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Twins and allotwins of basic mica polytypes: theoretical derivation and identification in the reciprocal space

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The geometry of the diffraction pattern from twins and allotwins of the four basic mica polytypes $(1M, 2M_1, 3T, 2M_2)$ is analysed in terms of the 'minimal rhombus', a geometrical asymmetric unit in reciprocal space defined by nine translationally independent reciprocal-lattice rows. The minimal rhombus contains the necessary information to decompose the reciprocal lattice of twins or allotwins into the reciprocal lattices of the individuals. The nine translationally independent reciprocal-lattice rows are divided into three types (S, D and X): rows of different type are not overlapped by the $n \times 60^{\circ}$ rotations about \mathbf{c}^* , which correspond to the relative rotations between pairs of twinned or allotwinned individuals. A symbolic representation of the absolute orientation of the individuals, similar to that used for layers in polytypes, is introduced. The polytypes 1M and $2M_1$ undergo twinning by reticular pseudo-merohedry with five pairs of twin laws: they produce twelve independent twins, of which nine can be distinguished by the minimal rhombus analysis. The $2M_2$ polytype has two pairs of twin laws by pseudo-merohedry, which give a single diffraction pattern geometrically indistinguishable from that of the single crystal, and three pairs of twin laws by reticular pseudo-merohedry, which give a single diffraction pattern different from that of the single crystal. The 3T polytype has three twin laws: one corresponds to complete merohedry and the other two to selective merohedry. Selective merohedry produces only partial restoration of the weighted reciprocal lattice built on the family rows and the presence of twinning can be recognized from the geometry of the diffraction pattern.

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1. Introduction

Micas, through their polytypes, give one of the most complex series of inorganic structures known to date (Smith & Yoder, 1956; Ross *et al.*, 1966; Takeda, 1967; Rieder, 1970; Takeda & Ross, 1995; Nespolo & Takeda, 1999; Kogure & Nespolo, 1999). Micas often form twins but the presence of twinning is not always evident through a morphological analysis. Oriented crystal associations of different polytypes with the same chemical composition (allotwins: Nespolo, Kogure & Ferraris, 1999) or with different chemical compositions (epitaxy/syntaxy) have also been reported (*e.g.* Babu, 1969; Rieder, 1970; Gresens & Stensrud, 1971; Tatekawa, 1975; Nespolo, Kogure & Ferraris, 1999).

Takano & Takano (1958) introduced the definitions of 'real polytypism' and 'apparent polytypism' to distinguish true polytypes from twins that can be mistaken for polytypes. Geometrical criteria for distinguishing between real and

apparent polytypism have recently been developed (Nespolo, 1999a) and applied to reinvestigate a previous finding of a long-period polytype of biotite (Nespolo & Takeda, 1999). The purpose of the present research is to obtain general criteria to decompose the composite weighted reciprocal lattice (w.r.l.) of a mica twin or allotwin, as it appears in the diffraction pattern, into the w.r.l. of the individuals, obtaining the relative rotations between individuals (preliminary account in Nespolo, Takeda, Ferraris & Kogure, 1997). Since more than one law may correspond to the same orientation of the individuals, the identification of the twin or allotwin laws requires a morphological analysis.

Micas are built by a layer (M layer) [C12/m(1)] in which an octahedral (O) sheet $[P(\bar{3})1m]$ is sandwiched between two

¹ The term 'individual' is here used to indicate one crystal in a twin and the term 'single crystal' to mean an untwinned crystal.

tetrahedral (T) sheets [P(6)mm] (Pabst, 1955; Durovič, 1994a) [layer-group notation after Dornberger-Schiff (1959); details given by Merlino (1990)]. In the O sheet, three translationally independent cation sites exist: the M1 site, on the mirror plane, and a pair of M2 sites, related by the mirror plane (Fig. 1a). Cleavage normally occurs between neighbouring M layers (Kogure, 1997). Several structural distortions are present in the real layer (Abbott & Burnham, 1988), which however mainly influence the diffraction intensities and much less the geometry of the diffraction pattern (Takeda & Ross, 1995; Nespolo, 1999a). The so-called 'Pauling model' (Pauling, 1930) neglects all the distortions, the most relevant of which is the ditrigonal rotation of the tetrahedra, which reduces the symmetry of the T sheets to P(3)1m (Fig. 1a) and is described by the so-called 'trigonal model' (Nespolo, Takeda, Kogure & Ferraris, 1999).

The thickness of the unit layer corresponds to the period along ${\bf c}$ for the 1M polytype $(c_{1M}\approx 10~{\rm \AA})$. The vertical distance between two closest interlayer cations is $c_0=c_{1M}\sin\beta_{1M}=1/c_1^*$ (Backhaus & Ďurovič, 1984; Nespolo, 1999a), where c_1^* (about $0.1~{\rm \AA}^{-1}$) is the reciprocal-space period along ${\bf c}^*$ for 1M. In the diffraction pattern of an N-layer polytype, N reflections appear in the c_1^* repeat along some reciprocal-lattice rows (nonfamily rows: see below) parallel to ${\bf c}^*$.

All mica polytypes have an (ideally) orthohexagonal axial setting based on two (pseudo)orthohexagonal axes (a, b) in the plane of the layer [(001) plane] (Figs. 1a and 1b), for which the relation

$$b = a3^{1/2}, \quad \gamma = 90^{\circ}$$
 (1)

is approximately obeyed. The a and b axes define the two shortest mutually orthogonal translations in the plane of the layer. A (pseudo)orthohexagonal setting based on (a, b) axes can be built, with period along \mathbf{c} equal to $N'c_0$, where

orthogonal polytypes:
$$N' = N$$

non-orthogonal polytypes: $N' = 3N$. (2)

This setting is labelled C_1 (Arnold, 1996; Nespolo, Takeda & Ferraris, 1997; Nespolo $et\ al.$, 1998)² and its cell coincides with the conventional cell of orthogonal polytypes, whereas it contains three conventional cells in the case of non-orthogonal polytypes. The C_1 setting is taken to coincide with Zvyagin's orthohexagonal setting (Zvyagin, 1997) and thus its a and b axes are directed up and left, respectively, in the plane of the drawing (Fig. 1b). All the reflection conditions and the definitions in reciprocal space given below implicitly include the C centring condition $h + k = 0 \pmod{2}$.

Among several kinds of symbols that have been introduced to describe mica polytypes (see Nespolo, Takeda, Kogure & Ferraris, 1999), those giving the absolute orientation and relative displacement of M layers are hereinafter used to derive a similar symbolism for twins. According to Zvyagin (1997), the whole M layer is described through a shortened

symbolism IJK...., where I,J,K,... can take six integral values (1 to 6) (Fig. 1b). These symbols have been called Z symbols (Nespolo, Takeda & Ferraris, 1997); their parity (odd or even) is called 'orientation parity' (Zvyagin, 1997). Z symbols indicate the absolute orientation of the (001) projection of the Z vectors, *i.e.* the vectors connecting two interlayer cations on the opposite sides of an M layer and passing through the origin of the O sheet, oriented along +c (Fig. 2).

Two types of classification of mica polytypes are necessary to develop general criteria for the interpretation of the composite diffraction pattern from mica twins. The OD classification (Backhaus & Ďurovič, 1984; Nespolo, 1999a) distinguishes polytypes on the basis of the equivalence of layer pairs. The reticular classification (Nespolo, Takeda & Ferraris, 1997; Nespolo *et al.*, 1998) emphasizes the metric relations in both direct and reciprocal spaces. A summary of these classifications is given below.

1.1. OD classification of mica polytypes

All polytypes of a substance built on the same structural principle belong to the same family and share a common fictitious structure, called family structure, in which all the possible positions of all OD layers are simultaneously realized (Durovič, 1994b). The symmetry and basis vectors of the family structure are obtained by completing the local symmetry operations of a space groupoid of any member of the family, to global symmetry operations of a space group (Fichtner, 1977). The Fourier transform of the family structure corresponds to a three-dimensional subset of reflections, called family reflections (Dornberger-Schiff & Fichtner, 1972), which are always sharp and are common to all polytypes of the same family. The remaining ones are called *nonfamily reflec*tions and are typical of each polytype; they can be sharp or diffuse, depending whether the polytype is ordered or not (Ďurovič & Weiss, 1986; Ďurovič, 1997, 1999). The family reflections correspond to a subgroup of translations of the polytype reciprocal lattice, whereas the basis vectors of the family structure correspond to a supergroup of translations of the polytype direct lattice. The lattices corresponding to the family structure are thus a superlattice in direct space and a sublattice in reciprocal space.⁵ Family rows are called the reciprocal-lattice rows corresponding to family reflections.

² Some authors, such as the OD school, adopted the C_2 cell, which has the a axis bisecting the hexagonal a_1 and a_2 axes (e.g. Ďurovič et al., 1984).

³ In the most general case, **Z** vectors are defined by pairs of vectors describing half-layers. Within the so-called homo-octahedral approximation (Nespolo, Takeda, Kogure & Ferraris, 1999), which assumes the origin of the O sheet at the M1 site, the two halves of a layer have the same orientation and thus the above shortened symbolism is adopted. For details, see Zvyagin (1997).

⁴ A *global* (also called *total*) operation is an ordinary space-group operation, *i.e.* a symmetry operation valid in the whole crystal space. A *local* operation is a space-groupoid operation, *i.e.* a symmetry operation valid in a subspace of the crystal space. A *partial* operation is a nonsymmetry operation valid in a subspace of the crystal space (Sadanaga *et al.*, 1980).

⁵ Since the lattice obtained from the original one by taking a subgroup (supergroup) of translations has a larger (smaller) unit cell, in some publications (e.g. Santoro & Mighell, 1972; Giacovazzo, 1992; Nespolo, Takeda, Kogure & Ferraris, 1999), the terms superlattice and sublattice are defined in the opposite way. Here we adopt the definition in terms of the group-subgroup relations, in agreement with International Tables for Crystallography, Vol. A, 5th ed. (Th. Hahn, personal communication).

For micas, within the Pauling model, the family reflections correspond to $h = 0 \pmod{3}$ and $k = 0 \pmod{3}$ and are common to all polytypes (Ďurovič *et al.*, 1984; Weiss & Wiewióra, 1986). Reflections in the c_1^* repeat obey the presence criterion $l = 0 \pmod{N'}$, where N' is defined in

equation (2). The family structure is ninefold (the subgroup of translations in the reciprocal space has order 9) and has symmetry P6/mmm. Within the trigonal model, the family reflections correspond to $k=0 \pmod{3}$ and the family structure is threefold (the subgroup of translations in the reciprocal

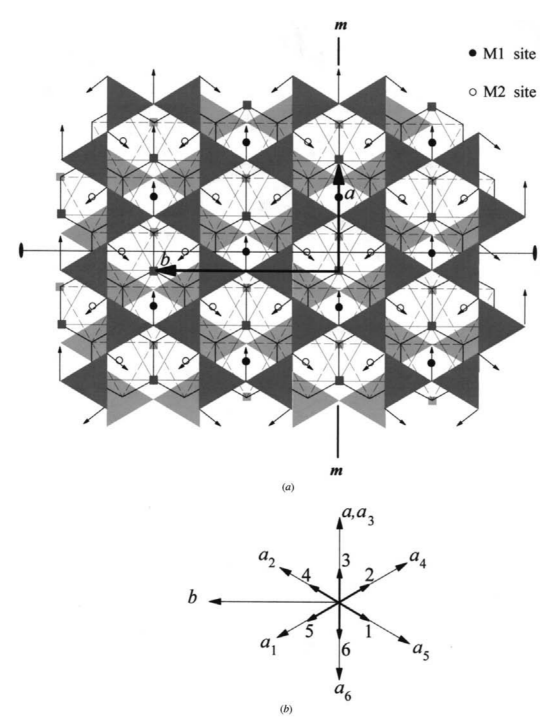


Figure 1
(a) Projection of the M layer onto the (001) plane. Dark- and light-grey T sheets are above and below the O sheet, respectively. Dark- and light-grey squares represent interlayer cations above and below the M layer, respectively. M1 sites are shown by black circles. Thin arrows indicate the direction of the ditrigonal rotation (shown only for the upper T sheet). The independent symmetry elements (twofold rotation axis and symmetry plane) of the 2/m point group are shown. (b) Axes (a, b) of the space-fixed reference $(C_1$ setting) and of the structure-related references in the six possible orientations (a_1-a_6) and corresponding Z symbols. The C axis of the C_1 setting is perpendicular to the layer. The six axes a_1-a_6 indicate the six possible directions of the structure-related references, and thus the six possible orientations of the layer (b_1-b_6) axes are not shown). The direction of the intralayer displacement is indicated by the C_2 symbol C_3 when the C_4 axis is parallel to the space-fixed C_4 axis (modified after Zvyagin, 1997).

space has order 3). Family reflections for the threefold structure are no longer common to all polytypes but depend upon the relative rotations between successive layers. Within the trigonal model, mica polytypes are thus classified in the following three kinds:

(i) Subfamily A polytypes: successive layers are rotated by $2n \times 60^{\circ}$ only and are described by Z symbols of the same parity. The threefold family structure has symmetry $H_R\bar{3}1m$ (the subscript R indicates that the smallest cell is rhombohedral). Reciprocal-lattice rows corresponding to $h \neq 0 \pmod{3}$ and $k = 0 \pmod{3}$ are family rows and show a single reflection in the c_1^* repeat, obeying the presence criterion $l_{C_1} = (N'h/3) \pmod{N'}$ if the parity of layers is odd (obverse setting of the family structure); instead, $l_{C_1} = (2N'h/3) \pmod{N'}$ if the parity of layers is even (reverse setting of the family structure).

(ii) Subfamily B polytypes: successive layers are rotated by $(2n+1)\times 60^\circ$ only and are described by Z symbols of alternating parity. The threefold family structure has symmetry $H6_3/mcm$. Reciprocal-lattice rows corresponding to $h \neq 0 \pmod{3}$ and $k = 0 \pmod{3}$ are family rows and show two reflections in the c_1^* repeat, with the presence criterion $l_{C_1} = 0 \pmod{N'/2}$.

(iii) Mixed-rotation polytypes: successive layers are rotated by both $2n \times 60^{\circ}$ and $(2n+1) \times 60^{\circ}$ and are described by Z symbols of mixed nonalternating parity; the only family rows are those corresponding to both h and $k = 0 \pmod{3}$. The only family structure is the ninefold one.

The period along \mathbf{c} of the family structure is c_0 in the Pauling model, whereas in the trigonal model it is $3c_0$ (subfamily A) or $2c_0$ (subfamily B) (Ďurovič $et\ al.$, 1984).

1.2. Reticular classification of mica polytypes

The reticular classification of mica polytypes is based upon the projection \mathbf{c}_n of the c axis onto the (001) plane. Mica polytypes are divided into three kinds:

- (i) orthogonal polytypes: $\mathbf{c}_n = (0, 0)$;
- (ii) Class a polytypes: $\mathbf{c}_n = (-1/3, 0)$;
- (iii) Class b polytypes: $\mathbf{c}_n = (0, -1/3)$.

The number of layers N in a mica polytype is suitably expressed as:

$$N = 3n(3K + L), \tag{3}$$

where *n* defines the *Series*, *L* the *Subclass* and *K* is an integer that appears in the transformation matrices relating different axial settings $(n \ge 0, L = 1, 2, K \ge 0)$.

The geometry of the diffraction pattern of a twinned mica polytype belonging to *Series n* is often similar to that of an untwinned mica polytype belonging to *Series n* + 1 (Nespolo, 1999a). Therefore, when the diffraction pattern of a mica crystal shows 3p equally spaced reflections in the c_1^* repeat, the possibility of apparent polytypism must be taken into account. The derivation of the possible twin diffraction patterns is thus a prerequisite for the study of polytypism.

The present geometrical analysis of mica twins and allotwins is devoted to the four most common polytypes, namely 1M (Z=3; subfamily A, Class a, Series 0), $2M_1$ (Z=24; subfamily A, Class a, Series 0), 3T (Z=351 or 315; subfamily A, orthogonal, Series 1) and $2M_2$ (Z=45; subfamily B, Class b, Series 0). These are by far the most common polytypes and are called $basic\ mica\ polytypes$ because the stacking sequence of longer-period polytypes is usually based on one of them

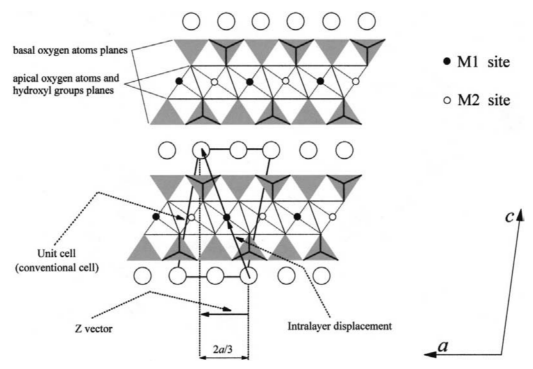


Figure 2
Projection along the b axis of the 1M mica structure, showing two layers with the same orientation. M1 and M2 cationic sites in the O sheet occupy crystallographically distinct positions (cf. Fig. 1). The scale along \mathbf{c} is compressed (modified after Nespolo, 1999b).

(Baronnet & Kang, 1989). The general approach derived henceforth can however be easily generalized to other polytypes.

2. Twin laws for mica polytypes

The basis for the derivation of the twin laws for mica polytypes in the general case is given in Nespolo (1999a). Here, the treatment is focused onto the four basic polytypes, and the derivation is performed through coset decomposition (Giacovazzo, 1992). Rotations in direct and reciprocal space are taken counterclockwise and clockwise, respectively. Symbols for the crystal systems of polytypes are given according to Guinier et al. (1984). In the case of twinning by merohedry, the twin laws are expressed through Shubnikov's two-colour group notation (the twin elements are dashed: Curien & Le Corre, 1958). Twins in which two or more twin laws have operated are called composite twins (Nespolo, Takeda, Ferraris & Kogure, 1997). The complete twin (Curien & Donnay, 1959) is a twin by merohedry or reticular merohedry in which the number of individuals generated from the original one is equal to the number of possible twin laws.

Mica polytypes can belong to 24 space groups of five crystal systems (Takeda, 1971): A, M, O, T and H. The pair of (pseudo)orthohexagonal axes (a, b) in the plane of the layer is common to all polytypes. All polytypes admit a hexagonal twin lattice based on a pair of orthohexagonal axes (a_H, b_H) parallel to (a, b): its symmetry corresponds to that of the ninefold family structure, namely P6/mmm (Dornberger-Schiff et al., 1982; Backhaus & Durovič, 1984). The cell of the twin lattice coincides with the C_1 cell for hexagonal and trigonal polytypes, and in this case a hexagonal cell can be chosen as well (Fig. 3). For all the others, equation (1) is obeyed by (a_H, b_H) but only approximately by (a, b). The angular (ε) (Donnay et al., 1964) and the linear (η) (Zvyagin & Drits, 1996) deviations from the orthohexagonality (Fig. 4a) are related to the twin obliquity. Moreover, for non-orthogonal polytypes, the length of the \mathbf{c}_n projection (§1.2) actually deviates from |a|/3 or |b|/3 by a quantity that depends mainly upon the crystal chemistry. Rieder (1970) described this deviation in terms of the monoclinic angle and showed that it is not negligible for dioctahedral and Li-Fe trioctahedral micas. The practical consequence is that the C_1 cell is no longer rigorously orthogonal and the reflections from twinned individuals may have a zigzag splitting around the position of the ideal row. This clearly indicates the presence of twinning. The relative rotations between individuals can be obtained also in this case by applying the geometrical treatment developed hereafter, provided the small splitting is neglected.

2.1. Monoclinic holohedral polytypes

Three of the four basic polytypes are monoclinic holohedral $(1M: C2/m; 2M_1 \text{ and } 2M_2: C2/c)$ and the C_1 cell contains three conventional cells. For 1M and $2M_1$ polytypes (Class a), twinning is by reticular pseudo-merohedry, with twin index 3. Lattice nodes are completely restored by the twin operation in

one plane out of three along \mathbf{c}^* , whereas they are not restored at all in the remaining two planes. The lattice of *Class b* polytypes, to which $2M_2$ belongs, admits a primitive cell close to rhombohedral (Nespolo, 1999a) and the classification of twin laws is based on this pseudo-symmetry. Point group 2/m is a subgroup of order 6 of 6/mmm, and thus six cosets can be obtained, but the unique axes do not coincide in the crystal lattice (*b* unique for *Class a* polytypes; *a* unique for *Class b* polytypes) and in the twin lattice (*c* unique). Following Donnay *et al.* (1964), the coset decomposition is here given through left cosets. Results are expressed in the C_1 setting. *Class a*:

$$6/mmm = \left\{1, 2_{[010]}, \bar{1}, m_{(010)}\right\} \cup \left\{6_{[001]}^{-}, 2_{[110]}, \bar{6}_{[001]}^{-}, m_{(130)}\right\} \\ \cup \left\{3_{[001]}^{-}, 2_{[310]}, \bar{3}_{[001]}^{-}, m_{(110)}\right\} \\ \cup \left\{2_{[001]}, 2_{[100]}, m_{(001)}, m_{(100)}\right\} \\ \cup \left\{3_{[001]}^{+}, 2_{[3\bar{1}0]}, \bar{3}_{[001]}^{+}, m_{(1\bar{1}0)}\right\} \\ \cup \left\{6_{[001]}^{+}, 2_{[1\bar{1}0]}, \bar{6}_{[001]}^{+}, m_{(1\bar{3}0)}\right\}.$$

$$(4a)$$

Class b:

$$6/mmm = \left\{1, 2_{[100]}, \bar{1}, m_{(100)}\right\} \cup \left\{3_{[001]}^{-}, 2_{[110]}, \bar{3}_{[001]}^{-}, m_{(130)}\right\} \\ \cup \left\{6_{[001]}^{-}, 2_{[310]}, \bar{6}_{[001]}^{-}, m_{(110)}\right\} \\ \cup \left\{2_{[001]}, 2_{[010]}, m_{(001)}, m_{(010)}\right\} \\ \cup \left\{6_{[001]}^{+}, 2_{[3\bar{1}0]}, \bar{6}_{[001]}^{+}, m_{(1\bar{1}0)}\right\} \\ \cup \left\{3_{[001]}^{+}, 2_{[1\bar{1}0]}, \bar{3}_{[001]}^{+}, m_{(1\bar{3}0)}\right\}.$$

$$(4b)$$

The first coset consists of the symmetry elements of 2/m point group and the other five give the possible twin laws. Because of the nonzero obliquity, the four operations in each coset corresponding to a twin law are not equivalent. The law of Mallard requires that twin elements are lattice planes or rows and that twin operations are crystallographic operations. Therefore, the rotations and rotoinversions about [001] are not possible twin operations because they slightly deviate from $n \times 60^{\circ}$ [equation (1) is only approximately obeyed]. Peacock & Ferguson (1943), Donnay et al. (1964) and Rieder (1970) concluded that the twin elements are the twofold axes in the (001) plane. However, when the composition plane is normal to (001) (see examples in Goldschmidt, 1918; Hurlbut, 1956; Sunagawa, 1964), the twin operation is a twin plane coinciding with the composition plane (Table 1).

2.2. 3*T* polytype

In the case of the 3T polytype (space group $P3_{1,2}12$), the twin lattice and the crystal lattice coincide.

The coset decomposition gives four cosets (indexing in C_1 setting):

$$6/mmm = \left\{1, 3_{[001]}^{+}, 3_{[001]}^{-}, 2_{[010]}, 2_{310}, 2_{3\bar{1}0}\right\}$$

$$\cup \left\{2_{[100]}, 2_{[1\bar{1}0]}, 2_{[110]}, 2_{[001]}, 6_{[001]}^{-}, 6_{[001]}^{+}\right\}$$

$$\cup \left\{m_{(100)}, m_{(130)}, m_{(1\bar{3}0)}, m_{(001)}, \bar{6}_{[001]}^{-}, 6_{[001]}^{+}\right\}$$

$$\cup \left\{m_{(010)}, m_{(110)}, m_{(1\bar{1}0)}, \hat{i}, \bar{3}_{[001]}^{+} + \bar{3}_{[001]}^{-}\right\}. \tag{5}$$

The first coset contains the symmetry elements of the 312 point group and the others the twin elements. These correspond to the three twin laws: 6'2'2; $\bar{6}'m'2$; $\bar{3}'12/m'$. The complete twin contains four individuals and has symmetry 6'/m'' 2'/m'' 2/m'''.

The six symmetry elements in each of the three cosets corresponding to a twin law are symmetrically equivalent under the operations of the point group 312: the twin law that actually operated appears in the crystal morphology and is called representative operation of the coset (Nespolo & Ferraris, 2000). The 6'2'2 and $\bar{6}'m'^2$ twin laws correspond to class IIA, according to the Nespolo & Ferraris (2000) definition: for them, the twin elements do not belong to the point group of the family structure. This situation corresponds to 'selective merohedry' (Nespolo, Ferraris & Durovič, 1999), which restores all the lattice nodes corresponding to the nonfamily reflections, but not all the lattice nodes corresponding to the family reflections of the threefold family structure. The presence of twinning is thus recognizable from the geometry of the diffraction pattern, since it shows violations of the non-space-group absences of the family reflec-

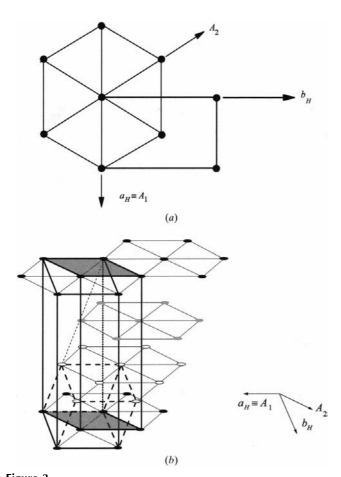


Figure 3

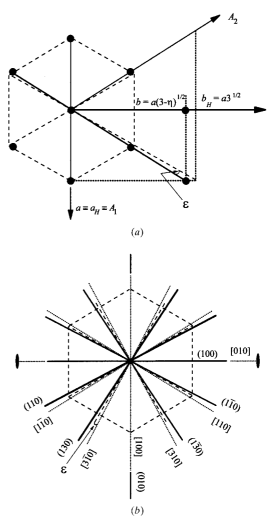
(a) Relation between hexagonal (A_1, A_2) and orthohexagonal (a_H, b_H) axes and location of the orthohexagonal cell C_1 . (b) The conventional monoclinic cell [(001) base dashed], the (pseudo)orthohexagonal cell (full lines), and the hexagonal cell (shaded) built overlapping three conventional cells. The scale along c is compressed (modified after Nespolo, Takeda & Ferraris, 1997).

tions. The $\bar{3}'12/m'$ twin law corresponds to class I (Catti & Ferraris, 1976; Nespolo & Ferraris, 2000): all the twin elements belong to the point group of the family structure and twinning is by complete merohedry (Nespolo, Ferraris & Ďurovič, 1999).

2.3. Summary

The twin elements for the four basic polytypes can be summarized as follows:

(i) 1M and $2M_1$ polytypes (Class a) undergo twinning by reticular pseudo-merohedry, with twin index 3. There are five pairs of twin elements: [310]/(110), $[3\bar{1}0]/(1\bar{1}0)$, [110]/(130), $[1\bar{1}0]/(1\bar{3}0)$ and [100]/(100).



Projection onto (001) of the C_1 cell (a) and location of the pseudosymmetry elements (b). A_1 , A_2 : hexagonal axes; a_H , b_H : orthohexagonal axes; a, b: pseudo-orthohexagonal axes. ε and η (exaggerated) are the angular and linear deviations from hexagonality. The figure is drawn for the case $b < b_H$. Black circles: lattice nodes of the crystal lattice; dashed lines: H cell of the twin lattice; dotted lines: C_1 cell built on the hexagonal and pseudo-hexagonal meshes. (b) Dashed lines: H cell of the twin lattice; full lines: pseudo-symmetry planes; dotted lines: twofold axes quasinormal to the planes. [010] and (010) are symmetry elements for the conventional cell in the case of monoclinic Class a polytypes. For monoclinic Class b polytypes, they become [100] and (100) (modified after Nespolo, Takeda & Ferraris, 1997).

 Table 1

 Twin laws for basic mica polytypes and corresponding relative rotations between pairs of twinned individuals.

Polytype	Point group	Twinning		Rotation twi	ns				Reflection tv	wins			
3 <i>T</i>	312	Complete merohedry Selective merohedry	Twin operation Rotation Twin operation Rotation	$[100]_{\pi}$ $(2n+1)\times60^{\circ}$					(010) $2n \times 60^{\circ}$ (100) $(2n+1) \times 60^{\circ}$				
$1M$, $2M_1$	2/ <i>m</i>	Reticular pseudo-merohedry	Twin operation	,	$[110]_{\pi}$ $300^{\circ}-2\varepsilon$	$[3\bar{1}0]_{\pi}$ $120^{\circ}+2\varepsilon$	$[1\bar{1}0]_{\pi}$ $60^{\circ}+2\varepsilon$	$[100]_{\pi}$ 180°	(110) $240^{\circ} + 2\varepsilon$	(130) 300°+2ε	$(1\bar{1}0)$ $120^{\circ}-2\varepsilon$	$(1\bar{3}0)$ $60^{\circ}-2\varepsilon$	(100) 180°
$2M_2$	2/ <i>m</i>	Reticular pseudo-merohedry	Twin operation Rotation	$[310]_{\pi}$ $300^{\circ}-2\varepsilon$		$[3\bar{1}0]_{\pi}$ $60^{\circ}+2\varepsilon$		$[010]_{\pi}$ 180°	(110) 300°+2ε		$(1\bar{1}0)$ $60^{\circ}-2\varepsilon$		(010) 180°
	2/ <i>m</i>	Pseudo-merohedry	Twin operation Rotation		$\begin{array}{c} [110]_{\pi} \\ 240^{\circ} - 2\varepsilon \end{array}$		$\begin{array}{c} [1\bar{1}0]_{\pi} \\ 120^{\circ} + 2\varepsilon \end{array}$			(130) 240°+2ε		$(1\bar{3}0)$ $120^{\circ}-2\varepsilon$	

- (ii) $2M_2$ polytype (*Class b*) has five pairs of twin laws too, namely: [310]/(110), $[3\bar{1}0]/(1\bar{1}0)$, [110]/(130), $[1\bar{1}0]/(1\bar{3}0)$ and [010]/(010). Of these, the two pairs [110]/(130) and $[1\bar{1}0]/(1\bar{3}0)$ are twin elements by pseudo-merohedry, whereas the remaining three pairs correspond to twinning by reticular pseudo-merohedry, with twin index 3.
- (iii) 3T polytypes have three twin laws, of which one $(\bar{3}'12/m')$ corresponds to complete merohedry and the other two to selective merohedry (6'2'2 and $\bar{6}'m'2$).

The five pairs of twin elements of the monoclinic polytypes are mutually almost, but not exactly, perpendicular in the crystal lattice. The deviation from the perpendicularity can occur both in the (001) plane ($\varepsilon \neq 0$, Fig. 4b) and in the (hk0) planes, which are not exactly perpendicular to the (001) plane (the C_1 cell is not exactly orthogonal). For the first four pairs of twin elements, common to both Classes, deviations in both planes occur. For the last pair, namely [100]/(100) for Class a and [010]/(010) for Class b, the deviation in the (001) plane does not exist ($\varepsilon = 0$) but the second one in general remains. It follows that all the five pairs of twin laws can produce the so-called reciprocal twins (Mügge, 1898) or corresponding twins (Friedel, 1904, 1926). These are however hardly distinguishable from their diffraction pattern because of the small value of the twin obliquity.

3. Symbolic description of the orientation of twinned mica individuals. Limiting symmetry

Two twin laws are said to be equivalent when the corresponding twin elements are related by symmetry operations of the crystal. In accordance to the law of Mallard, a twin element is restricted to a lattice element, and the twin operation to a symmetry operation for the lattice (Santoro, 1974).

The three basic monoclinic polytypes all correspond to the point group 2/m. Each of the rotational twin operations, premultiplied by a 180° rotation about the monoclinic symmetry axis, is equivalent to a rotation about the normal to (001). If it is assumed that $|\mathbf{c}_n| = |a|/3$ or |b|/3, the c axis of the C_1 cell is normal to (001). Only in the case of [100] (Class a) and [010] (Class b) does the angle between the twin element and the symmetry element (90°) not depend upon the twin obliquity, whereas for the other twin elements it corresponds to

 $n \times 30^{\circ} \pm \varepsilon$ (Fig. 4b). Two individuals twinned according to [100] (Class a) or [010] (Class b) are thus rotated by 180° around the normal to the layer but the other four rotations correspond to $n \times 60^{\circ} \pm 2\varepsilon$. For reflection twins, the same relative rotations are obtained through application of the (010) (Class a) or (100) (Class b) symmetry plane of the crystal but the deviations from $n \times 60^{\circ}$ rotation are in the opposite direction (Table 1). Since ε is normally very small (Donnay et al., 1964), the possible orientations of individuals in a twin are almost identical with the possible orientations of layers in a polytype, and the shortened Z symbols used for polytypes (written as Z_T , T indicating twin) can be applied also to twins. There are four main differences between Z and Z_T symbols:

- (i) Since in a twin there cannot be two individuals oriented in the same way, the sequence of Z_T never contains the same digit twice.
- (ii) The Z symbol of polytypes has to take into account the space group, whereas Z_T deals only with the symmetry of the point group. The orthohexagonal setting of the first individual is taken to coincide with that of the twin lattice: the first individual is always fixed in orientation $Z_T = 3$ (Fig. 1b), and the orientations of the others are determined by the twin laws.
- (iii) Rotation by 180° of the whole twinned edifice around the a axis of the space-fixed reference changes the $3IJ\dots PZ_T$ symbols into $(6-P)\dots(6-J)(6-I)3$; since the order of the individuals in the twin does not influence the diffraction pattern, this sequence is equivalent to $3(6-I)(6-J)\dots(6-P)$. The latter corresponds to inverting the direction of rotation of the individuals in the twin. Considering the effect on the lattice, the $3IJ\dots P\to 3(6-I)(6-J)\dots(6-P)$ transformation corresponds to reflecting the twin lattice across the (010) plane.
- (iv) The point group of the 3T polytype contains only proper motions. The orientation symbol of an individual produced by a twin operation corresponding to an improper motion is hereinafter distinguished by a small black circle (\bullet) after the Z_T symbol, which indicates a reflection across a symmetry plane for the lattice [e.g. (010)].

The number of independent orientations of the reciprocal lattice of an individual is determined by the lower symmetry between the crystal lattice and the family structure, which hereinafter is called *limiting symmetry*. For *Class a* polytypes, the lattice symmetry is monoclinic and the family structure is

Table 2 Orientation of the individuals building a twin in *Class a* mica polytype.

Angles in parentheses express the counterclockwise rotations of the whole twinned edifice. 'Shift' stands for the shift of the origin along c. (010) means reflection of the twin lattice across the (010) plane, which is equivalent to inverting the direction of rotation of the individuals in the twin, *i.e.* to the symbol transformation $3IJ \dots P \to 3(6-I)(6-J) \dots (6-P)$ (see §3).

Z_T	Equivalent to	Equivalent to	Equivalent to	Z_T	Equivalent to	Equivalent to
34	Unique	-	-	3456	Unique	_
35	Unique	_	_	3451	Unique	_
36	Unique	_	_	3452	4563 (60°)	3456 (shift)
31	53 (120°)	35 (shift)	_	3461	Unique	- ` `
32	43 (60°)	34 (shift)	_	3462	Unique	_
345	Unique	= ` ´	_	3412	5634 (120°)	3456 (shift)
346	Unique	_	_	3561	1345 (240°)	3451 (shift)
341	325 (010)	436 (60°)	346 (shift)	3562	4613 (60°)	3461 (shift)
342	453 (60°)	345 (shift)	_ ` `	3512	5134 (120°)	3451 (shift)
356	134 (240°)	341 (shift)	346	3612	6345 (180°)	3456 (shift)
351	Unique	_ ` `	_	34561	Unique	- ` `
352	463 (60°)	346 (shift)	_	34562	45613 (60°)	34561 (shift)
361	634 (180°)	346	_	34512	56134 (120°)	34561 (shift)
362	413 (60°)	341 (shift)	346	34612	61345 (180°)	34561 (shift)
312	534 (120°)	345 (shift)	- .	35612	13456 (240°)	34561 (shift)
	• /	` '		345612	Unique	- ` ′

rhombohedral: the limiting symmetry is thus monoclinic. Besides, since the unique axes do not coincide (b in the polytypes, c in the family structure), all the six orientations of the individual reciprocal lattice about the normal to (001) are independent. For a $2M_2$ ($Class\ b$) polytype, the lattice symmetry is (pseudo)rhombohedral and the family structure is hexagonal: the limiting symmetry is thus rhombohedral: only two orientations of the individual reciprocal lattice are independent, corresponding to the two parities of Z_T symbols. A common symbol is used for the three equivalent orientations with the same parity, namely U (uneven) and E (even). Finally, for the 3T polytype, the crystal lattice is hexagonal and the family structure is rhombohedral: the limiting symmetry is rhombohedral and the two independent orientations of the individual w.r.l. are again $Z_T = U$ and $Z_T = E$.

3.1. Class a polytypes

Since all the six possible orientations of the individuals correspond to independent orientations of the reciprocal lattice, the possible composite twins can be obtained by calculating the sequences of Z_T symbols for sets of individuals from two to six. The orientation of the first individual is fixed $(Z_T = 3)$ and five possible orientations remain on which m individuals $(1 \le m \le 5)$ have to be distributed. The number of twins is then:

$$N_T(m) = {5 \choose m} = \frac{5!}{m!(5-m)!}.$$
 (6)

Table 2 gives the corresponding 31 sequences of Z_T symbols; only 12 of them represent independent twins. The other 19 simply correspond to a rotation of the whole twinned edifice followed by a shift of the origin along \mathbf{c} , eventually coupled with the inversion of the direction of the rotation of the individuals in the twin [reflection of the lattice across (010)], as in the case of $Z_T = 341$.

3.2. $2M_2$ polytype

Twinning by pseudo-merohedry involves individuals with the same orientation parity of Z_T symbols and produces complete overlap of the individual reciprocal lattice. The reciprocal lattice of the twin is thus geometrically indistinguishable from the reciprocal lattice of the individual. The three twins $Z_T = 35$, $Z_T = 31$ and $Z_T = 351$ are all equivalent to the single crystal, when considering the geometry of their lattice, and are thus represented as $Z_T = U$. Instead, twinning by reticular pseudo-merohedry involves individuals with opposite orientation parity of the Z_T symbols and, considering the lattice only, they are all represented as $Z_T = UE$.

3.3. 3*T* polytype

If the first individual is fixed in orientation $Z_T = U$, the effect of the three twin laws on it is:

6'2'2, twin operation $[100]_{\pi}$ or equivalent: $Z_T = U \rightarrow Z_T = E$;

 $\bar{6}'m'2$, twin operation (100) or equivalent, which can be expressed as $(010) \cdot [100]_{\pi}$: $Z_T = U \rightarrow Z_T = E^{\bullet}$;

3'12/m', twin operation (010) or equivalent: $Z_T = U \rightarrow Z_T = U^{\bullet}$.

The first two twin operations correspond to selective merohedry (§2.2). Because of the non-space-group absences of the family structure (see above), along each family row with $h \neq 0 \pmod{3}$ and $k = 0 \pmod{3}$, two (one from each individual) out of three reflections appear, the third (ideally absent) always corresponding to $l = 0 \pmod{3}$. The third twin operation corresponds to complete merohedry.

4. Translationally independent reciprocal-lattice rows

Since the \mathbf{c}_n projection for non-orthogonal polytypes can ideally take only one of the two values |a|/3 (Class a) or |b|/3 (Class b), in the C_1 setting the metric relation in

reciprocal space is $l_{C_1} \pmod{3} = h \pmod{3}$ (Class a) or $l_{C_1} \pmod{3} = k \pmod{3}$ (Class b). By considering the $l_{C_1} \pmod{3}$ index of reciprocal-lattice nodes on rows related by $n \times 60^{\circ}$ rotations ($0 \le n \le 5$), it has been shown (Nespolo, Takeda, Ferraris & Kogure, 1997) that in the case of the 1M polytype there are only nine translationally independent rows parallel to \mathbf{c}^* (Fig. 5). For the other polytypes, the violation of the additional reflection conditions in principle may cancel the geometrical equivalence between reciprocal-lattice rows. However, the forbidden reflections are weak enough to be neglected and the same classification developed for the 1M polytype holds thus in general (Nespolo, 1999b).

The translationally independent rows are hereinafter indicated as R_i , $1 \le i \le 9$. In each R_i , the same distribution of 'present' and 'absent' reflections is repeated along \mathbf{a}^* and \mathbf{b}^* with 3p and 3q translations (p and q are integers of the same parity). R_i are defined in terms of h and k as: $[h_i \pmod{3}, k_i \pmod{3}, l]$ and are distributed along the edges and diagonals of a rhombus-shaped unit, called a *tessellation rhombus* (Fig. 5, solid lines), which can tessellate the whole reciprocal space by (3p, 3q) translations. A smaller unit, called *minimal rhombus*, can be drawn (Fig. 5, dotted lines), defined by the same R_i each taken only once. Opposite edges are different and, contrary to the tessellation rhombus, the minimal rhombus does not represent a translational unit.

The two rhombi have six possible orientations (Fig. 5), which however represent equivalent descriptions of the same reciprocal lattice: they just differ in the distribution of the R_i . Six equivalent rhombi are obtained by applying all the five rotations (besides the identity) to the h_i , k_i indices of each of

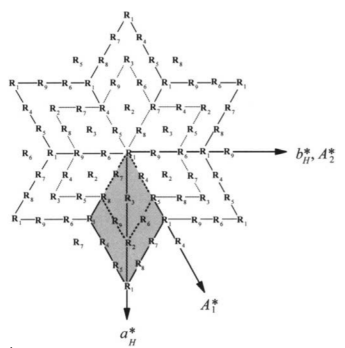


Figure 5 Minimal rhombus (dotted lines; in the foreground) and tessellation rhombus (solid lines) in the six orientations defining the star polygon. The nine translationally independent rows are distinguished by sequence numbers (R_1-R_9) . A_1^* , A_2^* : hexagonal axes; a_H^* , b_H^* : orthohexagonal axes (modified after Nespolo, Takeda, Ferraris & Kagure, 1997).

the nine R_i of the original rhombus and bringing the resulting R_i within the rhombus through a (3p, 3q) translation between equivalent rows. The rows that can be reached by rotating the original rhombus are within a star-polygon located by the six rhombi with a common origin (Fig. 5). The values of p and q to be taken into account are those connecting rows internal to the star-polygon but external to the original rhombus with rows internal to the original rhombus, *i.e.* $(0, \pm 2)$, $(1, \pm 1)$ and $(2, \pm 2)$.

5. Classification of reciprocal-lattice rows

In the c_1^* repeat of nonfamily rows, as many reflections appear as the number N of layers (some of which are very weak because of the additional reflection conditions), each separated by c_1^*/N . In the C_1 setting, the period along \mathbf{c}^* is c_1^*/N' [see equation (2)]. The shortest repeat common to all the four basic polytypes is $c_1^*/6$, which coincides with the period of $2M_1$ and $2M_2$ polytypes, and is a subperiod for 1M and 3T polytypes. By introducing a (pseudo)orthohexagonal setting with period $c_1^*/6$ along \mathbf{c}^* , the *l* indices of the nodes within a c_1^* repeat can be indexed in a common cell for the four basic polytypes. This setting is here called 'twin setting', and the corresponding l indices are indicated as l_T : $l_T(2M_1, 2M_2) = l_{C_1}(2M_1, 2M_2)$, but $l_T(1M, 3T) =$ $2l_{C_1}(1M, 3T)$. The twin setting of the twin lattice is space-fixed and parallel to C_1 , whereas that of the crystal lattice is crystalfixed for each of the individuals building a twin. Since the first individual of the twin is space-fixed ($Z_T = 3$ for Class a or $Z_T = U$ for $2M_2$ and 3T), its twin setting is parallel to C_1 .

Hereinafter, the 'row features' characterize the reciprocallattice rows parallel to \mathbf{c}^* as a whole and the 'node features' characterize each single reciprocal-lattice row, expressing the number and disposition of reciprocal-lattice nodes on them. If the deviation from orthohexagonality (ε, η) is neglected, all mica polytypes share the same row features, described by the regular tessellation $\{3, 6\}$ (Takeda & Donnay, 1965) and the nine R_i can be classified into three types (Fig. 6):

- (i) S (single) rows $[h = 0 \pmod{3}]$ and $k = 0 \pmod{3}$. They are family rows of the ninefold structure (Pauling model): the single reflection always corresponds to $l_T = 0 \pmod{6}$.
- (ii) D (double) rows $[h \neq 0 \pmod{3}]$ and $k = 0 \pmod{3}$. They are family rows of the threefold structure (trigonal model), but nonfamily rows for mixed-rotation polytypes. There are two independent D rows, labelled D_i : i = 1, 2; $h = i \pmod{3}$; $k = 0 \pmod{3}$.
- (iii) *X* (sextuple) rows $[k \neq 0 \pmod{3}]$. They are nonfamily rows for all polytypes. There are six independent *X* rows, labelled X_i : $1 \leq i \leq 6$; $h = i \pmod{3}$; $k = 2 \times (-1)^i \pmod{3}$.

The nine R_i rows are thus classified as: $R_1 = S$; $R_{2-3} = D_{1-2}$; $R_{4-9} = X_{1-6}$. Each of the three types of rows lies on non-

⁶ Sadanaga & Takéuchi (1961), dealing with polysynthetic twins of 1M, classified the reciprocal-lattice rows into S rows (S = single), twin rows and T rows (T = triple), recalling the number of reflections in a c_1^* repeat. That terminology is meaningful when dealing with 1M twins, but not in a general approach. The present classification is a general feature of the $\{3,6\}$ regular tessellation and does not depend on the polytype.

Table 3 Definition of the I_i for the four basic polytypes and their twins.

I indicates the number of nodes on the reciprocal-lattice row. The subscript j is a sequential number (see §5).

$\overline{I_j}$	11	12	1 ₃	21	22	23	24	25	26	31	32	33	34	35	36	37
l (mod 6)	0	2	4	0,2	0,4	0,3	1,4	2,5	2,4	0,2,4	0,1,4	0,2,3	0,2,5	0,3,4	1,2,4	2,4,5
I_{j}	41	42	4 ₃	4_4	4 ₅	4 ₆	51	52	5 ₃	6_1						
<i>l</i> (mod 6)	0,1,2,4	0,1,3,4	0,2,3,5	0,2,3,4	0,2,4,5	1,2,4,5	0,1,2,3,4	0,1,2,4,5	0,2,3,4,5	0,1,2,3,4,5						

intersecting circular orbits centred on c^* . An $n \times 60^\circ$ rotation overlaps rows belonging to the same type only. In Fig. 6, heavy and thin circles describe the orbits of family (S full lines and D dashed lines) and nonfamily (X) rows, respectively. For X rows, two kinds of orbits exist, connecting one (full lines) and two (dashed lines) sets of six rows. The $n \times 60^\circ$ rotations about \mathbf{c}^* lead to an alternate exchange of the two D-type R_i located on the long diagonal of the minimal rhombus, and they exchange the six X-type R_i on the edges of the minimal rhombus in six different ways.

The transformations for the h_i , k_i indices of the nine R_i in the six orientations of the minimal rhombus are shown in Fig. 7, where the distribution of the X_i is given by the arrows. To each of these six orientations, the reflection across the (010) plane, which corresponds to the $Z_T = 3IJ \dots P \rightarrow 3(6-I)(6-J)\dots(6-P)$ transformation (see §3), has to be applied. The second set is obtained from the first one through the sequence $X_4 \rightarrow X_5 \rightarrow X_6 \rightarrow X_1 \rightarrow X_2 \rightarrow X_3$ applied to

Figure 6
Rotational relation between reciprocal-lattice rows parallel to \mathbf{c}^* and definition of the rotational sequences (RS_i^P) . A_1^* , A_2^* : hexagonal axes; a_H^* , b_H^* : orthohexagonal axes. Because of the pseudo-hexagonal symmetry of the (001) plane, each type of row (S, D, X) lies on a circular orbit around \mathbf{c}^* . Heavy orbits: family rows (full lines: S rows; dashed lines: D rows); thin orbits: nonfamily (X) rows (full lines: one set of six rows; dashed lines: two sets of rows). The $n \times 60^\circ$ rotations, which correspond to the relative orientation of twinned or allotwinned mica individuals, do not mix different types of row (S, D, and X).

the original (0°) orientation. Two minimal rhombi that can be transformed into each other by application of one of these 12 operations are equivalent.

The node features (number and disposition of nodes on the reciprocal-lattice rows) are hereinafter identified by a symbol I_j , where I is the number of nodes within the c_1^* repeat and j is a sequence number. The definition of I_j is given in Table 3. Seven I_j are distributed on the nine R_i of the four basic polytypes (Table 4): 26 I_j are obtained by multiplying these seven I_j : they form a group, as shown in the multiplication table (Table 5).

6. Derivation of twin diffraction patterns

The reciprocal-lattice rows of a twin derive from the overlap of single-crystal rows (R_i) and are called 'composite rows'

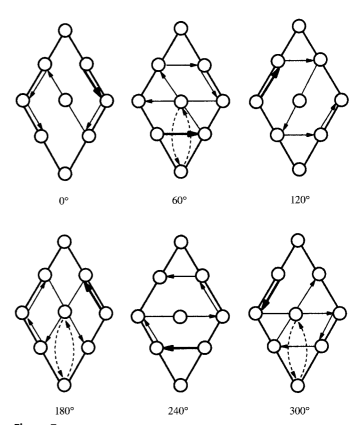


Figure 7 Distribution of the six X rows along the edges of the minimal rhombus as a function of the rotation of the minimal rhombus itself. The arrows indicate the reading sequence corresponding to the rotation angle given below the rhombus. The two D rows are unchanged for $\pm 120^{\circ}$ but exchanged for ± 60 or 180° rotations.

Table 4 R_i for the four basic polytypes (*cf.* Figs. 5 and 6).

For the definition of I_i , see Table 3.

				1 <i>M</i>		$2M_1$		$2M_2$		3T	
Single-crystal row	Type	h (mod 3)	k (mod 3)	l (mod 6)	I_j	l (mod 6)	I_j	l (mod 6)	I_j	l (mod 6)	I_j
R_1	S	0	0	0	11	0	11	0	11	0	11
R_2	D_1	1	1	2	12	4	13	0,3	2_{3}	2	12
R_3	D_2	2	2	4	13	2	12	0,3	2_{3}	4	13
R_4	X_1	1	1	2	1_2	1,4	2_{4}	1,4	2_{4}	0,2,4	31
R_5	X_2	2	2	4	13	2,5	25	2,5	25	0,2,4	3_1
R_6	X_3	0	1	0	1_1	0,3	23	1,4	2_{4}	0,2,4	31
R_7	X_4	1	2	2	1_2	1,4	2_{4}	2,5	25	0,2,4	31
R_8	X_5	2	1	4	1_3	2,5	25	1,4	2_{4}	0,2,4	31
R_9	X_6	0	2	0	1_1	0,3	2_{3}	2,5	25	0,2,4	31

 Table 5

 Multiplication table for composite rows.

An I_i in the body of the table is the product of the corresponding two I_i in the first rows (I_i^A) and in the first column (I_i^B) . For the definition of I_i see Table 3.

$I_j^A \ I_j^B$	61	53	52	51	4 ₆	4 ₅	4 ₄	4 ₃	42	41	3 ₇	3 ₆	35	3 ₄	33	32	31	26	25	2 ₄	2 ₃	22	21	1 ₃	12	11
11	61	53	52	51	52	45	44	43	42	41	45	41	35	34	33	32	31	31	34	32	23	22	21	22	21	11
1_2	6_1	53	52	5_1	46	45	4_{4}	4 ₃	5_1	4_1	37	3_{6}	4_{4}	3_{4}	3_3	4_1	3_1	2_{6}	2_{5}	3_{6}	3_3	3_1	2_1	2_{6}	1_2	
1_{3}	6_1	53	52	5_1	46	45	4_{4}	5_{3}	42	4_1	37	3_{6}	3_{5}	45	4_{4}	3_2	3_1	2_{6}	37	2_{4}	35	2_2	3_1	1_{3}		
2_1	6_1	53	5_2	5_1	52	45	4_{4}	4 ₃	5_1	4_1	45	4_1	4_{4}	3_{4}	3_3	4_1	3_1	3_1	3_{4}	4_1	33	3_1	2_1			
2_{2}	61	53	52	5_{1}	52	45	44	53	42	41	45	41	35	45	44	32	31	31	45	32	35	2_2				
23	61	53	6_1	51	61	5 ₃ 5 ₂	4_{4}	43	42	51	5 ₃ 4 ₆	51	$\frac{3}{4}$	4 ₃ 5 ₂	3 ₃ 5 ₁	42	44	44	4 ₃ 4 ₆	42	2_{3}					
24	61	6_1	52	51	46	5 ₂	51	61	42	$\frac{4_1}{5_2}$	4 ₆	36	4 ₂	52	5 ₁	32	4 ₁	36	4 ₆	2_{4}						
25	61	5 ₃ 5 ₃	5 ₂ 5 ₂	6_1	46	45	53	4 ₃ 5 ₃ 5 ₃	6_1	52	$\frac{3}{3}$	46	53	34	43	52	45	37	2_{5}							
26	61	5_3	52	5_{1}	46	45	44	5_{3}	5_{1}	41	37	36	44	45	44	41	31	2_{6}								
31	61	53	5 ₂ 5 ₂	5_{1}	5 ₂ 5 ₂	$4_5 \\ 5_2$	44	53	51	$4_1 \\ 4_1$	4 ₅ 5 ₂	41	44	45	4_{4}	41	3_1									
32	61	61	52	5_{1}	52	52	51	61	42	41	52	41	42	52	51	3_2										
33	61	53	6_{1}	51	6_1	53	4_{4}	43	51	51	53	5_{1}	44	43	3_3											
34	61	53	52	6_1	52	45	53	43	61	52	45	52	53	3_{4}												
35	61	53	6_1	5_{1}	61	53	44	53	42	51	53	51	3_{5}													
36	61	6_1	5_2	51	46	52	51	6_1	51	41	46	3_{6}														
37	61	53	5_2	6_1	46	45	53	53	6_1	52	37															
41	61	6_1	52	5_{1}	52	52	5_{1}	6_1	51	41																
42	61	6_1	61	51	61	6_1	51	61	4 ₂																	
43	61	5 ₃ 5 ₃	61	61	61	5 ₃ 5 ₃	53	4 ₃																		
44	61	23	61	51	61	33	4_{4}																			
45	61	53	52	61	52	4 ₅																				
46	61	61	52	61	4 ₆																					
51	61	61	61	5_1																						
52	61	61	52																							
53	61	53																								
61	6_1																									

(Sadanaga & Takéuchi, 1961). The analysis of R_i represents the starting point in the derivation of twin patterns.

The $n \times 60^{\circ}$ rotations about \mathbf{c}^* overlap only R_i belonging to the same type (S, D or X), but specific nodes may overlap or not depending upon which I_j corresponds to the rotationally related R_i . Each of the R_i is rotationally related to five other ones and along each of them a peculiar sequence of l_T indices is obtained, which has been called 'rotational sequence' (Nespolo, Takeda, Ferraris & Kogure, 1997). Each R_i generates one rotational sequence, which is shortened to $RS_i^P(n)$, where the superscript P indicates the polytype $(P = 1M, 2M_1, 2M_2, 3T)$; i is the same index defining R_i ; n points to each of the six digits of the RS. RS_1^P always corresponds to '000000' for all polytypes. The nth values of RS_1^P correspond to the l_T indices of the nodes on the row which is related to R_i by

 $(n-1) \times 60^{\circ}$ clockwise rotation. The two RS₁^P corresponding to *D*-type rows (R_{2-3}) on the one hand and the six RS₁^P corresponding to *X*-type rows (R_{4-9}) on the other can be transformed into each other by cyclic permutations. Since the orientations of the single-crystal lattices and of the twin lattice are fixed and determined by Z_T , also the starting point of each RS₁^P is fixed, and the nine RS₁^P are independent. The node features of the composite rows can be obtained from the corresponding RS₁^P by considering their relation with the Z_T symbols. A twin of *N* individuals $(2 \le N \le 6)$ is identified by N Z_T symbols. The I_T index of the qth node on the ith row from the jth individual is given by:

$$[l_T(i,j)]_q = [RS_i^P(n)]_q, \quad n = [(Z_T)_j + 4] \pmod{6}.$$
 (7)

Composite row	Type	$h \pmod{3}$	k (mod 3)	RS^{1M}	RS^{2M_1}	RS^{2M_2}	RS^{3T}
C_1	S	0	0	000000	000000	000000	000000
C_2	D_1	1	0	242424	424242	000000/333333	242424
$\tilde{C_3}$	D_2	2	0	424242	242424	000000/333333	424242
C_4	X_1	1	1	204402	102201/435544	121212/454545	000000/222222/44444
C_5	X_2	2	2	402204	201102/534435	212121/545454	000000/222222/44444
C_6	X_3	0	1	022044	011022/344355	121212/454545	000000/222222/444444
C_7	X_4	1	2	220440	110220/443553	212121/545454	000000/222222/444444
C_8	X_5	2	1	440220	220110/553443	121212/454545	000000/222222/444444
C_9	X_6	0	2	044022	022011/355344	212121/545454	000000/222222/444444

Table 6 Composite rows (C_i) and rotational sequences (RS_i^P) for the four basic polytypes (see §6 and Figs. 5 and 6).

The node features of composite rows are completely defined by the nine RS_1^P and Z_T symbols; therefore, there are only nine independent composite rows, for which the symbol C_i is adopted (Table 6). R_i and C_i share the same row features and thus the description of the reciprocal lattice in terms of the tessellation rhombus and of the minimal rhombus is the same for both the single-crystal lattice and the twin lattice.

Because of the metric relations (§1.2), the I_j of both C_i and R_i of the same type and belonging to the same central plane are related by:

$$[l_T(D_i)]_q = [6 - l_T(D_{3-i})]_q$$

$$[l_T(X_i)]_q = \{6 - l_T[X_{(9-i)(\text{mod } 6)}]\}_q.$$
(8)

With the I_j of one D-type C_i/R_i and three X-type C_i/R_i , the I_j of the remaining four C_i/R_i can be calculated. There are thus five truly geometrically independent C_i/R_i (one S-type, one D-type and three X-type), but nine translationally independent C_i/R_i .

The distribution of I_j on the C_i of a minimal rhombus is the information necessary to derive and identify the diffraction patterns of mica twins.

6.1. 1M polytype

The minimal rhombus analysis of 1M twins given in Nespolo, Takeda, Ferraris & Kogure (1997) is revised here in terms of Z_T symbols.

The c_1^* repeat coincides with the polytype period and along each R_i there is only one node, which obeys the relation $l_T = 2h \pmod{6}$. D-type R_i are either $1_2 \pmod{1}$ or $1_3 \pmod{2}$ and the RS_{2-3}^{1M} are 242424 and 424242. The $n \times 60^\circ$ rotations about \mathbf{c}^* produce the overlap of all the reciprocal-lattice nodes belonging to D-type R_i when n is even, but to their separation when n is odd. X-type R_i include all the three 1_1 , 1_2 and 1_3 and the RS_{4-9}^{1M} are the six cyclic permutations of 220440. On the basis of the relation between C_i and RS_i^{1M} (Table 6) and of the multiplication table (Table 5), seven different C_i appear in the twin lattice. Up to 2 reflections can appear on D-type C_i ($I_j = 1_2, 1_3$ or 2_6 ; I_T is never 0), whereas up to three reflections (all the seven possible I_j) can appear on X-type C_i . I_j define nine independent 1M twin patterns (Fig. 8).

Nespolo, Takeda, Ferraris & Kogure (1997) expressed the relative rotations between the first and the (j+1)th individual as a sequence of $n_j \pmod{60^\circ}$ integers. Those symbols are not space-fixed and in some respects are midway between the

Z symbols and the rotational RTW symbols (Ross *et al.*, 1966). The correspondence with Z_T symbols is simply $Z_T(1)=3$; $Z_T(j+1)=3+n_j$. Consideration of the relative rotations, instead of absolute orientations, revealed ten patterns, of which however one (corresponding to $Z_T=341$) turns out to be not independent when the lattice is reflected across (010) [symbol transformation $Z_T=3IJ\dots P\to 3(6-I)(6-J)\dots (6-P)$ (Table 2)].

As seen above, when three equally spaced reflections in the c_1^* repeat appear along nonfamily rows, in principle the

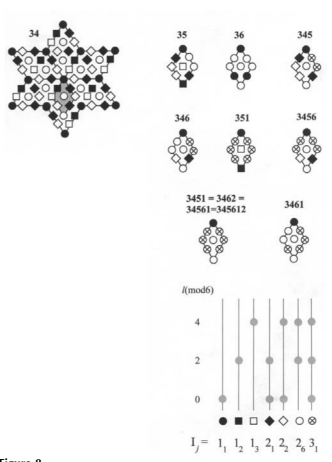


Figure 8 The nine independent patterns of 1M twins as expressed through the corresponding minimal rhombi. For the $Z_T = 34$ twin, the complete star polygon is given, with the minimal rhombus in it shaded. Inset on the lower right: $l \pmod 6$ indices of the nodes on reciprocal-lattice composite rows.

diffraction pattern may correspond either to a 1M twin (apparent polytypism) or to a three-layer polytype (real polytypism). The distinction is easily obtained by applying the geometrical criteria given in Nespolo (1999a). However, 1M twins with $Z_T = 351$ and the 3T polytype (Z = 351 or 315) cannot be distinguished geometrically (see also Nespolo & Kogure, 1998). In the case of dioctahedral and Li–Fe trioctahedral micas, this ambiguity is removed when weak reflections appear along family rows (Rieder, 1968, 1970) (see §8).

6.2. $2M_1$ polytype

Since the parity of layers is even (Z=24) for a $2M_1$ polytype but odd for a 1M polytype, the threefold family structure has opposite settings (reverse/obverse) and the corresponding family rows have different reflection conditions, namely $k=0 \pmod{3},\ l_T=2h \pmod{6}$ for 1M, but $k=0 \pmod{3},\ l_T=4h \pmod{6}$ for $2M_1$ (Nespolo, 1999a). One reflection appears in the c_1^* repeat along family R_i , but two along nonfamily R_i , with the following presence criteria:

$$\begin{cases} k \neq 0 \pmod{3} & \begin{cases} l_{T1} = h \pmod{6} \\ l_{T2} = (h+3) \pmod{6} \\ l_{T} = (h+k) \pmod{6}. \end{cases}$$
(9)

The node features of R_i are compared with those of the other basic polytypes in Table 4. D-type R_i are either 1_2 or 1_3 , as in the 1M case but, because of the opposite parity of the layers in the two polytypes, the positions of the two I_j and the two $RS_{2-3}^{2M_1}$ are inverted. X-type R_i have all the three possible pairs of values of I_T (mod 6): 0 and 3 ($I_j = 2_3$), 1 and 4 ($I_j = 2_4$), 2 and 5 ($I_j = 2_5$). For the X-type R_i , the sequence of $n \times 60^\circ$ rotations corresponds to a double sequence of I_T values: 011022/344355 or cyclic permutations, producing six independent double $RS_i^{2M_1}$ (Table 6). As in the case of the 1M polytype, the twelve composite twins produce nine different patterns, none of which can be mistaken for that of a 1M twin (Fig. 9).

6.3. $2M_2$ polytype

Being a *Class b* polytype, $2M_2$ has a markedly pseudorhombohedral lattice and two of the five pairs of twin laws, namely those corresponding to $\pm 120^{\circ}$ rotation about \mathbf{c}^* , correspond to pseudo-merohedry but the remaining three to reticular pseudo-merohedry. All the six $n \times 60^{\circ}$ rotations belong to the point group of the family structure (subfamily *B*), and thus the family sublattice of the individuals is always overlapped. The presence criteria in the twin setting are:

$$\begin{cases} k \neq 0 \pmod{3} \begin{cases} l_{T1} = k \pmod{6} \\ l_{T2} = (k+3) \pmod{6} \end{cases} \\ k = 0 \pmod{3} \begin{cases} h = 0 \pmod{3} : l_{T} = 0 \pmod{6} \\ h \neq 0 \pmod{3} \end{cases} \begin{cases} l_{T1} = 0 \pmod{6} \\ l_{T2} = 3 \pmod{6}. \end{cases}$$
(10)

 $RS_2^{2M_2}$ and $RS_3^{2M_2}$ both correspond to the double sequence 000000/333333, whereas the $RS_{4-9}^{2M_2}$ correspond to the cyclic permutations of the double sequence 121212/454545. There are only two kinds of patterns for $2M_2$ twins. Twinning by

pseudo-merohedry gives a pattern geometrically indistinguishable from that of the single crystal $(Z_T = U)$. The other pattern corresponds to twinning by reticular pseudo-merohedry $(Z_T = UE)$ and differs from the single-crystal pattern in the six X-type C_i , which are all $I_j = 4_6$ (see Tables 3 and 5). Neither of the two can be mistaken for any of the 1M or $2M_1$ polytypes or twins.

6.4. 3*T* polytype

The 3T polytype is an orthogonal subfamily A polytype, with odd parity of the layers (the same as 1M). D-type R_i and RS_i^{3T} are exactly the same as those of the 1M polytype, but the X-type R_i correspond to $I_j = 3_1$ and there is thus only one triple sequence of X-type RS^{3T} : 00000/222222/444444. The six orientations of the minimal rhombus are divided into two types, differing for the I_j corresponding to D_1 and D_2 . The $2n \times 60^\circ$ rotations belong to the point group of the individual and reproduce the same rhombus. On the other hand, $(2n+1) \times 60^\circ$ rotations do not belong to the point group of the individual and thus they exchange the two independent rhombi.

Twinning by complete merohedry ($Z_T = UU^{\bullet}$) by definition produces a diffraction pattern with the same geometrical appearance as the single crystal, which in its turn may be geometrically identical to the pattern of 1M twinned as $Z_T = 351$. On the contrary, in the case of twinning by selective merohedry ($Z_T = UE$, UE^{\bullet} , $UU^{\bullet}E$, $UU^{\bullet}E^{\bullet}$, $UU^{\bullet}EE^{\bullet}$), the two D-type C_i correspond to 2_6 . This is the same geometrical appearance of 1M twinned as $Z_T = 3451$. The distinction between 1M twins and the 3T polytype (twinned or

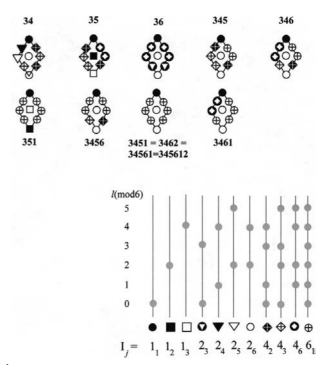


Figure 9 The nine independent patterns of $2M_1$ twins as expressed through the corresponding minimal rhombi. Inset on the lower right: $l \pmod{6}$ indices of the nodes on reciprocal-lattice composite rows.

untwinned) has to be reached by very careful examination of the violation of the additional reflection conditions (see §8).

7. Derivation of allotwin diffraction patterns

The oriented association of two (or more) individuals differing only in the polytypic character is called *allotwinning* [from the Greek $\alpha\lambda\lambda o\varsigma$, 'different', with reference to the individuals (Nespolo, Kogure & Ferraris, 1999)]. Since the building layer(s) are in principle identical, at least two of the three parameters – those in the plane of the layer – are identical. As in the case of triperiodic epitaxy (Royer, 1928, 1954), a three-dimensional common sublattice exists (allotwin lattice), whose cell is based on the two common parameters in the plane of the layer, the third one being the shortest parameter, outside the plane of the layer, common to the cells of both individuals.

The allotwin laws are the twin laws for each of the individuals building the allotwin, to which the symmetry operations of the crystal(s) point group(s) have to be added. By indicating the first individual with a superscript and the second one with a subscript, the allotwin $Z_T = {}^3_3$ has to be considered too, whereas the $Z_T = 33$ twin simply corresponds to a parallel intergrowth. Therefore, the number of possible laws becomes higher and depends upon the number of different polytypes undergoing allotwinning.

Since the geometrical appearance of the diffraction pattern of the 3T polytype and of its twins is the same as (ideal) 1M twinned as $Z_T = 351$ or 3451 (see §6.4), the contribution from 3T does not produce any independent pattern: it is not taken into account in the following systematic analysis.

The three basic monoclinic polytypes can produce three binary (two-portion) allotwins $(1M-2M_1; 1M-2M_2; 2M_1-2M_2)$ and one ternary (three-portion) allotwin $(1M-2M_1-2M_2)$. Binary (AB) and ternary (ABC) allotwins are henceforth indicated by ${}^A{}_B$ and ${}^A{}_B{}_C$, respectively, where A, B and C represent the Z_T symbols for each portion of the allotwin.

All these can be described on the following basis:

- (i) The allotwin can be thought of as built up by two (binary allotwin) or three (ternary allotwin) portions (A, B, C), each consisting only of individuals of the same polytype, which in their turn can be twinned; 1M is taken as the first portion (A) of the allotwin; when 1M is not involved (binary allotwin $2M_1-2M_2$), the portion A is $2M_1$.
- (ii) Since the individuals building the twin or allotwin are related by point-group operations, the A–B–C sequence has no influence on the composite lattice and the two or three portions can be described as juxtaposed and non-mixed; for example, $Z_T = {}^{3}_{4} {}^{5}_{6}$ is equivalent to $Z_T = {}^{35}_{46}$.
- (iii) Within each single portion (A, B, C), the restrictions on the possible orientations derived for the twins are retained, but they are not applicable when comparing individuals belonging to different portions.
- (iv) The first individual of the first portion (A) is fixed in the orientation $Z_T = 3$, but this restriction is not applicable for the first individual of the other portions. Therefore, the number of possible orientations for B and C portions has to be multiplied by the number of independent orientations of the minimal

rhombus, as determined by the limiting symmetry (§3), namely six for $2M_1$ and two for $2M_2$.

The minimal rhombi of the allotwins are calculated as combinations of the minimal rhombi of each portion, but the number of minimal rhombi to be considered depends upon the limiting symmetry. The minimal rhombi of two twins of 1M which are equivalent through an $n \times 60^{\circ}$ rotation about the normal to (001) can produce two independent minimal rhombi when combined with a minimal rhombus of $2M_1$. Therefore, in the derivation of the reciprocal lattice of $1M-2M_1$ allotwin, for both polytypes the minimal rhombi of all the 31 twins in Table 2 have to be considered. To these, the minimal rhombus corresponding to the single crystal has to be added. Moreover, if the minimal rhombi of 1M (first individual in orientation $Z_T = 3$) are kept fixed, the six independent orientations of each of the 32 minimal rhombi of $2M_1$ have to be considered. For the $2M_2$ polytype, there are only two independent orientations of the individual w.r.l. $(Z_T = U \text{ or } Z_T = E)$ and only one for the twin reciprocal lattice $(Z_T = UE)$. In deriving the reciprocal lattice of $1M-2M_2$ or $2M_1-2M_2$ allotwins, for Class a polytypes, only the minimal rhombus of the single-crystal and the minimal rhombi of the 23 twins related by $(2n + 1) \times 60^{\circ}$ rotations have to be combined with the three (U, E, UE)minimal rhombi of $2M_2$. The remaining eight minimal rhombi of Class a polytypes are related to some of the other 23 by $2n \times 60^{\circ}$ rotations, which are symmetry operations for the minimal rhombi of $2M_2$ and cannot produce any further independent allotwin minimal rhombus. Finally, for the ternary allotwins $1M-2M_1-2M_2$, the independent minimal rhombi of the binary allotwin $1M-2M_1$, and those related by $(2n+1) \times 60^{\circ}$ rotations, have to be combined with the three minimal rhombi of $2M_2$.

For each combination, the composite minimal rhombus obtained in this way, rotated by $n \times 60^{\circ}$ ($0 \le n \le 5$) and – for each of these rotations – reflected across (010), is compared with the ones calculated for the previous combinations and, if equivalent, is discarded. Because of their relatively high number, the resulting minimal rhombi for allotwins are shown in table format rather than in figure format. The number of independent minimal rhombi is 46 ($1M-2M_1$: Table 7), 13 ($1M-2M_2$: Table 8), 15 ($2M_1-2M_2$: Table 9) and 36 ($1M-2M_1-2M_2$: Table 10).

8. Effect of the violation of the additional reflection conditions

The above derivation of the composite reciprocal lattice of twins and allotwins has been obtained within the trigonal model. In the real structure, other distortions besides the trigonal rotation of the tetrahedra appear, which in principle result in the violation of the non-space-group absences of the family reflections and in additional weak reflections in the family rows. These are visible in dioctahedral and Li–Fe micas (Rieder, 1968, 1970) and may help to distinguish some patterns that are equivalent within the trigonal model. The D-type R_i of subfamily A polytypes are either 1_2 or 1_3 and the D-type C_i can be either 1_2 - 1_3 or 2_6 . No weak reflection appears

Table 7 Distribution of I_j on the nine C_i for the 46 distinguishable patterns of $1M-2M_1$ binary allotwins.

For the definition of I_j and C_i , see Table 3 and §6, respectively (cf. also Figs. 5 and 6).

Allotwin	D_1	D_2	X_1	X_2	X_3	X_4	X_5	X_6
3 3 4 3 5 3 6 34 3	26	26	36	37	23	36	37	23
3	12	13	33	35	32	36	3,	34
3	2_{6}^{2}	26	2.5	24	32	33	35	34
3	1_{2}^{-6}	1_3	2 ₅ 2 ₅	$\frac{-4}{2_4}$	$\frac{1}{2}$	25	24	23
34	26	26	4 ₁	45	33	36	37	35
3.4	$\frac{26}{26}$	$\frac{26}{26}$	34	32	4 ₁	3_{3}	3_{5}	45
5 35 3 35		26	2	2	2	<i>3</i> 3		45
35	26	26	36	37	33	41	45	35
4	1_2	13	44	44	41	41	45	45
1 26	26	26	4_{4}	4_{4}	34	34	32	32
36	26	26	4_4	4_4	32	36	3 ₇	34
34	2_{6}	2_{6}	51	53	42	36	37	43
3 45	26	$\frac{2_{6}}{2_{6}}$	43	42	$\frac{3_{2}}{4_{2}}$	51	5 ₃	34
3 56	26	26	25	2_{4}	42	43	4_{2}	43
3	26	2_{6}	46	46	42	51	53	43
3	12	13	43	42	42	46	46	43
3 51	2_{6}^{2}	26	43	4_{2}^{2}	5_{2}^{2}	43	42	52
3	26	26	4 ₆	46	2_3	46	46	2.
36	$\frac{26}{26}$	26	33	35	5_2	46	46	5_{2} 2_{3} 5_{2} 5_{2} 3_{5}
3 ⁴¹	∠ ₆	<u>∠</u> 6	<i>5</i> 3	55 5	<i>5</i> 2	5 ₁	4 6	<i>5</i> 2
42 345	12	13	51	53	52	J_1	53	32
345	26	26	41	45	33	41	45	35
4	26	26	44	44	41	41	45	45
4	26	26	4_4	4_4	41	36	37	45
35	26	2 ₆ 2 ₆ 2 ₆ 2 ₆	41	45	34	4_4	4_4	3 ₂ 5 ₃
34 34	26	2_{6}	51	53	51	36	37	53
34 34 45 34 35	2 ₆ 2 ₆	26	4 ₃ 5 ₂	42	4 ₁ 5 ₁	5 ₁ 5 ₁	53	45
34 35	26	26	52	52	51	51	53	53
57	26	26	$\overline{5}_{2}$	$\overline{5}_{2}$	52	33	35	52
	26	26	51	53	51	41	45	53
35 35 35 35 51	26	26	46	46	51	51	5 ₃	53
35	$\frac{26}{26}$	26	53	51	5_2	4 ₃	4 ₂	52
51 35	$\frac{26}{26}$	26	4_4	4_4	5_2 5_2	5 ₂	$\frac{4}{5}$	5_2
35 41 36 45	26	26	44	44	32	52	52	32
45 36	26	26	53	51	32	51	53	34
41	26	26	44	44	5 ₂ 4 ₂	46	46	52
345	26	26	61	61	42	5 ₁ 4 ₆	53	43
346	26	2_{6}	6_1	6_1	42	4 ₆	46	4 ₃
341	26	2 ₆ 2 ₆	51	6 ₁ 5 ₃	6_1	46	46	6_1
3 342	26	2 ₆ 2 ₆	51	53	6_1	$ \begin{array}{c} 5_1 \\ 6_1 \end{array} $	53	6_1
3 351	26	26	6_1	6_1	6_1	6_1	6_1	6_1
3 456	26	26	43	42	42	61	61	43
3	26	26	43	42	$\overline{5}_{2}^{2}$	61	61	52
3	26	26	61	6_1	52	51	53	5 ₂ 5 ₂
3	1_{2}^{26}	1_3	6_1	6_1	6_1	6_1	61	6_1
3	$\frac{1}{2}$	2.	6_1	6_1	4 ₂	6.	6.	4 ₃
3456 3	∠ ₆	2 ₆ 2 ₆ 2 ₆	6	6	-1 2	6 ₁ 5 ₁	6 ₁ 5 ₃	+ 3
3452 3	2 ₆ 2 ₆	∠ ₆	61	61	6_1	\mathcal{I}_1	J ₃	61
36 41 3 43 43 43 43 45 4 4512 4512	26	26	61	61	6_1	46	46	6_1
4512	26	26	6_{1}	61	52	6_{1}	6_1	52

in the case of the 1M polytype. For the $2M_1$ polytype, one weak reflection is expected at $l_T=5(D_2)$ or $l_T=4(D_1)$, transforming 1_2 , 1_3 and 2_6 into 2_5 , 2_4 and 4_6 , respectively. For the 3T polytype, two weak reflections are expected at $l_T=0$, $2(D_2)$ or $l_T=0$, $4(D_1)$, transforming both 1_2 and 1_3 into 3_1 . Composite patterns that are equivalent within the trigonal model can be in principle distinguished as follows.

- (i) 1M twinned as $Z_T = 351$ vs 3T untwinned or twinned by complete merohedry ($Z_T = UU^{\bullet}$). Two weak reflections are expected on both the D-type R_i and C_i when the 3T polytype is involved.
- (ii) 1M twinned as $Z_T = 3451 \text{ vs } 3T$ twinned by selective merohedry (involving both $Z_T = U$ and $Z_T = E$). One weak

Table 8 Distribution of I_j on the nine C_i for the 16 distinguishable patterns of $1M-2M_2$ binary allotwins.

For the definition of I_j and C_i , see Table 3 and §6, respectively (cf. also Figs. 5 and 6).

Allotwin	D_1	D_2	X_1	X_2	X_3	X_4	X_5	X_6
3 <i>U</i> 34	33	3 ₅	36	37	32	25	24	34
34 U	44	4_4	4_1	45	4_1	25	2_{4}	45
35 U	33	35	36	37	4_1	34	3_2	45
36 U	4_{4}	4_4	36	37	3_2	37	36	3_{4}
U	4_{4}	4_4	36	37	3_2	3_{4}	3_2	3_{4}
345 U	4_{4}	4_4	4_1	45	4_{1}	3_{4}	3_2	45
346 U	4_{4}	4_4	4_1	45	4_{1}	37	36	45
351 U	33	35	4_1	45	4_{1}	45	4_1	45
U 36 U 36 U 32 U 345 U 351 U 351 U 351 U 352 U 3456	44	4_{4}	36	37	4_{1}	34	3_2	45
3456 <i>U</i>	4_4	4_4	4_1	45	4_1	45	4_1	45
3 UE 34	33	35	46	46	52	46	46	52
34 UE 35	44	4_4	52	52	52	46	46	52
35 UE 36	33	35	46	46	52	52	52	52
36 UE	44	4_4	46	46	5_{2}^{-}	46	46	52
UE 345 UE 351	4_4	4_4	52	52	52	52	52	52
351 UE	3 ₃	35	52	52	$\overline{5_2}$	52	52	5_{2}^{-}

reflection corresponding to $l_T = 0$ is expected on both the D-type C_i when the 3T polytype is involved.

- (iii) $2M_1$ –3T allotwins vs 1M– $2M_1$ allotwins. In both cases, on the D rows one weak reflection, corresponding either to $l_T=1$ or to $l_T=5$, is expected from the $2M_1$ individual contribution. If 3T is involved, another weak reflection, corresponding to $l_T=0$, is expected, whereas if 1M is involved it does not appear. It is never possible to distinguish between a $2M_1$ –3T binary allotwin and a 1M– $2M_1$ –3T ternary allotwin.
- (iv) $2M_2$ –T allotwin vs 1M– $2M_2$ allotwin. If the 3T contribution is from a single crystal ($Z_T = U$ or $Z_T = E$) or from a twin by complete merohedry ($Z_T = UU^{\bullet}$), the two D-type C_i rows are 3_3 and 3_5 in the trigonal model, and the distinction is possible on the basis of the appearance of a weak reflection alternatively corresponding to $l_T = 4$ and $l_T = 2$. However, if the 3T contribution comes from a twin by selective merohedry (involving thus both $Z_T = U$ and $Z_T = E$), no distinction is possible, since all the three reflections corresponding to $l_T = 0$, 2 and 4 are present on the two D rows. The ternary 1M– $2M_2$ –3T allotwin cannot be distinguished, since the 1M features are hidden by the 3T ones.

9. Discussion

In this paper, it has been shown that the geometrical analysis of the reciprocal lattice permits one to derive and identify the relative rotations in twins or allotwins of mica polytypes in the case of twinning or allotwinning by reticular pseudo-merohedry and by selective merohedry. The analysis has been devoted to the four basic polytypes, but can be easily extended to other polytypes, by redefining the period along \mathbf{c}^* of the 'twin setting'. When dealing with a single polytype, the twin setting is best defined to coincide with the C_1 setting, which is based on the cell of the twin lattice. The adoption of a twin

Table 9 Distribution of I_j on the nine C_i for the 13 distinguishable patterns of $2M_1-2M_2$ binary allotwins.

For the definition of I_j and C_i , see Table 3 and $\S6$, respectively (cf. also Figs. 5 and 6).

Allotwin	D_1	D_2	X_1	X_2	X_3	X_4	X_5	X_6
3	35	33	24	25	42	46	46	43
U 34	4_4	4_{4}	42	43	$\frac{1}{4_{2}}$	46	46	43
$\begin{array}{c} & & & & & & \\ & & & & & \\ & & & & & \\ & & & & \\ & & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$	35	33	46	46	4_{2}^{2}	6_1	6_1	43
36 U	4_{4}	4_{4}	46	46	42	46	46	43
U	4_4	4_{4}	2_{4}	25	6_1	6_1	6_1	6_1
345 U	4_4	4_{4}	6_1	6_1	42	6_1	6_1	43
U	4_4	4_{4}	6_1	6_1	42	46	46	43
351 252 U	35	33	6_1	6_1	6_1	6_1	6_1	6_1
352 U	4_{4}	4_{4}	46	46	6_1	6_1	6_1	6_1
3451 U	4_{4}	4_{4}	6_1	6_1	6_1	6_1	6_1	6_1
UE	3 ₅	3_3	4 ₆	4 ₆	6_1	4_{6}	4 ₆	6_1
UE 35 UE 36	3 ₅	3_3	4 ₆	4 ₆	6_1	6_1	6_1	6_1
UE	4_4	4_4	4 ₆	4 ₆	6_1	4_{6}	4 ₆	6_1

setting with a period along \mathbf{c}^* different from that of the C_1 setting is useful to compare the geometry of the reciprocal lattice of polytypes with different period. In such a case, the twin setting is best defined to have the shortest period along \mathbf{c}^* in the C_1 setting among all the polytypes considered.

In the twin or allotwin lattices, nodes on planes parallel to (001) are either completely overlapped (twinning by merohedry or pseudo-merohedry, twin index 1) or are overlapped on one plane out of three (twinning by reticular pseudomerohedry, twin index 3). Other oriented crystal associations are known, in which complete restoration of the lattice nodes is not achieved in any plane. These oriented crystal associations are called 'plesiotwins' (Nespolo, Ferraris, Takeda & Takéuchi, 1999) and are built on larger hexagonal cells based on multiple two-dimensional meshes in the (001) plane. The plesiotwin elements are twofold axes in the plane of the layer with higher indices; the corresponding rotations about the normal to (001) express the relative rotations between individuals in a plesiotwin and are operations belonging to noncrystallographic point groups, even neglecting the angular deviation from orthohexagonality ε . On the other contrary, the relative rotations between individuals in both twins and allotwins belong to crystallographic point groups (neglecting ε). The derivation of the plesiotwin laws can be performed following the same procedure shown above for the twin laws. However, the row features no longer correspond to the regular tessellation {3, 6}, but to a compound tessellation $\{3, 6\}[n\{3, 6\}]$, where n is the multiplicity of the larger mesh in the (001) plane (Takeda & Donnay, 1965). Only a subset of the individual reciprocal-lattice rows, R_i , namely one out of n, overlaps in composite rows C_i . The diffraction pattern from the (hk0) plane thus shows a complex superstructure, corresponding to the overlap of two (eventually more) lattices rotated about the normal (Rang, 1958; Zvyagin & Gorshkov, 1966; Takéuchi et al., 1972; Sunagawa & Tomura, 1976), which can be interpreted by the geometrical theories of the coincidence-site lattice (e.g. Santoro & Mighell, 1973), whereas, in the case of twins and allotwins, the diffraction pattern from the

Table 10 Distribution of I_j on the nine C_i for the 36 distinguishable patterns of $1M-2M_1-2M_2$ ternary allotwins.

For the definition of I_j and C_i , see Table 3 and §6, respectively (cf. also Figs. 5 and 6).

Allotwin	D_1	D_2	X_1	X_2	X_3	X_4	X_5	X_6
$^{3}3_{U}$	4_4	4_4	36	37	42	46	46	43
34	33	35	51	53	32	46	46	3 ₄ 5 ₂ 5 ₂
4 _E	3_3	35	43	42	5_{2}	36	37	5_2
² 411F	33	35	6_1	6_1	52	46	46	5_2
5_{II}	4_{4}	4_{4}	46	46	3_2	43	42	3_{4}
³ 5 _E	4_4	4_4	25	2_{4}	52	5_{1}	53	5_2
35	4_4	4_{4}	46	46	52	6_1	6_1	52
37377	4_4	4_4	4_1	45	5_{1}	46	46	53
J-73	4_4	4_{4}	52	52	4 ₃	36	37	42
$^{34}5_{U}$	4_{4}	4_4	52	52	4_1	43	42	45
J45	4_{4}	4_4	3_{4}	32	52	51	53	5_2
	4_4	4_4	52	52	52	6_1	6_1	5 ₂ 5 ₃
$^{35}_{UE}$ $^{35}_{3U}$	4_4	4_4	36	37	51	52	52	53
$^{35}_{3E}$	4_4	4_4	46	46	4 ₃	4_1	45	4_{2}
JJ4	33	3 ₅	51	53	4 ₁	52	52	45
	33	35	6_1	6_1	52	52	5_{2}	52
411	4_4	4_{4}	5_1	53	32	46	46	3_{4}
3411	4_4	4_4	5_1	53	42	46	4_{6}	43
334 r	4_4	4_4	6_1	6_1	6_1	36	37	6_1
345	4_4	4_4	6_1	6_1	3_2	6_1	6_1	3_{4}
³ 45 -	4_4	4_4	43	42	52	5_1	53	$ \begin{array}{c} 5_2 \\ 5_2 \end{array} $
345	4_4	4_4	6_1	6_1	52	6_1	6_1	52
35 r	4_4	4_4	46	46	6_1	51	53	6_1
³ 51,,	4_4	4_4	6_1	6_1	52	43	42	52
³ 41,,	4_4	44	51	53	52	46	46	52
³ 41 _E	4_{4}	4_{4}	43	42	52	46	46	52
³ 42 ₁₁	3 ₃	35	51	53	52	6_1	6_1	52
2/12	33	35	6_1	6_1	52	6_1	6_1	52
³⁴⁵ 3 _U	4_{4}	4_{4}	4_1	45	5_{1}	52	52	53
3417	4_4	4_{4}	5_1	53	5_{1}	46	46	53
- 45,,,	4_4	4_4	6_1	6_1	4_1	6_1	6_1	45
3435,,	4_4	4_4	52	52	5_{1}	6_1	6_1	53
J+57	4_{4}	4_{4}	52	52	52	4 ₃	42	52
	4_{4}	4_4	52	52	52	51	53	5_2
3411	4_4	4_{4}	51	53	51	52	52	53
$^{3}345_{E}$	4_4	4_4	6_1	6_1	6_1	51	53	6_1

(hk0) plane shows the same pseudo-hexagonal symmetry of the single crystal. The coincidence-site lattice corresponding to the plesiotwin lattice is obtained by applying rotations that connect two X rows on the same orbit but belonging to two different sets (Fig. 6, thin dashed lines). The minimal rhombus approach applies to both twins and allotwins, but not to plesiotwins.

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