Interstratified Clays. I. Theoretical

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Abstract

A method for computing the diffraction effects from interlayered clay minerals is formulated, and the results are compared with those obtained using MacEwan's method. The method allows examination of individual layer transforms or of the mean for any given set and can calculate effects not readily obtained with MacEwan's method, e.g., three-component systems. Results from both methods are in good agreement. Differences in the results were caused by the particular composition distribution assumed for either method. Positions and shapes of diffraction maxima are determined by composition, composition distribution, the presence of one-layer spacings, and the distribution of the d values, in addition to the kind of interlayering. The kind of interlayering between I, a 10 Å repeat layer, and M, a 17 Å repeat layer, e.g., random or ordered, is most readily determined in the mid-composition range, that is, approximately 0.75 < pM < 0.25. The concept of IMII superlattice units formulated by Reynolds and Hower (1970) is reinterpreted. Patterns interpreted as due to IMII ordering can be explained by the existence of a montmorillonite layer deeply imbedded within an otherwise normal 10 Å crystallite. The latter interpretation may give some insight into differences between clay minerals termed illite and K-bentonites having similar expandable character.

Introduction

Most analyses of interstratified clays in recent years have utilized or have been based on the method formulated by MacEwan and co-workers (1956, 1958, 1959, 1961). Reynolds and Hower (1970; hereafter designated R&H) have extended the methods of MacEwan to computer analysis and have matched calculated and experimental diffraction patterns for a variety of cases. The favorable agreement for the majority of the comparisons reinforces one's belief in the corrections of the MacEwan approach. Certain of the conclusions of R&H, however, seem suspect. For example, they conclude that (a) virtually all illite/montmorillonites are ordered for pM ≤ 0.40, and (b) a special kind of ordering is present for pM ≤ 0.10. These conclusions raise questions such as: (1) Why should ordering be restricted to a particular structure/composition range? (2) How do these conclusions explain the nature of substances such as the type illite from Fithian and related clay minerals (Gaudette, Eades, and Grim, 1966)? (3) How do these conclusions fit with the “frayed-edge” concept of Bray (1937) and, more recently, of others? (4) What is the role played by the component having a single glycol layer sandwiched between two silicate layers? The search for answers to these and other questions led to the investigation of a method alternative to that of MacEwan for computing the X-ray diffraction effects from interstratified clay systems. Natural clays were analyzed, and laboratory experiments were performed which have a bearing on the conclusions of R&H; these are reported in Part II.

The main objective of this paper is to present this alternative computational method, to illustrate the results obtained by using it, and to compare it with the method and results obtained by using MacEwan's methods. Basically, the present study used the Fourier transform methods of Lipson and Taylor (1957), first applied to layer silicates by Ross (1968). Only the one-dimensional diffraction effects of the basal reflections of layer silicates were considered.
The essential correctness of the results obtained by using MacEwan’s methods is undoubted. However, it seemed desirable to explore a different method, if only to substantiate and verify what was already known. There are instances, however, in which MacEwan’s methods are not applicable or are difficult to apply. For example, the extension of his methods to an analysis of clay systems containing three or more types of layers, with precise attention paid to the incorporation of the scattering effects by the individuals, is a formidable problem even for the computer. Further, MacEwan’s method assumes that certain “consistency relations” (1956, p. 97) are obeyed. These relations do not hold for individual cases. The method used herein has the capability of circumventing these difficulties in the MacEwan approach, but it too has some inherent difficulties. Therefore, a secondary objective of this paper is to analyze and compare the results obtained from both methods. The equally important problem of the genesis of the mixed-layer clays or their thermodynamic status (Zen, 1962, 1967) was not considered.

Definitions and Symbols

The terminology and symbols used by MacEwan are adhered to herein. However, some definitions are desirable. N: the number of silicate layers per crystallite. NI: equals N-1, the number of interlayer regions or spacings. The identity of individual interlayer spacings is given by letters such as A and B or I and M. I: a 10 A repeat unit or, more precisely, a unit containing no ethylene glycol layers. M: a 17 A unit equivalent to montmorillonite or a unit containing two glycol layers. Component: a general term covering A, B, etc. Layer configuration or Case: a specific arrangement of A, B, and C, etc., AAABBAC equivalent to an “X-ray particle” or “crystallite.” \( n_A \), etc: the number of A layers in a given layer configuration. \( p_{AA} \), etc: the percentage of A in all of the cases under consideration, i.e., the overall proportion of A in the sample. \( AA \), etc: the number of A contacts in a given layer configuration.

MacEwan’s Method

The previously referenced papers fully describe MacEwan’s method so we shall simply point out those features appropriate for comparison with our method of analysis. The MacEwan method is not designed to compute the diffraction effects of a particular layer configuration. Rather, once \( p_A \), \( N \), and \( p_{AA} \) are chosen, assuming a two-component interstratification, the coefficients of each term in his series are fixed. Each of the terms corresponds to a particular interlayer spacing, and the coefficients correspond to the relative frequency of the particular spacing. A necessary consequence of this treatment and of the assumption that the “consistency relations” are valid is to superimpose a particular distribution of chemical compositions on the crystallites. To make this clear, the coefficients used in MacEwan’s series are listed in Table 1 for \( NI = 3 \), \( p_A = 0.7 \) for random \( (p_{AA} = 0.7) \) and ordered \( (p_{BB} = 0) \) interlaying. Looking at the random case, it can be seen that BBB is given a weight or coefficient of 0.027 which implies that “crystallites” composed solely of B occur with a probability of 0.027. Further, multiplying each of the coefficients, e.g., ABB, by the corresponding percentage of B in that particular term, i.e., 0.67, then summing, the overall proportion of B = 0.3 is obtained. In other words, a particular “distribution of chemical compositions” is assumed to be present.

Similarly, the MacEwan coefficients are set forth in Table 1 for regular interlaying. The “composition distribution” here is more restricted about the mean overall composition than for the random case. This is necessary since BB contacts are forbidden. The data in Table 1 further show that for a spacing represented by 2B’s and 1A, a larger probability is accorded to the array having B on the ends, i.e., AAB and BAA, rather than ABA. This is a consequence of the “consistency relations.” However, it is conceivable that clays may occur in nature that have B’s “on the inside” only. MacEwan’s method would not correctly represent such a situation. Similar analyses extended to other \( p_A \) values.

<table>
<thead>
<tr>
<th>Term</th>
<th>( p_{(Random)} )</th>
<th>( p_{(Regular)} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>AAA</td>
<td>0.343</td>
<td>0.22857</td>
</tr>
<tr>
<td>AAB</td>
<td>0.147</td>
<td>0.17143</td>
</tr>
<tr>
<td>ABA</td>
<td>0.147</td>
<td>0.304286</td>
</tr>
<tr>
<td>BA</td>
<td>0.147</td>
<td>0.17143</td>
</tr>
<tr>
<td>BBB</td>
<td>0.063</td>
<td>0.00</td>
</tr>
<tr>
<td>BBA</td>
<td>0.189</td>
<td>0.12857</td>
</tr>
<tr>
<td>BAA</td>
<td>0.063</td>
<td>0.00</td>
</tr>
</tbody>
</table>
enable one to demonstrate that the distribution of compositions is wider and generally has smaller coefficients for overall compositions in the mid-range, say $p_A \sim 0.5$, and is narrower and has generally larger coefficients for large values of $p_A$ (or $p_B$), e.g., $p_A = 0.9$. The distribution of compositions for regular interlayering is always more restricted about the mean than that of the corresponding random case, i.e., for a given $p_A$. These observations are true irrespective of $N$. In short, the MacEwan method tacitly assumes the existence of crystallites having a particular “composition distribution” once $p_A$ and $p_{AA}$ are fixed. However, given a single $N$, e.g., $N = 11$, for all crystallites in a given sample, then arbitrarily setting $p_A = 0.8$, one can envision a situation in which all crystallites have “identical compositions,” i.e., $8A$ and $2B$. Although highly restricted, this “narrow composition distribution,” i.e., all crystallites have a composition equivalent to the overall mean composition of the sample, may better approximate a given clay.

Along the same lines, the presumption that the kind of interlayering is specified by a knowledge of a single term such as $p_{AA}$ is a necessary consequence of the consistency relations. The existence of a single layer configuration with a given $n_A$ is thereby precluded. With the MacEwan method it may not be possible to distinguish between specific configurations such as $AABAAABAAA$ and $ABAAAABAAA$, whereas, in fact, these two layer configurations give distinct transforms. The reason for this is the spread of compositions inherent in the MacEwan calculation. A fixed arrangement of $B$’s and $A$’s and sharply restricted values of $n_A$ in all of the crystallites may more accurately describe some real clays. In such instances the consistency relations of MacEwan are not valid and the clay cannot be adequately described by any values of $p_{AA}$, etc. It seems likely that it is this feature inherent in the MacEwan method that led R&H to devise an artifice, i.e., an IMII superlative unit, which circumvents the MacEwan coefficients and in effect changes the composition distribution and also restricts the positions of the components within the crystallites. This extension of the MacEwan method may not be warranted since it leads to a conclusion of doubtful validity. We will show later that a different interpretation of “Kalkberg-type” diffraction patterns is possible.

Let us pursue the foregoing from a different viewpoint and examine the conclusion of R&H that virtually all illite/montmorillonite interlayered mixtures are ordered for $p_M \leq 0.4$. Employing the MacEwan method in the way they did means that the kind of interlayering can be specified by knowing any one of the probability coefficients $p_{AA}$, etc. Assuming a two-component system and $p_A > p_B$ one can examine $p_{AA}$ as a function of $p_A$ for the kinds of interstratification, mechanical $(p_{AA} = 1)$, random $(p_{AA} = p_A)$, and regular $[p_{AB} = 0$, therefore $p_{AA} = (p_A - p_B)/p_A]$. This examination leads to the conclusion that the $p_{AA}$ values are most different and, therefore, the kind of interlayering is easiest to specify, in the mid-composition range. For all practical purposes the $p_{AA}$ values for ordered and random interlayering are identical for $p_A \sim 0.9$ and they are similar even for $p_A = 0.8$. Real clays may differ not only in the kind of interlayering but also in size distribution of the crystallites, chemical composition and composition distribution, distribution of $d$ values, or in structural details such as inverted tetrahedra. These differences can act to reduce the distinction between clays resulting from different kinds of mixing. On the basis of this analysis it would appear that the kind of interstratification is most readily determined only in the mid-composition range, e.g., $0.75 < p_A < 0.25$.

**Method of Analysis**

A computer program was written for CuKα radiation to calculate the squared modulus of the layer transform in the $c^*$ direction for interstratified clay systems containing up to three components. All calculations included contributions from both real and imaginary terms. The resulting diagrams allow a comparison of various hypothetical mixed-layer clays whose sole differences are structural/compositional. The calculations are free of any additional assumptions concerning the pertinent $L_p$ factor, crystallite size distribution, or apparatus functions. They do not represent “complete diffraction patterns” such as those given by R&H. $G^2$ was computed as a function of $\theta$. The $d$ values shown above prominent maxima on the diagrams were determined to the nearest $0.05^\circ 2\theta$, this being the approximate precision with which one can read real diffraction diagrams for a goniometer scanning speed of $1^\circ 2\theta$/minute and a chart speed of 30 inches/hour. Com-

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1 For a copy of this program, order document AM-75-002 from the Business Office, Mineralogical Society of America, suite 1000 lower level, 1909 K Street, N.W., Washington, D.C. 20006. Please remit $1.00 for the microfiche.
putations were made for the two-component illite/montmorillonite (10 A/17 A) system and covered the entire composition range. As the study progressed and the utility of the method became apparent, the effect of (a) the one-glycol layer complex and (b) composition distribution was investigated. Each transformation was calculated with the origin taken at the plane of the octahedrally coordinated atoms in an end silicate layer. A montmorillonite unit was defined as a two-glycol layer interlamellar space, and an illite as an interlamellar space containing no glycol molecules. The scattering factors and other parameters used (Table 2) were quite similar to those used by R&H so as to facilitate comparison with their work. The same silicate layer was used for all components.

The detailed nature of the one-glycol layer complex has not been fully investigated. The parameters chosen to represent this component are in agreement with the scanty literature on the subject (Bradley et al., 1958; Brunton, Tettenhorst, and Beck, 1963) and the premise that it was similar to the 17 A component.

Transforms for particular layer configurations were generated one at a time for a given Ni and component array, e.g., Ni = 5 and IIIII, for which in this example 0.2. In addition all possible arrangements of 4I's and 1M were generated and a “mean” pattern calculated. An obvious advantage of this approach is that individual transforms can be examined. Potential disadvantages are (a) the composition of each crystallite is restricted to being identical with the overall composition, and (b) calculations are made for a single Ni. The former is easily overcome while the latter is not. Consequently, calculations were made for Ni = 5, Ni = 10, and Ni = 20 to determine the effect of thickness. The reason for choosing these Ni values was that a preliminary laboratory study showed that many natural illite/montmorillonites dehydrated for 1 day at 400°C give apparent Ni values in this range as determined from the measured breadths of the diffraction maximum at about 27° 2θ with CuKα (see Tettenhorst and Roberson, 1973, for details.) This procedure limits the particular compositions that can be investigated once Ni is fixed; i.e., p can vary only in 0.20 increments for Ni = 5, only in 0.10 increments for Ni = 10, etc. Generation of all possible individual transforms and computation of a mean transform is simple if Ni is small, say Ni = 5, since the number of layer configurations is given by (Ni)A/(nA!nB!). For Ni = 10 and nA = 9 (p = 0.9), ten arrays are possible, and only half of these need be calculated to produce a mean since identical transforms are generated by configurations that are symmetrical. For Ni = 10 the number

<table>
<thead>
<tr>
<th>Atom</th>
<th>Z(A)*</th>
<th>p</th>
<th>B(A°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Potassium</td>
<td>-5.00</td>
<td>0.3</td>
<td>1.68</td>
</tr>
<tr>
<td>Oxygen</td>
<td>-3.28</td>
<td>3.0</td>
<td>1.68</td>
</tr>
<tr>
<td>Silicon</td>
<td>-1.06</td>
<td>3.0</td>
<td>1.68</td>
</tr>
<tr>
<td>Aluminum</td>
<td>0.0</td>
<td>1.0</td>
<td>1.68</td>
</tr>
</tbody>
</table>

*Atomic coordinates of the opposite "end" layer were equal to

\[ z + (17 \cdot nA + 10 \cdot nI + 13.5 \cdot n_{13.5}) \]
of possible cases increases to 210 in the mid-composition region, for \( p_A = 0.6 \). For \( NI = 20 \) the number of cases is quite large at any composition, and it is simply impractical to compute so many individual cases in order to arrive at a mean transform. The power of the MacEwan method is that all such cases are considered with the additional advantage that \( p_A \) can be chosen equal to any value and not restricted to a composition such that \( p_A = n_A/NI \).

A specific example will be given to show how the computation of thousands of cases was circumvented. There are 38,760 ways of arranging 14A's and 6B's for \( NI = 20 \) (\( p_A = 0.7 \)). Some experimentation showed that the mean transform calculated from 100 randomly selected cases produced a reasonable approximation to the true mean. This was checked in two ways. First, a second batch of 100 cases randomly selected as before was generated, and the resultant mean transform was virtually identical to the first. This process was repeated for several \( p_A \) values with identical results. Further, these mean patterns compared very favorably with those calculated for the same \( p_A \) value for \( NI = 10 \), for which all possible cases could be calculated.

One cannot be certain that just any batch of 100 randomly selected layer configurations will provide a reasonably accurate mean transform. To ensure a reasonable agreement, the individual cases in sets of 25 were examined for AA and BB contacts and 4 such sets were put together to form the 100 to produce the “correct number” of AA and BB contacts. The “correct number” of AA contacts is given by the relation \( x_{AA} = p_A \times N_c \times (n_A - 1) \) for which \( N_c \) is the number of layer configurations (here = 100) and \( (n_A - 1) \) is the maximum number of AA contacts. The procedure is identical for the BB contacts. The reader may better understand this relationship with a simple example: Assume \( NI = 5 \) and \( p_A = 0.6 \), for which the total \( N_c = 10 \). Therefore, \( x_{AA} = 12 \) and \( x_{BB} = 4 \), as the reader can easily verify by listing all ten cases.

The procedure described was not necessary to compute a mean transform for a set of ordered configurations since these were always relatively small in number and all could be computed. Ordered is here taken to mean \( BB = 0 \) and \( AA = \) minimum for \( p_A > p_B \). In certain instances, however, mean transforms from a set of “partially ordered” patterns were calculated from a set of 100 individual cases, partial order being defined as \( BB = 0 \). To take an example, for \( NI = 20 \), and 12A-8B there are 1280 cases with \( BB = 0 \). Each of the 1280 cases was assigned a number, and 100 of these were selected randomly for the purpose of computing a mean transform.

**Discussion of Results**

**Two Components**

Representative patterns generated by the method described in the previous section are shown in Figures 1-4. Only those in the range \( p_B \leq 0.4 \) are included herein. Most of the transforms shown are for \( NI = 10 \); a few computed for \( NI = 20 \) are included for comparison. “Diffraction ripples” are present on all of these diagrams and are particularly pronounced at low 2\( \theta \) angles. MacEwan and Ruiz Amil (1959) noted that these diffraction ripples can be attenuated to produce a “smooth curve” by including a distribution of crystallite thicknesses, i.e., several \( N \) values are used in the computations. As will be shown later, “smoothing” was also accomplished here by including a distribution of crystallites having different chemical compositions. Although real clays likely represent a distribution in crystallite thickness and composition, their inclusion into the computations constitutes a guess on the part of the investigator as to the true situation.

Any particular pattern shown in Figures 1-4 represents the mean transform calculated for a number of crystallites all of which have the same \( NI \) and \( p_B \). For example, the curve shown at the bottom of Figure 2a is the mean of the 45 separate cases which have 8I and 2M layers, i.e., the 45 ways of arranging 8I and 2M layers. Nine of the

![Fig. 1. Calculated diffraction pattern, \( p_I = 1.0, NI = 10 \).](image)
45 cases have touching $MM$ layers, a proportion exactly equal to the overall percentage of $M$ in the “sample,” i.e., $MM = 0.2$. This “mean” transform appears to give a diagram strictly analogous to MacEwan’s “random” situation and will be so considered in the rest of this paper. Also shown are the mean transforms calculated for “fully ordered” situations, i.e., $MM = 0$ and $II$ contacts are minimum. For example, the curve in Figure 2c represents the mean of 21 cases in which the $M$ layers are separated from each other and likewise the $I$ layers. This “fully separated” mean transform appears analogous to MacEwan’s “ordered” situation.

The term “separated” appears preferable to “ordered” as it is more descriptive in those instances not in the mid-composition region $p_M \sim p_I$. The essential difference between those mean transforms computed with fully separated cases, e.g., Figure 2c, and those calculated only with the condition that $MM = 0$, e.g., Figure 2b, is that the latter contain some layer configurations having $M$’s on the ends of the crystallites.

**Interlayering Defined**

The kinds of interlayering as used in this study are defined in Table 3 in terms of $Q$ values. The $Q$ values are, for a representative set of layer configurations, the proportion of contacts of a given kind, say $AA$, compared to the maximum number of contacts of that type that would occur if all cases were completely segregated. The $Q$ values given in the table are valid only for $p_A > p_M$ and $NI \geq 3$. The expression for $Q_{AA}$ (separated) was obtained from its equivalent $(n_A - n_B - 1)/(n_A - 1)$, the numerator of which denotes the minimum number of $AA$ contacts and the denominator the maximum
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The proportion of $A$ contacts for random and ordered mixing relative to the maximum number of $A$ contacts if all cases were segregated vs $p_A$. (a) Random given by $---$, ordered $NI = 3$ given by $- - - - - - - - - - -$ ordered $NI = 10$ given by $- - - - - - - - - - -$, ordered $NI = \infty$ given by $- - - - - - - - - - -$. (b) Random and ordered ($NI = 10$) are redrawn assuming $p_A$ known to ±5 percent.

Fig. 5. The proportion of $A$ contacts for random and ordered mixing relative to the maximum number of $A$ contacts if all cases were segregated vs $p_A$. (a) Random given by $---$, ordered $NI = 3$ given by $- - - - - - - - - - -$, ordered $NI = 10$ given by $- - - - - - - - - - -$, ordered $NI = \infty$ given by $- - - - - - - - - - -$. (b) Random and ordered ($NI = 10$) are redrawn assuming $p_A$ known to ±5 percent.

Segregated interlayering as defined in Table 3 is not strictly equivalent to MacEwan's definition of mechanical mixing of individual crystallites. The latter implies individual crystallites while the former means segregation within a crystallite for which one contact exists between the two components. The computed transforms for either are quite similar, however.

Fig. 4. Calculated diffraction patterns, $p_t = 0.6$; (a) $MM = 0.4$, top $NI = 20$, bottom $NI = 10$, (b) $MM = 0$, $II = \text{minimum}$.

number of $AA$ contacts for a given composition using relations such as $p_A = n_A/(n_A + n_B) = n_A/NI$. Drawn in Figure 5a are continuous curves of $Q_{AA}$ vs $p_A$ for several $NI$ for ordered interlayering and the relation for random. A similar diagram was given by Gilkes and Hodson, 1971. As concluded previously when considering MacEwan's method, the distinction between ordered and random is best determined in the mid-composition range. This point is made clearer in Figure 5b where a conservative uncertainty in the value of $p_A$ of ±5 percent has been applied to both the random and the $NI = 10$ (separated) curves. The overlap of the two regions is severe at $p_A > 0.85$ but not at $p_A = 0.75$. Figure 5a further suggests that the distinction of random and ordered interlaying should be easier for smaller $NI$ since the curve for the $NI = 3$ separated case is the farthest from the random case. This relationship is not likely to be borne out in practice because of lower intensity and broader maxima with smaller $N$.

Segregated interlayering as defined in Table 3 is not strictly equivalent to MacEwan's definition of mechanical mixing of individual crystallites. The latter implies individual crystallites while the former means segregation within a crystallite for which one contact exists between the two components. The computed transforms for either are quite similar, however.

Some Observations

The average transforms generated herein for either random or ordered interlayering can be compared with those produced by using MacEwan's method (R&H) in a general way, particularly with

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Table 3. Definition of the Kinds of Interstratification as Employed in This Study

<table>
<thead>
<tr>
<th>Kind of interlayering</th>
<th>$Q_{AA}$</th>
<th>$Q_{BB}$</th>
<th>$Q_{AB}$ (or $Q_{BA}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Segregated</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Mean (Random)</td>
<td>$p_A$</td>
<td>$p_B$</td>
<td>$p_A \cdot p_B \cdot NI$</td>
</tr>
<tr>
<td>Separated (Ordered)</td>
<td>$2 \cdot NI \cdot p_A \cdot p_B \cdot NI - 1$</td>
<td>$0$</td>
<td>$p_B \cdot NI$</td>
</tr>
</tbody>
</table>

*Q values for $p_A > p_B$ and $NI > 3$
respect to the shape and 2θ position of prominent maxima. Comparison of intensity or diffracted energy is less reliable because the diagrams here lack certain of the factors assumed by R&H, as described earlier. Overall, the same basic information is obtained from either method. In general, the 2θ positions calculated agree well with those cited by R&H in their Table 3. The differences that do occur are confined to the low (<10°2θ) region (e.g., 9.6°2θ vs 9.43°2θ for R&H for \( p_m = 0.4 \) and \( MM = 0.4 \)). These differences are almost entirely a result of the restricted composition used here versus the distribution inherent in the MacEwan method. This observation suggests that “peak positions” alone cannot be used to determine the percentages of the various components in a natural mixed-layer clay with a very high degree of precision (say to within several percent) since the distribution of chemical compositions of the individual crystallites in real clays is not known.

Hower and Mowatt (1966) attributed the steep rise in the background intensity at low angles (<4°2θ) to signify the presence of interstratification (p. 847, their Fig. 8). Comparing 0.5 μ sized particles of muscovite and illite, as segregated by sedimentation methods, they noted that illite had a steeper background rise, and a broader (001) peak with \( d \) slightly greater than 10Å. The distinctive features of the illite were attributed to interstratification and, specifically, not to “particle size.” The present work does not support this conclusion as shown in the figures since the rate at which the background intensity rises at low angles is similar for all diagrams for a given \( N \). Contrary to Hower and Mowatt (1966), the distinctive features of their X-ray pattern of illite compared to muscovite can be attributed primarily to smaller mean crystallite thickness. Identical sedimentation-size-fractions of different layer silicates do not insure that identical crystallite thicknesses will be obtained for both materials (Tettenhorst and Roberson, 1973) nor that the morphological particle thicknesses will be the same since the particles in different samples may vary in their areal dimensions, specific gravity, extent of association with water molecules, etc. Along the same lines R&H have attached considerable significance to the match up of the rise in background at low angles for their calculated and experimental patterns. Their computed patterns contained the powder Lp factor. It is suggested that, if an experimental pattern is encountered whose background rise cannot be explained by \( N \), then attention should be given to the precise form of the Lp factor. It is possible that not all layer silicates or oriented interstratified clays behave like “powders” but that some may approximate a single crystal in scattering efficiency. MacEwan, Ruiz Amil and Brown (1961) drew a similar conclusion.

Layer Location within a Particle

Shown in Figure 6 are the five individual transforms and their mean, calculated for \( NI = 10 \) and \( 9I - 1M \), i.e., \( p_M = 0.1 \). As the position of the single \( M \) layer changes from the “outside of the crystallite” (position 1, Fig. 6a), to the “middle of the crystallite” (position 5, Fig. 6e), the diffraction pattern changes as follows: (a) the maximum at 8.8°2θ, which is rather skewed to low angles, splits into two sharper maxima one of which migrates to 9.0°2θ and the other develops at about 8.0°2θ; (b) the maximum at 17.6°2θ splits into two maxima of which the more prominent migrates to 17.2°2θ, the other to about 18.4°2θ. Note that the patterns in Figures 6d and 6e resemble the Kalkberg pattern of R&H (their Fig. 2, p. 30). Possibly a variation of thickness and, perhaps, of chemical composition.
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for the crystallites would give a “smooth” computed pattern that would match very well the experimental pattern of Kalkberg. It is concluded that patterns like the Kalkberg clay—rather than illustrating a special kind of ordering such as IMII at a composition, i.e., \( p_M \sim 0.1 \), where the kinds of interlayering are virtually impossible to distinguish—actually show that \( M \) layers are imbedded within the crystallites. Although the conclusion of R&H that Kalkberg-type patterns represent a special kind of ordering may be incorrect, their assignment of the quantity of expanded layers, i.e., \( p* \sim 0.1 \), in these interstratified clays appears to be reasonable.

Illites that give X-ray patterns such as Fithian or Grundite (Gaudette, Eades, and Grim, 1966) and the Strasburg “metabentonite” (API No. 37) are common. Glycolated patterns typically show an asymmetrical 001 peak at 10 Å, this being skewed to low angles, and a rather broad 002 peak at about 17.6°2θ. How are such clays to be interpreted? The presence of some expandable layers is indicated, but the disposition of these expanded layers must be different from that in Kalkberg-type clays. Although speculative it is suggested that the patterns of Fithian-type clays resemble that shown in Figure 6a and, therefore, the crystallites in Fithian may, more than not, terminate on expanded layers. The expanded layers would then define the limit or thickness of those crystallites. This suggestion is in broad agreement with the “core-rind” or “frayed-edge” hypothesis advanced for the structure of these clays by Gaudette, Grim, and Metzer (1966) on the basis of the sorption behavior of cesium. To carry this suggestion one step further, this interpretation might suggest that Fithian-type clays generally represent the effect of “degradation,” i.e., the “opening-up” of the particles at their edges or “surfaces” while Kalkberg-type patterns generally represent the effects of “aggradation,” the “closing-up” of the particles until the last remaining expandable layer is buried inside the individual crystallites.

Highly-Expandable Clays

Although not shown in the figures, two observations are noted concerning the patterns at the highly expandable end of the I/M scale, i.e., \( p_M \sim 1.0 \). First, compared with the patterns computed for \( p_M = 1.0 \), the introduction of 10 percent collapsed layers decreases the peak intensity of 001 to about half its value, its position being rather unaffected, and decreases the position of 002 by about 0.10°2θ. Second, the effect of a layer of glycol molecules on the surfaces or ends of the particles was calculated and compared with the “no-surface-glycol” situation (Tettenhorst and Reynolds, 1971) for both \( p_M = 1.0 \) and \( p_M = 0.9 \). The significant effects for both compositions compared to no-surface-glycol were (a) the intensity of the 001 peak decreased about 15 percent and the intensity of 002 increased by 15 percent and (b) the position of 002 increased about 0.1°2θ. Tettenhorst and Roberson (1973) noted that the relative positions of 001 and 002 from a group of montmorillonites were rather scattered and did not fit well with either a surface glycol or no-surface-glycol model. Interstratification was suggested as being responsible for the poor fit and this seems borne out by the present observations. Indeed, it may be difficult to decide from the diffraction patterns of individual montmorillonites what the relative contributions are between surface glycol or a small percentage of non-expanded layers. The transforms shown in this paper and those provided by R&H were calculated assuming no surface glycol.

Composition Distribution

The transforms discussed previously were the mean of some given number of layer configurations for which each individual layer configuration had the same composition. This is not necessary. The method used herein is capable of simulating any composition distribution, in particular, that used in the MacEwan method. The details for one such simulation are set forth in Table 4 for \( p_M = 0.2 \), \( MM = 0.2 \), and the resulting mean transform is shown in Figure 8a. The total number of layer configurations used to compute the mean transforms in these instances was 100. The MacEwan coefficient (column 2 in Table 4) was computed for a particular \( N_1 (= 10) \) and \( p_M \). This figure determined the number of layer configurations having a given composition, e.g., \( 7I - 3M \), that were included in the computation of the mean, i.e., 20 cases out of a possible 120. The specific cases used for calculating the mean were selected randomly. A final check was made to insure that the proper number of \( II, \ etc \) contacts was obtained for the 100 cases before performing the calculation. Several such mean transforms simulating MacEwan’s composition distribution were computed in this manner for various mean \( p_M \) values (Figures 7-9). The effect of ordering was also taken into account; two such
Table 4. Cases Used to Simulate the MacEwan Composition Distribution for \( N_I = 10, \ r_m = 0.2, \) and \( M_M = 0.2 \)

<table>
<thead>
<tr>
<th>Sequence</th>
<th>MacEwan Probability Coefficient</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>101</td>
<td>0.107</td>
<td>11</td>
</tr>
<tr>
<td>91 - 1M</td>
<td>0.268</td>
<td>27</td>
</tr>
<tr>
<td>81 - 2M</td>
<td>0.302</td>
<td>30</td>
</tr>
<tr>
<td>71 - 3M</td>
<td>0.302</td>
<td>20</td>
</tr>
<tr>
<td>61 - 4M</td>
<td>0.088</td>
<td>9</td>
</tr>
<tr>
<td>51 - 5M</td>
<td>0.026</td>
<td>3</td>
</tr>
</tbody>
</table>

Sum = 0.992

patterns for \( p_M = 0.2 \) are shown in Figures 8b and 8c.

Compared with the transforms shown previously, the most obvious difference is the smoothness of the curves and relative lack of diffraction ripples, except at angles below about 4°2\( \theta \). The smoothness is the result of the composition distribution and consequent variety of thickness for \( N_I = 10 \). A comparison of these with their equivalents computed by the MacEwan method shown in R&H indicates a close match of peak positions and peak shapes. Differences in the three transforms illustrated in Figure 8 for \( p_M = 0.2 \) are discernible, particularly the buildup of scattering at about 7°2\( \theta \) for the fully separated case. The presence of a peak in this 2\( \theta \) region along with a peak between 9–10°2\( \theta \) may be an indication of ordering as noted by Reynolds (1967). However, the form and position of peaks in this 2\( \theta \) range are also very susceptible to composition distribution as noted, and the presence
of one-layer spacings or d value distributions, as
will be shown in Part II.
A notable feature shown in Figures 4b and 9
for ordered or partly ordered cases is the prominent
superlattice peak that develops at about 3°2θ for
compositions given approximately by 0.35 ≤ psx
≤ 0.65. Its presence seems to be the surest indica-
tion of ordering, as might be expected. When this
superlattice peak is indistinct or represented merely
by a shoulder or break in the steep background rise
at low 2θ, the composition is given approximately
by psx = 0.30 – 0.35. For example, see the X-ray
pattern of A.P.I. No. 42 from High Bridge, Ken-
tucky, shown in Figure 4 of Hower, 1967.

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