Polymer/Layered Silicate Nanocomposites from Thermally Stable Trialkylimidazolium-Treated Montmorillonite

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The limited thermal stability of alkylammonium cations intercalated into smectite minerals (e.g., montmorillonite, MMT) and the processing instability of some polymers [polyamide-6 (PA-6) and polystyrene (PS)] in the presence of nanodispersed MMT have motivated the development of improved organophilic treatments for layered silicates. Success in this regard should enable the preparation of polymer/layersilicate nanocomposites from thermoplastic polymers that require high melt-processing temperatures or long residence times under high shear and from thermostable resins with high cure temperatures. Our efforts to address some of these issues focus on the use of new thermally stable imidazolium-treated layered silicates for the preparation of nanocomposites. Several trialkylimidazolium salt derivatives were prepared with propyl, butyl, decyl, and hexadecyl alkyl chains attached to the imidazolium through one of the nitrogens. These imidazolium salts were used to prepare the corresponding layered silicates. We report here that the use of 1-alkyl-2,3-dimethylimidazolium salts to replace the sodium in natural MMT gives organophillic MMT with a 100 °C improvement in thermal stability (in N2) as compared to the alkylammonium-treated MMT. The use of 1-alkyl-2,3-dimethylimidazolium salt in fluorinated synthetic mica (FSM) also gives a 100 °C improvement in thermal stability. The use of 1,2-dimethyl-3-hexadecylimidazolium-treated MMT gives an exfoliated PA-6 nanocomposite and, depending on processing conditions, either a partially exfoliated or an intercalated polystyrene nanocomposite.

Introduction

A new approach to addressing the ever-increasing demand for higher-performance polymeric products focuses on the use of mica-type layered silicates, such as montmorillonite (MMT), nanodispersed (exfoliated) in polymers.1 These “nanocomposites” exhibit the unusual combination of improved physical properties2,3 and reduced flammability.4–8

However, using the standard alkylammonium-treated MMT to prepare nanocomposites from a variety of polymers [polystyrene (PS), poly(ethylene-co-vinyl acetate) (PEVA), polypropylene (PP), polyamide-6 (PA-6), epoxies, and cyanate ester], we have observed less than the expected improvement in properties (physical and flammability); in some instances, a loss in properties was observed relative to the unmodified polymer.9,10 Indeed, the only known products on the market, after over two decades of research, that take advantage of the attributes of nanocomposites are those using low-melt-temperature resins such as polypropylene,11 poly(ethylene-co-vinyl acetate) polymers,12 and synthetic rubbers.13

The processing stability of both the polymer and the organically treated layered silicate has a significant influ-

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ence on the flammability performance of the nanocomposite. In a previous TGA–FTIR study of the pyrolysis of PS/MMT nanocomposites, aliphatic decomposition products were observed before the onset of PS degradation. This suggests that the organic treatment was degrading at a lower temperature in the MMT than in the PS. This might contribute to the shorter ignition times observed in PS/MMT nanocomposites during combustion experiments. Because flame spread rates are increased by reduced ignition times, this effect must be better understood and controlled. Gel permeation chromatography (GPC) analysis of the extruded PS nanocomposite samples also reveals some evidence of polymer degradation in the form of lower values of the number-average molecular weight \(M_n\). The molecular mass degradation occurs only if quaternary alkylammonium MMT is present, and it does not occur either if pure polymer is extruded or if the PS is extruded with sodium montmorillonite (NaMMT). It appears that the presence of the quaternary alkylammonium in the MMT somehow contributes to the degradation of the PS as well. The decrease of \(M_n\) can have a direct negative impact on the flame-retardant performance by reducing the melt viscosity. This counteracts the anti-dripping effect of nanocomposites, which is a critical aspect of the flammability reduction mechanism. We have observed one case in which the flame-retardant effect is completely negated when extensive degradation occurs.

In addition, these effects could also limit the improvements in other physical properties observed for PS/MMT nanocomposites.

High-melt-temperature engineering polymers, such as PA-6, PA-6,6, poly(ethylene terephthalate) (PET), and polycarbonate (PC), raise additional concerns when considering preparation of nanocomposites by melt extrusion or when processing (injection molding) the nanocomposites into final molded products. The concern is that most alkylammonium treatments for MMT have an onset of thermal decomposition at about 200 °C or below. The melt processing temperature of PA-6, PET, and PC are above this temperature, and to facilitate rapid manufacturing of filled PA-6, typical industrial melt-processing temperatures for PA-6 are in excess of 300 °C. The issue is whether the alkylammonium-treated layered silicates can survive short residence times (<200 s) in high-shear processing environments (extruder or other molding equipment) at 250 °C or higher.

(15) According to ISO 31-8, the term “molecular weight” has been replaced by “relative molecular mass”, symbol \(M_r\). Thus, if this nomenclature and notation were used here, \(M_r\) instead of the historically conventional \(M_n\) for the average molecular weight (with similar notations for \(M_w\), \(M_m\), and \(M_z\)) would be used. This quantity would be called the “number-average relative molecular mass”. The conventional notation, rather than the ISO notation, has been employed here.
(23) D. Paul, personal communication.

We have found that both alkylammonium and PA-6 itself can degrade during processing in the presence of nanodispersed MMT. Using NMR spectroscopy, we observed degradation of dimethyl-di(hydrogenated tallow)ammonium montmorillonite (DMDHT-MMT) during melt extrusion with PA-6. The NMR analysis showed significant concentrations of tertiary amine after extrusion. The concentration of tertiary amine was directly proportional to the residence time in the twin-screw extruder and appeared to be nearly independent of the thermal history in the absence of shear. It was found that as much as 80% (±10%) of the quaternary alkylammonium had degraded in the samples with extrusion residence times long enough to give delaminated nanocomposites.

This means that the nanocomposite was contaminated with up to a mass fraction of 3% tertiary amine. A comparison of another delaminated PA-6/MMT nanocomposite from the same study, prepared using dihydroxyethyl-methyl-hydrogenated tallow ammonium/MMT, showed an order of magnitude less degradation of the organic treatment. This sample, with no observable degradation products and similar extent of nanodispersion of the MMT, had superior mechanical properties compared to the PA-6/MMT nanocomposite with degradation products. Presumably, the better compatibility of this more polar treatment and the associated very short residence time necessary to exfoliate the treated MMT into the PA-6 prevented the degradation of the alkylammonium in this case.

In a separate study of PA-6/MMT nanocomposites prepared via in situ polymerization, we found large \(M_n\) decreases as high as 40% and significantly higher concentrations of caprolactam monomer after processing by injection molding at 300 °C. In the case of intercalated cyanate ester nanocomposites, the flammability of one of a series of materials was increased by 40% relative to that of the pure resin. In a series of epoxy nanocomposites cured at high temperature, the glass transition temperature \(T_g\) was depressed relative to that of the pure resin. For both of these cases, curing of the nanocomposites had to be done above the decomposition temperature of the alkylammonium-treated MMT. A possible explanation of the observed losses in properties is that the decomposition of the alkylammonium interfered with the cure chemistries. Several other groups have also found that these treatments have insufficient thermal stability to survive high-temperature melt-processing or curing conditions. Paul and co-workers at the University of Texas at Austin have also observed \(M_n\) reductions during compounding of PA-6/MMT nanocomposites, and they find a correlation between the degree of degradation and the structure of the alkylammonium MMT. Some solutions to this...
issue include the use of untreated layered silicate. In these approaches, either the monomer or the curing agent is made cationic in situ, and this renders the layered silicate organophilic. Other cations, such as phosphonium, pyridinium, and iminium, have also been used to treat the layered silicate because of their greater thermal stabilities. Thus, the need for organophillic-treated layered silicates that offer improved stability under processing conditions is well-documented. New thermally stable systems should enable the preparation of polymer/layered silicate nanocomposites from thermoplastic engineering polymers with high melt-processing temperatures (PA-6,6, PC, PET, s-PS, PEI) and from thermoset resins with high cure temperatures (cyanate esters, aromatic epoxies) without loss of properties due to the presence of degradation products, loss of molecular mass, or network structural defects. Our efforts to address some of these issues focus on the use of new thermally stable trialkylimidazolium-treated layered silicates for the preparation of nanocomposites.

**Experimental Section**

**Preparation of Trialkylimidazolium Salts.** Imidazolium salts (see Figure 1) were typically prepared as follows: The alkyl bromide or chloride (1 mol), 1,2-dimethylimidazole (distilled, 0.95 mol), and acetonitrile (50 mL) were combined in a round-bottom flask equipped with a reflux condenser. The mixture was refluxed for 7–10 days under nitrogen. After the reaction was complete, a large excess of ethyl acetate was added to precipitate the imidazolium salt. This solid was filtered and washed several times with ethyl acetate to remove the 1,2-dimethylimidazole. Residual solvent was removed from the resulting 1-alkyl-2,3-dimethylimidazolium salt under vacuum at 80 °C. In some cases, the salts were converted to the corresponding tetrafluoroborate salts to improve purity. The products were characterized via mass spectroscopy (MS), 1H and 13C NMR spectroscopy, and thermal gravimetric analysis (TGA); the characterization data are consistent with data published on similar dialkyimidazolium structures. Complete details of the synthesis and characterization of all of the imidazolium salts presented here, and of other imidazolium salts, will be published in a separate manuscript. The 1H and 13C NMR data for two of the trialkylimidazolium compounds are presented in Table 1.

**Preparation and Characterization of Ion-Exchanged Layered Silicates.** Standard ion-exchange procedures were employed for the preparation of the organic-treated layered silicates. The following trialkylimidazolium layered silicate complexes were prepared from the corresponding imidazolium chloride, bromide, or BF4 salt: 1-butyl-2,3-dimethylimidazolium montmorillonite (BDMIM-MMT), 1-decyl-2,3-dimethylimidazolium montmorillonite (DDMIM-MMT), 1,2-dimethyl-3-hexadecylimidazolium montmorillonite (DMHDIM-MMT), and 1,2-dimethyl-3-hexadecylimidazolium fluorinated synthetic mica (DMHDIM-FSM). For comparison purposes dimethyldioctadecylammonium montmorillonite (DDMDA-MMT) was prepared from the corresponding bromide (Aldrich), also using the standard ion-exchange methods, and dimethyl-di-(hydrogenated tallow)ammonium montmorillonite (DMHT-MMT) was obtained from Southern Clay Products (Gonzales, TX).

The imidazolium-treated layered silicate samples were characterized using X-ray diffraction (XRD) and TGA. In the case of DDMIM-MMT and DMHDIM-MMT, thermal desorption mass spectroscopy (TDMS) and residual chloride analysis were also conducted.

Thermal desorption mass spectrometry (TDMS) was performed on a Finnigan TSQ-70 triple quadrupole mass spectrometer. Samples were prepared as slurries in water and applied to the tip of a direct exposure probe, which consists of a small tungsten filament. The probe was then positioned in the ion source of the mass spectrometer. Samples were desorbed from the probe by passing a current through the wire and increasing the temperature of the probe tip. The desorption profile was generated by holding the current constant at 50 mA for 6 s, increasing the current (linearly) from 50 to 1000 mA over 45 s, and holding the current constant at 1000 mA for 6 s. Molecules desorbed from the probe were ionized with 70 eV electrons. Mass spectra were obtained at 0.5 s per scan.

**Trace Analysis of Chloride.** A cyclic Voltammogram (cathodic end only) was carried out on pure propylene carbonate, DDMIM-MMT, and DMHDIM-MMT. All work was conducted under N2. The organo-MMT samples were prepared as suspensions in propylene carbonate (1 g in 30 mL) by stirring

<table>
<thead>
<tr>
<th>1H chemical shift (ppm)</th>
<th>DDMIM+BF4</th>
<th>DDMIM+Cl</th>
</tr>
</thead>
<tbody>
<tr>
<td>N–CH3</td>
<td>3.65</td>
<td>3.75</td>
</tr>
<tr>
<td>C2–CH3</td>
<td>2.48</td>
<td>2.50</td>
</tr>
<tr>
<td>H(4,5)</td>
<td>7.22</td>
<td>7.40, 7.45</td>
</tr>
<tr>
<td>N–C1(CH3)</td>
<td>4.00</td>
<td>4.05</td>
</tr>
<tr>
<td>N–C2(CH3)</td>
<td>2.10</td>
<td>1.75</td>
</tr>
<tr>
<td>N–C3 to C14(CH2)n</td>
<td>1.25</td>
<td>–</td>
</tr>
<tr>
<td>N–C3 to C9(CH2)</td>
<td>–</td>
<td>1.25</td>
</tr>
<tr>
<td>N–C10(CH2)</td>
<td>–</td>
<td>0.85</td>
</tr>
<tr>
<td>N–C15(CH2)</td>
<td>1.70</td>
<td>–</td>
</tr>
<tr>
<td>N–C16 CH3</td>
<td>0.85</td>
<td>–</td>
</tr>
</tbody>
</table>

* The policy of the National Institute of Standards and Technology (NIST) is to use metric units of measurement in all of its publications and to provide statements of uncertainty for all original measurements. In this document however, data from organizations outside NIST are shown, which might include measurements in nonmetric units or measurements without uncertainty statements.
Polymer/ Layered Silicate Nanocomposites

Figure 2. XRD data for several 1M-MMt salts showing the d spacing between MMT layers. The standard uncertainty in the d spacing is ±0.03 nm (2σ).

Table 2. 13C NMR Data for 1,2-Dimethyl-3-hexadecylimidazolium Tetrafluoroborate (DMHDIM+ BF4−) and 1-Decyl-2,3-dimethylimidazolium Chloride (DDMIM+ Cl−)

<table>
<thead>
<tr>
<th>Chemical shift (ppm)</th>
<th>DMHDIM+ BF4−</th>
<th>DDMIM+ Cl−</th>
</tr>
</thead>
<tbody>
<tr>
<td>N−CH3</td>
<td>31.5</td>
<td>31.5</td>
</tr>
<tr>
<td>C2</td>
<td>144.5</td>
<td>144.5</td>
</tr>
<tr>
<td>C2−CH3</td>
<td>9.0</td>
<td>9.0</td>
</tr>
<tr>
<td>C4</td>
<td>121.0</td>
<td>121.0</td>
</tr>
<tr>
<td>C5</td>
<td>122.0</td>
<td>122.0</td>
</tr>
<tr>
<td>N−C1(CH3)</td>
<td>48.0</td>
<td></td>
</tr>
<tr>
<td>N−C2(CH3)</td>
<td>35.0</td>
<td></td>
</tr>
<tr>
<td>N−C3(CH3)</td>
<td>26.0</td>
<td></td>
</tr>
<tr>
<td>N−C4 to C8(CH2)5</td>
<td>−</td>
<td></td>
</tr>
<tr>
<td>N−C4 to C14(CH2)11</td>
<td>29.0</td>
<td></td>
</tr>
<tr>
<td>N−C9</td>
<td>−</td>
<td>22.2</td>
</tr>
<tr>
<td>N−C10</td>
<td>−</td>
<td>13.0</td>
</tr>
<tr>
<td>N−C15</td>
<td>23.0</td>
<td>−</td>
</tr>
<tr>
<td>N−C16</td>
<td>13.0</td>
<td>−</td>
</tr>
</tbody>
</table>

The samples were vigorously for 1 h. The suspension was filtered through a 0.45-μm polypropylene frit and placed in the cell. The cell was a 40-μL glass vial filled with a Teflon cap containing a 6.35-mm graphite rod counter electrode and a 4.6-mm-diameter Pt working electrode, with a 4.6-mm-diameter Pt wire serving as a pseudo-reference.

Preparation and Characterization of Nanocomposites. PS and PA-6 nanocomposites were prepared in a mini-twin-screw extruder (intermeshing, conical, DACA Corp.). Polymer- and organic-treated layered silicate were charged into the mini-extruder and typically mixed at 213 °C (200–300 rpm) for 3–5 min at 10 °C above the melting point of the polymer. The nanocomposites were characterized using XRD, TEM, and solid-state NMR spectroscopy (PA-6 nanocomposites only).

Transmission Electron Microscopy. All samples were ultra-thin sectioned with a diamond knife on a Leica Ultracut UCT microtome at either room temperature (PS) or −110 °C (PA-6) to give sections with a nominal thickness of 70 nm. The sections were transferred from water (room temperature) or dry conditions (−110 °C) to carbon-coated Cu grids of 200 mesh. Bright-field TEM images of nanocomposites (except for PA-6) were obtained at 120 kV, under low-dose conditions, with a Philips 400T electron microscope, using Kodak SO-161 film. Low-magnification images were taken at 2800× and 10 000×. High-magnification images were taken at 28 000× and 60 000×.

Solid-State NMR Spectroscopy. PA-6 nanocomposites prepared with DMHDIM-MMT were characterized using a solid-state NMR proton longitudinal relaxation rate (T1H) method recently developed by VanderHart et al.17 Solid-state T1H values were measured using a noncommercial NMR spectrometer operating at 2.35 T. T1H values were measured by the inversion recovery method using variable delays of 1 ms, 75 ms, 235 ms, 500 ms, 1.0 s, and 3.0 s. Proton polarization levels at each of these delays were monitored indirectly via 1H cross-polarization magic angle spinning (CPMAS) NMR spectroscopy at 25.19 MHz. 1H-CPMAS spectra were obtained using a 4.0-kHz magic angle spinning frequency, a 3-s recycle delay, a 1-ms CP period, and radio frequency (rf) levels corresponding to nutation frequencies of 60 and 64 kHz for

Table 3. Thermal Stability Data for Imidazolium- and Quaternary Alkylammonium-Treated MMTa

<table>
<thead>
<tr>
<th>Sample</th>
<th>Organic fraction (±0.05)</th>
<th>Onset decomposition temp (TGA) (°C)b</th>
<th>Peak decomposition temp (DTA) (°C)b</th>
<th>Change in d spacing vs NaMMT (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1,2-dimethyl-3-hexadecylimidazolium/MMT</td>
<td>0.25</td>
<td>343</td>
<td>406</td>
<td>0.72</td>
</tr>
<tr>
<td>1-decy-2,3-dimethylimidazolium/MMT</td>
<td>0.17</td>
<td>320</td>
<td>445</td>
<td>0.49</td>
</tr>
<tr>
<td>1-butyl-2,3-dimethylimidazolium/MMT</td>
<td>0.13</td>
<td>340</td>
<td>448</td>
<td>0.16</td>
</tr>
<tr>
<td>1,2-dimethyl-3-propylimidazolium/MMT</td>
<td>0.13</td>
<td>340</td>
<td>445</td>
<td>0.20</td>
</tr>
<tr>
<td>1,2-dimethyl-3-hexadecylimidazolium/FSM</td>
<td>0.24</td>
<td>392</td>
<td>400</td>
<td>0.63</td>
</tr>
<tr>
<td>dimethyl dioctadecyl ammonium bromide</td>
<td>−</td>
<td>221</td>
<td>233</td>
<td>−</td>
</tr>
<tr>
<td>dimethyl dioctadecyl ammonium bromide/MMT</td>
<td>0.36</td>
<td>280</td>
<td>308</td>
<td>1.49</td>
</tr>
<tr>
<td>dimethyl-di(hydrogenated tallow) ammonium/MMT</td>
<td>0.35</td>
<td>200</td>
<td>310</td>
<td>2.00</td>
</tr>
</tbody>
</table>

a MMT = montmorillonite. b Uncertainty for onset and peak T, values is ±1.2 °C (2σ).
protons and $^{13}$C nuclei, respectively. The total experiment time was 2 h per sample, with 320 scans for each of the 6 delay values.

Water content and thermal history, in addition to MMT dispersion quality, can significantly alter the $T_1^H$ values of PA-6/MMT nanocomposites. Therefore, samples were carefully

Figure 4. Thermal desorption electron ionization mass spectrum of DMHDIM-MMT and total ion plot of peak at 0.57 min.

Figure 5. Total ion plot of the peaks at (A) 0.49 and (B) 0.56 min and (C) thermal desorption electron ionization mass spectrum of DMHDIM $^{+}$Cl$^-$. 
prepared to normalize the effect of water and thermal history by $T_1^H$ and to allow quantitative determination of the extent of nanodispersion of MMT in PA-6.

**Thermal Gravimetric Analysis.** TGA was carried out using a TA Instruments Simultaneous TGA-DTA (SDT 2960) at 10 °C/min in nitrogen (samples of 5–10 mg each). Typically, three replicates were run for each sample, and the mean was reported. Both the onset (5% mass fraction loss) and peak mass loss rate have an uncertainty of 1.2 °C (2σ).

**Results and Discussion**

**Organo-Modified Montmorillonite: Imidazolium-Treated Layered Silicates.** It has been reported that the delocalized imidazolium cation has better thermal stability than the alkylammonium and pyridinium cations. The imidazolium salts shown in Figure 1 were used to treat NaMMT, via standard literature ion-exchange methods, to give a series of imidazolium-MMT (IM-MMT) materials. The NMR characterization data for the DDMIM$^+$Cl$^-$ and DMHDIM$^+$BF$_4^-$ salts are summarized in Tables 1 and 2. The IM-MMT were analyzed by XRD to determine whether the spacing between the layers ($d$ spacing) had changed. The XRD data showed that as the R group increased in length, the $d$ spacing increased (see Figure 2), with the exception of the butyl and propyl derivatives, which have nearly the same steric size and, hence, the same $d$ spacing. Using the same ion-exchange procedure, we...
also prepared and characterized 1,2-dimethyl-3-hexadecylimidazolium fluorinated synthetic mica (DMH-DIM-FSM). The XRD data for these materials are shown in Table 3.

As we emphasized above, the thermal stability of the organic treatment of the layered silicate is of prime importance; therefore, TGA was carried out on these materials. For comparison, two quaternary alkylammonium-treated MMT were evaluated as well: dimethyl dioctadecylammonium montmorillonite (DMDODA-MMT) and dimethyl-di(hydrogenated tallow)ammonium montmorillonite (DMDHT-MMT). The TGA data, recorded in a nitrogen atmosphere, comparing DMDODA-MMT and DMHDIM-MMT are shown in Figure 3. The TGA data for all of the trialkylimidazolium- and tetraalkylammonium-treated MMT and FSM samples are summarized in Table 3; they clearly show the improvements in thermal stability for all of the IM-LS samples compared to both DMDODA-MMT and DMDHT-MMT. Substitution of methyl at the methyne position of the imidazolium was found to be critical and to provide 50 °C higher thermal stability for the trisubstituted imidazolium compared to the disubstituted imidazolium (data not shown).³⁰

The enhancement in thermal stability that occurs when the imidazolium is intercalated into the MMT was probed by thermal desorption mass spectroscopy (TDMS). The greater thermal stability of the DMHDIM-MMT over the DMHDIM-Cl salt is evident by a comparison of Figures 4 and 5. Figure 4 shows that DMHDIM-MMT compared to both DMDODA-MMT and DMDHT-MMT.

Figure 10. (A and B) TEM images of DMHDIM-MMT/PA-6 (95:5) processed at 250 °C for 2 min.

Figure 11. (A and B) TEM images of DMHDIM-MMT/PA-6 (95:5) processed at 250 °C for 5 min.

produces primarily one desorption peak at 0.57 min, which is that for thermal desorption of the DMHDIM cation (321.7 amu). The TDMS data for the DMHDIM-Cl salt (Figure 5) reveal a major peak at 0.49 min and a minor peak at 0.56 min. The peak at 0.56 min appears to be the same as the 0.57-min peak in Figure 4, i.e., due to the desorption of the parent ion (321.7 amu). The first peak in the TDMS data for the DMHDIM-Cl salt at 0.49 min contains no parent ion and several main decomposition products. The first of these products results from the loss of one methyl group (presumably from the imidazolium moiety) from the parent cation (321.7 amu) to give the 306.7 amu peak; the second major decomposition product results from the loss of two methyls to give a 291.7 amu product. This difference in stability of the imidazolium cation is presumably due to the anion effect observed by others.27 Here, the chloride ion appears to be much more reactive with the imidazolium than the silicate anion present in the

Figure 12. (A and B) TEM images of DMHDIM-MMT/PA-6 (95:5) processed at 300 °C for 2 min.

MMT. These data raise the issue of the importance of purification after the ion-exchange process used to prepare the DMHDIM-MMT. If residual DMHDIM-Cl ion is present after this reaction, then the thermal stability of the DMHDIM-MMT will be lower. The imidazolium chloride must be completely removed from the MMT product. This is similar to the effects Vaia observed in alkylammonium-treated MMT.27 This chloride ion effect motivated our electrochemical analysis of the IM-MMT materials after the ion-exchange process to ensure complete ion exchange of the chloride by MMT and, hence, provide maximum thermal stability. We succeeded in complete removal of residual chloride by ion exchange to the BF₄ salt or by two or three recrystallizations from acetonitrile.

Although the improved thermal stability is of key importance in making polymer/clay nanocomposites, the organically treated MMT must also be compatible with
the polymeric matrix, monomer, or solvent used to prepare the nanocomposite; otherwise the material will not disperse in the polymer during processing to form a nanocomposite.

The decyl- and hexadecyl-functionalized imidazolium MMTs (DDMIM-MMT and DMHDIM-MMT) were each blended with PS in a mini-twin-screw extruder (5–10 min residence time, 180 °C). XRD of PS/DDMIM-MMT (95:5), with the results shown in Figure 6, reveals only a slight shift in the peak at 5.5–5.8° in the low-angle region of the data. This shift in low-angle peak to higher angles after melting and cooling of organic-modified MMT has been attributed to recrystallization of the treatment. Another explanation is the argument recently invoked to explain the same type of shift to higher angles in the XRD of imidazolium bilayer structures. Here, the shift is explained by a shift from a disordered extended bilayer to an interdigitated crystalline phase.

Presumably either, or both, of these processes could occur during extrusion. In any event, this indicates the poor miscibility of DDMIM-MMT with PS. TEM (Figure 7) confirms that the MMT is dispersed in the resin at the mesoscale (micron scale) but not delaminated. Many large multilayer tactoids, with small d spacings, remain, with very few single delaminated layers.

The miscibility of the hexadecyl-functionalized imidazolium MMT (DMHDIM-MMT) with PS, however, appears to be better than that of the decyl-functionalized imidazolium MMT. The XRD (Figure 6) of the PS/DMHDIM-MMT (95:5) sample processed under the same conditions (180 °C, 5 min) produced new peaks at 2.8 and 5.6° in the low-angle region. We propose the same recrystallization explanation as above for the peak at 5.6°; however, the peak at 2.8° is due to intercalation of PS. TEM (Figure 8) reveals good intercalation of the PS in the MMT. Many individual layers appear in the TEM images, as well as many smaller (3–7-layer) tactoids with expanded layer spacings. The same sample, PS/DMHDIM-MMT, was also prepared using solution blending in toluene with shear supplied via sonication. After solvent evaporation, characterization by XRD showed loss of the low-angle peak for the DMHDIM-MMT sample at 4.75°. The TEM image displayed in Figure 9 reveals considerable exfoliation of the MMT, with some double- and multilayer tactoids still remaining. Previously, Manias showed that, using quaternary alkylammonium-treated MMT to prepare PS/MMT nanocomposites, one obtains the same degree of exfoliation whether the mixing is done in an extruder or in toluene solution. This result leads to the conclusion that the extrusion conditions were not fully optimized for preparation of the DMHDIM-MMT/PS nanocomposites.

We also melt blended PA-6 with DMHDIM-MMT. Here, we obtained fully exfoliated nanocomposites. In these experiments, also done in a mini-extruder, four different processing conditions were evaluated to find the best conditions. The TEM images of these samples are shown in Figures 10–13. From these images, it is apparent that the DMHDIM-MMT exfoliated in the PA-6 matrix; however, it is difficult to differentiate which processing conditions produced the best nanocomposite. The XRD data shown in Figure 14 might lead one to conclude that the sample processed at 250 °C for 2 min is the best because no peak is observed; however, there might be other reasons, aside from exfoliation, for the lack of order and, hence, lack of a low-angle peak. The samples processed longer and/or at higher temperature do show a low-angle peak shifted to larger d spacings relative to that of DMHDIM-MMT. The TEM images of these samples clearly show the majority of

![Figure 14. XRD data for DMHDIM-MMT and DMHDIM-MMT/PA-6 (95:5) processed under four different sets of conditions.](image-url)
the MMT exfoliated in the PA-6 matrix; therefore, the
low-angle peaks (3–4° 2θ) might represent only a minor
fraction of the MMT in the nanocomposites or they
might also be higher-order reflections. Although there
are other emerging methods for characterizing the
degree of mixing in nanocomposites, we have had the
most direct experience utilizing a quantitative NMR
method recently developed by VanderHart et al.9 Be-
cause this method was developed using PA-6/MMT
nanocomposites we felt it was uniquely suited to char-
acterize these PA-6/MMT nanocomposites. Other meth-
ods such as small-angle X-ray scattering (SAXS)38 and
rheological measurements39,40 would also serve to com-
pliment the XRD and TEM data. This NMR method
focuses on the measurement of the proton longitudinal
relaxation rates (T1H) of the nanocomposites. The shorter
the T1H, the better the mixing of the MMT in the PA-6.
The data for these samples are shown in Table 4. From
the NMR data, it appears that the sample processed at
250 °C for 5 min is the most homogeneous, because it
has the shortest T1H.

<table>
<thead>
<tr>
<th>PA-6 and PA-6 nanocomposites</th>
<th>T1H (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DMHDIM-MMT/PA-6 at 250 °C/2 min</td>
<td>361</td>
</tr>
<tr>
<td>DMHDIM-MMT/PA-6 at 250 °C/5 min</td>
<td>343</td>
</tr>
<tr>
<td>DMHDIM-MMT/PA-6 at 300 °C/2 min</td>
<td>388</td>
</tr>
<tr>
<td>DMHDIM-MMT/PA-6 at 300 °C/5 min</td>
<td>381</td>
</tr>
<tr>
<td>pure PA-6a</td>
<td>529</td>
</tr>
<tr>
<td>medium exfoliation of MMT in PA-6a</td>
<td>400</td>
</tr>
<tr>
<td>very good exfoliation MMT in PA-6a</td>
<td>327</td>
</tr>
</tbody>
</table>

a Results from previous publications.9

The physical, thermal, and flammability properties of
these imidazolium-treated MMT nanocomposites are
in the process of being characterized. Furthermore, we
are also exploring the use of trialkylimidazolium-treated
layered silicates for preparing nanocomposites with
other resins with processing or curing temperatures
above 250 °C (PET, syndiotactic PS, cyanate esters, etc.).

Conclusions

This work has demonstrated the superior thermal
stability of trialkylimidazolium-treated layered silicates.
The hexadecyl-functionalized imidazolium (DMHDIM-
MMT) material shows excellent compatibility with PA-6
and, hence, forms high-quality PA-6/MMT nanocom-
posites. The DMHDIM-MMT also forms mixed interca-
lated and exfoliated nanocomposites with PS. Additional
efforts to explore the use of IM-treated layered silicates
might show them to be useful for the preparation of
nanocomposites from thermoplastic polymers with high
processing temperatures and for thermostet polymers
with high cure temperatures.

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