STRUCTURES, COMPOSITIONS, AND X-RAY DIFFRACTION IDENTIFICATION OF DIOCTAHEDRAL CHLORITES

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Abstract—Al-rich di,trioctahedral chlorite exists as the species cookeite and sudoite. Di,dioctahedral chlorite exists as the species donbassite. Cookeite has essential Li in its structure, sudoite has essential Mg, and donbassite has only small amounts of either element. To date, sudoite has been reported to have only IIb structural units and donbassite to have only Ia structural units. Cookeite is based primarily on Ia structural units, but IIb units are present in specimens from two localities. Most Al-rich chlorite species have regular-stacking "r" or "s" 2-layer stacking sequences, but 1-layer Ia-2 and Ia-6 polytypes also are known. The structural units (Ia or IIb) and the specific stacking sequences can be explained by a combination of local charge balance and minimization of cation-cation repulsion involving the interlayer and tetrahedral chlorite and to identify the type of structural unit present, but single crystal study is necessary to identify the 2-layer and 1-layer sequences with certainty.

Key Words-Chemical composition, Chlorite, Cookeite, Crystal structure, Donbassite, Sudoite, X-ray powder diffraction.

INTRODUCTION

Al-rich chlorite is being reported in the literature with increasing frequency. Three well-crystallized species have been defined to date-cookeite, sudoite, and donbassite. These species can be differentiated from trioctahedral chlorites by their compositions, smaller d(060) values (1.49 to 1.51 Å compared with 1.52 to 1.56 Å for trioctahedral species), slightly smaller d(001) values, and a distinctly more intense 003 reflection near 4.70 Å. The three species are more difficult to distinguish from one another by X-ray diffraction data alone. Cookeite and sudoite are di,trioctahedral chlorites, i.e., the 2:1 layer is dioctahedral, but the interlayer is trioctahedral. Donbassite is di,dioctahedral with two dioctahedral sheets. No examples are known as yet having a trioctahedral 2:1 layer and a dioctahedral interlayer. The present paper summarizes the compositions and several polytypic forms known to occur for these Al-rich species and presents indexed X-ray powder diffraction data to facilitate their identification.

COOKEITE

Cookeite is a di,trioctahedral chlorite commonly found as an alteration product of Li-bearing minerals in pegmatites and hydrothermal veins. The ideal composition is $Al_2(Si_3Al)O_{10}(OH)_2 \cdot (Al_2Li)(OH)_6$. Its essential Li content (~3-4 wt. % Li₂O) distinguishes it from sudoite and donbassite, which contain only minor amounts of Li. Černý (1970), in a study of the most reliable analyses of cookeite, found tetrahedral Si to be nearly constant at 3.0 Si per 4.0 positions, but with tetrahedral Al of some specimens substituted by small amounts of B or Be. Li ranges between 0.8 and 1.4 atoms per formula unit and is concentrated mainly in the interlayer sheet. Fluorine sometimes substitutes for OH in small amounts. Small amounts of Ca + Na + K are often reported in chemical analyses of dioctahedral chlorites as cations residing between the 2:1 layers and the interlayer sheet, but they are more likely to be impurities according to Peacor *et al.* (1988). Formula units of 13 cookeite specimens, modified from Černý (1970), are presented in Table 1. The total octahedral occupancy ranges between 4.8 and 5.3 atoms per 6.0 sites in these formulas, but if the Ca + Na + K totals are excluded as impurities, both the total occupancy and the individual atom values require small revisions.

By far the great majority of cookeite specimens are based on Ia structural units in the terminology of Bailey and Brown (1962). The symbol I indicates that the interlayer octahedral sheet has the same orientation or slant as the octahedral sheet within the 2:1 layer, whereas the symbol II indicates opposite orientations. The symbol a indicates that the interlayer sheet is positioned by long hydrogen bonds so that the interlayer cations project normal to the layers onto tetrahedral cations and the centers of 6-fold rings of the 2:1 layers above and below. The symbol b indicates lack of such superposition. A following integer (1 to 6) then indicates the precise location of the 6-fold rings of the upper 2: 1 layer relative to the symmetry plane of the layer plus interlayer below for regular-stacking polytypes. Several polytypic modifications are known. Vrublevskaja et al. (1975) described an unusual occurrence of cookeite in metamorphosed bauxite from Djalair in Middle Asia, where it formed as a result of interaction of Li-rich solutions with pyrophyllite-2M. The structure is unique also in that it is a regular-stacking 1-layer triclinic struc-

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Sample	Locality	Composition
1	Hebron, Maine	$(Na_{0.03}K_{0.015})(Al_{3.795}Fe^{3+}0.03}Li_{1.44})_{5.265}(Si_{3.045}Al_{0.955})O_{10}(OH)_{7.94}F_{0.06}$
2	Ward's Natural Science Estab- lishment, unknown locality	(Al _{3.825} Li _{1.365}) _{5.19} (Si _{3.165} Al _{0.835})O ₁₀ (OH) ₈
3	Buckfield, Maine	$(Na + K)_{0.075}(Al_{3.825}Fe^{3+}_{0.045}Li_{1.265})_{5.135}(Si_{3.06}Al_{0.94})O_{10}(OH)_{7.985}F_{0.015}$
4	Northwest U.S.S.R.	$(Na_{0.12}K_{0.075})(Al_{3.63}Fe^{3+}_{0.115}Fe^{2+}_{0.14}Mg_{0.05}Li_{1.22})_{5.155}(Si_{2.98}Al_{1.02})O_{10}(OH)_{8}$
5	Dobrá Voda, Czechoslovakia	$(Ca_{0.055}Na_{0.01})(Al_{3.80}Fe^{3+}_{0.015}Mg_{0.14}Li_{1.13})_{5.085}(Si_{3.04}Al_{0.96})O_{10}(OH)_{8}$
6	Muiane, Mozambique	$(Ca_{0.01}Na_{0.01})(Al_{4.025}Li_{1.12})_{5.145}(Si_{2.98}Al_{0.74}Be_{0.22}B_{0.06})O_{10}(OH)_{7.955}F_{0.045}$
7	Kalbinski Range, U.S.S.R.	$(Ca_{0.04}Na_{0.01}K_{0.01})(Al_{3.89}Fe^{2+}0.05}Mg_{0.025}Li_{1.11})_{5.075}(Si_{2.95}Al_{1.05})O_{10}(OH)_{8}$
8	Lipovka, Urals, U.S.S.R.	$(Ca_{0.025}Na_{0.19}K_{0.015})(Al_{3.75}Fe^{3+}_{0.005}Fe^{2+}_{0.015}Mg_{0.175}Li_{0.97})_{4.915}(Si_{3.135}Al_{0.865})O_{10}(OH)_{8}$
9	Manono, Katanga	$(Ca_{0.025}Na_{0.02})(Al_{3.94}Fe^{3+}_{0.005}Mg_{0.025}Li_{0.945})_{4.915}(Si_{3.09}Al_{0.905}B_{0.005})O_{10}(OH)_{8}$
10	North Little Rock, Arkansas	$(Ca_{0.01})(Al_{4.02}Fe^{3+}0.005}Mg_{0.005}Li_{0.86})_{4.89}(Si_{3.035}Al_{0.965})O_{10}(OH)_{8}$
11	Ogofau, Carmarthenshire, En- gland	$(Na_{0.01})(AI_{4.00}Fe^{3+}_{0.02}Fe^{2+}_{0.065}Li_{0.855})_{4.94}(Si_{3.035}AI_{0.965})O_{10}(OH)_{8}$
12	Radkovice, Czechoslovakia	$(Ca_{0.055})(Al_{3.90}Fe^{3+}) = Fe^{2+} = 0.01Mg_{0.11}Li_{0.815})_{4.855}(Si_{3.085}Al_{0.66}B_{0.255})O_{10}(OH)_{8}$
13	Wait-a-bit Creek, British Co- lumbia	$(Ca_{0.16}Na_{0.12}K_{0.01})(AI_{3.88}Mg_{0.11}Li_{0.77})_{4.76}(Si_{2.93}AI_{1.07})O_{10}(OH)_{8}$

Table 1. Structural formulae of cookeite specimens.

Modified from Černý (1970). See that author for references to samples.

ture of the Ia-6 type. The space group is $C\overline{1}$, and the crystallographic angle α is distorted to 90°33'. Most cookeite specimens are less well crystallized and are based on monoclinic 2-layer structures. Lister (1966) pointed out that most of the cookeite specimens observed in her survey of the globular rosettes in Libearing pegmatites appeared to be imperfect 2-layer structures similar to the monoclinic "s" structure derived by Mathieson and Walker (1954) in a study of vermiculite. The symbol of Lister and Bailey (1967) for this 2-layer structure is \bar{X}_1 -Ia-4: \bar{X}_1 -Ia-6. This symbol indicates that the a/3 shift within each 2:1 layer is directed along $-X_1$ and successive 2:1 layers are positioned so that the centers of the 6-fold rings are located alternately at points 4 and 6 on opposite sides of the symmetry plane below (related to one another by a b/3 shift). The 2-layer nature is only apparent by observation of weak diffuse reflections of index $k \neq 3n$ on single crystal patterns. These reflections do not show up well on Debye-Scherrer patterns (Table 2a).

Lister (1966) also identified a better crystallized 2-layer cookeite from Wait-a-bit Creek, British Columbia, and at several localities in the Magnet Cove district and around North Little Rock, Arkansas. The latter includes cookeite from the Jeffrey quarry, which is part of the Source Clays Repository of The Clay Minerals Society. The structure can be correlated with the monoclinic "r" and "q" structures of Mathieson and Walker (1954), which have polytypic symbols "r" = \bar{X}_1 -Ia-4: \bar{X}_3 -Ia-4 and "q" = \bar{X}_1 -Ia-6: \bar{X}_3 -Ia-2. Here the a/3shift is directed along $-X_1$ of the first 2:1 layer and along $-X_3$ of the second layer. Six-fold rings of successive 2:1 layers are positioned at points 4 and 4 for the "r" structure and at points 6 and 2 for the "q" structure. The two structures are enantiomorphic in space group Cc. Bailey (1975) summarized results of an incomplete structural refinement of one of the North Little Rock crystals. In this structure Li was found to

be ordered in one interlayer site, where it apparently achieves the best local charge balance. Interlayer Li+ lies on a line directly between a tetrahedral cation in the 2:1 layer below and a tetrahedral cation in the 2:1 layer above. The more highly charged interlayer Al³⁺ cations, however, only have a tetrahedral cation on one side and the center of a 6-fold ring on the other side. Although different T-O bond lengths were determined during structural refinement, the poor quality of the film data does not permit a firm conclusion as to ordering of tetrahedral Si and Al, nor has it been possible to obtain a better crystal to date. The powder pattern in Table 2b shows clearly $k \neq 3n$ reflections that identify the 2-layer nature of the specimen, but the more intense k = 3n lines are very similar to those of the more common but less well crystallized cookeite specimens.

Cerný et al. (1971) described a cookeite from a pegmatite at Dobrá Voda, Czechoslovakia, as having IIb structural units. The present writers have confirmed a second occurrence of cookeite-IIb as pink rosettes on elbaite from a pegmatite at Norway, Maine (University of Wisconsin Museum #6001/1). The powder pattern (Table 2c) is distinctly different from those of cookeite specimens based on Ia units. Single crystals were not available, but a comparison of observed and calculated intensities of $k \neq 3n$ reflections suggests 2-layer "r" or "q" structures.

Frank-Kamenetskii (1960) and Černý (1970) mentioned manandonite from Madagascar as a possible di,trioctahedral Al-Li chlorite having substantial tetrahedral boron ($B^{IV} \ge Al^{IV}$). Guggenheim *et al.* (1983) confirmed the high B^{IV} content of this material by ion probe analysis and stated that its X-ray powder pattern was that of a I*a* type chlorite. The type manandonite defined by Lacroix (1912, 1922), however, has since been shown to be a trioctahedral 1:1 layer silicate of approximate composition (Al₂Li)(Si₁₀Al_{0.5}B_{0.5})O₅(OH)₄ (A.-M. Fransolet, University Liége, Liége, Belgium, personal communication, 1988; Ranorosoa, 1986). This 1:1 type material is intermixed with a B-rich Al-Li "cookeite", which Strunz (1957) and later workers showed to be a chlorite and mistakenly called manandonite. This chloritic material will probably qualify as a new species when the details of its composition and structure are better known.

SUDOITE

Sudoite is also a di,trioctahedral chlorite, and has been identified in hydrothermal alteration zones, sediments, soils, fissure veins, and low-grade metamorphic rocks from many localities. The ideal composition is $Al_2(Si_3Al)O_{10}(OH)_2 \cdot (Mg_2Al)(OH)_6$. Tetrahedral Al is known to range from about 0.4 to 1.1 atoms per formula unit, octahedral Al from 2.5 to 3.4, and Mg from 1.2 to 2.5. Smaller amounts of Fe, Mn, and Li are also present. Total octahedral occupancy ranges from 4.5 to 5.1 atoms per 6.0 sites. Sudoite contains only a minor amount of Li and less Al and considerably more Mg than cookeite. Representative structural formulae for sudoite are listed in Table 3.

Stanton (1984) reported two groups of iron-rich "sudoites" at the Geco mine, Manitouwadge, Ontario, with $Fe^{2+} > Mg$ for one group. The materials are impure, however, and the average formulae of the two groups differ from those in Table 3 in several respects. Tetrahedral Al is low (0.1 to 0.3 atoms as recalculated by the present authors on the basis of 28 positive charges), the Ca + Na + K totals are moderately high (0.27 to 0.29 atoms), total octahedral occupancy is low (4.3 to 4.4 atoms per 6.0 sites, excluding Ca + Na + K), and the total octahedral charge is negative (rather than positive) relative to an ideal complement of anions. Purer material is needed to verify the nature of this mineral.

Eggleton and Bailey (1967) and Shirozu and Higashi (1976) attempted partial refinements of the structures of different fine-grained sudoite specimens on the basis of X-ray powder diffraction data. Their results confirmed that these specimens are based on IIb structural units in which the 2:1 layer is dioctahedral and the interlayer is trioctahedral. Drits and Lazarenko (1967) and Lin and Bailey (1985) used single crystals to show that specimens from the Ural Mountains and Belgium, respectively, have 2-layer structures of the "s" type. The powder pattern in Table 4 is indexed best by incorporation of a slight distortion to triclinic geometry. Di,dioctahedral and di,trioctahedral chlorites have d(001) values that are systematically smaller than those for trioctahedral specimens of similar tetrahedral compositions, thus invalidating the usage of existing d(001)graphs and equations for estimating tetrahedral Si,Al contents of such specimens. Lin and Bailey (1985) concluded from 1-dimensional electron density projections of sudoite that the smaller d(001) value is due primarily to a thinner dioctahedral sheet within each 2:1 layer.

DONBASSITE

Donbassite is defined as having two dioctahedral sheets and an ideal composition near Al₂(Si₃Al)O₁₀- $(OH)_2 \cdot Al_{2,13}(OH)_6$. For a variable amount (x) of tetrahedral substitution the formula can be written as $Al_2(Si_{4-x}Al_x)O_{10}(OH)_2 \cdot Al_{2+x}(OH)_6$. It has been reported from soils, sediments, hydrothermal alteration zones, and fissure veins. Substitution of Al for Si in the tetrahedral sheets of the best crystallized specimens ranges from 0.6 to 1.3 atoms per 4.0 positions. The octahedral cation total is always greater than 4.0 (excluding Ca + Na + K), ranging from about 4.2 to 4.5 atoms and indicating that the positive charge on the interlayer sheet arises primarily from the presence of Al + Mg + Fe cations in excess of 2.0 in that sheet, i.e., partly filling the "vacant" octahedron. Small amounts of octahedral Li are commonly present (Rozinova and Dubik, 1983). Representative structural formulae are listed in Table 5.

Whereas sudoite has only been identified to date as having IIb structural units, donbassite has only been identified as having Ia structural units. This statement takes into account the evolution of the definition of these names over time, i.e., some sudoite specimens as presently defined have been called donbassite in the past and vice-versa. Drits and Lazarenko (1967) identified a donbassite specimen from the Donbass region, U.S.S.R., as having a 2-layer structure like those of the equivalent "r" and "q" structures and consisting of Ia structural units. Aleksandrova et al. (1972) refined the crystal structure of a low-Li donbassite from Novaya Zemlya, U.S.S.R. They reported a regular-stacking 1-layer structure of space group C2, which is a distorted version of the Ia-2 polytype of ideal symmetry C2/m. The vacant octahedral site in the 2:1 layer lies on the symmetry plane of the layer, but the partly vacant octahedral site in the interlayer sheet lies off the projection of the mirror plane (thereby eliminating the mirror for the crystal as a whole). Adjacent layers adopt the mica configuration. Different mean T-O bond lengths observed in this refinement using film-collected data suggest ordering of tetrahedral Si and Al, but a more accurate refinement using counter-collected data is desired for proof of this point.

No accurate powder data for purified donbassite specimens are available for comparison with those of sudoite and cookeite. Observed d(060) values are between 1.49 and 1.50 Å for donbassite and between 1.49 and 1.51 Å for sudoite and cookeite. Table 6 lists a calculated XRD pattern for donbassite based on the atomic coordinates reported by Aleksandrova *et al.* (1972) for the Ia-2 structure. The strong k = 3n reflections are similar in positions and intensities to those

hki	1		d(obs.) (Å)	d(calc.) (Å)	hki	I	d(obs.) (Å)	d(calc.) (Å)
(a) Cookei	ite-Ia(s)				13,12	7	1.680	1.681
002	75		14.10	14.056	13,14	20	1.636	1.634
004	60		7.05	7.028	20,12	10	1.627	1.629
006	100		4.68	4.685	20,16	5	1.545	1.542
020	1 20			4.468	13,14	10	1.538	1.237
110	} 30		4.44	4.444	332 060 331	50	1.490	1.490
112	{		}	4.412	336		}	1.459
113	} 5		4.12	4.197	064 332	8	1.458 {	1.457
008	60		3 513	3 514	338	8	1.421	1.422
00.10	20		2.812	2.811	066	-		1.419
200.132	30		2.560	2.561	334		{	1.418
204	1 50		2505	2.510	$20,\overline{18}$			1.415
132	} 30		2.505	2.505	13,16 J	8	1.409	1.410
134			2 47	2.469	$13.\overline{18}:33.\overline{10}$)	6	1.375	1.373 Å
202	{ ³		2.77	2.465	068		1	1.371
206	} 1		2.37	2.377	20,16;336		1.372	1.370
134	{ ·			2.370	33,12		(1.319
130	> 90		2.317	2.319	06,10	5	1.314 {	1.316
204	{		<pre>></pre>	2.314	338		ļ	1.314
136	} 5		2.202	2.200	20,20	25	1.300	1.303
138	{		}	2.190	13,18 J		1	1.299
206	2		2.140	2.137	Engeimon from	Waisanan	Tomminan nit Gr	hoowna
20.10	{ _		<pre>></pre>	2.024	Maine has a	n imperfect	"s" structure Cell di	nensions.
138	} >		2.020	2.017	a = 5.163 h	= 8.935 c	= 78.34 Å $B = 97.7$	• Pattern
00,14	10		2.010	2.008	of crystal m	ounted in a	Gandolfi camera, 1	14.6-mm
13, 10	25		1.962	1.962	diameter, gr	aphite-mon	ochromatized FeK α	radiation.
208	12		1.953	1.956	Indexing by	comparison	with single crystal da	ta, inten-
00, 16	5		1.760	1.759	sities estima	ted visually.		
20,14	3		1.689	1.68/			<u></u>	·
(b) Cooke	ite-Ia(r)							
002	90		14.06	14.077	138	20	2.109	2.021
004	35		7.03	7.039	20,10		}	2.016
111) ⁶⁰		4.70	4.692	$\frac{208}{12}$	45	B 1.960	1.904
110	20		4.47	4.439	13,10		}	1.938
112	{		}	4 366	$\frac{20,10}{13,12}$ }	1	1.790	1.792
111	} 10		4.35 {	4 321	00.16	5	1 760	1.760
113	{		(4.17	4.184	13.12		n 1.686	1.685
112	> 10	В	{	4.118	20,14	10	B { 1.680	1.681
023	J		4.03	4.031	20,12	25	P 1.638	1.637
113		в	3.8 7	3.866	13,14	33	^B 1.630	1.632
024	f ³	D	3.77	3.770	13,14	20	1.542	1.541
115	1		3.66	3.674	$20,\overline{16}$	15	1.538	1.537
008	60		3.518	3.519	332			1.489
116	3		3.410	3.404	060	75	1.487	1.488
00.10	3		3.060	3.073	331			1.487
200	45		2.815	2.815	332	1.5	1 450	1.45/
132	60		2.560	2.361	064	15	1.456	1.455
132	{		}	2.338	330			1 4 20
$20\overline{4}$	80		2.505	2.503	224	15	1 4 1 9	1 418
202	{		}	2.469	066	15	1.417	1.418
134	3		2.465 {	2.464	13.16			1.414
134	1		2 2 70 }	2.372	20.18	10	1.412	1.411
206	$\int 2$		2.370	2.368	20.16	-	1 074	1.377
204	100		2 216	2.321	$13,\overline{18}$ }	7	(^{1.374}	1.373
136	۰ ^۲ (2.510	2.314	336	}	в {	1.373
02,11	1	1	2.220	2.220	068	7)	[1.371	1.370
136	-	} В		2.202	33,10	-		1.369
208	5	J	1,200	2.197	Specimen from	North Little	e Rock, Arkansas, has	a 2-layer
138	> 3		2.140	2.144	"r" structure	with Ia laye	rs. Cell dimensions: a	= 5.158,
02.12	ر ر ۱		2 070	2.138 2.077	b = 8.927, c	= 28.351 Å	$\beta = 96.8^{\circ}$. Condition	ons as for
· · · · · · · · · · · · · · · · · · ·	1		2.070	2.077	cookeite-Ia(s).		

Table 2. X-ray powder diffraction patterns of cookeite specimens.

hkl	I		d(obs.) (Å)	d(calc.) (Å)	hki	I		d(obs.) (Å)	d(calc.) (Å)
(c) Cookeite-IIb									
002	80		14.10	14.100	204,136	20		2.319	2.319
004	60		7.05	7.050	208,136	20		2.203	2.203
006	80		4.70	4.700	206,13;8	1		2.140	2.143
111)	65		A 45	∫ 4.463	20,10;138	15		2.021	2.022
110 🖌	65		4.45	4.445	208;13,10	75		1.962	1.962
112	5		4 25	∫ 4.373	208;12;13,10	12		1.848	1.847
111 ∫	3		4.33	4.321	20,10;13,12	40		1.792	1.791
112	5		4.13	4.117	00,16	5		1.762	1.762
024	5	р	2 72	∫ 3.775	20,14;13,12	15		1.686	1.686
115	5	D	5.75	3.684	20,12;13,14	3		1.638	1.636
008	40		3.52	3.525	20,16;13,14	55		1.542	1.542
026	1		3.22	3.239	060,332	90		1.490	1.490
118	1		2.92	2.915	064,332,33 6	15		1.458	1.458
00,10	30		2.820	2.820	066,334,33 8	20		1.420	1.420
119	5		2.705	2.700	20,16;13,18	40	р	1 275	∫ 1.376
118	1		2.62	2.630	068;336;33,10	40	Б	1.575	1.372
200,132	25	В	2.560	2.561	06,10;338;33,12	8		1.319	1.317
204,132	15		2.505	2.507	Specimen from	Norway M	Maine	Cell dimer	sions $a =$
202,134	100		2.468	2.468	$5161 \ b = 80$	38 c = 2	08 41 Å	$\beta = 97^{\circ}$	Conditions
206,134	80		2.373	2.373	as for cookeite-	Ia(s).	ю.ті <i>П</i>	, p 97.	Conditions

Table 2. Continued.

Table 3. Structural formulae of sudoite specimens.

Speci- men	Locality	Composition	Reference
1	Tracy mine, Michi- gan	$(Al_{2.7}Mg_{2.3})_{5.0}(Si_{3.3}Al_{0.7})O_{10}(OH)_8$	Bailey and Tyler (1960)
2	Colorado Plateau	$(R_{5,1})(Si_{3,4}Al_{0,6})O_{10}(OH)_8$	Schultz (1963)
3	Honko deposit, Ka- mikita mine, Ja- pan	$\begin{array}{l}(Ca_{0.11})(Al_{3.017}Fe^{3+}_{0.345}Fe^{2+}_{0.003}Mn_{0.004}Mg_{1.175})_{4.544}(Si_{3.261}Al_{0.739})-\\O_{10}(OH)_{8}\end{array}$	Hayashi and Oinuma (1964); Sudo and Sato (1966)
4	Furutobe mine, Ja- pan	$(Ca_{0.060}Na_{0.17}K_{0.159})(Al_{3.042}Fe^{3+}_{0.037}Mg_{1.594})_{4.673}(Si_{3.242}Al_{0.758})O_{10}(OH)_{8}$	Tsukahara (1964)
5	Kaiserbach Valley, Germany	$(Ca_{0.09})(Al_{2.70}Fe^{3+}{}_{0.09}Fe^{2+}{}_{0.04}Mg_{1.97})_{4.80}(Si_{3.43}Al_{0.57})O_{10}(OH)_8$	Frenzel and Schembra (1965)
6	Beriozovsk, Urals, U.S.S.R.	$(Ca_{0.03})(Al_{2.9}Fe_{0.22}Mg_{1.96})_{5.08}(Si_{2.86}Al_{1.14})O_{10}(OH)_{8}$	Drits and Lazarenko (1967)
7	Kazan Mineral Mu- seum, unknown location	$(Ca_{0.14})(Al_{3.04}Fe^{2+}{}_{0.18}Mg_{1.61})_{4.83}(Si_{3.02}Al_{0.98})O_{10}(OH)_{8}$	Vlasov and Drits (1967)
8	Niida mine, Japan	$(Na_{0.03}K_{0.03})(Al_{3.03}Fe^{3+}{}_{0.06}Mg_{1.78})_{4,87}(Si_{3.01}Al_{0.99})O_{10}(OH)_8$	Kimbara and Nagata (1974)
9	Uchinotai deposit, Kosaka mine, Ja- pan	$\begin{array}{l}(Ca_{_{0.06}}Na_{_{0.03}}K_{_{0.05}})\\(Al_{_{3,13}}Fe^{_{3+}}{}_{_{0.01}}Fe^{_{2+}}{}_{_{0.01}}Mg_{_{1.60}})_{_{4.75}}(Si_{_{3,19}}Al_{_{0.81}})O_{_{10}}(OH)_{_{8}}\end{array}$	Tsuzuki and Honda (1977)
10	Okunosawa deposit, Kamikita mine, Japan	$(Ca_{0.01}Na_{0.11}K_{0.01})(Al_{3.19}Fe_{0.03}Mg_{1.37})_{4.59}(Si_{3.08}Al_{0.92})O_{10}(OH)_{8}$	Honda (1975)
11	Hanaoka mine, Ja- pan	$(Al_{3,2}Mg_{1.6})_{4.8}(Si_{3,2}Al_{0.8})O_{10}(OH)_{8}$	Shirozu and Higashi (1976)
12	Rydnyy Altai, U.S.S.R.	$(Ca_{0.17}Na_{0.02}K_{0.06}(Al_{2.45}Fe^{3+}{}_{0.03}Fe^{2+}{}_{0.01}Mg_{2.54})_{5.03}(Si_{3.06}Al_{0.94})O_{10}(OH)_8$	Alysheva et al. (1977)
13	Venn-Stavelot Mas- sif, Ardennes, Bel- gium	(1) $(Al_{2.84}Fe^{3+}_{0.19}Fe^{2+}_{0.02}Mn_{0.02}Mg_{1.91})_{4.98}(Si_{3.01}Al_{0.99})O_{10}(OH)_{8}$ (2) $(Al_{2.86}Fe^{3+}_{0.17}Mn_{0.02}Mg_{1.94})_{4.99}(Si_{2.99}Al_{1.01})O_{10}(OH)_{8}$	Fransolet and Bour- guignon (1978)
14	Harz Mountains, German Demo- cratic Republic	(1) $(Al_{2.864}Fe^{3+}_{0.136}Fe^{2+}_{0.136}Mn_{0.032}Mg_{1.886})_{5.054}(Si_{3.028}Al_{0.972})O_{10}(OH)_{8}$ (2) $(Al_{2.871}Fe^{3+}_{0.129}Fe^{2+}_{0.136}Mn_{0.035}Mg_{1.875})_{5.046}(Si_{3.037}Al_{0.963})O_{10}(OH)_{8}$	Kramm (1980)

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	Table 4.	Powder pattern of	of sudoite-IIb, Ottre,	Belgium.
d(obs.)		d(calc.)		d(obs.)

hkl	1	d(obs.) (Å)				d(calc.) (Å)		hkl		1	d(obs.) (Å)				d(calc.) (Å)
002	40	14.2				14.16	•	13,10)					ſ	1.871
004	80	7.1				7.08		$2,0,\overline{12}$	}	5	1.870		В	{	1.871
006	100	4.72				4.721		1,3,10	}					ł	1.858
020)				(4.547		1,312	-	12	1.820				1.820
021	} 4	4.50				4.496		20,10	1	0	1 802			ſ	1.805
021	1				1	4.483		1,3,12	Ĩ	0	1.803			٦.	1.804
111	1				1	4.387		0,0,16	,	1	1.770			``	1.770
111	1 10	4.25			•	4.384		312)					ſ	1.717
$0\overline{2}2$	10	4.35			í	4.340		311			1.711)		- {	1.713
022						4.318		0.4.11	Į	10			n		1,712
114	1	4.01			}	4.024		1.3.12	1	10		1	в	1	1.706
114	} 10	4.01			1	4.004		153			1.705			ł	1.705
113	<u>۲</u>				1	3.915		314				,			1.705
113	} >	3.87			1	3.905		153	{	-		1		一 入	1.675
$0\overline{2}5$	{				}	3.561		245	~ ~	5	1.673			1	1.674
008	\$ 40	3.535			Į	3.540		154	1			ţ	в	- 2	1.650
025						3.531		246				1	-	i	1 650
115	{ .					3.361		2.0.12	}	1	1.650			í	1.647
115	} 1	3.36			1	3,350		13 14				,		1	1 646
076	{				}	3 789		245	{						1 600
026	2	3.28			{	3 260		$\frac{213}{157}$	}	1	1.600			1	1.600
116	{				>	3 102		1 3 14	{					- >	1 559
116	2	3.10				3 002		2016	}	15	1.560			- {	1.559
0.0.10	25	2 8 3 5			(2 0 2 2		1314	,	25	1 5/0			(1.539
202		2.055			(2.552		337	1	25	1,547			(1.545
130	10	2 615			J	2.019		060	l	50	1 515			J	1.516
132	$\int 10$	2.015)	2.010		337	ſ	50	1.515				1.510
204	{				- }	2.510		332	{					- }-	1 480
137	20	2 545			J	2.550		332							1.480
132	²⁰	2.545)	2.349		330	}	20	1.481			{	1.480
202	{				}	2.544		064							1.430
134	} 90	2.502				2.503		220	 					}	1.47
206	{				>	2.302		066							1.447
134		2 408			J	2.412		334							1.441
134	{ 35	2.400				2.412		220	}	10	1.422		В	- {	1.441
136	{				}	2.402		330							1.441
0.012	(15	2 3 5 8))	2.304		066	1						1 4 3 0
1111	$\int $	2.550)	2.300		000)	7	1 4 1 0			(1.439
204	{		<pre>}</pre>	В	- F	2.337		0,0,20	1	,	1.417			(1.410
136	L 10	2 345			J	2.340		3 3 10	}	20	1.398			{	1 208
1 1 1 1	$\int 10$	2.345)		Ì	2.347		3,310						}	1.390
1,1,11	{				ł	2.340		330	l	15	1 201			J	1.309
208	20	2.237				2.270		1512	ſ	15	1.364				1.309
208	{				<pre>}</pre>	2.209		1,5,15)					U	1.303
1 200	> 5	1.165			- {	2.163				-					
130	{				}	2.104									
2010	} 1	2.050			{	2.031									
2,0,10	J 10	2 0 2 0			L.	2.030		Specimen	ı fron	1 Ottro	e, Belgium.	a=5	5.247	, <i>b</i> =	9.094, <i>c</i> =
1210	10	2.020				2.023		28.557	Å, α	= 90.3	5°, $\beta = 97.3^{\circ}$	°, and	$1\gamma =$	89.9	^a . Structure
13,10	20	1.993			(1.990		is of th	ne s-ty	pe wi	th IIb layer	s. Co	nditi	ons a	as listed in
1,3,10	} 18	1.978			{	1.980		Table 2	2. B =	= broa	d.				

of cookeite-Ia. The strong 021 reflection at 4.269 Å is diagnostic of the Ia-2 polytype.

EVALUATION OF STRUCTURES

The dominance of the Ia structure in cookeite and donbassite and of the IIb structure in sudoite and trioctahedral chlorites is linked directly to the magnitude and location of the positive electrostatic charge in the interlayer sheet. Bailey and Brown (1962) and Shirozu and Bailey (1965) showed that the IIb structure is by far the most stable structural form of trioctahedral chlorite as a result of the complete absence of any cation-cation repulsion between sheets of the layered assemblage. The Ia and IIa structures, on the contrary, have exact vertical superposition of interlayer and adjacent tetrahedral cations with large resultant cationcation repulsion values. Nevertheless, the Ia structure would have a very high structural stability rating if the

Speci- men	Locality	Composition	Reference
1	Donbass region, U.S.S.R.	(1) $(Ca_{0.17}Na_{0.19})(AI_{3.92}Fe^{3+}_{0.05}Mg_{0.21}Li_{0.05})_{4.23}(Si_{3.09}AI_{0.91})O_{10}(OH)_{8}$ (2) $(Ca_{0.19}Na_{0.22})(AI_{3.88}Fe^{3+}_{0.08}Mg_{0.23})_{4.18}(Si_{3.06}AI_{0.94})O_{10}(OH)_{8}$ (3) $(AI_{4.14}Fe^{3+}_{0.07}Mg_{0.11})_{4.24}(Si_{1.10}AI_{0.92})O_{10}(OH)_{8}$	Lazarenko (1940)
2	Alberni soil, British Columbia	$(Ca_{0.02}K_{0.23}X_{0.24})$ $(A1_{22}Fe^{3+}_{0.03}R_{10,09}Mg_{0.76})_{4,45}(S1_{2,80}A1_{1,20})O_{10}(OH)_{8}$	Brydon et al. (1961)
3	Saint-Paul-de-Fe- nouillet, East Pyre- nees	$(Al_{4.3}Fe^{3+}_{0.03}Fe^{2+}_{0.02})_{4.35}(Si_{2.70}Al_{1.30})O_{10}(OH)_{8}$	Caillère et al. (1962)
4	Kesselberg, Federal Republic of Ger- many	(1) $(Al_{4.17}Mg_{0.10})_{4.27}(Si_{3.30}Al_{0.70})O_{10}(OH)_8$ (2) $(Al_{4.27})(Si_{3.2}Al_{0.8})O_{10}(OH)_8$	Müller (1961, 1963)
5	Nagol Tarasovki, Donbass, U.S.S.R.	$(Ca_{0.2})(Al_{4.0}Mg_{0.24})_{4.24}(Si_{3.12}Al_{0.88})O_{10}(OH)_{8}$	Drits and Lazarenko (1967)
6	Namivu, Alto Ligon- ho, Mozambique	$(Ca_{0.04})(Al_{3.96}Fe^{3+}_{0.02}Fe^{2+}_{0.016}Mn_{0.008}Mg_{0.04}Li_{0.27})_{4.314}$ $(Si_{3.45}Al_{0.55})O_{10}(OH)_8$	Gomes (1967)
7	Novaya Zemlya, U.S.S.R.	$(Al_{4.10}Fe^{3+}_{0.04}Fe^{2+}_{0.01}Mg_{0.08}Li_{0.26})_{4.49}(Si_{3.14}Al_{0.86})O_{10}(OH)_{8}$	Aleksandrova et al. (1972)
8	Donbass area, U.S.S.R.	$(Al_{3.96}Fe^{2+}{}_{0.08}Mg_{0.14}Li_{0.20})_{4.38}(Si_{3.36}Al_{0.64})O_{10}(OH)_{8}$	Rozinova and Dubik (1983)
9	Szabo Bluff, Scott Glacier, Antarctica	$(Al_{4.32}Fe^{3+}_{0.07})_{4.39}(Si_{2.83}Al_{1.17})O_{10}(OH)_{8}$	Ahn and Buseck (1988)

Table 5. Structural formulae for donbassite specimens.

cation-cation repulsion factor could be minimized. This is exactly what happens in vermiculite, which contains only a few exchangeable interlayer cations and for which the structural type has been shown to be Ia without exception. This favorable rating of the Ia structure carries over to dioctahedral chlorite species in which one of the three interlayer cation sites is partly vacant (donbassite) or has a low-charge Li⁺ cation present (cooke-ite).

In the regular-stacking 1-layer chlorite polytypes involving Ia and IIa layers, two different dispositions of the 2:1 layers exist on opposing sides of the interlayer sheet. The first, termed type A here, is found only in the monoclinic structures Ia-2 and IIa-1, in which successive 2:1 layers adopt the mica configuration with their 6-fold rings opposed. One interlayer cation site thus lies between the centers of these opposed fold rings, and the other two interlayer sites lie on vertical straight lines between tetrahedral cations of the two sets of rings. One form of donbassite adopts the Ia-2 structure. The structural refinement by Aleksandrova et al. (1972) places the partly vacant interlayer site M between one set of superimposed tetrahedral cations and an Al³⁺ cation between the other set of two tetrahedra. The second Al³⁺ interlayer cation lies between the centers of the 6-fold rings in vertical projection. This distribution of interlayer cations gives the minimum amount of cation-cation repulsion possible for the Ia-2 structure. The authors stated that repulsion is minimized further by tetrahedral cation ordering and local charge balance such that the source of positive interlayer charge is located between the sources of the tetrahedral negative charges, i.e., between Al-rich tetrahedra above and below. In this arrangement, the

source of positive interlayer charge is the partly vacant M octahedron. Thus, that cation-cation repulsion would be minimized even further by the opposite pattern of tetrahedral cation ordering in which the partly filled and low-charge M site would be between Si^{4+} -rich tetrahedra and the high-charge interlayer Al^{3+} would lie between Al-rich tetrahedra. Verification of the ordering pattern is desirable.

The other type A structure having a mica configuration of layers is IIa-1, which is not known in true chlorites, but is present in franklinfurnaceite, a hybrid chlorite-mica in which the chlorite is of the tri,dioctahedral type. In this mineral there is excellent local charge balance, with interlayer R^{3+} cations positioned midway between tetrahedral R^{2+} , interlayer R^{2+} between tetrahedral R^{4+} , and interlayer vacancies between the extra Ca that lie in the 6-fold ring openings (Peacor *et al.*, 1988).

A second disposition of 2:1 layers, termed here type B, exists in the 1-layer triclinic polytypes Ia-4 and IIa-3. Here, the unique interlayer site lies on a center of symmetry between tetrahedral sites above and below, and the other two interlayer sites are asymmetric, with a tetrahedral cation on one side and the center of a 6-fold ring on the other side. Cookeite in metamorphosed bauxite from Djalair, Middle Asia, adopts the Ia-6 = Ia-4 1-layer structure (Vrublevskaja *et al.*, 1975). Although the cation ordering scheme is not known for that specimen, logically the low-charge Li+ should exist on the center of symmetry between tetrahedral sites above and below and the two high-charge Al3+ cations should be in the asymmetric locations with a tetrahedral cation on only one side. Cookeite and donbassite also adopt either or both of the 2-layer Ia "s" and

hkl	I	d(calc.) (Å)	hkl	I	d(calc.) (Å)
001	60	14.127	207) (1.702
002	65	7.063	311		1.693
003	100	4.709	151	$\int 21 f$	1.688
020	54	4.478	136		1.683
111	35	4.395	151	1	1.670
021	49	4.269	152	<pre>> 11 </pre>	1.661
111	14	4.106	241	JU	1.657
112	93	3.983	137) (1.645
022	16	3.782	243	} 42 {	1.635
112	25	3.578	206	J	1.627
004	60	3.532			
113	11	3.444	208	16	1.557
113	28	3.058	137	$\int \int \int \int \int \partial f dx$	1.539
114	4	2.941	331	1 48 1	1.493
005	28	2.825	060	{ ^{**} }	1.493
024	5	2.773	062	} 6 {	1.460
114	, 4	2.619	331	{ `}	1.457
131	\$ 47	2.569	063] [1.423
200	{	} 2.563	332		1.418
202	52	2.522	227	2^{21}	1.414
131	{	2.508	138		1.413
132	\$ 4	2.479	00,10	1	1.413
201	{	2.463	335	$\frac{1}{7}$	1.382
203		2.392	064	$\{ \}$	1.375
025	} /	{ 2.390	20,10	31	1.315
132	ł	2.3/1	139) - (1.302
133	9 7	2.331			
202	{	2.310	Specim	en from	Novaya
204	> 10	2.223	Zemly	ya, U.S.S	.R., is a
133	{	2.200	Ia-2	polytype	. Pattern
134		2.155	calcul	ated from	n atomic
042	4	$\begin{cases} 2.134 \\ 2.122 \end{cases}$	coord	inates of	Aleksan-
203	{	$\left\{ \begin{array}{c} 2.132 \\ 2.042 \end{array} \right\}$	drova	<i>et al.</i> (19	772). Cell
203	6 20	$\int \frac{2.042}{2.018}$	dimer	isions: $a =$	≈ 5. 174, b
134	$\int 20$	2.018	= 8.9	56, c = 16	4.26 A, β
135	{	2.010	= 97.	85°. Space	e group =
204	} 49	1 052	C2.		
204	<u> </u>	(1.932			

Table 6. Calculated powder pattern of Ia-2 donbassite, Novaya Zemlya, U.S.S.R.

"r" structures (terminology of Mathieson and Walker, 1954) that have the same interlayer-layer type B configurations. The "r" structure has been reported for donbassite by Drits and Lazarenko (1967), although the cation-ordering pattern was not investigated. Both structures are known for cookeite. Our refinement of the cookeite "r" structure places the low-charge interlayer Li⁺ between tetrahedral cations on both sides and the two high-charge interlayer Al3+ cations in the asymmetric locations with a tetrahedral cation on only one side. The "s" structure has not been refined, but a similar local charge balance arrangement can be anticipated. The Ia "s" structure is also the most stable form of vermiculite (Shirozu and Bailey, 1966), in which the exchangeable cation lies vertically between tetrahedral cations above and below, with vacancies in the two asymmetric locations. At one time, the tetrahedral cation ordering was thought to place Al-rich tetrahedra

above and below the exchangeable cation to give optimum local charge balance (Shirozu and Bailey, 1966), but later work (de la Calle *et al.*, 1976) ruled out the existence of long-range tetrahedral cation ordering in vermiculite. Some evidence exists for short-range ordering in local domains (Thompson, 1984).

In contrast to the preceding species, sudoite has a trioctahedral interlayer sheet of composition (Mg_2AI), in which all of the cations have medium to high charges. And, just as in trioctahedral chlorites, the IIb structure is favored to avoid cation-cation repulsion present in an *a* position of the interlayer sheet. The presence of a vacancy in the dioctahedral 2:1 layer has little or no effect on the position of the interlayer.

It is not prudent to assume that all sudoite is of the IIb structural type and that all donbassite is of the Ia structural type, because study of more samples may show structural variations in each, just as it has in cookeite. Most cookeite and most donbassite specimens, however, are based on Ia structural units and most sudoite specimens are based on IIb units. These structural units can be identified by X-ray powder diffraction patterns. The Ia structural unit in a 2-layer dioctahedral chlorite is characterized by a strong d(204) XRD line near 2.32 Å. The IIb structural unit for a 2-layer dioctahedral chlorite is characterized by a strong d(202) line near 2.47–2.50 Å and a strong d($20\overline{6}$) line near 2.37-2.41 Å. Most dioctahedral chlorite specimens appear to be based on regular-stacking 2-layer structures of the "s" or "r" and "q" types, but regular 1-layer forms are known as well. Single crystal study is usually necessary to identify the specific stacking arrangement present.

Environment of occurrence also can be helpful in identification. The presence of essential Li in cookeite usually restricts its occurrence to Li-rich environments, rather than in sediments and soils. Essential Mg in sudoite is not very restrictive and leads to its ocurrence in many different environments. An analysis for MgO should be definitive. Donbassite lacks both substantial Mg and Li, and although this restricts its occurrence to very Al-rich environments, it does not necessarily rule out occurrences in soils, sediments, hydrothermal veins, metamorphosed aluminous sediments, etc. Donbassite is less abundant in nature than sudoite and cookeite, presumably because of its unique composition. X-ray diffraction study should be supplemented by chemical analysis for suspected occurrences of all Al-rich chlorites.

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