each isotope were calculated for all concordant scans. The precision of each isotopic ratio was calculated by using the background and signal errors for both isotopes. The fractionation factor for each data scan, corrected for the effect of accumulated α -damage, was weighted according to the ^{238}U or ^{232}Th signal value for that data scan; an overall weighted mean fractionation factor for all concordant data scans was used for final age calculation. If the number of concordant data scans for a spot was greater than zero, then either the $^{206}\text{Pb}/^{238}\text{U}$ or $^{207}\text{Pb}/^{206}\text{Pb}$ age was chosen as the preferred age, whichever exhibited the lower relative error.

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