



1 Expanding Limits of Laser-Ablation U-Pb Calcite Geochronology

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- 5 Abstract. U-Pb geochronology of calcite by laser-ablation inductively-coupled mass spectrometry (LA-
- 6 ICMPS) is an emerging field with potential to solve a vast array of geologic problems. Because of low
- 7 levels of U and Pb, measurement by more sensitive instruments, such as those with multiple collectors
- 8 (MC), is advantageous. However, whereas measurement of traditional geochronometers (e.g., zircon) by
- 9 MC-ICPMS has been limited by detection of the daughter isotope, U-Pb dating of calcite can be limited
- 10 by detection of the parent isotope, if measured on a Faraday detector. The Nu P3D MC-ICPMS employs a
- 11 new detector array to measure all isotopes of interest on Daly detectors. A new method, described herein,
- 12 utilizes the low detection limit and high dynamic range of the *Nu P3D* for calcite U-Pb geochronology,
- 13 and compares it with traditional methods. A model is created to explore the limits of U, Pb, and U/Pb
- ratios that can be measured by LA-ICPMS and can serve as a guide to evaluate potential candidate
- 15 materials for geochronology.

16 1. Introduction

- 17 Calcite U-Pb geochronology by laser-ablation inductively-coupled mass spectrometry (LA-ICPMS) is a
- 18 relatively new technique with untapped potential for solving numerous geochronologic problems from the
- timing of faulting (e.g., Roberts and Walker, 2016; Nuriel et al., 2017; Goodfellow et al., 2017), age of
- 20 ore deposits (Burisch et al., 2017) to paleoclimate, sedimentation, and diagenesis (e.g., Mangenot et al.,
- 21 2018; Rasbury et al., 1997; Hoff et al., 1995; Winter and Johnson, 1995; Wang et al., 1998; Rasbury et
- 22 al., 1998). Early studies focused on carbonates more likely to contain high concentrations of U, such as
- 23 speleothems (e.g., Richards et al., 1998) because the method employed—thermal ionization mass
- 24 spectrometry (TIMS)—required weeks to produce reliable ratios; samples with a low likelihood of
- success, that is, those with potentially low U contents, were ignored. With the advent of LA-ICPMS,
- 26 however, sample throughput and analytical costs have been greatly reduced, such that hundreds of





27	geoanalytical facilities can, at the very least, screen a large number of samples and choose those suitable
28	for geochronology in a relatively short period of time and for little cost. Actual analysis time and cost are
29	also reduced over TIMS—sample preparation is minimal and several samples can be analyzed in a day—
30	and dozens of labs worldwide have the capability to perform such analyses. LA-ICPMS also has the
31	advantage of sampling smaller volumes of material; it can thus take advantage of the heterogenous nature
32	of calcite with respect to U and Pb, using larger datasets to better constrain both the initial ²⁰⁷ Pb/ ²⁰⁶ Pb
33	compositions and the common Pb-corrected concordia ages. These isochron ages are calculated with ease
34	on a Tera-Wasserburg diagram similar to other common-Pb-bearing mineral chronometers like titanite
35	and apatite (e.g., Chew et al., 2014; Spencer et al., 2013).
36	For typical LA-ICPMS analyses, a 193 nm excimer laser is employed in conjunction with either a single-
37	collector (SC-ICPMS), or multi-collector (MC-ICPMS) sector-field instrument to take advantage of the
38	increased sensitivity over a quadrupole (Q-ICP-MS). Traditionally, an MC-ICPMS uses a series of
39	Faraday detectors on the high-mass side of the detector array to measure ²³⁸ U and ²³² Th, and either
40	Faraday cups or secondary electron multipliers (SEMs) on the low-mass side of the array to concurrently
41	measure Pb isotopes; SC-ICMPS instruments measure isotope count rates sequentially with a single SEM.
42	The SC and MC instruments have distinct advantages. Because there is only one SEM on a SC-ICPMS
43	instruments, there is no need to cross calibrate multiple detectors, yielding simpler data reduction and the
44	possibility for making 204- or 208-based common-Pb corrections. An MC-ICPMS, on the other hand, is
45	2-3 times more sensitive than the top SC-ICPMS instruments. This allows precise measurements of
46	samples with low levels of Pb (i.e., young and/or low common-Pb). Furthermore, its equivalent sensitivity
47	is even higher because it measures all masses at the same time. For example, a SC-ICPMS running only
48	masses 238, 207, and 206 (232, 208, and 204 are also typically measured) at equal dwell times measures
49	1/3 the counts over a given cycle than the count rate might suggest because only one mass can be
50	measured at a time; given that it is also 2-3x less sensitive than an MC-ICPMS, a laser spot must be ~6-9
51	times bigger to achieve the same precision on a SC-ICPMS. A further advantage of an MC-ICPMS is that





transient signals from changes in U and Pb concentration during ablation affect uncertainties in the 52 measured ²⁰⁷Pb/²⁰⁶Pb and ²⁰⁶Pb/²³⁸U less because all measurements are made concurrently. Finally, the 53 smaller dynamic range of the SEM can limit samples to a specific range of U concentrations; samples or 54 reference materials with high U contents can cause the detector to trip to a different measurement mode 55 (or trip off), yielding spurious results. Low U concentrations in calcite can also be a problem for an MC-56 ICPMS measuring 238 U with a Faraday cup because limits of detection are on the order of 10^4 cps. Even 57 58 though the SC-ICPMS has 2-3 times lower sensitivity than an MC-ICPMS, it can precisely measure count rates of $\sim 10^2$ cps by employing a secondary electron multiplier (SEM) for all masses. Because of its 59 60 reduced sensitivity, however, this equates to a very small range of samples. Fortunately, a recently introduced MC-ICPMS—the P3D—by Nu Instruments (Wrexham, UK) can 61 62 overcome both of these limitations. The instrument features a Daly detector array that allows for ion 63 counting on ²³⁸U and the Pb isotopes, and thus expands the range of calcite samples—those with lower U

64 concentrations—that can be precisely measured by LA-ICPMS. Not only does a Daly detector increase

the sensitivity of the instrument, but unlike a standard SEM, the Daly has a greater dynamic range

66 (approx. 10-fold over that of an SEM) and can thus be used with a larger range of U concentrations. This

67 contribution describes the analytical setup for LA-ICPMS using the new Nu P3D, comparing the two

68 modes with each other and with that of a SC-ICPMS, and thereby demonstrating the increased capability

69 of this new instrumentation to measure calcite U-Pb dates.

70 2 Experimental Setup

71 The analytical setup is described in Table 1. The instrumentation used in the study consists of a *Photon*

72 *Machines Excite* 193 nm excimer laser equipped with a HelEx cell, coupled to a *Nu Instruments P3D* for

- standard LA-ICPMS analyses. The Nu Plasma 3D (P3D) contains an array with 6 Daly detectors, 5 on the
- 74 low-mass side of the array and 1 on the high-mass side. A 14-Faraday array lies between the Daly
- detectors, and allows for measurement of ²³⁸U on either a Faraday or Daly detector, depending on the U
- concentration in the sample. Daly detectors are used to measure masses 202, 204, 206, 207, and 208, and





- ²³²Th is measured on a Faraday cup. Faraday backgrounds yield a 1SD of 0.04 mV, which implies a limit
- of detection (LOD) of ~0.1 mV or ~8000 cps; Daly backgrounds yield 1SD of 10–20 cps for isotopes of
- 79 Hg and Pb and 1 cps for ²³⁸U, corresponding to LODs of 30–60 and 3 cps, respectively.
- 80 In order to compare the difference between SC and MC analytical sensitivities and uncertainties, the laser
- 81 was used in conjunction with the *P3D* for two experiments, and an *Agilent 7700* Q-ICPMS for one
- 82 experiment. These 3 experiments were run with different spot sizes: *Experiment F*) a 65 μ m spot on the
- *P3D* using a Faraday for masses 238 and 232, and Daly detectors for masses 208–204 and 202 (110 total
- analyses); *Experiment D*) the same configuration, but with 238 measured on a Daly detector; and
- 85 *Experiment Q*) a 110 μm spot with the Q-ICPMS and cycle times of 0.06, 0.13, 0.1, and 0.1 s on masses
- 86 238, 207, 206, and 204 respectively. During each separate analytical run, each spot was located near the
- 87 corresponding spot from the other runs, to minimize uncertainty caused by grain homogeneity. For all
- experiments, the laser was run at 10 Hz for 15 seconds and a fluence of approximately 1 J/cm², yielding a
- spot depth of 10–15 μm. Analyses were preceded by two pre-ablation pulses, and 20 seconds of baseline
 measurement.
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Three calcite samples from the east coast of North America (courtesy of W. Amidon of Middlebury 92 93 College) were the main samples measured. These samples—C258, C304, and C273—are ca. 440, 110, and 80 Ma, respectively, and range in U concentration between a few ppb and a few ppm, with an average 94 95 of 120 ppb and a mode of ~20 ppb. Three further samples (C254, C283A, and C283B), were run in 96 experiments F and D and provide more data for uncertainty comparisons between the two instrumental 97 configurations (see Figure 1), but the data are described in less detail; they are ca. 440 Ma with variable 98 Cretaceous (?) (re)crystallization. Calcite and NIST614 reference materials (RMs) were interspersed 99 every 10 analyses, and a two-stage reduction scheme was employed. *Iolite v.3.0* (Paton et al., 2011) was used first used to correct the ²⁰⁷Pb/²⁰⁶Pb for mass bias, detector efficiency, instrumental drift etc., and to 100





- 101 correct the 238 U/ 206 Pb ratio for instrumental drift, using NIST614 as the primary reference material.
- 102 During this first data reduction, 2 seconds were removed from both the beginning and end of both the
- 103 RMs and the unknowns, yielding a total count time of 11 s. The 238 U/ 206 Pb ratio was then corrected using
- a linear correction in *Excel* such that the primary calcite RM, WC-1, yielded 254 Ma (Roberts et al.,
- 105 2017) on a Tera-Wasserburg (TW) diagram, anchored to a ²⁰⁷Pb/²⁰⁶Pb value of 0.85. Using this method,
- 106 we retrieved ages of 3.01 ± 0.15 (MSWD = 1.3; n = 30) and 65.9 ± 1.1 (MSWD = 1.2; n = 40) for
- secondary RMs ASH15 (2.96 Ma; Nuriel et al., in review) and Duff Brown Tank (64 Ma; Hill et al.,
- 108 2016), respectively. Analyses with large uncertainties (arbitrarily chosen as 50% for both ²³⁸U/²⁰⁶Pb and
- $109 \quad {}^{207}\text{Pb}/{}^{206}\text{Pb}$) were discarded; removing these data has little influence on the final age. The data from the
- unknowns are all a bit scattered for geological reasons, and were culled to yield single populations for
- ease of comparison. (Though beyond the scope of this manuscript, the Paleozoic samples are interpreted
- 112 to have suffered partial Pb loss or new crystal growth in the Cretaceous-Tertiary, and the older
- 113 Cretaceous sample likely (re)crystallized over an extended period.)

114 3 Results

- 115 Table 2 and Figure 2 shows the results for the 6 samples analyzed in the 3 experiments. *Experiment F*
- 116 (*P3D* 65 μm spot; U on a Faraday) yielded ~2.7 mV/ppm of mass 238 on NIST614 and was relatively
- 117 stable throughout the run. The sensitivity of *Experiment D* ($P3D 65 \mu m \ spot$; U on a Daly) was similar
- to that of *Experiment F*, but dropped approximately 25% during the analytical session to $\sim 2 \text{ mV/ppm of}$
- 119 mass 238 on NIST614. Experiment Q (Agilent Q-ICPMS 110 μm spot) yielded ~110 kcps/ppm of mass
- 120 238 on NIST614—equivalent to ~1.8 mV/ppm from a spot ~3 times larger than the 65 μm spot in
- 121 experiments 1 and 2—and was stable throughout the run.
- 122 For every sample, Experiment F yielded fewer analyses with uncertainties of <50% for ²⁰⁶Pb/²³⁸U, as well
- as the fewest spots available to make an isochron. These results are consistent with a higher average and
- 124 median U ppb; low U concentrations that were measured in Experiments D and Q went undetected or
- 125 yielded large uncertainties in Experiment F. Though samples with median ²³⁸U count rates of >10,000 cps





- 126 (C273C and C304A) returned fewer viable analyses and worse average ²³⁸U/²⁰⁶Pb uncertainties in
- 127 Experiment F, the uncertainty of the final age was similar for the higher-U samples on both
- 128 configurations on the *P3D*; both yielded lower uncertainties than the Q-ICPMS, despite the 3-fold volume
- increase in analyzed material on the Q-ICPMS.
- 130 When average count rates of 238 U were below ~8000 cps (near the detection limit of the Faraday detector
- 131 on the *P3D*), however, the number of viable analyses and final age uncertainty was significantly higher in
- 132 Experiment D (Table 2 and Figure 2). As an example, sample C258 yielded few viable data points (35%
- 133 of the 110 analyses) in Experiment F, fewer than half the number of good analyses in Experiments D and
- 134 Q. In addition, the resulting uncertainty in the final age calculation (~4%) is significantly larger than that
- 135 of Experiment D, and similar to the resulting uncertainty in Experiment Q (although the Q-ICPMS
- 136 yielded >2 times the number of viable spots). Samples C283A and C283C—which also contain low levels
- 137 of U—yielded ~50% fewer viable data, necessitated double the average count rates of ²³⁸U, and final
- uncertainties that were significantly greater in Experiment F than those of Experiment D.
- A summary of the precision vs. U count rate is shown in Figure 1, which shows the precision of $^{238}U/^{206}Pb$
- and 238 U on a single spot vs. the count rate of 238 U. While there is considerable overlap in the precision vs.
- 141 ²³⁸U cps of both ²³⁸U and ²³⁸U/²⁰⁶Pb at count rates above approx. 30,000 cps, data collected in Experiment
- 142 F yielded no better than a few kcps 2σ uncertainty on ²³⁸U; ²³⁸U/²⁰⁶Pb uncertainties consequently show a
- 143 similar deviation from the high-count-rate trend. Finally, though the Q-ICPMS shows similar gains in
- 144 precision for low-U analyses, the lower sensitivity of the Q-ICPMS results in a minor window of U
- 145 concentrations for which analyses have lower uncertainties than those run on the *P3D*.

146 4 Discussion

147 While there is a clear advantage of using the new Daly-only detector setup on the P3D for LA-based

148 calcite geochronology for some samples, the extent to which this advantage obtains for all samples is still

149 somewhat ambiguous. The samples that benefit most from the new instrumentation are not only low in U,





but also older. For most measurements of long-lived-isotope geochronology, the analytical limit is 150 151 determined by the detection limit of the daughter, not the parent, isotope. However, because older samples have more daughter product, they are-for samples with low U/Pbc ratios-analyses of those 152 153 samples are more likely to be limited by the count rate of the parent isotope. For samples run on a SC-ICPMS, this distinction is unimportant because the detection limit of ²³⁸U is in all cases lower than that 154 155 for Pb. However, because the MC-ICPMS has a large sensitivity and precision advantage over the SC-ICPMS, it is important to distinguish the limits of measurement between the Faraday–Daly and all-Daly 156 configuration. 157

158 4.1 Theoretical uncertainty of Tera–Wasserburg data

159 To explore the limits of precision for each analytical configuration, a dataset was created to represent different U/Pbc and ²³⁸U cps for samples of different ages. Figure 3 shows samples with ages of 440, 80, 160 161 and 15 Ma with error ellipses at U/Pbc ratios of 1, 2, 5, 10, 20, 100 and 200. The size of the ellipse is the maximum possible uncertainty (from counting statistics only) for a 10s analysis, given the limit of 162 detection of the instrument. For the all-Daly configuration, the limit of detection is determined by ²⁰⁷Pb 163 164 counts, the least abundant isotope of interest. For this example, 30 cps is assumed (the best achieved 165 LODs herein; Gerdes et al), but it is important to recognize that the LOD of Pb is based on the 166 background, which varies from lab to lab, and is also a function of the instrumental sensitivity. For the Faraday–Daly arrangement, the LOD is limited by ²³⁸U counts for samples with lower U/Pbc and by ²⁰⁷Pb 167 168 for samples with high U/Pb_c—and increasingly so as the sample age decreases. In this case, a minimum of 30,000 cps of ²³⁸U is considered—as opposed to the actual ca. 8000 cps LOD—for the Faraday, because 169 170 that is the count rate below which a distinct benefit in precision is gained by using the all-Daly 171 arrangement (see Figure 1 and discussion above). As depicted in Figure 3, older samples yield the greatest range of U/Pb_c ratios that could yield an advantage of measurement by ²³⁸U on an ion counter, whereas 172 173 the advantage of the Daly detector disappears at U/Pb_c ratios greater than ca. 500 and 250 for samples that are 80 and 15 Ma, respectively. As an example of the benefit of ²³⁸U measurement by Daly, an 80 Ma 174





- sample with a maximum U/Pbc ratio of 10 yields 1400 cps of ²³⁸U at the LOD of 30 cps ²⁰⁷Pb. Given a
- 176 limit of detection of 8000 cps for the Faraday detector, the signal size would need to be 6 times higher
- 177 before it could be measured by such means. Furthermore, as discussed above, and shown in Figure 1, the
- 178 benefit of the Daly extends to ca. 30,000 cps, or ~20 times the signal that can be measured by the
- 179 Faraday–Daly configuration. The benefit extends to 200 times for a U/Pbc ratio of 1; but some question
- arise as to the ability to measure ages at such low U/Pb_c values.
- 181 4.2 Choosing Samples and Instruments
- 182 One intention of this manuscript is to serve as a guide to determine whether any given calcite (or any
- 183 other Pb_c-bearing) sample is appropriate for U–Pb geochronology, and deciding which type of analytical
- 184 equipment to use. As such, the model above is expanded below to explore the U/Pb_c ratios and count rates
- 185 needed to produce a reliable age from a given number of analyses. These models are then compared with
- the natural results to determine best practices when selecting samples and instruments for analysis.
- 187 Calculating theoretical limits is complicated, however, because the uncertainty of an isochron depends on
- 188 the distribution of U, Pb_c, and thus the distribution of U/Pb and Pb/Pb ratios. For example, a sample with
- a given maximum U/Pb_c will yield a final precision that increases with the number of analyses, but this
- improvement depends on the distribution of the U/Pb_c ratios. The distribution of U and Pb, and thus
- 191 ²³⁸U/²⁰⁶Pb and ²⁰⁷Pb/²⁰⁶Pb, in calcite has not been a particular subject of study, but a cursory analysis of
- the reference materials and unknowns presented in this manuscript shows that U and Pb concentrations
- 193 follow normal distributions; RMs that contain sufficient U (WC-1 and Duff Brown Tank) display a near-
- 194 normal distribution of U, whereas the distribution of U concentration of samples and RMs with lower U
- contents (ASH15 and unknowns) are log normal (Figure 4). Like U, Hg-corrected ²⁰⁴Pb counts (a proxy
- 196 for common Pb) are normally distributed in RMs and unknowns; ²⁰⁸Pb counts are similar. The resulting
- 197 ²³⁸U/²⁰⁶Pb ratios of RMs are normally distributed, but unknowns vary and can be rather uniform (e.g.,
- 198 C273C). The manner by which the type of distribution affects the final uncertainty is demonstrated in
- 199 Figure 5. The precision of a T-W isochron is best defined by precisely defined end points with maximum





200	spread; as such, except for samples with extreme U/Pb _c , a uniform distribution of 238 U/ 206 Pb ratios results
201	in better final age precision than does a normal distribution. For example, a sample that is 440 Ma with
202	normally distributed data (and ratios $\pm 3\sigma$ from the mean) requires nearly 2 times as many points to
203	achieve the same precision as a sample with uniformly distributed data over the same U/Pb range (though
204	this also depends on the maximum U/Pbc). For normally distributed data with the same maximum U/Pbc,
205	but only 50% of the spread (i.e., more tightly clustered; Figure 5b), the number of necessary data points
206	increases further, excepting samples with extreme U/Pbc (these data would be less dependent on the
207	precision of the upper intercept).
208	To compare theoretical data with that obtained from this study—i.e., in order to best represent a natural
209	dataset—we present and discuss models with 100 uniformly distributed ²³⁸ U/ ²⁰⁶ Pb data points acquired for
210	10 s at 10 Hz, recognizing that, as stated above, this is likely a best-case scenario. We explore the
211	implications of varying maximum U/Pb _c ratios rather than 238 U/ 206 Pb ratios because the former are
212	independent of sample age. The results of the model are shown in Figure 6 Because the precision of
213	analyses in an ion-counter-only configuration is limited by the count rate of ²⁰⁷ Pb, we calculate the
214	maximum U/Pbc ratio that can be achieved for different concentrations of U. For example, a 440 Ma
215	sample with 10 ppb U run with a 65 μ m spot size will yield ~1500 cps of U. The maximum U/Pb _c that
216	could be achieved with this count rate will be ~13, because any higher values will yield too few counts of
217	207 Pb to be measured. Assuming constant U concentration and normally distributed 238 U/ 206 Pb ratios, the
218	best precision on the age of this sample is 0.6%—considerably better than expected for LA-ICPMS (e.g.,
219	Horstwood et al., 2016). As a comparison, sample C283A contains an average of 10 ppb U (and
220	maximum of 40 ppb) and thus yields a similar average count rate of 238 U. Its maximum U/Pb _c of 26 is
221	considerably less than the maximum theoretical value based on the concentration of that particular
222	analysis because its Pb concentration is well above detection. It should be no surprise then, that the age
223	uncertainty is higher than the theoretical value at that count rate, but it is also higher than the theoretical
224	value for a U/Pb_c of 26. Several factors may explain this: 1) though 100 analyses were measured, 32 were





- imprecise and rejected; 2) the distribution of ²³⁸U/²⁰⁶Pb ratios is not uniform; 3) laser instability, detector
 response time, laser-induced elemental fractionation (LIEF), signal instability, etc. add uncertainty
 beyond that based on counting statistics; and 4) low U/Pbc values likely have less U and Pb than in the
- 228 model.

229 Although optimistic, this model serves as a guide for the limitation of analyses of calcite by LA-ICPMS, 230 given U concentration, maximum U/Pbc, and spot size. First, for all but the youngest samples (<<15 Ma), 231 measurement with the P3D can be advantageous for samples with lower U or those necessitating small spot sizes (e.g., <150 ppb U and $<65 \mu m$; <50 ppb U and $<125 \mu m$). However, if, for example, the sample 232 contains concentrations >100 ppb U and the spot can be >100 μ m, there is no advantage to using the all-233 234 Daly configuration, and if there is significant material (i.e., spot size can be $>200 \,\mu$ m), any LA-ICPMS 235 will provide the best possible results (that is, the precision will be limited not by the count rate, but rather 236 other factors such as differences in LIEF, matrix effects etc.). Second, it is highly unlikely that even with 237 extreme spot sizes and rep rates, that samples with <<1 ppb U can be analyzed. Third, older samples— 238 when run on the P3D—reach their best possible uncertainty with U concentrations of 10–15 ppb; samples as young as 80 Ma require little more than 30 ppb U, and samples as young as 15 Ma require up to 150 239 240 ppb U at moderate spot sizes. Though 2% final uncertainty requires greater concentrations of U for younger samples (>2500 cps²³⁸U are needed for an 80 Ma sample, and >12,000 cps²³⁸U for a 15 Ma 241 242 sample), it should be noted that—at a given concentration, spot size and U/Pb_c—absolute uncertainty is relatively independent of age; for example, a sample with a 65 μ m spot and 10 ppb U yields an 243 244 uncertainty of just over 2 Ma, whether the sample is 15, 80, or 440 Ma. Finally, though not depicted directly in Figure 6, precise ages can be obtained from data with rather low U/Pbc values. For example, 245 246 100 spots with 2% uncertainty yields a final uncertainty of 5–15 Ma (2σ) for samples with U/Pb_c ratios as 247 low as 1–2. That said, data with such low U/Pbc ratios should be viewed with caution, as systematic 248 uncertainties-such as those introduced by inconsistencies in RM isotopic measurements-can lead to 249 large errors when extrapolating data clustered near the upper intercept.





250 **4.3 More spots, deeper spots, or bigger spots?**

251	The theoretical models discussed above use a 10 sec integration time to compare the models to the
252	empirical data. As discussed above, precision can be improved by increasing the number of analytical
253	spots, but each spot can also be ablated for longer or at a higher rep rate (i.e., making deeper pits rather
254	than more pits). One might imagine that these methods might be equally effective, however, there are two
255	important points to consider. First, individual spot precision is limited to the long-term reproducibility of
256	down-hole measurements, and is generally no better than 2%; this precision is more difficult to assess in
257	calcite because most known reference materials exhibit moderate isotopic heterogeneity (e.g., Roberts et
258	al., 2017). Thus, if increasing the depth of the pit yields analytical uncertainties <2%, then the excess pit
259	depth is wasted and overall uncertainty fails to improve. Second, whereas increasing the number of spots
260	leads to a linear increase in the total number of counts (and thus an increase in precision by \sqrt{n}), an
261	increase in pit depth does not lead to a linear increase in counts because ablation yields decrease with pit
262	depth. Thus, if an increase in total counts could yield better precision, that increase should come from
263	more, shallower laser pits, rather than fewer, deeper pits.
264	It is also possible to increase precision by increasing the spot size. In fact, an argument could be made
265	that a SC-ICPMS that measures 250 μm spots is just as effective as a MC-ICPMS that measures 100 μm
266	spots. Though this argument has merit, the downside is twofold; 1) some regions of interest are simply
267	not large enough to permit a spot 2.5X as wide, and 2) U and/or Pb (i.e., U/Pb _c) may be heterogeneous at
268	scales smaller than the spot size, mixing calcite of different age or reducing the range of isotopic ratios
269	that are used to construct an isochron. Figure 7 demonstrates that even though larger spots can yield a
270	better per-spot precision, analyzing the same volume of material with smaller spots can yield better age
271	precision because it can take advantage of the heterogeneous U and Pb concentrations typical of calcite.

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272 5 Conclusions

- 273 1) Unlike geochronometers with high U and little to no common Pb—such as zircon and monazite—U-Pb
- 274 dates of minerals with low U and significant common Pb can be limited by the count rates of the parent
- 275 U, rather than the daughter Pb.
- 276 2) Given a limit of detection of ~8000 cps for on a Faraday, and the sensitivity of the Nu P3D, samples
- with as low as 20 ppb U can be analyzed with a 100 µm spot at 10 Hz, and as low as 5 ppb for a 200 µm
- 278 spot. Even so, the Faraday is less precise than the Daly at count rates of <30,000 cps, corresponding to U
- concentrations of ca. 75 and 20 ppb, with the same respective spot sizes and rep rates.
- 280 3) When 238 U is analyzed on a Daly, the limit of detection drops by a factor of >1000, and the analytical
- capability is thus limited by the LOD of Pb—²⁰⁷Pb in almost all cases—and the ratio required for
- optimum precision. The typical LOD of ²⁰⁶Pb and ²⁰⁷Pb is ca. 50 cps; it is greater for higher sensitivity
- instruments, and those with a higher background of common Pb. For a desired U/Pbc ratio of ca. 5–10 for
- old and young samples, respectively, the required count rate of ²³⁸U would be 500–1000 cps or ca. 5–10
- times smaller than can be analyzed on a Faraday detector. The analysis of ²³⁸U on a Daly, therefore
- increases the analytical capability to ca. 0.5–2 ppb U for a 100–200 µm spot, respectively.
- 287 4) Although the % uncertainty that can be achieved with limited concentrations of U is considerably
- 288 different among samples with different ages, the absolute uncertainty is approximately the same. For
- example, samples with 1500 cps 238 U yield a maximum possible uncertainty of ca. 2 Ma, nearly
- 290 independent of age (older samples yield slightly higher absolute uncertainties). However, because most
- 291 LA-ICPMS facilities can achieve up to 2% precision on final age calculations, younger samples can yield
- better absolute uncertainties; these can only be achieved at high U concentrations, which limits the
- advantage of the *Nu P3D* for young samples.
- 5) Given enough material and analytical time, a SC-ICPMS, should, in theory, be capable of measuring
- samples with concentrations of approximately 2–10 times (i.e., 1–20 ppb U) that of the Nu P3D.





- 296 However, because of their lower cycle times and inability to make concurrent measurements, SC-ICPMS
- 297 instruments likely require considerably higher concentrations of U to obtain comparable date precision.

298 Figure Captions

- Figure 1. Relation between cps ²³⁸U and uncertainty of ²³⁸U (A), and ²⁰⁶Pb/²³⁸U (B). The 3 experiments
- show the same trend in uncertainty vs. cps at count rates above ~30 kcps 238, but below that, uncertainty
- 301 of measurements in Experiment F (238 on the Faraday) increase significantly compared to Experiments D
- and Q. Although Experiments D and Q (red and blue symbols) show the similar trends, the sensitivity
- 303 gain using the P3D leads to significant improvements in spot uncertainty (large symbols represent
- and expected uncertainties for a 100 um spot at 10 ppb U).
- Figure 2. Tera–Wasserburg concordia diagrams of the 3 unknown samples in each of the 3 experiments.See text for discussion.
- 307 Figure 3. Uncertainty ellipses for each Tera–Wasserburg plot depict the counting uncertainty for 10 s at a
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- 309 ellipse is the limit of detection for the all-Daly configuration, or any SC-ICPMS (limited by ²⁰⁷Pb counts).
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distribution over the upper 50% of the same range. The uniform distribution, shown in 5C, yields the 321 lowest uncertainties because there are more analyses at both the upper and lower intercepts. D shows how 322 the percent uncertainty decreases with number of analyses, depending on the type of $^{238}U/^{206}Pb$ 323 324 distribution depicted in A-C; data in D assumed the best case scenario of 2% uncertainty per data point and a U/Pbc ratio of 10 for samples of 440 Ma, 80 Ma, and 15 Ma. Best uncertainties are achieved with 325 uniform distributions and maximum spread. Although percent uncertainties are always better for older 326 samples, younger samples yield better absolute uncertainties for well distributed data. 327 Figure 6. 6A shows the count rate expected with the Nu P3D given for a given spot size at a laser energy 328 of ~1 J/cm2 and 10 Hz. Spots indicate analyses of unknowns in Experiment F (²³⁸U on the Faraday; color 329 330 represents the maximum U/Pbc ratio-taken from Table 2-and the size represents the uncertainty). Dark grey area is below the LOD for ²³⁸U on a Faraday; the light grey area represents ²³⁸U count rates favorably 331 measured on the Daly detector. Figures B, D, and F show the maximum possible U/Pbc ratio that can be 332 detected, given a minimum count rate of 30 cps for ²⁰⁷Pb at different sample ages. C, E, and G show the 333 334 best possible uncertainty at the given count rates and spot sizes for 100 analyses, all with the same U concentration but a uniform distribution of ²³⁸U/²⁰⁶Pb ratios. 335

Figure 5. A-C shows an example of the differing randomly generated distributions of 100 analyses with

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Figure 7. Tera–Wasserburg diagram representing the analysis of a heterogeneous medium using different
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Table 1.

Instrumental parameters of laser-ablation split-stream ICP-MS

r r	MC-ICP-MS	Q-ICP-MS		
Instrument model	Nu Plasma 3D	Agilent 7700x		
RF forward power	1300 W	1300 W		
RF reflected power	<10 W	<10 W		
Coolant gas	13 L/min	13 L/min		
Auxiliary gas	0.8 L/min	0.8 L/min		
Make up gas	~1 L/min	~1 L/min		
Monitored masses	²³⁸ U, ²³² Th, ²⁰⁸ Pb, ²⁰⁷ Pb,	²³⁸ U(0.06), ²⁰⁷ Pb (0.13), ²⁰⁶ Pb		
(dwell times listed for Agilent)	²⁰⁶ Pb, ²⁰⁴ Pb/ ²⁰⁴ Hg, ²⁰² Hg	(0.1), ²⁰⁴ Pb/ ²⁰⁴ Hg (0.1)		
²³⁸ U sensitivity, dry solution	0.5% (23 Mcps/ppb)	0.1% (4 Mcps/ppb)		
_	Laser-Ablation System			
Instrument model	Photon Machines Analyte			
	193			
Laser	ATLEX-SI 193nm ArF			
	excimer			
Fluence	~1 J/cm ²			
Repetition rate	10 Hz			
Excavation rate	~0.07 um/pulse			
Spot size	65–110 μm			
Delay between analyses	20 s			
Ablation duration	15 s			
Carrier gas (He) flow (cell; cup)	0.12; 0.06 L/min			





Table 2. Results from 3 experiments

sample #	C258	C273C	C304A	C283A	C283C	C254A			
Experiment F (P3D - 238 or	n Faraday; 65	µm, ~2.7 mV/	(ppm U)						
total spots	110	100	100	100	100	100			
$^{238}U/^{206}Pb \ 2\sigma < 50\%$	54%	76%	63%	29%	38%	63%			
spots for isochron	35%	76%	47%	21%	25%	n/a			
average U ppb	40	195	286	28	25	456			
median U ppb	30	73	96	25	27	55			
average cps 238	7100	33800	46800	4600	4100	73100			
median cps 238	5300	12700	15700	4100	4400	8800			
avg. ²³⁸ U/ ²⁰⁶ Pb 2σ	28%	17%	17%	32%	35%	24%			
maximum U/Pbc	49	145	54	27	17	n/a			
Age (Ma)	437 ± 18	80.9 ± 1.5	111.1 ± 2.1	453 ± 40	492 ± 81	n/a			
final 2 σ	4.1%	1.9%	1.9%	8.8%	16.5%	n/a			
Experiment D (P3D - 238 on Daly; 65 µm, ~2.1–2.7 mV/ppm U)									
total spots	100	100	100	100	100	100			
²³⁸ U/ ²⁰⁶ Pb 2σ <50%	96%	98%	97%	97%	93%	90%			
spots for isochron	75%	90%	84%	64%	68%	n/a			
average U ppb	24	144	196	11	18	232			
median U ppb	13	59	40	8	18	33			
average cps 238	3800	24500	27900	1600	2400	29300			
median cps 238	2100	10000	5700	1200	2400	4200			
avg. ²³⁸ U/ ²⁰⁶ Pb (20)	16%	10%	12%	17%	16%	19%			
maximum U/Pbc	30	205	79	26	12	n/a			
Age (Ma)	445 ± 11	83.5 ± 1.6	119.3 ± 2.3	430 ± 11	430 ± 14	n/a			
final 2σ	2.5%	1.9%	1.9%	2.6%	3.3%	n/a			
Experiment Q (Agilent 770	0 Q-ICPMS;	110 µm, ~1.8 n	nV/ppm U)						
total spots	110	100	100						
$^{238}U/^{206}Pb \ 2\sigma <50\%$	96%	87%	94%						
spots for isochron	75%	87%	94%						
average U ppb	28	126	371						
median U ppb	14	68	80						
average cps 238	3100	14400	39400						
median cps 238	1600	7800	8500						
avg. ²³⁸ U/ ²⁰⁶ Pb (20)	20%	14%	13%						
maximum U/Pbc	26	325	93						
Age (Ma)	460 ± 18	85.4 ± 2.0	118.1 ± 4.0						
final 2σ	3.9%	2.3%	3.4%						







Figure 1. Relation between cps 238U and uncertainty of 238U (A), and 206Pb/ 238U (B). The 3 experiments show the same trend in uncertainty vs. cps at count rates above ~30 kcps 238, but below that, uncertainty of measurements in Experiment F (238 on the Faraday) increase significantly compared to Experiments D and Q. Although Experiments D and Q (red and blue symbols) show the similar trends, the sensitivity gain using the P3D leads to significant improvements in spot uncertainty (large symbols represent expected uncertainties for a 100 um spot at 10 ppb U).







Figure 2. Tera–Wasserburg concordia diagrams of the 3 unknown samples in each of the 3 experiments. See text for discussion.









Figure 3. Uncertainty ellipses for each Tera–Wasserburg plot depict the counting uncertainty for 10 s at a given 238U count rate for different U/Pbc ratios of 1, 2, 5, 10, 20, 50, 100, 200. For each U/Pbc, the larger ellipse is the limit of detection for the all-Daly configuration, or any SC-ICPMS (limited by 207Pb counts). The smaller, red ellipse indicates the uncertainty at 30,000 cps 238U, the point at which the measurement of 238U on the Daly is no longer advantageous.







Figure 4. Left-hand plots show the difference in distribution of 238U/206Pb ratios in reference materials and unknowns; ratios are nomalized to the 238U/206Pb ratio of the age of the sample. Reference materials Duff Brown and WC-1 have the smallest variation in 238U/206Pb ratios, which correlates well with the distribution of their U and Pb contents (left-hand plots). Reference material ASH15 and unknown sample C283C still have a wider log-normal distribution, reflective of their larger distribution of U and Pb contents relative to Duff Brown and WC-1. Unknown sample C273C has a more uniform distribution of 238U/206Pb ratios, reflecting its largest distribution of U contents.







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Figure 7. Tera–Wasserburg diagram representing the analysis of a heterogeneous medium using different spot sizes. Though the bigger spot sizes yield smaller individual uncertainties, the smaller spots take advantage of the spread in U/Pbc ratios and thus yield a better overall uncertainty on the lower intercept age.