Synthesis and characterization of HDA/NaMMT organoclay

C YÜRÜDÜ, S İŞÇİ, C ÜNLÜ, O ATICI, Ö İ ECE and N GÜNGÖR*
Department of Physics, 1Department of Chemistry, 2Department of Geology, Istanbul Technical University, 34469, Maslak, Istanbul, Turkey

MS received 4 May 2005; revised 1 August 2005

Abstract. In this study, the rheologic and colloidal characterizations of sodium montmorillonite (NaMMT) were examined. Hexadecylamine (CH_{15}NH_{2}, HDA) was added to the bentonite water dispersion (2%, w/w) in different concentrations in the range $5 \times 10^{-3}$–$9.4 \times 10^{-4}$ m mol/l. The rheological and electrokinetic behaviour of aqueous montmorillonite dispersions was investigated as a function of solid content and HDA concentration. The basal spacings of the HDA/NaMMT composites were studied by X-ray diffraction. The FTIR spectra were obtained from the modified bentonite products, which revealed the characteristic absorbances after treatment with HDA.

Keywords. Montmorillonite; hexadecylamine; organoclay.

1. Introduction

Clays are widely used in industrial products and processes. The use of clay suspensions is important in ceramic industry as well as in the production of paper, detergent, and paints; foundry, civil engineering; and in drilling operations. The determination of the rheological characteristics such as viscosity and thixotropy of the clay–water dispersions is of great importance for the ceramic industry.

Bentonite, which is a kind of smectite group clay mineral, is almost made up of 80% of montmorillonite and contains a crystal lattice with three layers. It has a variety of uses because of its colloidal property. When it is mixed with water, it swells significantly. In water and in some organo environments, it produces a smooth homogeneous slurry having high plasticity. Layered smectite-type montmorillonites possess a negative surface charge which is compensated by exchange of cations, such as Na$^+$ or Ca$^{2+}$. Bentonite dispersions are sensitive to the exchangeable cations and pH. The valency of the exchangeable cations, monovalent or divalent, has a strong influence on the flow properties (Espinasse and Siffert 1979; Lagaly 1989; Angle and Hamza 1989; Permien and Lagaly 1994; Güngör 1998; Günister et al 2004). As organic cations exchange for exchangeable ions on the mineral surfaces, the cations are released into solution. The organic cations may also enter into ion-exchange reactions with exchangeable cations between the layers (Espinasse and Siffert 1979; Angle and Hamza 1989; Lagaly 1989; Alemdar et al 2005). The surface of the clay may be modified to make it strongly organophilic. Recently, nanoscale composites of polymers with organo-clay have been studied extensively. Montmorillonite is the most widely used layered silicate in polymer nanocomposite studies (Sheng et al 1996; Pan et al 1997; Chen et al 1998; Chang et al 2001, 2002; Chang and An 2002; Chen and Curliss 2003).

In this study, the Turkish natural bentonite sample has been identified as Ca-montmorillonite (CaMMT). The Na-activated montmorillonite (NaMMT) was obtained from natural montmorillonite by treating the clay with NaCl solution. The effect of HDA on the rheological properties and colloidal nature of the NaMMT dispersion is investigated. Besides, measurement of the colloidal and rheological properties, the NaMMT and HDA/NaMMT dispersions have been examined in detail with XRD and FTIR analysis.

2. Experimental

The clay sample was obtained from the bentonite deposits in Bigadiç–Balikesir, Turkey (courtesy of Unikim Ltd.). Ore samples have been identified as Ca-montmorillonite clay minerals using X-ray diffractometer (Philips model PW 1140, Holland), indicating dioctahedral structure of montmorillonite with minor amount of illite, quartz and feldspar. Natural montmorillonite is labeled as CaMMT and Na-activated clay as NaMMT.

The flow behaviour of the dispersions was measured in a Brookfield DVIII + type low-shear rheometer. The sample was dispersed in water (2% w/w) and shaken overnight. An adsorption time of 24 h was adopted for the surfactant. The rheological behaviour of the clay suspensions was obtained by shear stress–shear rate mea-
measurements within 0–350 s\(^{-1}\) shear rates. Rheological measurements were carried out in duplicate.

The zeta potential measurements were carried out using a zetasizer equipment (Malvern Instruments, Zetasizer 2000, UK). The optical unit of the zetasizer 2000 contains a 5 mW He–Ne (638 nm) laser. Before the measurements, all the dispersions were centrifuged at 4500 rpm for 30 min. Supernatants were then used for zeta potential measurements. To make an electrophoretic mobility measurement in this instrument, laser beams were crossed at a particular point in the cell. Particles in the cell were illuminated by these beams. At the crossing point of the beams, Young’s interference fringes formed. Particles moving through the fringes under the influence of the applied electric field scatter light whose intensity fluctuates with a frequency that is related to the particle velocity. The photons detected by the photomultiplier are fed to a digital correlator, the resulting function being analysed to determine the frequency spectrum, from which the mobility and hence the zeta potential are calculated. Measured electrophoretic mobilities were converted to zeta potential using established theories. Zeta potential was measured by injecting a small portion of the supernatant into the cell of the Zetasizer 2000 instrument at room temperature.

The obtained film samples of intercalated clay with different HDA concentrations were analysed by X-ray diffraction. X-ray diffraction (XRD) measurements were performed at room temperature using Ni-filtered Cu-K\(\alpha\) radiation. Infrared analyses (400–4000 cm\(^{-1}\)) were performed on a FTIR spectrophotometer (Jasco Model FT/IR-5300, Japan) using KBr pellets with a concentration of 1\%. Spectral outputs were recorded either in absorbance or in transmittance mode as a function of wave number. Hexa-decylamine (CH\(_3\) (CH\(_2\))\(_{15}\) NH\(_2\), M\(_\text{W}\) = 241-46) was purchased from Aldrich Chemical Co., USA and used as received.

3. Results

Figure 1 shows typical shear rate(\(\gamma\))–shear stress (\(\tau\)) curve for a 2% w/w Ca and Na-montmorillonite dispersions. The dispersions of the CaMMT exhibit nearly Newtonian behaviour. The viscosity of a Newtonian dispersion is the shear stress versus shear rate ratio. Yield value was not found and because of this we infer that the interaction between the particles of 2% dispersions is very small. The dispersions of NaMMT exhibit Bingham plastic flow behaviour. The apparent viscosity values at 55 s\(^{-1}\) of CaMMT and NaMMT have been found to be 1.25 and 15.27 mPas, respectively. This difference is caused by the change in swelling and dispersion behaviour of the neutral bentonite after the activation process. Exchangeable cations, which determine orientation of many physical properties of bentonitic clays such as particle size, surface charge, surface area, particle number per unit volume etc also have a strong influence on the dispersion mechanisms. The increase of viscosity and consistency of CaMMT dispersions after addition of sodium ion is a direct consequence of the opposite effect of Na\(^+\) and Ca\(^{++}\) ions. When clays are dispersed in water, NaMMT has unique crystal layer in very fine colloidal particles with sizes ranging from 10–100 \(\mu\)ms, because the electrostatic attraction among the crystal layers is weak. On the other hand, for the CaMMT, the mean particle size is kept at several \(\mu\)ms in water, because the electrostatic attraction among the crystal layers is strong.

Figure 1. The shear stress–shear rate rheogram for CaMMT and NaMMT dispersions.
Rheological parameters are sensitive to solid content. By increasing the solid content of the dispersions, the viscosity and yield values increase. The flow and apparent viscosity–shear velocity curves of the 1, 2, 3, 4, 5 and 6% clay dispersions that have been prepared with the NaMMT sample, have been drawn and the flow models determined. The shear stress versus shear rate for suspensions containing different concentrations of NaMMT samples are shown in figure 2. The dispersions (1, 2 and 3% w/w) of NaMMT exhibited Bingham plastic behaviour. The yield values have been increased by the solid ratio. The increase in the yield value with clay concentration (solid content) is due to the increase of number of clay particles in the dispersion (figures 2a and b). The Bingham yield value (extrapolated shear stress) is a criterion of particle–particle interaction and a better parameter for quantifying the structure of a flocculated system.

Hexadecylamine was used as an organophilic alkylamine in organo-clay. In figure 3, the yield value of bentonite dispersions was plotted as a function of increasing HDA concentrations. After the first addition of surfactant, the yield value of NaMMT dispersions increased. The flocculation of NaMMT dispersion was greatly increased by small additions of HDA and a maximum was reached at \(1.87 \times 10^{-3}\ \text{mol/l}\) HDA. Further addition of surfactant results in a decrease in the rheological parameters and the system begins to get deflocculated.

The degree of thixotropic or antithixotropic behaviour was measured by the area of the hysteresis loop (figure 4). The area of the hysteresis loop of the flow curves, which is a measure of the degree of hysteresis, is accepted to be positive if the behaviour of the system is thixotropic, and negative if the behaviour is antithixotropic. The results indicate a gradual increase in gelation with the first addition of HDA, which reaches a maximum at \(9.35 \times 10^{-4}\ \text{mol/l}\) surfactant. After that point a decrease is observed. Further addition of HDA results in an increase in the area of the hysteresis loop.

In figure 5, the flow curves of 2 wt.% bentonite dispersion and the modified organo-clay with \(5.91 \times 10^{-4}\)

![Figure 2](image2.png)  
**Figure 2.** Effect of clay concentration (w/w) on (a) the flow curves and (b) yield value.

![Figure 3](image3.png)  
**Figure 3.** The changes of the yield value of NaMMT dispersions with HDA surfactant additions.

![Figure 4](image4.png)  
**Figure 4.** The hysteresis loop area of 2% w/w bentonite dispersion as a function of HDA concentration.
$1.87 \times 10^{-3}$, $5.61 \times 10^{-3}$ and $9.35 \times 10^{-3}$ mmol/l HDA, respectively are displayed. Figure 5 shows an increase in gelation till the $1.87 \times 10^{-3}$ mol/l surfactant addition. However, at $5.6 \times 10^{-3}$ mol/l surfactant addition a clear decrease is observed. This curve agrees with the curves in figures 4 and 5 since the maximum yield value (figure 5) are seen at around $1.87 \times 10^{-3}$ mol/l surfactant addition. This point shows the maximum deflocculation; on further addition, the system begins to get deflocculated.

Figure 6 shows the change in zeta potential with HDA concentration for NaMMT dispersions. The zeta potential is an electrical potential in the double layer at the interface between a particle, which moves in an electric field, and surrounding liquid. The surface charge property can be characterized by zeta potential and the stability of a clay solution can be measured depending on its value. Increasing of zeta potential values after the first addition of HDA clearly shows an increase in electrostatic repulsion power between clay particles. However, two possible hypotheses can be proposed to explain the reason for increase in zeta potential. (i) Currently existing clusters between clay particles could be dispersed after the addition of HDA. Consequently, the system contains both large size clusters holding a number of particles together with much smaller clusters/particle sizes and therefore, electrostatic repulsion power and zeta potential will increase and (ii) positively charged edges of clay particles can make bonds with one of unbounded electrons of HDAs. Under these circumstances, electrostatic repulsion power will increase, because net charges of clay particles will be more negative. When the additive contents increase, it is possible to understand from the decrease in negative charges that some quantities of clays get

![Figure 5. The changes of flow curves of NaMMT dispersions with HDA additions.](image)

![Figure 6. The changes of the zeta potential of NaLB dispersions with HDA surfactant additions.](image)

![Figure 7. XRD patterns of NaLB and HDA/NaLB.](image)
attached on the surfaces of HDAs. However, surfactant does not cover completely the surface of clay minerals in the concentration range of this study. Dispersion shows lower thixotropic properties (figure 4) wherein zeta potentials have more negative values (figure 6). Even there is an increase in thixotropic values after last two additions of surfactants, but it never reaches the previous values of the system without surfactants. These observations are comparable with the idea of relationship between additional surfactants and deflocculation of the system (figures 3 and 5).

The clay sheets are bound with in-plane covalent bonds and therefore, their crystal structure is stable, but clay layers are held loosely with Van der Waals bonds and therefore, expansion of interlayers is commonly seen when water or organic molecules are introduced between the layers. Figure 7 shows the XRD patterns of NaMMT samples without organic modification and modified with HDA.

Figures 8a,b and 9 compare the FTIR spectra of hexadecylamine–montmorillonite, which were prepared using solution intercalation (HDA/NaMMT), with those of activated Balikesir montmorillonite (NaMMT) and hexadecylamine (HDA). After NaHCO₃ activation, the peak corresponding to structural hydroxyl stretching at 3628 cm⁻¹ is attributed to montmorillonite (figure 8a). The broad peak centred on 3464 cm⁻¹ is assigned to O–H stretching. HOH deformation peak centred at 1642 cm⁻¹ is also present in the FTIR spectrum of Na-montmorillonite (figure 8b). The absorption bands at 1032, 797, 619, 519 and 467 cm⁻¹ belong to Si–O–Si, Mg–Al–OH, Al–O, Si–O–Mg and Si–O–Fe vibrations, respectively (figure 9). The spectrum of Na-montmorillonite that adsorbed 8 g/l cationic surfactant HDA contains characteristic bands of all components (figures 8 and 9). Primary amine NH₃ asymmetrical and symmetrical stretching vibrations at 3123 and 3098 cm⁻¹, aliphatic C–H asymmetrical and symmetrical stretching vibrations at 2924 and 2853 cm⁻¹, C–H bendings at 1470, and 720 cm⁻¹ NH₃ asymmetrical and symmetrical bendings at 1541–1566 cm⁻¹ are assigned to the vibration modes of HDA. Absorbances due to structural O–H stretching around 3621 cm⁻¹ and Si–O bendings at 519 and 465 cm⁻¹ confirm the presence of Na-montmorillonite in the dispersion. In the bentonite HDA adsorption products, the O–H stretching peak was broadened and gave a maximum at 3428 cm⁻¹ (figure 8). The bonding of HDA also results in the weakening of H–O–H stretching vibration band, which can be explained by the replacement of metal ions by HDA via ion exchange. The Si–O stretching peak was also broadened and gave a maximum at 1030 cm⁻¹. Absorption bands of Mg–Al–OH vibration for HDA/NaMMT 801 cm⁻¹ are different than those for the NaMMT (figure 9). This is attributed to the relaxation of hydrogen bonding between Mg–Al–OH as

![Figure 8](image-url). FTIR spectra in the 2700–3800 cm⁻¹ and 1350–1750 cm⁻¹ regions. (a. Na-montmorillonite and b. 8 mmol/l HDA adsorbed on Na-montmorillonite).
well as to the hydrated water of exchangeable cationic metal ions on the montmorillonite surface, further supporting the presence of HDA.

4. Conclusions

The adsorbed HDA affects the rheological and electrokinetic properties of NaMMT dispersions. The degree of interaction between HDA and the bentonite particles depends on the HDA concentration in the suspension. Addition of the surfactant results in firstly the flocculant and then deflocculant effect on 2% w/w of clay dispersion.

The zeta potential measurements indicate that the HDA molecules hold onto the clay particle surfaces very weakly. XRD analyses results indicate that organic molecules did not enter sufficiently into the layers of clay structures. The comparison of FTIR spectra of the natural and HDA adsorbed montmorillonite has indicated that sorbed HDA molecules on montmorillonite are coordinated to exchangeable cations directly or indirectly through water bridges.

Acknowledgement

This research project is supported by the Istanbul Technical University, Research Fund (Project No: 30451).

References

Angle C W and Hamza H A 1989 Appl. Clay Sci. 4 263
Chen C and Curliss D 2003 Nanotechnology 14 643
Chen G, Han B and Yan H 1997 J. Colloid Interface Sci. 194 276
Espinasse P and Siffert B 1979 Clays and Clay Miner. 27 284
Güngör N 1998 Revue Roumaine de Chimie 43 121
Lagaly G 1989 Appl. Clay Sci. 4 105
Permien T and Lagaly G 1994 Clay Miner. 29 751
Sheng G, Xu S and Boyd S A 1996 Wat. Res. 30 1483

Figure 9. FTIR spectra in the 400–1180 cm\(^{-1}\) region (a. Na-montmorillonite and b. 8 mmol/l HDA adsorbed on Na-montmorillonite).