

# Evaluating Uncertainties in the Calibration of Isotopic Reference Materials and Multi-Element Isotopic Tracers (EARTHTIME Tracer Calibration Part II)

Noah M. McLean<sup>a,b,\*</sup>, Daniel J. Condon<sup>b</sup>, Blair Schoene<sup>c</sup>, Samuel A. Bowring<sup>a</sup>

<sup>a</sup>*Department of Earth, Atmospheric, and Planetary Sciences, Massachusetts Institute of Technology, Cambridge, MA 02139, USA*

<sup>b</sup>*NERC Isotope Geosciences Laboratory, British Geologic Survey, Keyworth, Nottinghamshire NG12 5GG, UK*

<sup>c</sup>*Princeton University, Department of Geosciences, Guyot Hall, Princeton, NJ 08544, USA*

---

## Abstract

A statistical approach to evaluating uncertainties in the calibration of multi-element isotopic tracers has been developed and applied to determining the isotopic composition of mixed U-Pb ( $^{202}\text{Pb}$ - $^{205}\text{Pb}$ - $^{233}\text{U}$ - $^{235}\text{U}$ ) tracers used for accurate isotope dilution U-Pb geochronology. Our experiment, part of the EARTHTIME initiative, directly links the tracer calibration to first-principles measurements of mass and purity that are all traceable to SI units, thereby quantifying the accuracy and precision of U-Pb dates in absolute time. The calibration incorporates new more accurate and precise purity measurements for a number of commonly used Pb and U reference materials, and requires inter-relating their isotopic compositions and uncertainties. Similar methods can be used for other isotope systems that utilize multiple isotopic standards for calibration purposes. We also detail the inter-calibration of three publicly available U-Pb gravimetric solutions, which can be used to bring the same first-principles traceability to in-house U-Pb tracers from other laboratories. Accounting for uncertainty correlations in the tracer isotope ratios yields a tracer calibration contribution to the relative uncertainty of a  $^{206}\text{Pb}/^{238}\text{U}$  date that is only half of the relative uncertainty in the  $^{235}\text{U}/^{205}\text{Pb}$  ratio of the tracer, which was historically used to approximate the tracer related uncertainty contribution to  $^{206}\text{Pb}/^{238}\text{U}$  dates. The tracer uncertainty contribution to  $^{206}\text{Pb}/^{238}\text{U}$  dates has in this way been reduced to <300 ppm when using the EARTHTIME and similarly calibrated tracers.

*Keywords:* tracer calibration, isotopic standards, inverse methods

---

## 1. Introduction

Our understanding of the rates and timing of events in Earth history depends on radioisotopic dating, or geochronology. An increasing demand for seamless integration of geochronologic data acquired with multiple radioisotopic dating methods, and with astrochronology, has motivated continued improvements in measurement precision, ongoing assessment of the accuracy of parent

---

\*Principal Corresponding Author  
email: noahm@bgs.ac.uk

6 radionuclide decay constants, and characterization of the compositions of reference materials used for  
7 calibrations and assessment of reproducibility. U-Pb dates are ultimately derived from experiments  
8 that utilize isotope dilution principles combined with isotope ratio mass spectrometry (Stracke et al.,  
9 2014, and references therein), an approach that permits the translation of isotope ratios measured  
10 on a mass spectrometer to relative elemental abundances with very high precision. In practical  
11 terms this is achieved through admixing a homogeneous isotopic tracer with a dissolved sample, so  
12 that the accuracy of U-Pb dates are relative to and therefore ultimately traceable back to a tracer  
13 solution. Other isotopic systems that utilize multi-element tracers include Rb-Sr (e.g., Nebel et al.,  
14 2011; Rotenberg et al., 2012), Sm-Nd (e.g., Wasserburg et al., 1981), and Lu-Hf (e.g., Vervoort  
15 et al., 2004).

16 This paper outlines the statistical methodology applied to determining the composition and  
17 associated uncertainties of a mixed ( $^{202}\text{Pb}$ - $^{205}\text{Pb}$ - $^{233}\text{U}$ - $^{235}\text{U}$ ) U-Pb tracer solution. The rare artificial  
18 isotope  $^{202}\text{Pb}$  is included in a limited quantity of this tracer, called ‘ET2535’, and omitted from the  
19 rest, separately denoted ‘ET535’. The methods described here can be applied to high-precision,  
20 high-accuracy calibration of other tracers, and the results can be used to compare and combine  
21 chronological information from multiple chronometers. The U-Pb tracer solution used here was  
22 created, calibrated and distributed to labs dedicated to high-precision U-Pb geochronology under  
23 the auspices of the EARTHTIME initiative. A companion paper to this one (Condon et al., in  
24 review) describes the metrologic traceability and mass spectrometry used to calibrate the tracer  
25 solution, detailing linked experiments that are underpinned by a number of primary determinations.  
26 Here we focus on building statistical models to determine maximum likelihood estimates of the  
27 isotopic composition of mixed-element tracers, using the EARTHTIME U-Pb tracer as an example.

28 The statistical models described here also quantify uncertainty contributions to radioisotopic  
29 dates. For the U-Pb tracer calibration, the isotopic composition of U and Pb isotopic standards that  
30 are utilized in multiple places in the calculations provide the largest contributions and the largest  
31 opportunity for improvement. Many calculated values (for instance, the U/Pb ratio and U and Pb  
32 ICs of the tracer) share this common uncertainty, so that quantifying the statistical correlation  
33 between the different experiments has become paramount in accurately estimating the uncertainty  
34 of U/Pb dates. These systematic uncertainties and others outlined in this paper, when considered  
35 with decay constant uncertainties and corrections for intermediate daughter excess/deficiencies,  
36 provide the present limit on the absolute uncertainties achievable by U-Pb geochronology. With  
37 fully traceable inputs and resulting uncertainties, this tracer calibration also provides a foundation  
38 for accurate comparison of U-Pb geochronology to other radioisotopic or non-radioisotopic dating  
39 methods similarly based on first-principles methods.

#### 40 *1.1. Isotope dilution mass spectrometry*

41 Isotope dilution (ID) isotope ratio mass spectrometry continues to be the most precise and  
42 accurate approach for measuring multi-element isotope ratios, such as the parent/daughter ratios  
43 used for geochronology, (e.g.,  $^{238}\text{U}$  to  $^{206}\text{Pb}$ ,  $^{147}\text{Sm}$  to  $^{143}\text{Nd}$ ). Isotope dilution entails adding a tracer  
44 solution with one or more synthetic or artificially enriched isotopes to a sample. The concentration  
45 of a sample isotope can be estimated using the known amount of the tracer isotope and the measured  
46 sample/tracer isotope ratio. This process can be repeated for other targeted elements in the sample  
47 and tracer, and the ratio of two different elements in the sample (e.g., parent/daughter) can then  
48 be calculated. Uncertainties in the concentrations of parent and daughter elements in the tracer  
49 largely cancel, meaning the uncertainty in the desired isotope ratio or date depends on the accuracy

50 and precision of the parent/daughter element ratio in the tracer. Estimation of this ratio, or tracer  
51 calibration, is carried out by mixing the tracer with gravimetric reference solutions, which are  
52 usually created by dissolving large quantities (to minimize weighing errors) of isotopic or elemental  
53 reference materials that have well-determined purity and isotopic composition (Tilton et al., 1955;  
54 Wetherill, 1956; Wasserburg et al., 1981). Although the reference materials (e.g., NBS 981 for Pb,  
55 CRM 112a for U) are widely used and easily obtained, their isotopic composition and purity have  
56 uncertainties that must be quantified for a complete assessment of a tracer calibration’s accuracy.

57 Another concern for generating accurate and precise data is correcting for intra-element isotopic  
58 fractionation that occurs during mass spectrometric analysis. To correct for short timescale  
59 fluctuations requires using a double spike, composed of two artificially enriched isotopes of the same  
60 element that is being analyzed. There are two commonly used double spike formulations. In the  
61 first, the spike isotopes are also present in the sample, but with different relative abundances (Galer,  
62 1999; Rudge et al., 2009), for instance a  $^{204}\text{Pb}$ - $^{207}\text{Pb}$  tracer. These double spikes require analyzing  
63 both a spiked and an unspiked aliquot of the same sample, limiting their utility for very small  
64 samples. A second type of double spike is composed of two synthetically produced isotopes that  
65 do not naturally occur in the sample. Examples include  $^{202}\text{Pb}$ - $^{205}\text{Pb}$  tracers (Amelin and Davis  
66 (2006) and this work), and the  $^{233}\text{U}$ - $^{236}\text{U}$  tracer (Verbruggen et al., 2008) used to inter-calibrate U  
67 reference materials below. A third type of hybrid double-spike is enriched in a synthetic isotope and  
68 a naturally occurring isotope, for instance the  $^{233}\text{U}$ - $^{235}\text{U}$  tracer described here, which can be used  
69 to monitor instrumental isotopic fractionation when the isotopic composition of the analyte is well  
70 constrained (e.g., Krogh, 1964; Hofmann, 1971).

71 It is common that data generated in different laboratories are determined relative to tracers whose  
72 calibrations are not relatable, in that they may not be calibrated against the same reference materials  
73 or using the same methodology. This makes it necessary to account for tracer calibration uncertainty  
74 when comparing dates determined in different laboratories using different tracers. Though published  
75 geochronologic data often include an estimate of the uncertainty in the tracer calibration, this  
76 uncertainty is often comparable to or exceeds a date’s analytical uncertainty. The tracer calibration  
77 uncertainty therefore limits the resolution of timelines constructed using data from multiple labs.  
78 For this reason, we have mixed and calibrated a ( $^{202}\text{Pb}$ - $^{205}\text{Pb}$ - $^{233}\text{U}$ - $^{235}\text{U}$ ) U-Pb tracer that is being  
79 utilized in a number of laboratories dedicated to high-precision isotope dilution U-Pb geochronology  
80 by thermal ionization mass spectrometry (TIMS). We also provide new traceable and inter-relatable  
81 isotopic compositions and purities for isotopic reference materials and gravimetric solutions derived  
82 from them that together can be used to calibrate other U-Pb tracers, such that their accuracy is  
83 relatable to the EARTHTIME tracer calibration. Widespread distribution and use of these U-Pb  
84 tracers removes interlaboratory bias related to tracer calibration and provides the opportunity to  
85 engage in a well-documented, transparent tracer calibration experiment that is traceable to SI units.  
86 We have attempted to identify all sources of random and systematic uncertainties that should be  
87 considered in any multi-element tracer calibration, and provide an estimate for the total tracer  
88 calibration uncertainty to be included in comparisons of dates between different dating methods.

## 89 *1.2. Application of Inverse Methods*

90 Accurately and precisely relating the isotopic composition of a tracer to first principles measure-  
91 ments benefits from a large quantity of data: over  $10^5$  measured isotope ratios are used to calibrate  
92 the EARTHTIME U-Pb tracer. Each measured isotope ratio contains information about the mixture  
93 of components being analyzed, which include small but unavoidable Pb and U contamination

94 known as laboratory blank, along with the tracer, an isotopic/elemental reference solution, or both.  
95 Unraveling the isotopic compositions (ICs) and relative proportions of each component is made  
96 more difficult by isotopic fractionation, or the preferential evaporation, ionization and/or detection  
97 of lighter isotopes over heavier ones, which changes in magnitude during the course of a typical  
98 TIMS analysis. A measured ratio may therefore be expressed as the amount ratio of its summed  
99 components, modified by a correction factor for isotopic fractionation. The system of equations that  
100 relates each measured ratio to the relative abundances and isotopic compositions of its components  
101 is known as the measurement model, which is derived and described in Appendix B.

102 To determine the relative abundances of the tracer isotopes, the inverse methods of Tarantola  
103 (2005) are applied to a series of linked experiments that represent the complete tracer calibration ex-  
104 periment. Each of these experiments is solved as a separate inverse problem, and a subset of the model  
105 (free) parameters in the first experiments is used to constrain solutions of the following experiments.  
106 For example, the tracer  $^{233}\text{U}/^{235}\text{U}$  is a free model parameter in the U IC experiment in Section 4,  
107 the results of which are used as well-constrained ‘systematic’ uncertainties in Section 6. Combined,  
108 the following experiments represent the complete tracer calibration algorithm, illustrated in Figure 1:

- 109
- 110 Section 2) Inter-calibrate Pb and U ICs of all isotopic reference materials used for tracer cali-  
111 bration such that their covariance is determined;
  - 112 Section 3) Determine the Pb IC of the tracer and the blanks, and their covariance;
  - 113 Section 4) Use critical mixtures of the tracer and multiple U isotopic reference materials to estimate  
114 the U IC of the tracer and its covariance when the natural U IC of a sample is related;
  - 115 Section 5) Determine the U/Pb ratios of the gravimetric solutions using measurements of mass,  
116 purity, and the ICs of their constituent Pb and U reference materials;
  - 117 Section 6) Combine results from all previous calculations to estimate the  $^{235}\text{U}/^{205}\text{Pb}$  and  $^{202}\text{Pb}/^{205}\text{Pb}$   
118 ratio of the tracer using mixtures of the tracer with three gravimetric solutions.

## 119 2. Inter-calibration of Pb and U Isotope Reference Materials

120 In order to determine the U/Pb ratio of the EARTHTIME tracers, each was mixed with  
121 three gravimetric solutions that have U and Pb concentrations determined by weighing and a  
122 gravimetrically traceable isotopic composition. The solutions are composed of three different Pb  
123 reference materials, NBS 981, NBS 982 and ‘Puratronic Pb’, along with two different U reference  
124 materials, CRM 112a and CRM 115. An additional U reference material, U500, was used during  
125 evaluation of the uranium IC of the tracers using critical mixtures; see (Condon et al., in review)  
126 for a more complete description of each of these reference materials. The tracer uranium IC is then  
127 used in the fractionation correction equations for the gravimetric-tracer mixture data reduction  
128 (Fig. 1).

129 Because all six isotopic reference materials are used to determine the tracer U/Pb ratio, their  
130 uncertainties all contribute to the final uncertainty budget of the tracer IC, and any correlation  
131 between the uncertainties in their ICs must be considered when averaging the results from the  
132 three gravimetric solutions. Uncertainty correlations result from relating the ICs of the reference  
133 materials to SI units, accomplished for both U and Pb by measuring the isotope ratios of the  
134 reference materials against a single independent gravimetrically calibrated isotope ratio, a process  
135 known as inter-calibration.

136 *2.1. Pb Isotope Reference Material Inter-calibration*

137 Unlike the well-characterized IRMM-3636(a) U isotopic reference solution (see below), there  
138 is no gravimetric mixture of synthetically produced isotopes against which Pb isotopic reference  
139 materials can be measured. However, the NBS 981, 982, and 983 reference materials have been  
140 calibrated by sample-standard bracketing with mixtures of high-purity  $^{208}\text{Pb}$  and  $^{206}\text{Pb}$  that were  
141 gravimetrically mixed to mimic the  $^{208}\text{Pb}/^{206}\text{Pb}$  of the reference materials (Catanzaro et al., 1968).  
142 As such, these determinations may be considered traceable to SI units.

143 Modern mass spectrometric measurements are significantly more precise than those of Catanzaro  
144 et al., and modern laboratory protocols have significantly lower contamination levels, or laboratory  
145 blanks, to bias the measurements. Because the original high-purity  $^{208}\text{Pb}$  and  $^{206}\text{Pb}$  are no  
146 longer available, the absolute uncertainties of Pb reference materials cannot be improved with new  
147 measurements, but the relative uncertainties between them may be significantly refined. For this  
148 purpose, the ICs of Pb isotopic standards have been revisited multiple times since their original  
149 certification (e.g., Todt et al., 1996; Doucelance and Manhès, 2001; Baker et al., 2004), all of which  
150 base their results on an assumed value of  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 981 or 982 from Catanzaro et al.  
151 (1968). In these studies and in ours, fractionation lines for the standard measurements lie within  $2\sigma$   
152 uncertainties of the certified ICs of NBS 981 and 982.

153 The isotopic composition of NBS 981 is closer to that of modern terrestrial Pb, and therefore  
154 laboratory blank Pb, so that blank corrections for even small samples of NBS 981 result in negligible  
155 changes in blank-corrected IC. We therefore choose to base our calibration on the gravimetrically  
156 traceable Catanzaro et al. (1968) isotopic composition of NBS 981, and assume the value and total  
157 uncertainty of its  $^{208}\text{Pb}/^{206}\text{Pb}$  of  $2.1681 \pm 0.0008$  ( $2\sigma$ ). The reported  $2\sigma$  uncertainty is derived from  
158 a 95% confidence interval calculated with linear, rather than quadratic addition, and is therefore  
159 likely to be a conservative estimate of the true precision of the original measurement. The Pb  
160 isotopic composition of NBS 981, NBS 982 and Puratronic Pb reported in Tables 3 and 4 are  
161 calculated relative to the  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 981, and do not represent absolute measurements of  
162 their Pb isotopic composition.

163 To inter-calibrate the Pb isotopic reference materials used to create the gravimetric solutions,  
164 we used the raw data for measurements of NBS 981, NBS 982, and Puratronic Pb reported in  
165 Amelin and Davis (2006), along with several analyses of the same reference materials from the  
166 same laboratory using the same methods (Fig. 2), with all data compiled and reported in (Condon  
167 et al., in review). Given one of the goals of this intercalibration is to account for mass independent  
168 fractionation, which requires measuring large aliquots of solution, the Amelin and Davis (2006) data  
169 are used instead of separate measurements utilizing ET2535 in order to avoid a large expenditure of  
170 relatively scarce  $^{202}\text{Pb}$ . Smaller loads of the gravimetric solutions with ET(2)535 were thus used  
171 to calibrate the U/Pb ratio of the tracer using accurate Pb reference material ICs derived from  
172 published data.

173 *2.2. U Isotope Reference Material Inter-calibration*

174 To relate the isotopic compositions of the U reference materials to one another, we measured each  
175 against IRMM 3636(a), which was created by weighing highly pure  $^{233}\text{U}$  and  $^{236}\text{U}$  and mixing them  
176 in a 1:1 ratio (Verbruggen et al., 2008). Because the artificial isotopes that comprise IRMM 3636(a)  
177 have been weighed against an in-house kilogram reference, its precisely determined  $^{233}\text{U}/^{236}\text{U}$  is  
178 traceable to the SI system, and ICs that have been measured against it are relatable to the SI

179 through their measurement uncertainties and to one another by tracing each measurement back to  
180 SI units.

181 In this way, the  $^{238}\text{U}/^{235}\text{U}$  ratios and uncertainties for the reference materials SRM U500 and  
182 CRM 112a can be related using the supplementary data from Condon et al. (2010), which reports  
183 the derivative, or linear dependence, of each measured IC with respect to IRMM 3636(a). New data  
184 for the U reference material CRM115 is provided in the supplementary data of this publication, and  
185 is reduced using the same algorithms as Condon et al. (2010).

186 No correlations between the isotope ratio uncertainties are reported on the IRMM 3636 certificate  
187 of analysis, so they are assumed to be uncorrelated. The uncertainty in the IC of IRMM 3636(a)  
188 is treated here as a systematic uncertainty among the reference material analyses. CRM 115 and  
189 CRM U500 are both synthetic reference materials and thus contain  $^{236}\text{U}$ . The  $^{233}\text{U}/^{236}\text{U}$  values for  
190 each were measured for un-spiked aliquots, and then this source of  $^{236}\text{U}$  was subtracted before using  
191 the  $^{233}\text{U}/^{236}\text{U}$  of IRMM 3636(a) to determine the magnitude of isotopic fractionation.

### 192 2.2.1. Algorithm

193 The six equations in the system (B.5) describe the anticipated outcome of measuring a mixture of  
194 Pb isotopic reference material with laboratory blank and a  $^{202}\text{Pb}$ - $^{205}\text{Pb}$  tracer that is undergoing both  
195 mass-dependent and mass-independent fractionation in the presence of a  $\text{BaPO}_2$  isobaric interference.  
196 Several of the variables on the right hand side of equations (B.5), along with their uncertainties,  
197 can be constrained *a priori*. For instance, the IC of the non-enriched tracer components, along  
198 with an average loading blank mass and their uncertainties, can be estimated with the algorithm  
199 in Section 3. The approximate  $\text{BaPO}_2$  IC is calculated from the approximate natural abundances  
200 of its component elements (Böhlke et al., 2005), which are assumed to fractionate by 0.1% per u  
201 during analysis, and are assigned a 2% prior uncertainty. The relative uncertainties in the isotopic  
202 masses of the Pb isotopes are at the ppb level (Audi et al., 2003), and their uncertainties are not  
203 propagated here. Finally, a single gravimetric Pb reference material isotope ratio is required to  
204 calibrate *r25t*, the  $^{202}\text{Pb}/^{205}\text{Pb}$  ratio of the tracer, which can then be used to fractionation-correct  
205 the remaining two gravimetric reference material Pb ICs.

206 Parameters treated as unknowns include the  $^{202}\text{Pb}/^{205}\text{Pb}$  ratio of the tracer, and the Pb ICs of  
207 the gravimetric solutions, excluding the single assumed ratio. Although the gravimetric solution and  
208 tracer masses were weighed prior to mixing, a far more precise estimate of their relative abundance,  
209 represented by the ratio of  $^{206}\text{Pb}$  in the gravimetric solution to the  $^{202}\text{Pb}$  in the tracer, can be  
210 calculated using mass spectrometer measurements, and so this parameter is treated as an unknown.  
211 The three parameters that describe an instantaneous state of mass-independent fractionation,  $\gamma_{205}$ ,  
212  $\gamma_{207}$ , and  $\beta$  (see Appendix B.1), are also treated as unknowns. Thus, multiple measurements of  
213 several Pb reference materials mixed with the same  $^{202}\text{Pb}$ - $^{205}\text{Pb}$  tracer define a system of equations,  
214 and the model in (B.5) relates the measured values to the physical parameters of interest.

### 215 2.2.2. Results

216 The highest-precision data reported in Amelin and Davis (2006) were used for each reference  
217 material: ten NBS 981, nine NBS 982, and four Puratronic Pb analyses. Each is reported as several  
218 block means and standard errors, with isotope ratios relative to  $^{206}\text{Pb}$ , as in equations (B.5), and  
219 the measured ratio uncertainties are assumed to be uncorrelated. To avoid uncorrected isobaric  
220 interferences, which usually occur at the beginning or end of an analysis and significantly displace the  
221 block mean from the trend defined by the majority of the data, all block data for each analysis were

222 plotted in  $^{202}\text{Pb}/^{206}\text{Pb}$  -  $^{204}\text{Pb}/^{206}\text{Pb}$  -  $^{207}\text{Pb}/^{206}\text{Pb}$  -  $^{208}\text{Pb}/^{206}\text{Pb}$  coordinates, and any outliers  
 223 were rejected from further consideration. Plots of the included and excluded block data are shown  
 224 in the electronic supplement.

225 After outlier rejection, there are 160 blocks each of NBS 981 and NBS 982, and 36 blocks of  
 226 Puratronic Pb, each consisting of six measured ratios and uncertainties, for a total of 2136 isotope  
 227 ratio measurements. This is the length of the vector  $\mathbf{d}$  in the model described in Appendix A. These  
 228 data can be used to constrain the 824 model parameters in the vector  $\mathbf{m}$  needed to describe them:  
 229 three Pb ratios for each gravimetric Pb reference material, five Pb ratios for the IC of the tracer used,  
 230 three  $\text{BaPO}_2$  isotope ratios, three Pb blank ratios, unique values to quantify the ratios of gravimetric  
 231 solution and laboratory blank to the isotopic tracer and the mass-independent fractionation of  $^{205}\text{Pb}$   
 232 and  $^{207}\text{Pb}$  for each of the 23 analyses and a unique magnitude of mass-dependent fractionation ( $\beta$ )  
 233 and level of  $\text{BaPO}_2$  interference for each of the 356 included blocks. This defines an over-determined  
 234 system that can be solved by conventional non-linear least squares techniques. We employed an  
 235 iterative method known as preconditioned gradient descent (Tarantola, 2005) to minimize the  
 236 misfit function in equation (A.1), which was executed in MATLAB. Initial values for the model  
 237 parameters were calculated using a linearized form of equations (B.5), and the derivatives required  
 238 were calculated analytically for each iteration. The MATLAB code is provided in the electronic  
 239 supplement.

240 In this model, the assumed  $^{208}\text{Pb}/^{206}\text{Pb}$  ratio of NBS 981, as well as the blank, and  $\text{BaPO}_2$   
 241 ICs and the ratios of  $^{204}\text{Pb}$ ,  $^{206}\text{Pb}$ ,  $^{207}\text{Pb}$ , and  $^{208}\text{Pb}$  to  $^{205}\text{Pb}$  in the tracer act as systematic  
 242 uncertainties, whose values and uncertainties are known *a priori*. The measurement uncertainties  
 243 are calculated by approximating the nonlinear function  $G(\hat{\mathbf{m}})$  with its Jacobian matrix evaluated at  
 244 the solution, denoted  $G$ . The matrix  $G$  has 2136 rows and 824 columns that contain the derivative  
 245 of each of the 2136 predicted values with respect to the 824 model parameters. The measured  
 246 uncertainties are estimated using equation (A.2).

247 In order to estimate the component of uncertainty arising from systematic effects, the system  
 248 was solved for 5000 Monte Carlo realizations of these systematic parameters, created with a  
 249 pseudorandom number generator to have a multivariate normal distribution with a covariance  
 250 structure corresponding to their assumed uncertainties. Thus the system with 2136 measurements  
 251 and 824 model parameters was solved 5000 times, once with each Monte Carlo realization of the  
 252 systematic parameters, which were given extremely small prior uncertainties to ensure that the  
 253 least-squares solution converged to their input values. The uncertainty in the unknown model  
 254 parameters calculated from the inverse model solution using the best estimates of the systematic  
 255 parameters, for instance with the  $^{208}\text{Pb}/^{206}\text{Pb}$  of 981 equal to 2.1681 with infinitesimal uncertainties,  
 256 defines their measured uncertainties.

257 For the model parameters treated as unknowns, the distribution of the resulting 5000 solutions  
 258 defines the probability distribution of the model parameters resulting from the input systematic  
 259 uncertainties. The data plotted in Fig. 4 for the  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 982 confirms that the modeled  
 260 output data is well-approximated by a Gaussian distribution, and the same holds for the other  
 261 isotope ratios of the gravimetric Pb reference materials. The results of all Monte Carlo trials are  
 262 included in the electronic supplement.

263 The systematic uncertainties in the model parameters may therefore be estimated by evaluating  
 264 the mean and covariance matrix of the 5000 estimates of  $\hat{\mathbf{m}}$ , denoted  $\tilde{C}_M^{sys}$ . The total systematic  
 265 and measurement uncertainties, estimated with equations (A.2) and (A.3) are given in Table 3, and  
 266 the correlation coefficients between them, derived from  $\tilde{C}_M^{tot}$ , are provided in Table 4.

267 **3. Determining the Pb IC of the Tracer and Blank**

268 It is not possible to accurately measure the isotopic composition of the tracer and the loading  
 269 blank independently. For the TIMS method, all measurements involve a “loading blank” derived  
 270 from mixing tracer with a silica gel activator, which has a finite Pb blank, with the possibility  
 271 of introducing additional blank by pipetting the combined solution onto the filament and later  
 272 exposing the loaded filament to atmosphere. Mass spectrometry by ICP-MS is similarly affected by  
 273 Pb blank contributions from reagents and carrier gases, as well as interferences from other elements  
 274 such as Hg and Tl. The IC of the Pb blank is also problematic to analyze alone because small  
 275 (e.g. <0.3 pg) blanks produce weak ion beams (e.g. less than ca. 20 cps  $^{204}\text{Pb}$  and 800 cps  $^{208}\text{Pb}$ )  
 276 that limit measurement precision and accuracy. In addition to the average IC of the Pb blank, its  
 277 variability from load to load is an important uncertainty contribution to both the tracer calibration  
 278 exercise and analyses of geologic materials (McLean et al., 2011). The practice of combining several  
 279 Pb blanks into a single analysis averages out and therefore underestimates this blank IC variability.

280 However, since the loading blank and tracer are measured together, a linear regression algorithm  
 281 can be used to estimate each. Loading and analyzing several different masses of tracer using  
 282 approximately the same amount of silica gel establishes a two-component mixing line between the  
 283 blank IC and the tracer IC. When plotted in  $^{204}\text{Pb}/^{205}\text{Pb}$  -  $^{206}\text{Pb}/^{205}\text{Pb}$  -  $^{207}\text{Pb}/^{205}\text{Pb}$  -  $^{208}\text{Pb}/^{205}\text{Pb}$   
 284 coordinates, the tracer IC occupies a unique point on this line and the blank IC is defined by its  
 285 slope. The correct tracer and blank IC together are important for accurately interpreting U-Pb  
 286 data, especially data with lower Pb\*/Pbc ratios (e.g. Rioux et al., 2012).

287 *3.1. Determining the mixing line parameters*

The algorithm presented in McLean (2014) calculates the best fit line through data with correlated  
 uncertainties in two or more dimensions, and is well-suited to a four-dimensional mixing model. In  
 this model, a measured mixture of tracer and blank falls on a mixing line that follows the equation

$$\begin{bmatrix} \left(\frac{^{204}\text{Pb}}{^{205}\text{Pb}}\right)_{mix} \\ \left(\frac{^{206}\text{Pb}}{^{205}\text{Pb}}\right)_{mix} \\ \left(\frac{^{207}\text{Pb}}{^{205}\text{Pb}}\right)_{mix} \\ \left(\frac{^{208}\text{Pb}}{^{205}\text{Pb}}\right)_{mix} \end{bmatrix} = \begin{bmatrix} \left(\frac{^{204}\text{Pb}}{^{205}\text{Pb}}\right)_{tr} \\ \left(\frac{^{206}\text{Pb}}{^{205}\text{Pb}}\right)_{tr} \\ \left(\frac{^{207}\text{Pb}}{^{205}\text{Pb}}\right)_{tr} \\ \left(\frac{^{208}\text{Pb}}{^{205}\text{Pb}}\right)_{tr} \end{bmatrix} + \begin{bmatrix} 1 \\ \left(\frac{^{206}\text{Pb}}{^{204}\text{Pb}}\right)_{bl} \\ \left(\frac{^{207}\text{Pb}}{^{204}\text{Pb}}\right)_{bl} \\ \left(\frac{^{208}\text{Pb}}{^{204}\text{Pb}}\right)_{bl} \end{bmatrix} \cdot \tau_{Pb} \quad (1)$$

288 where *mix*, *tr*, and *bl*, correspond to the mixture, tracer, and blank components, respectively, and  
 289  $\tau_{Pb}$  is the ratio of the moles of  $^{204}\text{Pb}$  contributed from the blank to the moles of  $^{205}\text{Pb}$  from the  
 290 tracer, which varies from analysis to analysis. This is a parametric equation for a line of the form  
 291  $\mathbf{p}_i = \mathbf{a} + \mathbf{v} \tau_i$ , where the vector  $\mathbf{a}$  is a point on the line corresponding to the tracer IC, as above, and  
 292 the vector  $\mathbf{v}$  describes the direction in which the tracer composition is perturbed by the addition  
 293 of loading blank. Each measured mixture IC vector  $\mathbf{p}_i$  is assigned an uncertainty in the form of a  
 294 covariance matrix that incorporates measurement and fractionation-correction effects.



295 Using the measurement uncertainties to weight the line fit assumes that the deviation of each  
296 measurement from the best fit line is due only to the assigned measurement uncertainty and that  
297 the blank has a single, constant IC. In a dataset of measured tracer-blank mixes, the IC of the blank  
298 is expected to vary because it reflects a mixture of Pb from several sources. For this experiment, the  
299 Pb blank combines Pb in the silica gel emitter, Pb obtained from the phosphoric and hydrochloric  
300 acid used to dry down the tracer, Pb on the surface of the beaker used for drying and the inner  
301 surface of the pipette used for loading, as well as any particulate matter from the inside of the  
302 laminar flow bench on which the sample was dried down. Although these often sum to  $<0.3$  pg  
303 of total Pb blank (e.g., Rioux et al., 2012), the ICs of the sources are likely variable, as are their  
304 relative contributions, resulting in a Pb blank with variable IC. This would create scatter from the  
305 mixing line between the tracer IC (assumed constant) and the mean blank IC beyond that expected  
306 from measurement uncertainties alone.

307 A dataset comprising 22 measurements of loading blank-tracer mixtures exhibits overdispersion  
308 with a mean square weighted deviation of 39. To correctly account for the blank IC variability as an  
309 additional source of scatter, an uncertainty term must be added to each point that is proportional  
310 in magnitude to the amount of blank present. Thus, a point on the mixing line close to the IC of  
311 the tracer would be perturbed minimally, while a point farther away would be more sensitive to  
312 variation in the blank IC. The overdispersion will affect the measured  $^{206}\text{Pb}/^{205}\text{Pb}$ ,  $^{207}\text{Pb}/^{205}\text{Pb}$ ,  
313 and  $^{208}\text{Pb}/^{205}\text{Pb}$  relative to the measured  $^{204}\text{Pb}/^{205}\text{Pb}$ , and these effects will be correlated: the  
314 blank IC is expected to be variable, but to generally trend between more and less radiogenic Pb ICs.

315 To calculate the variability in the blank IC from the measured data, a trial tracer IC was first  
316 determined by fitting a line to the measurement data using the measurement uncertainties only. This  
317 trial tracer IC was subtracted from each measured mixture, and the measurement and fractionation  
318 correction uncertainties were propagated to calculate  $^{206}\text{Pb}/^{204}\text{Pb}$ ,  $^{207}\text{Pb}/^{204}\text{Pb}$ , and  $^{208}\text{Pb}/^{204}\text{Pb}$   
319 ratios and covariance matrices for each measurement. The scatter in the resulting estimated blank  
320 ICs cannot be explained by measurement uncertainties alone (Fig. 6).

321 Neglecting the measurement uncertainties and calculating the  $2\sigma$  covariance ellipse for the  
322 discrete ratio data will overestimate the true variability of the tracer, since it does not consider  
323 the extra scatter caused by measurement uncertainties. Alternatively, the scatter in the Pb blank  
324 ICs may result from the sum of two different multivariate normal distributions: the individual  
325 measurement uncertainty unique to each data point and a blank IC variability that affects all  
326 data points. The maximum likelihood estimate of the covariance matrix for this ‘extra scatter’  
327 (Vermeesch, 2010) is shown in green in Figure 6. It is smaller than the discrete data covariance  
328 ellipse, and the lower correlation coefficients demonstrate that part of the high correlation of the  
329 measured data is due to the high correlation of the measured uncertainties.

330 In order to account for the excess scatter from the variable Pb blank IC, the overdispersion  
331 covariance matrix must be added to the uncertainty in each measured data point, multiplied by a  
332 factor proportional to the distance from the point to the tracer IC. This factor increases linearly  
333 with distance along the line from the tracer IC, where it is zero. A new line was fit to the data with  
334 these increased uncertainties, a tracer IC chosen, and the overdispersion calculated; iterating these  
335 steps quickly converges on the tracer IC, and the uncertainties in the line parameters now reflect all  
336 sources of scatter.

337 *3.2. Results*

338 Estimates for the tracer and blank ICs are given in Table 5. Although the tracer IC is constrained  
 339 to be on the mixing line (Fig. 5), there are only two physical constraints on its location. First, the  
 340 isotope ratios that comprise it must not be negative. Second, the proposed IC must have less of  
 341 the common Pb components than the analysis with the highest ratio of tracer to blank. These two  
 342 endpoints define a line segment along which the tracer IC must lie, and it will be shown that the  
 343 location chosen does not influence the value or uncertainty budget of an analysis that has been  
 344 corrected for both tracer and blank. The tracer IC in Table 5 has been arbitrarily chosen to have a  
 345 composition halfway between the two possible endpoints of this line segment, and its  $2\sigma$  uncertainty  
 346 is set to the half-length of the segment (Fig. 5).

347 The line fit algorithm of McLean (2014) outputs uncertainties for a point on the line (the tracer  
 348 IC) and the slope of the line (the blank IC), given in Table 5. The uncertainties in the tracer  
 349 IC ratios are strongly correlated, as are the blank IC ratios, with correlation coefficients given  
 350 in Table 6. The magnitude of the uncertainties in the tracer IC and their correlation define an  
 351 uncertainty envelope around the tracer IC that is parallel to the tracer-blank mixing line.

352 *3.3. Application to TIMS Pb measurements*

353 Although the blank and tracer ICs may be expressed separately, both components must be  
 354 subtracted from routine analyses. Therefore it is their sum, which is constrained to fall along the  
 355 mixing line, that is of interest. There are two approaches to blank subtraction currently employed  
 356 for ID-TIMS analyses, used when analyses contain  $^{204}\text{Pb}$  masses consistent with total procedural  
 357 blank measurements, for example chemically abraded zircon, or contain initial common Pb, for  
 358 example titanite, apatite, or perovskite (Corfu and Dahlgren, 2008; McLean et al., 2011; Schmitz  
 359 and Schoene, 2007).

360 In the first approach, all  $^{205}\text{Pb}$  comes from the tracer and all  $^{204}\text{Pb}$  from the tracer and the blank.  
 361 The measured and fractionation-corrected  $^{204}\text{Pb}/^{205}\text{Pb}$  defines a unique location on the tracer-blank  
 362 mixing line. The corresponding  $^{206}\text{Pb}/^{205}\text{Pb}$ ,  $^{207}\text{Pb}/^{205}\text{Pb}$  and  $^{208}\text{Pb}/^{205}\text{Pb}$  on the line are those of  
 363 the tracer-blank mixture, which are subtracted from the measured, fractionation-corrected ratio  
 364 (McLean et al., 2011). Both the uncertainties in the mixing line parameters and the measured  
 365  $^{204}\text{Pb}/^{205}\text{Pb}$  are propagated into the tracer- and blank-corrected ratios.

In the second approach to common Pb correction, the mass of the Pb blank is assumed based  
 upon some prior knowledge, and the estimated masses of tracer and blank are subtracted together  
 from the fractionation-corrected measured IC of the sample. The rest of the common Pb (including  
 $^{204}\text{Pb}$ ) is considered part of the sample, so the measured  $^{204}\text{Pb}/^{205}\text{Pb}$  is no longer restricted to  
 tracer and blank components. To determine the moles of a given isotope in the sample, for instance  
 $^{206}\text{Pb}$ , first the moles of total procedural blank is determined (equation 15 of McLean et al., 2011),

$$\text{moles}(^{206}\text{Pb})_{tpb} = \text{moles}(^{205}\text{Pb})_{tr} \cdot \left[ \left( \frac{^{206}\text{Pb}}{^{205}\text{Pb}} \right)_{fc} - \left( \frac{^{206}\text{Pb}}{^{205}\text{Pb}} \right)_{tr} \right] \quad (2)$$

and the total procedural blank is subtracted from a subsequent analysis

$$\text{moles}(^{206}\text{Pb})_{spl} = \text{moles}(^{205}\text{Pb})_{spl} \cdot \left[ \left( \frac{^{206}\text{Pb}}{^{205}\text{Pb}} \right)_{fc} - \left( \frac{^{206}\text{Pb}}{^{205}\text{Pb}} \right)_{tr} \right] - \text{moles}(^{206}\text{Pb})_{tpb} \quad (3)$$

366 where *fc* denotes a measured, fractionation-corrected ratio, *tr* the tracer, *spl* the sample, and *tpb*  
 367 the total procedural blank.

However, if the tracer IC was on the measured tracer-blank mixing line, but its location was chosen incorrectly, then its (incorrect) IC could be expressed as

$$\left(\frac{^{206}\text{Pb}}{^{205}\text{Pb}}\right)'_{tr} = \left(\frac{^{206}\text{Pb}}{^{205}\text{Pb}}\right)_{tr} + \left(\frac{^{206}\text{Pb}}{^{204}\text{Pb}}\right)_{bl} \cdot (\Delta\tau) \quad (4)$$

where  $(\Delta\tau)$  is proportional to the incorrect displacement along the tracer-blank mixing line. Calculating equations (2) and (3) with  $(^{206}\text{Pb}/^{205}\text{Pb})_{tr}'$ , the resulting difference between the calculated values of  $moles(^{206}\text{Pb})_{spl}$  is

$$\Delta moles(^{206}\text{Pb})_{spl} = [moles(^{205}\text{Pb})_{spl} - moles(^{205}\text{Pb})_{tpb}] \cdot (\Delta\tau) \cdot \left(\frac{^{206}\text{Pb}}{^{204}\text{Pb}}\right)_{bl} \quad (5)$$

368 According to equation (5), an incorrect choice of the location for the tracer IC on the tracer-blank  
 369 mixing line incurs an error that is proportional to the difference in the moles of  $^{205}\text{Pb}$  added to  
 370 the sample and to the total procedural blank, or for several total procedural blanks, their mean.  
 371 If the same quantity of tracer is used for spiking the total procedural blank measurements and  
 372 the samples, then the first term on the right-hand side of equation (5) approaches zero, and the  
 373 arbitrary choice of the location of the tracer IC on the tracer-blank mixing line will have a minimal  
 374 affect the reduced data.

#### 375 4. A Model for Determining the U ‘Double Spike IC’ and Uncertainty

376 The U isotopic composition of the tracer was determined using the critical mixture approach  
 377 (Krogh, 1964; Hofmann, 1971; Condon et al., in review) using the uranium isotopic reference materials  
 378 CRM U500 and CRM 112a as end-member mixture components. Although the critical mixture  
 379 concept was formulated using a linear fractionation law, it applies to an exponential fractionation  
 380 law as well. The trend predicted by variable-magnitude exponential fractionation, which deviates  
 381 from its linear approximation less than 10ppm over observed isotopic fractionation values, is very  
 382 close to parallel to the mixing line in the vicinity of the critical mixture.

383 In the same way that data from several constraints on the IC of Pb reference materials were  
 384 confederated to calculate the best estimate of several model parameters simultaneously in Section 2.1,  
 385 many measurements of the tracer, measured both on its own and mixed in critical mixture proportions  
 386 can be combined to constrain the U IC of the tracer. The solution and minimization take a similar  
 387 form.

##### 388 4.1. Algorithm and Results

389 To solve for the U IC of ET(2)535, we use multiple critical mixture IC measurements with both  
 390 SRM U500 and CRM 112a as isotopic reference materials, as well as several measurements of pure  
 391 tracer alone. All U loads were estimated to be 500 ng, and U blanks 0.1 pg, which is assigned a  
 392 100% prior relative uncertainty to account for loading blank mass variability. The  $^{238}\text{U}/^{235}\text{U}$  of  
 393 the blank is estimated to be 137.82, an average of terrestrial sources relative to IRMM 3636 (Hiess  
 394 et al., 2012) with no uncertainty assigned because the magnitude of the blank subtraction is nearly  
 395 negligible.

396 As with the Pb reference material inter-calibration, a least squares inverse solution to the large  
 397 system of overdetermined equations in (B.13) created by several measurements is reached by a

398 preconditioned gradient descent method. The unknown variables the U IC of the tracer, as well  
 399 as the sample/spike ratio of each measurement and the magnitude of isotopic fractionation are  
 400 assigned large prior relative uncertainties. This analysis utilizes 14 measurements of SRM U500, 12  
 401 of CRM 112a, and 7 of pure tracer made at NIGL and MIT.

402 The uncertainties in the ICs of the U reference materials used in the critical mixtures are treated  
 403 as systematic uncertainties. To determine their contribution to the total uncertainty in the tracer U  
 404 IC, a Monte Carlo algorithm is employed, which is used to test the assumption that the system is  
 405 locally linear at its least-squares solution. Monte Carlo realizations of the  $^{238}\text{U}/^{235}\text{U}$  of SRM U500  
 406 and CRM 112a are generated by a pseudorandom number generator with the distribution given in  
 407 Tables 1 and 2, and the uncertainty of each realization is assumed to be infinitesimally small. The  
 408 least squares solution is then calculated for  $10^4$  Monte Carlo realizations of the U reference material  
 409 ICs, and their variability represents the systematic uncertainty contribution from the U reference  
 410 material ICs. This estimate is combined with the measurement uncertainties derived by solving the  
 411 system at the maximum likelihood estimate of the U reference material ICs to calculate the total  
 412 uncertainty in the U IC of the tracer (equation A.3).

413 The results of the critical mixtures experiment are presented in Table 8. The null hypothesis that  
 414 the Monte Carlo realizations used to calculate the systematic uncertainty contribution are normally  
 415 distributed is accepted by a K-S test with a p-value of 0.9 for both the  $^{233}\text{U}/^{235}\text{U}$  and  $^{238}\text{U}/^{235}\text{U}$  of  
 416 the tracer. This result permits addition of the measurement and systematic uncertainty covariance  
 417 matrices, the multivariate analog of quadratic uncertainty addition.

#### 418 4.2. Correlation with Sample $^{238}\text{U}/^{235}\text{U}$

419 In an important development for high-precision U-Pb geo- and cosmochronology, the  $^{238}\text{U}/^{235}\text{U}$   
 420 of samples has been found to vary beyond measurement precision (Condon et al., 2010; Hiess et al.,  
 421 2012; Stirling et al., 2007; Weyer et al., 2008). In order to determine the  $^{238}\text{U}/^{235}\text{U}$  of each sample,  
 422 its IC can be measured against IRMM 3636. Thus, in the same way that the U IC of the tracer  
 423 can be traced to the IC of IRMM 3636, so can a precisely determined  $^{238}\text{U}/^{235}\text{U}$  value, and the  
 424 uncertainties of both are therefore correlated. Since uncertainties in both the tracer and sample U  
 425 IC are used to determine the total uncertainty budget for a U-Pb date, this correlation must be  
 426 calculated and included in the uncertainty propagation.

427 To assess their degree of correlation, the derivative of the mean  $^{238}\text{U}/^{235}\text{U}$  for the sample and  
 428 the tracer IC must be evaluated relative to the IC of IRMM 3636. For the specific accessory phases  
 429 measured in Hiess et al. (2012), as well as the reported average zircon  $^{238}\text{U}/^{235}\text{U}$  of  $137.817 \pm 0.045$ ,  
 430 these derivatives are found in the Supporting Online Materials. For the tracer U ratios, the  
 431 derivatives are reported in Table 7, along with the derivatives of the tracer  $^{235}\text{U}/^{205}\text{Pb}$  derived in  
 432 Section 6. The covariance between the tracer and sample ICs can be calculated with a Jacobian  
 433 matrix, or matrix of partial derivatives, and a covariance, using the equation

$$\Sigma_{ts} = J^T \Sigma_{3636} J \quad (6)$$

434 where  $J$  is a Jacobian matrix that contains the derivatives of the tracer and sample IC with respect  
 435 to the IC of IRMM 3636, like that presented in Table 7, and  $\Sigma_{3636}$  is the covariance matrix for  
 436 the IRMM 3636 IC. The resulting tracer-sample covariance matrix  $\Sigma_{ts}$  contains, as its off-diagonal  
 437 components, covariance terms that relate the tracer and sample IC, which can be used in a U-Pb  
 438 uncertainty propagation algorithm such as McLean et al. (2011).

### 4.3. Sample Fractionation Correction with a $^{233}\text{U}$ - $^{235}\text{U}$ Tracer

The true IC of the mixture is known to lie on a mixing line between the ICs of the tracer and sample, both of which are required for this calculation. For a finite fractionation factor  $\beta$ , the measured IC lies off of the mixing line, along a fractionation line from the true IC on the mixing line to the measured IC of the sample. Geometrically, fractionation correction entails finding the point on the mixing line whose fractionation line goes through the measured datum.

However, at the critical mixture IC used above, the fractionation line is parallel to the mixing line. Any error in the measurement or IC of the sample or tracer (even within arbitrarily small uncertainties) will result in a discrepancy between the mixing line and the measured data that cannot be corrected back to the mixing line along the parallel fractionation line. For measured ICs close to the critical mixture IC, fractionation lines are sub-parallel to the mixing line, and small errors (for instance, within measured uncertainties), can result in large, erroneous extrapolation distances back to the mixing line. For this reason,  $^{233}\text{U}$ - $^{235}\text{U}$  tracers are not optimized for ICs near the critical mixture, and the  $^{233}\text{U}/^{235}\text{U}$  of the mixture should be chosen to avoid these sample-tracer ratios. For ET(2)535, uranium ICs near the critical mixture occur for very young under-spiked samples. For example, a 3 Ma sample with a  $^{206}\text{Pb}/^{205}\text{Pb}$  near 4.25 yields measured  $^{238}\text{U}/^{235}\text{U}$  ratios near 55, magnifying small errors in the measurement or estimate of the sample  $^{238}\text{U}/^{235}\text{U}$ . The simplest practical solution is to disregard the double-spike fractionation determination and instead use an average U fractionation value based on past determinations with smaller  $^{238}\text{U}/^{235}\text{U}$  values.

A perceived weakness of the ‘hybrid’  $^{233}\text{U}$ - $^{235}\text{U}$  tracer is that it cannot simultaneously be used to determine the  $^{238}\text{U}/^{235}\text{U}$  of the sample, as it instead relies upon a presumed value for this ratio. For routine U-Pb geochronologic analyses, however, there is not enough uranium available to measure the small  $^{235}\text{U}$  abundance with sufficient precision to detect the epsilon-level natural variation in  $^{238}\text{U}/^{235}\text{U}$ . For instance, the measurements presented in Hiess et al. (2012) comprise weighted means of multiple microgram-mass uranium samples, whereas routine U-Pb geochronology involves samples with a few nanograms or less.

## 5. U/Pb Ratios of the Gravimetric Solutions

In order to determine the U/Pb ratio of each gravimetric solution, three aliquots of high-purity Pb and U reference materials were weighed then dissolved in acid in three independent labs, creating solutions whose U-Pb ratios are gravimetrically calibrated. The uncertainty in the U/Pb ratio of each gravimetric solution is a function of the uncertainties in the masses of the reference materials and their purities. The procedures by which the three gravimetric solutions were mixed in independent laboratories is included in (Condon et al., in review).

The  $^{235}\text{U}/^{205}\text{Pb}$  and  $^{202}\text{Pb}/^{205}\text{Pb}$  ratio of the tracer were determined against three separate gravimetric solutions for several reasons. First, the uncertainties due to the mass measurements of the Pb and U metal used to make the solutions, as well as their purities, are major contributions to the uncertainty in the U/Pb ratio of the gravimetric solutions, and therefore the U/Pb ratio of the tracer. Evaluating the mean tracer U/Pb ratio over multiple independently mixed solutions averages out some of this uncertainty. Also, the different  $^{206}\text{Pb}/^{238}\text{U}$  ratios of the solutions, ranging from 0.094 for the RP solution to 0.017 for the ET solution, allow for varied sample/tracer ratios for each element when mixed with a tracer with a constant  $^{235}\text{U}/^{205}\text{Pb}$ . Using multiple sample/tracer

481 ratios and Pb and U ratios for internal fractionation correction provides an additional check for  
482 internal consistency between results.

### 483 *5.1. Uncertainty in Mass Determinations*

484 The masses of the Pb and U reference materials were determined before the metals were dissolved  
485 to create the gravimetric solutions. Although care was taken to remove oxidation and surface  
486 contamination before weighing on precise balances, each mass measurement has finite uncertainty.  
487 Because the ultimate parameter of interest is the U/Pb ratio of the solutions, any scale bias that is  
488 linearly proportional to the measured mass will divide out. Thus we propagate only the uncertainty  
489 determined from the reproducibility of successive measurements of calibration weights and the Pb  
490 and U metals. Additional corrections were made to account for buoyancy effects, but the magnitude  
491 and systematic nature of these corrections resulted in a negligible contribution to the U/Pb ratio  
492 uncertainty.

### 493 *5.2. Uncertainty in Purity of Pb Isotopic Reference materials*

494 Although the purity, or the assay, of both NBS reference and assay materials and the Puratronic  
495 Pb are certified, these measurements are often dated, contain no supporting information, and are  
496 quoted with conservative uncertainties (e.g. “>99.9%” for NBS 981 and 982). A purity quotation in  
497 this form is unhelpful because there is no expected value or probability density function (pdf) from  
498 which to construct confidence intervals or perform uncertainty analysis. In order to better quantify  
499 the purity of the Pb isotopic reference materials used here, the purities of NBS 981, 982, 983, and  
500 Puratronic Pb were measured by glow discharge mass spectrometry at the GD-MS facility of the  
501 National Research Council (NRC) of Canada. The raw data appears in the electronic supplement of  
502 Condon et al. (in review).

503 Elemental concentrations from GD-MS are reported in two formats. If a significant isobaric  
504 interference exists at the same mass to charge ratios as the element of interest, observed as an  
505 elevated baseline in the mass scan, then its concentration is reported as <X ppb. This result may  
506 reasonably be interpreted as a uniform probability distribution function with limits at zero and X  
507 ppb. If no significant isobaric interference is detected, then the concentration is reported without  
508 the less than symbol, and repeated reference material analyses indicate the concentration of each  
509 element may be represented as with a triangular pdf (Fig. 7). The triangular distribution has a  
510 mean and mode at the stated concentration, and upper and lower limits at  $\pm 50\%$  of the measured  
511 value.

512 The purity of each Pb reference material is defined as the difference between unity and the sum  
513 of all the impurities. Since the pdfs for the impurities are not Gaussian, uncertainty propagation  
514 by quadratic summation is not applicable. Instead, we employ a Monte Carlo approach, where a  
515 randomly generated realization from the pdf of each element’s concentration is summed to produce  
516 a model value for the total impurity concentration. This process was iterated  $10^7$  times for each  
517 reference material, and the resulting distribution of purities, normalized to unity, is an accurate  
518 estimate of the pdf for the purities. The pdf is closely approximated by a histogram of the Monte  
519 Carlo realizations with small bin sizes, illustrated in Figure 7. Because the pdfs of each elemental  
520 concentration are symmetric, the pdf of the total impurities is also symmetric, and the maximum  
521 likelihood estimate and 95% confidence interval for each reference material may be expressed as a  
522 symmetric range about the mean, listed in Table 9.

523 *5.3. Purity of U Isotopic Reference materials*

524 The purities of the U reference materials used to make the gravimetric solutions, CRM 112a and  
 525 CRM 115, have been recently certified to significantly higher precision than the older Pb reference  
 526 materials. In a Sept. 2010 revision of the CRM 112a certificate of analysis by the New Brunswick  
 527 Laboratory, the total impurity concentration is reported as 223  $\mu\text{g/g}$  U, which equates to a purity  
 528 of 0.999777 g U/g metal, and the total uncertainty in the assay is reported as 0.00006 g U/g metal,  
 529 expressed as an approximate 95% confidence interval calculated with a  $k = 2$  coverage factor.  
 530 The coverage factor and confidence interval width together imply that the modeled distribution  
 531 is Gaussian, with a standard deviation of 0.00003 g U/g metal. Likewise, the CRM 115 purity is  
 532 reported (April 2012 certification) as  $0.999770 \pm 0.000046$ , with the same coverage factor and stated  
 533 confidence interval, translating to a standard deviation of 0.000023.

534 *5.4. Gravimetric U/Pb ratio*

535 The U/Pb ratio of the gravimetric solution is most conveniently expressed as its  $^{206}\text{Pb}/^{238}\text{U}$   
 536 ratio, which can be determined from the total moles of each isotope present. The moles of  $^{206}\text{Pb}$  in  
 537 the gravimetric solution is

$$\text{moles}(^{206}\text{Pb}) = \frac{\text{mass}(\text{Pb}_{\text{grav}}) \cdot \text{purity}(\text{Pb}_{\text{grav}})}{M_{204} \cdot \left(\frac{^{204}\text{Pb}}{^{206}\text{Pb}}\right)_{\text{gr}} + M_{206} + M_{207} \cdot \left(\frac{^{207}\text{Pb}}{^{206}\text{Pb}}\right)_{\text{gr}} + M_{208} \cdot \left(\frac{^{208}\text{Pb}}{^{206}\text{Pb}}\right)_{\text{gr}}} \quad (7)$$

538 and likewise the moles of  $^{238}\text{U}$  in the solution is

$$\text{moles}(^{238}\text{U}) = \frac{\text{mass}(\text{U}_{\text{grav}}) \cdot \text{purity}(\text{U}_{\text{grav}})}{M_{234} \cdot \left(\frac{^{234}\text{U}}{^{238}\text{U}}\right)_{\text{gr}} + M_{235} / \left(\frac{^{238}\text{U}}{^{235}\text{U}}\right)_{\text{gr}} + M_{238}} \quad (8)$$

539 The  $^{206}\text{Pb}/^{238}\text{U}$  ratio of the gravimetric solution is simply the quotient of equations (7) and (8).

540 **6. Gravimetric-Tracer Mixtures**

541 In order to determine the U/Pb ratio of the tracer, expressed as the ratio of two of its enriched  
 542 isotopes,  $^{235}\text{U}/^{205}\text{Pb}$ , as well as its  $^{202}\text{Pb}/^{205}\text{Pb}$ , the tracer was mixed with the series of three  
 543 gravimetric solutions described in Section 5.2. Measuring the U/Pb ratio of the tracer against  
 544 the known U/Pb ratio of a gravimetric solution is the inverse of measuring a sample U/Pb ratio  
 545 with the tracer: the known ICs of the gravimetric solution Pb and U components can be used to  
 546 fractionation-correct the measured Pb and U tracer/sample ratios, then the known U/Pb ratio of  
 547 the gravimetric solution is used to determine the U/Pb ratio of the tracer. The small contributions  
 548 of laboratory blank and non-enriched tracer isotopes complicate the calculation, but as with the Pb  
 549 reference material inter-calibration and U critical mixtures, the resulting mixture can be represented  
 550 by a system of non-linear equations, and the best estimate of the tracer parameters reached with a  
 551 non-linear least squares approach.

552 *6.1. Algorithm and Results*

553 A total of 46 paired Pb and U measurements were analyzed at NIGL and MIT: 19 with the RP  
 554 solution, 14 with ET, and 13 with JMM. Of these mixtures, 15 used ET2535 and 31 used ET535.  
 555 The Pb and U for each mix were loaded together on Re filaments, with the Pb run as a metal and U  
 556 as  $\text{UO}_2^+$  polyatomic ion, described in Condon et al. (in review). Between 100 and 600 independent  
 557 isotope ratios were measured for each Pb and U solution, provided in the electronic supplement.  
 558 The mean of each block, defined as 20 consecutive ratio measurements, was evaluated along with  
 559 the multivariate analog of its squared standard error, the covariance matrix for all isotope ratios  
 560 divided by  $n - 1 = 19$  degrees of freedom. A total of 212 blocks of Pb were measured with ET2535,  
 561 579 with ET535, and 644 blocks of U, for a total of  $212 \times 4 + 579 \times 3 + 644 \times 2 = 3873$  ratio means.

562 The most important model parameters in the system described above are the  $^{235}\text{U}/^{205}\text{Pb}$  and  
 563 the  $^{202}\text{Pb}/^{205}\text{Pb}$  of the tracer. In order to solve for these, the following variables must be determined  
 564 for each analysis: the mass-independent fractionation parameters  $\gamma_{205}$  and  $\gamma_{207}$ , and the ratio of  
 565 gravimetric solution to tracer, parameterized by  $^{238}\text{U}_{gr}/^{235}\text{U}_{tr}$ . These variables are assigned initial  
 566 values based on a linearized solution to the model and assigned diffuse priors. Additionally, the  
 567  $^{207}\text{Pb}/^{206}\text{Pb}$  and  $^{208}\text{Pb}/^{206}\text{Pb}$  of the laboratory blank are known to vary between loads, along with  
 568 the mass of the Pb and U blanks themselves.

569 The Pb blank IC for each bead is assigned an IC calculated using data from MIT and NIGL,  
 570 and its the prior uncertainty derived in Section 3. The Pb and U blank mass initial values are  
 571 0.3 and 0.1 pg, respectively, and given a 100% relative uncertainty, which allows the least-squares  
 572 algorithm to determine the blank mass for each load that best fits the data. To ensure that tracer  
 573 IC estimates are not dependent on the choice of Pb blank IC, full calculations were performed  
 574 with a range of ICs measured at MIT and NIGL. The discrepancies were well within our reported  
 575 numerical precision, smaller than the significant digits reported in data tables herein. Finally, an  
 576 initial value for the magnitude of isotopic fractionation for each block,  $\beta_j^i$ , was estimated for each  
 577 Pb and U block using a simple linearized model and assigned a diffuse prior. In total, there are 1759  
 578 unknowns constrained by 3873 isotope ratio measurements, so the system itself is overdetermined.

579 For the gravimetric mixtures, the uncertainties in the gravimetric solution Pb and U ICs, the  
 580 tracer minor isotope IC, and the U/Pb ratios of the gravimetric solutions are all treated as systematic  
 581 uncertainties, since they are not expected to vary from mixture to mixture. To propagate their  
 582 uncertainties, Monte Carlo simulations of each were created using the probability distribution  
 583 functions for the Pb and U ICs of the reference materials, as well as their purities and mass  
 584 determinations. The non-linear least squares model with the data and model parameters described  
 585 above was solved for 2000 Monte Carlo realizations of these systematic effects.

586 Monte Carlo realizations of the gravimetric solution  $^{206}\text{Pb}/^{238}\text{U}$  values include the non-Gaussian  
 587 probability distribution functions for the Pb purities derived from the GD-MS measurements, and  
 588 thus themselves cannot be assumed to be perfectly normally distributed. Testing the distribution of  
 589 the Monte Carlo realizations of the  $^{235}\text{U}/^{205}\text{Pb}$  and the  $^{202}\text{Pb}/^{205}\text{Pb}$  of the tracer, the null hypothesis  
 590 that the distributions are Gaussian is accepted with a p-value of 0.89 and 0.97, respectively. Although  
 591 the distribution of the Pb reference material purities, which are included in this calculation, are not  
 592 Gaussian, they are overwhelmed by other uncertainties, notably the  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 981 and  
 593 the  $^{233}\text{U}/^{236}\text{U}$  of IRMM 3636, both of which are assumed to be normally distributed. Therefore, the  
 594 systematic uncertainties may be combined using equation (A.2) with the measurement uncertainties  
 595 for the  $^{235}\text{U}/^{205}\text{Pb}$  and the  $^{202}\text{Pb}/^{205}\text{Pb}$  using equation (A.3). The results are given in Table 10.



596 **7. Discussion: Impact of the Tracer Calibration model on U-Pb dates**

597 The highest precision dates produced by U-Pb geochronology, used to calibrate the geologic  
 598 timescale from the Paleozoic through the Cenozoic, are  $^{206}\text{Pb}/^{238}\text{U}$  dates. For samples younger  
 599 than about 500 Ma, both  $^{238}\text{U}$  and  $^{206}\text{Pb}$  are more abundant than  $^{235}\text{U}$  and  $^{207}\text{Pb}$ , and thus the  
 600 uncertainty in the  $^{207}\text{Pb}/^{206}\text{Pb}$  date is dominated by the blank correction to the less abundant  
 601 amount of ingrown  $^{207}\text{Pb}$ . Analysis of the total contribution of the tracer calibration uncertainties  
 602 to the uncertainty in a  $^{206}\text{Pb}/^{238}\text{U}$  date is performed with a simplified data reduction scheme.  
 603 The only isotopes considered are  $^{202}\text{Pb}$ ,  $^{205}\text{Pb}$ , and  $^{206}\text{Pb}$  along with  $^{233}\text{U}$ ,  $^{235}\text{U}$ , and  $^{238}\text{U}$ . Using  
 604 the tracer parameters calculated above along with reasonable sample/tracer ratios and a range of  
 605 frequently observed fractionation factors, hypothetical measured ratios corresponding to a range of  
 606 dates were calculated. For instance, a 100 Ma sample with a sample  $^{238}\text{U}/^{235}\text{U}$  ratio of 137.814 and  
 607 a sample/tracer ( $^{238}\text{U}_{spl}/^{235}\text{U}_{tr}$ ) ratio of 1 will have a measured  $^{206}\text{Pb}/^{205}\text{Pb}$  ratio of about 1.56  
 608 and  $^{238}\text{U}/^{235}\text{U}$  ratio of 0.988, assuming typical magnitudes of isotopic fractionation of 0.2% per u  
 609 for Pb and 0.1% per u for U.

610 Using the measured values calculated in this way, a  $^{206}\text{Pb}/^{238}\text{U}$  ratio can be calculated by  
 611 fractionation-correcting the measured  $^{206}\text{Pb}/^{205}\text{Pb}$  and  $^{238}\text{U}/^{235}\text{U}$  ratios using the  $^{202}\text{Pb}/^{205}\text{Pb}$  and  
 612  $^{233}\text{U}/^{235}\text{U}$  ratios, then utilizing the isotope dilution formula

$$\left(\frac{^{206}\text{Pb}}{^{238}\text{U}}\right)_{spl} = \left(\frac{^{206}\text{Pb}}{^{205}\text{Pb}}\right)_{fc} / \left[ \left(\frac{^{238}\text{U}}{^{235}\text{U}}\right)_{fc} \cdot \left(\frac{^{235}\text{U}}{^{205}\text{Pb}}\right)_{tr} \right] \quad (9)$$

613 where *fc* denotes a measured, fractionation-corrected ratio. The  $^{206}\text{Pb}/^{238}\text{U}$  can then be used to  
 614 calculate a date.

615 There are three pertinent tracer parameters whose uncertainty must be considered: the  
 616  $^{202}\text{Pb}/^{205}\text{Pb}$ ,  $^{233}\text{U}/^{235}\text{U}$ , and  $^{235}\text{U}/^{205}\text{Pb}$  ratios. Because both the  $^{235}\text{U}/^{205}\text{Pb}$  and the  $^{202}\text{Pb}/^{205}\text{Pb}$   
 617 derive much of their uncertainty from the *a priori* uncertainty of the  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 981,  
 618 their uncertainties are highly correlated. The result is a relatively large, negative correlation coeffi-  
 619 cient, which indicates that a positive error in the  $^{202}\text{Pb}/^{205}\text{Pb}$  is likely correlated with a negative  
 620 error in the  $^{235}\text{U}/^{205}\text{Pb}$ , and vice versa. The two effects partially cancel one another: a higher  
 621  $^{202}\text{Pb}/^{205}\text{Pb}$  results in a smaller fractionation correction and therefore less apparent  $^{206}\text{Pb}$ , but the  
 622 lower  $^{235}\text{U}/^{205}\text{Pb}$  increases the apparent  $^{206}\text{Pb}/^{238}\text{U}$  (equation 9). Likewise, the  $^{235}\text{U}/^{205}\text{Pb}$  and  
 623  $^{233}\text{U}/^{235}\text{U}$  both depend on the  $^{233}\text{U}/^{236}\text{U}$  of IRMM 3636, and therefore their uncertainties are also  
 624 correlated. In this case, the correlation is positive, and an increase in the tracer  $^{233}\text{U}/^{235}\text{U}$  results  
 625 in a smaller fractionation correction and less apparent  $^{238}\text{U}$ , which is partially offset by the larger  
 626 likely  $^{235}\text{U}/^{205}\text{Pb}$ .

627 In this way, the uncertainty correlations between the  $^{235}\text{U}/^{205}\text{Pb}$  and both the  $^{202}\text{Pb}/^{205}\text{Pb}$   
 628 and  $^{233}\text{U}/^{235}\text{U}$  act to decrease the effective uncertainty in the  $^{206}\text{Pb}/^{238}\text{U}$  date. As Figure 8  
 629 shows, ignoring all covariance terms between the tracer ratios results in an overestimation of the  
 630 tracer uncertainty contribution by a factor of almost two. The only published U-Pb uncertainty  
 631 propagation algorithm to include the required covariance terms is (McLean et al., 2011), which  
 632 permits uncertainty correlations between all tracer parameters. Current version of the ICs and  
 633 uncertainties of both ET535 and ET2535 are available for download into the associated U-Pb\_Redux  
 634 software package through the EARTHTIME initiative.

635 **8. Conclusions**

636 Correct tracer uncertainty propagation is essential to accurate, precise U-Pb isotope dilution  
637 geochronology. This contribution, combined with the data and metrological traceability outlined  
638 in Condon et al. (in review) presents a measurement model that links first-principles mass and  
639 purity measurements to a complete description of the EARTHTIME ( $^{202}\text{Pb}$ - $^{205}\text{Pb}$ - $^{233}\text{U}$ - $^{235}\text{U}$ ) tracer  
640 isotopic composition, using a series of mixtures between the tracer and gravimetric solutions with  
641 known U/Pb ratios and isotopic compositions. The foundation of the tracer calibration depends  
642 upon two sets of measurements: the gravimetrically determined  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 981 and the  
643  $^{233}\text{U}/^{236}\text{U}$  of IRMM 3636, and the weights and purities of the three Pb reference materials and two  
644 U reference materials used to make three independently calibrated gravimetric solutions. Because  
645 the tracer ratios with the strongest influence on U-Pb dates, the  $^{202}\text{Pb}/^{205}\text{Pb}$ ,  $^{233}\text{U}/^{235}\text{U}$ , and  
646  $^{235}\text{U}/^{205}\text{Pb}$  ratios, are mutually dependent on the first-principles measurements, their uncertainties  
647 are significantly correlated. This correlation acts to decrease the overall uncertainty contribution to  
648 U-Pb dates due to tracer calibration.

649 The improvement presented here over the commonly assumed tracer calibration uncertainty of  
650 ca. 0.1% to <300 ppm represents a significant increase in the accuracy and transparency of U/Pb  
651 determinations and resultant calculation of U-Pb dates. Other labs using the same algorithms with  
652 calibration experiment underpinned by appropriate isotopic and assay reference materials should  
653 be able to collaboratively combine U-Pb data at the sub per-mil level, which approaches modern  
654 measurement uncertainties. The converse is also true: a precise tracer calibration is capable of  
655 revealing sub-per-mil variation between collaborating laboratories that may be due to previously  
656 unrecognized instrument calibration or laboratory blank biases. Finally, by establishing U-Pb dates  
657 (i.e.,  $^{206}\text{Pb}/^{238}\text{U}$  and  $^{207}\text{Pb}/^{235}\text{U}$ ) in an absolute reference frame, this tracer calibration combined  
658 with the  $^{235}\text{U}$  and  $^{238}\text{U}$  decay constant data (Jaffey et al., 1971) legitimizes further efforts at  
659 inter-calibrating the U decay constants (e.g. Schoene et al., 2006; Mattinson, 2010; Cheng et al.,  
660 2013) as well as the U-Pb system with other radio-isotopic dating systems through the analyses of  
661 assumed coeval geological materials (e.g. Nebel et al., 2011; Renne et al., 2010) and the accurate  
662 calibration of geologic time through U-Pb geochronology.

663 **Acknowledgements**

664 This research was supported by NSF Award EAR 0451802 (the EARTHTIME project), NERC  
665 Fellowship NE/C517909/1, NERC grant NE/I013814/1, and NIGFSC award (IP/1028/0508). We  
666 thank Ralph Sturgeon for guidance interpreting glow discharge mass spectrometry data, and Yuri  
667 Amelin for kindly sharing Pb standard data.

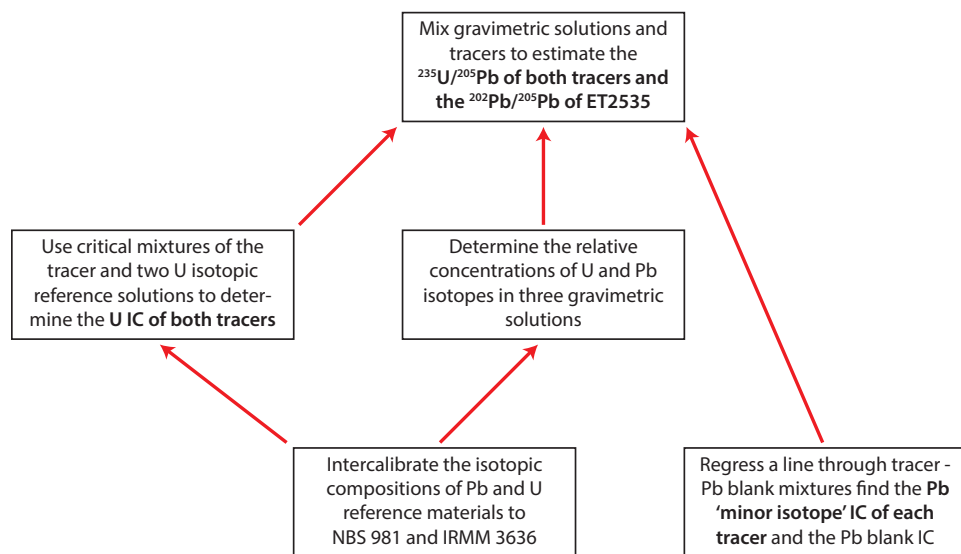


Figure 1: Flow diagram of U-Pb tracer calibration, which consists of five linked experiments that determine the isotopic and elemental composition (Pb IC, U IC, and Pb/U) of mixed U-Pb tracers. The flow proceeds from the bottom to the top of the figure.

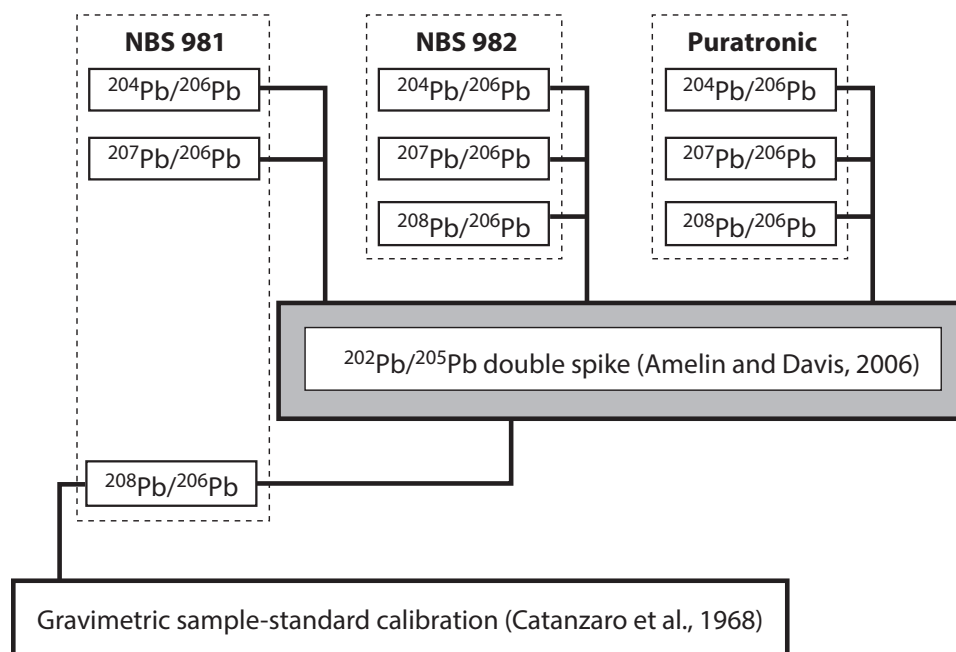


Figure 2: Diagram showing the traceability relationships between the isotopic compositions of the three inter-calibrated Pb reference materials. Note that the  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 981 is assumed to be  $2.1681 \pm 0.0008 (2\sigma)$ , and all other measured isotopic compositions are derived relative to this value.

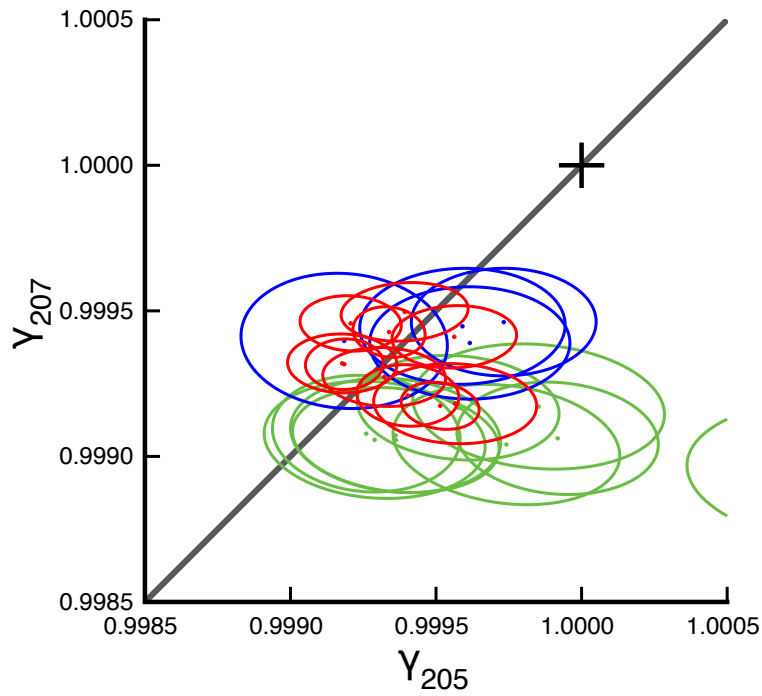


Figure 3: Measurements of the magnitude of MIF for the mixtures of Pb isotopic reference materials and a  $^{202}\text{Pb}$ - $^{205}\text{Pb}$  tracer. The parameters  $\gamma_{205}$  and  $\gamma_{207}$  of McLean (2014) quantify the degree of mass-independent fractionation, empirically modifying the conventional exponential fractionation equation (B.4). Data is from Amelin and Davis (2006) along with newer measurements with the same tracer. The red data is from mixtures of NBS 981, green NBS 982, and blue Puratronic Pb. The black cross on the gray 1:1 line is the predicted behavior using a mass-dependent exponential fractionation law. Ellipses are  $2\sigma$  or  $\sim 86\%$  confidence intervals, with all sources of uncertainty propagated.

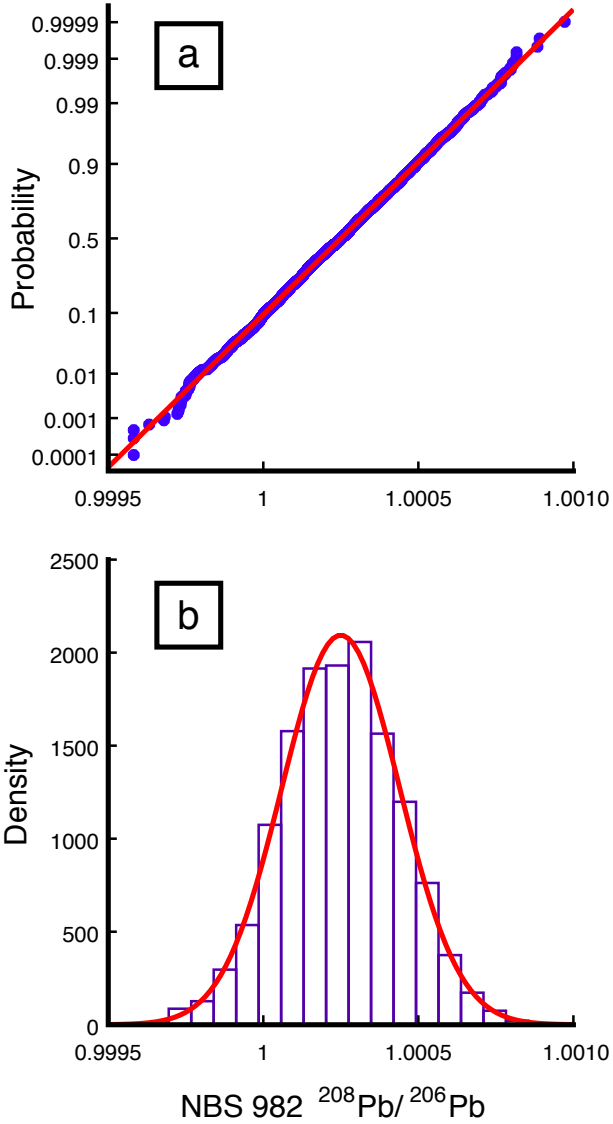


Figure 4: Five thousand Monte Carlo simulations of the effect of systematic uncertainties on the best-fit value of  $^{208}\text{Pb}/^{206}\text{Pb}$  of NBS 982. a) Probability plot of all Monte Carlo solutions. An ideal Gaussian distribution with the mean and standard deviation of the Monte Carlo solutions should plot on the diagonal red line. b) Histogram of Monte Carlo solutions with overlaid Gaussian distribution (red) with the observed mean and standard deviation. Both plots demonstrate that the Monte Carlo solutions are well-approximated by a Gaussian distribution, confirmed by a one-sided K-S test.

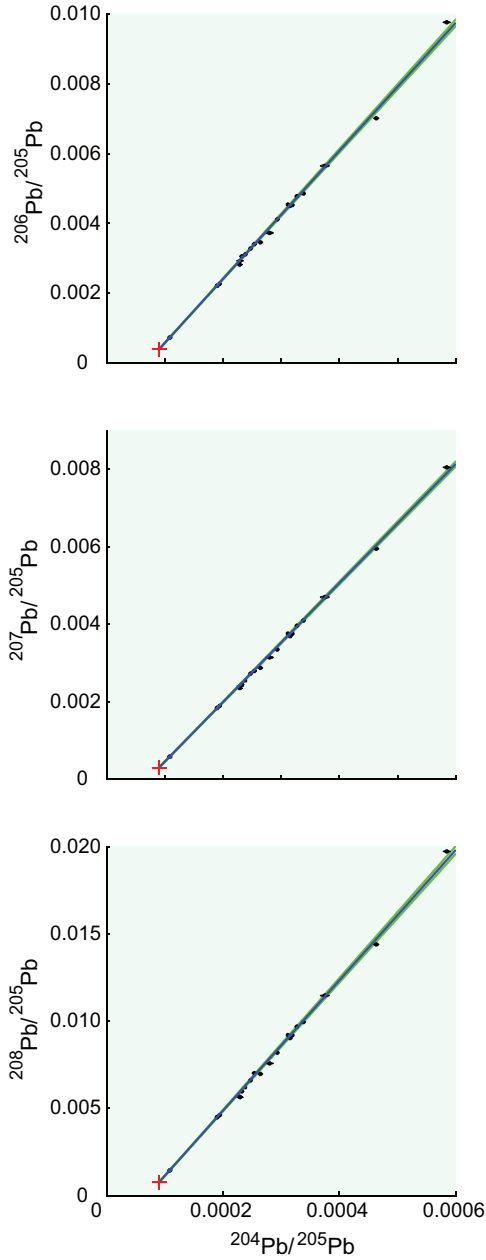


Figure 5: Results of multi-dimensional linear regression through the observed mixing line between ET535 and 22 total procedural Pb blank measurements. The best fit line is shown in blue, with its  $2\sigma$  uncertainty envelope in green, and uncertainties in each data point are generally smaller than their marker size. The isotopic composition of ET535 is constrained to be on this line, contain less common Pb than the cleanest observed analysis, and have positive isotope ratios. Our estimate of the isotopic composition, shown in red, is therefore chosen to be halfway between the cleanest analysis and the location at which the mixing line leaves positive ratio space, with a  $2\sigma$  uncertainty that spans this distance. The slope of the mixing line defines the average isotopic composition of the Pb blank: for instance, in the  $^{206}\text{Pb}/^{205}\text{Pb}$  vs.  $^{204}\text{Pb}/^{205}\text{Pb}$  projection of the four-dimensional line, the slope is the  $^{206}\text{Pb}/^{204}\text{Pb}$  of the Pb blank.

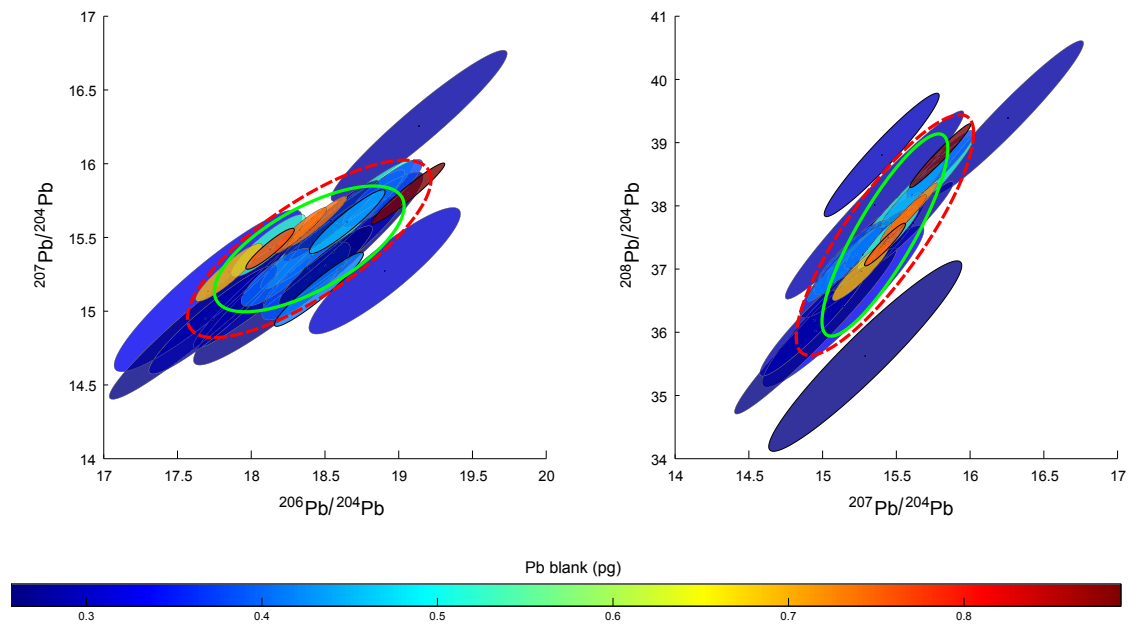


Figure 6: Measurements of the blank isotopic composition after subtracting the final tracer IC. All ellipses are  $2\sigma$ , or  $\sim 86\%$  confidence intervals, including ion counting and fractionation correction uncertainties, and color coded by the blank mass. The covariance ellipse for the discrete dataset of blanks is represented by the large red dashed ellipse, which does not account for the scatter in the blank ICs due to measurement uncertainty. The green ellipse is termed the overdispersion, which separates the positively correlated variability in the blank IC from the even more correlated measurement uncertainties (Vermeesch, 2010).

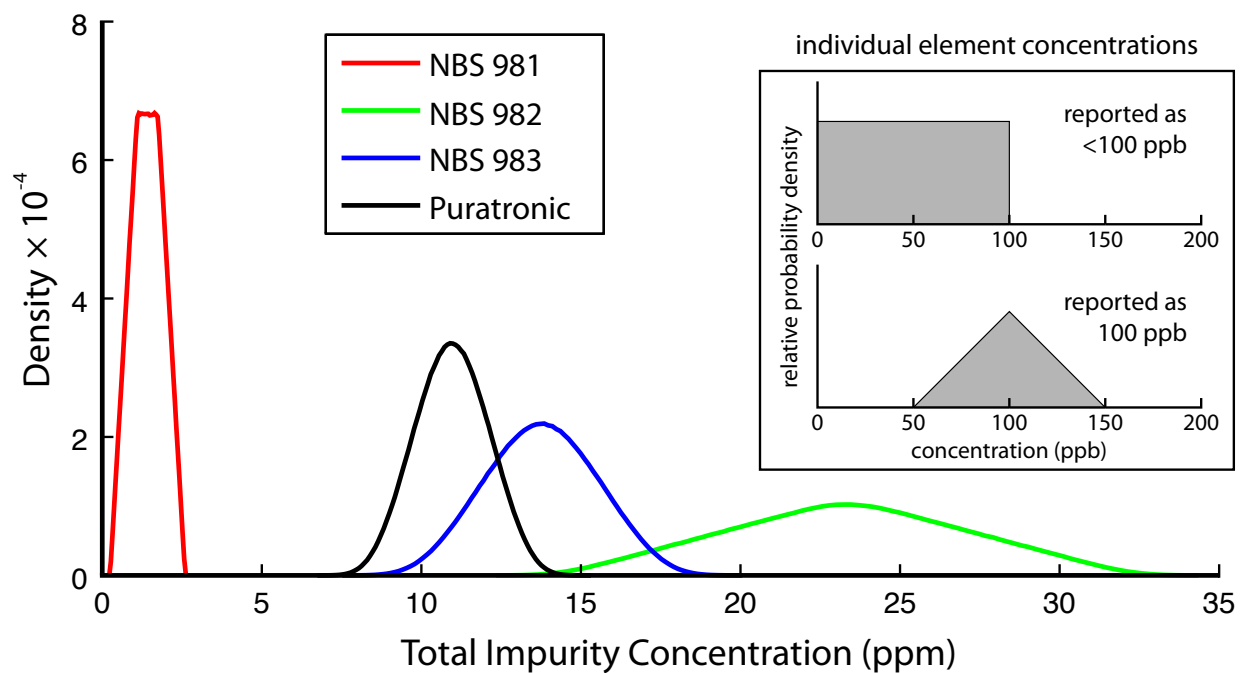


Figure 7: Probability distribution functions for the purity of four commonly used Pb reference materials, derived by summing elemental abundances measured by GD-MS. The inset shows the probability distribution functions assumed for the individual elements measured. In the presence of isobaric interferences, the true concentration is assumed to be between the measured value and zero, with equal relative probability along this interval. For a measurement free of isobaric interference, the true concentration is assumed to be within  $\pm 50\%$  of the measured value, with the measured value being the most probable.



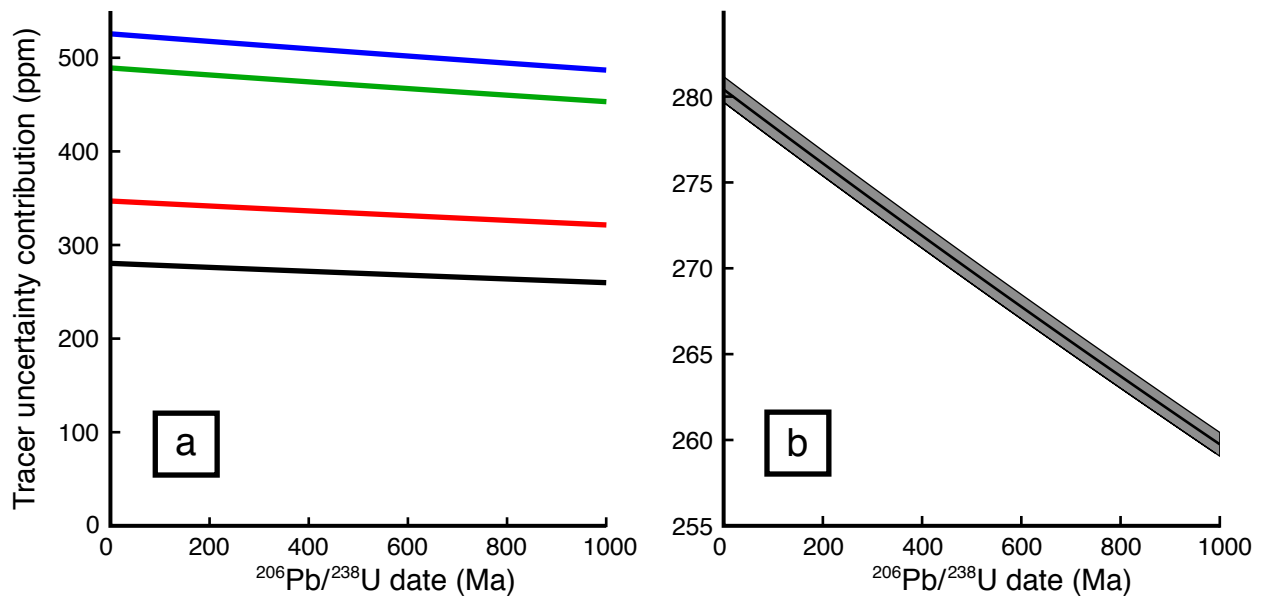


Figure 8: Relative uncertainty contribution ( $2\sigma$ , ppm) to the  $^{206}\text{Pb}/^{238}\text{U}$  date from uncertainty in the tracer IC. a) The black line indicates the correct relative uncertainty contribution as a function of  $^{206}\text{Pb}/^{238}\text{U}$  date. The red, green, and blue lines show the calculated uncertainty contribution if the covariance terms that belong to the tracer  $^{233}\text{U}/^{235}\text{U}$ ,  $^{202}\text{Pb}/^{205}\text{Pb}$ , or all tracer variables, respectively, are neglected. b) The tracer contribution to the  $^{206}\text{Pb}/^{238}\text{U}$  date uncertainty varies with the magnitude of isotopic fractionation. The shaded region encloses commonly observed values: Pb from 0.1 to 0.3% per u, and U from 0.0 to 0.2% per u.

Table 1: Weighted mean  $^{238}\text{U}/^{235}\text{U}$  values and their random, and combined random and systematic uncertainties for the U isotopic standards CRM 112a, CRM 115, and CRM U500 used for tracer calibration. Reported ratios are all fractionation-corrected using the isotopic composition of IRMM 3636(a) (Verbruggen et al., 2008).

	$^{238}\text{U}/^{235}\text{U}$	$\pm 2\sigma^a$	$\pm 2\sigma^b$	MSWD	n (beads)
CRM 112a <sup>c</sup>	137.841	0.011	0.024	1.4	7
CRM 115	491.548	0.039	0.086	0.7	4
CRM U500 <sup>c</sup>	0.999781	0.000077	0.00017	1.0	35

<sup>a</sup> propagating only components of uncertainty arising from random effects during measurement

<sup>b</sup> propagating components of uncertainty arising from systematic and random effects

<sup>c</sup> data from Condon et al. (2010)

Table 2: Matrix of correlation coefficients between the combined random and systematic uncertainties reported in Table 1

	CRM 112a	CRM 115	CRM U500
CRM 112a	1		
CRM 115	0.825	1	
CRM U500	0.914	0.897	1

Table 3: Least squares solutions, random, and combined random and systematic uncertainties for the the Pb isotopic standards NBS 981, NBS 982, and Puratronic Pb used for tracer calibration, calculated with data reported in Amelin and Davis (2006)

		wtd. mean	$\pm 2\sigma^a$	$\pm 2\sigma^b$	n (blocks)
981	$^{204}\text{Pb}/^{206}\text{Pb}$	0.0590074	0.0000016	0.000022	160
	$^{207}\text{Pb}/^{206}\text{Pb}$	0.914683	0.000011	0.00015	
	$^{208}\text{Pb}/^{206}\text{Pb}^c$	<i>2.1681</i>	–	<i>0.0008</i>	
982	$^{204}\text{Pb}/^{206}\text{Pb}$	0.0272058	0.0000021	0.000010	160
	$^{207}\text{Pb}/^{206}\text{Pb}$	0.466967	0.000026	0.00008	
	$^{208}\text{Pb}/^{206}\text{Pb}$	1.000249	0.000056	0.00039	
Pur.	$^{204}\text{Pb}/^{206}\text{Pb}$	0.0548861	0.0000044	0.000021	36
	$^{207}\text{Pb}/^{206}\text{Pb}$	0.856720	0.000046	0.00015	
	$^{208}\text{Pb}/^{206}\text{Pb}$	2.10227	0.00011	0.00079	

<sup>a</sup> Propagating only components of uncertainty arising from random effects during measurement

<sup>b</sup> Propagating components of uncertainty arising from systematic and random effects

<sup>c</sup> Value from Catanzaro et al. (1968). All other isotope ratios are calculated relative to this value.

Table 4: Matrix of correlation coefficients between the combined random and systematic uncertainties in the Pb ICs used for tracer calibration. Correlation coefficients close to -1 or 1 indicate a high degree of correlation. The shared sources of systematic uncertainty largely responsible for the high correlation coefficients are the ratios being used for the fractionation correction.

		NBS 981			NBS 982			Puratronic Pb		
		$\frac{^{204}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{207}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{208}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{204}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{207}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{208}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{204}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{207}\text{Pb}}{^{206}\text{Pb}}$	$\frac{^{208}\text{Pb}}{^{206}\text{Pb}}$
981	$^{204}\text{Pb}/^{206}\text{Pb}$	1								
	$^{207}\text{Pb}/^{206}\text{Pb}$	-0.974	1							
	$^{208}\text{Pb}/^{206}\text{Pb}$	-0.990	0.991	1						
982	$^{204}\text{Pb}/^{206}\text{Pb}$	0.939	-0.960	-0.958	1					
	$^{207}\text{Pb}/^{206}\text{Pb}$	-0.875	0.809	0.846	-0.753	1				
	$^{208}\text{Pb}/^{206}\text{Pb}$	-0.965	0.919	0.950	-0.883	0.943	1			
Pur.	$^{204}\text{Pb}/^{206}\text{Pb}$	0.974	0.958	-0.973	0.933	-0.848	-0.942	1		
	$^{207}\text{Pb}/^{206}\text{Pb}$	-0.941	0.939	0.948	-0.909	0.815	0.909	-0.948	1	
	$^{208}\text{Pb}/^{206}\text{Pb}$	-0.982	0.975	0.988	-0.947	0.850	0.950	-0.986	0.961	1

Table 5: Results of linear fit for tracer - blank mixing line

	ET535		ET2535		Pb blank	
	value	$\pm 2\sigma$	value	$\pm 2\sigma$	value	$\pm 2\sigma$
$^{204}\text{Pb}/^{205}\text{Pb}^a$	0.000090	0.000018	0.000130	0.000050	$^{206}\text{Pb}/^{204}\text{Pb}$	18.41 0.48
$^{206}\text{Pb}/^{205}\text{Pb}$	0.00039	0.00034	0.00093	0.00092	$^{207}\text{Pb}/^{204}\text{Pb}$	15.41 0.29
$^{207}\text{Pb}/^{205}\text{Pb}$	0.00030	0.00028	0.00077	0.00077	$^{208}\text{Pb}/^{204}\text{Pb}$	37.61 1.13
$^{208}\text{Pb}/^{205}\text{Pb}$	0.00074	0.00070	0.0019	0.0019		

<sup>a</sup> This value is chosen arbitrarily to be half the distance from a null composition to the analysis with the highest ratio of tracer to blank (see Section 3.2).

Table 6: Correlation coefficient matrix for ET535 and loading blank Pb IC.

		ET535 Pb IC				blank Pb IC		
		$\frac{^{204}\text{Pb}}{^{205}\text{Pb}}$	$\frac{^{206}\text{Pb}}{^{205}\text{Pb}}$	$\frac{^{207}\text{Pb}}{^{205}\text{Pb}}$	$\frac{^{208}\text{Pb}}{^{205}\text{Pb}}$	$\frac{^{206}\text{Pb}}{^{204}\text{Pb}}$	$\frac{^{207}\text{Pb}}{^{204}\text{Pb}}$	$\frac{^{208}\text{Pb}}{^{204}\text{Pb}}$
ET535	$^{204}\text{Pb}/^{205}\text{Pb}$	1						
	$^{206}\text{Pb}/^{205}\text{Pb}$	0.980	1					
	$^{207}\text{Pb}/^{205}\text{Pb}$	0.989	0.992	1				
	$^{208}\text{Pb}/^{205}\text{Pb}$	0.974	0.987	0.992	1			
blank	$^{206}\text{Pb}/^{204}\text{Pb}$	0	-0.136	-0.076	-0.114	1		
	$^{207}\text{Pb}/^{204}\text{Pb}$	0	-0.103	-0.100	-0.135	0.755	1	
	$^{208}\text{Pb}/^{204}\text{Pb}$	0	-0.099	-0.086	-0.156	0.729	0.864	1

Table 7: Derivatives of the U IC of ET(2)535 with respect to IRMM 3636, used to determine the covariance between a measured  $^{238}\text{U}/^{235}\text{U}$  and the tracer, for propagating uncertainty in U-Pb dates.

		ET(2)535		
		$\frac{^{233}\text{U}}{^{235}\text{U}}$	$\frac{^{238}\text{U}}{^{235}\text{U}}$	$\frac{^{235}\text{U}}{^{205}\text{Pb}}$
IRMM3636	$^{233}\text{U}/^{236}\text{U}$	0.6707	0.002907	99.79
	$^{234}\text{U}/^{236}\text{U}$	-0.001262	$4.209 \times 10^{-6}$	-0.1951
	$^{235}\text{U}/^{236}\text{U}$	-0.8686	0.005051	-121.8
	$^{238}\text{U}/^{236}\text{U}$	0.4466	-0.001505	68.93

Table 8: The U isotopic composition of ET(2)535 from the critical mixture experiment. The tracer was mixed with CRM 112a and SRM U500, whose ICs, uncertainties, and correlation coefficient are given in Section 2.2.

	MLE	$\pm 2\sigma^a$	$\pm 2\sigma^b$	$\rho^c$
$^{233}\text{U}/^{235}\text{U}$	0.995062	0.000009	0.00011	-0.599
$^{238}\text{U}/^{235}\text{U}$	0.00307993	0.00000064	0.00000080	

<sup>a</sup> propagating only components of uncertainty arising from random effects during measurement

<sup>b</sup> propagating components of uncertainty arising from systematic and random effects

<sup>c</sup> correlation coefficient between  $^{233}\text{U}/^{235}\text{U}$  and  $^{238}\text{U}/^{235}\text{U}$ , with all uncertainties propagated

Table 9: Purities of Pb isotopic standards measured by glow discharge mass spectrometry, with estimated symmetric 95% confidence intervals

	purity	95%CI
NBS 981	0.9999986	$\pm 0.0000009$
NBS 982	0.9999767	$\pm 0.0000072$
NBS 983	0.9999862	$\pm 0.0000033$
Puratronic	0.9999890	$\pm 0.0000047$

Table 10: Results of the gravimetric solution - tracer mixtures.

	MLE	$\pm 2\sigma^a$	$\pm 2\sigma^b$	$\rho^c$
$^{202}\text{Pb}/^{205}\text{Pb}$	0.999239	0.000019	0.00053	-0.915
$^{235}\text{U}/^{205}\text{Pb}$	100.2329	0.0022	0.047	

<sup>a</sup> propagating only components of uncertainty arising from random effects during measurement

<sup>b</sup> propagating components of uncertainty arising from systematic and random effects

<sup>c</sup> correlation coefficient between  $^{202}\text{Pb}/^{205}\text{Pb}$  and  $^{235}\text{U}/^{205}\text{Pb}$ , with all uncertainties propagated

668 **Appendix A. Application of inverse methods to tracer calibration**

669 The parameters in the measurement model, such as the ICs and relative proportions of the  
 670 tracer, isotopic reference materials, and blank, as well as the magnitude of isotopic fractionation,  
 671 fall into three categories, depending on our state of knowledge at the beginning of the experiment.  
 672 1) Parameters measured by first principles, which underpin the tracer calibration. One example  
 673 is the  $^{233}\text{U}/^{236}\text{U}$  ratio of the U isotopic standard IRMM3636, whose value and uncertainty are  
 674 traceable to first-principles measurements of mass (Verbruggen et al., 2008). Uncertainties in these  
 675 parameters are not improved by the tracer calibration measurements, and can therefore be considered  
 676 ‘systematic.’ 2) Parameters estimated from prior experience, such as the mass of the laboratory Pb  
 677 blank in an analysis, which may be further constrained by the isotope ratio measurements. The  
 678 values of these parameters are allowed to change within the available *a priori* constraints. 3) Free  
 679 model parameters, such as the IC of the tracer, which have essentially no *a priori* information.

680 When formulated as an inverse problem (e.g. Tarantola and Valette, 1982a,b; Tarantola, 2005),  
 681 the model may be expressed as  $\mathbf{d} = \mathbf{G}(\mathbf{m})$ , where  $\mathbf{d}$  is a vector of data, in this case isotope ratio  
 682 measurements, and  $\mathbf{G}(\mathbf{m})$  is the function of the model parameters  $\mathbf{m}$  that explains the measured  
 683 data. A simple example would be fitting a line ( $y = ax + b$ ) to  $n$  measurements of a parameter ( $y$ )  
 684 made at well-known positions ( $x$ ) assumed to have zero uncertainty. In this case, the data vector  
 685  $\mathbf{d}$  is a column vector containing the  $n$  measured  $y$ -values  $[y_1, y_2, \dots, y_n]$ , and the model parameter  
 686 vector  $\mathbf{m}$  would have only two elements,  $[a, b]$ . The measurement model is the system of equations  
 687  $y_i = ax_i + b$  that relates  $\mathbf{d}$  to  $\mathbf{m}$ , so that  $\mathbf{G}(\mathbf{m})$  is the linear function  $\mathbf{G}([a, b]) = ax_i + b$ .

688 *Appendix A.1. Solving the Inverse Problem*

689 For the linked tracer calibration experiments, the  $\mathbf{G}(\mathbf{m})$  functions found in equations (B.5),  
 690 (B.13), and (B.8) are nonlinear, and because there are more measured isotope ratios in  $\mathbf{d}$  than  
 691 parameters to solve for in  $\mathbf{m}$ , the measurement model is overdetermined. There is no unique solution  
 692 for the model parameters  $\mathbf{m}$ , but the best choice of the parameter values  $\hat{\mathbf{m}}$  minimizes the misfit  
 693 between the observed values of the isotope ratios ( $\mathbf{d}$ ) on the left hand side of  $\mathbf{d} = \mathbf{G}(\hat{\mathbf{m}})$  and  
 694 the values predicted by best fit model parameters  $\hat{\mathbf{m}}$  on the right-hand side. This ‘misfit’ is the  
 695 weighted sum of two differences, the first between the data and the model predictions,  $\mathbf{d} - \mathbf{G}(\hat{\mathbf{m}})$ ,  
 696 which should fall within the uncertainties of the measured data, and the second between any prior  
 697 constraints on the model parameters and the best fit values,  $\mathbf{m}_{\text{prior}} - \hat{\mathbf{m}}$ . These differences are  
 698 weighted by the measured uncertainties for the data, embodied in the covariance matrix  $\mathbf{C}_D$ , and the  
 699 *a priori* uncertainties in the model parameters, embodied in the covariance matrix  $\mathbf{C}_M$ . Free model  
 700 parameters such as the isotopic composition of the tracer, considered unknowns, are given initial  
 701 estimates with diffuse priors, or large initial uncertainties in  $\mathbf{C}_M$  that ensure the initial estimate  
 702 only negligibly affects the outcome.

703 The numerical value of the misfit is given by the objective function, whose minimum is the  
 704 solution to the inverse problem (Tarantola, 2005),

$$S = (\mathbf{d} - \mathbf{G}(\hat{\mathbf{m}}))^T \mathbf{C}_D^{-1} (\mathbf{d} - \mathbf{G}(\hat{\mathbf{m}})) + (\mathbf{m}_{\text{prior}} - \hat{\mathbf{m}})^T \mathbf{C}_M^{-1} (\mathbf{m}_{\text{prior}} - \hat{\mathbf{m}}) \quad (\text{A.1})$$

705 where  $\mathbf{C}_D$  and  $\mathbf{C}_M$  are the covariance matrices of the measured data and the prior model parameter  
 706 estimates, respectively. To minimize  $S$ , a stable linear preconditioned steepest descent method is  
 707 employed (Tarantola, 2005). The algorithm calculates the gradient, or multi-dimensional direction  
 708 in which the value of  $S$  is most rapidly decreasing, then uses an estimate of the curvature of  $S$

709 to determine a trial minimum value. At the trial minimum, a new gradient and curvature are  
 710 estimated, and the procedure is iterated until convergence. The result is the maximum likelihood  
 711 estimate of the model parameter values.

712 To evaluate the measurement uncertainties, or the joint conditional density function of the model  
 713 parameters treated as unknowns given the best estimates of systematic variables, the overdetermined  
 714 inverse problem was solved using near-zero prior uncertainties for the systematic variables. The  
 715 measurement uncertainties are then estimated by approximating the system of equations with its  
 716 Jacobian matrix evaluated at the best fit solution, denoted  $G$ . The matrix  $G$  has a row for each  
 717 measured isotope ratio and a column for each model parameter, and contains the derivative of each  
 718 of the predicted measured isotope ratios with respect to each of the model parameters, so that  
 719  $G(i, j) = (\partial \mathbf{d}_i / \partial \mathbf{m}_j)$ . The posterior covariance matrix, which contains the measured uncertainties,  
 720 is estimated using

$$\tilde{C}_M^{meas} = C_M - C_M G^T (G C_M G^T + C_D)^{-1} G C_M \quad (\text{A.2})$$

721 which can be derived from equation (A.1), the objective function (Tarantola, 2005).

## 722 *Appendix A.2. Evaluating Systematic Uncertainties*

723 In order to evaluate the component of uncertainty arising from systematic effects, the entire  
 724 nonlinear inverse problem can be solved for  $M$  Monte Carlo realizations of the systematic parameters,  
 725 created with a pseudorandom number generator to have the desired probability density function,  
 726 here a multivariate Gaussian. The value of  $M$  used for the inverse problems described here ranges  
 727 from  $10^4$  to  $10^7$ , depending on the computational difficulty of the calculation. The systematic  
 728 parameters, such as the assumed  $^{208}\text{Pb}/^{206}\text{Pb}$  ratio of NBS 981, can either be given infinitesimally  
 729 small prior uncertainties to ensure that the model converges to the input value, or can be omitted  
 730 from  $\mathbf{m}$  entirely.

731 For the free model parameters treated as unknowns, the distribution of the  $M$  resulting solutions  
 732 defines the probability distribution of the model parameters resulting from the input systematic  
 733 uncertainties. Because the model is nonlinear, even if the uncertainties in the measured data and  
 734 systematic variables all have Gaussian distributions, the calculated uncertainties in the other output  
 735 model parameters, such as the  $^{235}\text{U}/^{205}\text{Pb}$  of the tracer, may be significantly non-Gaussian. The  
 736 departure from the normal distribution depends on both the degree of non-linearity of the system at  
 737 the value of the solution and the size of the input uncertainties. Solving the least squares system for  
 738 many Monte Carlo realizations of the systematic variables provides a way to evaluate the probability  
 739 distribution of the output model parameters without assuming that the model is locally linear. The  
 740 normality of the Monte Carlo-modeled solutions can be checked by plotting the data as a histogram  
 741 or a Q-Q plot, or with a Kolmogorov-Smirnov (K-S) test, which compares the observed Monte-Carlo  
 742 distribution with the theoretically predicted normal distribution with the same mean and standard  
 743 deviation.

744 If the probability distributions of the free model parameters from the  $M$  Monte Carlo solutions  
 745 are confirmed to be normally distributed, they can be estimated by evaluating the mean and  
 746 covariance matrix of the  $M$  estimates of  $\hat{\mathbf{m}}$ , denoted  $\tilde{C}_M^{sys}$ . The total systematic and measurement  
 747 uncertainty can then be expressed as the sum of the measured and systematic covariance matrices:

$$\tilde{C}_M^{tot} = \tilde{C}_M^{meas} + \tilde{C}_M^{sys} \quad (\text{A.3})$$

748 **Appendix B. Measurement models**

749 *Appendix B.1. Pb Isotope Standards Inter-calibration Model*

To inter-calibrate the Pb isotopic reference materials against a  $^{202}\text{Pb}/^{205}\text{Pb}$  tracer, while monitoring the effect of a  $\text{BaPO}_2$  isobaric interference, six masses must be measured. The contributions to each measured mass may be broken down as follows,

$$\begin{aligned}
 201_{tot} &= {}^{201}\text{BaPO}_2 \\
 202_{tot} &= {}^{202}\text{Pb}_{tr} + {}^{202}\text{BaPO}_2 \\
 204_{tot} &= {}^{204}\text{Pb}_{gr} + {}^{204}\text{Pb}_{bl} + {}^{204}\text{Pb}_{tr} + {}^{204}\text{BaPO}_2 \\
 205_{tot} &= {}^{205}\text{Pb}_{tr} + {}^{205}\text{BaPO}_2 \\
 206_{tot} &= {}^{206}\text{Pb}_{gr} + {}^{206}\text{Pb}_{bl} + {}^{206}\text{Pb}_{tr} \\
 207_{tot} &= {}^{207}\text{Pb}_{gr} + {}^{207}\text{Pb}_{bl} + {}^{207}\text{Pb}_{tr} \\
 208_{tot} &= {}^{208}\text{Pb}_{gr} + {}^{208}\text{Pb}_{bl} + {}^{208}\text{Pb}_{tr}
 \end{aligned} \tag{B.1}$$

750 where *gr* denotes a contribution from the Pb reference material used in the gravimetric U-Pb  
 751 solution, *tr* from the tracer, and *bl* from the laboratory blank.

752 Assuming the signal at mass 201 is entirely  $^{201}\text{BaPO}_2$ , it can be used to subtract the iso-  
 753 baric interferences from underneath masses 202, 204, and 205. The ratios  $^{202}\text{BaPO}_2/^{201}\text{BaPO}_2$ ,  
 754  $^{204}\text{BaPO}_2/^{201}\text{BaPO}_2$ , and  $^{205}\text{BaPO}_2/^{201}\text{BaPO}_2$ , calculated by considering all permutations of the  
 755 constituent isotopes weighted by their relative abundance in nature (Böhlke et al., 2005), are calcu-  
 756 lated to be  $1.4055 \times 10^{-3}$ ,  $2.2259 \times 10^{-6}$ , and  $4.2206 \times 10^{-6}$ , respectively. Expressing, for instance,  
 757 the  $^{202}\text{BaPO}_2$  contribution as the signal at mass 201 multiplied by the  $^{202}\text{BaPO}_2/^{201}\text{BaPO}_2$ , the  
 758  $\text{BaPO}_2$  contributions may be subtracted from each measured mass.

The moles of each isotope of Pb can be normalized to  $^{202}\text{Pb}$ , which occurs only in the tracer. Likewise, the  $\text{BaPO}_2$  can be normalized to the polyatomic ion with molecular weight 205.

$$\begin{aligned}
 201_{tot}/^{202}\text{Pb}_{tr} &= {}^{201}\text{BaPO}_2/^{202}\text{Pb}_{tr} \\
 202_{tot}/^{202}\text{Pb}_{tr} &= 1 + {}^{202}\text{BaPO}_2/^{202}\text{Pb}_{tr} \\
 204_{tot}/^{202}\text{Pb}_{tr} &= {}^{204}\text{Pb}_{gr}/^{202}\text{Pb}_{tr} + {}^{204}\text{Pb}_{bl}/^{202}\text{Pb}_{tr} + {}^{204}\text{Pb}_{tr}/^{202}\text{Pb}_{tr} + {}^{204}\text{BaPO}_2/^{202}\text{Pb}_{tr} \\
 205_{tot}/^{202}\text{Pb}_{tr} &= {}^{205}\text{Pb}_{tr}/^{202}\text{Pb}_{tr} + {}^{205}\text{BaPO}_2/^{202}\text{Pb}_{tr} \\
 206_{tot}/^{202}\text{Pb}_{tr} &= {}^{206}\text{Pb}_{gr}/^{202}\text{Pb}_{tr} + {}^{206}\text{Pb}_{bl}/^{202}\text{Pb}_{tr} + {}^{206}\text{Pb}_{tr}/^{202}\text{Pb}_{tr} \\
 207_{tot}/^{202}\text{Pb}_{tr} &= {}^{207}\text{Pb}_{gr}/^{202}\text{Pb}_{tr} + {}^{207}\text{Pb}_{bl}/^{202}\text{Pb}_{tr} + {}^{207}\text{Pb}_{tr}/^{202}\text{Pb}_{tr} \\
 208_{tot}/^{202}\text{Pb}_{tr} &= {}^{208}\text{Pb}_{gr}/^{202}\text{Pb}_{tr} + {}^{208}\text{Pb}_{bl}/^{202}\text{Pb}_{tr} + {}^{208}\text{Pb}_{tr}/^{202}\text{Pb}_{tr}
 \end{aligned} \tag{B.2}$$

The molar Pb ratios may now be recast in terms of the ratio of gravimetric solution and laboratory blank Pb to that in the tracer and the ICs of the three components. Likewise, the  $\text{BaPO}_2$  contribution to each measured mass may be recast in terms of the ratio of  $\text{BaPO}_2$  to tracer

and the BaPO<sub>2</sub> IC. This is accomplished by defining  $r62gt$  and  $r62bt$  as the ratio of the <sup>206</sup>Pb contribution from the gravimetric solution and the laboratory blank to the <sup>202</sup>Pb contribution from the tracer, respectively, and  $r52BaPb$  as the ratio of the <sup>205</sup>BaPO<sub>2</sub> to the <sup>202</sup>Pb contribution from the tracer. The tracer IC is then expressed as the isotope ratios  $r25t$ ,  $r42t$ ,  $r62t$ , and  $r72t$ , representing  $(^{202}\text{Pb}/^{205}\text{Pb})_{tr}$ ,  $(^{204}\text{Pb}/^{202}\text{Pb})_{tr}$ , and so on, and likewise for  $r46b$ ,  $r76b$ ,  $r86b$  for the blank IC;  $r46g$ ,  $r76g$ ,  $r86g$  for the gravimetric Pb reference material IC; and  $r15Ba$ ,  $r25Ba$ , and  $r45Ba$  for the ratios of BaPO<sub>2</sub> components. The measured signal at each mass can then be written as

$$\begin{aligned}
201_{tot}/^{202}\text{Pb}_{tr} &= r15Ba \cdot r52BaPb \\
202_{tot}/^{202}\text{Pb}_{tr} &= 1 + r25Ba \cdot r52BaPb \\
204_{tot}/^{202}\text{Pb}_{tr} &= r46g \cdot r62gt + r46b \cdot r62bt + r42t + r45Ba \cdot r52BaPb \\
205_{tot}/^{202}\text{Pb}_{tr} &= 1/r25t + r52BaPb \\
206_{tot}/^{202}\text{Pb}_{tr} &= r62gt + r62bt + r62t \\
207_{tot}/^{202}\text{Pb}_{tr} &= r76g \cdot r62gt + r76b \cdot r62bt + r72t \\
208_{tot}/^{202}\text{Pb}_{tr} &= r86g \cdot r62gt + r86b \cdot r62bt + r82t
\end{aligned} \tag{B.3}$$

759 Finally, the left hand side of equations (B.3) can be expressed as measured isotope ratios with  
760  $206_{tot}$  in the denominator by dividing each equation by the fifth equation in the system above,  
761 thereby reducing the number of equations to six. To equate the measured isotope ratios to the true  
762 IC of the sample, isotopic fractionation, or the preferential evaporation, ionization, and/or detection  
763 of light isotopes over heavier ones, must be considered as well. We use a modified exponential  
764 fractionation law (Russell et al., 1978), which has been shown empirically to closely model measured  
765 Pb isotopic data analyzed with a silica gel emitter (McLean, 2014).

766 Pb has also been observed to exhibit mass-independent fractionation (MIF), with odd-numbered  
767 isotopes preferentially evaporating and/or ionizing relative to the mass-dependent trend predicted  
768 by exponential fractionation (Doucelance and Manhès, 2001; Amelin et al., 2005; McLean, 2014).  
769 For large loads effectively free of isobaric interferences, this effect has been observed to remain  
770 constant throughout the analysis of a sample loaded on a single filament, and can be parameterized  
771 by a factor  $\gamma$  that is unique to each isotope and is expected to be 1 for even-mass number isotopes.  
772 The modified exponential equation for two isotopes a and b thus takes the form (McLean, 2014)

$$\left(\frac{a}{b}\right)_m = \left(\frac{a}{b}\right)_{true} \cdot \left(\frac{M_a \cdot \gamma_a}{M_b \cdot \gamma_b}\right)^\beta \tag{B.4}$$

773 where  $M_a$  and  $M_b$  are the atomic masses of isotopes  $a$  and  $b$ , and  $(a/b)_m$  and  $(a/b)_{true}$  are the  
774 measured and true (fractionation-corrected) isotope ratios, respectively. For clarity, the  $\beta$  in equation  
775 (B.4) has the opposite sign but same absolute value as the conventional exponential fractionation  
776 equation.

Assuming  $\gamma_{202} = \gamma_{204} = \gamma_{206} = \gamma_{208} = 1$  and adding the modified exponential fractionation term



to the resulting Pb isotope ratios produces the system of equations

$$\begin{aligned}
r_{16}m_j^i &= \frac{r_{15}Ba \cdot r_{52}BaPb_j^i}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \\
r_{26}m_j^i &= \frac{1}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \left( \frac{M_{202}}{M_{206}} \right)^{\beta_j^i} + \frac{r_{25}Ba \cdot r_{52}BaPb_j^i}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \\
r_{46}m_j^i &= \frac{r_{42}t + r_{46}g \cdot r_{62}gt^i + r_{46}b \cdot r_{62}bt^i}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \left( \frac{M_{204}}{M_{206}} \right)^{\beta_j^i} + \frac{r_{45}Ba \cdot r_{52}BaPb_j^i}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \\
r_{56}m_j^i &= \frac{1/r_{25}t}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \left( \frac{M_{205} \cdot \gamma_{205}^i}{M_{206}} \right)^{\beta_j^i} + \frac{r_{52}BaPb_j^i}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \\
r_{76}m_j^i &= \frac{r_{72}t + r_{76}g \cdot r_{62}gt^i + r_{76}b \cdot r_{62}bt^i}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \left( \frac{M_{207} \cdot \gamma_{207}^i}{M_{206}} \right)^{\beta_j^i} \\
r_{86}m_j^i &= \frac{r_{82}t + r_{86}g \cdot r_{62}gt^i + r_{86}b \cdot r_{62}bt^i}{r_{62}t + r_{62}gt^i + r_{62}bt^i} \left( \frac{M_{208}}{M_{206}} \right)^{\beta_j^i}
\end{aligned} \tag{B.5}$$

777 where the superscripted index  $i$  refers to an analysis and the subscripted index  $j$  refers to an  
778 individual measured value. Thus each measurement must be corrected by a unique  $r_{52}BaPb$ , or  
779 magnitude of  $BaPO_2$  isobaric interference, and  $\beta$ , or magnitude of isotopic fractionation, which both  
780 change during the course of the analysis. Each analysis, which consists of multiple measurements,  
781 has a unique value for  $r_{62}gt$  and  $r_{62}bt$ , which express the relative quantities of tracer, blank, and  
782 gravimetric solution, as well as  $\gamma_{205}$  and  $\gamma_{207}$ , which quantify the degree of MIF for the odd-mass  
783 number Pb isotopes. Variables without indices, such as the tracer, Pb reference material, blank,  
784 and  $BaPO_2$  ICs, are assumed to be constant for all analyses of the same gravimetric Pb reference  
785 material.

#### 786 *Appendix B.2. Gravimetric-Tracer Mixture Pb Equations*

Unlike the data used in the Pb isotopic reference material inter-calibration in Section 2, the gravimetric-tracer Pb measurements, made at both NIGL and MIT, do not monitor the  $BaPO_2$  isobaric interference. Because the Pb and U were loaded together and run on the same filament, Pb as  $Pb^+$  at lower temperature, then U as  $UO_2^+$  at higher temperature, the Pb was not run to the high temperatures at which  $BaPO_2$  becomes a significant isobaric interference. Additionally, because the  $^{204}Pb$  abundance is always less than an order of magnitude smaller than the other Pb isotopes, it does not contribute meaningfully to tracer calibration calculations, and has been ignored here. These omissions result in a simpler isotopic contribution budget than that used for the Pb

reference material inter-calibration,

$$\begin{aligned}
^{202}\text{Pb}_{tot} &= ^{202}\text{Pb}_{tr} \\
^{205}\text{Pb}_{tot} &= ^{205}\text{Pb}_{tr} \\
^{206}\text{Pb}_{tot} &= ^{206}\text{Pb}_{gr} + ^{206}\text{Pb}_{bl} + ^{206}\text{Pb}_{tr} \\
^{207}\text{Pb}_{tot} &= ^{207}\text{Pb}_{gr} + ^{207}\text{Pb}_{bl} + ^{207}\text{Pb}_{tr} \\
^{208}\text{Pb}_{tot} &= ^{208}\text{Pb}_{gr} + ^{208}\text{Pb}_{bl} + ^{208}\text{Pb}_{tr}
\end{aligned} \tag{B.6}$$

787 where *tr*, *gr*, and *bl* correspond to the tracer, gravimetric, and blank components, respectively,  
788 which sum to the total abundance of each isotope, denoted *tot*.

Normalizing equations (B.6) to  $^{205}\text{Pb}$ , which is present in both ET535 and ET2535 tracers but not in the gravimetric solutions, yields

$$\begin{aligned}
\frac{^{202}\text{Pb}_{tot}}{^{205}\text{Pb}_{tr}} &= \left( \frac{^{202}\text{Pb}_{tr}}{^{205}\text{Pb}_{tr}} \right) \\
\frac{^{206}\text{Pb}_{tot}}{^{205}\text{Pb}_{tr}} &= \left( \frac{^{206}\text{Pb}_{gr}}{^{205}\text{Pb}_{tr}} \right) + \left( \frac{^{206}\text{Pb}_{bl}}{^{205}\text{Pb}_{tr}} \right) + \left( \frac{^{206}\text{Pb}_{tr}}{^{205}\text{Pb}_{tr}} \right) \\
\frac{^{207}\text{Pb}_{tot}}{^{205}\text{Pb}_{tr}} &= \left( \frac{^{207}\text{Pb}_{gr}}{^{205}\text{Pb}_{tr}} \right) + \left( \frac{^{207}\text{Pb}_{bl}}{^{205}\text{Pb}_{tr}} \right) + \left( \frac{^{207}\text{Pb}_{tr}}{^{205}\text{Pb}_{tr}} \right) \\
\frac{^{208}\text{Pb}_{tot}}{^{205}\text{Pb}_{tr}} &= \left( \frac{^{208}\text{Pb}_{gr}}{^{205}\text{Pb}_{tr}} \right) + \left( \frac{^{208}\text{Pb}_{bl}}{^{205}\text{Pb}_{tr}} \right) + \left( \frac{^{208}\text{Pb}_{tr}}{^{205}\text{Pb}_{tr}} \right)
\end{aligned} \tag{B.7}$$

The isotope ratios on the right hand side of equations (B.7) can be re-cast in terms of the isotope ratios of the gravimetric, tracer, and blank components. For instance, the ratio of  $^{207}\text{Pb}$  in the blank to  $^{205}\text{Pb}$  in the tracer can be expressed as the  $^{207}\text{Pb}/^{206}\text{Pb}$  of the blank, derived in Section 3, multiplied by the ratio of the moles of  $^{206}\text{Pb}$  in the blank to the moles of  $^{205}\text{Pb}$  in the tracer. Additionally, the isotope ratios on the left hand side of equations (B.7) can be expressed as measured isotope ratios when the ‘true’ isotope ratios on the right hand side are modified by fractionation correction factor (equation B.4). With these two substitutions, equations (B.7) become

$$\begin{aligned}
\left( \frac{^{202}\text{Pb}}{^{205}\text{Pb}} \right)_m^i &= \left( \frac{^{202}\text{Pb}}{^{205}\text{Pb}} \right)_{tr} \left( \frac{M_{202}}{M_{205} \cdot \gamma_{205}^i} \right)^{\beta_j^i} \\
\left( \frac{^{206}\text{Pb}}{^{205}\text{Pb}} \right)_m^i &= \left[ \left( \frac{^{206}\text{Pb}_{gr}}{^{205}\text{Pb}_{tr}} \right)^i + \left( \frac{^{206}\text{Pb}_{bl}}{^{205}\text{Pb}_{tr}} \right)^i + \left( \frac{^{206}\text{Pb}}{^{205}\text{Pb}} \right)_{tr} \right] \left( \frac{M_{206}}{M_{205} \cdot \gamma_{205}^i} \right)^{\beta_j^i} \\
\left( \frac{^{207}\text{Pb}}{^{205}\text{Pb}} \right)_m^i &= \left[ \left( \frac{^{207}\text{Pb}}{^{206}\text{Pb}} \right)_{gr} \left( \frac{^{206}\text{Pb}_{gr}}{^{205}\text{Pb}_{tr}} \right)^i + \left( \frac{^{207}\text{Pb}}{^{206}\text{Pb}} \right)_{bl} \left( \frac{^{206}\text{Pb}_{bl}}{^{205}\text{Pb}_{tr}} \right)^i + \left( \frac{^{207}\text{Pb}}{^{205}\text{Pb}} \right)_{tr} \right] \left( \frac{M_{207} \cdot \gamma_{207}^i}{M_{205} \cdot \gamma_{205}^i} \right)^{\beta_j^i} \\
\left( \frac{^{208}\text{Pb}}{^{205}\text{Pb}} \right)_m^i &= \left[ \left( \frac{^{208}\text{Pb}}{^{206}\text{Pb}} \right)_{gr} \left( \frac{^{206}\text{Pb}_{gr}}{^{205}\text{Pb}_{tr}} \right)^i + \left( \frac{^{208}\text{Pb}}{^{206}\text{Pb}} \right)_{bl} \left( \frac{^{206}\text{Pb}_{bl}}{^{205}\text{Pb}_{tr}} \right)^i + \left( \frac{^{208}\text{Pb}}{^{205}\text{Pb}} \right)_{tr} \right] \left( \frac{M_{208}}{M_{205} \cdot \gamma_{205}^i} \right)^{\beta_j^i}
\end{aligned} \tag{B.8}$$

789 where *i* denotes a variable that changes from load to load and *j* denotes a variable that additionally  
790 changes from block to block. In this system, the variables corresponding to the blank, tracer,

791 and gravimetric solution Pb ICs are all treated as known variables, with maximum likelihood  
 792 estimates and uncertainties reported above. The ratio of the blank to the tracer, embodied in the  
 793  $^{206}\text{Pb}_{bl}/^{205}\text{Pb}_{tr}$  can be estimated from tracer mass used and total procedural blank measurements.  
 794 Following McLean (2014) the mass-independent fractionation parameterized with the  $\gamma$  terms is  
 795 assumed to remain constant for each load but vary between loads, and the absolute magnitude of  
 796 fractionation,  $\beta$  changes during the course of each analysis. For gravimetric solution mixtures with  
 797 ET535, the first equation that includes  $^{202}\text{Pb}$  is ignored.

798 For the corresponding U measurement, the same system used for the critical mixtures in equations  
 799 (B.13) applies. Because the magnitude of isotopic fractionation changes dramatically during the  
 800 long gravimetric-tracer mixture analyses, the parameter  $\beta$  is allowed to change from block to block.  
 801 The paired Pb and U measurements combine to form a large overdetermined system of equations,  
 802 with each block of data contributing five or six measured ratios, depending on whether  $^{202}\text{Pb}$  is  
 803 present. Using the gravimetric solution  $^{206}\text{Pb}/^{238}\text{U}$  calculated from equations (7) and (8), and the  
 804  $(^{238}\text{U}_{gr}/^{235}\text{U}_{tr})$  from equations (B.13), the term  $(^{206}\text{Pb}_{gr}/^{205}\text{Pb}_{tr})$  can be recast as

$$\left(\frac{^{206}\text{Pb}_{gr}}{^{205}\text{Pb}_{tr}}\right)^i = \left(\frac{^{235}\text{U}}{^{205}\text{Pb}}\right)_{tr} \cdot \left(\frac{^{206}\text{Pb}}{^{238}\text{U}}\right)_{gr} \cdot \left(\frac{^{238}\text{U}_{gr}}{^{235}\text{U}_{tr}}\right)^i \quad (\text{B.9})$$

805 Substituting the right-hand side expression (B.9) into equations (B.8), along with equations (B.13)  
 806 for U measurements, creates a system of five equations for ET535 measurements, or six equations  
 807 for ET2535 measurements that relate measured Pb and U isotope ratios to the gravimetric and  
 808 tracer solution ICs and their U/Pb ratios.

### 809 *Appendix B.3. U Model*

The isotopic composition of all critical mixtures and tracer ICs were measured as  $\text{UO}_2$  (Condon  
 et al., in review). Isobaric interference corrections were made on each cycle, and the mean and  
 standard error of the resulting U ratios represent the best estimate of the fractionated IC of the  
 sample. Expressing the components that make up a mixture of the ET(2)535 tracer with a reference  
 material, such as SRM U500 or CRM 112a,

$$\begin{aligned} 233_{tot} &= 233_{tr} \\ 235_{tot} &= 235_{st} + 235_{bl} + 235_{tr} \\ 238_{tot} &= 238_{st} + 238_{bl} + 238_{tr} \end{aligned} \quad (\text{B.10})$$

810 where *st* denotes the contribution from an isotopic reference material, *tr* from the tracer, and *bl*  
 811 from the blank.

Normalizing all the components to the moles of  $^{235}\text{U}$  in the tracer results in

$$\begin{aligned} 233_{tot}/235_{tr} &= (233_{tr}/235_{tr}) \\ 235_{tot}/235_{tr} &= (235_{st}/235_{tr}) + (235_{bl}/235_{tr}) + (235_{tr}/235_{tr}) \\ 238_{tot}/235_{tr} &= (238_{st}/235_{tr}) + (238_{bl}/235_{tr}) + (238_{tr}/235_{tr}) \end{aligned} \quad (\text{B.11})$$

Recasting the mixed component ratios ( $235_{bl}/235_{tr}$ ) as  $(238_{bl}/235_{tr}) / (238_{bl}/235_{bl})$  and  $(235_{st}/235_{tr})$  as  $(238_{st}/235_{tr}) / (238_{st}/235_{st})$  yields

$$\begin{aligned} 233_{tot}/235_{tr} &= r35t \\ 235_{tot}/235_{tr} &= r85st/r85s + r85bt/r85b + 1 \\ 238_{tot}/235_{tr} &= r85st + r85bt + r85t \end{aligned} \tag{B.12}$$

where  $r35t$  and  $r85t$  are the tracer parameters that are being determined,  $r85b$  and  $r85s$  are the  $^{238}\text{U}/^{235}\text{U}$  of the blank and the isotopic reference material, respectively, and  $r85st$  and  $r85bt$  are the moles of  $^{238}\text{U}$  in the reference material and blank, respectively, relative to the moles of  $^{235}\text{U}$  in the tracer.

Dividing the first and third equations by the second equation yields the U ratios as commonly measured. In order to equate the true and measured ratios, an exponential fractionation term is added as well.

$$\begin{aligned} r35m^i &= \frac{r35t}{1 + r85bt^i/r85b + r85st^i/r85s} \left( \frac{M_{233}}{M_{235}} \right)^{\beta^i} \\ r85m^i &= \frac{r85bt^i + r85st^i + r85t}{1 + r85bt^i/r85b + r85st^i/r85s} \left( \frac{M_{238}}{M_{235}} \right)^{\beta^i} \end{aligned} \tag{B.13}$$

where  $M_x$  denotes the isotopic mass of  $x$  and the superscript  $i$  denotes variables that change from one analysis to the next. These two equations apply to each measurement of a critical mixture made, regardless of the isotopic reference material used. For measurements of the tracer only, which also include a loading blank, the  $r85s$  and  $r85st$  terms may be dropped.

## Bibliography

- Amelin Y., Davis D.W. and Davis W.J. (2005) Decoupled fractionation of even- and odd-mass isotopes of Pb in TIMS. *Geochimica et Cosmochimica Acta* **69**(10), A215.
- Amelin Y. and Davis W.J. (2006) Isotopic analysis of lead in sub-nanogram quantities by TIMS using a  $^{202}\text{Pb}$ - $^{205}\text{Pb}$  spike. *Journal of Analytical Atomic Spectrometry* **21**(10), 1053–1061.
- Audi G., Wapstra A.H. and Thibault C. (2003) The Ame2003 atomic mass evaluation: (II). Tables, graphs and references. *Nuclear Physics A* **729**(1), 337–676.
- Baker J., Peate D., Waight T. and Meyzen C. (2004) Pb isotopic analysis of standards and samples using a  $^{207}\text{Pb}$ - $^{204}\text{Pb}$  double spike and thallium to correct for mass bias with a double-focusing MC-ICP-MS. *Chemical Geology* **211**(3-4), 275–303.
- Böhlke J.K., de Laeter J.R., Bièvre P.D., Hidaka H., Peiser H.S., Rosman K.J.R. and Taylor P.D.P. (2005) Isotopic compositions of the elements, 2001. *Journal of Physical and Chemical Reference Data* **34**(1), 57–67.
- Catanzaro E.J., Murphy T.J., Shields W.R. and Garner E.L. (1968) Absolute isotopic abundance ratios of common, equal-atom, and radiogenic lead standards. *Journal of Research of the National Bureau of Standards - A. Physics and Chemistry* **72A**(3), 261–267.

- 836 Cheng H., Lawrence Edwards R., Shen C.C., Polyak V.J., Asmerom Y., Woodhead J., Hellstrom  
837 J., Wang Y., Kong X., Spötl C., Wang X. and Calvin Alexander Jr. E. (2013) Improvements in  
838  $^{230}\text{Th}$  dating,  $^{230}\text{Th}$  and  $^{234}\text{U}$  half-life values, and U–Th isotopic measurements by multi-collector  
839 inductively coupled plasma mass spectrometry. *Earth and Planetary Science Letters* **371–372**(0),  
840 82–91.
- 841 Condon D.J., McLean N., Noble S.R. and Bowring S.A. (2010) Isotopic composition ( $^{238}\text{U}/^{235}\text{U}$ )  
842 of some commonly used uranium reference materials. *Geochimica et Cosmochimica Acta* **74**,  
843 7127–7143.
- 844 Condon D.J., Schoene R.B., McLean N.M., Parrish R. and Bowring S.A. (in review) Metrology and  
845 traceability of U–Pb isotope dilution geochronology (EARTHTIME tracer calibration Part I),  
846 submitted to *Geochem. Geophys. Geosyst.*
- 847 Corfu F. and Dahlgren S. (2008) Perovskite U–Pb ages and the Pb isotopic composition of alkaline  
848 volcanism initiating the Permo-Carboniferous Oslo Rift. *Earth and Planetary Science Letters*  
849 **265**(1–2), 256–269.
- 850 Doucelance R. and Manhès G. (2001) Reevaluation of precise lead isotope measurements by  
851 thermal ionization mass spectrometry: comparison with determinations by plasma source mass  
852 spectrometry. *Chemical Geology* **176**(1–4), 361–377.
- 853 Galer S.J.G. (1999) Optimal double and triple spiking for high precision lead isotopic measurement.  
854 *Chemical Geology* **157**(3–4), 255–274.
- 855 Hiess J., Condon D.J., McLean N. and Noble S.R. (2012)  $^{238}\text{U}/^{235}\text{U}$  systematics in terrestrial  
856 U-bearing minerals. *Science* **335**(6076), 1610–1614.
- 857 Hofmann A. (1971) Fractionation corrections for mixed-isotope spikes of Sr, K, and Pb. *Earth and*  
858 *Planetary Science Letters* **10**(4), 397–402.
- 859 Jaffey A.H., Flynn K.F., Glendenin L.E., Bentley W.C. and Essling A.M. (1971) Precision measure-  
860 ment of half-lives and specific activities of  $^{235}\text{U}$  and  $^{238}\text{U}$ . *Phys. Rev. C*(4), 1889–1906.
- 861 Krogh T.E. (1964) Strontium isotopic variation and whole rock isochron studies in the Grenville  
862 Province of Ontario. *MIT Annual Progress Reports (AEC Contract AT(30-1)-1381)* **12**, 73.
- 863 Mattinson J.M. (2010) Analysis of the relative decay constants of  $^{235}\text{U}$  and  $^{238}\text{U}$  by multi-step  
864 CA-TIMS measurements of closed-system natural zircon samples. *Chemical Geology* **275**(3–4),  
865 186–198.
- 866 McLean N.M. (2014) Straight line regression through data with correlated uncertainties in two or  
867 more dimensions, with an application to kinetic isotope fractionation. *Geochimica et Cosmochimica*  
868 *Acta* **124**(0), 237–249.
- 869 McLean N.M., Bowring J.F. and Bowring S.A. (2011) An algorithm for U–Pb isotope dilution data  
870 reduction and uncertainty propagation. *Geochem. Geophys. Geosyst.* **12**.
- 871 Nebel O., Scherer E.E. and Mezger K. (2011) Evaluation of the  $^{87}\text{Rb}$  decay constant by age  
872 comparison against the U–Pb system. *Earth and Planetary Science Letters* **301**(1–2), 1–8.

- 873 Renne P.R., Mundil R., Balco G., Min K. and Ludwig K.R. (2010) Joint determination of  $^{40}\text{K}$   
874 decay constants and  $^{40}\text{Ar}^*/^{40}\text{K}$  for the Fish Canyon sanidine standard, and improved accuracy  
875 for  $^{40}\text{Ar}/^{39}\text{Ar}$  geochronology. *Geochimica et Cosmochimica Acta* **74**(18), 5349–5367.
- 876 Rioux M., Johan Lissenberg C., McLean N.M., Bowring S.A., MacLeod C.J., Hellebrand E. and  
877 Shimizu N. (2012) Protracted timescales of lower crustal growth at the fast-spreading East Pacific  
878 Rise. *Nature Geoscience* **5**(4), 275–278.
- 879 Rotenberg E., Davis D.W., Amelin Y., Ghosh S. and Bergquist B.A. (2012) Determination of the  
880 decay-constant of  $^{87}\text{Rb}$  by laboratory accumulation of  $^{87}\text{Sr}$ . *Geochimica et Cosmochimica Acta*  
881 **85**(0), 41–57.
- 882 Rudge J.F., Reynolds B.C. and Bourdon B. (2009) The double spike toolbox. *Chemical Geology*  
883 **265**(3–4), 420–431.
- 884 Russell W.A., Papanastassiou D.A. and Tombrello T.A. (1978) Ca isotope fractionation on the  
885 Earth and other solar system materials. *Geochimica et Cosmochimica Acta* **42**(8), 1075–1090.
- 886 Schmitz M.D. and Schoene B. (2007) Derivation of isotope ratios, errors, and error correlations  
887 for U-Pb geochronology using  $^{205}\text{Pb}$ - $^{235}\text{U}$ -( $^{233}\text{U}$ )-spiked isotope dilution thermal ionization mass  
888 spectrometric data. *Geochem. Geophys. Geosyst.* **8**(8).
- 889 Schoene B., Crowley J.L., Condon D.J., Schmitz M.D. and Bowring S.A. (2006) Reassessing  
890 the uranium decay constants for geochronology using ID-TIMS U–Pb data. *Geochimica et*  
891 *Cosmochimica Acta* **70**(2), 426–445.
- 892 Stirling C.H., Andersen M.B., Potter E.K. and Halliday A.N. (2007) Low-temperature isotopic  
893 fractionation of uranium. *Earth and Planetary Science Letters* **264**(1–2), 208–225.
- 894 Stracke A., Scherer E. and Reynolds B. (2014) 15.4 - application of isotope dilution in geochemistry.  
895 In *Treatise on Geochemistry (Second Edition)* (eds. H.D. Holland and K.K. Turekian), pp. 71–86,  
896 Elsevier, Oxford.  
897 **URL:** <http://www.sciencedirect.com/science/article/pii/B9780080959757014042>
- 898 Tarantola A. (2005) *Inverse Problem Theory and Methods for Model Parameter Estimation*. SIAM,  
899 Philadelphia, Pennsylvania.
- 900 Tarantola A. and Valette B. (1982a) Generalized nonlinear inverse problems solved using the least  
901 squares criterion. *Review of Geophysics and Space Physics* **20**, 219–232.
- 902 Tarantola A. and Valette B. (1982b) Inverse problems = quest for information. *Journal of Geophysics*  
903 **50**, 159–170.
- 904 Tilton G.R., Patterson C., Brown H., Inghram M., Hayden R., Hess D. and Larsen E. (1955) Isotopic  
905 composition and distribution of lead, uranium, and thorium in a Precambrian granite. *Geological*  
906 *Society of America Bulletin* **66**(9), 1131–1148.
- 907 Todt W., Cliff R., Hanser A. and Hofmann A. (1996) *Evaluation of a  $^{202}\text{Pb}$ - $^{205}\text{Pb}$  Double Spike for*  
908 *High-Precision Lead Isotope Analysis*, vol. 95, Earth Processes: Reading the Isotopic Code of  
909 *Geophysical Monographs*, pp. 429–437. American Geophysical Union.

- 910 Verbruggen A., Eykens R., Kehoe F., Kühn H., Richter S. and Aregbe Y. (2008) *Preparation and*  
911 *Certification of IRMM-3636, IRMM-3636a and IRMM-3636b*. Tech. rep., JRC Scientific and  
912 Technical Reports.  
913 **URL:** <http://publications.jrc.ec.europa.eu/repository/handle/111111111/4450>
- 914 Vermeesch P. (2010) Helioplot, and the treatment of overdispersed (U-Th-Sm)/He data. *Chemical*  
915 *Geology* **271**(3-4), 108–111.
- 916 Vervoort J.D., Patchett P.J., Sderlund U. and Baker M. (2004) Isotopic composition of yb and the  
917 determination of lu concentrations and lu/hf ratios by isotope dilution using mc-icpms. *Geochem.*  
918 *Geophys. Geosyst.* **5**(11), Q11002–.
- 919 Wasserburg G., Jacousen S., DePaolo D., McCulloch M. and Wen T. (1981) Precise determination of  
920 Sm/Nd ratios, Sm and Nd isotopic abundances in standard solutions. *Geochimica et Cosmochimica*  
921 *Acta* **45**(12), 2311–2323.
- 922 Wetherill G.W. (1956) Discordant uranium-lead ages, I. *Transactions, American Geophysical Union*  
923 **37**, 320–326.
- 924 Weyer S., Anbar A.D., Gerdes A., Gordon G.W., Algeo T.J. and Boyle E.A. (2008) Natural  
925 fractionation of  $^{238}\text{U}/^{235}\text{U}$ . *Geochimica et Cosmochimica Acta* **72**(2), 345–359.