The electron-diffraction pattern for a sample of biotite (1M-trioctahedral) shows weak diffractions violating the C-centring of the most commonly reported space-group C2/m. Al-Si ordering in trioctahedral micas, for which Al:Si is 1:3, is likely on a unit-cell scale only in the space groups P2, P1, P1' and P21. There are two distinct Al-Si ordered primitive centrosymmetric triclinic structures, here designated P1 and P1'. Taking into account the principle of aluminum avoidance and other crystal-chemical considerations, the order of these space groups, from most favorable to least, is P21, P1, P2, = P1'. There are four complexions of the P21 structure. Domains of the different P21 complexions would be related by reflection twinning, rotation twinning or inversion twinning. If the domains are very small, X-ray-diffraction and even electron-diffraction patterns may suggest the pseudosymmetry C2/m.

Keywords: biotite, trioctahedral micas, Al-Si ordering.

INTRODUCTION

The space group for most 1M trioctahedral micas has been reported as C2/m (Deer et al. 1963); it is the basis for several structural refinements (Hazen & Burnham 1973, McCauley et al. 1973, Takeda & Morosin 1975, Takeda & Ross 1975). Biotite from a pegmatite, whose locality is unknown, was examined in this study by transmission-electron microscopy (TEM). The composition of the biotite is reported in Table 1. The c-axis diffraction pattern (Fig. 1) shows several violations of the C-centring in space group C2/m. The violations are weak, diffuse diffractions of the type hk0 where h + k is odd, including (320), (110), (340), (540), (270), (470), (670), (870), (1,10,0), (3,10,0) (5,10,0) and the mm2 symmetrically related diffractions. The strongest of these are the types (270), (340), (110) and (1,10,0), in decreasing order of intensity. Except for the (270), even these are difficult to see in Figure 1, but are clearly visible on the negatives. The diffraction pattern shown in Figure 1 is not precisely centred, but by examining several near c-axis patterns, it appears that the Laue symmetry is mm2, consistent with the space groups P21/m, P21/m, P2/a, P2/a, P2, Pm, Pa or P21. It is the purpose of this paper to con-

<table>
<thead>
<tr>
<th>Table 1. Composition of Biotite1 (Average of 7 Analyses)</th>
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</thead>
<tbody>
<tr>
<td>Wt. % oxide</td>
</tr>
<tr>
<td>SiO2</td>
</tr>
<tr>
<td>TiO2</td>
</tr>
<tr>
<td>Al2O3</td>
</tr>
<tr>
<td>MgO</td>
</tr>
<tr>
<td>MnO</td>
</tr>
<tr>
<td>MgO</td>
</tr>
<tr>
<td>CaO</td>
</tr>
<tr>
<td>K2O</td>
</tr>
<tr>
<td>Na2O</td>
</tr>
<tr>
<td>TOTAL</td>
</tr>
</tbody>
</table>

1The analyses were performed by H.Y. McSween, Jr. on an MCA 400-S electron microprobe at the University of Tennessee, Knoxville.
2Standard deviation (1 sigma).
3All Fe reported as FeO.

(Traduit par la Rédaction)

Mots-clés: biotite, micas trioctahédriques, degré d'ordre Al-Si.
consider the possible consequences of reduced symmetry, especially with regard to Al–Si order in the biotite just discussed and in trioctahedral micas in general.

The problem is addressed by introducing an Ordering Factor, OF, for the purpose of discriminating between different schemes of Al–Si ordering. The OF is simply calculated from the cell parameters and is purely geometrical. The bases for discriminating between different ordering schemes are the aluminum-avoidance principle (Loewenstein 1954) and charge-balance considerations discussed by Güven (1971).

Loewenstein's (1954) aluminum-avoidance principle asserts that two tetrahedral sites sharing an oxygen atom in a structure are not likely to be both occupied by aluminum. Furthermore, the distance between tetrahedral sites occupied by aluminum tends to be maximized. The principle follows from Pauling's (1960) rules and is satisfied in many structures, including feldspars, nepheline, kalsilite, margarite and sillimanite. For micas, Güven (1971) argued further that two apical oxygen atoms of Al-tetrahedra along the same shared octahedral edge would be especially unfavorable with respect to local balancing of electrostatic charge.

Gatineau (1964) and Gatineau & Mérie (1966) have reported non-Bragg diffractions and streaked diffractions for muscovite, phlogopite and biotite. They suggested short-range ordering of tetrahedrally co-ordinated cations as a possible cause.

**UNIT-CELL GEOMETRY**

Figure 2a is an idealized (001) projection of the tetrahedral sites in one unit cell of a 1M mica (Pabst 1955). The octahedral cation sites between the sheets of tetrahedra and the large cation sites (K, Na, Ca) have been omitted in order to emphasize the geometrical relationship between the sheets of tetrahedra. The maximum symmetry is C2/m, in which all tetrahedral sites (labeled 1, 2, 3, 4 in the upper sheet and 1′, 2′, 3′, 4′ in the lower sheet) are symmetrically equivalent. For the C2/m trioctahedral micas such as biotite or phlogopite, in which Al:Si is 1:3, the Al and Si must be disordered over the 8 equivalent tetrahedral sites, such that the probability of finding an Al atom on any one site is 25%.

Figures 2b, c and d illustrate other C-centred structures compatible with the diffraction symmetry C2/m. In Cm, there are two types of tetrahedral sites, but all sites in one sheet are equivalent, T1 = T2 = T3 = T4 and T1′ = T2′ = T3′ = T4′. Al–Si ordering in this structure would necessarily place more aluminum in one sheet than in the other, a possibility for which there is very little justification. The 2C2 structure has two sites, T1 = T1′ = T3 = T3′ and T2 = T2′ = T4 = T4′. The structure is compatible with the aluminum-avoidance principle. In the 2M1 dioctahedral mica margarite, the Al and Si (Al:Si = 1) are ordered in this way in each tetrahedral-octahedral-tetrahedral unit layer (Guggenheim & Bailey 1975). Figure 2d illustrates another possibility, CT, compatible with the principle of aluminum avoidance. For 1M micas in which Al:Si equals 1:3, the structures represented by Figures 2a, b, c and d can only account for the ‘average’ or statistical distribution of Al and Si atoms. These space groups cannot be used to illustrate the actual distribution of Al and Si atoms in any one cell volume of a structure.
Fig. 2. Projection of the ideal 1M mica structure onto the (001) plane. Only the tetrahedral cation sites are shown. The tetrahedra in the upper sheet are numbered 1, 2, 3, 4; the tetrahedra in the lower sheet are numbered 1', 2', 3', 4'. a. The holosymmetric space-group C2/m. b, c, d. Other C-centred structures. e, f, g, h, i, j. Primitive unit-cell structures. Except for C2/m, in each structure, one set of symmetrically equivalent sites is stippled. For C2/m, all sites are equivalent.
Within a single unit-cell volume, the distribution of Al and Si atoms must correspond to one of the primitive structures \( P2 \), \( P\bar{1} \), \( P2_1 \), or \( P\bar{1}' \) (Fig. 2g, h, i, j). Note that there are two distinct types of primitive centrosymmetric triclinic Al–Si ordered structures, here designated \( P\bar{1} \) and \( P\bar{1}' \). The \( P2/m \) structure (Fig. 2e), in which there are only two types of sites, \( T1 = T1' = T4 = T4' \) and \( T2 = T2' = T3 = T3' \), can be discounted for two reasons. Firstly, the multiplicities of the two sites (4 each) are not compatible with an Al:Si ratio of 1:3 in 1M trioctahedral micas. \( P2/m \) cannot represent the arrangement of Al and Si atoms in one unit-cell volume in which Al:Si equals 1:3. Secondly, the structure precludes compliance with the aluminum-avoidance principle, except where the space group represents a statistical average of different orientations of some lower-symmetry structure such as \( P\bar{1} \), \( P2 \) or \( P\bar{1}' \). The \( P2_1/m \), \( P2/a \) and \( P2_1/a \) structures (not shown in Fig. 2) can be discounted for the same reasons. The unit-cell type \( Pm \) (Fig. 2f) is unlikely because if the Al atoms are in any of the four types of sites \( (T1 = T4, T2 = T3, T1' = T4' \) or \( T2' = T3' \) ) the resulting structure of \( Pm \) cells alone violates the aluminum-avoidance principle. The \( Pa \) structure (not shown in Fig. 2) would also violate the aluminum-avoidance principle. The four remaining unit-cell symmetries, \( P2 \), \( P\bar{1} \), \( P2_1 \) and \( P\bar{1}' \) (Fig. 2g, h, i, j)

Fig. 3. The complexions of the symmetry types \( P2_1, P2, P\bar{1}, P\bar{1}' \). In each case, one set of symmetrically equivalent Al' sites is stippled.
Fig. 4a (left). The four vectors of the type \((T-T')\), where \(T'\) is a tetrahedral site in the lower sheet and \(T\) is a tetrahedral site in the upper sheet. Formulæ for calculating the \((T-T')\) distances, \(e, f, g\) and \(h\) are presented in Table 1. b, c, d, e. The \((T=T')\) distances for each of the symmetry types \(P2, P'1', P2_1,\) and \(P1\). The stippled tetrahedra are occupied by Al.
are possible, but not necessarily equally likely, as will be demonstrated in the next section.

Figure 3 shows, in matrix form, the four complexions of each unit-cell type $P_2$, $P_1$, $P_2$, and $P_1'$. Each complexion of each unit-cell type is designated by the tetrahedral sites occupied by the 2 Al atoms in the unit cell. Hence, $P_2$ can be represented by $T1-T1'$, $T2-T2'$, $T3-T3'$, $T4-T4'$, the principal diagonal of the matrix. $P_1$ is represented by $T4-T1'$, $T3-T2'$, $T2-T3'$ or $T1-T4'$. $P_1'$ is represented by $T2-T1'$, $T1-T2'$, $T4-T3'$ or $T3-T4'$. And $P_2'$ is represented by $T3-T1'$, $T1-T3'$, $T4-T2'$ or $T2-T4'$. Complexions of the same symmetry are related to one another by at least one symmetry element ($A_2$, $m$ or $i$) with or without a translation. For instance, $P_1'$ ($T4-T1'$) is related to $P_1'$ ($T1-T4'$) by rotation, whereas $P_1'$ ($T1-T4'$) is related to $P_1'$ ($T2-T3'$) by rotation plus translation $(\frac{1}{2}a + \frac{1}{2}b)$.

Because all complexions of a given symmetry-type have the same energy, the four complexions of a given symmetry-type are equally likely in an actual structure composed of domains of one symmetry-type. The domains would be related by twinning or antiphase boundaries. If the domains are sufficiently small, the diffraction symmetry may show the average of the four complexions, which is $C2/m$.

**THE MOST FAVORABLE SYMMETRY-TYPE**

Although all complexions of a symmetry type have the same energy and are, therefore, equally likely, it does not follow that all symmetry types are energetically the same. This should be obvious, if for no other reason than the distances between tetrahedrally co-ordinated aluminum atoms differ from one symmetry type to another.

Other factors considered equal, however small the energy differences, the symmetry types are distinguished on the basis of the distance between an Al atom in one sheet and the nearest Al atom in the other sheet in the same unit-layer. Regardless of the effectiveness of screening by octahedrally co-ordinated cations of the electrostatic interaction between $T$ and $T'$ sites, the energy differences between the four symmetry types must be related, at least indirectly, to the relative positions of the Al atoms. According to the principle of aluminum avoidance (Loewenstein 1954) and charge-balance considerations (Güven 1971), the most stable structure maximizes the distances between tetrahedrally co-ordinated Al atoms. Because the $T$-$T'$ interactions are ultimately electrostatic, it may be assumed that the importance of a particular interaction is inversely proportional to the distance between the $T$ and $T'$ cations. It is convenient to design an ordering factor, OF, that behaves like an energy function and is minimized for the most favorable (i.e., most stable) Al-Si arrangement. For this reason, the OF is formulated in terms of inverse interatomic distances in the following way for one unit cell having the ideal Al:Si ratio of 1:3.

$$\text{OF} = (\text{Al-\text{Al}}')^{-1} - \Sigma_{\text{Si}} (\text{Si-\text{Al}}')^{-1}$$

The $(\text{Al-\text{Al}}')^{-1}$ term is the inverse of the distance between a reference Al' in the 'lower' sheet and the closest Al in the 'upper' sheet. Each $(\text{Si-\text{Al}}')^{-1}$ term is the inverse of the distance between the reference Al' and one of the three Si atoms in the upper sheet. The formulation behaves like an energy function wherein the unfavorable Al-Al' interaction has a positive contribution and the favorable Si-Al' interactions have negative contributions.

**TABLE 2. EQUATIONS FOR T-T' AND EVALUATION OF O.F. FOR AN IDEAL IM MICA**

<table>
<thead>
<tr>
<th>EXAMPLE MICA</th>
<th>$a = 5.3$ Å, $b = \sqrt{2} \times 5.3$ Å, $c = 10.15$ Å, $\beta = 100^\circ$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T - T'$</td>
<td>$f = \sqrt{(0.56 \text{ c sin}B)^2 + (0.35 \text{ a} + 0.56 \text{ c cos}B)^2 + (b/6)^2}$</td>
</tr>
<tr>
<td>$e = \sqrt{(0.56 \text{ c sin}B)^2 + (0.35 \text{ a} + 0.56 \text{ c cos}B)^2}$</td>
<td></td>
</tr>
<tr>
<td>$g = \sqrt{(0.56 \text{ c sin}B)^2 + (0.35 \text{ a} + 0.56 \text{ c cos}B)^2 + (b/3)^2}$</td>
<td></td>
</tr>
<tr>
<td>$h = \sqrt{(0.56 \text{ c sin}B)^2 + (0.35 \text{ a} + 0.56 \text{ c cos}B)^2 + (b/2)^2}$</td>
<td></td>
</tr>
<tr>
<td>$\text{OF}(P_T') = f^{-1} - e^{-1} - g^{-1} - h^{-1} = -0.288$ Å$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>$\text{OF}(P_2') = e^{-1} - f^{-1} - g^{-1} - h^{-1} = -0.288$ Å$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>$\text{OF}(P_T) = g^{-1} - e^{-1} - f^{-1} - h^{-1} = -0.326$ Å$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>$\text{OF}(P_{2}) = h^{-1} - e^{-1} - f^{-1} - g^{-1} = -0.354$ Å$^{-1}$</td>
<td></td>
</tr>
<tr>
<td>$\text{OF}(P_{2'}) = e^{-1} - f^{-1} - g^{-1} - h^{-1} = -0.288$ Å$^{-1}$</td>
<td></td>
</tr>
</tbody>
</table>
Figure 4a illustrates the relevant $T-T'$ distances. Table 2 gives the formulae for calculating the different $T-T'$ distances using the atomic co-ordinates for the ideal $1M C2/m$ mica of Pabst (1955). For some complexions of each symmetry type, it is necessary to select a new set of lattice points, translated by $\frac{1}{2}a + \frac{1}{2}b$, in order that the shortest possible (Al-Al') distance be within one unit cell and all (T-Al') distances be of the types shown in Figure 4a and listed in Table 2. Figures 4b and e show complexions of $P2$ and $P1$, respectively, for which the lattice must be translated by $\frac{1}{2}a + \frac{1}{2}b$ (dashed unit-cells) for the shortest (Al-Al') to be within a unit cell. For the purpose of this paper, the values $a = 5.3$ Å, $b = 3.6\times5.3$ Å and $d(001) = 10$ Å were used for computing the example values of $T-T'$ and $(T-T')^{-1}$ in Table 2. Values of OF are also reported for the symmetry types $P2_1$, $P1$, $P1'$ and $P2$. On the basis of the shortest Al-Al' interaction alone or the Si-Al' interactions alone or the OF, the order of the symmetry types is $P2_1$, $P1$, $P2 \approx P1'$ in order of decreasing stability.

The OF demonstrates that, whereas there is not an order-of-magnitude difference between any two symmetry types, the differences are significant in comparison with the absolute values of OF. Whereas it is practically obvious by visual inspection of Figure 3 that $P2_1$ is in better agreement with the principle

![Diagram](https://via.placeholder.com/150)

**Fig. 5.** The different kinds of twinning involving domains of different complexions of $P2_1$. a. The complexions $(T1-T3')$ and $(T4-T2')$ are related by (010) reflection twinning. b. The complexions $(T1-T3')$ and $(T3-T1')$ are related by rotational twinning. c. The complexions $(T1-T3')$ and $(T2-T4')$ are related by inversion twinning.
of aluminum avoidance and Güven's (1971) rule than the other symmetry types, it is not obvious just how much better is the agreement. The OF provides an objective basis for comparing different states of ordering.

The departure of actual structures from Pabst's (1955) ideal 1M C2/m mica is small (Radoslovich & Norrish 1962, Hazen & Burnham 1973, McCauley et al. 1973, Takeda & Morosin 1975) and is such that the relative position of the tetrahedral sites remains practically ideal. Most of the departure from ideality affects the oxygen atoms in the structure. The equations given in Table 2 for calculating T-7, should apply to any 1M mica, ideal or nonideal.

The OF is easily adapted to micas in which Al:Si equals 2:2, such as a hypothetical 1M margarite. In the case of 1M margarite, there are two space groups compatible with Al-Si ordering, C2 and C1 (Fig. 2c, d). The OF predicts that the C2 structure is more stable than C1. Interestingly, Guggenheim & Bailey (1975) have shown that individual tetrahedral-octahedral-tetrahedral layers in margarite-2M, are ordered, like in the C2 arrangement.

P2₁ DOMAINS

Figures 5a, b and c illustrate three kinds of boundaries separating different complexions of the symmetry type P2₁. T₁-T₃' and T₄-T₂' domains are related by (010) reflection twinning. T₁-T₃' and T₃- T₁' domains are related by rotation twinning. T₁-T₃' and T₂-T₄' domains are related by inversion twinning. Because the four complexions are energetically identical, all four are likely to occur in natural micas. Hence, the three types of twin boundaries should occur, but may not be equally developed because the twin boundaries are not energetically equal.

In the statistically averaged 1M C2/m structures, each unit-cell volume must be ordered according to one of the four symmetry types of Figure 3, most likely as the P2₁ symmetry type. The four complexions of P2₁ would occur as domains, however small they may be. If sufficiently small, the domain model for 1M micas would provide the observed C2/m diffraction-symmetry, which is the combination of the twinning operations and P2₁.

CONCLUSIONS

The most stable unit-cell symmetry type for 1M micas is P2₁. There are four complexions of this symmetry type. The different complexions occur in domains, however small, related by three types of twinning operations.

An ordering factor OF, like the one proposed in this paper, may be useful in predicting the most likely schemes of ordering in other minerals. The OF has the advantage of being purely geometrical, yet it behaves like an energy function, such that the most stable ordered arrangement has the lowest OF. Whereas this approach is not as rigorous as molecular orbital theory, the OF provides adequate results, is simple to calculate and has a sound theoretical basis.

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