Аl₃₀-ПИЛЛАРНЫЙ МОНТМОРИЛЛОНИТ С УЛУЧШЕННЫМИ ТЕКСТУРНЫМИ СВОЙСТВАМИ ОБУСЛОВЛЕННЫМИ ПРЕДВАРИТЕЛЬНОЙ МЕХАНИЧЕСКОЙ ОБРАБОТКОЙ
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Изучено влияние механической обработки природного монтмориллонита в планетарно-центробежной мельнице на эффективность интеркаляции полигидроксокомплексов алюминия [Al₃₀O₈(OH)₅(H₂O)]₁₈⁶⁺ при формировании пилларных структур. Измерения методом фотометрии показали, что в монтмориллонитовой матрице после интеркаляции полигидроксокомплексов Al₃₀ с использованием механической обработки наблюдалось увеличение содержания ионов Al³⁺ на 13%. По данным электрофоретического рассеяния света, размер частиц суспензии диспергированного в воде механоактивированного монтмориллонита составил около 100 нм. Исходный, механоактивированный и пилларный монтмориллонит охарактеризованы методами малоугловой дифракции рентгеновских лучей, сканирующей электронной микроскопии и низкотемпературной адсорбции-десорбции азота. Показано, что предварительная механоактивация исходного субстрата увеличивает базальное расстояние d₀₀₁ и существенно (примерно на 45-50%) повышает удельную площадь поверхности и суммарный объем пор Al₃₀-пилларного монтмориллонита; при этом возрастает как мезо-, так и микропористость, а размеры пор уменьшаются приблизительно на 12%. Установлено особая важность размеров частиц монтмориллонита при интеркаляции и дальнейшем формировании пилларной структуры. Уменьшение размера частиц монтмориллонита при механической обработке увеличивает площадь межфазной границы, через которую происходит ионный обмен. Показано, что малые размеры такождодов (около 100 нм) в водной суспензии играют ключевую роль в увеличении катионобменной способности механоактивированного монтмориллонита. В меньшей степени на эффективность интеркаляции влияют процессы дефектообразования и связанные с этим изменения электрических свойств силикатных слоев монтмориллонита. Исходя из особенностей структурных свойств, полученные пилларные материалы могут быть рекомендованы для использования в качестве селективных адсорбентов, молекулярных сит и катализаторов.
Ключевые слова: пилларный монтмориллонит, механическая активация, полигидроксокомплексы алюминия, ионы Кеггина, интеркаляция

Aл₃₀-PILLARED MONTMORILLONITE WITH ENHANCED TEXTURAL PROPERTIES DUE TO PRELIMINARY MECHANICAL TREATMENT
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The effect of mechanical treatment of natural montmorillonite in a planetary-centrifugal mill on the efficiency of intercalating aluminum polyhydroxocomplexes \([\text{Al}_{13}\text{O}_{5}(\text{OH})_{24}(\text{H}_2\text{O})_{26}]^{18+}\) in the formation of pillared structures was studied. Measurements made using the photometry method showed that in the montmorillonite matrix, after intercalation of the \(\text{Al}_{13}\) polyhydroxocomplexes using mechanical treatment, an increase in the content of \(\text{Al}^{3+}\) cations by 13% was observed. According to the electrophoretic light scattering data, the particle size for the suspension of mechanically activated montmorillonite dispersed in water was about 100 nm. The raw, mechanically activated and pillared montmorillonites are characterized by the methods of low-angle X-ray diffraction, scanning electron microscopy, and low-temperature nitrogen adsorption-desorption. It was shown that preliminary mechanical activation of the initial substrate increases the basal distance \(d_{001}\) and significantly (approximately by 45–50%) increases the specific surface area and the total pore volume of \([\text{Al}_{13}]\)-pillared montmorillonite; in this case, both meso- and microporosity increase, and the pore size decreases by about 12%. The special importance of the size of montmorillonite particles during the intercalation and further formation of the pillared structure is shown. A decrease in the size of the montmorillonite particles during mechanical treatment increasing the area of the interphase boundary through which ion exchange takes place. It was shown that the small sizes of tactoids (about 100 nm) in an aqueous suspension play a key role in increasing the cation exchange capacity of mechanically activated montmorillonite. To a lesser extent, the efficiency of intercalation is influenced by the processes of defect formation and the related changes in the electrical properties of the silicate layers of montmorillonite. Based on the structural properties, the obtained pillared materials can be recommended for use as selective adsorbents, molecular sieves and catalysts.

**Key words:** pillared montmorillonite, mechanical activation, aluminum polyhydroxocomplexes, Keggin-type ions, intercalation

**INTRODUCTION**

Recently, methods for obtaining new environmentally friendly polyfunctional nanomaterials on the basis of various layered systems have been intensively developed [1]. Intercalated layered systems are of great interest for the synthesis of sorbents and catalyst carriers, superionic conductors, optical and photoactive materials, nanomagnets, adsorbents, electrodes and membranes [2-3].

In particular, montmorillonite (MM) intercalated by polyhydroxocomplexes of various metals from solutions and sols enables obtaining pillared structures [4-6] by annealing: nanoparticles of oxides of various metals (pillars) are relatively uniformly distributed in the interlayer space and immobilized by crosslinking with silicate layers. Due to its increased basal distance \(d_{001}\) and regular distribution of nanoparticles pillared montmorillonite has unique textural and physico-chemical properties such as developed surface area, large volume of micro- and mesopores, thermal stability, and the presence of active centers of various nature.

Intercalation of the \([\text{Al}_{13}\text{O}_{5}(\text{OH})_{24}(\text{H}_2\text{O})_{26}]^{18+}\) ions (abbreviated as \(\text{Al}_{13}\)) [2, 4, 5, 7], which represent Keggin-type structures \([\text{XM}_{12}\text{O}_{40}]^{n-}\), is most well studied. Introducing \(\text{Al}_{13}\) ions into the interlayer space leads to an increase in the basal distance \(d_{001}\) of \(\text{Al}_{13}\)-pillared MM by a few angstroms compared to source MM and a significant increase in specific surface area and porosity [2, 4, 5]. It has been recently shown by us [8-9] and Jianxi et al. [10-11] that these characteristics can be further increased by intercalating "giant" aluminum polyacations \([\text{Al}_{30}\text{O}_{5}(\text{OH})_{50}(\text{H}_2\text{O})_{26}]^{18+}\) (\(\text{Al}_{30}\)) (consisting of two \(\text{Al}_{13}\) δ isomers and a bridge of four \(\text{AlO}_6\) octahedra) obtained by high-temperature hydrolysis of \(\text{Al}_{13}\).

It is obvious that textural characteristics of pillared materials can be improved by increasing the efficiency of intercalating polyhydroxocomplexes of metals. On the one hand, this can be achieved by using...
physical methods to activate intercalation, in particular, microwave radiation and ultrasonically [12-13]. Another approach proposed in the present work concerns preliminary mechanical activation of the layered substrate itself. The use of mechanoactivation in this case should be aimed at increasing, first, the specific surface area of clay particles and, second, their cation exchange capacity. In this case, a short-term mechanical treatment is necessary, which does not lead to a significant change in crystal structure of MM [14]. In our previous study [15] it was shown that preliminary mechanical treatment of natural MM increased substantially the textural properties of Al$_{13}$-pillared MM. A key role in increasing the capacity of the cation exchange of mechanically activated montmorillonite is played by the small dimensions of the tactoids (about 100 nm) in an aqueous suspension. To a lesser extent, the efficiency of intercalation is influenced by the processes of defect formation and associated changes in the electrical properties of silicate montmorillonite layers.

The goal of this work is to study the effect of preliminary mechanical treatment of natural MM on the efficiency of intercalating Al$_{30}$ ions and textural properties of the resulting pillared MM. For the sake of comparison with Al$_{13}$-pillared MM, all the conditions of mechanical treatment were kept the same as in [15].

EXPERIMENTAL

Source materials

The montmorillonite used in this work is the same as in [16] and has the following chemical composition, mass, %: SiO$_2$ – 57.70; TiO$_2$ – 1.04; Al$_2$O$_3$ – 13.75; Fe$_2$O$_3$ – 5.36; FeO – 0.20; CaO – 2.49; MgO – 3.13; Na$_2$O – 1.74; K$_2$O – 0.24; P$_2$O$_5$ – 0.16; SO$_3$ – 0.65; BaO – 0.08; loss on ignition – 13.46. The main "impurity" minerals are cristobalite, quartz, plagioclase, calcite and gypsum. The composition of exchangeable cations (mg-eq/100 g): Ca$^{2+}$ – 24.69; Mg$^{2+}$ – 22.74; (Na$^+$ + K$^+$) – 51.33, total – 98.76.

Mechanical treatment

Mechanically activated MM (hereafter referred to as AMM) was produced in an AGO-2C planetary centrifugal mill (Russia) for 3 min using high-strength zirconium grinding media at a constant rotor speed of 1500 rpm; the weight ratio for MM and grinding media is 7.5:1.

Preparation of intercalated and pillared samples

The solution containing Al$_{30}$ polycations was obtained by a hydrothermal technique, in which the solution containing Al$_{13}$ was kept at 127 °C for 5 h. The solution containing Al$_{13}$ polycations was prepared by hydrolysis of aluminum chloride: 0.2 M NaOH solution (Sigma Aldrich) was added to 0.2 M solution of AlCl$_3$·6H$_2$O (Fluka) until an [OH$^-$]/[Al$^{3+}$] molar ratio of 2.4 was obtained at pH 4.3-4.7 and room temperature. The solution was then aged for 24 h at 60 °C, resulting in the formation of Al$_{13}$ polyhydroxo complexes [5, 17-18]. Deionized water was used in all cases. The formation of the polycations was confirmed by the variety of methods as described in [9].

Intercalation of MM and AMM by Al$_{30}$ polycations was carried out by means of ion exchange in a 1% aqueous suspension by introducing an intercalating solution (3 mmol Al$^{3+}$/1 g MM) dropwise, while agitating intensively with a magnetic stirrer for 2 h at 80 °C. After 12 h of coagulation at room temperature, the suspension was washed to remove Cl$^-$ ions, centrifuged and dried in a drying oven at a temperature of 60 °C.

Pillared materials were obtained by annealing intercalated samples in an oven at 350 °C for 3 h. The designations used for these, like for Al$_{13}$ case [15], are as follows: Al$_{30}$-PMM, and Al$_{30}$-PAMM.

Research methods

Monitoring the efficiency of intercalating aluminum polyhydroxo complexes into studied materials was performed by a UV-Vis U-2010 spectrophotometer (Hitachi, Japan): the photometric technique is based on the ability of aluminum ion to yield an orange-red colored complex compound upon treatment with aluminon (tri(aminomethyl)amine) in a molar ratio [9] of 1:30 in a 0.65% aqueous solution. Measuring the basal distance $d_{001}$ of the samples by X-ray patterns was performed using a Bruker D8 Advance diffractometer (Bruker-AXS, Germany) with CuK$\alpha$ radiation ($\lambda = 0.154056$ nm) at 40 kV. Porosimetric measurements were carried out by low-temperature nitrogen adsorption-desorption using an ASAP 2020 specific surface area and porosity analyzer (Micromeritics, USA); prior to measurements, the samples were degassed at 180 °C and a residual pressure of 5-10 Pa for 3.5 h.

RESULTS AND DISCUSSION

The intercalation efficiency was evaluated by the photometric method, which makes it possible to determine the concentration of aluminum ions in the intercalating solution before and after completion of the intercalation and, accordingly, the aluminum content in the sample. Table 1 reveals that mechanoactivation increases the gross capacity of the cation exchange of montmorillonite.
Table 1

<table>
<thead>
<tr>
<th>Al³⁺ content</th>
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<tbody>
<tr>
<td>in the solution, mg/L</td>
</tr>
<tr>
<td><strong>Before intercalation</strong></td>
</tr>
<tr>
<td>MM</td>
</tr>
<tr>
<td>AMM</td>
</tr>
<tr>
<td><strong>After intercalation</strong></td>
</tr>
<tr>
<td>MM</td>
</tr>
<tr>
<td>AMM</td>
</tr>
</tbody>
</table>

Fig. 1 shows small-angle diffractograms of the investigated montmorillonite samples. A decrease in intensity and broadening of the reflex (001) after mechanical treatment of MM indicates some amorphization of the AMM structure and destabilization of the basal plane [14]. The increase in $d_{001}$ by about 0.3 nm in AMM (1.56 nm) compared to MM (1.26 nm) is due to the peeling of single layers after mechanoactivation [19]. Moreover, we believe that one should not rule out the diffusion of charge-compensating small-radius ions from the interlayer space into partially deformed and broken units of silicate layers, which should also result in an increase in $d_{001}$.

In Table 2 the basal distances of the Al₃₀-PMM and Al₃₀-PAMM are given along with those for previously studied Al₁₃-PMM and Al₁₃-PAMM [15] for the comparison. One can see that for both types of Keggin ions intercalated the values of $d_{001}$ are larger in the case of mechanical treatment. The Al₃₀-PAMM demonstrates the largest spacing of silicate layers, which is by 0.12 nm larger than that in Al₃₀-PAMM.

Fig. 2 shows the isotherms of low-temperature nitrogen adsorption-desorption for the obtained pillared materials. For all samples, nitrogen adsorption isotherms are of type IV according to the IUPAC classification and are characterized by a capillary-condensation hysteresis loop, which is typical for mesoporous materials. The shape of the hysteresis loop is of type H3, which is characteristic of porous materials with slit-shaped and plane-parallel pore [20]. Smaller hysteresis loop on the isotherms of Al₁₃-PAMM compared to Al₃₀-PAMM indicates the greater number of mesopores between the silicate layers. Al₃₀-PAMM shows higher adsorption capacity than Al₃₀-PMM.

![Fig. 1. Low-angle diffractograms: a) 1 - MM, 2 – Al₁₃-PMM; 3 – AMM, 4 – Al₃₀-PAMM](image1)

![Fig. 2. Nitrogen adsorption/desorption isotherms: 1 – Al₁₃-PMM, 2 – Al₃₀-PAMM](image2)

The specific surface area ($S_{\text{BET}}$) and total pore volume ($V_{\text{tot}}$) in both Al₁₃- [15] and Al₃₀-pillared samples are presented in Table 2. One can see that after mechanical treatment these characteristics increase substantially. From the data in Table 2 it follows that mechanical treatment in the Al₁₃-PAMM case resulted in significant increase in both micro- ($V_{\text{mic}}$) and mesoporosity ($V_{\text{meso}}$). In the Al₃₀-PAMM case the increase in mesoporosity due to mechanical treatment is even more pronounced whereas the volume of micropores somewhat decreases in comparison with untreated samples.
The pore size distribution curves for all the samples are shown in Fig. 3. Preliminary mechanical treatment of montmorillonite does not fundamentally change the unimodal pore size distribution, however, in this case the curve maximum shifts towards smaller pore sizes. This observation is in accordance with the notion that an increase in the number of intercalated polycations should lead to a decrease in the pore size.

### Table 2

<table>
<thead>
<tr>
<th>Sample</th>
<th>(d_{101}), nm</th>
<th>(S_{BET}), m²/g</th>
<th>(V_{mp}), cm³/g</th>
<th>(V_{imp}), cm³/g</th>
<th>(\sum V_{por}), cm³/g</th>
<th>(D_{av}), nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al₃⁺-PMM [15]</td>
<td>1.63</td>
<td>108</td>
<td>0.029</td>
<td>0.137</td>
<td>0.166</td>
<td>8.42</td>
</tr>
<tr>
<td>Al₃⁺-PAMM [15]</td>
<td>1.76</td>
<td>169</td>
<td>0.040</td>
<td>0.210</td>
<td>0.250</td>
<td>5.84</td>
</tr>
<tr>
<td>Al₃⁺⁻PMM</td>
<td>1.69</td>
<td>125</td>
<td>0.035</td>
<td>0.138</td>
<td>0.173</td>
<td>8.04</td>
</tr>
<tr>
<td>Al₃⁺⁻PAMM</td>
<td>1.88</td>
<td>182</td>
<td>0.031</td>
<td>0.231</td>
<td>0.262</td>
<td>7.11</td>
</tr>
</tbody>
</table>

**CONCLUSION**

The possibility of increasing the efficiency of intercalating the Al₃⁺ ions in the preparation of pillared structures by means of preliminary mechanical treatment of natural montmorillonite has been studied. In the case of mechanical treatment, an increase in the content of the Al³⁺ ions by 13% in the montmorillonite matrix after intercalating Al₃⁺ was observed. It has been established that preliminary mechanical treatment of montmorillonite allows one to significantly (approximately by 50%) enhance textural characteristics, in particular, to increase the specific surface area and total porosity of pillared materials; this increases mostly mesoporosity, and pore sizes decrease by approximately 12%. This result demonstrates the particular importance of the size of intercalated montmorillonite particles for the resulting porosity of the pillared material. A decrease in the size of the MM particles during mechanical treatment enables, first of all, increasing the area of the interphase boundary through which ion exchange takes place. Apparently, this factor becomes decisive in the process of intercalation and further formation of the pillared structure.

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