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Study of dehydroxylated-rehydroxylated smectites by SAXS

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Abstract: Montmorillonite and beidellite are dioctahedral 2:1 phyllosilicates. The weakness of the bonding between layers allows the intercalation of water molecules (disposed in layers) in the interlayer space. The samples studied are constituted of cv layers (cv for vacant octahedral sites in cis positions). They have been dehydroxylated. This is accompanied by the migration of the octahedral cations from former trans-octahedra to empty cis-sites therefore the layers become tv (vacant site in trans position). To characterize the stacking of the layers, SAXS (Small Angle X-ray Scattering) analyses have been investigated in natural (N) and after a dehydroxylation-rehydroxylation cycle (R) states. The SAXS pattern modelisation for Na -exchanged samples in the N state shows that the layers stack in particles with well defined interlayer distances $d_{001}$ corresponding to 0 water layer, 1 water layers and 2 water layers. The dehydroxylation-rehydroxylation cycle increases the proportion of interlayer distances with zero water layer and the disorder in the stacking. The decreasing of the disorder parameter with the proportion of tetrahedral charge in the N and R sample shows that the distribution of the water layers depend on the localization of the deficit of charge.

Résumé: Lea montmorillonites et les smcctites sont des phyllosilicates 2:1 dioctadétriques. Les liaisons entre feuilles sont suffisamment faibles pour permettre l'insertion, dans l'espace interfolialaire, de molécules d'eau qui se disposent en couches. Les échantillons étudiés ont des feuilles cis-vacants (le site octadétrique inoccupé est en une des deux positions "cis") Ils ont été dehydroxylés. Ceci s'accompagne d'une migration cationique, à l'intérieur des couches octadétriques, des sites trans vers les sites cis et le feuillet devient trans-vacant. Des expériences de Diffusion X aux Petits Angles (DPA) ont permis de caractériser l'empilement des feuilles. La modélisation des diagrammes de DPAX met en évidence, pour les échantillons sodium non traités, des emplacements de feuilles formant des particules avec des distances interlaminaires à 0, 1 et 2 couches d'eau. Après le cycle de déshydroxylation-réhydroxylation, la proportion de feuilles avec une distance interlaminaire correspondante à zéro couche d'eau et le désordre dans l'empilement des feuilles augmentent. La décroissance du paramètre de désordre avec la proportion de charges tétraédriques montre que l'organisation des couches d'eau dépend de la localisation du déficit de charge.

1. INTRODUCTION

Montmorillonites and Beidellite (smectite family) are dioctahedral 2:1 phyllosilicates of idealized structural formula:

$$M_{x+y}^{+} (Mg, Fe^{II})_x (Al, Fe^{III})_{2-x} Si_{4-y}Al_y O_{10}(OH)_2$$

where $M^+$ is the interlayer cation which compensates the negative charge of the layer arising from cation substitutions inside the layer. The 2:1 notation means that layers are formed of two sheets of tetrahedra sandwiching a sheet of octahedra (Fig. 1a). Dioctahedral means that in the octahedral sheet, only two of the three sites per one half unit-cell are occupied.

The relative weakness of the bonding between layers, through the interlamellar cations (Na, Ca, ...) authorizes intercalation of various molecules and in particular water molecules. The water molecules are organized in layers in the interlayer space and generally montmorillonite and beidellite minerals consist in interstratified samples in which the number of water layers characterize each type of layer (Fig. 1b). For example, in air dried state, the Wyoming Na-montmorillonite correspond of hydration states of 0, 1 or 2 water layers.
Generally, in the natural montmorillonite or beidellite, the vacant site is in one of the two cis positions (cv1 or cv2 layer, $C12(I)$ space-group, Fig. 2a, [1]). After a treatment consisting of dehydroxylation (replacement of the adjacent hydroxyls of the octahedral sheet by a residual oxygen atom $O_r$: $2(OH) \rightarrow H_2O(T) + O_r$) and rehydroxylation (regain of the hydroxyls: $O_r + H_2O \rightarrow 2(OH)$), the layers become trans vacant (tv layer, $C12/m(I)$ space-group, Fig. 2b, [2, 3]). This transformation results of the migration of the octahedral $A1^{3+}$ cations during the dehydroxylation [1]. In addition to the changing in the position of the vacant octahedral site (trans to cis), the dehydroxylation-rehydroxylation cycle induces two other structural effects. First is the increasing of the OH-OH distance, which is longer in cv layers in comparison with tv ones (see Fig. 2). Second is the changing of the apparent interlayer distance $d_{sp}(001)$ [3, 4].

Recent studies [3] have shown that, in Wyoming Na-montmorillonite, the distribution of water layer in the interlayer spaces changes in small proportion after a dehydroxylation-rehydroxylation cycle which modify the hydration properties of the sample. The aim of this work is to determine the number and proportion of water layers in the natural state (N) and in the rehydroxylated state (R) of three samples differing by the localization of the deficit of charge.

![Figure 1: Schematic representations of a 2:1 layer (a) and a stacking of interstratified layers (b).](image1)

![Figure 2: Schematic representation of an octahedral sheet in the case of cis-vacant layers (a) and trans-vacant layers (b).](image2)

2. METHOD

To determine the water contents in the interlayer space a simulated method has been used. The model is based on the notion of particle and of particle internal structure (Fig. 3, [4, 5]). The basic components of all 2:1 phyllosilicate structures are the layers. One particle consists of a stacking of $M$ parallel layers in which interlayer spaces can be different from one layer to another (Fig. 1b).
In the case of a heterogeneous clay system with water content, a particle can be consisted with two swelling phases in which interlayer spaces can be different from one layer to another. The distance distribution inside a phase is totally random. If we assume that \([4, 6] : 1\) layers have a great lateral extension and particles have all possible orientations in space, \(2\) all the layers in a given phase have the same structure factor, then the matrix expression of the scattered intensity is \([4, 7]\):

\[
I(s) = \frac{1}{s^2} \sum_p \alpha(M) S_{\text{pur}} \times \left[ \text{Re} \left\{ [F][W] \mathbb{I} \right\} + 2 \sum_n \left( \frac{M - n}{M} \right) \mathbb{Q}^n \right] \tag{1}
\]

where \(s\) is the modulus of the diffraction vector \((s = 2\sin\theta/\lambda, \lambda = \text{wavelength})\); \(\alpha(M)\) the distribution of the number of layers per particle; \(\text{Re}\) means the real part of the final matrix; \(S_{\text{pur}}\), the sum of the diagonal terms of the real matrix; \(M\) the number of parallel layers per stack; \(n\), an integer varying between \(1\) and \((M-1)\), \([F]\) the structure factors matrix; \([I]\) the unit matrix; \([W]\) the diagonal matrix of the proportions of the two phases, and \([\mathbb{Q}]\) the matrix representing the interference phenomena between adjacent layers. \([\mathbb{Q}],[F]\) and \([W]\) have the following forms:

\[
[\mathbb{Q}] = \begin{bmatrix}
P_{A,A} \sum p(A), \exp(2\pi isd(A),) & P_{A,B} \sum p(A), \exp(2\pi isd(A),) \\
P_{B,A} \sum p(B), \exp(2\pi isd(B),) & P_{B,B} \sum p(B), \exp(2\pi isd(B),)
\end{bmatrix} \tag{2}
\]

\[
[F] = \begin{bmatrix}
F_A^* F_A & F_A^* F_B \\
F_B^* F_A & F_B^* F_B
\end{bmatrix}; \quad [W] = \begin{bmatrix}
W_A & 0 \\
0 & W_B
\end{bmatrix} \tag{3}
\]

where \(W_A\) and \(W_B\) are the abundances of the two phases; \(F_A\) and \(F_B\) the structure factors related to the two phases; \((d(A), (d(B))\) the possible distances between the layers of the phase A (B); \(p(A), (p(B))\) the probabilities of going from one layer of a thickness \(d(A), (d(B))\) in the A (B) phases to another in same phase; \(P_{A,B}\) the probability of going from one layer in the phase A to another in the phase B \((P_{A,A} P_{B,A}\) and \(P_{B,B}\) are defined in the same way).

The relationships between proportions of the two phases and probabilities are given by:

\[
W_A + W_B = 1; P_{A,A} + P_{A,B} = 1; P_{B,A} + P_{B,B} = 1; W_A P_{A,B} = W_B P_{B,A} \tag{4}
\]
In this study we consider a random interaction between swelling phases which are limited to 0,1 and 2 water layers. In this case, the abundance \( w_i (w_j) \) of one interlayer distance in a given phase can be written:

\[
\begin{align*}
\sum_i p(A)_i &= 1; \sum_j p(B)_j = 1 \\
\end{align*}
\]  

(5)

It is possible to define the mean interlayer distance :

\[
\bar{d} = \sum_i w_i d(A)_i + \sum_j w_j d(B)_j
\]

(7)

and a disorder parameter [7] :

\[
\frac{\bar{d}^2}{\bar{d}^2} = \sum_i w_i (d(A)_i - \bar{d})^2 + \sum_j w_j (d(B)_j - \bar{d})^2
\]

(9)

### 2.2 Water content calculations

Numerous works [5, 6, 8, 9] have shown that different types of water could be defined. We will call "total internal water" the water contained in the swelling phase of clay particles. It is the sum of the water contained in the interlayer space (water layers) and the water contained in the internal porosity (inside the particle, Fig. 3). The external water is all the remaining water (outside the particle).

The probability of finding \( k \) water layers between 2 adjacent layers is given by the total abundance of layers with \( k \) water layers \( w_k \).

The water content \( WC_k \) (expressed in water gram by clay gram) in an interlayer space corresponding the probability \( w_k \) i.e. to a interlayer distance \( d_k \) is given by [8] :

\[
WC_k = \frac{Sp(tot) \times (M - 1) \times w_k \times (d_k - d_{col}) \times 10^{-4}}{2M}
\]

(10)

where \( Sp(tot) = 800 \text{m}^2/\text{g} \) [6] is the total specific area and \( d_{col} \) the distance between two layers in a collapsed phase (\( d_{col} = 9.6 \, \text{Å}, [6] \)).

### 3. EXPERIMENTAL

The smectites sample submitted to the dehydroxylation-rehydroxylation process were Na-montmorillonites sample from Wyoming (Wy) and Camp Berthaux (CB) and a beidellite sample from Unterruppsroth (Un) (Table 1). To get the dehydroxylated state, the natural (N) sample was heated with a 100°C/hour heating rate to 750°C, kept at this temperature during 1 hour, and then cooled in air. For rehydroxylation a gold capsule was filled with 50mg of the dehydroxylated sample plus 15% in weight of water. The capsule was placed in a Tuttle type pressure vessel at 350°C, 1 kbar for 5 days. After removal from the pressure vessel, the rehydroxylated (R) sample was heated at 120°C during 4 hours to extract excess water. The R sample has been studied in air dried state.

Thermogravimetric analyses were performed using a Setaram TGA 92 microanalyser, with a heating rate of 10°C/minute, using 30mg of sample for each measurement. The derived curves of the TGA patterns were drawn to compare the weight loss, assigned to hydroxyl water, in the natural and rehydroxylated states.
X-ray diffraction intensity data were recorded on a SAXS setup using a RIGAKU rotating anode (12 kW) with a curved germanium monochromator (CuKα = 1.5405 Å). The monochromatic beam is convergent in the horizontal plane and diverging in the vertical plane. It is possible to obtain a punctual collimation simply by limiting the height of the beam by a pair of horizontal tantalum slits. Under such consideration, the direct beam dimensions at the focalization point are 300 μm by 500 μm. Alignment and calibration are performed using the 001, 002 and 003 reflections of tetradecanol powder [10]. The cell used is made of aluminium, with mica windows. The surface of the irradiated sample is 0.5 mm by 1 mm. All the experiments had to be carried out at room temperature.

<p>| Table 1. Cation composition of the sample under study calculated per O10(OH)2. |</p>
<table>
<thead>
<tr>
<th>Sample</th>
<th>Interlamellar cation</th>
<th>Tetrahedral cations</th>
<th>Octahedral cations</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Na</td>
<td>Si</td>
<td>AlIV</td>
</tr>
<tr>
<td>Wy</td>
<td>0.29</td>
<td>3.96</td>
<td>0.04</td>
</tr>
<tr>
<td>CB</td>
<td>0.34</td>
<td>3.99</td>
<td>0.01</td>
</tr>
<tr>
<td>Un</td>
<td>0.43</td>
<td>3.55</td>
<td>0.44</td>
</tr>
</tbody>
</table>

4. RESULTS

4.1 Thermogravimetric analyses

The thermo-gravimetric analysis (TGA) shows a change in the dehydroxylation temperature which decreases strongly from the natural specimens to the rehydroxylated samples. This shift is clearly related to the change in the OH-OH distance, which is longer in cv layers (Fig. 2a) in comparison with tv ones (Fig. 2b) [2, 3]. The dehydroxylation-rehydroxylation process keeps the original periodicity of the samples but induces a transformation of the layers into tv ones [2, 3]. An example is given in the Fig.4; the DT curve of the Na-Wyoming montmorillonite sample contains one strong maximum at 700°C. It corresponds to the dehydroxylation of cv 2:1 layers [1, 2, 3]. In contrast, the dehydroxylation of the R sample takes place at a lower temperature (540°C) and corresponds of tv layers [3].

![Temperature (°C)](image)
4.2 SAXS patterns analyses

An examination of the experimental SAXS patterns (Fig. 5a and b) shows some modulations of the intensity situated between 5 and 8 °2θ. These results indicate that the N or R samples have a structure of hydrated solids in which occur short-distance interactions between first neighbors [4, 6, 8]. The sample consists of a set of particles, connected or not, created by the more or less disordered stacking of hydrated layers with a limited number of interlayer water layers (0, 1, 2 or 3). The apparent interlayer distances (\(d_{ap}\)) before and after the dehydroxylation-rehydroxylation cycle are very similar. The width of the modulation increase in this R specimens in comparison with the N ones. These result confirm the previous studied on this subject [3]. Note the presence of a relatively fine additional reflection at \(d_{ap} = 12.5 \text{ Å}\) in the CB R sample.

![Figure 5: SAXS experimental and calculated patterns for the three samples in the natural (a) and rehydroxylated (b) states.](image)

Table 2 presents the value of the different structural parameters \(\bar{d}\), \(\bar{M}\) and disorder, determined by adjusting the theoretical spectra (obtained from the structural model described above) to the experimental ones (Fig. 5). The \(d_{dc}\) values used for the \(\bar{d}\) calculations are given by Berend [11] and Ben Brahim [12].

<table>
<thead>
<tr>
<th>Interlayer hydration</th>
<th>Structural parameter</th>
<th>Water contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>(w_1c)</td>
<td>(w_2c)</td>
<td>(d(A))</td>
</tr>
<tr>
<td>Wy N</td>
<td>0.35</td>
<td>0.49</td>
</tr>
<tr>
<td>Wy R</td>
<td>0.55</td>
<td>0.18</td>
</tr>
<tr>
<td>CB N</td>
<td>0.47</td>
<td>0.24</td>
</tr>
<tr>
<td>CB R</td>
<td>0.48</td>
<td>0.13</td>
</tr>
<tr>
<td>Un N</td>
<td>0.17</td>
<td>0.71</td>
</tr>
<tr>
<td>Un R</td>
<td>0.28</td>
<td>0.58</td>
</tr>
</tbody>
</table>

\(WC_{t}\), total water content
5. DISCUSSION AND CONCLUSIONS

For the three samples, the treatment is responsible of a perceptible increasing of the disorder parameter (Table 2, Fig. 6). These results can be predicted because the increasing of the width of the reflections (Fig. 5). Table 2 shows that, after a dehydroxylation-rehydroxylation cycle, the number of layers with zero water layer increase. The behaviour of the CB sample is distinguishable from others; \( w_{\text{BC}} \) slightly differs between the N (0.47) and the R (0.48) states in comparison with the two others samples (see Table 2). The main characteristic of the CB sample is the exclusive localization of the deficit of charge in the octahedral sheet (Table 2). This can be associated to the small variations of \( w_{\text{BC}} \) in comparison with samples containing no negligible deficit of charge in tetrahedral sites. The weak evolution of \( w_{\text{BC}} \) in the CB sample can explain the increasing of the total water content in the interlayer space which evolve in a opposite direction in comparison with Wy and Un samples. This results permit to assume that the treatment efficiency for the hydration properties is connected to the localization of cation substitutions in the tetrahedral sheets.

The decreasing of the disorder parameter with the proportion of tetrahedral charge (Fig.6) shows that, in particle, the distribution of water layers depend on the localization of the deficit of charge. Localization of cation substitutions in the tetrahedral sheets is a propitious factor for the order in the organization of the mineral.

![Image](Figure 6: Evolution of the disorder parameter with the proportion of the tetrahedral charge in the natural (N) and rehydroxylated (R) states)

Evolution du paramètre de désordre avec la proportion de charge tétraédriques dans les états naturel (N) et réhydroxyté (R).

References