

## Interlamellar Silylation of Montmorillonite with 3-Aminopropyltriethoxysilane

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H-montmorillonite was modified by interlayer surface silylation using 3-aminopropyltriethoxysilane and dodecylamine in ethanol without a pre-swelling step. Dodecylamine acts as a gallery expander and silylation catalyst. The evaporation of ethanol from the dispersion yields well-ordered silylated montmorillonites with large basal spacing between 1.50 and 4.20 nm. Solid-state <sup>29</sup>Si CP MAS NMR of the silylated samples showed Q<sup>2</sup> and Q<sup>3</sup> signals as well as T<sup>2</sup> and T<sup>3</sup> signals. The increase in the relative intensity of Q<sup>3</sup> for Q<sup>2</sup> and the appearance of T<sup>2</sup> and T<sup>3</sup> signals was attributed to the grafting of 3-aminopropyltriethoxysilane to the interlayer surface silanol groups.

**Key Words :** Montmorillonite, Silylation, Intercalation, 3-Aminopropyltriethoxysilane

### Introduction

Chemical modification of silica surface is an area of intense interest from both fundamental and practical points of view.<sup>1-5</sup> The covalent grafting of organic units on inorganic surfaces have been attempted mainly for chromatographic application.<sup>6</sup> In recent years, the chemical modification into the interlayer surface of layered materials found increasing interests.<sup>7-17</sup> Organochlorosilanes, used widely for the modification of silica surfaces, have also been employed for interlayer modification of layered silicates,<sup>11,12</sup> leading to covalent grafting on interfaces. The covalent grafting of ethylene glycols<sup>13</sup> and silylation of various alkylchlorosilanes<sup>14-16</sup> into organo-layered silicates was reported.

This paper reports on the grafting of 3-aminopropyltriethoxysilane in interlayer surface of acid treated montmorillonite (HMMT). The silylation mechanism of 3-aminopropyltriethoxysilane into amorphous silica and alumina surface was well elucidated.<sup>18</sup> Interlamellar silylation of 3-aminopropyltriethoxysilane into montmorillonite was also conducted by refluxing in dry toluene.<sup>19</sup> Our synthetic strategy was focused on the simultaneous intercalation of 3-aminopropyltriethoxysilane and dodecylamine into HMMT in ethanol without a pre-swelling step. Intercalation and silylation was easily achieved by direct evaporation of ethanol from the dispersion of HMMT in 3-aminopropyltriethoxysilane, dodecylamine and ethanol. Attached amine groups on the surface can offer reactive sites for chemical bonds with epoxy, nylon, imide, and urethane polymers. In particular, layered materials with attached amine groups also may be used as adsorbents of heavy metal ions, supports of drugs or enzymes and photosensitive devices.

### Experimental Section

**Materials.** The silane coupling agent was APS [3-aminopropyltriethoxysilane: NH<sub>2</sub>-CH<sub>2</sub>-CH<sub>2</sub>-CH<sub>2</sub>-Si(OC<sub>2</sub>H<sub>5</sub>)<sub>3</sub>] of special grade (Aldrich, USA). Dodecylamine (DDA) was used special grade (Aldrich, USA). HCl and 95%-ethanol were used. MMT (montmorillonite) was purchased from YAGURI (Japan). The commercial MMT was purified using a sedimentation method. 2 wt% MMT suspension was prepared in deionized water and stirred overnight at room temperature, and then allowed to settle down for 2 h. The sediment was removed by decantation. The purified MMT was then recovered by centrifuging the decanted upper slurry, washed and oven dried for 24 h at 100 °C.

HMMT was prepared by an ion exchange of exchangeable cations for H<sup>+</sup> in a 0.1 N HCl solution. The suspension composed of purified MMT (40 g) and deionized water (500 mL) was slowly titrated with a 0.1 N HCl solution to a final pH of 1.8 and then maintained at the same pH value for an additional 24 h. The sample was recovered by filtering, washed with deionized water to be Cl<sup>-</sup>-free, and then dried in air for 24 h at 100 °C. The replacement of exchangeable cations by H<sup>+</sup> in the layered silicate produces organophilic properties in the interlayer surfaces because the silanol groups produce hydrogen-bonding sites.

**Sample preparation.** APS solutions were prepared to be 0.01 M-0.1 M in ethanol (95%). HMMT (1.0 g) was dispersed in 10 mL of these solutions (ultrasound for 20 min) at room temperature. The dispersion was evaporated to dryness for 24 h at 50 °C (APS-MMT). To investigate swelling property of APS-MMT, 1.0 g samples were dispersed in 0.5 M DDA-ethanol solution (10 mL) and dried by evaporating for 24 h at 50 °C (APS-MMT-DDA).

DDA-APS mixed solutions was also prepared by dissolving APS and DDA in ethanol (95%). The concentrations of APS in solution was 0.01 M-0.1 M, concentration of DDA was 0.1 M-0.5 M. HMMT (1.0 g) was then dispersed in 10 mL of these solutions (ultrasound, 20 min, at room

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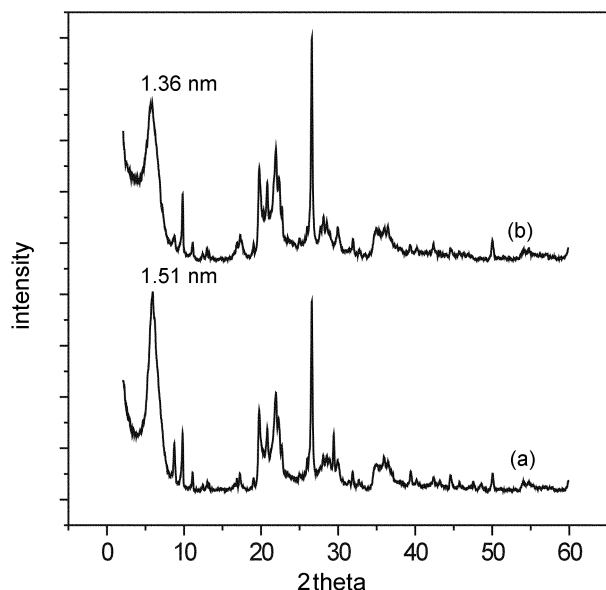
temperature) and evaporated to dryness for 24 h at 50 °C (DDA-APS-MMT).

**Characterization.** Powder X-ray diffraction measurements were recorded on a Rigaku Rotaflex 200B diffractometer equipped with  $\text{CuK}\alpha$  X-ray radiation and a curved crystal graphite monochromator. Solid-state  $^{29}\text{Si}$  MAS NMR spectra were recorded on a Bruker CXP-100 spectrometer at a resonance frequency of 19.89 MHz with a 45° pulse and a recycle delay of 7 s. Energy dispersive X-ray spectrometer analysis (EDS, Link system AS1000-85S) and thermo gravimetric analysis (TGA, 10 °C/min to 900 °C, 100 cc/min  $\text{N}_2$  purge) were performed.

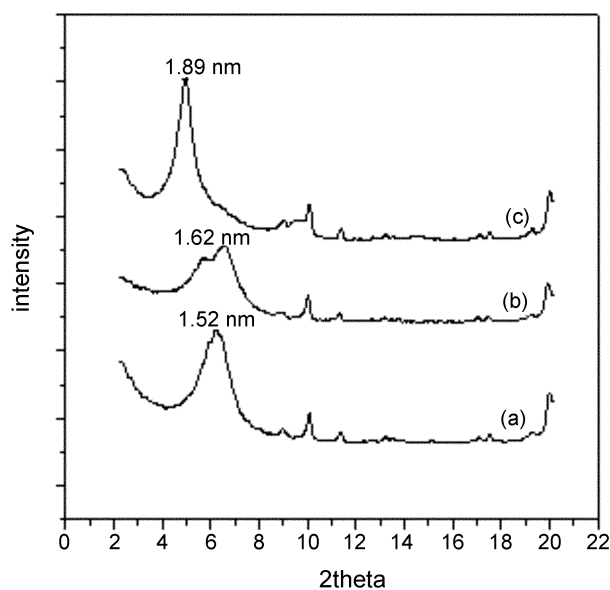
### Results and Discussion

The X-ray powder diffraction pattern of purified MMT exhibits several (001) reflections, indicating a basal spacing of 1.51 nm (Figure 1a). This is larger than that (1.23 nm) of commercial MMT. It is attributed to solvation of interlayer cations by water molecules. Gillery<sup>20</sup> reported that the basal spacing of clay minerals is closely related to the relative humidity. Air-dried HMMT is very similar to MMT except for a little decrease in basal spacing (Figure 1b). This indicates that the original structure of MMT does not change by the acid treatment. Decrease in the basal spacing indicates a loss of the interlayer  $\text{H}_2\text{O}$  upon the replacement of exchangeable cations for  $\text{H}^+$ . Analysis of EDS showed that the intrinsic Na and K peak in purified MMT disappeared in HMMT. This proves that the acid treatment of MMT results in complete exchange of exchangeable cations for  $\text{H}^+$ .

Figure 2 shows the XRD patterns for HMMT treated with 0.01, 0.05 and 0.1 M APS-ethanol solutions. During this process, the APS molecules were intercalated and grafted to interlayer surface silanol groups. The basal spacing of the APS-MMT increases 1.52 nm to 1.89 nm with increased



**Figure 1.** XRD pattern of (a) purified MMT and (b) HMMT.

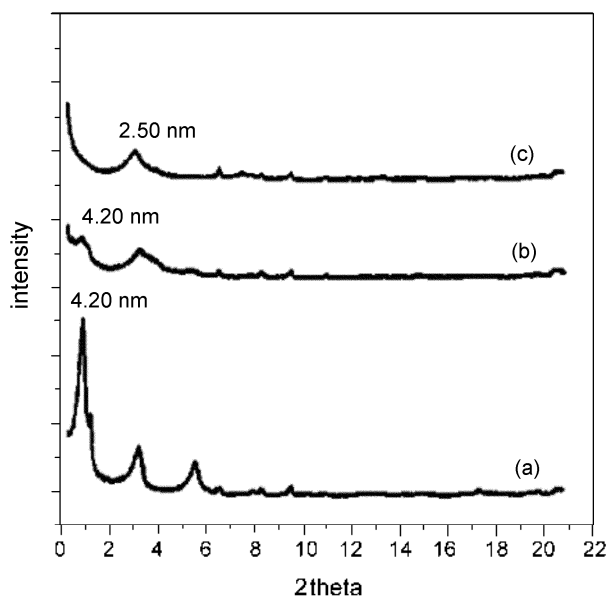


**Figure 2.** XRD patterns of HMMT treated with APS-ethanol solution. (a) 0.01M APS-MMT. (b) 0.05M APS-MMT and (c) 0.1M APS-MMT.

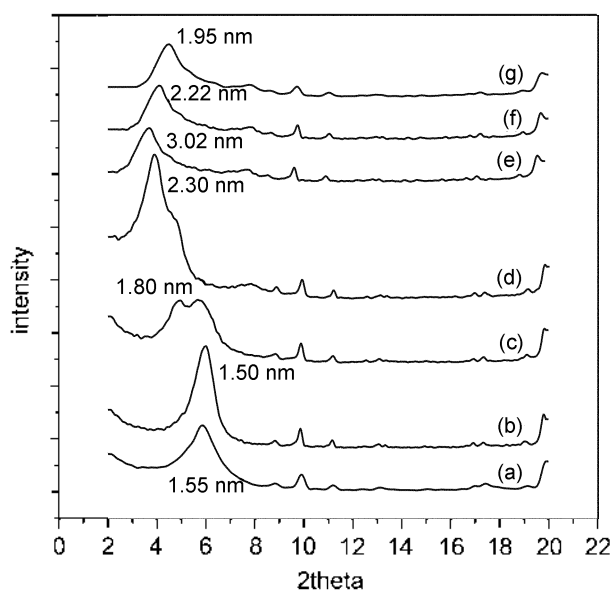
APS concentration, allowing large expansion (from 0.11 nm to 0.53 nm) of interlayer space. This indicates that the APS molecules arrange in monolayers or paraffin-type between the aluminosilicate layers. The interlayer space, at lower concentration (0.01 M and 0.05 M), cannot intercalate enough to fill with APS molecules, resulting in only monolayer or double layer arrangement. In case of this, APS can build a bridge between adjacent layers by silylating with silanol groups in adjacent interlayer surface (interlayer condensation). This interlayer condensation may lose swelling property of APS-MMT. However, at higher concentration (0.1 M), the interlayer space expands to molecular length (~0.7 nm) of APS because APS molecules fill with enough to allow paraffin-type. This shows that APS grafted-MMT with large basal spacing can be prepared simply by evaporation of solvent from the dispersion of HMMT in APS-ethanol solution. In general, grafting of APS on the silica surface was conducted by filtering after reaction for several hours in toluene comprising excess APS.

Figure 3 shows the swelling property of APS-MMT in 0.5 M DDA-ethanol solution. HMMT in DDA-ethanol solution showed large increase of the basal spacing to 4.6 nm after drying. This indicates that DDA molecules are arranged in a bilayer structure. APS-MMT (0.01 M) showed excellent swelling properties in DDA, resulting in well ordered intercalates with a basal spacing of 4.2 nm. Swelling by DDA of APS-MMT implies that interlayer condensation does not happen between adjacent interlayer surfaces. However, increasing concentration of APS (0.05 and 0.1 M) broadens the reflections and decreases the basal spacing because a higher degree of grafting reduces the amount of intercalated DDA.

The treatment of HMMT with APS-DDA mixed solution ensures successful silylation of APS. The dried powders

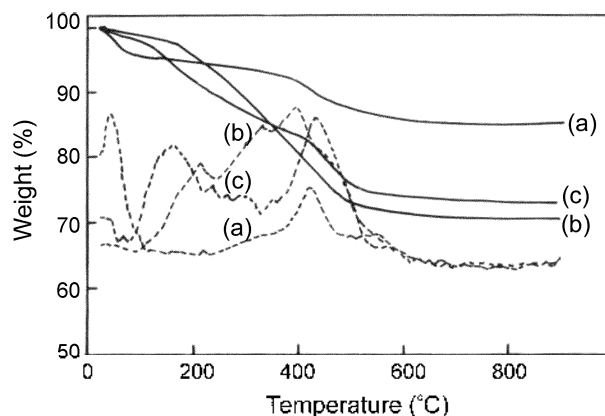


**Figure 3.** XRD patterns of APS-MMT treated with 0.5 M DDA-ethanol solution. (a) 0.01APS-MMT, (b) 0.05APS-MMT and (c) 0.1APS-MMT.



**Figure 4.** XRD patterns of HMMT treated simultaneously with APS and DDA in ethanol. (a) 0.01DDA-0.01APS-MMT, (b) 0.05DDA-0.05APS-MMT, (c) 0.1DDA-0.1APS-MMT, (d) 0.3DDA-0.01APS-MMT (e) 0.5DDA-0.01APS-MMT (f) 0.5DDA-0.05APS-MMT, and (g) 0.5DDA-0.1APS-MMT.

consist of HMMT with intercalated APS and DDA. Figure 4 shows that the basal spacing of DDA-APS-MMT samples exhibits large difference of 1.55 nm to 3.02 nm, depending upon concentration of DDA and APS. Increasing concentration of DDA results in large increase in basal spacing (a, d and e; b and f). In particular, the intercalation of DDA is influenced by co-intercalate APS (e, f and g). More ordered and larger basal spacing (e) result from the sample (0.5DDA-0.01APS) prepared in solution with the largest

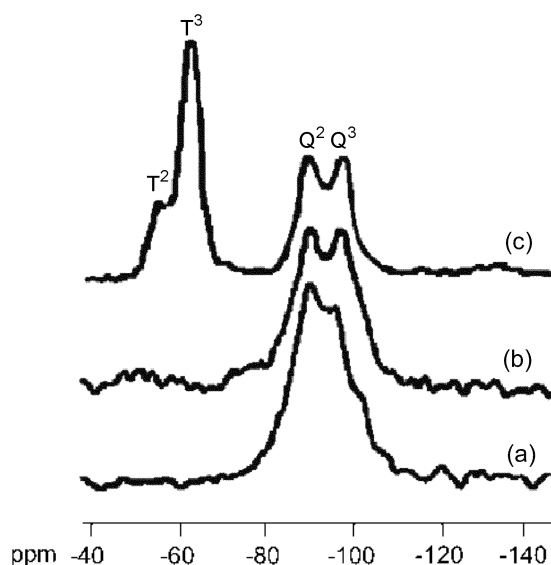


**Figure 5.** The thermogravimetric analysis of (a) 0.1APS-MMT, (b) 0.5DDA-MMT, and (c) 0.5DDA-0.1APS-MMT.

difference in relative concentration of DDA and APS. However, increasing concentration of APS (c, g) broadens the reflections and decreases the basal spacing because a higher degree of grafting reduces the amount of intercalated DDA.

The TGA data obtained from 0.5DDA-MMT, 0.1APS-MMT and 0.5DDA-0.1APS-MMT are illustrated in Figure 5. 0.5DDA-MMT appears weight losses at broad range between 100 °C and 500 °C, caused by desorption and combustion of intercalated DDA. However, APS grafted-MMT exhibits two-step weight loss. The weight loss below 100 °C, in 0.1APS-MMT, is from desorption of ethanol molecules and the loss occurring after 400 °C is from combustion of the silylated APS. Two-step weight loss, in 0.5DDA-0.1APS-MMT, is attributed to different bonding environment of DDA and APS in the gallery. The initial weight loss below 200 °C is from desorption of non-intercalated or intercalated DDA molecules and loss after 400 °C is from combustion of the silylated APS.

The solid-state  $^{29}\text{Si}$  CP MAS NMR spectra for silylated HMMT show  $\text{Q}^2$  (---90 ppm),  $\text{Q}^3$  (---96 ppm),  $\text{T}^2$  (---50 ppm) and  $\text{T}^3$  (---60 ppm) signals due to diverse environments of silicon (Figure 6). In general, MMT exhibits a resonance in the range from -75 to -95 ppm, which are typical of several additional silicon environments.<sup>21,22</sup> Each Si atoms in  $\text{SiO}_4$  structures have two or three Si atoms as nearest neighbors ( $\text{Q}^2$ ,  $\text{Q}^3$ ) and remaining nearest neighboring atom/atoms being either Al or H.  $\text{Q}^2$  is attributed to central Si in  $(\text{Si-O})_2\text{Si}(\text{O-Al})\text{OH}$  and  $\text{Q}^3$  in  $(\text{Si-O})_3\text{Si}(\text{O-Al})$ . The increase of  $\text{Q}^3/\text{Q}^2$  ratio compared with HMMT clearly indicates the grafting of APS molecules to surface silanol groups. New peaks ( $\text{T}^2$  and  $\text{T}^3$ ), near ---50 to ---60 ppm, in 0.3DDA-0.01APS-MMT are attributed to the Si atoms of the grafted APS molecules. The Si signals for APS silylated on silica appeared in the range of -49 ppm to -68 ppm depending upon bonding type.<sup>23</sup> Si signals for alkyltri-chlorosilane grafted on layered silicates was also appeared at ---56 to ---65 ppm.<sup>16,17</sup> Here, all the APS molecules may not silylate with silicate surface because APS could hydrolyze in water.



**Figure 6.**  $^{29}\text{Si}$  CP MAS NMR spectra of (a) HMMT, (b) 0.1APS-MMT and (c) 0.3DDA-0.01APS-MMT.

Above results show that the silylation by APS and intercalation of DDA proceeded more effectively by evaporation of the solvent from the dispersion of HMMT, APS, DDA and ethanol. Amine groups of APS and DDA molecules are very accessible to silanol groups in interlayer surface. Solvent also solvates interlayer Si-OH groups and makes easy intercalation of DDA and APS into interlayer. The solvation of DDA and APS within gallery results in additional expansion in gallery. DDA and APS molecules in gallery can have fully stretched arrangement. Evaporation of ethanol starts at the outside of particle, resulting in a higher concentration of DDA and APS on the external surface. This promotes intercalation of APS and DDA. The evaporation of residual ethanol in the gallery results in silylation of APS. The water molecules contained in ethanol catalyze interlayer surface silylation of APS during the evaporation of ethanol. The condensation of alkoxy silane is known to catalyze by base. DDA as gallery expander can also have a role as catalyst for the grafting of APS into the surface silanol groups. This evaporation process should be very efficient method for intercalation and silylation of silane coupling agents into layered materials and mesoporous materials.

### Conclusions

MMT with attached amine group in interlayer surface was

prepared successfully. The process proceeded more effectively by evaporation of the solvent from the dispersion of HMMT, APS, DDA and ethanol. APS was intercalated directly into interlayer space of acid-treated MMT and silylated with interlayer silanol groups, resulting in the grafting of APS on interlayer surfaces. DDA could have a role as gallery expander or silylation catalyst. Ethanol could control quantitatively the amount of APS and DDA needed to the gallery silylation and the water contained in ethanol catalyze interlayer surface silylation.

### Reference

1. Plueddemann, E. P. *Silane Coupling Agents*; Plenum: New York, 1982.
2. Sung, C. S. P.; Lee, S. H.; Sung, N. H. *Polym. Sci. Technol.* **1980**, *12B*, 757.
3. Kulkarni, R. D.; Goodard, E. D. *Int. J. Adhes. Adhes.* **1980**, *1*, 73.
4. Furukawa, T.; Fib, N. K.; Mittal, K. L.; Anderson, H. R. *J. Colloid Interface Sci.* **1983**, *96*, 322.
5. Alexander, J. D.; Gent, A. N.; Henriksen, P. N. *J. Chem. Phys.* **1985**, *83*, 5981.
6. Pesck, J. J. In *Chemically Modified Oxide Surfaces*; Leyden, D. E., Collins, W. T., Eds.; Gordon & Breach: New York, U. S. A., 1990; Vol. 3, p 93.
7. Kwon, O. Y.; Park, K. W. *Bull. Korean Chem. Soc.* **2003**, *24*, 1561.
8. Kwon, O. Y.; Park, K. W. *Bull. Korean Chem. Soc.* **2004**, *25*, 25.
9. Kim, J. W.; Liu, F.; Choi, H. J. *J. Ind. Eng. Chem.* **2002**, *8*, 399.
10. Kwon, O. Y.; Park, K. W. *J. Ind. Eng. Chem.* **2004**, *10*, 252.
11. Ruiz-Hitzky, E.; Rojo, J. M. *Nature* **1980**, *287*, 28.
12. Ruiz-Hitzky, E.; Rojo, J. M.; Lagaly, G. *Colloid Polym. Sci.* **1985**, *263*, 1025.
13. Tunny, J. M.; Detellier, C. *Chem. Mater.* **1993**, *5*, 747.
14. Ogawa, M.; Okutomo, S.; Kuroda, K. *J. Am. Chem. Soc.* **1998**, *120*, 7361.
15. Isoda, K.; Kuroda, K.; Ogawa, M. *Chem. Mater.* **2000**, *12*, 1702.
16. Shimojima, A.; Mochizuki, D.; Kuroda, K. *Chem. Mater.* **2001**, *13*, 3603.
17. Thiesen, P. H.; Beneke, K.; Lagaly, G. *J. Mater. Chem.* **2002**, *12*, 3010.
18. Impens, N. R. E. N.; Vort, V. P.; Vansant, E. F. *Microporous and Mesoporous Mater.* **1999**, *28*, 217.
19. Mukkanti, K.; Subba Rao, Y. V.; Choudary, B. M. *Tetrahedron Letters* **1989**, *30*, 251.
20. Gillery, F. H. *Am. Mineralog.* **1959**, *44*, 806.
21. Thompson, A. R.; Botto, R. E. *Energy & Fuels* **2001**, *15*, 176.
22. Carrado, K. A.; Xu, L.; Gregory, D. M.; Song, K.; Seifert, S.; Botto, R. E. *Chem. Mater.* **2000**, *12*, 3052.
23. Caravajal, G. S.; Leyden, D. E.; Quinting, G. R.; Maciel, G. E. *Anal. Chem.* **1988**, *60*, 1776.