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Development of Microporous Structure and its Application to Optical Film for Cellulose Triacetate Containing Diisodecyl Adipate

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Abstract

Phase separation in plasticized cellulose triacetate (CTA) films is investigated to produce a microporous film that can be used in optical devices. Hot-stretched CTA films containing diisodecyl adipate (DIDA) show negative orientation birefringence similar to the hot-stretched pure CTA. After extracting DIDA from the stretched films by immersion into an organic solvent, however, the films exhibit positive birefringence. Moreover, the magnitude of the birefringence increases with the wavelength, known as extraordinary dispersion, which is an essential property in the preparation of an ideal quarter-wave plate. Numerous ellipsoidal pores with micro-scale were detected in the film after the immersion, indicating that DIDA were segregated and formed ellipsoidal domains in the CTA matrix during annealing and stretching. These results indicate that extraordinary wavelength dispersion is given by the combinations of orientation birefringence from CTA and form birefringence from micropores. Furthermore, it was found that annealing time and stretching condition affect the phase separation as well as the shape and size of pores.

Keywords: Cellulose triacetate; Optical Film; Birefringence; Microporous structure; Blends
1. Introduction

Microporous materials have been studied intensively these days because of their application in the industry and their future potential of additional functions for various products (Silverstein et al., 2011). In general, micropores are responsible for some advantages such as increase in the surface area and penetration distance from one surface to another, besides weight saving. Therefore, microporous materials are used as absorbent, catalyst, and so on (Wakao & Smith, 1964; Arenas & Crocker, 2010). Moreover, microporous materials are expected to expand the applications recently, because advanced functions attributed to micropores have been discovered. As an example of their application in thermal insulation, it is desired that the pore size should be smaller than the mean free path of air, i.e., ca. 60 nm (Clyne et al., 2006). Moreover, micropores having appropriate size are also required for biomaterials such as vehicle for drug and gene delivery (Selvam et al., 2009). In the case of the Lithium ion battery, a separator has to have microporous structure, which should provide the self-shuttering function at high temperature (Love, 2011; Phulkerd et al., 2013). These applications indicate that well-controlled micropores have a great potential to produce functional materials, and it is necessary to control the size and amount of micropores in order to be used in such applications.

There are several methods to produce porous materials: fiber bonding, solvent casting/particulate leaching, gas foaming, plastic deformation, and phase separation/emulsification (Mikos & Temenoff, 2000; Selvam et al. 2009; Phulkerd et al., 2013; Samthong et al., 2014). In the phase separation method, a component of dispersed phase is removed after separation to produce
pores. Therefore, the miscibility is important to control the structure.

As well known, free energy of mixing $\Delta G_{\text{mix}}$ is expressed as,

$$\Delta G_{\text{mix}} = \Delta H_{\text{mix}} - T\Delta S_{\text{mix}}$$

(1)

where $\Delta H_{\text{mix}}$ is the mixing enthalpy and $\Delta S_{\text{mix}}$ is the mixing entropy. Equation (1) demonstrates that the temperature and the interaction parameter which decides $\Delta H_{\text{mix}}$ are the basic parameters to be considered. These factors are important for the formation of pores. For example, phase separation should occur by spinodal decomposition in order to obtain large number of fine pores (Olmsted et al., 1998; Matsuba et al. 1999). In addition, phase separation can occur by several factors other than temperature change and interaction of materials (Rangel-Nafaile et al., 1984; Lee et al., 2004).

Specifically, the orientation of polymer chains by stretching causes the phase separation originating from entropy loss of miscible system, i.e., the so-called flow-induced phase separation (Yanase et al., 1991). On the contrary, some polymer blends show flow-induced mixing by the reduction of concentration fluctuation (Mazich & Carr, 1983; Hindawi et al., 1990). In order to control the pore growth, the effect of these factors must be revealed.

Here, we focus on the mechanism of phase separation to produce the porous material. Cellulose triacetate (CTA), one of the general-purpose polymers for optical films, is used as a matrix polymer. Because CTA exhibits excellent transparency and heat resistance (Songsurang et al., 2012), the films are used for optical applications in industry. In this paper, the micropores in an oriented CTA film are provided using a plasticizer which has poor miscibility with CTA, and this film can be employed as a
multi-band wave plate, one of the functional optical films required for an advanced display. In this study, the microporous structure using heat-induced and flow-induced phase separation is investigated.

2. Experimental

2.1. Materials

Cellulose triacetate (CTA) (Daicel, LT-35) was employed as the matrix polymer. The degree of acetylation per a pyranose unit is 2.9. The weight-average $M_w$ and the number-average molecular weights $M_n$ were $1.3 \times 10^5$ and $3.5 \times 10^5$, determined by a gel permeation chromatography (Tosoh, HLC-8020) with TSK-GEL® GMHXL as a polystyrene standard. Commercially available plasticizers such as diethyl phthalate (DEP) and diisodecyl adipate (DIDA), which are involatile liquid, were used in this study without further purification. Both of them were produced by Daihachi Chemical Industry. The structure and properties of CTA and plasticizers are shown in Figure 1 and Table 1. The values of refractive indices at 633 nm of the plasticizers are close to that of CTA at room temperature.

The solubility parameters calculated by the group contribution method (Grulke, 2003) indicate that DIDA has poor miscibility with CTA as compared with DEP.

[Figure 1] [Table 1]
2.2. Film Preparation

The films were prepared by a solution-cast method. CTA and plasticizers with 10/1 weight ratio were dissolved into mixture solvent of dichloromethane and methanol with 9/1 weight ratio. It was confirmed that the solvent dissolves CTA rapidly. The solution containing 4 wt% of CTA was evaporated at room temperature to obtain the films with 70 µm thickness.

The films were uni-axially stretched at various strain rates of 0.01, 0.05, and 0.10 s\(^{-1}\) by a tensile machine (UBM, DVE-3) with a temperature controller. The hot-stretching was performed after holding at the drawing temperature (\(T_{\text{draw}}\)) for 10 min. \(T_{\text{draw}}\), shown in Table 2, was decided to obtain the same stress level for all samples. The hot-stretched films were immediately quenched by cold air blowing at a draw ratio of 1.5 to avoid relaxation of molecular orientation. Furthermore, the annealing treatment of non-stretched films was performed at 208 °C, i.e., \(T_{\text{draw}}\) for CTA/DIDA, for 1, 10, and 20 min to clarify the effect of thermal history on the morphology development.

The films were immersed into methanol for 6 hours to remove plasticizers. Then, they were dried in vacuo at room temperature. The removal of plasticizers was confirmed from spectra of infrared absorption spectrometry (Perkin Elmer, Spectrum 100).

2.3. Measurements

Dynamic mechanical analysis (DMA) for the solution-cast films of CTA and plasticized CTA was performed to measure oscillatory tensile modulus at 10 Hz as a function of temperature using a
dynamic mechanical analyzer (UBM, DVE-E4000) from -100 to 250 °C with a heating rate of 2 °C min⁻¹. The glass transition temperature $T_g$ was estimated from a peak temperature of tensile loss modulus $E''$ and shown in Table 2.

The birefringence $\Delta n$ of films was measured by an optical birefringence analyzer (Oji Scientific Instruments, KOBRA-WPR) as a function of wavelength by changing color filters. Prior to the measurement of birefringence, the films were kept in a humidic chamber (Yamato, IG420) at 25 °C and 50 %RH for one night to avoid the moisture effect on the optical properties (Abd Manaf et al., 2011a). Birefringence is calculated by the following equation,

$$\Delta n = n_x - n_y$$  \hspace{1cm} (2)

where $x$ is the stretching direction and $y$ is the vertical direction from $x$ in plane.

The light transmittance of films was measured by an ultraviolet-visible absorption spectroscopy (Perkin Elmer, Lambda 25) with the wavelength from 200 nm to 700 nm at room temperature. The total transmittance in the visible wavelength, i.e., 400-700 nm, was calculated to evaluate the transparency of the films.

The morphology of the films was observed by a scanning electron microscope (SEM) (Hitachi, S4100) with an acceleration voltage of 20 kV. Prior to the SEM observation, the surface of films was coated with Pt-Pd by an ion-sputter machine (Hitachi, E1010).
3. Results and Discussion

3.1. Miscibility of plasticizers with CTA

The miscibility of CTA and the plasticizers was examined by the dynamic mechanical properties. Figure 2 shows the temperature dependence of tensile storage moduli $E'$ and $E''$ for the solution-cast films of pure CTA and CTA with 10 phr of the plasticizer. The dynamic mechanical properties of CTA/DEP demonstrate that $T_g$ of CTA is shifted to lower temperature by DEP addition, suggesting the plasticizing effect of DEP. On the other hand, DIDA hardly affects $T_g$. Considering that the solidification temperature of DIDA is lower than that of DEP, the result indicates that DIDA is immiscible with CTA. Furthermore, it should be mentioned that the CTA/DIDA film becomes opaque after the measurement of the dynamic mechanical properties owing to the light scattering originated from the phase separated morphology, although the virgin sample without thermal history, i.e., the solution-cast film, is transparent. These results indicate that the CTA/DIDA blend has lower critical solution temperature. During the measurement, DIDA is segregated and thus forms domains at high temperature. In other words, DIDA is dissolved into CTA in the molecular scale prior to the thermal history of the dynamic mechanical measurement. As a result, the magnitude of $E'$ in the glassy region for the CTA/DIDA film is almost identical to that for the CTA/DEP film. Although there is a slight possibility that a small amount of DIDA is segregated at the final stage of evaporation for CTA/DIDA, the domains, if there, could be significantly small.

[Figure 2]
3.2. Orientation birefringence of CTA and plasticized CTA

The films were stretched above $T_g$. As well known, the stretched films of polymeric materials show the orientation birefringence originated from polarizability anisotropy by molecular orientation. In order to compare the orientation birefringence of stretched films, the stress at the hot-stretching was adjusted to be the same by adjusting the drawing temperature. This is reasonable because the orientation birefringence $\Delta n_o$ of a polymer is proportional to the stress $\sigma$ as follows (Treloar, 1958),

$$\Delta n_o = C \sigma$$

where $C$ is the stress-optical coefficient.

The stress-strain curves of CTA and plasticized CTA at a strain rate of 0.05 $s^{-1}$ are shown in Figure 3. All films show the same stress level, indicating that the orientation of CTA chains in the blend is identical to that in the pure CTA.

The birefringence of the stretched films is shown in Figure 4. The hot-stretched CTA/plasticizer films show negative birefringence with ordinary wavelength dispersion, as similar to that of the pure CTA film; the absolute value of the birefringence decreases with wavelength. Furthermore, the addition of DEP greatly affects the birefringence. This is attributed to the Nematic interaction that occurs in a miscible system (Abd Manaf et al., 2011b). In the case of DIDA, this phenomenon is not so obvious as compared with DEP. This is reasonable because DIDA is not fully miscible with CTA.
at the drawing temperature. Instead of the cooperative molecular orientation, the segregated DIDA droplets are deformed to the stretching direction. As a result, form birefringence, originated from anisotropy of refractive indices in anisotropic structure, is expected (Born & Wolf, 1964). However, the refractive indices of CTA and plasticizers are almost the same as shown in Table 1. Therefore, the form birefringence originated from ellipsoidal DIDA droplets hardly affects the total birefringence of a stretched film. Moreover, this result indicates that both plasticizers cannot change the wavelength dispersion of CTA from ordinary wavelength dispersion.

3.3. Removal of plasticizers

The plasticizers were extracted from the stretched films by immersion into methanol. Although the transparency barely changes by the methanol immersion followed by the drying process, which is shown later, numerous prolonged pores to the stretching direction are detected by SEM in the CTA/DIDA film as shown in Figure 5(a). It is confirmed by the weight measurements and FT-IR spectra that DIDA is fully extracted by the solvent. Therefore, the film in Figure 5(a) is composed of pure CTA with the voids. Considering that voids are hardly detected in the CTA/DEP film after methanol immersion as shown in Figure 5(b), the voids in Figure 5(a) are the dispersed phase of DIDA before immersion.
It is known that the homogeneous distribution of ellipsoidal dispersions with different refractive index provides the form birefringence. Although the form birefringence prior to the methanol immersion can be neglected due to the small difference in the refractive indices, it becomes a large value after the extraction process. Figure 6 shows the birefringence of the stretched film before/after the extraction. Birefringence of a pure CTA film is not affected by the immersion, suggesting that the orientation of CTA chains is not relaxed by the process. In the case of the miscible system, the birefringence approaches that of pure CTA as discussed in the previous paper (Abd Manaf et al., 2011b).

On the contrary, in the case of the immiscible CTA/ DIDA blend, the birefringence of the film dramatically changes from negative to positive with extraordinary wavelength dispersion; birefringence increases with the wavelength. This result indicates the great contribution of form birefringence originated from anisotropic pores as illustrated in Figure 7. Since the extraordinary wavelength dispersion is required in industry for advanced optical devices such as organic electroluminescence (EL) display, optical pick-up units, and three dimensional (3D) display (Yamaguchi et al., 2009), the results in Figure 6 should be noted.

Assuming that the refractive index of the void, i.e., air, is unity, the form birefringence $\Delta n_F$ ascribed to voids is given by the following equation (Richter et al., 1995),

$$\Delta n_F = A - \frac{n}{A}$$ \hspace{1cm} (4)

$$A = \left[(1 - f)n^2 + f\right]^{1/2}$$ \hspace{1cm} (5)
where $n$ is the refractive index of CTA and $f$ is the filling factor of voids, i.e., the ratio of the void length to the distance between neighbor voids in the stretching direction.

Since the stretching condition affects the shape of voids, i.e., the filling factor, it has to be considered seriously to control the birefringence.

3.4. Mechanism of phase separation

The growth of DIDA droplets at the heating and/or stretching processes is investigated by SEM observation using the films after the DIDA extraction. As shown in Figure 8(a), pores are not found in the cast-film, which corresponds with the high level of transparency. After annealing at the drawing temperature, small pores are detected in the film with a low volume fraction, indicating that the phase separation occurs at high temperature to some degree. Moreover, the pores become larger after stretching with the deformation to the stretching direction. The enlargement of each void and the total volume fraction of voids indicates the flow-induced phase separation (Yanase et al., 1991). Therefore, the thermal history and the stretching rate should be considered at the hot-stretching to control the porous structure, and thus, the birefringence.

The effect of heating and stretching on phase separation is investigated by light transmittance using the films before/after extracted DIDA. Table 3 shows the total transmittance in the range of
400-700 nm of UV-Vis spectra of the annealed films. As shown here, the transmittance of CTA/DIDA decreases with the annealing time, as compared with that of pure CTA. It suggests that DIDA is segregated from CTA and forms domains during heating.

[Table 3]

Figure 9 shows the stress-strain curves of pure CTA and CTA/DIDA films with various strain rates: 0.01, 0.05, and 0.10 s\(^{-1}\) after 10 min annealing. As increasing the strain rate, the stress level increases, suggesting that the chain orientation of CTA is enhanced. The total transmittance of the stretched films after extraction is around 87-88%, irrespective of the strain rates. The value is slightly lower than that for the unstretched film, which corresponds with SEM observation.

[Figure 9]

Figure 10 shows the birefringence of the stretched film after extraction. Although the stress level is different, the pure CTA shows almost the same orientation birefringence. Because polarizability anisotropy of the main chain of CTA is small as compared with the acetyl group in side chain (Yamaguchi et al., 2009), the orientation birefringence is hardly affected by the orientation of main chains. In contrast, birefringence of the extracted CTA/DIDA films increases with the strain rate, although it hardly affects the transparency. The enhanced form birefringence will be attributed to the prolonged voids, which is determined by the chain orientation of CTA.

[Figure 10]
4. Conclusion

We focus on microporous structure formation in CTA films utilizing phase separation of a plasticizer in the polymer. The hot-stretched CTA film containing plasticizers shows negative birefringence with ordinary wavelength dispersion, as similar to that of the pure CTA film. However, the birefringence of the films with an immiscible plasticizer, DIDA, changes from negative to positive after the removal of the plasticizer. In addition, the film exhibits extraordinary wavelength dispersion. The SEM observation reveals that the film contains numerous ellipsoidal pores with nanoscale, which provide the form birefringence as a positive value. This is a new material design of the film showing extraordinary wavelength dispersion, i.e., high performance retardation film.

Further, the important phenomena to produce the microporous structure are demonstrated such as phase diagram (LCST) and flow-induced phase separation.

References


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Figure Captions

Figure 1  Chemical structure of (a) CTA, (b) DEP, and (c) DIDA.

Figure 2  Temperature dependence of dynamic tensile moduli such as storage modulus $E'$ and loss modulus $E''$ at 10 Hz for pure CTA and plasticized CTA films with 10 phr of DEP and DIDA.

Figure 3  Stress-strain curves of the films stretched at $T_{\text{draw}}$. The strain rate is 0.05 s$^{-1}$.

Figure 4  Wavelength dependence of birefringence for the pure CTA and plasticized CTA films with a draw ratio of 1.5.

Figure 5  SEM pictures of cut surface of the films after extraction; (a) CTA/DEP and (b) CTA/DIDA.

Figure 6  Wavelength dependence of birefringence for the stretched films before/after extraction; (a) CTA, (b) CTA/DEP and (c) CTA/DIDA. The films were stretched at a draw ratio of 1.5 at a strain rate of 0.5 s$^{-1}$.

Figure 7  Birefringence of the stretched film with ellipsoidal pores.

Figure 8  Growth of porous structure in the CTA/DIDA film; Phase separation occurs at heating and stretching processes.

Figure 9  Stress-strain curves at various strain rates; (a) CTA and (b) CTA/DIDA.

Figure 10 Wavelength dependence of birefringence (a) CTA films after extraction and (b) CTA/DIDA films before/after extraction. The films were stretched at a draw ratio of 1.5 at various strain rates.
<table>
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<tr>
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<th>Plasticizer</th>
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<td></td>
<td></td>
<td>DEP</td>
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<tr>
<td>Solubility parameter ((\text{MPa}^{1/2}))</td>
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<td>20.7</td>
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<tr>
<td>Refractive index ((-))</td>
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<td>1.50</td>
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Table 2  Glass transition temperature and drawing temperature for the films.

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<th>CTA/DEP</th>
<th>CTA/DIDA</th>
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<tbody>
<tr>
<td>$T_g$ (°C)</td>
<td>195</td>
<td>151</td>
<td>192</td>
</tr>
<tr>
<td>$T_{draw}$ (°C)</td>
<td>213</td>
<td>178</td>
<td>208</td>
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Table 3  Total transmittance of annealed CTA/DIDA films with various annealing time.

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<tr>
<td></td>
<td>0</td>
</tr>
<tr>
<td>CTA</td>
<td>92.38</td>
</tr>
<tr>
<td>CTA/DIDA</td>
<td>92.55</td>
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</table>
(a) \[
\left(\begin{array}{c}
\text{OR} \\
\text{OR} \\
\text{OR}
\end{array}\right)
\]
\[R = \text{COCH}_3\]

(b) [Diagram of structure (b)]

(c) [Diagram of structure (c)]
Figure 2

The graph shows the variation of modulus with temperature for different samples:

- **CTA**
- **CTA/DEP**
- **CTA/DIDA**

The modulus is represented on a log scale (log [E' / Pa], log [E'' / Pa]), and the temperature range is from 100 to 250 °C. The graph indicates a decrease in modulus with increasing temperature.
Figure 4

Wavelength /nm

$\Delta n \times 10^{-4}$

- CTA
- CTA/DEP
- CTA/DIDA
Figure 5

(a) CTA/DEP

(b) CTA/DIDA
\[ \Delta n_{total} = \Delta n_F + \Delta n_O \]

Figure 7
Figure 8

- Cast-film
- Annealed film
- Stretched film

- Heating 10 min
- Stretching x 1.5
- Methanol immersion

Images:
- 5 µm
- 5 µm
- 5 µm
Figure 9

213 °C

Strain (MPa)

Strain

208 °C

Strain (MPa)

Strain

CTA 0.01 s⁻¹
CTA 0.05 s⁻¹
CTA 0.1 s⁻¹
CTA/DIDA 0.01 s⁻¹
CTA/DIDA 0.05 s⁻¹
CTA/DIDA 0.1 s⁻¹
Figure 10