

Preparation and cell refinement of mica microsamples

Autor(en): **Stern, W.B.**

Objekttyp: **Article**

Zeitschrift: **Schweizerische mineralogische und petrographische Mitteilungen
= Bulletin suisse de minéralogie et pétrographie**

Band (Jahr): **71 (1991)**

Heft 1

PDF erstellt am: **26.04.2024**

Persistenter Link: <https://doi.org/10.5169/seals-54353>

Nutzungsbedingungen

Die ETH-Bibliothek ist Anbieterin der digitalisierten Zeitschriften. Sie besitzt keine Urheberrechte an den Inhalten der Zeitschriften. Die Rechte liegen in der Regel bei den Herausgebern.

Die auf der Plattform e-periodica veröffentlichten Dokumente stehen für nicht-kommerzielle Zwecke in Lehre und Forschung sowie für die private Nutzung frei zur Verfügung. Einzelne Dateien oder Ausdrucke aus diesem Angebot können zusammen mit diesen Nutzungsbedingungen und den korrekten Herkunftsbezeichnungen weitergegeben werden.

Das Veröffentlichen von Bildern in Print- und Online-Publikationen ist nur mit vorheriger Genehmigung der Rechteinhaber erlaubt. Die systematische Speicherung von Teilen des elektronischen Angebots auf anderen Servern bedarf ebenfalls des schriftlichen Einverständnisses der Rechteinhaber.

Haftungsausschluss

Alle Angaben erfolgen ohne Gewähr für Vollständigkeit oder Richtigkeit. Es wird keine Haftung übernommen für Schäden durch die Verwendung von Informationen aus diesem Online-Angebot oder durch das Fehlen von Informationen. Dies gilt auch für Inhalte Dritter, die über dieses Angebot zugänglich sind.

Frau Prof. Dr. Emilie Jäger gewidmet

Preparation and cell refinement of mica microsamples

by *W. B. Stern*¹

Abstract

Least squares refinement of X-ray powder diffractograms of mica microsamples is possible when preferred orientation is minimized electrostatically. Their relative 1-sigma errors for a, b, c and cell volume are 0.06, 0.04, 0.02 and 0.05% respectively. The correlation of a and b parameters is within the statistical error of data, but a, b versus c scatter far more. When literature cell parameters of pure end members muscovite-phengite-paragonite are combined in a triangular grid and used for a diffraction analysis of white 2M1-mica, the results obtained correspond trendwise only with chemical analyses performed on the same micas. One may conclude that even qualitative chemical analysis on microsamples (e.g. by energy-dispersive X-ray fluorescence) enables more reliable results than high-quality diffraction data. Though cell parameters a and b of white mica are certainly linked with "phengite"-, and c with "paragonite"-content, the interdependence seems to be more complex than expected from literature.

Keywords: 2M1-mica, cell refinement, X-ray diffraction, microsample, chemical composition.

Zusammenfassung

Zellverfeinerungen pulverdiffraktometrischer Aufnahmen von Mikroproben sind im Falle von Hellglimmer möglich, wenn Orientierungseffekte elektrostatisch reduziert werden. Typische 1-sigma-Relativfehler für a, b, c und Zellvolumen sind 0,06, 0,04, 0,02 und 0,05% für 2M1-Hellglimmer, von denen 53 unterschiedlicher Herkunft diffraktometrisch und röntgenfluoreszenzanalytisch untersucht worden sind. Werden aus der Literatur die b- und c-Zellparameter von Muskowit, Phengit und Paragonit in einem Dreiecksdiagramm kombiniert und zur diffraktometrischen «Analyse» von Hellglimmer verwendet, so zeigen die gefundenen Werte zwar eine ungefähre Korrelation mit den an denselben Proben erhobenen chemischen Daten; von einem quantitativen Zusammenhang kann aber – trotz der kleinen Relativfehler (XRD) – keine Rede sein. Offensichtlich sind die Beziehungen zwischen Gittergrösse und Chemismus komplexer, als z.B. durch die Bezeichnung «Phengit» zum Ausdruck kommt, bei dem Mg, Fe²⁺ und Fe³⁺ einen unterschiedlichen und wohl auch gegenläufigen Einfluss auf die a- und b-Parameter haben können.

Introduction

Lattice dimensions of crystals are essential parameters, but not easy to determine on low-symmetry flaky minerals like mica. When chemical data, based on bulk mineralogical (powdered) specimens are correlated with lattice parameters, the latter should be determined on powdered samples as well (not on single crystals), preferably on the same specimen, from which chemical information was obtained.

A recently developed preparation method for diffraction and chemical investigation on micro-

samples (HANDSCHIN, STERN, 1990) was tested by studying dioctahedral micas from schists, gneisses, granites and pegmatites of various origin.

Procedures and results

X-RAY DIFFRACTION (XRD)

When microsamples have to be investigated by X-ray diffraction or -fluorescence, the size of sample surface is important in order to get statistically relevant signals. Thus, 30 mg of powdered mica

¹ Mineralogisch-Petrographisches Institut der Universität, Bernoullistrasse 30, CH-4051 Basel.

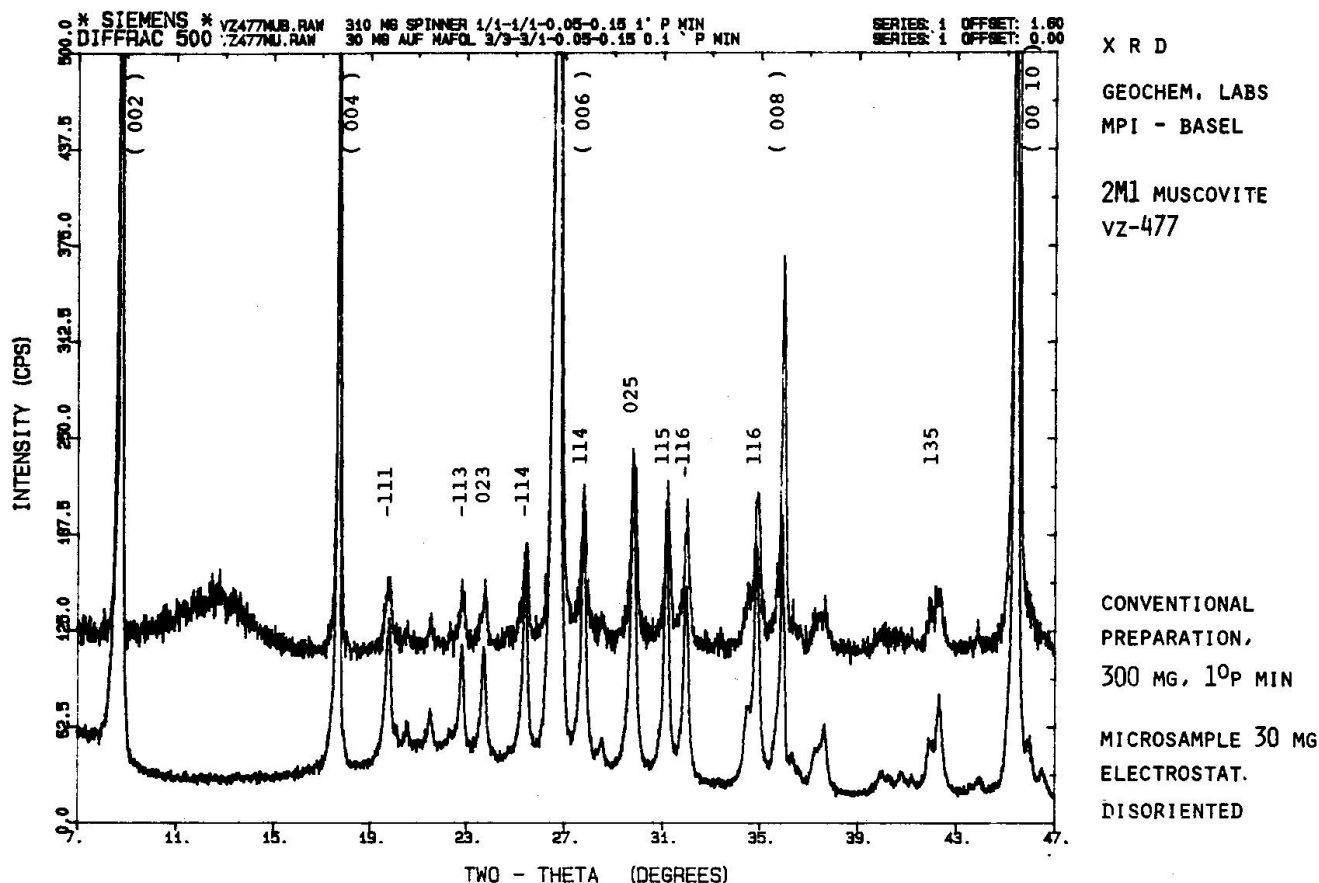


Fig. 1 X-ray diffraction pattern of powdered 2M1 – muscovite (Vz 477): Comparison between conventional preparation (300 mg in sample cup) and electrostatically disoriented microspecimen (30 mg) on stretched foil. Non-basal reflections of the disoriented microsample display double net intensities.

(10 mg would do as well for diffraction work) were distributed evenly on a stretched foil of 40 mm diameter, and fixed to it with 0.25 ml Griltex solution. In order to minimize preferred orientation, the mica flakes were disordered electrostatically by moving a plexiglass rod close to the surface of the drying mica / Griltex film.

In contrast to conventional mounting (e.g. 300 mg powdered mica in a sample cup or smear slides) not only basal spacings (00l) are prominent, but also random hkl-reflections (Fig. 1). Since least squares cell refinement (APPLEMAN, EVANS, 1973) of a monoclinic structure needs around 20 linearly independent strong reflections, it is evident that only disoriented samples can be used. Even these diffraction patterns may lead to plausible, but nevertheless erroneous cell parameters (STERN, 1987), when statistics of signal/peak ratios are not appropriate.

In order to combine suitable resolution and signal statistics, the samples were run with an angular goniometer speed of 0.1 degrees 2 θ only (around 12 hours per exposure); least squares refinements were performed on-line after data reduction, the results transferred to a Lotus worksheet (reg. trade

name LOTUS 1-2-3, version 3.0), (Fig. 2, Tab. 1). The statistical average errors are ± 0.003 Å for a, 0.004 for b and c, and 0.4 Å³ for the cell volume respectively.

The correlation of a and b parameters is fair and corresponds to literature data, e.g. from BORG and SMITH, 1969 (Fig. 2).

X-RAY FLUORESCENCE (XFA)

During the past few years a large amount of micas and "coexisting" mica pairs was re-examined with improved wavelength- and energy-dispersive X-ray fluorescence methods (WD-XFA, ED-XFA) for main constituents and trace elements, part of these unpublished data were used here to correlate them with XRD data.

Quantitative WD-XFA is still (STERN, 1979) executed on fused minerals for main element analysis, and on pressed powders for trace element analysis, both routines taking advantage of fully automated procedures optimized for intensity (matrix-) corrections. All data were automatically transferred to a Lotus worksheet file for further processing and graphical display.

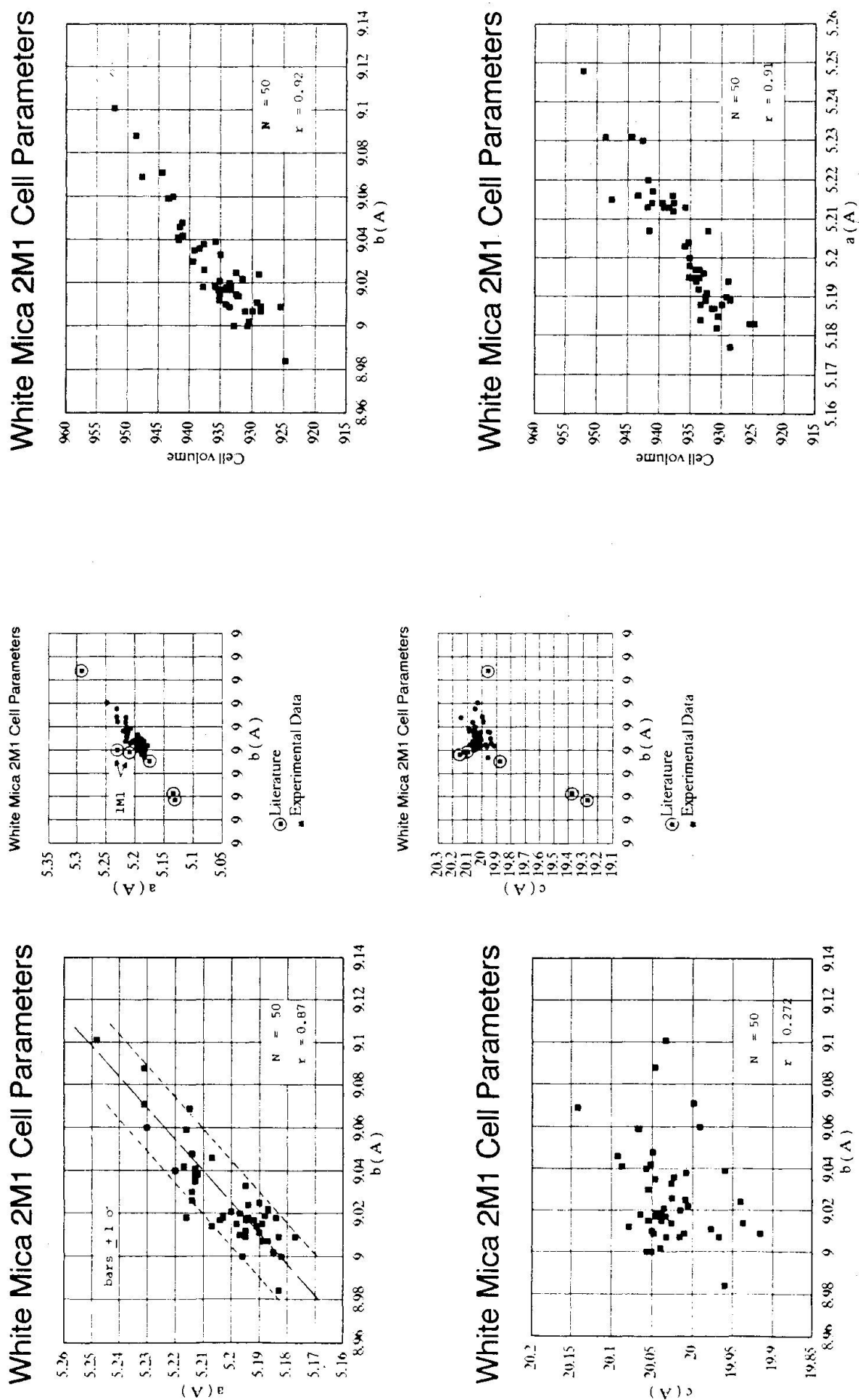


Fig. 2 Experimental X-ray powder diffraction data, and values from literature (for key see Fig. 4). Correlation of lattice parameters a versus b , and a , b versus cell volume are highly significant, in contrast to b versus c . Position and slope of the a versus b regression line corresponds well with data from literature (small inset). Note that 1M polytypes tend to plot outside the a/b correlational field of 2M1 white mica.

Tab. 1a Data White Mica (XRD) and selected chemical data (XFA).

Tab. 1a

Data White Mica (XRD) and selected chemical data (XFA).

2M1 Sample	Origin	Host Grp	a (Å)	b (Å)	c (Å)	error	beta	volume	Std. err.	log-ratios ex-XFA K/Rb, Ba Si/Al Al/Fe, K/Rb Hk=Oxide	Fe/Mg	ED-XFA Fe203t	Specimen no.	Calculated (XFA) Pheng Si/Al Na/Na+ %	
Ad41b	Bernardino	Sh	5.213	0.002	9.039	0.003	19.960	0.002	0.3	0.012	-0.051	0.236	0.610	40.1	59.9
Cal26	Calanca	Sh	5.200	0.002	9.021	0.002	20.035	0.001	0.3	0.012	-0.029	0.182	0.820	27.0	4.6
Gli11	Bernardino	Sh	5.231	0.004	9.071	0.005	19.999	0.003	0.7	0.016	-0.009	0.255	0.446	44.7	0.0
Ha225	Laengtal	Sh	5.190	0.003	9.011	0.004	19.977	0.006	0.5	0.019	-0.133	0.373	0.660	73.4	13.1
KAW0165	Eisten	Gn	5.230	0.003	9.060	0.004	19.991	0.003	0.5	0.015	-0.003	0.285	0.360	52.0	0.7
Vals1	Vals	Gn	5.194	0.002	9.024	0.003	19.940	0.003	0.3	0.015	-0.046	0.265	0.569	47.1	52.9
Bedr107a	Bedretto	Gn	5.207	0.003	9.014	0.008	19.937	0.001	0.7	0.011	-0.013	0.247	0.580	42.8	0.7
Blen49	Blenio	Sh	5.217	0.005	9.042	0.009	20.052	0.004	0.9	0.022	-0.010	0.138	0.951	16.3	83.7
Cr22d	Levent	Gn	5.185	0.003	8.992	0.006	19.887	0.030	0.6	0.016	-0.076	0.093	1.410	5.3	14.9
Gli04	Chiavenna	G	5.182	0.004	9.000	0.005	20.056	0.002	0.6	0.022	-0.010	0.128	1.180	13.9	1.1
Hal99	Tosafall	Gn	5.213	0.001	9.036	0.003	20.023	0.001	0.4	0.015	-0.009	0.190	0.690	28.9	7.8
KAW0160	Leberdun	Gn	5.212	0.002	9.038	0.004	20.008	0.002	0.4	0.017	-0.006	0.192	0.600	29.4	0.7
KAW0207	Binnental	Gn	5.248	0.002	9.101	0.002	20.033	0.004	0.4	0.022	-0.008	0.275	0.420	49.6	0.4
Hul132	Antigorio	Sh	5.189	0.003	9.007	0.004	19.967	0.001	0.5	0.012	-0.043	0.121	0.860	12.2	7.6
HuSt900	Bavona	Gn	5.214	0.004	9.026	0.005	20.025	0.002	0.6	0.018	-0.022	0.233	1.036	39.4	4.1
KAW0082	Croppo	Gn	5.214	0.003	9.048	0.011	20.049	0.002	0.9	0.018	-0.007	0.144	0.920	17.7	1.6
KAW0083	Beura	Gn	5.216	0.004	9.018	0.005	20.040	0.001	0.5	0.016	-0.004	0.171	0.810	24.3	0.3
Vz691	Verzasca	Sh	5.183	0.001	8.984	0.002	19.960	0.002	0.3	0.017	-0.410	0.110	1.175	9.5	3.8
Gli10	Prato	P	5.203	0.003	9.019	0.003	20.044	0.001	0.3	0.008	-0.018	0.142	0.804	17.3	2.3
Vz297	Lavertezzo	P	5.195	0.001	9.009	0.002	20.048	0.001	0.2	0.016	-0.015	0.155	1.010	2.32	2.4
Vz477	Odno	P	5.216	0.003	9.059	0.002	20.066	0.004	0.5	0.019	-0.008	0.159	0.700	4.1	7.7
Vz501	Lavertezzo	P	5.187	0.001	9.007	0.001	20.032	0.001	0.2	0.011	-0.025	0.142	0.930	2.62	2.9
WS62a	Morbobbia	P	5.198	0.004	9.015	0.004	20.055	0.003	0.5	0.020	-0.014	0.126	1.156	1.79	1.7
WS72b	Novate	G	5.185	0.002	9.002	0.003	20.039	0.002	0.3	0.014	-0.009	0.115	1.030	1.80	2.7
81.200	Irkutsk	P	5.213	0.004	9.035	0.006	20.046	0.002	0.5	0.019	-0.032	0.117	0.950	1.83	2.35
83.330	Rustum	P	5.190	0.003	9.025	0.002	20.009	0.002	0.4	0.015	-0.020	0.213	0.432	4.66	8.52
86.321	Kali-b	P	5.196	0.003	9.000	0.003	20.051	0.001	0.4	0.011	-0.036	0.107	1.126	8.7	6.9
86.323	Sharda	P	5.195	0.004	9.033	0.004	20.026	0.002	0.5	0.017	-0.030	0.149	0.782	1.28	1.94
86.326	Badani	P	5.191	0.002	9.014	0.003	20.026	0.001	0.3	0.011	-0.021	0.117	0.985	2.68	4.11
88.003gr	Morocco	P	5.184	0.002	9.018	0.003	20.064	0.003	0.4	0.017	-0.031	0.108	1.185	1.67	2.14
88.005ru	Morocco	P	5.194	0.004	9.018	0.004	20.038	0.003	0.6	0.019	-0.031	0.128	0.936	2.88	3.35
88.008	Argentina	P	5.204	0.002	9.017	0.001	20.033	0.002	0.3	0.015	-0.029	0.149	0.681	3.82	5.50
88.010	Shivsh	P	5.195	0.003	9.017	0.003	20.046	0.004	0.3	0.011	-0.033	0.099	1.172	1.75	1.70
89.001	Sitarama	P	5.197	0.001	9.020	0.003	20.015	0.002	0.2	0.012	-0.040	0.117	1.050	1.82	2.02
89.003	Menakshi	P	5.187	0.002	9.022	0.004	20.006	0.002	0.4	0.017	-0.041	0.131	0.860	2.30	3.51
89.005gr	PVSR-India	P	5.231	0.002	9.088	0.003	20.046	0.006	0.4	0.018	-0.008	0.174	0.532	2.30	3.51
89.006	Vandana	P	5.195	0.004	9.012	0.005	20.078	0.004	0.6	0.019	-0.015	0.101	1.230	6.47	7.90
90.009	Tucuman	P	5.216	0.003	9.059	0.011	20.067	0.004	0.9	0.028	-0.013	0.111	0.949	2.07	3.10
90.010	Argentina	P	5.220	0.002	9.040	0.004	20.057	0.002	0.3	0.019	-0.010	0.075	0.982	1.80	3.13
90.011	Zimbabwe	P	5.214	0.002	9.030	0.003	20.055	0.002	0.3	0.017	-0.010	0.064	1.059	2.81	3.10
89.009	Tanzania	P	5.215	0.004	9.069	0.014	20.141	0.009	1.6	0.038	-0.023	0.118	0.996	1.30	2.23
Shilling	Sudan	P	5.177	0.003	9.009	0.006	20.010	0.003	0.7	0.019	-0.027	0.149	0.861	2.14	3.00
086.010	Malaysia	G	5.197	0.002	9.010	0.005	20.050	0.005	0.6	0.020	-0.035	0.228	0.839	3.32	4.5
094.004	Malaysia	G	5.213	0.002	9.041	0.003	20.037	0.005	0.3	0.015	-0.017	0.133	0.910	5.19	3.64
094.008	Malaysia	G	5.207	0.001	9.046	0.006	20.092	0.004	0.3	0.020	-0.010	0.150	0.900	2.47	3.47
094.009	Malaysia	G	5.188	0.001	9.019	0.005	20.045	0.003	0.5	0.023	-0.017	0.150	0.990	2.41	2.68
094.010	Malaysia	G	5.203	0.002	9.018	0.003	20.044	0.001	0.3	0.015	-0.013	0.144	0.960	4.09	2.91
094.014	Malaysia	G	5.189	0.001	9.015	0.003	20.038	0.004	0.1	0.014	-0.013	0.152	0.840	4.11	3.95
094.015	Malaysia	G	5.192	0.002	9.017	0.003	20.045	0.003	0.4	0.013	-0.006	0.159	0.880	2.61	3.37
103.001	Malaysia	G	5.188	0.003	9.007	0.007	20.016	0.011	0.8	0.033	-0.019	0.232	0.900	3.67	3.11
Polytypes other than 2M1															
KAW0041	Nalps	P	8.878	0.010	5.249	0.003	20.420	0.003	0.8	0.021	-0.007	0.134	1.110	1.98	2.30
Vz690	Verzasca	Sh	5.169	0.005	8.991	0.003	10.145	0.003	0.5	0.007	-0.068	0.167	1.098	0.83	1X962

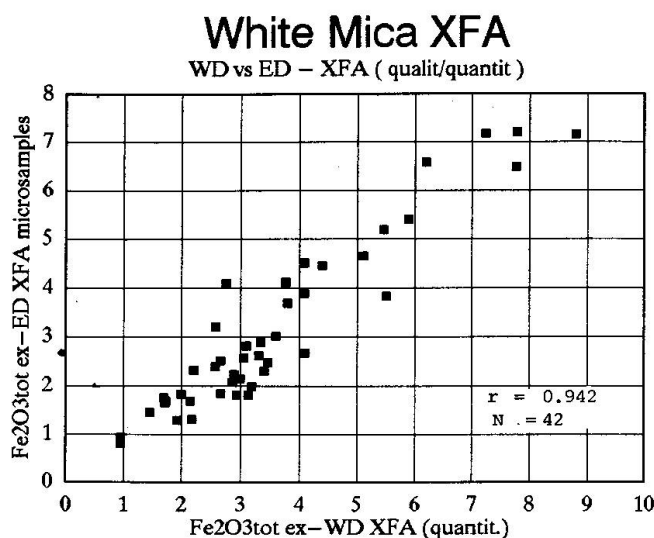


Fig. 3 Comparison of quantitative and qualitative X-ray fluorescence data (Fe_2O_3 total).

Qualitative: 30 mg powdered mica on stretched foil (as for x-ray diffraction), ED-XFA

Quantitative: fusion with $\text{Li}_2\text{B}_4\text{O}_7$ (Tab. 2), WD-XFA

The correlation coefficient ($r=0.942$, $N=42$) is better than the one obtained by phengite analysis XRD versus XFA.

Some microsamples which were examined by XRD were analyzed qualitatively by ED-XFA (STERN, 1985) in order to examine the reliability of this fast and efficient method of non-destructive simultaneous instrumental analysis (Fig. 3).

The analyzed mica concentrates cover a wide, though not the complete compositional field of dioctahedral mica. They represent various types of host rock, like Alpine schists, gneisses, granites and pegmatites, Triassic granites, and Pre-cambrian pegmatites from Argentina, India, Sudan and Tanzania.

Discussion

Cell parameters of white mica display a large variation, much larger than the error of data determination (ref., see Fig. 4):

	phengite	muscovite	paragonite
a	5.291	5.183	5.135
b	9.169	8.990	8.993
c	19.947	20.152	19.270

MUSCOVITE S.S.

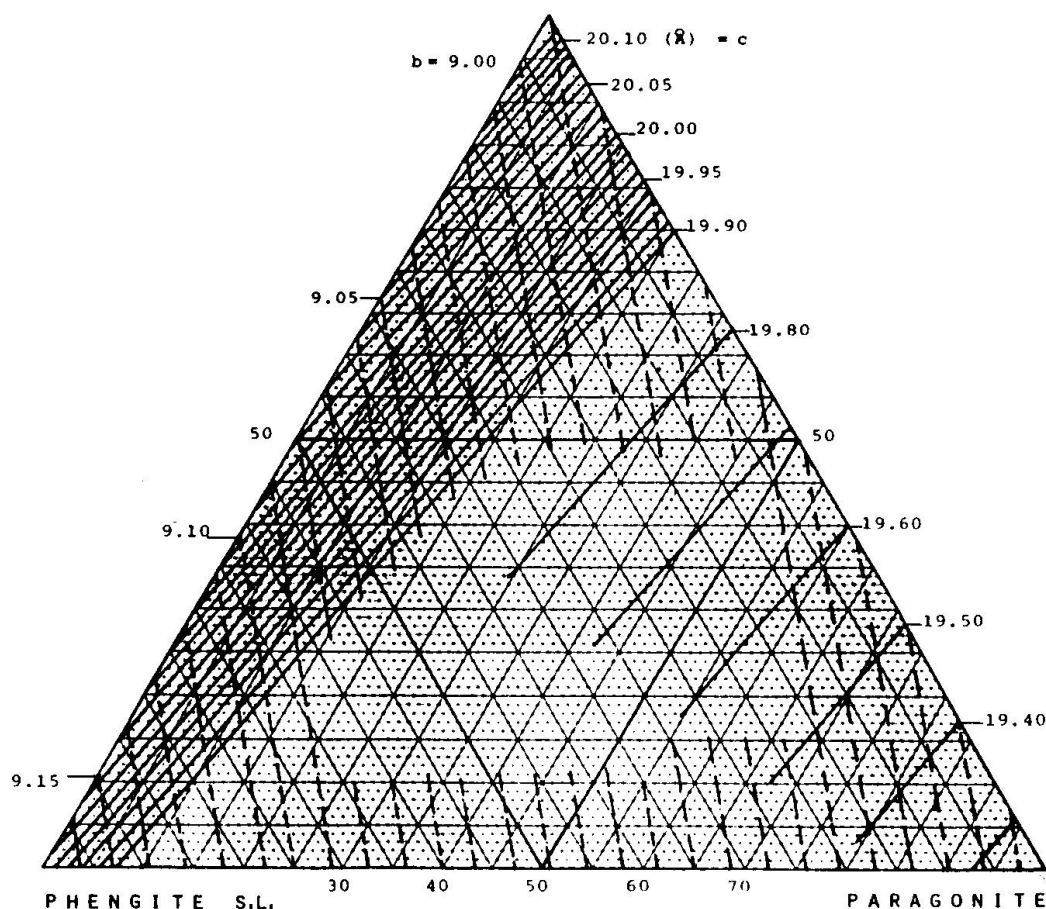


Fig. 4 Triangular plot of 2M1 dioctahedral micas muscovite – phengite – paragonite. Cell parameters b and c taken from literature: muscovite b, c after CHATTERJEE, JOHANNES, 1974; paragonite b, c after BORG and SMITH, 1969; phengite b after GUIDOTTI et al., 1989; phengite c after BORG and SMITH, 1969. The diagram enables – theoretically – the deduction of mica composition (% muscovite, paragonite, phengite) from experimentally determined cell parameters b and c. Far more reliable, and commonly used, is the direct chemical analysis.

The values for *a* and *b* have been correlated with the exchange of octahedral Al by Fe²⁺ and Mg combined with exchange of tetrahedral Al³⁺ by Si (Tschermak substitution, see e.g. GUIDOTTI et al., 1989). The value for *c* has been attributed to the exchange of interlayer K by Na (paragonite substitution, see e.g. CIPRIANI et al., 1968) – among other, less reported substitutions.

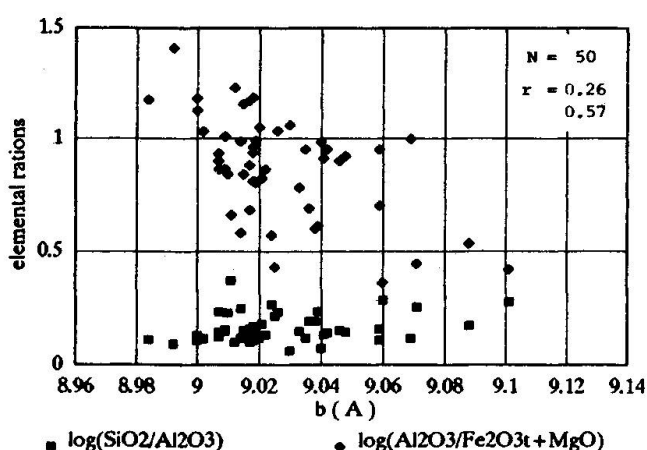
Corresponding correlation data published so far display a trendwise interdependence between diffraction and relevant chemical data, but the uncertainty of the correlation has been such that a quantitative use of *b*- or *c*-parameters for phengite or paragonite analysis seems hardly possible.

When *b*- and *c*-parameters of pure end members phengite, paragonite and muscovite (taken from literature) are combined in a triangular grid, it

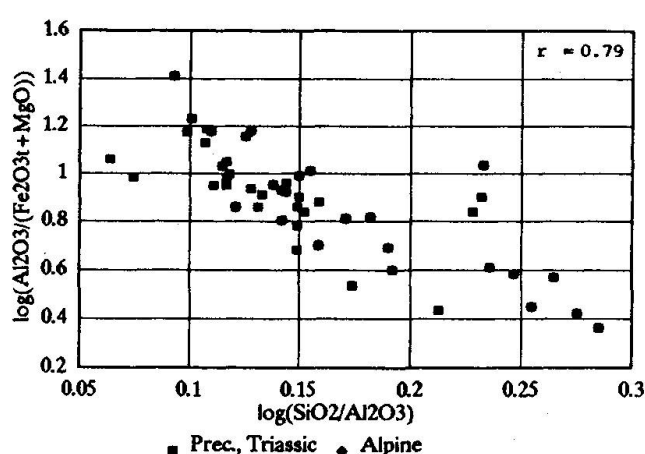
is – theoretically – possible to determine e.g. the phengite or paragonite content of an unknown white mica (Fig. 4). If results of such a procedure, however, are controlled by chemical analyses, the agreement of diffractometric and chemical data is not too encouraging:

	phengite		paragonite		muscovite s.s.		
	XRD	XFA	XRD	XFA	XRD	XFA	(%)
Gli-04	8	14	7	1	85	85	
Gli-11	19	45	20	0	61	65	
KAW-083	21	24	6	0	73	75	
KAW-160	29	29	8	1	63	70	
KAW-165	42	52	6	1	52	47	
KAW-207	60	50	0	0	40	50	
Vz-477	35	21	0	2	65	77	
WS-72b	10	11	6	1	84	88	

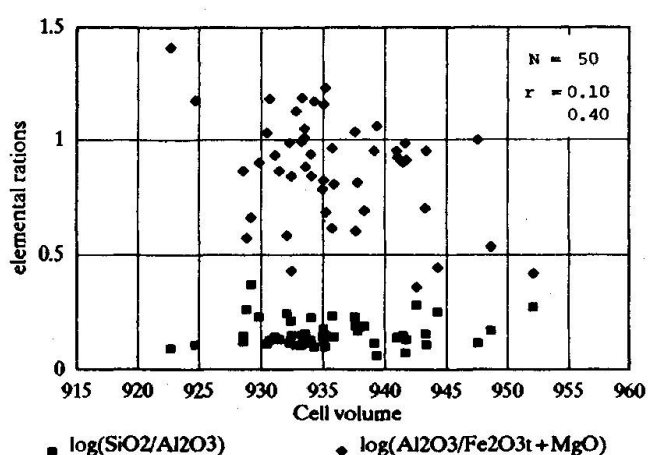
White Mica 2M1 XRD vs XFA



White Mica calculated "Phengite"



White Mica 2M1 XRD vs XFA



Alpine Mica 2M1 XRD vs XFA

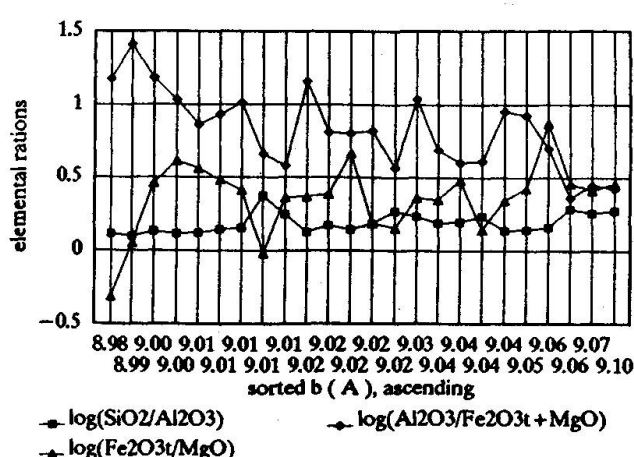


Fig. 5 Cell parameter *b* (XRD, experimental) and "phengite" content (XFA, experimental). Two elemental ratios are indicative, Si/Al connected with phengite content, and Al/Fe + Mg with ferrimuscovite and phengite content. Though a trendlike correlation between *b* and phengite content (XFA) exists, the interdependence is far too weak to be used for analytical purpose (i.e. phengite analysis by XRD).

The question arises (NÄEF, STERN, 1982), whether lacking accuracy of diffractometrical cell determination has been the reason for this poor correlation, or the complexity of chemical substitutions occurring.

The present powder diffraction data on disoriented microsamples display relatively small errors, being obviously not the reason for the large scatter around a regression curve e.g. a versus c or b versus volume; the slope of the regression line, again, corresponds well with the situation expected from literature (BORG and SMITH, 1969).

The correlation found between cell data and relevant chemical data (Fig. 5) resembles much from what is known from the literature and is too weak to be used as a calibration function. Since it can not be explained by errors of measurement, opposite effects of chemical substitutions have to account for it. Certain assumptions have been too simple:

- phengite consists of at least two different species, ferro- and picrophengite; the effect of Fe^{2+} and Mg on a and b parameters probably being different (Fig. 5);
- ferric iron is always present and may influence a and b by either replacing tetrahedral or octahedral Al;
- interlayer Na is scarce in phengitic mica and therefore hardly responsible for large deviations of a and b (Fig. 6);
- the size of c certainly depends on the Na-content of mica but on eventual Ca, Ba, Rb as

well, the latter often being neglected in chemical analysis, as is F whose influence on mica cell parameters is virtually unknown.

An intriguing fact is the obviously poor correlation of the c parameter with any of the chemical variables tested (Fig. 6). The use of basal spacings for paragonite quantification – as is common practice in petrographic literature – can therefore not be recommended. They are, however, helpful when paragonite has to be identified in presence of muscovite.

Acknowledgements

Mica samples have been provided by Dr. H.H. Klein, Schweiz. Isolawerke, Breitenbach; by Mr. Askury Abd. Kadir, Geol. Survey of Malaysia, which is gratefully acknowledged. For critical reading I have to thank Prof. Dr. M. Frey, Mineralogical Institute, Basel University, and Dr. H.H. Klein, and for laboratory support Mrs S. Wüthrich and Mr. R. Handschin, both of Geochemical Laboratorium Mpl Basel.

References

- APPLEMAN, D.E. and EVANS, H.T., Jr. (1973): Indexing and least-squares refinement of powder diffraction data. Report PB 216188, U.S. Dept. Commerce, National Techn. Inf. Service, 5285 Port Royal Rd., Springfield, VA 22151.
- BORG, I.Y. and SMITH, D.K. (1969): Calculated X-ray powder patterns for silicate minerals. Mem. 122, Boulder CO, Geol. Soc. Amer.
- CIPRIANI, C., SASSI, F.P. and VITERBO-BASSANI, C. (1968): La composizione delle miche chiare in rapporto con le costanti reticolari e col grado metamorfico. *Rend. Soc. Ital. Mineral. Petrol.*, 24, 153–187.
- CHATTERJEE, N.D. and JOHANNES, W. (1974): Thermal stability and standard thermodynamic properties of synthetic 2M1 muscovite $\text{KAl}_2(\text{AlSi}_3\text{O}_{10})(\text{OH})_2$. *Contrib. Mineral. Petrol.* 48, 89.
- CHATTERJEE, N.D. (1974): X-ray powder pattern and molar volume of synthetic 2M-paragonite: a refinement. *Contrib. Mineral. Petrol.* 43, 25.
- GUIDOTTI, C.V., SASSI, F.P. and BLENCOE, J.G. (1989): Compositional controls on the a and b cell dimensions of 2M1 muscovite. *Eur. J. Mineral.* 1, 71–84.
- HANDSCHIN, R. and STERN, W.B. (1990): Ein Präparationsverfahren zur röntgendiffraktometrischen und röntgenfluoreszenzspektrometrischen Analyse von Mikroproben. *Siemens Analysentechn. Mitt.* 319.
- NÄEF, U. and STERN, W.B. (1982): Some critical remarks on the analysis of phengite and paragonite components in muscovite by X-ray diffractometry. *Contrib. Mineral. Petrol.* 79, 35.
- SCHWANDER, H., HUNZIKER, J. and STERN, W. (1968): Zur Mineralchemie von Hellglimmern in den Tessiner Alpen. *Schweiz. Mineral. Petrogr. Mitt.* 48, 357.
- STERN, W.B. (1979): Probleme der quantitativen röntgenspektrometrischen Analyse von Hauptkomponenten und Spuren in geologischen Proben. *Schweiz. Mineral. Petrogr. Mitt.* 59, 83.

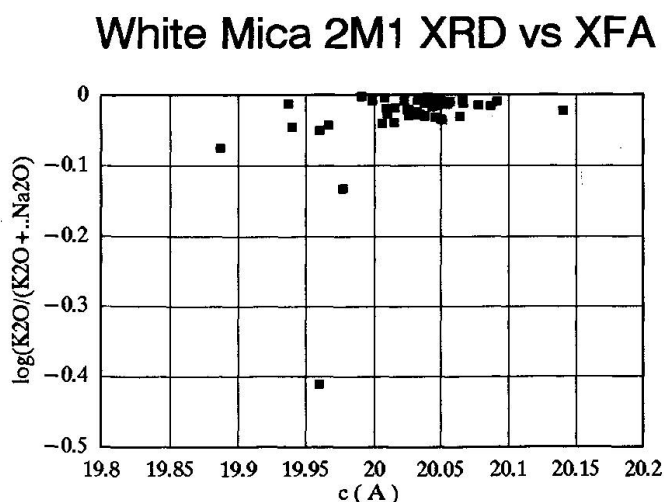


Fig. 6 Cell parameter c and paragonite content (XFA). Compositional variation of the paragonite content is 0 to 25%, main part of points plotting between 0 and 10% paragonite. No correlation between paragonite percentage and cell parameter c is statistically ascertained (Alpine muscovites s.l. $N = 23$, $r = 0.50$; Precambrian, Triassic muscovites s.l. $N = 19$, $r = 0.15$).

STERN, W.B. (1985): Zur Simultananalyse von Silicaten (Hauptkomponenten, Spuren) mittels energiedispersiver Röntgenfluoreszenz-Spektrometrie (EDS-XFA). *Fresenius Z. Anal. Chem.* 320, 6–14.

STERN, W.B. (1987): Determination of mica cell parameters by X-ray powder diffractometry – a case study. *Powder Diffraction*, 2, 249.

Manuscript received August 15, 1990; revised manuscript accepted December 11, 1990.

Appendix: Instrumental conditions

A. X-ray diffraction (XRD)

apparatus	D-500, Siemens GFR
excitation	Cu 40 kV, 30 mA, no primary filter, secondary graphite monochromator
apertures	automatic divergence slit set at 3, 3–3 entrance aperture, 1–0.05–0.15 secondary side
exposure	0.1 2 θ P min., angular increment 0.02 2 θ angular limits 3–73 2 θ
software	DIFFRAC-500 least squares refinement after APPLEMAN, EVANS, 1973, 3 basal reflections pre-indexed, minimum error for rejection 0.05 2 θ ; only reflections > 10 cps were taken for refinement, low scaling factor of 0.1 for reflections below 20 2 θ
specimen	30 mg powdered mica on Makrofol (KG, Bayer) foil 40 mm ϕ electrostatically disoriented (HANDSCHIN, STERN, 1990)

B. X-ray fluorescence (XFA); wavelength-dispersive (WD-XFA)

apparatus	SRS-303, Siemens GFR
excitation	Rh-end window tube, variable according to chemical element; 40 to 60 kV, 70 to 40 mA, 10 to 100 sec
analyzers	In Sb for Si, P, S, Cl multilayer OVO-55 for Al, Mg, Na, F LiF for K- and L-lines of heavy elements fitted background correction for traces
software	SPECTRA/AT (Siemens) routines QUANTXV, QUANTIX data management with LOTUS 1–2–3
specimen	main constituents: fused glass beads consisting of 150 mg ignited sample powder + 2350 mg Li ₂ B ₄ O ₇ annealing in 95 Pt–5 Au crucible, diameter 32 mm, inductive furnace (STERN, 1979); trace elements: 800 mg dried sample powder pressed into Al-rings, diameter 20 mm, elvacite binder

energy-dispersive (ED-XFA)

apparatus	Spectrace-5000, Tracor X-ray, U.S.A.
excitation	W-tube (127 microns Be window), 6 to 50 kV, 0.35 to 0.20 mA, integration time per procedure 200 sec, dead time kept below 40%
analyzer	solid state detector (Li)–Si, with ultrathin Be-window, 7.6 microns
software	Tracor X-ray, running on IBM AT 80-311 314 Mb, 2 Mb RAM
specimen	30 mg on stretched Makrofol foil, as described under section XRD