kaolinite Interlamellar amino functionalization of

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analysis, ¹³C CP/MAS NMR spectroscopy, and elemental analysis, a structural model is proposed in which every third interlayer **Abstract**: The interlamellar surface of kaolinite has been modified with molecules possessing amino functionalities. Either the dimethyl sulfoxide intercalate of kaolinite (Kao/DMSO) or the *N*-methylformamide intercalate of kaolinite (Kao/NMF) were used as starting materials. One of the products, an ethanolamine functionalized kaolinite (Kao–EOA) was resistant to thermal an amino-functionalized ordered two-dimensional organo-mineral assembly. groups that point away from this surface are each keyed into the $-(SiO_{-})_{6}$ macro-rings of the adjacent silicate surface, resulting in group or is strongly H-bonded to an aminoalcohol molecule. A mixture of both types of linkages could coexist. The amino surface hydroxyl group on the aluminol (Al-OH) surface of kaolinite is either replaced with an interlayer Al-OCH $_2$ CH $_2$ NH $_2$ decomposition in both air and N₂ atmospheres up to temperatures greater than 150°C. Based on results from thermal analysis, IR

Key words: kaolinite, halloysite, organo-mineral nanocomposites, clay functionalization, supramolecular assemblies

(Kao/NMF). Sur l'un des produits, une kaolinite fonctionnalisée à l'éthanolamine (Kao–EAO) est résistante à la décomposition thermique, jusqu'à des températures supérieures à 150°C, dans des atmosphères tant d'air que de N₂. En se basant sur les résultats d'analyses thermiques, d'analyses élémentaires, de spectroscopies IR et RMN ¹³C avec polarisation croisée/rotation à adjacente de silicate qui conduit à un arrangement organominéral aminofonctionnalisé ordonné d'une façon bidimensionnelle. **Résumé**: On a modifié la surface interlamellaire de la kaolinite avec des molécules portant des fonctions amino. Comme produits de départ, on a utilisé de la kaolinite intercalée de diméthylsulfoxyde (Kao/DMSO) ou de N-méthylformamide Les groupes amino qui sont orientés vers l'extérieur de cette surface sont insérés dans les macrocycles de -(SiO-)₆ de la surface fortement liés, par liaisons hydrogènes, à une molécule d'aminoalcool. Il est possible que les deux types de liaisons coexistent. l'intercouche de l'aluminol (Al-OH) de la kaolinite sont soit remplacés par une intercouche de groupes Al-OCH₂CH₂NH₂ soit angle magique, «CP/MAS», on propose un modèle structural selon lequel un sur trois groupes hydroxyles de la surface de

supramoléculaires. Mots clés: kaolinite, halloysite, nanocomposites organominéraux, fonctionnalisation de l'argile, arrangement

[Traduit par la rédaction]

Introduction

It has previously been shown that it is possible to rigidly fix small alcohol moieties onto the interlamellar aluminate surface of kaolinite $(Al_2Si_2O_5(OH)_4)$, through the grafting reaction of the alcohol group to the Al-O-H surface of the kaolinite (1–3). These interesting materials are organo-mineral nanocomposites with organic units covalently attached to the

Received May 16, 1997

This paper is dedicated to Professor Brian E. Conway on th occasion of his 70th birthday.

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- With the diol-based organo-kaolinites, the interlayer cohesive forces for these new materials prevented easy layer swelling (1, 2). This was presumably due to the strong hydrogen bonding between the pendant hydroxyl groups of the diols and the silicate surface of the kaolinite layers. As a result, even though the alcohol functionality of the diol unit offers an

as polyethylene glycol of average molecular weight 3400, into the interlayers of kaolinite by first preswelling with either DMSO or NMF (17). This suggests that kaolinite could be systems than the previously reported alcohol/diol organoused as a non-centrosymmetric inorganic organizing medium. it is nonetheless possible to intercalate large molecules, such the slow kinetics of many kaolinite intercalation reactions (8), kaolinite system (1-3). It has been demonstrated that despite the interlayer grafting chemistry of kaolinite to include other orientation (1-3, 7-16). It would thus be of interest to expand kaolinite (4-6) may act to orient molecules in a preferred dipole moment that exists rigid inorganic supramolecular organizing medium, since the inorganic substrate in an ordered stacking of organo-mineral layers. The non-centrosymmetric kaolinite layers may act as a inorganic asymmetric kaolinite layers. non-centrosymmetric organo-mineral materials based on rigid One may thus envisage rationally assembling sophisticated across the interlayer space of

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consequently greater swelling properties, are required if one wishes to use them as the first step in the development of more sophisticated structures. Part of the appeal in attaching molecules with pendant amine groups in the interlayer space of kaolinite lies with the possibility of using the free amine group Amines could, for example, provide an ideal site for the attachment of a carbonyl-functionalized moiety. to attach larger molecules to the interlayer surface kaolinite. kaolinites, preferably with weaker interlayer cohesion and inaccessible to reactant molecules. Other types of organoattractive site for further interlayer modification reactions, it is

kaolinite to hydrazine vapours or to a hydrazine solution. Despite its ease of formation, Kao/Hz is much less stable than either Kao/DMSO or Kao/NMF, decomposing in ambient conditions within less than an hour. This ease of formation and inherent instability allows Kao/Hz to be used to advantage as hmite materials in which they proposed that the organic units were attached to the boehmite via $Al-O-(CH_2)_nNH_2$ bonds. et al. (24) prepared a series of aminoalcohol-based organoboean intermediate in the preparation of other organo-kaolinites via the displacement method (21-23). In related work, Inoue (Kao/Hz) (8, Organo-kaolinites with organic guests possessing amino groups have previously been reported. Kaolinite-hydrazine 18-21) is rapidly formed by exposing untreated

Experimental

before proceeding with the purification procedure. For both kaolinite and halloysite, purification by standard sedimentation procedures to remove the >2 µm size fraction was required (25–27). Even after purification, in the case of kaolinand were found to be consistent with literature data for purified natural kaolinite and halloysite (28, 29). clays were then characterized by XRD, FTIR, and TGA/DSC responding to the (101) reflection of anatase. The purified detected by XRD as a sharp weak reflection at d = 3.52 Å, corite small amounts of anatase (TiO2) impurities could be (Rochester, N.Y.) in the form of hard rocklike chunks. This was obtained from Ward's Natural Science Establishment Inc. Halloysite no. 13 from Eureka, Utah), a polytype of kaolinite, tory of the Clay Mineral Society (Department of Geology, University of Missouri, Columbia, Mo.). Halloysite (coded as tallized kaolinite, was obtained from the Source Clay Reposi-All chemicals used were of reagent grade quality and were not further purified unless otherwise specified. KGa-1, well-cryshad to first be ground up to the consistency of a coarse powder

Characterization
XRD powder patterns were measured on a Philips PW 3710 automated diffractometer using Cu Kα radiation (wavelength pension to dry on the glass disk. Some samples were mounted dry by first smearing a small quantity of vaseline to the glass 1 mL of methanol, sonicating for 10 s, and allowing the suscircular glass disk by first dispersing 20-30 mg of sample with without employing any mask. Samples were mounted on a automatic divergent slit and a 0.1 mm receiving slit were used step. The sample was spun during pattern acquisition, and an of 0.04° 20 was used with a dwell time of 0.5 seconds per = 1.5416 Å). All measurements were taken using a generator voltage of 45 kV and a generator current of 40 mA. A step size

> (001) reflections of the modified organo-kaolinite phase and the unexpanded kaolinite phase (8). The c-spacing or basal spacing is calculated based on the indexed (00*l*) reflections also given in parentheses. intense reflection. The assigned indexing of the reflection is kaolinite are identified with the symbol K. All d-values are given in Ångstrom units (Å), with the relative intensities given (8). Reflections due to the presence of residual unreacted ification, was calculated from the relative intensities of the disk and then placing the powder directly on this. The interca-lation ratio (I.R.), which is an indication of the extent of modin parentheses on a scale where 100 is the intensity of the most

or air (20–90 cm³/min) and a heating rate of 20°C/min. Approximately 10 mg of sample was used for each run using alumina sample and reference pans. residual mass (as a percentage of the starting mass) of the product after calcination at 1100°C for 3 h in air atmosphere. Thermal analyses (TGA and DSC) runs were performed on a Polymer Labs 1500H instrument under either flowing nitrogen ple holder, with the scans referenced to a mixture of SiO_2 and ments to measure the Si/Al ratio were performed on a Philips PW 2400 fluorimeter equipped with a Philips PW 2510 sam-0.50% by weight in KBr). X-ray fluorescence (XRF) measure-100 FTIR spectrometer using 30–50 averaged scans at 4 cm⁻¹ resolution. The samples were prepared as KBr pellets (0.25– Al_2O_3 . Ceramic yields were determined by measuring the Infrared spectra were obtained on a Bomem Michelson MB

¹³C cross polarization/magic angle spinning (CP/MAS) spectra (50.33 MHz) were performed on a Bruker ASX-200 instrument with a magic angle spinning rate of 3–4 kHz. The ¹³C CP/MAS spectra were referenced to hexamethylbenzene at 16.9 ppm. Solution state NMR (¹H and ¹³C) were performed on either a Varian XL-300 or a Bruker AMX-500, and both ¹H and ¹³C chemical shifts are reported relative to TMS in CDCl₃.

Starting materials

based on standard literature procedures (1, 2, 8). Briefly, this involves allowing 20 g of the unexpanded purified kaolinite to remain in contact with DMSO or NMF in an enclosed jar for at least 2 months at room temperature, followed by washing with 1,4-dioxane to remove excess DMSO or NMF, and air drying. After this time, both Kao/DMSO and Kao/NMF are found by XRD to be formed with an intercalation ratio (I.R.) exceeding 90%. XRD patterns, IR spectra, and TGA were all consistent with literature data (8). The DMSO intercalate of this case, no residual unexpanded halloysite was observed by halloysite (Hal/DMSO) was formed in an identical manner. In starting materials have been described elsewhere and are The preparations of both Kao/DMSO and Kao/NMF and XRD (8)

Reactions with amines

Kao/DMSO + ethanolamine(Rxn I)

This was then filtered and washed with 1,4-dioxane to remove excess EOA. Air drying at 40°C for 2 h yielded 0.76 g of an distilled ethanolamine (EOA) for 22 h under N₂ atmosphere. resulted. The following is a typical example: 1.0 g of Kao/DMSO (I.R. = 0.97) was refluxed with 50 mL of freshly number of times and in all cases nearly identical materials The reaction of Kao/DMSO with ethanolamine was repeated a

off-white powder. XRD: 10.64 (100, *d*₀₀₁); 7.14 (1.5, *d*₀₀₁K); 5.34 (0.1, *d*₀₀₂); 4.46 (1.4); 4.34 (2.3); 4.28 (2.1); 4.20 (2.0); 4.15 (1.4); 3.72 (1.2); 3.56 (16.5, *d*₀₀₃); 3.32 (0.5); 3.19 (1.2); 3.14 (0.9); 2.84 (0.1); 2.67 (0.7, *d*₀₀₄); 2.13 (1.2, *d*₀₀₅); 1.78 (0.2, *d*₀₀₆); 1.487 (0.3, *d*₀₀₆). *d* = 10.67 ± 0.03 Å, I.R. = 0.985. FTIR (cm⁻¹), ν(O-H): 3693 (s), 3628 (s), 3558 (m), 3486 (s) ν(NH): 3354 (m), 3311 (m); ν(C-H): 3074 (w, br), 2964 (νw), 2946 (w), 2935 (νw), 2889 (w); δ(HOH): 1663 (w); δ(C-H, N-H): 1613 (w), 1525 (w), 1476 (w), 1353 (w), 1197 (w); Si-O vibrations: 1125 (s), 1092 (s), 1045 (vs), 1016 (vs); δ(AIOH): 974 (m), 908 (s); other bands: 2792 (vw), 2691 (w), 2612 (w), 1385 (w, impurity), 874 (w), 796 (w), 740 (w), 683 (m), 604 (w, sh), 552 (s), 471 (s), 433 (s). ¹³C CP/MAS NMR (Bruker CXP-180, spinning speed = 6 kHz, 45.27 MHz, δ(ppm)): 42 (-CH₂NH₂), 60 (-CH₂O-); no dipolar dephasing signal detected after 40 μs dephasing time for either resonance. Elemental analysis: %C 8.06, %N 4.12; %H 2.72, ceramic yield 72.5%. Calculated for Al₂Si₂O₅(OH)_{3,0}(OCH₂CH₂NH₂)_{1,0}: %C 7.52, %N 4.39, %H 3.45, ceramic yield 69.6%. Calculated for Al₂Si₂O₅(OH)_{3,0}(OCH₂CH₂NH₂)_{1,0}: %C 7.97, %N 4.65, %H 2.99, ceramic yield 73.8%.

Kao/NMF + ethanolamine (Rxn 2)

excess ethanolamine. Air drying yielded 20 g of an off-white powder. The FTIR and XRD patterns proved to be nearly identical to product (1). XRD (vaseline mounted, vacuum dried at 40°C for 3 h): 10.36 (100, d_{001}); 7.02 (3.7, $d_{001}\text{K}$); 4.41 (7.5); 4.31 (9.8); 4.25 (11); 4.15 (9.1); 4.00 (5.6); 3.69 (5.9); 3.53 (23, d_{003}); $d = 10.5 \pm 0.2 \text{ Å}$; I.R. = 0.96. Kao/NMF $(2.0\,\mathrm{g})$ (I.R. = 0.91) was refluxed with 60 mL of eth-anolamine (freshly distilled) for 23 h under N₂ atmosphere. This was then filtered and washed with 1,4-dioxane to remove

Kaolinite + ethanolamine (Rxn 3)

 $0.9~{\rm g}$ of a light grey product. The FTIR pattern of this product was indicative of the well-crystallized kaolinite starting mate-Kaolinite $(1.0\,\mathrm{g})$ was refluxed with 50 mL of ethanolamine in a round-bottom flask fitted with a reflux condenser for 17 h mixture with methanol, the product was air dried, yielding under N₂ atmosphere. After filtering and washing the reaction

amine. Air drying yielded 0.70 g of an off-white powder. XRD: $10.61 (100, d_{001})$; $3.55 (13, d_{003})$. $d = 10.6 \pm 0.1$ Å; I.R. = 1.0. FTIR (cm⁻¹), ν (O-H): 3695 (s), 3630 (s), 3558 (s), 3491 (s); ν (NH): 3356 (m), 3313 (m); ν (C-H): 3076 (w, br), 2948 (w), 2889 (w); ν (C = O): 1614 (m); δ (C-H, N-H): 1478 (vw), 1355 (vw); Si-O vibrations: 1125 (s), 1035 (vs); δ (A1OH): 908 (s); other bands: 2620 (w), 744 (w), 680 (m), 551 (s), 468 (s), Hal/DMSO + ethanolamine (Rxn 4) Hal/DMSO (10 g) was refluxed with 40 mL of ethanolamine (freshly distilled) for 20 h under N_2 atmosphere. This was then filtered and washed with ethanol to remove excess ethanol-

round-bottom flask fitted with a reflux condenser for 18 h Kao/DMSO + ethylene diamine (Rxn 5) Kao/DMSO (1.2 g) (I.R. = 0.97) was refluxed with 60 mL reagent grade ethylene diamine (Aldrich, BP 118°C) in a mixture with 1,4-dioxane, the product was air dried, yielding under N₂ atmosphere. After filtering and washing the reaction

the treatment of amines with kaolinite. Table 1. Summary of the organo-kaolinite products formed from

	NR	8	Nitroaniline	Kao/DMSO
	NR	7	DL-1-Amino-2-propanol	Kao/DMSO
0.58	11.1	6	3-Amino-1-propanol	Kao/DMSO
	NR	Ŋ	Ethylene diamine	Kao/DMSO
1.0	10.6	4	Ethanolamine	Hal/DMSO
	NR	ယ	Ethanolamine	Kaolinite
0.96	10.5	2	Ethanolamine	Kao/NMF
0.98	10.7	1	Ethanolamine	Kao/DMSO
I.R.	Basal spacing ^b (Å)	Rxn code"	Reactant	Starting material

0.8 g of a light grey product. The FTIR pattern of this product was indicative of partially collapsed Kao/DMSO.

Kao/DMSO + 3-amino-1-propanol (Rxn 6)

patterns were indicative of a partially modified product. XRD: 11.07 (90, d_{001}), 7.10 (66, d_{001} K). I.R. = 0.58. FTIR (cm⁻¹), ν (O-H): 3698 (s), 3652 (m), 3622 (s), 3626 (w, br); ν (NH): 3364 (vw); ν (C-H): 2941 (w), 2878 (w); δ (C-H, N-H): 1574 (w), 1482 (w), 1322 (vw); Si-O vibrations: 1110 (s, sh), 1085 (s), 1046 (vs), 1032 (vs), 1013 (vs); δ (Al-OH): 914 (s); other bands: 791 (w), 752 (w), 685 (m), 539 (s), 472 (s), 431 (s). 3-amino-1-propanol (Aldrich, NH₂(CH₂)₃OH, bp 187–188°C) for 19 h under N₂ atmosphere. Work-up consisted of cooling, filtering, and washing with methanol; 0.82 g of an off-white powder was recovered after air drying for 1 h. XRD and FTIR Kao/DMSO (1.0 g) (I.R. = 0.97) was refluxed with 50 mL of

Kao/DMSO + pt-1-amino-2-propanol (Rxn 7)

pt-1-amino-1-propanol (Aldrich, NH₂CH₂CH₂OHCH₃, bp 160°C) for 19 h under N₂ atmosphere. Work-up consisted of cooling, filtering, and washing with methanol; 0.72 g of an off-white powder was recovered after air drying for 1 h. FTIR indicated that no modification had occurred, and the Kaol DMSO starting material had reverted to a collapsed kaolinite. Kao/DMSO (1.0 g) (I.R. = 0.97) was refluxed with 50 mL of

were both consistent with a collapsed kaolinite structure. excess organic material as possible. FTIR and XRD patterns reaction, the mixture slowly turned black. Work-up consisted of repeated washings with methanol to remove as much of the interlayer space of kaolinite was made by reacting 1.0 g of Kao/DMSO in 10 g of melted nitroaniline (mp 149°C) at 175°C for 40 h under N₂ atmosphere. During the course of Kao/DMSO + nitroaniline (Rxn 8)
A preliminary attempt to intercalate nitroaniline into the

Results and discussion

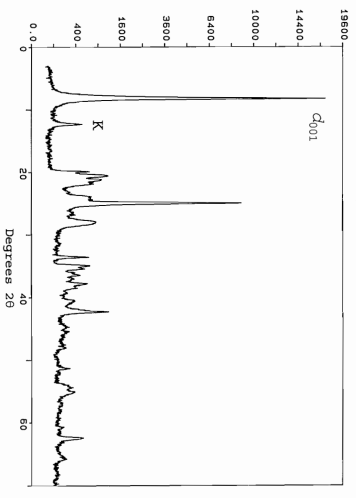
A series of amines were reacted with Kao/DMSO and Kao/ NMF (Table 1). These included the aminoalcohols ethanol-

phase. NR, no reaction. 'Intercalation ratio. "See experimental section for details.

"This refers to the basal spacing of the principal modified product

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Fig. 1. XRD pattern (2°-70° 20) for Kao-EOA. K indicates (001) reflection from residual kaolinite



and deformation bands at 1574 (w), 1482 (w), and 1322 cm⁻¹. contributed to a weaker host-guest interaction. may have also made interlayer diffusion more difficult and kaolinite before structural collapse could occur. Alternately, the presence of an end methyl group in 1-amino-2-propanol rapid diffusion of 3-amino-1-propanol into the interlayers of sibly be due to the higher reaction temperature for Kao-3,1-AP formation (187°C vs. 160°C), which permitted a more kaolinite, whereas DL-1-amino-2-propanol did not, may plau-The fact that 3-amino-1-propanol formed an intercalate with 3364 (vw) cm⁻¹, ν (CH) bands at 2941 (w) and 2878 (w) cm⁻¹ example, faint $\nu(NH)$ bands were detected at 3426 (vw) and attributable to the 3-amino-1-propanol guest species. For starting material has been altered, and one observes bands 11.1 Å, and the intercalation ratio (I.R.) was only 0.58. IR results for this material definitely show that the Kao/DMSO For Kao-3,1-AP (6), the modified phase had a basal spacing of 11.1 Å, and the intercalation ratio (I.R.) was only 0.58. IR lapsed unmodified kaolinite and Kao/DMSO starting material. and nitroaniline (8) were unsuccessful and yielded only colinvolving ethylene diamine (5), DL-1-amino-2-propanol (7), formed new organo-kaolinite (EOA) and 3-amino-1-propanol (3,1-AP), which intercalates. The reactions

In view of the fact that ethylene diamine may intercalate into halloysite to form an 11.7 Å intercalation complex (30, 31), it is surprising that an organo-kaolinite product was not formed via the displacement of DMSO. Reaction with *para*-nitroaniline was also unsuccessful, presumably due to the hydrophobic nature of *para*-nitroaniline and the weaker basicity of *para*-nitroaniline compared to primary amines.

ity of para-nitroaniline compared to primary amines. Clearly, the greatest success was achieved using ethanolamine as the reactant, and this product (Kao-EOA) will be examined in some detail. It was possible to form nearly identical Kao-EOA products from both Kao/DMSO and Kao/NMF starting materials but not from unexpanded kaolin-

ite. This indicates that the intercalating agent (DMSO or NMF) that was used to provide access to the interlamellar space of kaolinite plays a passive role and is not directly involved in the actual chemical reaction between ethanolamine and the interlamellar surface of kaolinite. DMSO and NMF are required only to provide access to the clay interlayers.

layers.

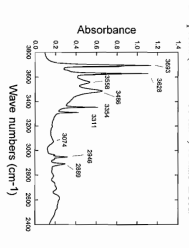
The Kao–EOA product exhibited a remarkable crystallinity as exhibited by the strong reflections of the XRD pattern (Fig. 1). The main product had a basal spacing of 10.67 Å. This product was found to be stable with respect to vacuum drying over phosphorus pentoxide for 24 h at 25°C, but was unstable when washed with water overnight, reverting slowly to the parent kaolinite 7.2 Å phase.

It was also possible to form the halloysite analogue of Kao–EOA (Hal–EOA), by reacting Hal/DMSO with ethanolamine (4). These materials exhibited similar basal spacings and IR patterns. Hal–EOA was much less crystalline than Kao–EOA, as evidenced by the very broad diffraction peaks. This was presumably due to the turbostratic nature of halloysite and Hal–EOA compared to kaolinite and Kao–EOA. No residual unexpanded parent mineral could be detected for the Hal–EOA, indicating that 100% modification was achieved. This was not the case for Kao–EOA, where a reflection at 7.2 Å representing 1.5% relative intensity indicates the presence of a small amount of residual unexpanded kaolinite.

To our knowledge, an ethanolamine derivative (intercalate or otherwise) of kaolinite has not been previously reported in the literature. Ethanolamine has, however, been reported intercalated in halloysite (31), forming a material with a basal spacing of 10.7 Å, which is similar to what is reported here (4). This corresponded to a layer expansion of 3.5 Å, which is less than the 4.0 Å minimum clearance space that was calculated

Fig. 2. IR spectra (3800-2400 cm⁻¹) of Kao-EOA.

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some keying in of the guest (1-3,lates that show a smaller than expected basal spacing due to for ethanolamine. This is in accord with many other interca-<u>∞</u>

anolamine units (Fig. 2). mally found at 3695, 367 between the kaolinite host and ethanolamine guest. Kao-EOA. These last two bands represent significant red shifts, indicating the effects of increased hydrogen bonding replaced by additional N-H stretching bands could be seen due to the eth-anolamine units (Fig. 2). The $\nu(OH)$ bands of kaolinite norparticular, the O-H stretching region was greatly perturbed and evidence of modification of the respective host minerals. In The IR patterns of Hal-EOA and Kao-EOA showed strong at 3695, 3670, 3652, and 3620 cm⁻¹ have b bands at 3693, 3628, 3558, and 3486 cm⁻¹ been

ity of kaolinite (20). A partial collapse of the hydrazine inter-C-H stretching bands were observed at 2946 and 2889 cm⁻¹ of two sharp N-H stretching bands at 3354 and 3311 cm evidence for ethanolamine modification includes the presence of kaolinite. In addition to the O-H stretching region, further the -NH₂ group of ethanolamine into the (Si-O-)₆ macro-ring band, therefore, provides some evidence for the keying in of phenomenon. This 8 cm^{-1} shift of the inner hydroxyl $\nu(\text{OH})$ the -NH₂ moiety of hydrazine into the siloxane ditrigonal cavintercalated kaolinite was previously attributed to keying of non for an evacuated (0.01 Torr; 1 Torr = 133.3 Pa) hydrazine The inner hydroxyl kaolinite peak at 3620 cm⁻¹ peak has been blue shifted to 3628 cm⁻¹. A similar blue shift phenomefrom 10.4 ō 9.6 was associated with this

tions region. The very strong in-plane Si-O-Si stretching vibrations of kaolinite at 1010 and 1033 cm⁻¹ (26) are shifted to higher frequencies in Kao-EOA to 1016 and 1045 cm⁻¹. The δ (Al-OH) band of the inner hydroxyl of kaolinite is shifted from 915 to 908 cm⁻¹, and the δ (Al-OH) band of the tions between kaolinite and the ethanolamine moiety. inner surface kaolinite hydroxyls at 938 cm⁻¹ is replaced by a band at 974 cm⁻¹. All this indicates strong host-guest interacmaterial also showed signs of perturbation in the lattice vibraband at 1353 cm⁻¹ could all be observed. The kaolinite host In the deformation region (Fig. 3), an NH bending band at 1613 cm⁻¹, a δ (CH) band at 1476 cm⁻¹, and a C-N stretching

The solid state ¹³C CP/MAS NMR spectrum (45.3 MHz) of Kao-EOA showed two broad resonances, one at 43 ppm and

Fig. 3. IR spectra (1800–1200 cm⁻¹) of Kao-EOA

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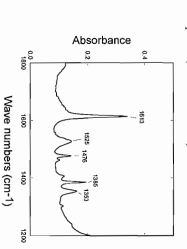
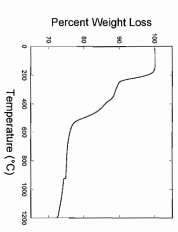


Fig. 4. TGA (20-1200°C) for Kao-EOA. Flowing nitrogen (40 cc/min); the heating rate was 20°C/min.



intercalated, but held rigidly in place by the inorganic host. minate interlamellar surface via \equiv Al-O-CH₂CH₂-NH₂ linkrigidly bound ethanolamine unit, plausibly grafted to the aludisappearance of both signals, indicating the existence of iment (32) (40 μs interrupted decoupling) led to the complete units in the interlayers of kaolinite. A dipolar dephasing experthe other at 60.5 ppm corresponding, respectively, to the 13 C-NH₂ carbon and the 13 C-O- carbon of the ethanolamine unit. was consistent with the insertion of the ethanolamine Alternatively, the ethanolamine molecule may be

Thermal analysis

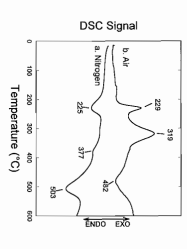
Thermal analysis of Kao-EOA shows that decomposition does not occur until about 150°C, and then there appears to be a three-step decomposition ending at 600°C (Fig. 4). The total almost immediately upon heating. liminary decomposition step (7.6% weight loss) beginning 27.5%. Hal-EOA differs from Kao-EOA in that there is a precalcination weight loss after heating to 1100°C in air for 3 h is

The thermal decomposition in nitrogen between 150 and 600°C is characterized by endothermic peaks at 225, 377, and 503°C (Fig. 5a). The peaks at 225 and 377°C are assigned to the endothermic pyrolysis of the interlayer organic material, mic events corresponding to the decomposition of interlayer organic material, two exotherms with peaks at 229 and 319°C the kaolinite host material. However, instead of two endotherendothermic peak at 482°C assigned to the dehydroxylation of the kaolinite host. In air atmosphere (Fig. 5b), one observes an and the peak at 503°C is assigned to the dehydroxylation of

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flowing air (40 cc/min). The heating rate was 20°C/min. Fig. 5. DTA for Kao-EOA under flowing nitrogen (40 cc/min), and



interlayer organic material had occurred. of the interlayer organic material. When the TGA/DSC run are observed. They are assigned to the exothermic combustion 1100°C was white, indicating that complete combustion of the was done in air atmosphere, the product after heating to

reaction of the alcohol group of the ethanolamine with the interlamellar Al-O-H groups of kaolinite. This is consistent one assumes that grafting has occurred via the condensation assign the formula $\mathrm{Al_2Si_2O_5(OH)_{3.0}(OCH_2CH_2NH_2)_{1.0}}$. Here (2.72%) values for the%H of this last assignment suggests that the chemical formula based on the grafting of ethanolamine units $(Al_2Si_2O_5(OH)_{3.0}(OCH_2CH_2NH_2)_{1.0})$ may be a better a chemical formula of Al₂Si₂O₅(OH)₄(CH₂OHCH₂NH₂)_{1.0}. amino group to key into one (Si-O-)6 macro-ring of the silicate representation of the structure. The discrepancy between the calculated (3.45%) and found lamellar surface of kaolinite without grafting, one could assign sider an ethanolamine unit that is intercalated into the intersurface of the adjacent layer. Alternately, if one were to con-Al₂Si₂O₅(OH)₄ mites of Inoue et al. (24) and with the methoxy-functionalized with the model for the aminoalcohol-derivatized organoboeh-Based on elemental analysis of Kao-EOA (1), one can structural unit of kaolinite, allowing each There is one ethanolamine unit per

Conclusions

each other in an ordered non-centrosymmetric manner. However, on the basis of the basal spacing (10.7 Å) and of the lesser ular assembly of dipolar organo-mineral layers, stacked upon also consistent with this structure, representing a supramolecthermal stability of Kao-EOA. Elemental analysis and IR are accounts for both the rigidity of the ethanolamine units observed from ¹³C DD/MAS NMR results, and the relative (SiO-)6 macro-ring of the adjacent silicate surface. This model valent linkages, with the pendant NH2 group keyed into the lamellar surface of kaolinite via \equiv Al-O-CH₂CH₂NH₂ cowhere ethanolamine units are attached to the aluminate inter-DMSO and Kao/NMF. A model in agreement with the characprepared from the reaction of ethanolamine with both Kao/ The well-ordered organo-kaolinite material, Kao-EOA, thermal and hydrolytic stability than the corresponding methterization data can be suggested: an organo-kaolinite material was

> oxy (3) and ethylene glycol derivatives (2), one can not unambigously disprove a structure when EOA units would be rigidly intercalated with EOA units keyed into the silicate the two previously reported ethylene glycol - kaolinite derivand grafted EOA-kaolinite. A similar mixture could be could be a mixture of both types of compounds, intercalated macro-rings and the aluminol sheets. The prepared material atives (1, obtained under certain experimental conditions in the case of

expected on the basis that the parent materials show this same difference in long-range ordering (8, 29). arranged in a turbostratic manner whereas Kao-EOA appears calate appears to be similar to Kao-EOA. The long-range ordering, as evidenced from XRD, indicates that Hal-EOA is to have some three-dimensional ordering. Regarding Hal-EOA, the short-range structure of this inter-

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