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LA-ICP-MS/MS-based Rb–Sr isotope mapping for geochronology

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This study introduces a new approach for *in situ* Rb–Sr dating that utilizes rapid line scans instead of static spot ablation, enabling the creation of two-dimensional $^{87}\text{Rb}/^{86}\text{Sr}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ isotope ratio and Rb–Sr age maps. The data acquisition is conducted utilizing an ICP-MS/MS instrument with N_2O as the reaction gas, coupled to a 193 nm excimer laser *via* a low-aerosol-dispersion interface. This configuration allows for high repetition rates (>100 Hz) and sensitivities, enabling data acquisition at a high scanning speed and small laser beam size (3–4 μm). Notably, this approach requires just about 1/30 of the sample volume typically utilized in conventional spot ablation mode, while achieving similar levels of precision and accuracy. Line scan ablation is tested and compared to spot ablation on age-homogeneous crystalline muscovite and biotite, for which reference Rb–Sr age data is acquired through ID-TIMS. Results show that a key requirement for accurate Rb–Sr ages based on line scan analyses is matrix correction using chemically matched crystalline mica. By presenting Rb–Sr age maps of three naturally deformed mica samples, we highlight the potential of Rb–Sr mapping for extracting age data from rocks that exhibit complex metamorphic-metasomatic histories and microscale dynamic recrystallization. Additionally, we show that quantitative elemental information (Al, Fe, Si, Li) can be collected alongside Rb–Sr isotope data. This advancement offers a distinctly more insightful assessment of isotope mobility in natural systems, the timing of element enrichment processes and enables, in high-Rb/Sr rock systems, precise and accurate isotopic dating of intricate geological processes at small scales.

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1. Introduction

Rb–Sr dating relies on the beta decay of ^{87}Rb to ^{87}Sr with an ^{87}Rb half-life of 49.61 ± 0.16 billion years.¹ This decay system has been discovered already in 1937, and was later extensively used for isotopic dating of almost any kind of Rb-bearing geologic materials. The attractiveness of the Rb–Sr geochronometer lies in its wide applicability, as both elements occur in most crustal lithologies and are present in $\mu\text{g g}^{-1}$ -levels in the majority of rock forming minerals. The contrasting geochemical behavior of Rb and Sr often leads to highly contrasting Rb/Sr ratios between minerals of a given rock, and, in consequence, to large inter-mineral contrasts of Sr isotopic compositions that facilitate precise dating of geologic processes. Rb–Sr mineral data have successfully been used to date crystallization of magmatic rocks as well as high-grade metamorphic processes like eclogitization of subducted crust.^{2,3} Moreover, Rb–Sr dating can also provide age constraints on secondary events such as hydrothermal alteration, hydrothermal vein formation, or deformational events that can cause isotope mobility during fluid-mediated recrystallization or recrystallization as a response to

intracrystalline deformation during accommodation of strain in ductile shear zones.^{4–8} Rb–Sr isotopic dating mainly relies on minerals with high Rb/Sr elemental ratios. These minerals are typically K-rich phases, like the phlogopite–annite solid solution series (biotite), different types of potassic white mica, and K-feldspar. In the classic approach, either whole rock powders or mineral separates are dissolved, followed by wet-chemical separation of Rb and Sr from each other and from the matrix, to avoid the isobaric interference of ^{87}Rb and ^{87}Sr during mass spectrometric analysis of Sr isotopic compositions. Only recently laser ablation-ICP-MS-based analytical methodologies have been developed for *in situ* dating of Rb-rich mineral phases such as mica or K-feldspar.^{9–11}

Unlike the well-established *in situ* ^{40}Ar – ^{39}Ar dating technique, which focuses on the same minerals,^{12–14} *in situ* Rb–Sr dating offers several distinct advantages. Notably, it eliminates the need for access to irradiation facilities, which can be both time-consuming and costly, and involves handling radioactive materials. In contrast, *in situ* Rb–Sr dating is significantly less demanding in terms of instrumentation and sample preparation, making it a cost-effective and versatile option that can be implemented in various laboratory settings. Additionally, this method allows for a rapid sample turnaround, enhancing its potential as a widely accessible dating tool suitable for a diverse range of geological targets. The two essential components of

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suitable laboratory equipment comprise (1) a laser for *in situ* sampling of the target material and (2) a mass spectrometer capable of (quasi) simultaneously analysing a minimum of three nuclides: ^{85}Rb or ^{87}Rb , ^{87}Sr , and ^{86}Sr or ^{88}Sr throughout the sample ablation process. Notably, achieving the necessary mass resolution for effectively distinguishing between ^{87}Rb and ^{87}Sr ($M/\Delta M \sim 300\,000$) surpasses the capabilities of any commercially accessible mass spectrometer, so that the use of a collision/reaction cell (CRC) is required. Within the CRC, reactive gases (O_2 , SF_6 , N_2O , CH_3F) form Sr-molecule ions such as SrF^+ or SrO^+ that can be measured free of the ^{87}Rb -interference.^{9,15–18} Interference from mononuclide ions on mass shifted Sr-molecule ions (e.g. $^{104}\text{Pd}^+$ on $^{88}\text{Sr}^{16}\text{O}^+$) is in turn avoided by applying a pre-CRC mass filter (either quadrupole-based or through Wien filtering). This technique obviates the need for time-consuming offline chemical separation of Rb and Sr by classic cation-exchange chromatography. However, this approach currently reaches its limits when it comes to dating of geologically young (Cenozoic) assemblages or to K-phases with particularly low Rb/Sr ratios, like white mica populations rich in non-radiogenic Sr. Here, precise age information depends upon the extremely precise determination of Sr isotopic compositions not only for mica but also for accompanying mineral phases, a task that requires mass spectrometers featuring multi-collection (TIMS, MC-ICP-MS).

Traditionally, Rb–Sr ages are obtained by constructing multi-point isochrons through analyses of co-genetic phases with contrasting Rb/Sr ratios.¹⁹ It is also possible to obtain model ages with an accuracy and precision of 1–3% from single spot *in situ* analyses using laser ablation (LA), provided that either (1) the grains show high Rb/Sr ratios and correspondingly very high $^{87}\text{Sr}/^{86}\text{Sr}$ ratios or (2) a reasonable guess of the initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratio can be made.²⁰ Most recently, the isochron approach was applied to single spot dating using high-precision LA-MC-ICP-MS/MS analyses, which allows precise single spot dating (2–4%) without prior knowledge of the initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratio.²¹ So far, *in situ* Rb–Sr dating relies on placing rather large single laser spots (50–100 μm)^{20,21} on the phase of interest. To obtain a reasonable level of uncertainty for $^{87}\text{Rb}/^{86}\text{Sr}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (typically around 1–3% and 0.5–1% when using quadrupole instruments), the ablation interval is chosen long enough to enable the recording of multiple mass sweeps ($n > 100$). Due to the destructive nature of the ablation process, this results in ablation pits that are a few tens of μm deep. However, large ablations depths are problematic in fine grained rocks or when aiming to date small (μm scale) secondary features, like reaction rims on larger grains, local shear bands, or localized domains of fluid-induced recrystallization.

In this contribution, we present a novel technique that enables *in situ* Rb–Sr dating with minimal sample material extraction. This approach uses line rasters instead of conventional single spot ablation, resembling the well-established image-based dating techniques utilized in carbonate and zircon U–Pb dating.^{22–24} By applying a low-aerosol dispersion system, sampling at significantly greater repetition rates (>100 Hz) is enabled, so that large Rb–Sr maps can be produced from multiple line scans in a reasonable amount of time and, most

importantly, at μm -scale resolution and high precision and accuracy but with only a fraction of the material that is used for conventional spot dating. By using examples of naturally deformed mica we show that this technique opens up new avenues to study isotope mobility in nature, to derive precise and accurate ages from rocks showing textural disequilibria, and to date particularly small-scale features like localized dynamic recrystallization in shear bands at the micrometer scale.

2. Material and methods

2.1. Single crystal mica reference materials

To correct for elemental and mass-dependent isotopic fractionation, reference materials (RM) with a well-known and homogeneous $^{87}\text{Rb}/^{86}\text{Sr}$ and $^{87}\text{Sr}/^{86}\text{Sr}$ distribution are applied. For *in situ* Rb–Sr dating, NIST 610 glass²⁵ and MicaMg-NP^{9,26} are the most widely used.^{15,20,21,27,28} The latter is a finely ground pressed powder of phlogopite from a pyroxenite from southern Madagascar that displays a homogeneous age signal (519.4 ± 6.5 Ma).⁹ Although these RM are frequently used in geochronological applications, none provides a proper matrix match to single-crystal mica, which is also the case for the recently tested fused mica mineral glasses.²⁹ It has been shown that a lack of matrix match between RM and sample may negatively affect dating accuracy. Systematic matrix age biases of as much as 10% have been observed.^{29–31}

We therefore produced secondary reference materials with well-known ages, determined by isotope dilution thermal ionization mass spectrometry (ID-TIMS) at the GFZ Potsdam. The aim was to establish an additional set of mica reference materials with good matrix match to both natural K-white mica (muscovite–phengite) and natural K-(Mg, Fe^{2+}) solid solution series (phlogopite–annite, ‘biotite’)

2.1.1. Phalaborwa biotite. The here characterized biotite from Phalaborwa (23.99°S, 31.13E, RSA) is separated from a boulder of biotite apatite. The rock is coarse-grained, with crystals of apatite up to 5 cm and biotite up to 2 cm in diameter, has an igneous-pegmatitic appearance with absence of ductile deformation textures combined with presence of cm-sized rounded xenoliths of fine-grained, blackish mafic rock. The biotite is homogeneously reddish-brown, whereas the apatite is greenish and transparent, indicating that no post-crystallizational overprint occurred. The Rb–Sr age can thus be expected to be identical to the magmatic intrusion age of the Phalaborwa main phoscorite (2062 ± 2 Ma (ref. 32)).

2.1.2. Harvard 98973 muscovite. The Harvard 98973 Muscovite is a Li-rich muscovite from the mineral collection of the Mineralogical Museum, Harvard University, Cambridge, MA. It was recovered from a granitic pegmatite in the Lord Hill Quarry, Stoneham, Oxford Co., Maine, USA, is commercially available *via* IAGEO Ltd, serves as a reference material for light elements and is described in more detail elsewhere.³³ It has not yet been directly dated; however, for the regional pegmatite field ages between 250 and 270 Ma are envisaged.³⁴

2.1.3. SagaB biotite. The SagaB biotite is a crystal that was collected by one of the authors (JG) in 2017 in remnants of



a large, nepheline-bearing syenite pegmatite body in the former Saga 1 larvikite quarry (59.0439°N, 9.8289°E), Sagåsen, Porsgrunn, Norway. The pegmatite is extremely coarse grained, with dm-sized biotite crystals, and has, in places, gradational contacts to the host larvikite (an alkali monzonite). Formation of the pegmatite body is thus expected to be late-magmatic with respect to its host intrusion that is dated by U–Pb zircon and baddeleyite ages of 293.2 ± 1.3 and 294.3 ± 1.2 Ma, respectively.³⁵

2.2. Natural mica from metamorphic rocks

Rb–Sr age mapping was undertaken on natural muscovite and biotite from three locations (Fig. 1).

2.2.1. GÄD08. Sample GÄD08 is a mylonitic mica schist from the Middle Seve Nappe near Ankarede, Jämtland, Sweden (64.7923°N, 14.2912°E) with an assemblage comprised of muscovitic white mica, quartz, plagioclase, biotite, epidote and garnet. The muscovite-based Rb–Sr age for mylonitic ductile deformation is 431.7 ± 3.1 Ma.³⁶

2.2.2. GER2B. Sample GER2B is a metamorphosed, ductily deformed granitic pegmatite from the Western Bohemian Massif in NE Bavaria, Germany. Sampling location is the abandoned feldspar quarry ‘Gertrud’, 1.5 km NE of Obersdorf, 49.7752°N, 12.1288°E. Age estimates for the primary, magmatic crystallization are provided by a 476 ± 2 Ma garnet U–Pb age and 482 ± 5 Ma (recalculated using the revised ^{87}Rb decay constant⁴) TIMS-based Rb–Sr muscovite-feldspar age based on undeformed, cm sized, primary muscovite and feldspar crystals.⁵

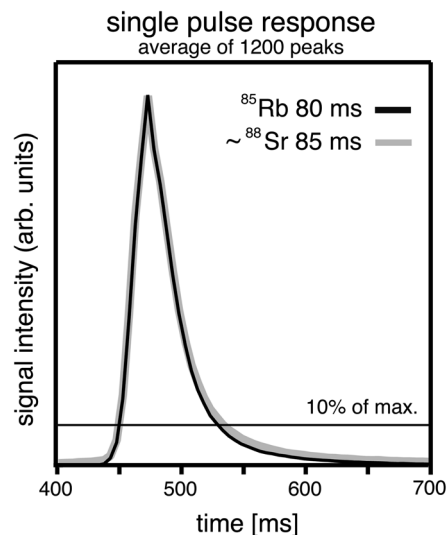


Fig. 2 Average intensity profile ($n = 1200$) of a single laser shot for mass channels $m/z = 85$ (Rb) and $m/z = 104$ ($\sim^{88}\text{Sr}$). The single pulse response (SPR) is defined as the full width at 10% of the maximum intensity and is similar for ^{85}Rb and $\sim^{87}\text{Sr}$. Signal intensities are normalized based on their integral.

2.2.3. WIL1A. Sample WIL1A originates from an abandoned feldspar mine (49.8010°N, 12.0439°E), ca. 1.5 km west of Wildenreuth, NE Bavaria, Germany. The sample is a metamorphosed, in parts ductily deformed granitic pegmatite. It originates from the same pegmatite field as sample GER2B. A previous, TIMS-based Rb–Sr age determination for primary, magmatic muscovite and cogenetic feldspar provided a (recalculated) age of 484 ± 5 Ma.⁵

2.3. LA-ICP-MS/MS isotope ratio, Rb–Sr age and elemental maps

Elemental and isotope ratio mapping on mica from mylonite and metapegmatites were performed at the MAGMA Lab of the TU Berlin during three sessions (GÄD08, WIL1A and GER2B, corresponding to the sample names) using an Agilent 8900 ICP-MS/MS coupled to an Analyte Excite 193 nm excimer laser (Teledyne Photon Machines). Prior to analysis, the reaction cell is flushed for at least 12 hours with N_2O (25%) to reduce instrumental drift.⁹ For introduction of N_2O into the Agilent 8900 CRC, the 4th cell gas mass flow controller is used, where a flow rate of 100% corresponds to ~ 1.07 ml min^{-1} . The aerosol rapid introduction system ARIS (Teledyne Photon Machines) was used in combination with a low-volume adapter from Glass Expansion. Helium (99.999%) was used as a carrier gas with a total flow rate of 0.95 l min^{-1} (0.5 l min^{-1} cell flow, 0.45 l min^{-1} cup flow) and small amounts (3 ml min^{-1}) of high-purity N_2 (99.999%) were admixed to increase sensitivity. After plasma ignition and initialization of He flow, $m/z = 42$ (e.g., $^{14}\text{N}^{14}\text{N}^{14}\text{N}^+$) and $m/z = 31$ (e.g., $^{15}\text{N}^{16}\text{O}^+$) were recorded to monitor the amount of air entrained in the interface tubing and ablation cell. After count rates were stable and below 30 000 cps for $m/z = 31$ and 42, the Ar nebulizer gas and He carrier gas

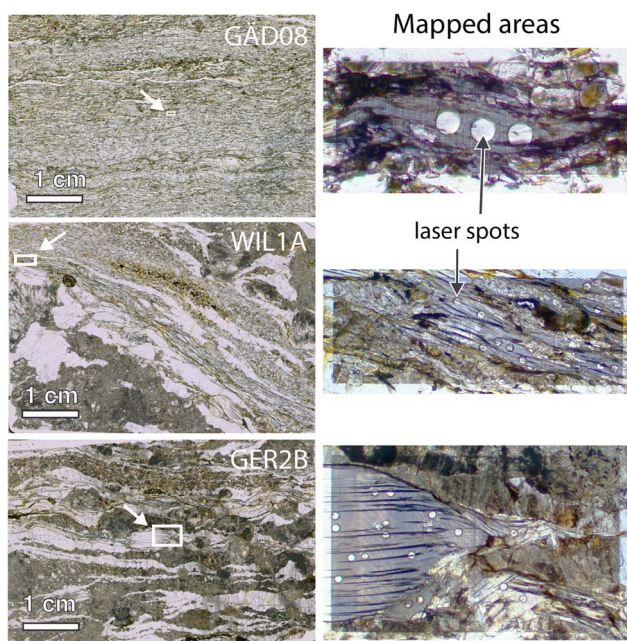


Fig. 1 Thin sections scans of the samples under investigation. GER2B and WIL1A are metapegmatites from the Western Bohemian Massif in NE Bavaria (Germany). GÄD08 is a mylonitic mica schist from the Middle Seve Nappe near Ankarede, Jämtland (Sweden). Selected regions for Rb–Sr data acquisition are shown as they appear after both line scan and spot mode analyses. Diameter of laser spots is 65 μm .



flows were tuned to achieve $m/z = 248/232$ ratios of $<0.1\%$ (ThO/Th) and $m/z = 232/238$ ratios of $100 \pm 1\%$ (Th/U), while ablating NIST 610 in line scans and in single quadrupole mode. An automatic lens tune was performed on NIST 610 to maximize sensitivity in the mid-mass range with $m/z = 88$ (Sr) as the sensitivity monitor. Laser parameters for the sensitivity and gas flow tuning were 50 Hz repetition rate, 5 μm circular spot size, and 4 J cm^{-2} fluence.

After sensitivity and carrier gas flow tuning, the ICP was switched to MS/MS mode. To resolve the isobaric interference of ^{87}Rb and ^{87}Sr , Sr isotopes were measured as SrO^+ molecular ions in mass shift mode using an N_2O -filled reaction cell. Throughout this paper we use the notation “ $\sim^y\text{X}$ ” with y being the atomic mass of element X to refer to a +16 oxygen mass shift of that particular isotope (e.g. $\sim^{88}\text{Sr}$, refers to $^{88}\text{Sr}^{16}\text{O}$). Flow rates of N_2O were tuned to yield maximum counts on $m/z = 104$ ($\sim^{88}\text{Sr}$) and were between 15 and 19%. Typically, the $\sim^{88}\text{Sr}/(^{88}\text{Sr} + \sim^{88}\text{Sr})$ ratio was 0.97. Counts on $m/z = 101$ ($\sim^{85}\text{Rb}$) were checked and suggested negligible formation of RbO^+ (typically 1 count per second). After a second sensitivity tune in MS/MS mode the single pulse response (SPR), *i.e.*, the combined wash-in and wash-out time, was determined by (1) ablating NIST 610 at 1 Hz and dosage 1 in line scan mode and recording

$m/z = 85$ (Rb) and $m/z = 104$ ($\sim^{88}\text{Sr}$) at a dwell time of 5 ms and (2) averaging the full width at 10% of the maximum of 1200 single pulses. Usually, ^{85}Rb and $\sim^{88}\text{Sr}$ yielded similar SPR (Fig. 2 and Table 1). The last tuning step was tuning of pulse-analog (P/A) factors to correct for the non-linear response across the pulse to analog switch of the detector, which appears between 1 and 2 million counts per second (cps) or, more precisely, at count rates of 10 million cps multiplied by the respective P/A factor (J. Hansmann, Agilent, personal communication). P/A tuning is done by switching the ICP from time-resolved analysis (TRA) to spectrum mode, opt-in “independent P/A factor” in the mass hunter configuration and “P/A factor adjustment” and then running several line scans with different spot diameter as calibration standards on NIST610. Resulting P/A factors were between ~ 0.15 and 0.17 , varied between sessions and also displayed a certain relative difference between Sr isotopes (typically 1–2% difference). It should be noted that for all mappings all $\sim^y\text{Sr}$ nuclides are acquired in pulse mode, whereas ^{85}Rb is acquired in pulse mode in NIST610, Phalaborwa biotite, GER2B, WIL1A and GÄD08 and in analog mode in SagaB biotite and Harvard 98973 muscovite (<https://doi.org/10.26434/chemrxiv-2023-273lw-v2>).

Table 1 Parameters for data acquisition during individual measurements sessions^a

Session	Line scan (mapping)			Spots				
	GÄD08	GER2B	WIL1A	GÄD08	GE2B/WIL1A	Homogeneity		
Date	22/03/2022	28/06/2022	29/06/2022	30/05/2023	08/08/2022	23/05/2022		
Spot size (μm)	3	4	3	65	65	65		
Fluence (J cm^{-2})	5.0	5.0	5.0	5.0	5.0	5.0		
He (l min^{-1})	0.95	0.95	0.95	0.95	0.95	0.95		
N_2 (ml min^{-1})	3.0	3.0	3.0	3.0	3.0	3.0		
SPR ^{85}Rb (ms)	136	80	80	—	—	—		
SPR $\sim^{88}\text{Sr}$ (ms)	112	85	85	—	—	—		
Dosage	10	10	10	—	—	—		
Repetition rate (Hz)	50	118	118	10 Hz	10 Hz	10 Hz		
Scan speed ($\mu\text{m s}^{-1}$)	15.0	47.2	35.4	—	—	—		
Mapping time (h)	1.5	8.0	7.0	60 s per shot	60 s per shot	60 s per shot		
Element	m/z Q1	m/z Q2	Dwell time (ms)		Dwell time (ms)			
Li	7	7	—	10	20	1	1	1
Na	23	23	—	—	—	1	1	1
Mg	24	24	—	—	—	1	1	1
Al	27	27	—	1	1	1	1	1
Si	28	60	1	1	1	1	1	1
K	39	39	—	—	—	1	1	1
Ca	44	44	—	—	—	1	1	1
Ti	47	63	—	—	—	1	1	1
Fe	57	57	—	1	1.8	1	1	1
Rb	85	85	10	10	10	50	50	50
Sr	86	102	—	—	—	70	70	100
Sr	87	103	109.7	35	24.6	70	70	100
Sr	88	104	10.4	10	10	70	70	—
Total sweep time (ms)			140.10	85.00	85.40	294	294	283

^a Dosage refers to the number of laser shots applied within the diameter of the laser beam during its vertical movement. Q1 and Q2 denote the quadrupoles before and after the collision-reaction cell, respectively.



Table 2 ICP-MS setting for each of the measurement sessions

Session	Line			Spots		Homogeneity
	GÄD08	GER2B	WIL1A	GÄD08	GER2B/WIL1A	
Plasma						
RF power (W)	1450	1400	1400	1400	1450	1400
RF matching (V)	0.86	0.88	0.90	0.86	0.87	0.87
Sample depth (mm)	6.0	6.0	6.0	6.0	6.0	5.0
Nebulizer gas (l min ⁻¹)	0.6	0.6	0.6	0.6	0.55	0.8
Plasma gas (l min ⁻¹)	15	15	15	15	15	15
Auxiliary gas (l min ⁻¹)	0.9	0.9	0.9	0.9	0.9	0.9
Lenses						
Extract 1 (V)	-11.5	-12.0	-12.0	-9.6	-12.0	-12.0
Extract 2 (V)	-250	-250	-250	-245	-250	-240
Omega bias (V)	-135	-130	-145	-140	-145	-140
Omega lens (V)	6.8	7.0	6.2	7.4	6.2	6.8
Q1 entrance (V)	-50	-50	-50	-50	-50	-50
Q1 exit (V)	0.0	-1.0	0.0	0.0	0.0	0.0
Cell focus (V)	2.0	2.0	2.0	3.0	4.0	4.0
Cell entrance (V)	-40	-40	-40	-40	-40	-40
Cell exit (V)	-60	-60	-60	-60	-60	-60
Deflect (V)	-1.2	-0.8	-0.8	-0.8	-0.4	-0.8
Plate bias (V)	-50	-50	-50	-50	-50	-50
Q1						
Q1 bias (V)	-1.0	-2.0	-2.0	-2.0	-1.0	-1.0
Q1 prefilter bias (V)	-5.5	-9.5	-6.0	-8.5	-6.0	-5.5
Q1 postfilter bias (V)	-9.0	-7.0	-9.0	-3.0	-9.0	-5.0
Cell						
N ₂ O gas flow (%)	15	19	19	18	14	15
OctP bias (V)	-8.0	-8.0	-8.0	-8.0	-8.0	-8.0
OctP RF (V)	140	150	180	180	180	180
Axial acceleration (V)	2.00	2.0	2.0	2.0	2.0	2.0
Energy discrimination (V)	-7.0	-7.0	-7.0	-7.0	-7.0	-7.0

Table 3 Elemental and isotopic homogeneity of the single crystal mica reference materials^a

	Phalaborwa biotite (n = 29)		SagaB biotite (n = 28)		Harvard 98973 (n = 30)		Dyar <i>et al.</i> ³³ (2001)	LOD (ppm)
	Mean	2s	Mean	2s	Mean	2s		
SiO ₂ (wt%)	42.28	0.40	38.44	0.28	45.80	0.22	45.67 (0.28)	1.02
Al ₂ O ₃ (wt%)	7.66	1.60	11.56	0.36	32.83	0.38	30.58 (0.34)	0.05
TiO ₂ (wt%)	0.70	8.52	2.98	0.78	0.03	9.50	0.08 (0.02)	0.13
FeO (wt%)	10.80	1.98	27.41	0.92	4.91	2.18	6.65 (0.48)	0.85
MgO (wt%)	22.27	1.12	4.38	1.02	0.01	4.08	0.02 (0.02)	0.04
CaO (wt%)	<LOD		0.07	53.20	<LOD		0.00 (0.00)	151
Na ₂ O (wt%)	0.037	5.40	0.42	6.70	0.52	3.50	0.38 (0.08)	0.29
K ₂ O (wt%)	11.26	1.52	9.75	0.70	10.90	0.34	10.14 (0.17)	0.70
Li (µg g ⁻¹)	9.77	7.10	0.44	2.96	1886	6.68	1998 (18.61)	0.15
Rb (µg g ⁻¹)	791.8	1.16	3862	0.58	3826	1.20		0.0070
⁸⁶ Sr (µg g ⁻¹)	0.023	28.38	0.073	7.30	0.009	10.02		0.00004
⁸⁷ Sr (µg g ⁻¹)	5.83	0.92	4.069	0.66	3.600	1.14		0.00008
⁸⁷ Sr/ ⁸⁷ Rb	0.02596	0.39	0.00379	0.32	0.00341	0.33		
Expected age (Ma)	2062 ± 2		295.4 ± 1.4		266.8 ± 1.6			

^a Major element composition for Harvard 98973 data of Dyar *et al.* (2001)³³ are from GFZ electron microprobe session, except Li₂O, which is from the PIGE session. The limit of detection (LOD) for LA-ICP-MS data is calculated after Longerich *et al.* (1996).⁴⁵ Elemental and isotopic concentrations normalized to SiO₂ + Al₂O₃ + TiO₂ + FeO + MgO + CaO + Na₂O + K₂O = 95%. Expected age of Phalaborwa biotite is from Wu *et al.* (2011).³² SagaB and Harvard 98973 expected ages are based on ID-TIMS analyses presented in this study.



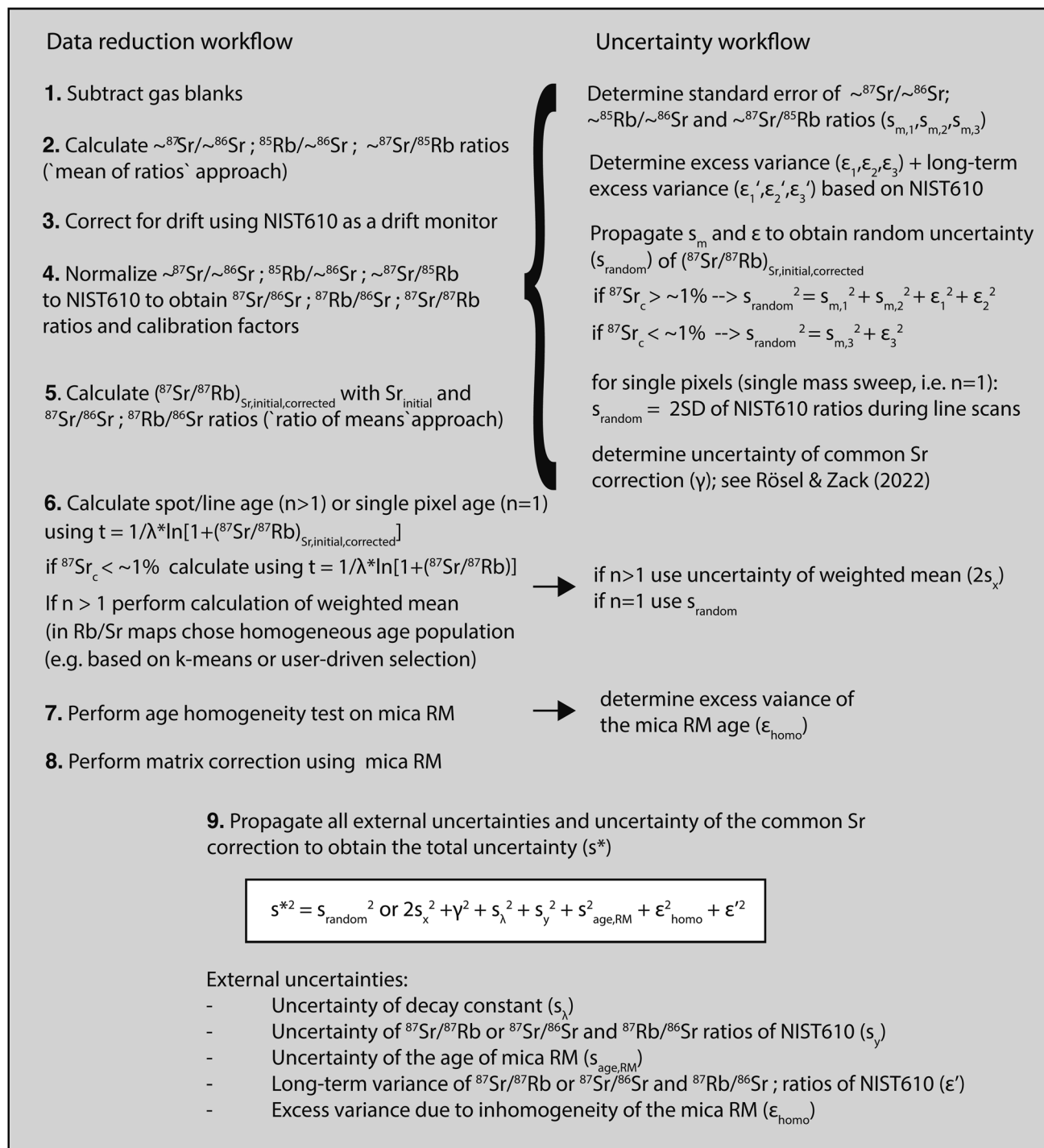


Fig. 3 Schematic illustration of the data reduction and uncertainty propagation workflow. For further information see text.

For mapping sessions, the following mass channels were recorded: $m/z = 7$ (Li), $m/z = 27$ (Al), $m/z = 60$ ($\sim^{28}\text{Si}$), $m/z = 57$ (Fe), $m/z = 85$ (Rb), $m/z = 103$ ($\sim^{87}\text{Sr}$), $m/z = 104$ ($\sim^{88}\text{Sr}$). Dwell times, repetition rate and scan speed (Table 1) were optimized on test line scans on target grains to suppress aliasing and carry-over effects and to obtain a signal/noise ratio > 10 for each mass channel. Grains were mapped with a 3–4 μm round spot at a fluence of 5.0 J cm^{-2} by using a raster of unidirectional scans

with no overlap between lines and a 2 s pause after each line. The total penetration depth of the laser is estimated to between 1 and 2 μm based on the 10-fold degree of spot overlap (dosage in Table 1) and assuming 100–200 nm ablation per laser shot. At regular intervals (\sim every 120 minutes), a set of reference materials (RM) was ablated in 30–40 s long line scans (total of 6–15 scans per RM and session). Reference materials were NIST610 as the primary calibration standard and the in-house



Table 4 Rb–Sr data of the mica RM obtained by ID-TIMS analyses

Sample	Rb [$\mu\text{g g}^{-1}$]	Sr [$\mu\text{g g}^{-1}$]	$^{87}\text{Rb}/^{86}\text{Sr}$	$\pm 2s$ (%)	$^{87}\text{Sr}/^{86}\text{Sr}$	$\pm 2s_m$ (%)
SagaB: 295.4 \pm 1.4 Ma; Sr ini. = 0.7049 \pm 0.0014; MSWD = 1.7						
Biotite	3043	5.22	5331	0.5	22.7375	0.0024
Feldspar	402	55	21.32	1.5	0.793036	0.0017
Bt + feldspar	2502	24.2	340.0	1.5	2.12442	0.0020
Harvard 98973: 266.8 \pm 1.6 Ma; Sr ini. = 0.720 \pm 0.011; MSWD = 0.12						
Musc 1	3666	3.98	52 860	1.5	197.872	0.0405
Musc 2	4268	4.57	54 575	2.4	205.646	0.0160
Musc 3	3639	3.96	67 847	0.74	254.067	0.0620
Assumed ini.			0.0001	100	0.72	Assumed 1.5
Phalaborwa: 2058.2 \pm 7.0 Ma; Sr ini. = 0.707671 \pm 0.000035; MSWD = 2.6						
Apatite	0.03	3751	0.00002	1.5	0.707672	0.0012
Biotite 1	654	12.5	262.48	0.5	8.34328	0.0040
Biotite 2	635	21.1	116.59	0.5	4.11875	0.0019

secondary RM, which are single crystal mica, namely SagaB biotite, Phalaborwa biotite and Harvard 98973 muscovite. In run GÄD08, the pressed mica nanopowder MicaMg^{9,26} has been measured in addition. A comprehensive overview of the essential ICP and laser settings used in each session is presented in Tables 1 and 2.

2.4. LA-ICP-MS/MS isotope ratios, Rb–Sr and concentrations by single spot ablation

Following the initial mapping process, the mica samples underwent a re-investigation using conventional analysis, employing static spot ablation to validate the accuracy of the mapped age distribution (Fig. 1). This verification procedure was conducted in two separate sessions, namely “GER2B/Wil1A spot” and “GÄD08 spot”. Moreover, the single crystal reference materials were analyzed during an additional session by randomly placing 28–30 spots across the single crystal to assess both the composition and the uniformity of the Rb–Sr age signal. This session is referred to as “homogeneity”. The same interface tubing is used for the aerosol transport, but another higher volume glass adapter from Glass Expansion is installed before the ICP plasma inlet as a signal smoothing device. The tuning procedure was identical to the mapping sessions and the following mass channels were recorded $m/z = 7$ (Li), $m/z = 23$ (Na), $m/z = 24$ (Mg), $m/z = 27$ (Al), $m/z = 39$ (K), $m/z = 44$ (Ca), $m/z = 57$ (Fe), $m/z = 60$ ($\sim^{28}\text{Si}$), $m/z = 63$ ($\sim^{47}\text{Ti}$), $m/z = 85$ (Rb), $m/z = 102$ ($\sim^{86}\text{Sr}$), $m/z = 103$ ($\sim^{87}\text{Sr}$), $m/z = 104$ ($\sim^{88}\text{Sr}$) with dwell times compiled in Table 1. While ^{85}Rb was acquired in analog mode, $\sim^{86}\text{Sr}$ and $\sim^{87}\text{Sr}$ were always acquired in pulse mode. In NIST610, $\sim^{88}\text{Sr}$ is acquired in analog mode but in pulse mode for all mica samples, hence being the only non-P/A matched nuclide during spot analysis (<https://doi.org/10.26434/chemrxiv-2023-273lw-v2>). Spot analyses of GER2B, Wil1A, and GER2B were bracketed by analyses of NIST610, SagaB, Harvard 98973, Phalaborwa and MicaMg RMs. Spot size was 65 μm at a repetition rate of 10 Hz and a fluence of 5.0 J cm^{-2} . One spot analysis comprised 30 seconds of gas blank and a 30-second period to allow for complete wash-out of the

aerosol. The length of the actual ablation interval was 40 s during “GER2B/Wil1A spot”, and “GÄD08 spot” and 60 s during “homogeneity”.

2.5. Data reduction strategy for elemental concentrations and phase assignment

Elemental and isotopic concentrations are processed using the software HDIP (version 1.6, Teledyne Photon Machines). The background was fitted using a cubic spline function for each mass channel and drift is corrected linearly using NIST610 as a drift monitor. For the elemental concentration maps, each mass sweep is transformed into a quadratic pixel and is calibrated using NIST610 line scan analyses. The assignment of pixels to specific mineral phases (muscovite, biotite, quartz, plagioclase, K-Fsp) is achieved using the labeled segments tool within HDIP. This tool provides flexibility in pixel selection, allowing manual selection, cluster analyses based on a k -means algorithm, or user-defined thresholds using logical operators. In the case of the k -means algorithm, a predefined number of clusters is necessary, which we determined based on the expected mineral phases. For instance, in the GER2B dataset, we used 3 clusters for quartz, plagioclase, and muscovite, while in the GÄD 08 dataset, 4 clusters were utilized to represent quartz, plagioclase, muscovite, and biotite. The k -means algorithm partitions the data points into clusters by assigning each point to the cluster whose mean (centroid) is closest to the point. This iterative process refines cluster assignments by recalculating centroids and reassigning data points until the clusters' positions converge and remain stable. To account for mixed analyses occurring at grain boundaries, an additional cluster per map was added. The clusters were generated by taking into account specific mass channels (^{28}Si , ^{85}Rb , ^{88}Sr , ^{87}Sr for GÄD08 and ^7Li , ^{27}Al , ^{28}Si , ^{85}Rb , ^{88}Sr , ^{87}Sr for GER2B). However, in the case of the WIL1A mapping, the compositional variability of muscovite and the appearance of K-Fsp as an additional phase posed challenges for the k -means filter. Therefore, we opted for the logical operators tool, where specific thresholds were defined for each of the four phases.

Muscovite: $^{87}\text{Sr} < 5 \mu\text{g g}^{-1}$, $\text{Rb} > 50 \mu\text{g g}^{-1}$, $\text{Fe} < 15\,000 \mu\text{g g}^{-1}$.



Table 5 Rb/Sr isotope ratios and calibration factors obtained from analyses of NIST610 in line mode

Session	$\sim^{87}\text{Sr}/\sim^{88}\text{Sr}$	$^{87}\text{Sr}/^{86}\text{Sr}$	2SE (%)	$^{85}\text{Rb}/\sim^{88}\text{Sr}$	$^{87}\text{Rb}/^{86}\text{Sr}$	2SE (%)	$\sim^{87}\text{Sr}/^{85}\text{Rb}$	$^{87}\text{Sr}/^{87}\text{Rb}$	2SE (%)	
GÄD08 (40 s line scan)	0.9014	0.7130	2.56	0.7004	2.4051	1.62	1.28698	0.29637	1.63	
	0.8971	0.7096	2.24	0.6788	2.3309	1.38	1.32160	0.30434	1.59	
	0.8931	0.7064	2.48	0.6600	2.2664	1.48	1.35318	0.31162	1.69	
	0.8995	0.7115	2.23	0.6767	2.3237	1.32	1.32924	0.30610	1.62	
	0.8965	0.7091	2.13	0.6752	2.3186	1.31	1.32775	0.30576	1.51	
	0.8959	0.7086	2.17	0.6801	2.3354	1.39	1.31731	0.30336	1.54	
	Mean	0.8973			0.6785			1.32268		
	2SE (%)	0.27			1.56			1.33		
	Expected	0.7097			2.3300			0.30459		
	Calibration factor	0.7910			3.4339			0.23028		
GER2B (40 s line scan)	0.2941	0.7145	0.62	0.6313	2.3246	0.92	0.46586	0.30731	0.60	
	0.2894	0.7031	0.67	0.6147	2.2635	0.80	0.47080	0.31057	0.48	
	0.2918	0.7089	0.69	0.6386	2.3515	0.81	0.45694	0.30143	0.51	
	0.2911	0.7072	0.53	0.6293	2.3172	0.65	0.46258	0.30515	0.41	
	0.2913	0.7077	0.51	0.6298	2.3191	0.63	0.46253	0.30511	0.39	
	0.2913	0.7077	0.52	0.6298	2.3191	0.66	0.46253	0.30511	0.44	
	0.2916	0.7084	1.76	0.6555	2.4137	3.16	0.44485	0.29345	1.63	
	0.2919	0.7091	0.49	0.6276	2.3110	0.69	0.46511	0.30681	0.47	
	0.2922	0.7099	0.54	0.6312	2.3242	0.66	0.46293	0.30538	0.41	
	0.2921	0.7096	0.55	0.6324	2.3286	0.65	0.46189	0.30469	0.42	
	0.2919	0.7091	0.57	0.6313	2.3246	0.69	0.46238	0.30502	0.42	
	0.2919	0.7091	0.52	0.6297	2.3187	0.71	0.46355	0.30579	0.50	
	0.2934	0.7128	0.51	0.6356	2.3404	0.64	0.46161	0.30451	0.42	
	0.2932	0.7123	0.55	0.6332	2.3316	0.66	0.46304	0.30545	0.57	
	0.2948	0.7162	0.50	0.6416	2.3625	0.62	0.45948	0.30310	0.41	
	Mean	0.2921			0.6328			0.46174		
	2SE (%)	0.24			0.70			0.62		
	Expected	0.7097			2.3300			0.30459		
	Calibration factor	2.4294			3.6822			0.65966		
WIL1A (50 s line scan)	0.2054	0.7082	0.60	0.6391	2.3234	0.61	0.32139	0.30479	0.58	
	0.2055	0.7085	0.57	0.6393	2.3241	0.60	0.32145	0.30484	0.58	
	0.2055	0.7085	0.57	0.6400	2.3266	0.67	0.32109	0.30451	0.95	
	0.2059	0.7099	0.49	0.6423	2.3350	0.57	0.32057	0.30401	0.56	
	0.2056	0.7088	0.54	0.6403	2.3277	0.61	0.32110	0.30452	0.50	
	0.2057	0.7092	0.49	0.6415	2.3321	0.54	0.32065	0.30409	0.50	
	0.2057	0.7092	0.53	0.6424	2.3354	0.54	0.32021	0.30367	0.52	
	0.2057	0.7092	0.53	0.6417	2.3328	0.57	0.32055	0.30400	0.54	
	0.2061	0.7106	0.54	0.6416	2.3324	0.55	0.32123	0.30464	0.53	
	0.2059	0.7099	0.56	0.6425	2.3357	0.56	0.32047	0.30392	0.59	
	0.2064	0.7116	0.57	0.6442	2.3419	0.62	0.32040	0.30385	0.71	
	0.2057	0.7092	0.55	0.6415	2.3321	0.59	0.32065	0.30409	0.58	
	0.2065	0.7120	0.51	0.6448	2.3441	0.60	0.32025	0.30371	0.54	
	0.2065	0.7120	0.56	0.6445	2.3430	0.65	0.32040	0.30386	0.55	
	0.2056	0.7088	0.51	0.6282	2.2837	0.65	0.32728	0.31038	0.52	
	Mean	0.2058			0.6409			0.32118		
	2SE (%)	0.09			0.31			0.28		
Expected	0.7097			2.3300			0.30459			
Calibration factor	3.4477			3.6354			0.94835			

Plagioclase: $^{88}\text{Sr} > 3 \mu\text{g g}^{-1}$, $\text{Rb} < 20 \mu\text{g g}^{-1}$, $\text{Fe} < 10\,000 \mu\text{g g}^{-1}$.

Biotite: $\text{Fe} > 15\,000 \mu\text{g g}^{-1}$.

K-Fsp: $^{88}\text{Sr} > 40 \mu\text{g g}^{-1}$, $\text{Rb} > 10 \mu\text{g g}^{-1}$.

In a last step, all pixels assigned to mica were normalized to SiO_2 concentrations of 45.2 wt%, corresponding to $\text{KAl}_3\text{Si}_3\text{O}_{10}(\text{OH})_{1.8}\text{F}_{0.2}$ stoichiometry. The mean normalization factor from all mica pixels was also applied to all non-mica pixels. This procedure introduces a systematic error to the biotite composition, whose SiO_2 concentration deviates slightly from muscovite (e.g. 41.6 wt% considering

$\text{K}(\text{Mg}_{2.5}\text{Fe}_{0.5})\text{AlSi}_3\text{O}_{10}(\text{OH})_{1.8}\text{F}_{0.2}$ stoichiometry). However, elemental and isotopic ratios or relative compositional changes within a map remain unaffected.

Concentrations obtained during spot analyses were calibrated on NIST 610 and normalized to a sum of $\text{SiO}_2 + \text{Al}_2\text{O}_3 + \text{TiO}_2 + \text{FeO} + \text{MgO} + \text{CaO} + \text{Na}_2\text{O} + \text{K}_2\text{O} = 95\%$, assuming $\text{H}_2\text{O} + \text{F} = 5 \text{ wt}\%$. The compositions obtained for Phalaborwa biotite, Harvard 98973 muscovite and SagaB biotite during session homogeneity are displayed in Table 3.



2.6. Data reduction strategy and uncertainty propagation for LA-ICP-MS/MS based Rb–Sr ages

The data reduction and uncertainty propagation strategy for line, map and spot derived Rb–Sr ages largely follows the recommendations by Horstwood *et al.* (2016)³⁷ and Rösel & Zack (2022).²⁰ An overview of the workflow is summarized in Fig. 3.

For line scan/mapping analyses the background was subtracted for each mass channel using a cubic spline function in HDIP. Drift is linearly corrected using NIST610 as a drift monitor. Background and drift correction for spot analyses is performed using an in-house Excel spreadsheet. Here, the background of the Rb and Sr mass channels was fitted for each individual spot by subtracting the mean gas blank signal obtained in the time interval 1–26 s before ablation was initiated. No downhole fractionation correction was applied.

After transferring each channel from counts to counts per second based on the applied dwell times (Table 2), “raw” isotope ratios were calculated for each individual mass sweep (*e.g.* $\sim^{87}\text{Sr}/\sim^{88}\text{Sr}$ and $\sim^{85}\text{Rb}/\sim^{88}\text{Sr}$). Mean isotope ratios and their respective standard deviations (SD) and standard errors (SE; $\text{SE} = \text{SD}/\sqrt{n}$ with n being the number of mass sweeps) are then calculated based on a “mean of ratios” approach.³⁸ For spot analyses all mass sweeps within the first ~ 10 seconds of the ablation interval were discarded to allow for stabilization of the signal. This interval was held constant for each individual spot during a particular session. No such cut-off was applied for line scan analyses. Calibration factors are derived based on replicate analyses (either spot or line scans) on NIST610 glass to correct for mass bias. “Raw” isotope ratios (*e.g.* $\sim^{87}\text{Sr}/\sim^{88}\text{Sr}$ and $\sim^{85}\text{Rb}/\sim^{88}\text{Sr}$) were transferred into “true” isotope ratios (*e.g.* $^{87}\text{Sr}/^{86}\text{Sr}$ and $^{87}\text{Rb}/^{86}\text{Sr}$ ratios) by applying the following and analogous equations:

$$\left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{sample,true}} = \left[\frac{\left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{NIST 610,true}}}{\left(\frac{\sim^{87}\text{Sr}}{\sim^{88}\text{Sr}} \right)_{\text{NIST 610,raw}}} \right] \left(\frac{\sim^{87}\text{Sr}}{\sim^{88}\text{Sr}} \right)_{\text{sample,raw}} \quad (1)$$

For the true Rb–Sr isotope ratios of NIST 610 the following working values are used: $^{87}\text{Rb}/^{86}\text{Sr} = 2.3300$ and $^{87}\text{Sr}/^{86}\text{Sr} = 0.709699$.^{15,25}

Ages are calculated based on the isochron equation:¹⁹

$$\text{Age} = \frac{1}{\lambda} \ln \left\{ 1 + \left[\left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{sample,true}} - \left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{initial}} \right] \times \left/ \left(\frac{^{87}\text{Rb}}{^{86}\text{Sr}} \right)_{\text{sample,true}} \right. \right\} \quad (2)$$

with $\lambda = 1.3892 \pm 0.0045 \times 10^{-11} \text{ a}^{-1}$ as the value for the ^{87}Rb decay constant.¹ This equation can be simplified:

$$\text{Age} = \frac{1}{\lambda} \ln \left[1 + \left(\frac{^{87}\text{Sr}}{^{87}\text{Rb}} \right)_{\text{Sr,initial,corrected}} \right] \quad (3)$$

Given that:

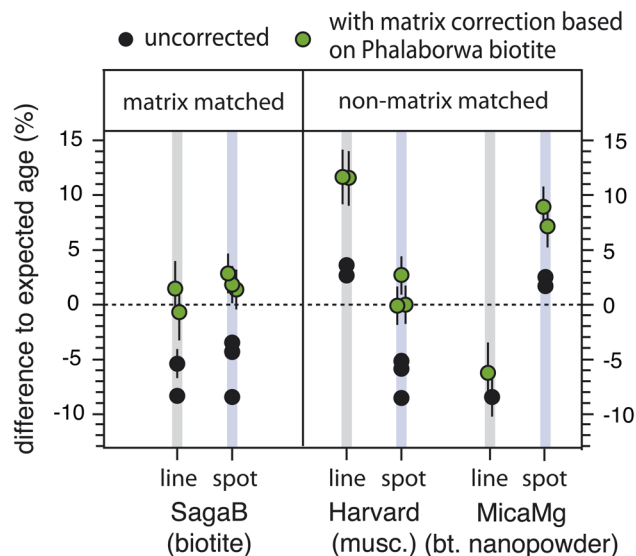


Fig. 4 Accuracy of Rb–Sr ages of single crystal mica RM and MicaMg pressed nanopowder with and without matrix correction. “Line” indicates data acquisition in line scan mode used for mapping purposes and “spot” indicates conventional spot acquisition. Matrix correction is based on Phalaborwa biotite. Only Phalaborwa and SagaB are considered structurally and chemically matrix matched since both are crystalline biotite. In line scan mode a chemical and structural matrix match is pivotal for accurate results, whereas in spot mode only a structural matrix match is required.

$$\left[\left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{sample,true}} - \left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{initial}} \right] / \left(\frac{^{87}\text{Rb}}{^{86}\text{Sr}} \right)_{\text{sample,true}} = \left[1 + \left(\frac{^{87}\text{Sr}}{^{87}\text{Rb}} \right)_{\text{Sr,initial,corrected}} \right] \quad (4)$$

with $\left(\frac{^{87}\text{Sr}}{^{87}\text{Rb}} \right)_{\text{Sr,initial,corrected}}$ being the $^{87}\text{Sr}/^{87}\text{Rb}$ ratios corrected for the initial Sr-isotope composition $\left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{initial}}$. To calculate $\left(\frac{^{87}\text{Sr}}{^{87}\text{Rb}} \right)_{\text{Sr,initial,corrected}}$ a “ratio of means” approach

is applied.³⁸ To obtain the random uncertainty (s_{random}), the standard error of $\left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{sample,true}}$ and $\left(\frac{^{87}\text{Rb}}{^{86}\text{Sr}} \right)_{\text{sample,true}}$ ratios (denoted s_{m}) and their excess variance (ϵ) were added in quadrature (Fig. 3).

An essential aspect during data reduction is the uncertainty of the common Sr correction (γ), which corrects for the amount of common ^{87}Sr ($^{87}\text{Sr}_{\text{c}}$) that did not form *via in situ* radioactive decay of ^{87}Rb . We implemented the procedure proposed by Rösel & Zack (2022),²⁰ treating γ as a systematic uncertainty that is calculated by weighting the uncertainty of $\left(\frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_{\text{initial}}$ to the magnitude of the common Sr correction [eqn (9) in Rösel & Zack (2022)].²⁰ In case the fraction of $^{87}\text{Sr}_{\text{c}}$ is low enough ($< \sim 1\%$), a common Sr correction is not required and ages are directly calculated from the $\left(\frac{^{87}\text{Sr}}{^{86}\text{Rb}} \right)_{\text{true}}$ ratios:



Table 6 Rb/Sr isotope ratios and calibration factors obtained from analyses of NIST610 in spot mode

	$^{87}\text{Sr}/^{86}\text{Sr}$	$^{87}\text{Sr}/^{86}\text{Sr}$	2SE (%)	$^{85}\text{Rb}/^{86}\text{Sr}$	$^{87}\text{Rb}/^{86}\text{Sr}$	2SE (%)	$^{87}\text{Sr}/^{85}\text{Rb}$	$^{87}\text{Sr}/^{87}\text{Rb}$	2SE (%)
GER2B/WIL1A (spots)	0.72853	0.7096	0.63	8.25120	2.3445	0.72	0.08836	0.30343	0.69
	0.72888	0.7099	0.67	8.18057	2.3244	0.90	0.08924	0.30647	0.99
	0.72764	0.7087	0.68	8.16421	2.3197	0.82	0.08921	0.30635	0.81
	0.72717	0.7083	0.96	8.27992	2.3526	1.44	0.08794	0.30199	1.27
	0.73017	0.7112	1.11	8.26125	2.3473	1.50	0.08852	0.30399	1.41
	0.73008	0.7111	1.06	8.20448	2.3312	1.14	0.08896	0.30549	1.16
	0.72607	0.7072	0.94	8.15324	2.3166	1.28	0.08888	0.30522	1.21
	0.72807	0.7091	0.99	8.16681	2.3205	1.28	0.08899	0.30559	1.32
	0.72567	0.7068	1.03	8.16689	2.3205	1.40	0.08874	0.30474	1.42
	0.73381	0.7147	1.44	8.18843	2.3266	2.24	0.08974	0.30816	2.09
	0.72715	0.7082	1.26	8.20666	2.3318	1.92	0.08849	0.30387	1.76
	0.73112	0.7121	1.09	8.20293	2.3307	1.58	0.08880	0.30495	1.46
	0.72937	0.7104	0.93	8.23217	2.3391	1.24	0.08791	0.30187	1.25
	0.72815	0.7092	0.90	8.13048	2.3102	1.22	0.08883	0.30504	1.21
	0.72782	0.7089	0.90	8.21518	2.3342	1.20	0.08787	0.30173	1.20
Mean	0.72865			8.20029			0.08870		
2SE (%)	0.15			0.27			0.31		
Expected	0.7097			2.3300			0.30459		
Calibration factor	0.97400			0.28414			3.43406		
GÄD08 (spots)	0.72528	0.7125	1.03	5.63831	2.3437	1.49	0.12912	0.30432	1.46
	0.71951	0.7068	1.14	5.57035	2.3154	1.51	0.12968	0.30564	1.41
	0.72135	0.7086	0.82	5.53829	2.3021	1.31	0.13072	0.30808	1.36
	0.72498	0.7122	0.88	5.63313	2.3415	1.36	0.12915	0.30439	1.42
	0.72325	0.7105	0.90	5.67350	2.3583	1.27	0.12773	0.30105	1.17
	0.72197	0.7092	0.86	5.62590	2.3385	1.01	0.12846	0.30276	1.07
	0.72421	0.7114	0.57	5.59656	2.3263	0.92	0.12948	0.30516	0.91
	0.71996	0.7073	0.73	5.60574	2.3301	1.01	0.12863	0.30315	1.02
	0.71901	0.7063	0.77	5.57798	2.3186	1.22	0.12940	0.30498	1.23
	0.71986	0.7072	1.15	5.61409	2.3336	1.59	0.12883	0.30362	1.49
	0.72475	0.7120	0.90	5.60244	2.3287	1.44	0.12994	0.30624	1.45
	0.72435	0.7116	1.05	5.59543	2.3258	1.42	0.12970	0.30567	1.29
	0.72262	0.7099	1.06	5.61006	2.3319	1.37	0.12902	0.30408	1.36
	0.72295	0.7102	1.11	5.61255	2.3329	1.53	0.12922	0.30454	1.52
	0.72243	0.7097	0.92	5.58758	2.3226	1.28	0.12949	0.30519	1.22
Mean	0.72243			5.60546			0.12924		
2SE (%)	0.15			0.29			0.28		
Expected	0.7097			2.3300			0.30459		
Calibration factor	0.98238			0.41567			2.35684		
Homogeneity (spots)	0.7228	0.7087	0.48	6.131607	2.3218	0.82	0.11817	0.30554	0.94
	0.7239	0.7097	0.49	6.179168	2.3399	0.87	0.11747	0.30371	0.97
	0.7247	0.7105	0.42	6.196586	2.3464	0.86	0.11727	0.30320	0.92
	0.7262	0.7120	0.49	6.208785	2.3511	0.75	0.11717	0.30294	0.83
	0.7261	0.7119	0.48	6.155889	2.3310	0.85	0.11826	0.30576	1.00
	0.7213	0.7071	0.46	6.131273	2.3217	0.94	0.11796	0.30499	0.94
	0.7238	0.7096	0.49	6.193627	2.3453	0.90	0.11710	0.30277	0.94
	0.7261	0.7119	0.47	6.130510	2.3214	0.89	0.11867	0.30682	0.91
	0.7245	0.7103	0.55	6.132414	2.3221	0.96	0.11841	0.30614	0.95
	0.7237	0.7096	0.38	6.109771	2.3136	0.69	0.11849	0.30635	0.73
	0.7252	0.7110	0.53	6.101410	2.3104	0.73	0.11888	0.30736	0.77
	0.7227	0.7085	0.41	6.086960	2.3049	0.81	0.11885	0.30728	0.90
	0.7229	0.7087	0.49	6.190817	2.3443	0.96	0.11696	0.30240	1.06
	0.7225	0.7083	0.44	6.099260	2.3096	0.89	0.11857	0.30657	0.95
	0.7208	0.7067	0.50	6.127270	2.3202	0.89	0.11772	0.30437	0.90
	0.7237	0.7095	0.52	6.149925	2.3288	0.86	0.11768	0.30426	0.84
	0.7232	0.7090	0.50	6.132032	2.3220	0.82	0.11795	0.30497	0.89
	0.7278	0.7135	0.70	6.160059	2.3326	0.95	0.11825	0.30574	1.11
	0.7226	0.7084	0.44	6.243558	2.3642	0.96	0.11580	0.29941	1.00
	0.7238	0.7096	0.52	6.155113	2.3307	0.93	0.11767	0.30422	1.06
	0.7234	0.7092	0.50	6.200097	2.3478	0.90	0.11667	0.30165	0.91
Mean	0.72389			6.15315			0.11781		
2SE (%)	0.10			0.29			0.29		
Expected	0.7097			2.3300			0.30459		
Calibration factor	0.98039			0.37867			2.58548		



$$\text{Age} = \frac{1}{\lambda} \ln \left[1 + \left({}^{87}\text{Sr}/{}^{87}\text{Rb} \right)_{\text{sample,true}} \right] \quad (5)$$

This way ages are calculated for each of the spot analyses and line scans, both of which consist of multiple mass sweeps. For line scan analyses on NIST610 the number of mass sweeps is between 280 and 580, depending on dwell times and length of the line scans (Table 5). For spot analysis, the number of mass sweeps is either 97 (sessions “GER2B/WIL1A” and “GÄD08”) or 151 (session “homogeneity”).

In both line and spot mode acquisition, the NIST610 calibrated ages of the mica RM were offset with respect to their expected ages by up to ~11% (Fig. 4 and Table 7). This bias is in the order of magnitude that has been observed in other studies,^{29–31} where it is ascribed to matrix effects resulting from the use of a non-matrix matched primary reference material (*e.g.* NIST610 glass or pressed nanopowder Mica-Mg). It has been suggested to correct for that matrix-induced bias by co-analyzing a matrix-matched, *i.e.* crystalline mica with known age.³⁰ We adapted this approach by calculating matrix correction factors (MCF):

$$\text{MCF} = \frac{\text{age}_{\text{RM,expected}}}{\text{age}_{\text{CRM,measured}}} \quad (6)$$

that were then used to correct the obtained mean single spot or line scan ages *via*:

$$\text{Age}_{\text{matrix corrected}} = \text{MCF} \times \text{age}_{\text{NIST calibrated}} \quad (7)$$

The total uncertainty (s^*) of a single spot or line scan is calculated by adding the random uncertainty and all systematic uncertainties in quadrature:

$$2s^* = s_{\text{random}}^2 + \gamma^2 + s_{\lambda}^2 + s_y^2 + s_{\text{age, RM}}^2 + \varepsilon_{\text{homo}}^2 + \varepsilon'^2 \quad (8)$$

with s_{λ} being the uncertainty of the decay constant (0.3%),¹ s_y being the uncertainty of NIST610 ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ and ${}^{87}\text{Rb}/{}^{86}\text{Sr}$ ratios (0.003% and 0.21%),^{15,25} $s_{\text{age, RM}}$ being the uncertainty of the expected age of the secondary mica RM used for matrix correction, $\varepsilon_{\text{homo}}$ being the excess variance obtained during the LA-ICP-MS homogeneity test of the mica RM and ε' being the long term excess variance on ${}^{87}\text{Rb}/{}^{86}\text{Sr}$, ${}^{87}\text{Rb}/{}^{86}\text{Sr}$, and ${}^{87}\text{Rb}/{}^{86}\text{Sr}$ ratios obtained on NIST610 across all sessions referring to a particular measurements mode, *i.e.* spot or line scan. The values for the long-term excess variance are available *via* the electronic supplement (<https://doi.org/10.26434/chemrxiv-2023-273lw-v2>). If multiple single spot or line scan analyses are pooled ($n > 1$), $2s_{\text{random}}$ is replaced by $2s_x$, which is the $2s$ value obtained from the weighted mean statistics calculated using Isoplot.³⁹

Two-dimensional Rb–Sr age maps were created using HDIP. During this procedure, the ${}^{87}\text{Rb}/{}^{86}\text{Sr}$ and ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ ratios, which had been calibrated with NIST610 and obtained in a single mass sweep, were converted into quadratic pixels. The pixel sizes were adjusted to correspond to the laser spot dimensions (3 μm for session GÄD08/WIL1A and 4 μm for session GER 2B). The

numerical values of the pixels were then assigned to a color scale for visualization (Fig. 5). We employed the viridis color scheme, which features blue and yellow sequences, making it suitable for both color-blind individuals and black and white presentations. Finally, for each pixel which corresponds to a single mass sweep, an age is calculated by applying the Rb–Sr isochron equation (eqn (2)), which has been implemented in the HDIP channel calculator tool. The random uncertainty (s_{random}) of a single mass sweep (or pixel) is estimated by the standard deviation (SD) of the ${}^{87}\text{Rb}/{}^{86}\text{Sr}$ and ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ ratios obtained during a single line scan on NIST610 (Fig. 5 and Table 5). For the GÄD08 session we have obtained a mean ratio uncertainty of 39% (2SD) on ${}^{87}\text{Rb}/{}^{86}\text{Sr}$ and 24% (2SD) on ${}^{87}\text{Sr}/{}^{86}\text{Sr}$. For the subsequent sessions GER2B and WIL1A, NIST610 was polished with 1 μm diamond paste, significantly improving the uncertainty (mean 2SD uncertainty on ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ = 12–13% and mean 2SD uncertainty on ${}^{87}\text{Rb}/{}^{86}\text{Sr}$ = 14.4–15.2%). Both of the latter isotope ratio uncertainties are similar to the uncertainties obtained in spot ablation mode (Table 6).

To further improve the measurement uncertainty on Rb–Sr ages, multiple pixels were pooled. This is accomplished through a meticulous user-driven selection of pixel segments employing the Rb–Sr age map as a reference to identify regions that represent homogeneous age populations. This way the mixed analyses of multiple age populations, as seen for example in the GER2B map (Fig. 6) is avoided. In sample GÄD08, which consists of co-genetic and apparently age homogeneous muscovite and biotite, we additionally analyzed segments comprising all biotite and muscovite pixel as assigned by the *k*-means algorithm (Fig. 7, see Section 2.5).

To finally extract the ages from the resulting pixel segments we applied two approaches (Fig. 8 and Table 9). The first one is calculating the weighted mean age of all pixels using the IsoplotR software⁴⁰ and taking into account the estimated random uncertainty of a single mass sweep (see above). The second approach is to create histograms from all single pixel ages within a segment and fitting a Gaussian probability distribution to it, with the age corresponding to the Gaussian mean. Both histograms and Gaussian fitting are performed using the Igor Pro software (version 8.04) and applying the Freedman–Daiconis algorithm to calculate the bin width. The resulting ages were again matrix corrected, and all systematic uncertainties were added in quadrature (see eqn (8)). Since GER2B and WIL1A are muscovites, a MCF based on RM Harvard 98973 is applied. For GÄD08, which contains muscovite and biotite, a Phalaborwa biotite-based MCF is used for matrix correction.

The initial Sr-isotope values for GÄD08, GER2B and WIL1A are obtained by averaging the ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ over several small segments within plagioclase contained in the same mapping (Fig. 6 and 7). To derive the initial ${}^{87}\text{Sr}/{}^{86}\text{Sr}$, the plagioclase segment with the smallest ${}^{87}\text{Sr}/{}^{86}\text{Sr}$ ratio is chosen (Table 8) to account for post-crystallization mobility of radiogenic Sr in pegmatites,^{5,41} which is of particular importance for rocks with high Rb/Sr ratios. The final $({}^{87}\text{Sr}/{}^{86}\text{Sr})_{\text{initial}}$ is calculated from that segment by correcting for the amount of radiogenic ${}^{87}\text{Sr}$ based on the expected ages and the measured ${}^{87}\text{Rb}/{}^{86}\text{Sr}$ using IsoplotR.⁴⁰ For all mica RM, the fraction of common ${}^{87}\text{Sr}$ is low



Table 7 Rb–Sr dating of single crystal mica reference materials recorded in line scan and spot mode^a

Line scan (this study)			
Session	Rb–Sr age (Ma)	2s _x (%)	Rb–Sr age (Ma) matrix corrected 2s* (%)
Phalaborwa (age = 2062 ± 2 Ma) ⁸⁷Rb/⁸⁶Sr = 350–42 000			
GÄD08 (<i>n</i> = 6)	2013	0.8	
Accuracy (%)	–2.38		
GER2B (<i>n</i> = 15)	1902	0.5	
Accuracy (%)	–7.76		
WIL1A (<i>n</i> = 15)	1919	1.2	
Accuracy (%)	–6.94		
SagaB (age = 295.4 ± 1.4 Ma) ⁸⁷Rb/⁸⁶Sr = 32 000–270 000			
GER2B (<i>n</i> = 15)	270.2	0.8	293.0 2.4
Accuracy (%)	–8.53		–0.81
WIL1A (<i>n</i> = 15)	278.8	1.2	299.5 2.5
Accuracy (%)	–5.62		1.38
Harvard 98973 (age 266.8 ± 1.6 Ma) ⁸⁷Rb/⁸⁶Sr = 11 000–44 000			
GER2B (<i>n</i> = 15)	273.9	0.6	296.9 2.3
Accuracy (%)	2.66		11.30
WIL1A (<i>n</i> = 15)	276.6	0.8	297.2 2.4
Accuracy (%)	3.67		11.39
MicaMg (age = 519.4 ± 6.5 Ma) ⁸⁷Rb/⁸⁶Sr = 154.6			
GÄD08 (<i>n</i> = 6)	475.3	1.7	486.9 2.8
Accuracy (%)	–8.49		–6.26
Single spot (this study)			
Session	Rb–Sr age (Ma)	2s _x (%)	Rb–Sr age (Ma) matrix corrected 2s* (%)
Phalaborwa (age = 2062 ± 2 Ma) ⁸⁷Rb/⁸⁶Sr = 350–42 000			
GER2B/WIL1A (<i>n</i> = 15)	1939.3	0.5	
Accuracy (%)	–5.95		
GÄD08 (<i>n</i> = 15)	1960.9	0.4	
Accuracy (%)	–4.90		
Homogeneity (<i>n</i> = 29)	1834.2	0.4	
Accuracy (%)	–11.05		
SagaB (age = 295.4 ± 1.4 Ma) ⁸⁷Rb/⁸⁶Sr = 32 000–270 000			
GER2B/WIL1A (<i>n</i> = 15)	282.8	0.5	300.7 1.7
Accuracy (%)	–4.27		1.79
GÄD08 (<i>n</i> = 15)	284.5	0.7	299.2 1.9
Accuracy (%)	–3.69		1.29
Homogeneity (<i>n</i> = 28)	270.4	0.3	304.0 1.8
Accuracy (%)	–8.45		2.92
Harvard 98973 (age = 266.8 ± 1.6 Ma) ⁸⁷Rb/⁸⁶Sr = 11 000–44 000			
GER2B/WIL1A (<i>n</i> = 15)	250.7	0.4	266.5 1.7
Accuracy (%)	–6.05		–0.11
GÄD08 (<i>n</i> = 15)	253.9	0.4	267.0 1.8
Accuracy (%)	–4.97		0.07
Homogeneity (<i>n</i> = 30)	243.9	0.3	274.2 1.8
Accuracy (%)	–8.79		2.79
MicaMg (age = 519.4 ± 6.5 Ma) ⁸⁷Rb/⁸⁶Sr = 154.6			
GÄD08 (<i>n</i> = 15)	529.2	0.7	556.5 1.9
Accuracy (%)	1.89		7.14
GER2B/WIL1A (<i>n</i> = 15)	532.7	0.7	566.4 1.9
Accuracy (%)	2.56		9.05

Table 7 (Contd.)

Single spot (Rösel & Zack 2022) ²⁰		
Session	Rb–Sr age (Ma)	2s _x (%)
Högsbo (age = 1029 Ma) ⁸⁷Rb/⁸⁶Sr = 3440–24 000		
191 003 (<i>n</i> = 12)	1037.8	0.7
Accuracy (%)	0.86	
191 128 (<i>n</i> = 9)	1037.0	0.7
Accuracy (%)	0.78	
203 021 (<i>n</i> = 14)	1002.8	0.7
Accuracy (%)	–2.54	
La Posta (age = 91.6 Ma) ⁸⁷Rb/⁸⁶Sr = 408–1239		
180 220 (<i>n</i> = 27)	88.7	1.5
Accuracy (%)	–3.17	
191 128 (<i>n</i> = 7)	90.0	2.1
Accuracy (%)	–1.75	
203 021 (<i>n</i> = 7)	91.0	2.3
Accuracy (%)	–0.66	
Mt Dromedary (age = 100.4 Ma) ⁸⁷Rb/⁸⁶Sr = 55–171.1		
191 003 (<i>n</i> = 7)	99.1	3.0
Accuracy (%)	–1.28	
191 128 (<i>n</i> = 9)	100.3	2.5
Accuracy (%)	–0.10	
McClure Mt (age = 523.98 Ma) ⁸⁷Rb/⁸⁶Sr = 110–940		
191 003 (<i>n</i> = 7)	511.5	1.0
Accuracy (%)	–2.38	
191 128 (<i>n</i> = 8)	513.7	1.1
Accuracy (%)	–1.96	
200 321 (<i>n</i> = 10)	518.5	1.0
Accuracy (%)	–1.05	

^a For Harvard 98973, SagaB and Phalaborwa, ages are calculated without a common Sr correction due to the high amount of ⁸⁷Sr using eqn (3). Ages for MicaMg are calculated using eqn (2) with (⁸⁷Sr/⁸⁶Sr)_{initial} = 0.72607 ± 0.0007.⁹ Single spot ages of Högsbo, La Posta, Mt Dromedary and McClure Mountain are from Rösel & Zack (2022).²⁰ All ages are indicated as weighted means. 2s_x denotes the uncertainty on the weighted mean calculated using random uncertainties. 2s* denotes the total uncertainty resulting from propagating all random and systematic uncertainties. Matrix correction for single spot and line scan ages presented in our study is based on Phalaborwa biotite. Expected ages for Phalaborwa are from Wu *et al.* (2011);³² for MicaMg from Högalm *et al.* (2017);⁹ for Högsbo from Romer & Smeds (1994);⁴³ for La Posta from Zack & Högalm (2016);¹⁵ for Mt Dromedary from Williams *et al.* (1982);⁴⁶ for McClure Mountain from Schoene & Bowring (2006).⁴⁷ For SagaB and Harvard 98973 expected ages are based on ID-TIMS data presented in this study.

enough (0.2–1.3%) to neglect a common Sr correction. For mica-Mg pressed nanopowder (⁸⁷Sr/⁸⁶Sr)_{initial} is 0.72607 ± 0.0007.⁹

2.7. ID-TIMS dating of SagaB, Harvard 98973 and Phalaborwa mica RM

For the secondary reference materials Phalaborwa and SagaB, kg-sized samples of the original rock were available. Mica and apatite or feldspar used for analysis were parts of particularly clear domains of individual crystals. In the case of Harvard 98973, we used different aliquots of white mica, combined with a small turbid feldspar crystal that was left from the original



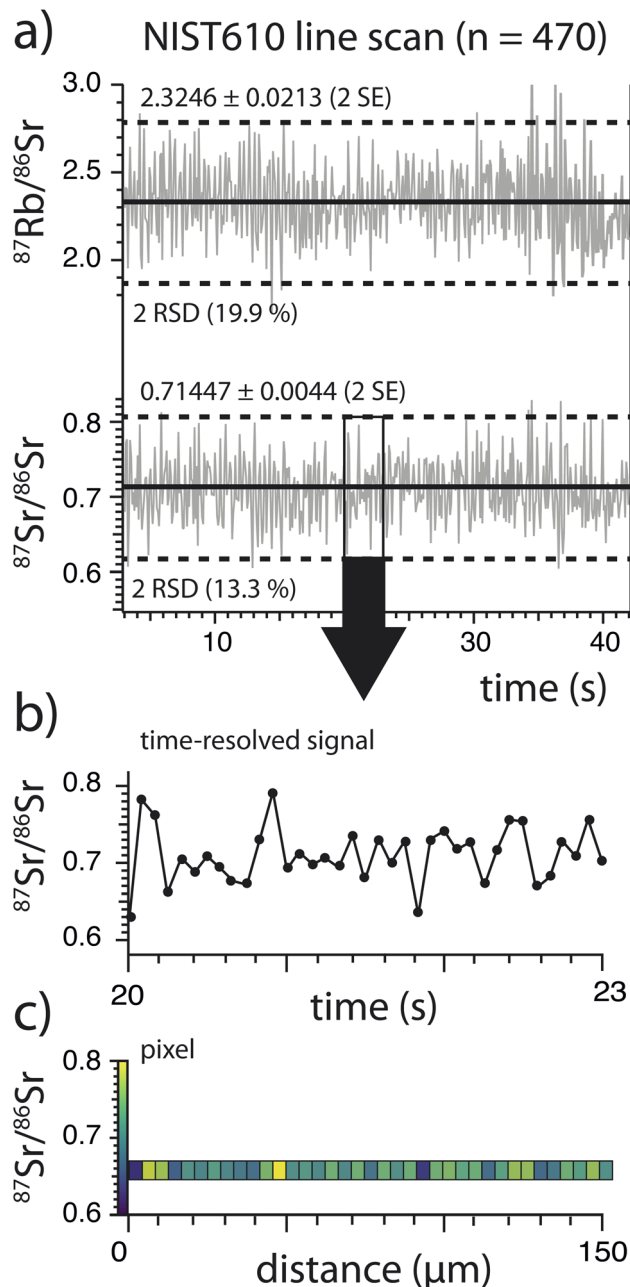


Fig. 5 (a) Single line scan on NIST610 acquired during the mapping session of GER2B (analyses #1 in Table 5). (b) Blow-up of $^{87}\text{Sr}/^{86}\text{Sr}$ ratios within a short time segment (3 s) within that line scan. Each point within the time-resolved signal corresponds to a single mass sweep ($n = 470$, for the whole line scan). (c) To obtain 2D maps of $^{87}\text{Sr}/^{86}\text{Sr}$, $^{87}\text{Rb}/^{86}\text{Sr}$ isotope ratios and ultimately ages, these values were transferred to colored pixel using the viridis color scheme.

preparation of reference material. Rb–Sr isotopic data were generated at GFZ Potsdam using a Thermo Scientific TRITON thermal ionization mass spectrometer and a set of mixed ^{87}Rb – ^{84}Sr tracers. Sr was measured in dynamic multicollection mode, and Rb isotope-dilution analysis was done in static multicollection mode. The $^{87}\text{Sr}/^{86}\text{Sr}$ obtained for NIST SRM 987 reference material during the period of analytical work was 0.710245 ± 0.000008 (2 s; $n = 10$). For age calculation, in case of

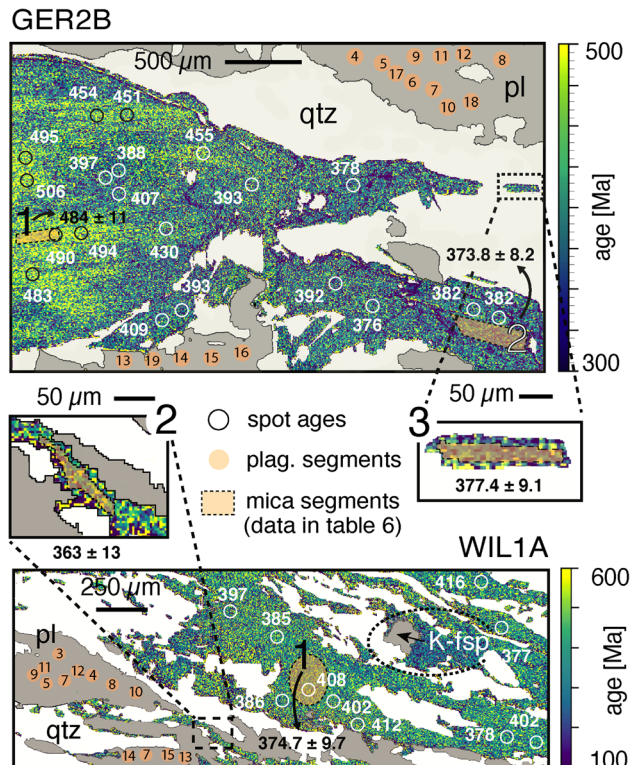


Fig. 6 Rb–Sr age maps of deformed muscovite from two meta-pegmatites (GER2B and Wil1A) from the Western Bohemian Massif in NE Bavaria (Germany) consisting mainly of white mica and plagioclase. Different segments have been defined for plagioclase and mica for which isotope and age information is obtained (Tables 8 and 9). Image pixel size is $3 \mu\text{m}$ for WIL1A and $4 \mu\text{m}$ for GER2B. Values adjacent to the black and white circles indicate the single spot ages that have been acquired post mapping. For segments referring to mica, ages and uncertainties (95% conf.) in Ma are indicated. All ages are with matrix correction based on Harvard 98973 muscovite.

feldspar and apatite, standard 2 s uncertainties of $\pm 0.005\%$ for $^{87}\text{Sr}/^{86}\text{Sr}$ and of $\pm 1.5\%$ for $^{87}\text{Rb}/^{86}\text{Sr}$ ratios were assigned to the results. Individual analytical uncertainties were generally smaller than these values. For mica $^{87}\text{Rb}/^{86}\text{Sr}$ ratios, uncertainty estimates are based on the standard error of several (3 to 6) independent isotope dilution analyses of Rb concentrations, combined with uncertainties of the Rb instrumental mass fractionation correction factor and of the Sr isotope dilution analysis. For mica $^{87}\text{Sr}/^{86}\text{Sr}$ ratios, we used either a standard uncertainty of $\pm 0.005\%$ or, if higher, the individual analytical uncertainties. Analytical procedures for sample dissolution and separation of Rb from Sr are described in more detail elsewhere.⁴² Uncertainties of isotope and age data are quoted at 2 s. The program Isoplot³⁹ was used to calculate regression lines. The ^{87}Rb decay constant was the one recommended by Villa and co-workers.¹

Age results are as follows: for Phalaborwa, we obtained an age of 2058.2 ± 7.0 Ma (MSWD = 2.6; initial $^{87}\text{Sr}/^{86}\text{Sr} = 0.707671 \pm 0.000035$), based on data for apatite and two biotite crystal fragments. For SagaB, the Rb–Sr age based on biotite, a biotite-feldspar aggregate and feldspar is 295.4 ± 1.4 Ma (MSWD = 1.7,



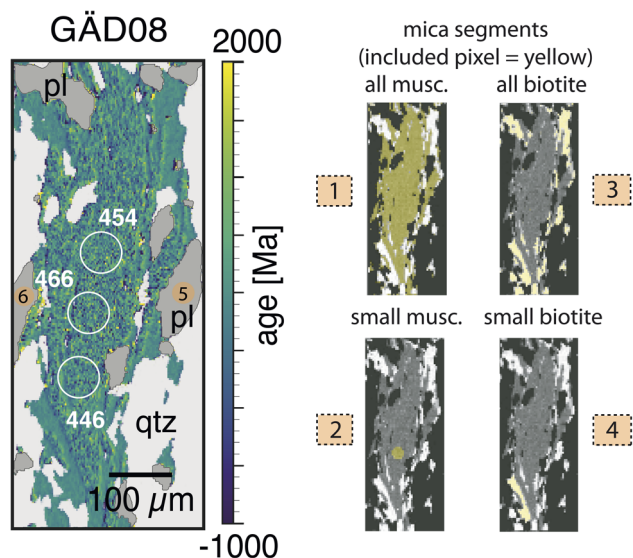


Fig. 7 Rb–Sr age map of a mylonitic mica schist (GÄD08) from the Middle Seve Nappe near Ankarede, Jämtland (Sweden) consisting mainly of muscovite, biotite and plagioclase. Image pixel size is 3 μm . Different segments have been assigned for mica and biotite, with segment #1 and #3 comprising all muscovite and biotite pixel, respectively. Segments #2 and #4 comprise a smaller subset. Two segments #5 and #6 are assigned to plagioclase. White circles indicate Rb–Sr ages obtained from single spot analyses with matrix correction based on Harvard 98973. Isotopic and age data for the segments is compiled in Tables 8 and 9.

initial $^{87}\text{Sr}/^{86}\text{Sr} = 0.7049 \pm 0.0014$). Both these results are in agreement with published U–Pb data (better than $\pm 1\%$; see Section 2.1 for references). The model age determined here for RM Harvard 98973 is 266.8 ± 1.6 Ma (MSWD = 0.12; assumed initial $^{87}\text{Sr}/^{86}\text{Sr} = 0.720 \pm 0.011$), an age that is well within the range of ages of other pegmatite bodies in its regional pegmatite field. The turbid feldspar crystal that we analyzed alongside the Harvard 98973 mica fractions calculates to an impossibly low, apparently sub-primordial initial Sr isotopic composition, an effect that indicates some secondary overprint of that feldspar. Therefore, this analysis is not taken into further consideration. Calculation of the Rb–Sr model age for Harvard 98973 is, due to the extremely radiogenic $^{87}\text{Sr}/^{86}\text{Sr}$ signature of the mica, almost insensitive to the model initial $^{87}\text{Sr}/^{86}\text{Sr}$ ratio. We therefore consider the above Harvard 98973 model age as equally robust as the Rb–Sr ages obtained for SagaB and Phalaborwa. All ID-TIMS results are compiled in Table 4.

3. Results & discussion

3.1. Isotopic homogeneity of the secondary mica RM

An integral part of the data reduction scheme involves matrix correction by co-analysis of a matrix matched single crystalline mica RM (SagaB biotite, Harvard 98973 muscovite and Phalaborwa biotite). Hence, it is crucial to assess the isotopic homogeneity of the mica RM. This was done by randomly placing 28–30 LA-ICP-MS/MS spot analyses across the epoxy

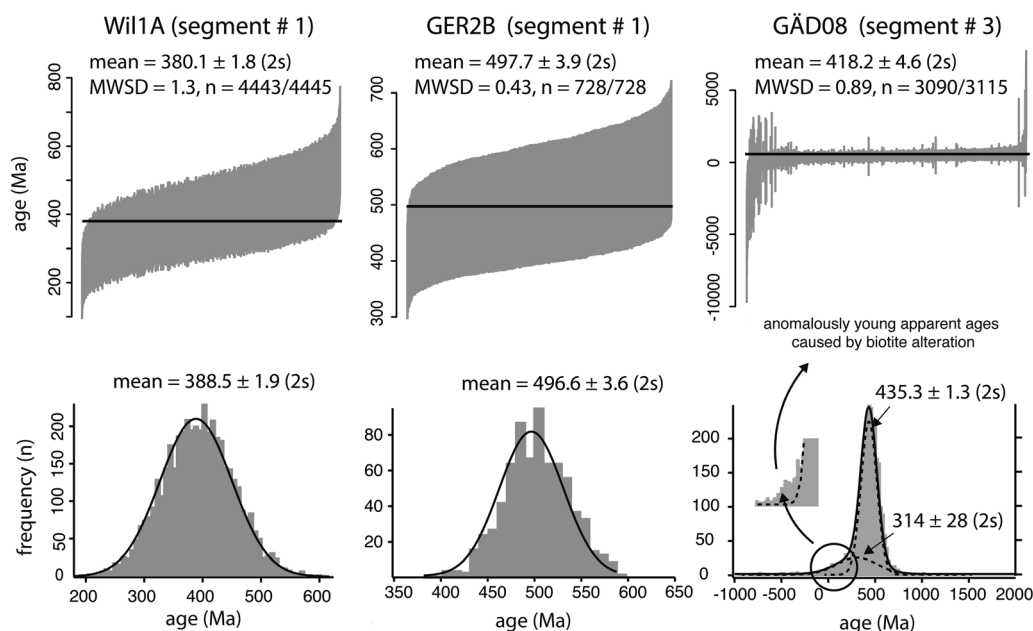


Fig. 8 Rb–Sr ages of selected mica segments within the Rb–Sr age maps (Fig. 5 and 6) expressed as either the weighted means (top) or the mean of a Gaussian fit to the age histograms (bottom). The latter have been generated using the Igor Pro software (version 8.04) and applying the Freedman–Daiconis method to calculate the bin width. The $(^{87}\text{Sr}/^{86}\text{Sr})_{\text{initial}}$ is anchored to the values obtained from plagioclase analyses (Table 8). Ages displayed are without matrix correction. Matrix corrected ages are compiled in Table 9. For WIL1A (segment #1) and GER2B (segment #3) both approaches yield similar means and uncertainties. For GÄD08 segment #3, which comprises all biotite-assigned pixel, the Gaussian mean yields a higher age compared to the weighted mean. This is explained by the presence of anomalously low apparent ages caused by biotite alteration, visible as a shoulder on the lower-age side of the main age peak.



Table 8 Rb–Sr isotope ratios of segments defined in Fig. 5 and 6

Sample	Segment #	Description	<i>n</i> pixel	⁸⁷ Rb/ ⁸⁶ Sr	2SE (abs)	⁸⁷ Sr/ ⁸⁶ Sr	2SE (abs)
GÄD08	1	All muscovite	11 880	22.6977	0.2574	0.9009	0.0026
	2	Small muscovite	247	16.2712	0.1494	0.8463	0.0074
	3	All biotite	3115	283.9290	3.6264	2.5282	0.0253
	4	Small biotite	528	301.9315	6.6887	2.6483	0.0524
	5	Plagioclase	91	0.0209	0.0034	0.73982	0.0141
	6	Plagioclase	33	0.0415	0.0124	0.73849	0.0202
GER2B	1	Muscovite old core domain	728	364.5206	0.9001	3.2830	0.0102
	2	Muscovite new (large)	2515	416.9389	1.0787	2.9717	0.0069
	3	Muscovite new (small)	200	476.5435	4.8434	3.3480	0.0275
	4	Plagioclase	311	0.2086	0.0494	0.74828	0.0103
	5	Plagioclase	311	0.5803	0.094	0.71875	0.0096
	6	Plagioclase	330	3.0187	0.514	0.74263	0.0141
	7	Plagioclase	332	2.2074	0.5278	0.74739	0.0134
	8	Plagioclase	311	2.7003	1.0446	0.75779	0.0163
	9	Plagioclase	311	2.2078	0.5622	0.73119	0.0129
	10	Plagioclase	311	0.6851	0.0938	0.73138	0.0108
	11	Plagioclase	311	0.5853	0.1332	0.72526	0.0138
	12	Plagioclase	332	1.2544	0.3022	0.73971	0.0162
	13	Plagioclase	311	1.0809	0.2174	0.74634	0.0101
	14	Plagioclase	311	0.1868	0.039	0.7216	0.0089
	15	Plagioclase	332	0.1599	0.0228	0.72169	0.0095
WIL1A	1	Muscovite shear zone (large)	4445	150.5048	0.9742	1.5222	0.0094
	2	Muscovite shear zone (small)	216	185.4582	0.8914	1.7341	0.0056
	3	Plagioclase	373	0.19798	0.04556	0.74537	0.0142
	4	Plagioclase	373	1.6672	0.31774	0.73652	0.0152
	5	Plagioclase	397	0.07997	0.04546	0.72325	0.0128
	6	Plagioclase	347	0.03425	0.0314	0.73985	0.0143
	7	Plagioclase	366	0.72295	0.16136	0.72975	0.015
	8	Plagioclase	311	0.64183	0.17836	0.74466	0.0161
	9	Plagioclase	311	0.18682	0.0448	0.74264	0.0153
	10	Plagioclase	311	7.02967	0.69448	0.76065	0.0225
	11	Plagioclase	311	0.31599	0.1363	0.73881	0.0149
	12	Plagioclase	345	0.6681	0.14396	0.73282	0.0156
	13	Plagioclase	311	0.10586	0.02704	0.73492	0.0139
	14	Plagioclase	311	0.04974	0.01838	0.7353	0.0152
	15	Plagioclase	311	0.01382	0.02084	0.71214	0.0153

mounted crystal/rock fragments, which were at least 1 × 1 cm in size (images available at <https://doi.org/10.5281/chemrxiv.8245448>). The variability in both ⁸⁷Sr/⁸⁶Sr is ⁸⁷Rb/⁸⁶Sr is rather large (13–28%, 2SE, <https://doi.org/10.26434/chemrxiv-2023-273lw-v2>) and reflects the Rb/Sr variability within natural mica. For comparison, the isotope ratio variability for NIST610 obtained in session “homogeneity” is 0.1% (2SE) for ⁸⁷Sr/⁸⁶Sr and 0.3% (2SE) for ⁸⁷Rb/⁸⁶Sr. Hence, the mica RM are not suited to correct for mass bias on ⁸⁷Sr/⁸⁶Sr and ⁸⁷Rb/⁸⁶Sr ratios. In contrast, the ⁸⁷Sr/⁸⁷Rb ratios are reproducible within <1% (2SE) in each of the three mica RM (<https://doi.org/10.26434/chemrxiv-2023-273lw-v2>). Since SagaB, Harvard 98973 and Phalaborwa are characterized by a high proportion of radiogenic ⁸⁷Sr (98.7–99.8%; Table 3), precise knowledge of the initial Sr isotope ratio is not required and a common Sr correction can be suspended.²⁰ Hence, the good reproducibility on ⁸⁷Sr/⁸⁷Rb directly implies a good reproducibility of the Rb–Sr age signal

(see Section 2.6). The uncertainty of the weighted mean ⁸⁷Sr/⁸⁷Rb ratio is below 0.4% (2*s_x*, Table 3), if outliers are accepted (0 for Harvard 98973, 1 for SagaB and 2 for Phalaborwa). It is noteworthy that this uncertainty (Table 7) is similar to the one obtained by Rösel & Zack (2022),²⁰ which can be considered a benchmark study for single-spot Rb–Sr dating. They reported 0.7% (2*s_x*, *n* = 9–14) for the Högsbo muscovite (⁸⁷Rb/⁸⁶Sr = 3440–24 000), which has an expected age of 1029 ± 1.7 Ma (ref. 43) and shares a similar Rb/Sr range to the mica RM analyzed in this study (Table 7). However, excess scatter has been detected (1.2% in Harvard 98973, 1.4% in SagaB and 1.7% in Phalaborwa biotite; <https://doi.org/10.26434/chemrxiv-2023-273lw-v2>), suggesting some degree of inhomogeneity in ⁸⁷Sr/⁸⁷Rb ratios beyond the analytical uncertainty. In the uncertainty workflow this excess variance (ϵ_{homo}) is treated as a systematic uncertainty. It is incorporated in quadrature into the uncertainties associated



Table 9 Rb–Sr ages of segments defined in Fig. 6 and 7^a

Sample	Segment #	Description	Gaussian mean				Weighted mean			
			Age (Ma)	2s _x (%)	Matrix corrected	2s* (%)	Age (Ma)	2s _x (%)	Matrix corrected	2s* (%)
GER2B	1	Muscovite old core	496.6	0.7	483.7	2.3	497.7	0.8	484.8	2.2
(⁸⁷ Sr/ ⁸⁶ Sr) _{initial}	2	Muscovite new (large)	383.7	0.4	373.8	2.2	382.7	0.4	372.8	2.1
0.7148 ± 0.0094	3	Muscovite new (small)	387.4	1.0	377.4	2.4	392.8	1.5	382.6	2.5
WIL1A	1	Muscovite shear zone (large)	388.5	0.5	374.7	2.6	380.1	0.5	366.6	2.5
(⁸⁷ Sr/ ⁸⁶ Sr) _{initial}	2	Muscovite shear zone (small)	376.1	2.4	362.8	3.5	375.6	1.9	362.3	3.1
0.712 ± 0.015										
GÄD08	1	All muscovite	424.7	0.7	435.0*	6.8	397.5	2.5	407.2*	7.1
(⁸⁷ Sr/ ⁸⁶ Sr) _{initial}	2	Small muscovite	389.2	26.5	398.7*	27.3	378.5	24.0	387.7*	24.9
0.739 ± 0.011	3	All biotite	Peak 1: 435.3	0.3	445.9	2.5	418.2	1.2	426.2	2.4
			Peak 2: 314	8.9	321.6	9.2				
	4	Small biotite	427.2	1.4	437.6	2.8	414.8	2.8	424.9	3.5

^a 2s_x denotes the uncertainty on the weighted/Gaussian mean. 2s* denotes the total uncertainty resulting from propagating all random and systematic uncertainties. The initial Sr-isotope composition is obtained from plagioclase analyses within the same sample. *Matrix correction based on Phalaborwa biotite, *i.e.*, here there is no chemical matrix match between mica RM and sample.

with all ages calculated using the MCF derived from any of the three secondary mica RM.

3.2. Measurement uncertainty and accuracy by line scan vs. static spot ablation

This study presents Rb–Sr age maps generated using an innovative line scan ablation technique, employing small beam sizes, high scanning speeds, repetition rates, and shallow ablation depth. To evaluate the effectiveness of this method in comparison to the conventional static spot mode ablation, it is crucial to analyze and compare both measurement uncertainty and accuracy. The following assessment is based on repeated analyses of the secondary mica RM during the three spot and three line mode sessions and considers the propagation of all random and systematic uncertainties, as detailed in Section 2.6 and illustrated in Fig. 4 and Table 7.

In this study, we define accuracy as the degree of similarity between the measured ages and the expected ages, quantified as the relative deviation (in %). For all mica RM investigated in this study, high-precision ID-TIMS data is available (Table 4), providing reliable expected ages. In all spot ablation sessions, an improved accuracy of <3.0% is obtained for dating of Harvard 98987 muscovite (−0.1–2.9%) and SagaB biotite by applying a Phalaborwa biotite-based MCF (Fig. 4 and Table 7) compared to the accuracy obtained for non-matrix matched ages (−3.7% to −11.05%). In the case of MicaMg-pressed nanopowder, the matrix correction has the opposite effect and leads to a decrease in accuracy from <2.6% (without matrix correction) to <9.1% (with Phalaborwa-based matrix correction). This observation supports earlier conclusions on the general matrix mismatch between crystalline mica and pressed mica nanopowders.^{21,27,29}

For line scan ablation-based age determinations, application of a Phalaborwa biotite-based MCF leads to a good accuracy for SagaB biotite (−0.8–1.4%, Fig. 4 and Table 7), whereas for Harvard 98973 muscovite, the accuracy even decreases

compared to the non-matrix corrected ages (from 2.7–3.7% to 11.3–11.4%). This implies that both a structural AND a chemical matrix match between the secondary mica RM and unknown sample material are crucial. This is somehow different to spot mode ablation, where accurate ages seem to mainly rely on a structural matrix match, *i.e.* muscovite secondary RM can be equally well used for matrix correction of biotites (Fig. 4).

The total propagated uncertainty on matrix corrected ages obtained during line scan analyses is slightly worse compared to spot mode (1.7–1.9% vs. 2.3–2.8%, Table 7). This is mainly owed to the higher ⁸⁷Sr/⁸⁷Rb ratio uncertainty in line compared to spot mode (2s_x = 0.5–1.7% vs. 0.3–0.7%, Table 7) and the higher long-term excess variance on ⁸⁷Sr/⁸⁷Rb ratios (1.4% in line mode vs. 0% in spot mode; <https://doi.org/10.26434/chemrxiv-2023-273lw-v2>). In contrast, the average accuracy is ~1% better in line mode compared to spot mode when employing matrix correction with chemically matched mica RM. The level of accuracy achieved remains unaffected by the non P/A-matched acquisition of ⁸⁵Rb. Specifically, the Phalaborwa biotite-based MCF, when utilizing ⁸⁵Rb in analog mode, consistently provides the anticipated Rb–Sr age for SagaB biotite, for which ⁸⁵Rb has been acquired in pulse mode. Therefore, the P/A tuning is regarded accurate for ⁸⁵Rb. It is important to note that data acquisition in line scan mode only requires a fraction of the sample volume. A single 50 s line scan on NIST610 during the WIL1A mapping session consists of 5900 shots with a 3 μm beam, while a single spot analysis during session “GER2B/WIL1A” consists of 400 shots with a 65 μm beam. Assuming a constant ablation depth per shot, this results in a ~31.8 times more sample volume during a single spot analysis as opposed to line scan analyses.

3.3. LA-ICP-MS/MS-based Rb–Sr age mapping

3.3.1. GER2B. The timing of the magmatic crystallization and metamorphic (partial) recrystallization of muscovitic mica samples GER2B and WIL1A is well constrained by ID-TIMS



analyses.⁵ Both are samples of pegmatites that formed at ~ 485 to 480 Ma and experienced deformation during a secondary metamorphic event at ~ 382 to 377 Ma. The Rb–Sr age map of sample GER2B (Fig. 6) shows the age distribution within a large deformed mica porphyroclast and in newly crystallized syndeformational muscovite (Fig. 1). The large mica crystal displays old domains with orientations subparallel to the mica cleavage (Fig. 6). While the oldest domains are restricted to the innermost core, patches of partially reset mica reach out all the way to the interface to neighboring quartz (Fig. 6). The elemental maps that have been acquired alongside Rb–Sr isotopes (Fig. 9), show a rather similar and homogenous distribution of Al and Fe in deformed and newly formed muscovites. However, the deformed primary mica is clearly depleted in Rb with respect to the newly crystallized mica, with the latter showing a core-rim zonation with higher Rb concentrations observed in their cores (Fig. 9). Lithium displays a particularly striking behaviour with concentrations that almost perfectly correlate with the age distribution, *i.e.* older age domains display higher Li concentrations (Fig. 9).

For a more detailed assessment of the element and age distribution we inspected a ~ 1.3 mm transect across the core region of the large deformed muscovite (Fig. 10). In this context, we observe two distinct types of domains exhibiting partial age reset. One is marked by a noticeable rise in Rb concentration, with an increase of 50 – $100 \mu\text{g g}^{-1}$ in Rb and a corresponding $\sim 20 \mu\text{g g}^{-1}$ decrease in Li concentration (depicted as blue regions in Fig. 10). In contrast, the other domain experiencing age reduction shows a slight decrease in Rb concentration coupled with an increase in Li concentrations. Given that both Rb and Li have a strong affinity for fluids, we propose that the former domains, exhibiting a significant age reset, record element and isotope mobility induced by fluid activity. The latter type of partial age reset, we tentatively attribute to element and isotope mobility occurring during dry volume diffusion.

To constrain and compare the total age range displayed in the Rb–Sr age map to the ID-TIMS-based ages, we defined three segments: one in the oldest domains of the large mica core, one containing a larger ($200 \times 500 \mu\text{m}$) and one a smaller ($50 \times 150 \mu\text{m}$) newly crystallized mica. For the old segment, both matrix-corrected weighted mean and Gaussian mean approaches yield comparable results (485 ± 11 Ma vs. 484 ± 11 Ma, Table 9) and are in agreement with the ID-TIMS age of the pegmatite crystallization of GER2B (482 ± 5 Ma,⁵ Fig. 11). The Gaussian and weighted mean ages for the latter two segments also agree within their uncertainty covering a range of 373–383 Ma and are in line with the ID-TIMS age of the deformation event (377 to ~ 382 Ma (ref. 5)). As anticipated, the uncertainty in the weighted means ($2\sigma_x$) rises proportionally with the square root of the number of pixels (2515 pixels result in 0.4% uncertainty, while 200 pixels yield 1.0–1.5% uncertainty). Nevertheless, after accounting for all external uncertainties, the enhancement in measurement precision is somewhat mitigated (2515 pixels lead to 2.1–2.2% uncertainty, whereas 200 pixels result in 2.4–2.5% uncertainty).

3.3.2. WIL1A. The Rb–Sr mapping of sample WIL1A probes a white mica aggregate within a high-strain zone running through rather undeformed pegmatite (Fig. 1 and 5). The

formation of the shear zone is most likely related to a regional Devonian-age amphibolite-facies metamorphic event.⁵ This is tested by defining one large (4445 pixel) and one small (216 pixel) segment within muscovite (Fig. 6). Again, the results obtained with Gaussian and weighted mean approaches yield similar Rb–Sr ages (Table 9). However, whereas the large muscovite segment agrees with the anticipated age of the deformation event as constrained by ID-TIMS (Table 9), the small muscovite segment appears slightly younger (363 ± 13 Ma). Since the ID-TIMS ages are interpreted to rather represent maximum age estimates for the waning stages of deformation, it is plausible to consider slightly younger *in situ* ages in the more extensively deformed high-strain zone probed by the WIL1A mapping.

Some regions within the Rb–Sr map display anomalously low ages, in particular around a larger K-feldspar crystal, which is

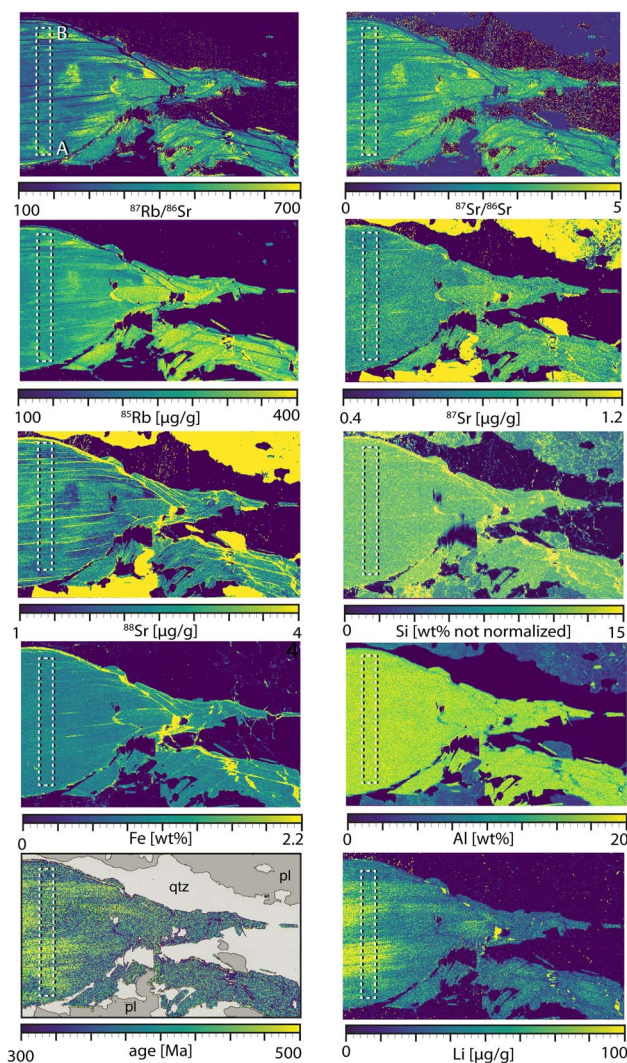


Fig. 9 Elemental, isotopic, isotope ratio and Rb–Sr age distribution of sample GER2B. All concentration maps have been normalized to Si (see Methods section). Hence for Si, only non-normalized concentrations are shown. The dotted square denotes the location of the transect shown in Fig. 10.



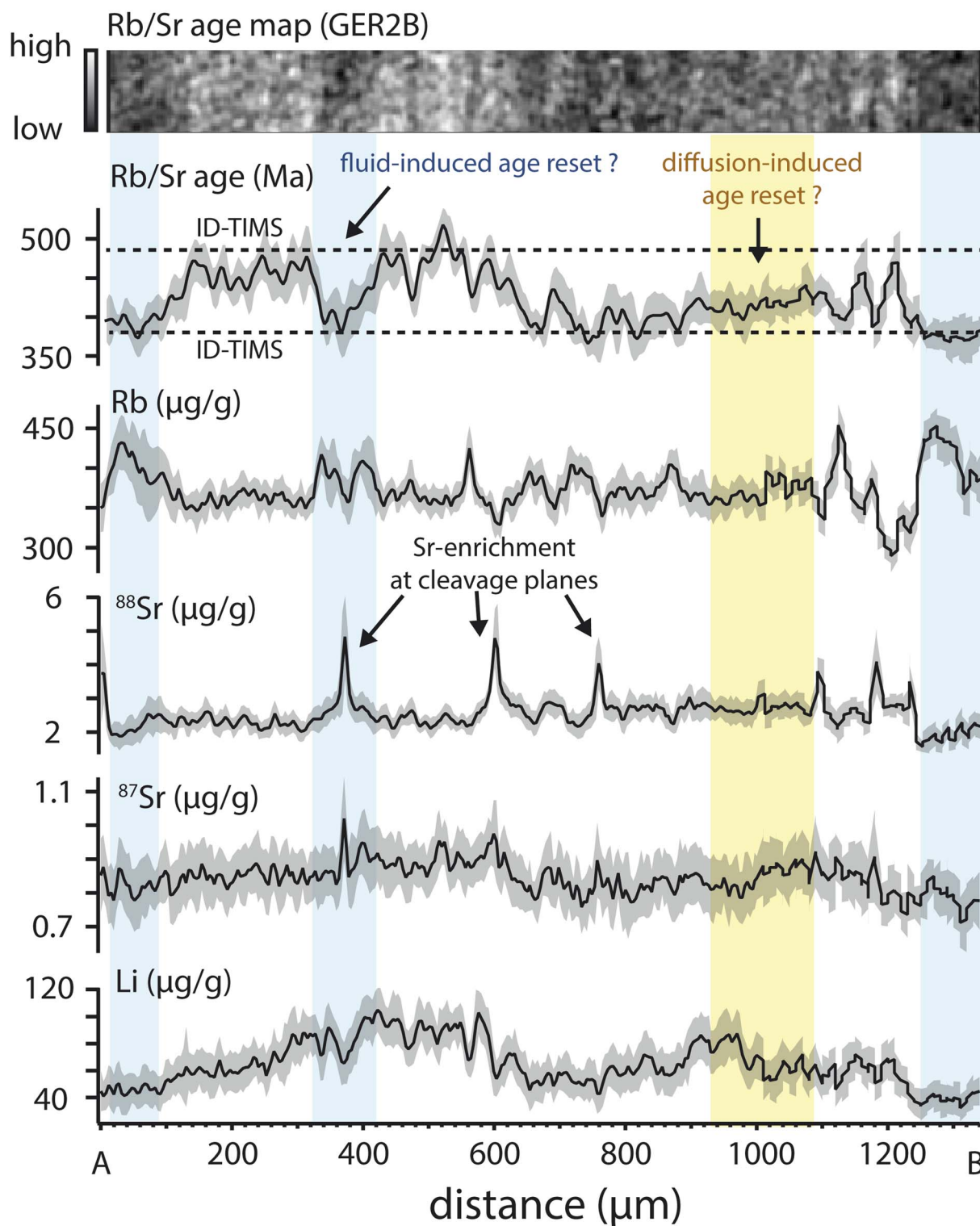


Fig. 10 Distribution of Rb–Sr age, selected elements and isotopes along a transect (A and B, see Fig. 9) perpendicular to the cleavage of the large deformed muscovite crystal displayed in Fig. 6 and 9. Shown are mean values \pm 1SD of all pixel perpendicular to the transect. Blue regions are interpreted as domains affected by fluid-induced age reset. The yellow domain highlights age reset likely caused by dry volume diffusion (see text). The lower and upper age range as constrained by ID-TIMS analyses is indicated by dotted lines. Sr occasionally shows strong enrichment along cleavage planes also visible in the elemental distribution maps in Fig. 9.



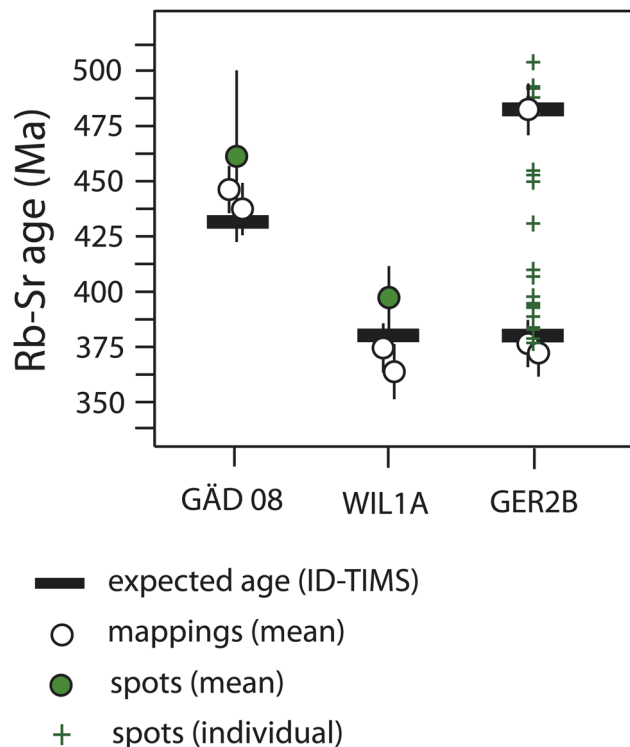


Fig. 11 Comparison of matrix corrected Rb–Sr mica ages obtained from the segments defined in Fig. 5 and 6, matrix corrected Rb–Sr spot ages and expected ages as based on ID–TIMS Rb–Sr multiminerall isochron data (see Section 2.1). All data is based on muscovite analyses, except mapping data for GÄD08, which is based on biotite analyses.

identified by the quantitative chemical mapping (Fig. 6, <https://doi.org/10.26434/chemrxiv-2023-2731w-v2>). This is interpreted as an artefact from mixed K-Fsp/mica analyses, for which mica-based matrix correction does not correctly account for the Rb/Sr fractionation behavior of K-feldspar.

3.3.3. GÄD08. Sample GÄD08 displays a metamorphic mica aggregate comprised of cogenetic biotite and muscovite formed by dynamic recrystallization in the waning stages of ductile deformation at 431.7 ± 3.1 Ma.³⁶ Four segments have been defined to assess the accuracy and precision of Rb–Sr ages as a function of the mica matrix: two segments comprise all muscovite ($n = 11\,880$) and biotite ($n = 3115$) pixel and two segments are smaller subsets of these populations ($n = 247$ for muscovite and $n = 528$ for biotite, Fig. 7). In general, the uncertainty on the weighted mean ($2s_x$) for muscovite (26.5% for 247 pixel) is considerably higher compared to biotite (1.4% for 528 pixel). The higher analytical age uncertainty for muscovite is due to its lower $^{87}\text{Sr}/^{86}\text{Sr}$ ratios. During common Sr correction, a constant value for $(^{87}\text{Sr}/^{86}\text{Sr})_{\text{initial}}$ is subtracted from $(^{87}\text{Sr}/^{86}\text{Sr})_{\text{sample,true}}$ ratios of individual pixels (eqn (4)). As the value of $(^{87}\text{Sr}/^{86}\text{Sr})_{\text{sample,true}}$ approaches $(^{87}\text{Sr}/^{86}\text{Sr})_{\text{initial}}$ the relative variability in the $(^{87}\text{Sr}/^{86}\text{Rb})_{\text{Sr,initial,corrected}}$ ratios—and consequently the Rb–Sr ages—becomes more amplified (see eqn (3)). This is also reflected by the higher age noise displayed by muscovite pixel (Fig. 7). Nevertheless, by pooling a larger number of pixels, e.g. all 11.880 muscovite pixels, we are able to

attain a reasonable weighted mean uncertainty of 0.7% (Table 9). However, it's worth noting that the total uncertainty is significantly influenced by the common Sr correction (γ), which tends to be higher for lower $^{87}\text{Sr}/^{86}\text{Sr}$ ratios.²⁰ As a result, even when all muscovite pixels are included, the total uncertainty is reduced but remains at 6.8% (Table 9). For both muscovite and biotite, matrix correction is based on Phalaborwa biotite and hence is not matrix matched with respect to the muscovite in GÄD08. However, within the limits of uncertainty, both muscovite segments match the ID–TIMS derived Rb–Sr ages (Table 9).

For biotite, improved total uncertainties are achieved and both, small and large segments, yield ages in agreement with ID–TIMS data (425 ± 15 Ma and 426 ± 10 Ma for the weighted mean approach). Interestingly, the age distribution of the “all biotite” segment exhibits an intriguing skewness towards younger ages (Fig. 8). To account for the asymmetry of the histogram, two Gaussians probability distributions were used for fitting. The main age peak displays an age of 446 ± 11 Ma and is assigned to the main deformation event at 431.7 ± 3.1 Ma.³⁶ The peak that fits the population of anomalously young apparent ages at 322 ± 30 Ma is interpreted to reflect fluid alteration leading to chloritization of biotite and hence partially reset ages.⁴⁴ This is supported by Rb–Sr data presented by Bender and co-workers,³⁶ which yield lower mean ages for the apatite plus biotite isochron (421.7 ± 6.2 Ma) compared to the age defined by the feldspar and muscovite fractions (431.7 ± 3.1 Ma). The detection of a younger age population within the age distribution displayed by biotite in the Rb–Sr map is likely a result of the small laser beam size of $3\ \mu\text{m}$. This beam size enables precise sampling of altered biotite domains. In contrast, both ID–TIMS analyses and conventional spot ablation with larger beam sizes more likely lead to mixed analyses of pristine and altered biotite, which imposes problems in interpreting the obtained ages.

3.4. Single spot Rb–Sr ages

Since in single spot mode, both $\sim^{86}\text{Sr}$ and $\sim^{88}\text{Sr}$ were recorded instead of only $\sim^{88}\text{Sr}$ in line scan mode, ages obtained using either Sr-isotope can be compared. The age range displayed by the single spots on GER2B based on $\sim^{86}\text{Sr}$ (youngest age 376 ± 22 ; oldest age: 505 ± 28 Ma, Fig. 11) encompasses the age range constrained by ID–TIMS. However, if $\sim^{88}\text{Sr}$ is used for age calculation, individual spot ages are up to 5–18 Ma years older (Table 10), indicating an adverse impact on accuracy. This is also exemplified by the weighted mean age of all spot analyses obtained for WIL1A and GÄD08, for which only $\sim^{86}\text{Sr}$ -based ages (396 ± 14 Ma and 462 ± 39 Ma) are in agreement with the expected ages ($377\text{--}382$ Ma and 431.7 ± 3.1 Ma).^{5,36} Since in spot mode, the acquisition of $\sim^{88}\text{Sr}$ is not P/A matched in NIST610 and muscovite as opposed to $\sim^{86}\text{Sr}$, the observed bias points to a failure of the employed P/A tuning procedure (see Methods section) for $\sim^{\text{J}}\text{Sr}$ nuclides. This is somewhat disappointing given the better total uncertainty of $\sim^{88}\text{Sr}$ vs. $\sim^{86}\text{Sr}$ -based single spot ages (Table 10), due to the higher natural abundance of ^{88}Sr as compared to ^{86}Sr .



Table 10 Rb–Sr single spot ages^a

Sample	^b Nuclides: ⁸⁵ Rb, ^{~86} Sr, ^{~87} Sr			Nuclides: ⁸⁵ Rb, ^{~88} Sr, ^{~87} Sr		
	Age (Ma)	Age matrix corrected (Ma)	2s* (%)	Age (Ma)	Age matrix corrected (Ma)	2s* (%)
GER2B	460	489	6.9	474	505	2.8
	464	493	5.3	469	499	2.3
(⁸⁷ Sr/ ⁸⁶ Sr) _{ini}	454	483	5.8	466	496	2.1
0.7148 ± 0.0094	465	495	5.4	482	513	2.3
	475	505	5.5	490	522	2.6
	364	388	6.3	380	404	2.3
	382	406	5.0	386	411	2.6
	373	397	6.0	387	411	2.2
	426	454	5.8	440	468	2.7
	423	451	5.8	435	463	2.3
	404	430	4.9	409	435	2.5
	428	455	5.2	434	462	2.4
	370	393	5.1	380	405	2.3
	368	392	5.9	379	403	2.6
	353	376	5.7	365	389	2.6
	355	378	5.3	365	388	2.4
	359	382	5.1	366	389	2.2
	359	382	5.0	374	398	2.2
	369	393	6.4	379	404	2.2
	384	409	5.8	390	415	2.6
WIL1A	378	402	4.1	393	418	3.3
	384	408	3.8	395	420	3.0
(⁸⁷ Sr/ ⁸⁶ Sr) _{ini}	362	385	4.6	369	392	3.7
0.712 ± 0.015	373	397	4.9	388	412	3.8
	378	402	4.3	395	420	3.5
	355	378	4.2	373	397	3.0
	354	377	5.3	371	395	3.7
	390	415	4.3	408	434	4.0
	362	386	4.5	389	414	3.8
	387	412	4.5	405	431	4.1
Weighted mean (WIL1A)		396.3			411	
MWSD		2.5			3.7	
2s _x (%)		2.5			2.5	
2s* (%)		3.4			3.3	
GÄD08	434	456	7.0	494	519	6.0
	463	487	6.8	527	554	5.7
(⁸⁷ Sr/ ⁸⁶ Sr) _{ini}	444	466	8.2	538	566	6.6
0.739 ± 0.011	437	460	7.3	494	519	6.5
	448	471	6.5	503	528	5.7
	422	455	8.1	500	525	6.7
	429	451	9.5	543	570	7.3
	432	454	10.5	556	585	7.9
*	432	454	10.5	556	585	7.9
*	424	446	8.6	498	523	7.1
*	443	465	10.1	570	599	7.6
Weighted mean (GÄD08)		462.0			543.0	
MWSD		0.42			1.6	
2s _x (%)		2.4			2.4	
2s* (%)		8.4			6.9	

^a (⁸⁷Sr/⁸⁶Sr)_{initial} is anchored to the indicated value, which is obtained from plagioclase analyses within the same sample. Matrix correction is based on Harvard 98973 muscovite. Spot locations are indicated in Fig. 6 and 7. There, only ages based on ^{~86}Sr are shown. ^b Nuclides used for calculating the single spot ages. * spots obtained within the mapped area. The other spots are obtained on two adjacent mica grains. 2s_x denotes the uncertainty on the weighted/Gaussian mean. 2s* denotes the total uncertainty resulting from propagating all random and systematic uncertainties.

As suggested by Rösler & Zack (2022)²⁰ the total uncertainty for both individual and pooled spot ages is a function of the ⁸⁷Sr/⁸⁶Sr ratio. Muscovite in GÄD08 exhibits the lowest ⁸⁷Sr/⁸⁶Sr ratios (0.85–0.90) and hence also the highest single spot age

uncertainty (6.5–10.5%). In contrast, in Wil1A and GER2B (⁸⁷Sr/⁸⁶Sr = 1.5–3.3) the single spot uncertainty is considerably lower (3.8–6.9%).



4. Conclusions & outlook

This study demonstrates that precise and accurate Rb–Sr ages can be obtained based on Rb–Sr isotope mapping with μm -scale spatial resolution using much smaller sample volumes than required for conventional spot ablation. This technique opens new possibilities for geochronological applications. It allows for the dating of μm -sized shear zones and small crystals that are either tightly intergrown with other phases or too small for conventional spot analyses. This includes the analysis of usually small detrital mica grains, which is a promising approach for sedimentary provenance analysis.²⁰

Another notable strength of Rb–Sr mapping lies in its capacity to identify and pool homogeneous age populations, which is essential for meaningful weighted mean age determinations.³⁷ This is particularly important, given the presence of sharp intracrystalline age gradients (~ 100 Ma over 20–30 μm) and larger domains displaying partial age reset (Fig. 10). These μm -scale age variations correspond to the identification of multiple age populations in natural biotite, as observed in the time-resolved signal of a single spot ablation interval.²¹ While increasing the size of a particular age population segment improves the weighted mean uncertainty, the obtainable total uncertainty of the Rb–Sr age is limited by the propagation of the uncertainty of the initial Sr correction and other systematic uncertainties. In this respect, the availability of matrix-matched, homogeneous, and well characterized RM as well as an accurate and precise determination of the initial $^{87}\text{Sr}/^{86}\text{Sr}$ is crucial. Furthermore, it is imperative to ensure a P/A match for mass-shifted Sr-ions, given the absence of a viable software-based P/A tuning method. Developing alternative routines for P/A factor determination and correction for P/A mismatch will advance analytical accuracy and precision.

As shown, the quadrupole-based mapping approach not only provides age data but also offers valuable quantitative elemental information. This dual capability enhances our understanding of the formation and timing of economically important ore deposits, encompassing even light elements like lithium. Analyzing these lighter elements alongside Rb/Sr poses challenges, particularly when employing a Wien field to restrict the mass range transmitted to the CRC.²¹ Rb–Sr age maps are also highly promising for thermochronology applications, as they allow to constrain diffusion profiles and to distinguish diffusion (at static conditions) from fluid-induced age resetting or dynamic recrystallization. This capability is pivotal for the much-needed re-evaluation of true diffusion-related closure temperatures in natural biotite and muscovite.

Data availability

Supplementary data can be accessed through the ChemRxiv® server (<https://doi.org/10.26434/chemrxiv-2023-273lw-v2>). Here both raw and corrected isotope ratios of reference materials and samples of all six sessions are available as well the uncertainty components and the P/A configuration for ^{85}Rb and ^{87}Sr nuclides. Furthermore, an excel sheet including the Rb/Sr isotope ratios and Rb–Sr ages of every individual pixel

within the mica segments outlined in the mapping is included. Additionally, high-resolution, unedited elemental, isotope ratio, and Rb–Sr age maps of all mappings are provided.

Conflicts of interest

There are no conflicts to declare.

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