Real-time investigation of crystallization in nylon 6-clay nano-composite probed by infrared spectroscopy

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Via time-resolved FTIR, we examined the real-time investigation of the structural change in molecular chain of nylon 6 during crystallization of neat nylon 6 and the corresponding nano-composite (N6C3.7) having fully exfoliated structure. The neat nylon 6 predominantly formed α-phase in the crystallization temperature (Tc) range of 155–195 °C. For N6C3.7 crystallization at low Tc range of 150–168 °C, the network structure formed by the dispersed clay particles still affected chain folding of nylon 6, the formation of the γ-phase was dominant. The crystallization took place so rapidly (less than 1 s) without induction time of crystallization. At high Tc range (≈177–191 °C), the stable growth of the α-phase crystal coexisting with γ-phase occurred in N6C3.7 crystallization. The growth mechanism in the subsequent crystallization processes (amides IIIα and IIIγ) was virtually the same in both N6C3.7 and neat nylon 6.


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1. Introduction
Nylon 6 was crystallized extensively in the γ-form in the nanocomposites [1–13]. The phase is preferentially crystallized in the γ polymorph in the presence of the montmorillonite (MMT) and/or the organically modified MMT (organo-clay). They reported that similar crystal lattices between clay and the γ polymorph, and the large flat surface of the clay are the key factors to interact between polymer and inorganic materials. The epitaxial crystallization was also revealed from the transmission electron microscopic images [9,12]. That is, the clay surface induces kinetically favored γ-phase formation. In the pseudo-hexagonal γ-form crystals, the molecular chains have to twist away from the zigzag planes to form hydrogen bonds among the parallel chains in the crystals giving rise to lesser inter-chain interactions compared with the monoclinic α-form, which are all-trans and packed in more stable antiparallel-chain arrangement of hydrogen bonds [13].

Besides the nucleating effect, the clay surfaces (silicate layers) can also clearly retard the crystal growth process [6,14]. The clay-induced reduction of the crystallization rate is due to the disruption of lamella growth by the well dispersed MMT particles [6]. Homminga et al. [15] suggested that the silicate layers hinder the diffusion of polymer chains to the crystal growth front and impurity migration away from the growth front takes place. Mittner et al. [16] reported that the interaction between nylon 6 and MMT reduce the mobility of the polymer chain which benefits to form the γ-phase, depending on the nature of the organo-clay (MMT-intercalant). Furthermore, once a percolated network of the dispersed MMT particles is formed, the networks retard crystalline capability as reported by our previous paper [12] and Wang et al. [17].

In our previous paper [12], we have shown that the lamellar growth of the γ-phase crystal took place on both sides of the dispersed MMT particles. The overall crystallization rate, including the nucleation, on nylon 6 increased after nano-composite formation as compared with that of the neat nylon 6.

The formation of the network structure owing to the dispersed MMT particles affected the lamellar growth of the γ-phase crystal.

Despite extensive studies of the polymer crystallization in nylon 6-based nano-composites, the correlation between the nucleating effect and the growth mechanism of the different polymorphism (γ-phase) underlying these observation is not very well explored in the literature. In this regard, we need to pin down the crystallization kinetics of the nylon 6-clay nano-composites.

To better understand the kinetics, time-resolved Fourier transform infrared spectroscopy (FTIR) is a very powerful tool on the basis of the change in the molecular conformation of the nylon 6 during isothermal crystallization. It motivated us to investigate the time development of the different polymorph in the nylon 6 upon nano-composite formation. Recently, a full analysis of the crystallization of nylon 6 in nano-composites has been done by using FTIR [10], but not by time-resolved FTIR analysis.

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In the present paper, via time-resolved FTIR, we examined the real-time investigation of the change in the molecular conformation of the nylon 6 during crystallization of neat nylon 6 and the corresponding nano-composite. Knowledge of such a comparison should also be useful in assessing how does the nano-filler control the crystallization behavior and the different polymorphism of nylon 6 matrix in the clay-based nano-composites?

2. Experimental section

2.1. Materials

The nylon 6-based nano-composite was kindly supplied by Dr. A. Usuki of Toyota Central R&D Labs., Inc. It was synthesized by in-situ polymerization of ε-caprolactam in lauryl ammonium intercalated MMT in the presence of small amount of 6-aminocaproic acid (the number- and weight-average molecular weights \( M_w \) and \( M_n \) of N6C3.7 are 19.7 \( \times 10^3 \) g/mol, and 45.0 \( \times 10^2 \) g/mol, respectively) [9].

The amount of inorganic montmorillonite (MMT) part, obtained from the burning out of organic part, was 3.7% and was designated as N6C3.7. The \( M_w \) and \( M_n \) of pure nylon-6 polymerized in bulk were 21.7 \( \times 10^3 \) g/mol and 44.0 \( \times 10^2 \) g/mol, respectively.

The nano-composite structure of N6C3.7 exhibited well-exfoliated structure. Details regarding the structure analysis can be found in our previous paper [12].

2.2. Differential scanning calorimetry (DSC)

The crystallized specimens were characterized by using temperature-modulated DSC (TMDSC) (TA 2920; TA Instruments) at the heating/cooling rate of 5 °C/min with a heating/cooling cycle of the modulation period of 60s and an amplitude of +/-0.769 °C, to determine the crystallization temperature with cooling from melt (\( T_c \)), the melting temperature (\( T_m \)) and heat of fusion (\( \Delta H_f \)), the DSC was calibrated with Indium before experiments. For the measurement of degree of crystallinity (\( \chi_c \)) prior to TMDSC analysis, the extra heat absorbed by the crystallites formed during cooling had to be subtracted from the total endothermic heat flow due to the melting of the whole crystallites. This can be done according to the principles and procedures described in our previous paper [18]. In the TMDSC experiments, the endothermic heat flow \( \Delta H_{\text{difference}} \) of the initially existing crystallites can be easily calculated as \( \Delta H_{\text{difference}} = \Delta H_{\text{rev}} - \Delta H_{\text{nonrev}} \) where \( \Delta H_{\text{rev}} \) is the endothermic melting (reversible) enthalpy from the reversing heat flow profile and \( \Delta H_{\text{nonrev}} \) is the exothermic ordering/crystallization (nonreversible) enthalpy from the nonreversing heat flow profile appearing in the temperature range of 150–230 °C.

By considering the melting enthalpy of 100% crystalline nylon 6 (\( \gamma \)-phase as 241 J/g and \( \alpha \)-phase as 239 J/g, respectively [19]), we have estimated the value of the \( \chi_c \), calculated as \( \Delta H_{\text{difference}}/\Delta H_{100\%} \) of neat nylon 6 and N6C3.7, and these values are also presented in Table 1.

2.3. Fourier transform infrared spectroscopy (FTIR)

FTIR spectra were collected at 2 cm\(^{-1}\) nominal resolution using a Varian FTS7000 spectrometer equipped with an MCT detector in transmission mode. The spectra were obtained by averaging 32 scans with a mean collection length of 0.6 s per spectrum. The background spectra used for reduction were collected at the same crystallization temperature (\( T_c \)) with the sample. The homogenous mixture of KBr powder and nylon 6 (powder) or N6C3.7 (powder) in the weight ratio 98:2 was prepared. The mixtures were then converted into disks with a thickness of ~0.4 mm by pressing. We have used homemade heating/cooling chamber with a cooling rate of ~1000 °C/min. The disks were placed in a homemade environmental heating chamber, which allowed to reach the desired \( T_c \), in a very short time (~2 s). Each sample was kept at 250 °C for 5 min to erase any thermal history, and it was immediately cooled to \( T_c \). After attaining the \( T_c \), a time-resolved FTIR measurement was carried out in the temperature range of 150–195 °C. The collected data were processed by soft ware (Grams/Al\(^{TM}\), Thermo Galactic Co., USA).

3. Results and discussion

3.1. Crystallization behavior

Fig. 1 shows the TMDSC thermograms of neat nylon 6 and N6C3.7 during cooling from melt state (~260 °C) and 2nd heating with a rate of 5 °C/min. For N6C3.7, the evolution of the crystallization peak is sharp and its maximum is at 186 °C. For the neat nylon 6, this peak is much broader and it appears at 176 °C. The dispersed MMT particles in N6C3.7 promote the crystallization possibly due to the nucleation effect. The estimate values of \( \chi_c \) are shown in Table 1. The presence of MMT particles cause a slight shift in the melting peak to slightly lower temperature (215.4 °C) and lower total crystallinity. In our previous observation [9], the lower \( T_m \) has been attributed to an increased fraction of \( \gamma \)-phase crystallites present in N6C3.7. As seen in Table 1, the higher \( T_c \) value for N6C3.7 (\( T_c = 186.0 °C \)) suggests that the nano-composite is more easily crystallizable as compared with neat nylon 6 (\( T_c = 176.2 °C \)) during the TMDSC cooling process.

3.2. Spectral evolution during crystallization

Fig. 2(a–c) and Fig. 3(a–c) show a typical time-dependent spectral variations of neat nylon 6 and the corresponding nano-composites during crystallization.

![Fig. 1. TMDSC scan for neat nylon 6 and its nano-composite (N6C3.7).](image-url)
composites (N6C3.7) in the region of 3500–400 cm⁻¹ during isothermal crystallization at 180 °C for neat nylon 6 and at 158 °C for N6C3.7. The frequencies and the vibrational assignments for the neat nylon 6 with α-phase and nano-composites with γ-phase are reported in the literature [10]. In Fig. 2(a), a band located at 3450 cm⁻¹, which is assigned to the hydrogen bond-free N–H stretching mode, arise and their intensities decrease until the crystallization is complete (≈1200 s). The time variation of the band at 3335 cm⁻¹ (attributed to the hydrogen bonded N–H stretch: some times called amide A) provides valuable information. This amide A band shifts to the lower frequency side and changes to sharper peak with crystallization time, reflecting the formation of the intermolecular hydrogen bonding. At the same time, hydrogen bond-free band at 3450 cm⁻¹ decreases in intensity. The band of around 3070 cm⁻¹, which is overtone of the amide bonding, shifts oppositely to the higher frequency side, indicating the enhancement of the intermolecular hydrogen bond strength. That is to say, the crystallization takes place.

The bands at 2935 (asymmetric methylene stretching mode: νas(CH₂)) and 2860 cm⁻¹ (symmetric methylene stretching mode: νs(CH₂)) are not related to crystalline fraction [10]. These bands are discussed in terms of the conformational change and lateral chain–chain interaction [20].

The α-phase shows characteristic absorbance peaks at ca. 1660 (amide I: ν(C–O)), 1530 (amide II: coupling of C–N stretching and N–H in plane bending), 1262 cm⁻¹ (amide III) and 1200 cm⁻¹ as seen in the region from 900 to 1800 cm⁻¹ (Fig. 2(b)). The bands at 1370 and 925 cm⁻¹ are CH₂ wagging vibration and intrinsic band to CH₂CO group, respectively. The region from 400 to 900 cm⁻¹ is shown in Fig. 2(c). The amide V(N–H) wagging mode; υ(N–H)) and amide VI ((C=O) wagging mode; υ(C=O)) bands appear at 685 and 580 cm⁻¹ respectively, for the characteristic of α-phase. The band at 730 cm⁻¹ corresponds to the rocking mode of CH₂. Both α- and γ-phases show this band at the same wave number, indicating the same conformation of CH₂ segments when the different hydrogen bonding schemes are taken place.

Qualitatively, the spectra of the γ-phase look similar, but the peak occurs at 625 cm⁻¹ (amide VI), which is shifted from 580 cm⁻¹ for α-phase (Fig. 3(c)). An interesting feature is the contribution to the amide III, which comes from bands at 1232 cm⁻¹ for the γ-phase and 1262 cm⁻¹ for the α-phase [10] (Fig. 3(b)). These results suggest that major γ-phase crystal coexist with small α-phase is formed in the nano-composite systems, while neat nylon 6 predominantly forms α-phase despite of the evolution of the sequence in the amide III. Another interesting feature is the amide A band. The amide A band for the α-phase is sharper than the
The absorbance of the 2220 cm\(^{-1}\) band is regarded as an internal standard because no change was observed during crystallization process. Therefore, the intensity of crystalline-sensitive bands and the internal standard is used to depict the crystalline structure development.

For neat nylon 6 at \(T_c = 167\,^\circ\text{C}\) (supercooling \((\Delta T) = T_m - T_c\) (\(\approx 54.2\,^\circ\text{C}\))) we notice that for the amide A band, the intensity growth and frequency shift to the lower side are very rapid as compared with the intensity changes of the hydrogen bond-free and amide III bands (Fig. 4(a)). This feature is also observed in the temperature range of 155–195 °C for neat nylon 6 crystallization. After the temperature drop to \(T_c\), in an early stage the stronger hydrogen bonding interactions are formed as revealed by a \(\nu(N-H)\) band intensity increment. The formation of \(\alpha\)-phases appear at much later timing. The neat nylon 6 predominantly forms \(\alpha\)-phase in the \(T_c\) range of 155–195 °C.

3.3. Time-dependent spectral evolution

Figs. 4 and 5 show typical examples of the time variation of the reduced intensities for the characteristic bands taken at 167 °C and 158 °C for neat nylon 6 and for N6C3.7, respectively. The upper panels in each figure indicate the temperature drop profile during crystallization of the sample. During the temperature drop of the sample, a very rapid temperature change is observed till the achievement of \(T_c\) (\(~1500\,^\circ\text{C/min}\)). The \(T_c\)s were chosen by referring to the equilibrium melting temperature \((T_m)\). The values of \(T_m\) by Hoffman-Weeks [22] extrapolation based on plotting \(T_m\) versus \(T_c\) were 221.2 °C for neat nylon 6 and 215.8 °C for N6C3.7 [12].

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![Fig. 4. Time variation of intensity for crystalline-sensitive bands of (a) neat nylon 6 taken at 167 °C and (b) N6C3.7 taken at 158 °C. Upper panels indicate temperature drop profile during crystallization of the sample. The arrows indicate the induction time (\(t_0\)) of crystallization and the crystallization half-time (\(t_{1/2}\)). The broken lines indicate onset time of temperature drop (\(t_d\)) and onset time of isothermal crystallization (\(t_c\))(see text for details).](image-url)
For N6C3.7 at $T_c = 158\,^\circ C$ ($\Delta T \equiv 57.8\,^\circ C$) (Fig. 4(b)), just after the temperature drop, the intensity growth for all characteristic bands is more rapid than that of neat nylon 6. In the changes in $t_0$ with $\Delta T (\equiv 24.8–38.8\,^\circ C)$, the N6C3.7 shows no induction time, especially at large $\Delta T (\equiv 47.8–65.8\,^\circ C)$, suggesting that the dispersed MMT particles have a significant contribution to enhance the nucleation of the $\gamma$-phase crystal as mentioned above. This is due to the formation of the network structure owing to the dispersed MMT particles affecting the mobility of nylon molecular chains. At $T_c = 158\,^\circ C$ ($\Delta T \equiv 57.8\,^\circ C$), the N6C3.7 shows no induction time especially at large $\Delta T (\equiv 47.8–65.8\,^\circ C)$ (see text for details).

As shown in Fig. 4(a), we estimated the induction time ($t_0$) and the crystallization half-time $t_{1/2}$ at which the reduced intensity reaches 1/2, we define $1/t_{1/2}$ as a measure of the overall crystallization rate at each characteristic band [23].

In Fig. 6, at all $T_c$ measured here the $t_0$ value of $\nu(N-H)$ band of N6C3.7 is lower than that of neat nylon 6. In the changes in $t_0$ with $\Delta T (\equiv 24.8–38.8\,^\circ C)$, the N6C3.7 shows no induction time especially at large $\Delta T (\equiv 47.8–65.8\,^\circ C)$, suggesting that the dispersed MMT particles have a significant contribution to enhance the nucleation of the $\gamma$-phase crystal as mentioned above. This is due to the formation of the network structure owing to the dispersed MMT particles affecting the mobility of nylon molecular chains. At

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**Fig. 5.** Time variation of intensity for crystalline-sensitive bands of N6C3.7 taken at (a) 158 \(^\circ\)C and (b) 180 \(^\circ\)C. Upper panels indicate temperature drop profile during crystallization of the sample.

<table>
<thead>
<tr>
<th>Temperature ((^\circ)C)</th>
<th>Intensity</th>
<th>Time (s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>140</td>
<td>0.24</td>
<td>0</td>
</tr>
<tr>
<td>160</td>
<td>0.26</td>
<td>50</td>
</tr>
<tr>
<td>180</td>
<td>0.27</td>
<td>150</td>
</tr>
<tr>
<td>200</td>
<td>0.29</td>
<td>250</td>
</tr>
<tr>
<td>220</td>
<td>0.31</td>
<td>350</td>
</tr>
</tbody>
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**Fig. 6.** $\Delta T$ dependence of induction time of $\nu(N-H)$ band for neat nylon 6 and N6C3.7. N6C3.7 shows no induction time especially at large $\Delta T (\equiv 47.8–65.8\,^\circ C)$ (see text for details).
In order to analyze the crystallization growth rate, we assume heterogeneous nucleation and apply the Hoffman–Lauritzen growth rate equation of slow kinetics [25,26] even if we are not using the analysis to determine linear growth rates. The growth rate of crystal can be written as

\[
\frac{1}{T_c DT_f} = \frac{1.71}{10^{2} C^{0.4} K^{2.5}}\text{ (1)}
\]

Overall rate \(\propto \beta_k \exp \left[ -\frac{K_g}{T_c (\Delta T)^f} \right] \)

where \(\beta_k\) is a mobility term, which describes the transportation rate of crystallizable molecules to the growth front; \(f\) is the correction factor given by \(2T_c/(T_m + T_c)\); and \(K_g\) is the nucleation constant, which depends on the crystallization regimes (regime I, single nucleation; regimes II and III, multiple nucleation) [25,27]. At small supercoolings and over limited temperature range, the temperature dependence of growth is determined predominantly by the nucleation term. Accordingly, assuming that the \(1/T_{1/2}\) is a satisfactory measure to the rate of crystallization, we plot the data as \(1/T_{1/2}\) versus \(1/(T_c \Delta T)\): see Fig. 8.

The overall rate of amide IIIa band for N6C3.7 is enhanced for every temperature of measurement as compared with that of amide III for neat nylon 6. The result obviously implies that the number of nucleation sites is larger in N6C3.7, but the growth mechanism in the subsequent crystallization processes are virtually the same in both nano-composite and neat nylon 6 because of almost same slope (~4 × 10^4 K^2), reflecting the apparent nucleation constant. As discussed in the beginning in Fig. 7, unfortunately, we could not discuss the crystallization kinetics of amides III\(\gamma\) and VI for N6C3.7 for a small \(\Delta T\) range.

For neat nylon 6 crystallization, a change in slope at \(1/(T_c \Delta T)^{0.44} \approx 1.71 \times 10^{-4}\text{ K}^{-2}\) corresponding to \(\Delta T \approx 34.2\text{ °C}\) (\(T_c \approx 187\text{ °C}\)) is obvious. The ratio of the two slopes is about 2, suggesting a transition from regime III to regime II at \(T_c \approx 187\text{ °C}\) [27].

4. Conclusions

We have investigated the change in the molecular conformation of the nylon 6 during crystallization of neat nylon 6 and the corresponding nano-composite (N6C3.7) having well-exfoliated structure. The neat nylon 6 predominantly formed \(\alpha\)-phase in the crystallization temperature (\(T_c\)) range of 155–195 °C. N6C3.7 exhibited mainly \(\gamma\)-phase crystal coexisting with \(\alpha\)-phase at low \(T_c\) range (~150–168 °C). It was shown that for the bands of amides III\(\alpha\), III\(\gamma\) and VI for N6C3.7, the crystallization took place so rapidly (less than 1 s) without induction time of crystallization as compared with that of amide III for neat nylon 6 at low \(T_c\) range of 150–168 °C (\(\Delta T = 47.8–65.8\text{ °C}\)). Here, the network structure still hindered chain folding, resulting in the formation of the \(\gamma\)-phase, but the chain mobility at elevated temperatures was comparable to neat nylon 6 as demonstrated by the temperature dependence of the overall crystallization rate.

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