1	Calibration methods for laser ablation Rb–Sr geochronology:
2	comparisons and recommendation based on NIST glass and
3	natural reference materials
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9	
10	Abstract

In-situ Rb-Sr geochronology using LA-ICP-MS/MS technology allows rapid dating of K-rich 11 12 minerals such as micas (e.g. biotite, muscovite, phlogopite) and K-feldspar. While many studies 13 have demonstrated the ability of the method, analytical protocols vary significantly and to date no 14 studies have provided an in-depth comparison and synthesis in terms of precision and accuracy. 15 Here we compare four calibration protocols based on commonly used reference materials for Rb-16 Sr dating. We demonstrate that downhole fractionation trends (DHF) for natural biotite, K-feldspar 17 and phlogopite contrast with that for the commonly used Mica-Mg nano-powder reference 18 material. Consequently, Rb-Sr dates calibrated to Mica-Mg can be up to 5% inaccurate and the 19 degree of inaccuracy appears to be unsystematic between analytical sessions. Calibrating to Mica-20 Mg also introduces excess uncertainty that can be avoided with a more consistent primary 21 calibration material. We propose a calibration approach involving (1) NIST-610 glass as the 22 primary reference material (PRM) for normalization and drift correction and (2) a natural mineral 23 with similar DHF characteristics to the analysed samples as matrix correction RM (MCRM) to 24 correct the Rb/Sr ratio for matrix-induced offsets. In this work, MDC phlogopite (the source 25 mineral for Mica-Mg nano-powder) was used as the MCRM, consistently producing accurate Rb-26 Sr dates for a series of natural biotites and K-feldspars with well-characterized expected ages. 27 However, biotite from the Banalasta Adamellite, Taratap Granodiorite and Entire Creek pegmatite are also suitable RMs for Rb/Sr ratio calibration purposes with consistently <1.5% fully 28 29 propagated uncertainties in our methodological approach. Until calibration using isochronous 30 natural standards as the primary RM becomes possible in data-reduction software, the two-step 31 calibration approach described here is recommended.

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Keywords: reaction-cell ICP-MS; in-situ geochronology; Rb–Sr reference materials; calibration
 standards

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36 **1. Introduction**

Rubidium-Strontium (Rb–Sr) geochronology using laser ablation – inductively coupled plasma – tandem mass spectrometry (LA-ICP-MS/MS) has become a popular method to constrain the formation or cooling age of potassium-bearing minerals (Gorojovsky and Alard, 2020; Hogmalm et al., 2017; Jegal et al., 2022; Kirkland et al., 2023; Larson et al., 2023; Laureijs et al., 2021; Li et al., 2020; Liebmann et al., 2022; Olierook et al., 2020; Redaa et al., 2021; Rosel and Zack, 2022; Sengun et al., 2019; Tillberg et al., 2021; Tillberg et al., 2020; Wang et al., 2022; Zack and Hogmalm, 2016). In contrast to traditional Rb–Sr dating involving column-chemistry in 44 specialized laboratories, the laser-ablation method allows rapid acquisition of Rb–Sr dates directly 45 from thin sections or rock blocks with minimal sample preparation. The method involves the use of an ICP-MS/MS, equipped with a reaction cell where isobaric isotopes can be chemically 46 47 separated due to their significant differences in reactivity with an introduced reaction gas (Balcaen 48 et al., 2015 and references therein). Applied to Rb–Sr geochronology, CH₃F, SF₆, O₂ and N₂O 49 have been used as reaction gasses (e.g. Hogmalm et al., 2017; Moens et al., 2001; Zack and Hogmalm, 2016), with the latter being the most widely used for quadrupole ICP-MS/MS due to 50 51 its high reactivity. However, published analytical methodologies for LA-ICP-MS/MS Rb-Sr 52 dating vary significantly beyond the applied reaction gas (Table 1). Reported laser conditions 53 (fluence and repetition rate) are largely laser-wavelength dependent with common conditions being either $\sim 5 - 7$ J.cm⁻² / 10 Hz for 213nm lasers, especially during initial development work 54 55 (e.g. Hogmalm et al., 2017; Laureijs et al., 2021; Rosel and Zack, 2022; Sengun et al., 2019; Tillberg et al., 2020; Zack and Hogmalm, 2016) or ~2 - 4 J.cm⁻² / 5 Hz for 193nm lasers (e.g. 56 Kirkland et al., 2023; Larson et al., 2023; Li et al., 2020; Liebmann et al., 2022; Olierook et al., 57 58 2020; Redaa et al., 2021). The applied calibration protocols for mass discrimination and elemental 59 fractionation, however, vary more significantly.

- 60 We define three types of reference materials (RM) in this manuscript:
- (1) The Primary RM (PRM) has a homogenous isotopic composition and is used for
 normalisation and drift correction;
- 63 (2) The matrix correction RM (MCRM) has a heterogenous isotopic composition but well64 known age and is used to correct the Rb/Sr ratio for systematic matrix-induced off-sets
 65 between the PRM and mineral samples.

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(3) The secondary RM (SRM) has a well-known age and a similar composition to the analysed samples and is used to verify the accuracy of the calibration protocol.

68 Most published work uses a glass reference material as PRM, with NIST-610 being most popular 69 to correct for drift and calibrate the Sr isotopic ratios. Rb/Sr ratios are most commonly calibrated 70 against Mica-Mg, a phlogopite prepared as a pressed nano-powder pellet, regardless of the 71 analysed mineral (micas in most published work). However, the approach varies, with some 72 methods directly calibrating to Mica-Mg as the PRM (e.g. Gorojovsky and Alard, 2020; Hogmalm 73 et al., 2017; Li et al., 2020; Redaa et al., 2021; Rosel and Zack, 2022; Sengun et al., 2019; Wang 74 et al., 2022) and others using NIST-610 as the PRM followed by a correction for matrix-dependent fractionation against Mica-Mg as MCRM (e.g. Liebmann et al., 2022; Olierook et al., 2020). 75 76 Secondary RMs, used to verify the accuracy of obtained dates, are either glass reference materials 77 (e.g. Larson et al., 2023; Laureijs et al., 2021; Rosel and Zack, 2022) or in-house natural materials 78 such as the La Posta biotite (Zack and Hogmalm, 2016), the MDC phlogopite (Redaa et al., 2021), 79 or the CK001 biotite (Olierook et al., 2020).

80 In addition, laser-induced down-hole fractionation (DHF) can occur during ablation and aerosol 81 condensation processes and is most apparent when ratioing elements with contrasting volatilities 82 (e.g. Jackson and Günther, 2003; Košler et al., 2005; Longerich et al., 1996). Elemental Sr is more 83 refractory than the volatile Rb and hence has a high potential to fractionate during laser ablation (Zack and Hogmalm, 2016). A small number of studies have directly compared different 84 85 calibration approaches and have described differences in Rb-Sr DHF behaviour between 86 commonly used reference materials (e.g. Redaa et al., 2021; Wang et al., 2022). However, 87 systematic comparisons between data reduction protocols, tested with natural materials, are limited 88 in the current literature. Here, we compare four different calibration approaches for a series of

89	natural biotite and K-feldspar samples. The samples were taken from quickly cooled igneous rocks,
90	eliminating potential diffusion-related issues when comparing dates of different minerals. Hence,
91	the well-constrained igneous crystallization ages are the expected reference ages for the analysed
92	samples and one of the biotite samples has previously been dated by the Rb–Sr ID-TIMS method.
93	The calibration approaches we compare are:
94	(A) NIST-610 as the PRM for both ⁸⁷ Rb/ ⁸⁷ Sr and ⁸⁷ Sr/ ⁸⁶ Sr ratios plus MDC phlogopite as MCRM;
95	(B) NIST-610 as the PRM for both ⁸⁷ Rb/ ⁸⁷ Sr and ⁸⁷ Sr/ ⁸⁶ Sr ratios plus Mica-Mg pressed pellet as
96	MCRM;
97	(C) Mica-Mg as the PRM for 87 Rb/ 87 Sr ratios and NIST-610 as the PRM for 87 Sr/ 86 Sr ratios;
98	(D) Mica-Mg as the PRM for both ⁸⁷ Rb/ ⁸⁷ Sr and ⁸⁷ Sr/ ⁸⁶ Sr ratios
99	We discuss the differences between these approaches in terms of accuracy and precision, and
100	highlight the importance of monitoring and correcting down-hole fractionation with appropriate
101	reference materials.
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103	2. Sample descriptions
104	2.1. MDC phlogopite and Mica-Mg nano powder
105	Mica-Mg nano-powder is used as a reference material for Rb-Sr dating. It consists of crushed
106	phlogopite from Bekily (Madagascar) with a high Rb (1300 \pm 40 μ g.g ⁻¹) and low Sr (27 \pm 3 μ g.g ⁻¹
107	¹) concentration (Redaa et al., 2023 and references therein). MDC is natural phlogopite, which was
108	sourced from the same locality as Mica-Mg (Redaa et al., 2021). The reference age for both

materials is 519.4 ± 6.5 Ma and the initial 87 Sr/ 86 Sr ratio is 0.72607 ± 0.0007 (2SE uncertainties),

constrained from a diopside (low-Rb mineral) that occurs in the same location (Hogmalm et al.,
2017). However, for Mica-Mg some pellet to pellet variation in both Rb/Sr and Sr/Sr ratios has
been observed (Jegal et al., 2022; Redaa et al., 2023).

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2.2. Entire Creek pegmatite

114 The Entire Creek sample was taken from a deformed pegmatite in the Harts Range meta-igneous 115 complex of central Australia, in the same location as described by Mortimer et al. (1987). The 116 pegmatite cross-cuts folded and foliated amphibolites, is composed of coarse-grained quartz, 117 plagioclase, alkali feldspar and biotite, with the latter defining a strong axial-plane foliation to 118 folds outlined by the pegmatite. Biotite and whole-rock Rb/Sr and Sr/Sr isotope ratios, obtained 119 by ID-TIMS at the University of Adelaide, are reported in Mortimer et al. (1987) and define a 7-120 point (3 biotite and 4 whole rock analyses) isochron age of $312.1 \pm 1.8 / 5.1$ Ma (95% confidence 121 uncertainties, without / with overdispersion), recalculated in IsoplotR (Vermeesch, 2018), using the Villa et al. (2015) Rb–Sr decay constant of $1.3972 \pm 0.0045 \times 10^{-11} a^{-1}$ (Appendix 1). 122

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2.3. Banalasta Adamellite (Bundarra Suite)

124 The S-type Banalasta Adamellite forms the southern end the Bundarra Batholith in the Southern 125 New England Orogen in eastern Australia (e.g. Flood and Shaw, 1975; Jeon et al., 2012; 126 Rosenbaum et al., 2012; Shaw and Flood, 1981). The Bundarra Batholith is an elongate north-127 south trending magmatic suite, spanning approximately 200 km. The Banalasta Adamellite is 128 approximately 40 km in diameter and has sharp contacts with surrounding metasediments with a 129 contact metamorphic aureole characterised by fine-grained cordierite-bearing assemblage at the 130 pluton margin grading out to regional prehnite-pumpellyite metagreywacke assemblages over a distance of approximately 3 km (Flood and Shaw, 1977). Internally the granite is massive, coarse-131

132 grained granitoid containing approximately equal proportions of K-feldspar and plagioclase, 133 together with accessory apatite, zircon and monazite. Biotite predominantly occurs in multi-grain 134 clots together with quartz, plagioclase, magnetite, zircon and apatite. In rare cases they contain 135 relic garnet, suggesting they formed from hydration of garnet entrained from the granitic source 136 region.

137 Melt-precipitated zircon from the Banalasta Adamellite gives zircon U-Pb ages of 286.2 ± 2.2 Ma 138 (Black, 2007), 289.2 ± 1.7 Ma (Jeon et al., 2012) and 282 ± 4 Ma (Phillips et al., 2011). Whole 139 rock Rb–Sr data from the Bundarra Suite give an age of 285 ± 15 Ma (n = 6/7, MSWD = 0.4). 140 When additional feldspar Rb–Sr data are included in the isochron, the isochron age is 283 ± 10 Ma 141 (n = 9/10, MSWD = 0.24) (Appendix 2). Both isochron dates were recalculated using the data from 142 Flood and Shaw (1977) and the decay constant from Villa et al. (2015). Additionally, Hensel et al 143 (1995) reported a model whole rock Rb–Sr age of 287 ± 10 Ma for a group of 16 samples from the 144 Bundarra Suite. Overall, it is evident that Rb-Sr age data are similar to the ages of melt 145 precipitated zircon, consistent with the lack of evidence for extended fractional crystallisation 146 (Jeon et al 2012). The samples used in this study come from the same location as Black (2007) 147 that has a granitic zircon of 286.2 ± 2.2 Ma, as well as a second location approximately 800 meters 148 away.

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2.4. Taratap Granodiorite

151 The Taratap Granodiorite in the Delamerian Orogenic belt in southern Australia is classified as S-152 type, calc-alkaline with a composition dominated by microcline megacrysts (c. 3–4 cm in length), 153 which define a NNE-trending magmatic fabric in a coarse-grained groundmass of plagioclase, quartz, K-feldspar and biotite, with accessory zircon, apatite, and monazite. Low-temperature alteration is evident in thin section by the presence of chlorite–muscovite–titanite and minor allanite (Burtt and Abbot, 1998). The sample was chosen for analysis because the timing of emplacement is tightly constrained by a zircon U-Pb ID-TIMS age of 497.11 \pm 0.56 Ma (²⁰⁶Pb/²³⁸U weighted mean age, 95% confidence interval uncertainty, MSWD = 1.8, n = 6) and an apatite Lu-Hf age of 497.1 \pm 5.5 Ma (MSWD = 1.1, n = 38) (Glorie et al., 2023 and references therein).

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161 **3. Analytical methods**

162 All Rb-Sr analyses were conducted at Adelaide Microscopy, University of Adelaide, using an 163 Agilent 8900x ICP-MS/MS, coupled to a RESOlution-LR ArF excimer (193 nm) laser ablation 164 system. A squid mixing device (Laurin Technic) was used to smooth the pulsed laser signal 165 between the laser and the mass-spectrometer. The instrument parameters follow those reported in 166 Redaa et al. (2021), with ablation in a He atmosphere (350 mL.min⁻¹), mixed with Ar (890 mL.min⁻¹) 167 ¹) as the carrier gas and N₂ (3.5 mL.min⁻¹) added before the ICP torch to enhance the signal sensitivity. N₂O (0.37 mL.min⁻¹) was used as the reaction gas to separate ⁸⁷Sr from ⁸⁷Rb. The ⁸⁶Sr 168 and ⁸⁷Sr isotopes were measured as their oxide reaction products (e.g. ⁸⁷Sr¹⁶O) with a mass shift 169 170 of 16 amu between the two quadrupole mass analysers (e.g. Q1 = 87 m/z, Q2 = 103 m/z). Despite 171 the high reaction efficiency of ⁸⁷Sr, residual unreacted Sr prevents direct measurement of ⁸⁷Rb. Instead, ⁸⁵Rb was measured as a proxy for ⁸⁷Rb and calculated assuming natural isotopic 172 173 abundance. The samples and reference materials were ablated using a circular laser beam of 67 µm diameter, a fluence of 3.5 J.cm⁻², and repetition rate of 5 Hz. Further details are presented in 174 175 Table 2. A total of three analytical sessions were conducted, with largely identical instrumental 176 parameters between the different sessions. The ICP-MS was tuned to a sensitivity which kept Rb

in pulse mode in Mica-Mg (the material with the highest Rb concentration), negating the
requirement for additional pulse-analogue (P/A) corrections.

179 For each analytical session, NIST-610, Mica-Mg and MDC were used as reference materials for 180 calibration purposes. All data were processed in LADR (Norris and Danyushevsky, 2018) using 181 an in-built data reduction algorithm that calculates error correlations (Pearson correlation 182 coefficient) from the raw isotopic ratios for each sweep in an analysis, in the same way as for U-183 Pb data reduction. Isotope ratios were calculated by: (1) background subtraction, (2) correcting 184 down-hole fractionation (DHF) against the PRM, (3) averaging the DHF corrected ratios of each 185 sweep in the analysis, and then (4) normalising to the PRM to correct for matrix independent 186 instrument mass bias and drift. LADR applies a robust uncertainty propagation using the total 187 uncertainty budget of the measured quantified ratios. An example of an `uncertainty tree`, which 188 can be queried for every analysis, is given in Appendix 3. The reader is referred to the LADR 189 software manual (https://norsci.com/?p=ladr-support) for further details.

190 Normalisation of the measured Rb/Sr and Sr/Sr ratios was conducted with two different reference 191 materials (NIST-610 and Mica-Mg), following the four analytical protocols outlined above (A-D). The reference ${}^{87}\text{Rb}/{}^{87}\text{Sr}$ and ${}^{86}\text{Sr}/{}^{87}\text{Sr}$ ratios used for Mica-Mg were 83.4 ± 1.0 and $0.53981 \pm$ 192 0.00070, respectively (Hogmalm et al., 2017). For NIST-610, the ⁸⁷Rb/⁸⁷Sr was calculated from 193 194 concentration data (GeoREM preferred values) as 3.28 ± 0.03 and for the 86 Sr/ 87 Sr ratio, the 195 reference value of 1.409048 ± 0.000036 was used (Woodhead and Hergt, 2001). For each normalization protocol, DHF corrections of the ⁸⁷Rb/⁸⁷Sr ratios were applied based on the DHF 196 behaviour of the applied PRM. No DHF correction was applied to the ⁸⁶Sr/⁸⁷Sr ratios. Where 197 198 NIST-610 was used as the PRM, MDC phlogopite or Mica-Mg were used as MCRM to correct the ⁸⁷Rb/⁸⁷Sr ratios for matrix-dependant fractionation (cfr. Roberts et al., 2017 for U/Pb ratios;
Simpson et al., 2022 for Lu/Hf ratios).

All mica samples (including biotite samples and MDC phlogopite) were ablated with the laser ablating parallel to cleavage. The Bundarra and Taratap samples were analysed in thin section and optical microscopy (birefringence) was used to only select ablation targets with upright (\pm 10°) cleavage. The coarse Entire Creek biotites were mounted as mica-books using a vice to prevent air-gaps between individual mica sheets, with the mica sheets mounted upright exposing multiple cleavage planes perpendicular to the surface.

207 For each sample and reference material, inverse isochron Rb–Sr dates (Li and Vermeesch, 2021) were calculated in IsoplotR (Vermeesch, 2018), based on the processed ⁸⁷Rb/⁸⁷Sr and ⁸⁶Sr/⁸⁷Sr 208 209 ratios, their 2SE uncertainties, and the calculated error correlations. Reported inverse isochron uncertainties are fully propagated 95% confidence intervals, including the uncertainty on the decay 210 211 constant and added uncertainty for overdispersion where required (calculated in IsoplotR). The 212 exceptions are the inverse isochron dates for MDC and Mica-Mg when used as MCRMs, which 213 are used to correct the Rb/Sr ratios after calibrating to NIST-610. For these cases the reported 214 uncertainties are 95% confidence uncertainties without external uncertainties (as the external 215 uncertainties would otherwise be applied twice to the isochron dates of the analysed samples). Session-dependant correction factors (CF) were calculated from the measured ⁸⁷Rb/⁸⁷Sr ratio for 216 217 MDC and Mica-Mg (after drift corrections) and compared to the reference value (calculated from 218 the published age for both MDC and Mica-Mg of 519.4 ± 6.5 Ma; Hogmalm et al., 2017; Redaa 219 et al., 2021). These CF values (= measured ratio/expected ratio) were subsequently applied to each 220 unknown analysis when calibrated to either MDC or Mica-Mg. Finally, the uncertainties on the

MDC and Mica-Mg dates are propagated to the reported Rb–Sr isochron uncertainties for each
 NIST-610 calibrated sample using the quadratic addition of the relative uncertainties.

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4. Results

4.1. Downhole fractionation trends

In this section, we compare the downhole fractionation (DHF) trend of the ⁸⁷Rb/⁸⁷Sr ratio between 226 227 the analysed feldspars and micas and the reference materials (NIST-610 and Mica-Mg) (Fig. 1). The obtained fractionation trends do not vary significantly between different sessions; however, 228 229 data from analytical session 3 are presented as this session contains data for all analysed samples 230 presented in this paper. The DHF trends were calculated in LADR and individual scatter plots can 231 be found in Appendix 4. As shown, The DHF trends for the analysed biotite, phlogopite and K-232 feldspar samples are internally consistent, showing $\sim 10\%$ increase in Rb/Sr ratio over the first 20 233 s of ablation, followed by a flatter trend in the subsequent 20 s. NIST-610 shows a similar trend 234 of increasing Rb/Sr ratio with ablation time, however the amplitude of the DHF curve is more 235 subdued compared to the natural samples (~3.5% increase in the first 20 s ablation). In contrast, 236 the DHF pattern for Mica-Mg shows an oscillating trend, increasing for the first ~10 s of ablation 237 and then dropping for the subsequent ~ 30 s of ablation (Fig. 1).

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239 4.2. Within-session reproducibility of ⁸⁷Rb/⁸⁷Sr and ⁸⁶Sr/⁸⁷Sr ratios

Figure 2 shows the within-session variability (prior to drift correction) of the ⁸⁷Rb/⁸⁷Sr and ⁸⁶Sr/⁸⁷Sr ratios for both PRMs NIST-610 and Mica-Mg in analytical session 3. The reference materials are considered homogenous in both isotopic ratios, meaning that any variations are

243 purely due to differences in the ablation characteristics from spot to spot. As shown, the measured ⁸⁷Rb/⁸⁷Sr ratios and ⁸⁶Sr/⁸⁷Sr ratios are significantly more consistent for NIST-610 compared to 244 245 Mica-Mg (both measured using the same analytical conditions and spot size). The maximum within-session variability (=min-max range) in the 87 Rb/ 87 Sr ratio is < 3% for NIST-610, compared 246 to > 8% for Mica-Mg. The ⁸⁶Sr/⁸⁷Sr ratio is more consistent for both RMs, however, the uncertainty 247 248 on individual analyses is approximately 3× larger for Mica-Mg compared to NIST-610. ICP-MS 249 mass-bias drift is minimal for both isotope ratios in NIST-610, with only a slight increase in the 250 Rb/Sr ratio over the first 2-3 hours of analysis. As both Mica-Mg and NIST-610 were analysed sequentially in the same analytical session, the apparent 'drift' in the Mica-Mg ⁸⁶Sr/⁸⁷Sr ratios are 251 252 due to variations in ablation rather than changes in the ICP-MS mass bias.

253 4.3. Isochron Rb–Sr dates for natural K-feldspars and micas

254 Inverse isochron plots and resulting Rb-Sr dates are presented for each analytical protocol in Appendix 5. Summary plots are shown in Figure 3. The data table with the input ⁸⁷Rb/⁸⁷Sr and 255 ⁸⁶Sr/⁸⁷Sr ratios is accessible from Figshare via the link in the data availability section. For the 256 257 Bundarra samples, the biotite isochrons are anchored to apatite Rb/Sr ratios, given that the apatites 258 commonly occur as inclusions within biotite. For the K-feldspars, the isochrons are anchored to 259 plagioclase, given that the analysed K-feldspars often show minor exsolution with plagioclase. 260 However, the choice of anchoring mineral gives no difference in the obtained biotite and K-261 feldspar inverse isochron Rb/Sr dates. For the Taratap sample, isochron anchoring was conducted 262 to a combination of plagioclase and apatite in session 1, but only plagioclase in sessions 2 and 3, 263 given the limited occurrence of apatite in thin section. For the Entire Creek biotite sample, anchoring was conducted to whole-rock ⁸⁶Sr/⁸⁷Sr ratios from Mortimer et al. (1987). The MDC 264

and Mica-Mg isochrons were anchored to an initial 86 Sr/ 87 Sr ratio of 1.3773 \pm 0.0013 and calibrated to the published age of 519.4 \pm 6.5 Ma (Hogmalm et al., 2017; Redaa et al., 2021).

The summary of obtained inverse Rb–Sr dates is presented in Table 3. As shown, there is only marginal variation in the absolute biotite dates between the three analytical protocols involving Mica-Mg, either as the PRM for Rb/Sr ratios (protocols C & D) or as a MCRM (protocol B). Hence, in order to evaluate the accuracy of the obtained Rb–Sr dates against the expected references ages for each sample, we only compare the first two analytical protocols (NIST-610 as the PRM and either: (A) MDC or (B) Mica-Mg as MCRM).

273 Figure 4 compares the obtained Rb-Sr inverse isochron dates to the expected ages for the three 274 samples, that were analysed over two or three analytical sessions. The uncertainties for the K-275 feldspar dates are not shown as they are too large to be useful (due to the relatively low radiogenic 276 nature of typical K-feldspar versus micas), here we only compare the accuracy of the absolute 277 dates. As shown, analytical protocol (A) involving NIST-610 as PRM and MDC phlogopite as 278 MCRM consistently gives the most accurate Rb–Sr dates across all different analytical sessions. 279 For this analytical protocol, the Rb–Sr biotite dates for the Bundarra samples are 287.1 ± 2.4 Ma, 280 284.7 ± 3.0 Ma, 287.7 ± 2.3 Ma and 285.7 ± 2.6 Ma (between two samples over two analytical 281 sessions), which are in excellent agreement with the published zircon U-Pb age of 286.2 ± 2.2 Ma 282 (Black, 2007) from the same outcrop. The K-felspar dates of 290 ± 14 Ma, 285 ± 15 Ma, 290 ± 37 283 Ma and 288 ± 37 Ma are in excellent agreement as well but are less useful to evaluate age 284 accuracies given their larger uncertainties. Similarly for the Taratap sample, the obtained biotite 285 Rb–Sr dates of 499.4 ± 3.6 Ma and 495.7 ± 4.0 Ma as well as the (imprecise) K-feldspar Rb–Sr 286 dates of 500 ± 30 Ma, 501 ± 50 Ma and 495 ± 35 Ma are in excellent agreement with the zircon 287 U-Pb ID-TIMS age of 497.1 \pm 0.6 Ma as well as the apatite Lu-Hf age of 497.1 \pm 5.5 Ma for the

288 same sample (Glorie et al., 2023). Hence, the combined dataset suggests that the biotite, K-feldspar 289 and zircons record the same (crystallization) age for both the Bundarra and Taratap samples. The 290 Entire Creek biotite gave consistent Rb–Sr dates of 310.7 ± 1.5 Ma and 311.6 ± 3.1 Ma, in excellent 291 agreement with the ID-TIMS age of $312.1 \pm 1.8 / 5.1$ Ma (95% confidence uncertainties, without 292 / with overdispersion), based on the Rb/Sr ratios in Mortimer et al. (1987), recalculated with the 293 Villa et al. (2015) Rb–Sr decay constant.

294 5. Discussion

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5.1. Downhole fractionation corrections

296 Few previous studies have reported Rb-Sr DHF trends for a series of artificial reference materials 297 (i.e. glass standards and pressed pellets; Redaa et al., 2021; Wang et al., 2022). However, to the 298 best of our knowledge, DHF trends have not been evaluated for natural materials with the 299 exception of phlogopite MDC (Redaa et al., 2021). In our experiments, DHF is more pronounced 300 in natural micas and K-feldspar than observed for the NIST-610 glass and Mica-Mg pressed pellet, 301 when ablated under the same analytical conditions (Fig. 1). Comparatively, Mica-Mg appears least 302 appropriate to correct the analysed samples for DHF, given its systematically different DHF trend. 303 NIST-610 shows less DHF compared to the analysed micas and K-feldspars but its trend is more 304 systematic (similar shape with lower amplitudes). Thus, correcting for DHF against NIST-610 305 reduces the observed DHF for the analysed samples, while Mica-Mg significantly under-corrects 306 for DHF or accentuates it when applied to minerals. MDC biotite would be the most appropriate 307 choice for DHF correction as it behaves very similar to the analysed mica and K-feldspar samples. However, as with most natural materials, MDC is not sufficiently homogenous in ⁸⁷Rb/⁸⁷Sr ratio 308 309 to be used as a PRM. While the shape or slope of DHF trends can vary depending on laser 310 conditions (spot size, frequency and fluence), it cannot be eliminated for elements with contrasting 311 volatilities such as Rb and Sr. However, based on the presented data, the use of NIST-610 is the 312 more appropriate reference material to correct for DHF and Mica-Mg would exacerbate instead of 313 reduce the effects of DHF.

314 If no DHF correction is applied, accurate data can only be achieved if exactly the same signal 315 interval is selected in both the RM and sample. If there is a residual DHF slope on the sample 316 Rb/Sr ratios that is different to the RM (e.g. crystalline material versus Mica-Mg), then selecting 317 a shorter signal interval can significantly bias Rb/Sr ratios and hence the apparent age.

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319 5.2. Mica-Mg vs NIST-610 and MDC as calibration standards

320 **5.2.1.** Uncertainty comparisons

321 The contributions to the propagated uncertainties of individual analyses from the reference 322 materials (average signal precision and calibration curve misfit) are much larger when calibrating to Mica-Mg compared to NIST-610 for both ⁸⁷Rb/⁸⁷Sr and ⁸⁶Sr/⁸⁷Sr ratios (Fig. 5). For example, 323 in analytical session 1, the obtained uncertainties for individual ⁸⁷Rb/⁸⁷Sr ratios for the Entire 324 325 Creek biotite sample are more than double when using Mica-Mg compared to NIST-610 as the 326 PRM (Fig. 5). As a result, the choice of Mica-Mg instead of NIST-610 as the PRM will increase 327 the uncertainties on each analysis, and might consequently mask the presence of multiple data 328 populations. It will also introduces excessive uncertainties onto the calculated isochron dates.

The use of Mica-Mg as calibration standard for ⁸⁶Sr/⁸⁷Sr ratios most significantly affects the isochron precision of low-radiogenic samples such as K-feldspar samples. As shown in Table 3 and Figure 5, the uncertainty on the K-feldspar isochron dates can be up to $2 \times$ larger, compared to other calibration methods. Furthermore, the resulting MSWD values on the isochron regressions are consistently < 0.3 (Table 3), suggesting excessive uncertainties on individual data points. For the more radiogenic biotite samples, the larger uncertainties in ⁸⁶Sr/⁸⁷Sr ratios have negligible
effects to the precision on the isochron dates.

The precision of the calibrated ⁸⁷Rb/⁸⁷Sr ratios is more important to the isochron uncertainty of 336 337 highly radiogenic materials, such as most types of micas. Calibrating to NIST-610 versus Mica-338 Mg yields either more precise biotite isochron dates or identical precision. However, when NIST-339 610 is used as the PRM, uncertainty propagation from the MCRM (MDC or Mica-Mg) leads to 340 either identical or slightly worse isochron uncertainties compared to using Mica-Mg as PRM (Fig. 341 5; Table 3). The difference relates to the degree of overdispersion. The larger uncertainties on the 342 Rb/Sr ratios when using Mica-Mg as PRM result in lower MSWD values, reducing the uncertainty 343 on the isochron regression. This excess uncertainty when calibrating to Mica-Mg might mask 344 meaningful geological scatter in Rb/Sr ratios and it is, therefore, advisable to produce isochrons 345 based on data with the best possible analytical precision.

In summary, Mica-Mg should not be used as calibration standard for ⁸⁶Sr/⁸⁷Sr ratio calculations 346 347 for low-radiogenic samples as it introduces excessive uncertainties to age calculations. For high-348 radiogenic samples, using Mica-Mg as the PRM also introduces larger uncertainties to individual 349 data points compared to using NIST-610, but there is no significant difference in propagated 350 uncertainty after secondary correction to either MDC or Mica-Mg. For Rb/Sr ratio calibrations, 351 NIST-610 is more consistent, resulting in lower uncertainties on individual Rb/Sr ratios. When 352 there is no overdispersion, this results in better isochron age precision. However, overdispersion 353 can be masked by the increased uncertainties on Rb/Sr ratios, resulting in better apparent precision 354 when data are calibrated to Mica-Mg.

355 5.2.2. Accuracy comparisons

356 It has been observed previously that Rb-Sr dates are offset from their expected ages when 357 calibrated to the NIST-610 reference material (e.g. Gorojovsky and Alard, 2020; Wang et al., 358 2022). In contrast, Mica-Mg seems to better reproduce expected ages, although the significant 359 uncertainties obtained for natural materials in previous studies render appropriate accuracy testing 360 difficult. For example, Wang et al. (2022) compares measured to expected Rb–Sr dates for three 361 samples with known ages. The best achieved uncertainty in their experiment was $\sim 2.6\%$ for one 362 sample, while for the other samples the reported uncertainties are \sim 5.6 and 6.3 %. Similarly, the 363 accuracy comparisons in Gorojovsky and Alard (2020) use the Monastery phlogopite, with a 364 precision of ~4% when calibrated to Mica-Mg. Both papers report data normalized to NIST-610 365 but do not apply a secondary correction for matrix-dependent fractionation.

366 For the biotites analysed in our study, the fully propagated 95% confidence interval uncertainties 367 ranges between 0.8 and 1.6% when calibrated to Mica-Mg and between 1.0% and 1.4% when 368 calibrated to NIST-610 and corrected to MDC (depending on the sample and analytical session; 369 Table 3), allowing for more detailed accuracy comparisons. Figure 4 illustrates that using NIST-370 610 and MDC as calibration reference materials produces the most accurate results, compared to 371 the expected references dates. For the biotite results, the obtained Rb-Sr dates are within 0.5% 372 accuracy compared to the expected ages. The K-feldspar dates are accurate within 1%, except for 373 session 2, where accuracy is within 1.5%. When the same data are calibrated against Mica-Mg 374 (either using NIST-610 as the PRM and Mica-Mg as MCRM, or directly using Mica-Mg as the 375 PRM), the results are significantly offset from their expected ages. For the biotite results calibrated 376 to Mica-Mg, accuracy is within 2% for sessions 2 and 3 and there is 5% age off-set in session 1. 377 For the K-felspars, the age offset is up to 2.5% in sessions 2 and 3 and 6% in session 1. While the 378 age offsets in sessions 2 and 3 might be regarded as `acceptable`, given the obtained precision, the 379 more significant inaccuracy in session 1 renders Mica-Mg to be less desirable as a PRM.

380 The difference in accuracy between session 1 and sessions 2 and 3 can be explained by the 381 difference in measured dates for the MDC and Mica-Mg reference materials, normalised to NIST-382 610. For sessions 2 and 3, MDC and Mica-Mg produced similar isochron dates (2.3 and 1.9% 383 difference respectively) (Table 3; Fig. 6). For session 1, however, MDC gives a significantly 384 different age (494 ± 4 Ma) compared to Mica-Mg (469 ± 4 Ma). These differences in accuracy (ca. 385 5 % in session 1 and ca. 2 % in sessions 2 and 3) are in line with the observed age off-sets between 386 the measured dates and reference dates for the biotite and K-feldspar samples, calibrated to Mica-387 Mg.

388

5.2.3. Long-term comparison between MDC and Mica-Mg as secondary calibration standards

391 Given that the accuracy of the Rb–Sr method appears to be significantly dependent on the applied 392 calibration reference materials, and that the measured Rb-Sr dates of these calibration standards 393 fluctuate significantly between analytical sessions when compared to NIST-610, the long-term 394 behaviour of the MDC and Mica-Mg reference materials needs to be evaluated. Figure 7 presents 395 2.5 years of measured Rb-Sr dates for MDC and Mica-Mg, both calibrated to NIST-610 as the 396 PRM. All data in this plot have been processed identically. The Rb-Sr dates for Mica-Mg are 397 generally more consistent, ranging between ca. 462 and 479 Ma, with a standard deviation of 4.5 398 Ma, while the MDC dates show more variation, ranging between ca. 465 and 494 Ma, with a 399 standard deviation of 7.7 Ma. In all but two sessions, MDC produces an older Rb–Sr date compared 400 to Mica-Mg. The analytical sessions discussed above are highlighted in Figure F and encompass

401 the maximum variability in measured Rb–Sr dates for MDC. With the premise that calibration to 402 NIST-610 and MDC produces accurate Rb-Sr dates (as discussed in section 5.2.2), the difference 403 between the measured MDC and Mica-Mg dates (Fig. 6, 7) can be regarded as an estimate of the 404 degree of inaccuracy when data are calibrated to Mica-Mg. While some sessions reveal very little 405 off-set between both standards, using Mica-Mg as calibration standard can lead to up to 5% 406 inaccuracy in Rb–Sr dates. The cause of the observed variability is currently unknown, however, 407 in the second-to-last session with a significantly older Mica-Mg date compared to MDC, the 408 analysed samples might have received a lower effective laser fluence compared to other sessions 409 as the glass between the laser beam and samples was not cleaned prior to analysis. The lower 410 fluence could change the effective matrix bias between NIST-610, Mica-Mg and MDC, however, 411 calibration of biotite against MDC produces accurate results as demonstrated in section 5.2.2. In 412 contrast, although Mica-Mg produces more consistent Rb-Sr dates between analytical sessions, 413 these dates are unreliable given the variable and unsystematic degree of inaccuracy between 414 sessions.

415

416 **6.** Conclusions

417 Based on our observations, the use of Mica-Mg as calibration reference material is not418 recommended, for the following reasons:

(1) The down-hole fractionation (DHF) trend for Mica-Mg is not comparable with the DHF
trends of natural bitote, phlogopite and K-feldspar (Fig. 1). Using Mica-Mg to correct DHF
would exacerbate instead of reduce DHF in those minerals;

422 (2) Given the relatively poor reproducibility of ⁸⁷Rb/⁸⁷Sr ratios and significant uncertainty on
423 individual ⁸⁷Sr/⁸⁶Sr measurements (Fig. 2), Mica-Mg as PRM or MCRM introduces excess
424 uncertainty that can be avoided using a more consistent PRM such as NIST-610;

425 (3) We demonstrated that calibrating to Mica-Mg may lead to up to 5% inaccuracy in Rb–Sr

426 age (Fig. 4, 6, 7) and that the degree of inaccuracy is unsystematically session-dependant.

We suggest a different approach, involving (1) calibration of the ⁸⁷Rb/⁸⁷Sr and ⁸⁷Sr/⁸⁶Sr ratios to a 427 428 primary reference material with high Rb and Sr concentrations and homogenous isotopic ratios 429 such as NIST-610 glass, including DHF correction of the Rb/Sr ratios, followed by (2) a correction of the ⁸⁷Rb/⁸⁷Sr ratio to a natural mineral MCRM with a similar DHF trend as the samples to be 430 431 analysed. In our observations with a 67µm spot-size, there are no significant differences in matrix 432 effects comparing biotite, phlogopite and K-feldspar, suggesting that any of these natural minerals 433 as MCRM can produce accurate dates for K-rich minerals. We have used MDC phlogopite as 434 MCRM and demonstrate accurate Rb-Sr dates for a range of biotites and K-feldspars with well-435 established age constraints. For the biotite dates, the fully propagated uncertainties are <1.5 %, 436 allowing accuracy verifications at high analytical precision. The K-feldspar dates have relative 437 high uncertainties (ca. 5-10%) and, therefore, the accuracy of the calibration cannot robustly be 438 tested. However, absolute values agree with biotite dates and for a given sample, biotite and K-439 feldspar analyses statistically constitute a single isochron.

440 Finally, while this two-step calibration protocol is currently recommended due to current 441 constraints with data processing software, new developments involving calibrating to isochronous 442 reference materials might become the desired approach in the future.

443

444 Data availability

445 The Rb–Sr dataset used in this manuscript is freely available on figshare at 446 https://doi.org/10.25909/23996484.

447 448

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453

454

455 Author contributions

- 456 SG: Conceptualization, investigation, writing original draft, methodology, funding acquisition,
 457 visualisation, formal analysis
- 458 SEG: Conceptualization, investigation, writing review and editing, methodology
- 459 MH: Conceptualization, investigation, writing review and editing, resources
- 460 JCL: Conceptualization, investigation, writing review and editing, formal analysis
- 461

462 **Competing interests**

- 463 The authors declare that they have no conflict of interest.
- 464
- 465 **Ethical statement**
- 466 This manuscript is an original work that is not submitted or published elsewhere.

468 Tables

	React. gas (ml.min ⁻¹)	Laser wavel. (nm)	Fluence (J.cm ⁻²)	Rep. rate (Hz)	Spot (µm)	Rb-Sr calibration	Sr-Sr calibration	DHF	Err. corr.
Zack and Hogmalm 2016	O ₂ (0.25)	213	7	10	80	Pl: NIST-610; Ksp: BCR-2G; Bt: La Posta	NIST-610	No	No
Hogmalm et al. 2017	$\begin{array}{c} O_2 \ (0.25) \\ N_2 O \ (0.16) \\ SF_6 \ (0.04) \end{array}$	213	O ₂ : 7 N ₂ O: 6-8 SF ₆ : 6-8	10 4-5 10	80 50 50	Mica-Mg	NIST-610	No	No
Tillberg et al. 2020	N2O (?)	213	?	?	50	BCR-2G (Sec: Mica- Mg/La Posta)	NIST-610	No	Yes
Rösel and Zack 2022	N ₂ O (0.18- 0.20)	213	5-7	10	50- 60	Mica-Mg (sec: NIST- 610 / BCR-2G)	Mica-Mg	No	No
Gorojovsky and Alard 2020	N ₂ O (0.25)	193 and 213	7.8	5	85	Mica-Mg	Mica-Mg, NIST-610, BHVO-2G	No	No?
Larson et al 2023	N ₂ O (0.37)	193	4	10	50	Mica-Mg (Sec: Mica-Fe)	NIST-610	Yes?	Yes
Laureijs et al. 2021	CH ₃ F (10%)	213	6	10	50	ATHO-G, T1- G, StHs6/80-G	NIST-612	No	Yes
Li et al. 2020	N ₂ O (0.35)	193	3.5	5	74	Mica-Mg Sec: MDC	Mica-Mg	No	No
Liebmann et al. 2022	N ₂ O (?)	193	2.5	5	64	NIST-610 + Mica-Mg Sec: CK001 bt	NIST-610	No	Yes
Olierook et al. 2020	N ₂ O (0.25)	193	2.5	5	64- 87	NIST-610 + Mica-Mg Sec: CK001 bt	NIST-610	No	No
Redaa et al. 2021	N ₂ O (0.37)	193	3.5	5	74	Mica-Mg Sec: MDC	Mica-Mg	mon itore d	No
Sengun et al. 2019	N2O (?)	213	5.7	10	50	Mica-Mg	NIST-610	No	No
Tilberg et al. 2021	N ₂ O (?)	213	?	?	50	Mica-Mg / NIST 610	NIST-610	No	Yes
Wang et al. 2022	N ₂ O (0.25)	193	7	5	85	Mica-Mg	NIST-610, BHVO- 2G, BCR-2G	mon itore d	No
Kirkland et al. 2023	N ₂ O (0.25)	193	2	5	64	Mica-Mg Sec: CK001 bt	NIST-610	No	No

- 469 **Table 1:** Summary of published analytical conditions and protocols for LA-ICP-MS/MS Rb–Sr
- 470 dating. Rep. rate = laser repetition rate; Sec = secondary reference material; Bt = biotite; ksp = K-
- 471 feldspar; Pl = plagioclase; Err. Corr. = error correlation calculated (in most cases based on
- 472 calculated uncertainties after data-reduction rather than during data-reduction); In case of method
- 473 development work `best conditions` are quoted.

Laboratory Laser ablation system ICP-MS instrument Analytical method – materials Sample preparation Plasma Settings	Adelaide Microscopy – The University of Adelaide RESOlution-LR ArF excimer laser Agilent 8900x ICP-MS/MS <i>in situ</i> Rb-Sr - biotite, K-feldspar, plagioclase (anchor), apatite (anchor) mineral separates in 1-inch resin mounts and thin sections
-	1350 W
RF power Sample Depth	5.0 mm
	0.89 L/min
Ar carrier gas	0.38 L/min
He carrier gas N2 addition	4 mL/min
Lens Parameters	4 IIIL/IIIII
	1 5 37
Extract 1	1.5 V
Extract 2	-80 V
Omega Bias	-85 V
Omega Lens	5.0 V
Q1 entrance	-10 V
Q1 exit	-2.0 V
Cell focus	-2.0 V
Cell Entrance	-90 V
Cell Exit	-120 V
Deflect	10 V
Plate Bias	-80 V
Q1 bias	-2.0 V
Q1 Prefilter Bias	-10.0 V
Q1 Postfilter Bias	-10.0 V
N2O gas flow	0.37 mL/min
Octopole bias	-6.0 V
Axial Acceleration	2.0 V
Octopole RF	180 V
Energy Discrimination	-7.0 V
Analysis Parameters	
Laser Wavelength	193 nm
Laser fluence	3.5 J/m2
Sample laser diameter	67 μm
Laser repetition rate	5 Hz
Background duration	30 s
Analysis duration	40 s
Isotopes measured & dwell times (ms) 475	²³ Na (2), ²⁴ Mg (2), ²⁷ Al (2), ²⁹⁺¹⁶ Si (2), ³¹⁺¹⁶ P (2), ³⁹ K (2), ⁴³⁺¹⁶ Ca (2), ⁵⁵ Mn (2), ⁵⁶⁺¹⁶ Fe (2), ⁸⁵ Rb (10), ⁸⁶⁺¹⁶ Sr (50), ⁸⁷⁺¹⁶ Sr (50), ⁸⁸⁺¹⁶ Sr (50), ⁸⁹⁺¹⁶ Y (5), ⁹⁰⁺³² Zr (5), ⁹³⁺³² Nb (5), all ^{x+16} REE (5), ²³²⁺¹⁵ Th (5), ²³⁸⁺¹⁶ U (5)

Table 2: Analytical conditions for the three LA-ICP-MS/MS sessions in this paper.

			(A) NIST-610 + MDC		(B) NIST610 + Mica-Mg		(C) Mica-Mg & NIST-610		(D) Mica-M	g
Sample	S	n	Age	MS	Age	MS	Age	MS	Age	MS
(exp. age)			(± 2σ) [Ma]	WD	(± 2σ) [Ma]	WD	(± 2σ) [Ma]	WD	(± 2σ) [Ma]	WD
Ent Crk Bt	1	24	310.7 ± 1.5/2.5/3.1	1.1	327.8 ± 1.7/2.7/3.2	0.96	327.6 ± 3.3/3.9	0.27	328.8 ± 3.4/4.0	0.25
(312.1 ± 1.8^{1})	3	20	311.6 ± 3.1/3.8/4.5	2.5	317.6 ± 3.2/3.8/4.6	2.4	316.1 ± 3.2/3.8	0.85	316.2 ± 3.2/3.8	0.84
Bund1b Bt	2	44	287.1 ± 1.6/2.4/3.4	1.6	280.3 ± 1.5/2.4/3.2	1.7	280.1 ± 1.6/2.4	0.97	280.2 ± 1.6/2.4	0.82
(286.2 ± 2.2^2)	3	22	284.7 ± 2.4/3.0/3.8	1.0	290.1 ± 2.5/3.1/3.8	1.1	288.4 ± 4.1/4.5	0.7	288.2 ± 4.3/4.7	0.28
Bund1b Ksp	2	57	290 ± 14/14/14	0.87	284 ± 14/14/14	0.88	284 ± 14/14	0.84	280 ± 23/23	0.3
(286.2 ± 2.2^2)	3	53	287 ± 15/15/15	0.88	292 ± 15/15/15	0.88	290 ± 15/16	0.86	283.4 ± 38/38	0.19
Bund6a Bt	2	38	287.7 ± 1.3/2.3/3.4	1.4	280.9 ± 1.3/2.2/3.1	1.5	279.5 ± 1.5/2.3	0.71	279.5 ± 1.5/2.3	0.7
(286.2 ± 2.2^2)	3	22	285.7 ± 1.9/2.6/3.4	0.74	291.2 ± 1.9/2.7/3.6	0.72	288.7 ± 3.5/4.0	0.54	288.8 ± 3.6/4.0	0.34
Bund6a Ksp	2	45	290 ± 37/37/37	0.69	283 ± 36/36/36	0.69	281 ± 36/36	0.68	283 ± 75/75	0.16
(286.2 ± 2.2^2)	3	40	288 ± 37/37/37	0.65	293 ± 38/38/38	0.65	294 ± 39/39	0.65	296 ± 93/93	0.11
Taratap Bt	2	30	499.4 ± 1.8/3.6/5.6	1.2	487.7 ± 1.7/3.5/5.2	1.2	489.6 ± 2.8/4.2	0.67	489.6 ± 2.8/4.2	0.63
(497.1 ± 0.6^3)	3	16	495.7 ± 2.5/4.0/5.5	1.2	505.1 ± 2.6/4.1/5.8	1.2	504.0 ± 5.4/6.3	0.51	504.0 ± 5.5/6.3	0.52
Taratap Ksp	1	54	500 ± 30/30/30	0.53	527 ± 31/31/31	0.53	527 ± 32/32	0.50	539 ± 58/58	0.14
(497.1 ± 0.6^3)	2	20	501 ± 50/50/50	0.58	490 ± 49/49/49	0.58	492 ± 50/50	0.56	494 ± 106/106	0.12
	3	18	495 ± 35/35/35	0.95	504 ± 36/36/36	0.95	490 ± 39/39	1.1	511 ± 63	0.3

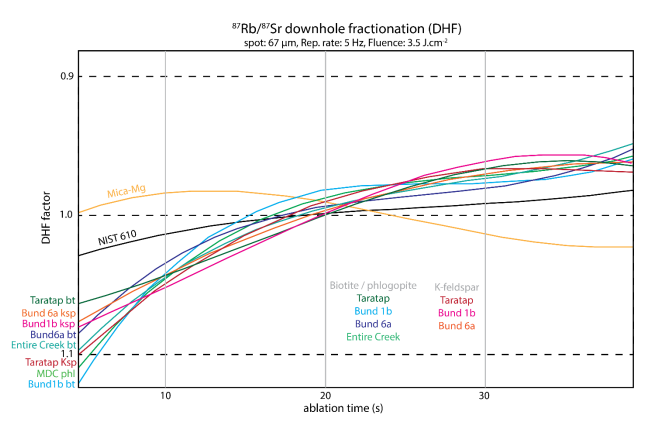
478

			NIST-610 as PRM		
MCRM	S	n	Age	MS	
			(± 2σ) [Ma]	WD	
MDC	1	34	494.4 ± 3.0	1.4	
	2	21	464.5 ± 4.0	1.3	
	3	30	470.6 ± 3.6	1.1	
Mica-Mg	1	35	468.6 ± 2.5	2.8	
	2	21	475.7 ± 3.7	3.5	
	3	20	461.8 ± 3.7	2.7	

481	Table 3: Summary table of Rb–Sr dates obtained in this study. $S =$ session number, $n =$ number of analysed grains, exp. age = expected
482	reference age (see below). All uncertainties are 95% confidence intervals and are reported as (1) excluding external uncertainty (on the
483	decay constant) / (2) including external uncertainties / (3) with propagated uncertainty from the correction standard (for methods (A)
484	and (B) only). (A) NIST-610 as PRM and MDC as MCRM to calibrate Rb/Sr ratios; (B) NIST-610 as PRM, Mica-Mg as MCRM to
485	calibrate Rb/Sr ratios; (C) Rb/Sr ratios calibrated to Mica-Mg as PRM and Sr/Sr ratios calibrated to NIST-610 as PRM; (D) Mica-MG
486	as PRM for both Rb/Sr and Sr/Sr ratios. ¹ Rb-Sr TIMS age from Mortimer et al. (1987), recalculated with Villa et al. (2015) decay
487	constant in IsoplotR (Vermeesch, 2018). The reported uncertainty is 95% confidence interval but does not take overdispersion into
488	account. ² Zircon U-Pb age for the Banalasta Adamellite in the Bundarra Suite, from Black (2007). ³ Zircon U-Pb TIMS age for the
489	Taratap Granodiorite, reported in Glorie et al. (2023).

491 Figures

492



494 **Figure 1:** ⁸⁷Rb/⁸⁷Sr downhole fractionation profiles for the analysed reference materials Mica-Mg

496 (red-pink lines) samples in analytical session 3, calculated in LADR (Norris and Danyushevsky,

(yellow line) and NIST-610 (black line), the biotite / phlogopite (green-blue lines) and K-felspar

497 2018). The DHF factor is calculated relative to the average ratio of the ablation signal (i.e. DHF

- 498 factor of $1 = \text{average } {}^{87}\text{Rb}/{}^{87}\text{Sr}$ ratio of the downhole signal).
- 499

493

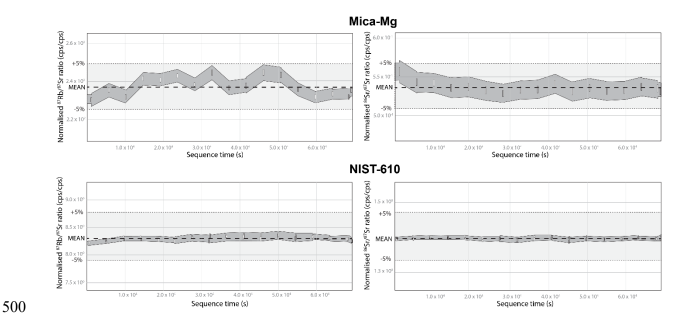
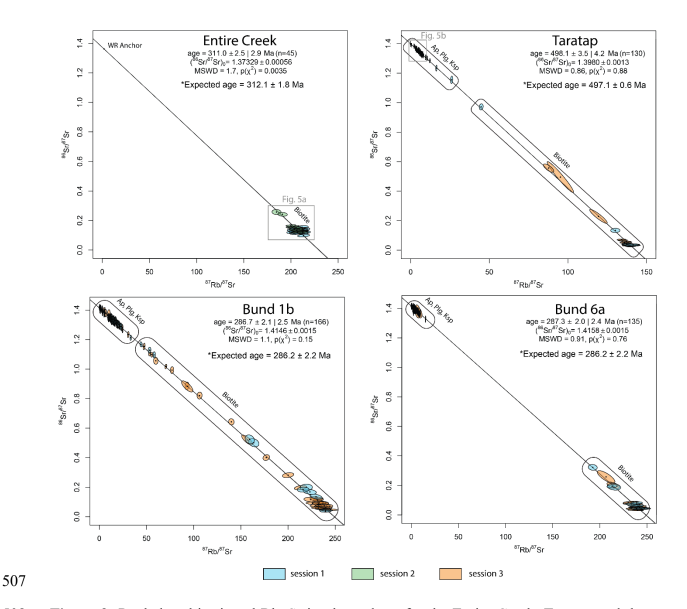


Figure 2: Variability of the 87 Rb/ 87 Sr and 86 Sr/ 87 Sr ratios for the analysed reference materials NIST-610 and Mica-Mg over the total duration of analytical session 3 (prior to drift correction). All plots are scaled equally to \pm 5% variation of the mean to aid visual comparisons. The vertical bars are \pm 1 standard deviation. The gray envelopes models \pm 2 standard deviation (note that for NIST-610 each standard was measured twice at each standard bracket).



508 Figure 3: Pooled multi-mineral Rb–Sr isochron dates for the Entire Creek, Taratap and the two 509 Bundarra samples (Bund 1b and Bund 6a). The data were calibrated against NIST-610 as PRM 510 and MDC as MCRM (see text for details). The colour-code refers to the analytical session in which 511 the data was obtained. Biotite analyses plot towards the radiogenic lower-intercept of the inverse 512 isochrons, while feldspar and apatite anchor Rb/Sr ratios plot towards the low-radiogenic end of 513 the isochron regression. All plots were calculated in IsoplotR (Vermeesch, 2018), reporting 95% 514 confidence interval uncertainties (including the uncertainty on the decay-constant) with and 515 without propagated uncertainty from the MDC MCRM. Expected ages are the recalculated Rb-Sr

age from Mortimer et al. (1987) with the Villa et al. (2015) decay constant for the Entire Creek
sample; the zircon U-Pb ID-TIMS age reported in Glorie et al. (2023) for the Taratap sample, and
the Zircon SHRIMP U-Pb age from Black (2007) for the Bundarra samples (see text for further
details).

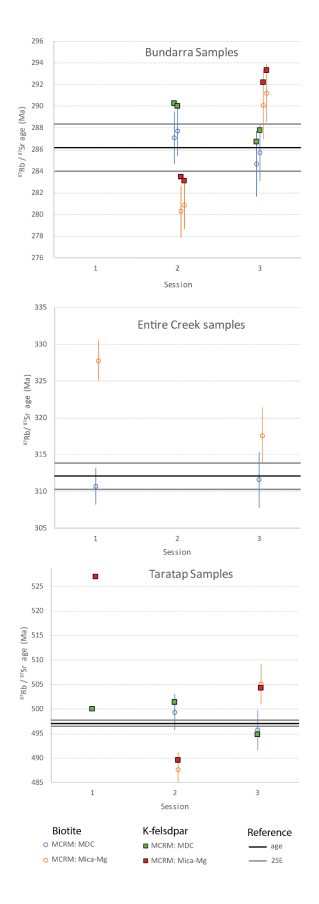
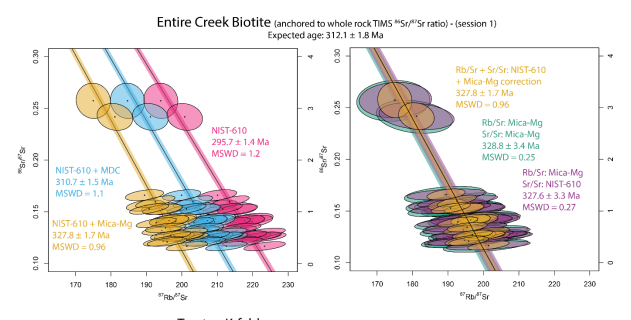
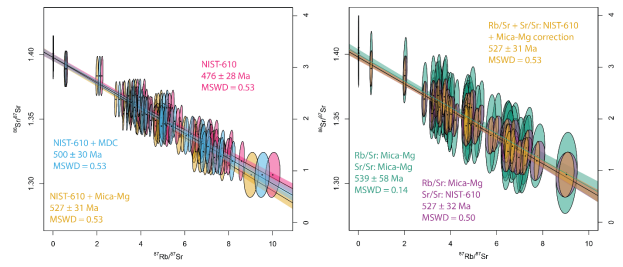


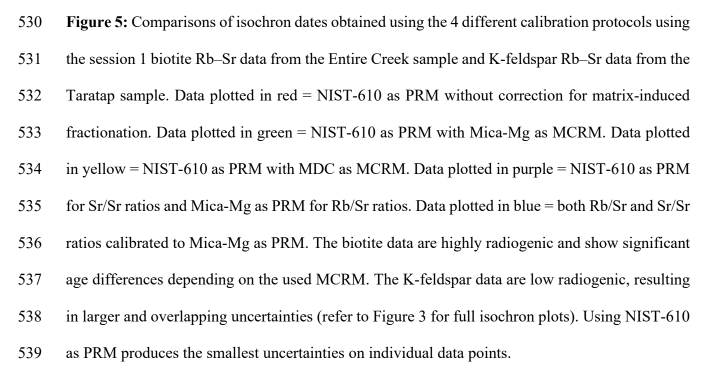
Figure 4: Comparisons of Rb–Sr dates over three analytical sessions, calibrated to either MDC or Mica-Mg as MCRM, with respect to the expected ages for each sample (black line with gray 2SE uncertainty bars). In all cases, NIST-610 was used as PRM. Biotite data are plotted as open circles (blue = calibrated to MDC as MCRM, orange = calibrated to Mica-Mg as MCRM). K-feldspar data are plotted as filled squares (green = calibrated to MDC as MCRM, red = calibrated to Mica-Mg as MCRM).





Taratap K-feldspar (anchored to plagioclase + apatite) - (session 1) Expected age: 497.1 ± 0.6 Ma







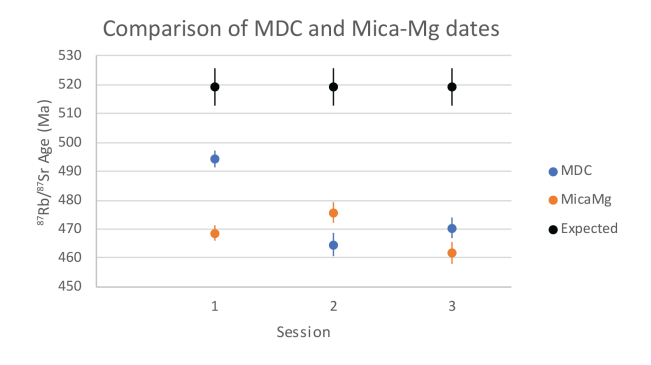
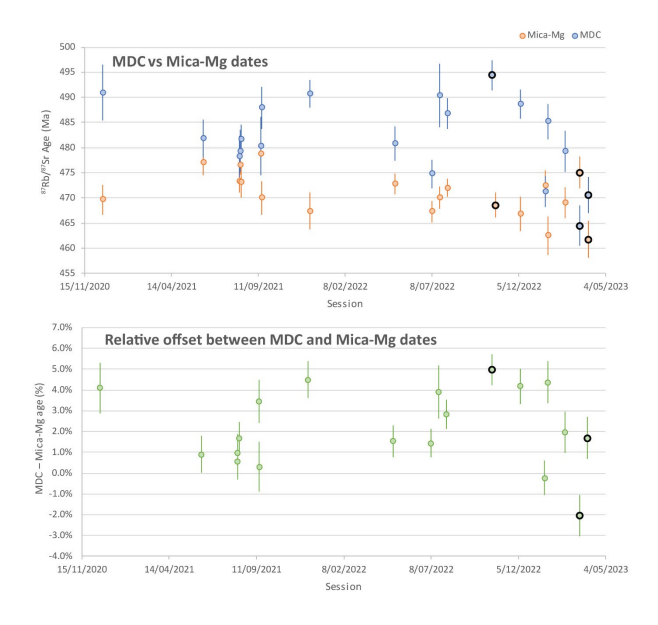


Figure 6: Rb–Sr dates for MDC and Mica-Mg calibrated to NIST-610 over the three analytical
sessions used in this paper. The off-set of the Rb–Sr age with respect to the reference age is used
to calculate the correction factor on the Rb/Sr ratios. Uncertainties are 2SE.

545



547 Figure 7: Long-term (2.5 years) Rb–Sr age data for Mica-Mg and MDC for the lab (Adelaide 548 Microscopy). All uncertainties are 2SE. The top plot shows absolute dates and the bottom plot 549 shows the percentage difference between the MDC and Mica-Mg dates. All data were processed

- 550 in the same way using NIST-610 as PRM. The three analytical sessions previously discussed are
- 551 highlighted by black rims and capture the most extreme differences obtained in our lab to date.
- 552 Given that MDC as MCRM produces consistently accurate data, the plot indicates that Mica-Mg
- as PRM can lead to up to 5% inaccuracy in Rb–Sr age calculations.
- 554

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