

Title	Effect of acetyl substitution on the optical anisotropy of cellulose acetate films
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Citation	Cellulose, 25(8): 4453-4462
Issue Date	2018-06-09
Type	Journal Article
Text version	author
URL	http://hdl.handle.net/10119/16121
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**Effect of acetyl substitution on the optical anisotropy
of cellulose acetate films**

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1 Abstract

2 The effect of acetyl substitution on the optical properties of cellulose acetate (CA) was
3 investigated in the present study. During hot-stretching, the absolute orientation
4 birefringence increased with tensile stress. The stretched films with high acetyl
5 substitution had negative orientation birefringence, whereas those with low acetyl
6 substitution had positive orientation birefringence. Furthermore, there was a great
7 reduction in the rate of decrease in orientation birefringence following the cessation of
8 hot-stretching. The slow relaxation of crystal orientation was responsible for the
9 anomalous optical anisotropy, as confirmed by two-dimensional X-ray diffraction.
10 Moreover, the slow relaxation of orientation birefringence would greatly benefit the
11 preparation of CA optical retardation films by hot-stretching, because it would simplify
12 the precise control of retardation. The stress-optical coefficient in the glassy state was
13 also evaluated, and was found to decrease with the degree of acetyl substitution.

14

15 **Keywords:** Cellulose acetate; Birefringence; Orientation; Refractive index

16

1 **Introduction**

2 Because cellulose esters usually have high optical transparency and high heat resistance,
3 they are often used for optical films such as polarizer protective films and retardation
4 films (Edgar et al. 2004; Sata et al. 2004; Gleisser 2004; Kamide 2005; Necula et al.
5 2010; Yamaguchi & Mohd Edderozey 2013). Cellulose acetate (CA), which is produced
6 by the acetylation of cellulose, is one of the most popular cellulose esters used as optical
7 films because it is inexpensive and has excellent heat resistance. As a consequence of
8 the rapid progression of display systems, advanced CA retardation films are currently
9 produced by hot stretching. Multi-band waveplates, in which birefringence is
10 proportional to wavelength, are one of the best-known examples. Mohd Edderozey et al.
11 (2011a) prepared a multi-band quarter-wave plate using cellulose triacetate (CTA) with
12 a conventional plasticizer (tricresyl phosphate). Nobukawa et al. (2014) and Shimada et
13 al. (2015) also prepared multi-band waveplates by exploiting the form birefringence of
14 cellulose ester films.

15 The degree of acetyl substitution in CA has a major impact on its optical
16 properties. Necula et al. (2010) reported that optical transparency and the refractive
17 index are highly dependent on the degree of acetyl substitution. Yamaguchi et al. (2009)
18 found that the degree of acetyl substitution affects the wavelength dispersion of
19 orientation birefringence. El-Diasty et al. modified the orientation birefringence of CTA
20 by irradiation (2007). Modification by the addition of biomass-based materials has also
21 been proposed as a means of controlling retardation (Nobukawa et al. 2015; Soeta et al.
22 2017). However, the effects of the hot-stretching processing conditions remain unclear.
23 The crystallinity of CTA increases during hot-stretching (Songsurang et al. 2012;
24 Shimada et al. 2017). Therefore, the optimum conditions for hot-stretching should be

1 clarified. The crystallinity of CA approximately corresponds to its acetyl content (Edgar
 2 et al. 2004; Sata et al. 2004). Therefore, in the present study the relationship between
 3 the degree of acetylation and orientation birefringence was examined by the
 4 simultaneous measurement of stress and birefringence during hot-stretching, and after
 5 the cessation of stretching. Structural change was further characterized by
 6 two-dimensional X-ray diffraction analysis. Finally, photoelastic birefringence caused
 7 by stress in the glassy state—which is an important property of optical films—was
 8 evaluated to confirm the effect of acetyl substitution.

9

10 **Experimental**

11 **Samples and Film preparation**

12 Four CA samples with various degrees of acetyl substitution were provided by
 13 the Daicel Corporation. The details of the characteristics are summarized in Table 1.
 14 The numerals in the sample code represent the degree of acetyl substitution.

15

16 Table 1 Characteristics of polymers

	Degree of acetyl substitution	$M_n \times 10^{-4}$	$M_w \times 10^{-4}$
CA2.9	2.93	10.0	31.8
CA2.8	2.85	11.1	35.8
CA2.4	2.44	8.3	20.7
CA2.2	2.18	7.4	20.8

17 Molecular weights: poly(methyl methacrylate) standard

18

19 The films were prepared by a solution-casting method as follows. Each sample
 20 was dissolved in a mixed solvent comprising dichloromethane (CH_2Cl_2) and methanol

1 (CH₃OH) in a 9-to-1 weight ratio, and stirred for 24 h at room temperature. The
2 solutions were poured onto glass plates and left overnight to allow the solvent to
3 evaporate. The obtained films (100 μm thick) were stored in a temperature- and
4 humidity-controlled environment at 25°C and 50% relative humidity (RH) before
5 obtaining the measurements.

6

7 **Measurements**

8 The water content of each sample film was obtained with a moisture meter
9 (Mitsubishi Chemical, CA-200) using the Karl Fischer method, in which the Karl
10 Fischer reagent comprising iodine, sulfur dioxide, a base, and a solvent reacts
11 quantitatively and selectively with water (MacLeod 1991).

12 The refractive index of each sample film was measured as a function of
13 wavelength using a DR-M2 Abbe refractometer (Atago). A halogen lamp with various
14 interference filters was used, which enabled changes in wavelength, and
15 α-bromonaphthalene was used as a contact liquid.

16 The temperature dependencies of tensile storage modulus E' and loss modulus
17 E'' between 30 and 250°C were determined using an E-4000 dynamic mechanical
18 analyzer (UBM). The frequency and heating rate were 10 Hz and 2°C/min, respectively.

19 Rectangular specimens (5-mm wide) were cut from solution-cast films, and
20 uniaxially stretched beyond their glass transition temperatures; stress was measured
21 using a tensile machine with a temperature controller (UBM, DVE-3 S1000). The initial
22 distance between the clamps was 10 mm, and the stretching speeds were 1.0, 0.5, and
23 0.1 mm/min. Therefore, the initial strain rates were 0.1, 0.05, and 0.01 s⁻¹, respectively.
24 The hot-stretching temperature was determined to show similar stress levels for all films

1 at a constant strain rate of 0.05 s^{-1} . Optical retardation was simultaneously evaluated by
2 measuring the light intensities under crossed polars using a polarized laser beam (632.8
3 nm) during stretching; the procedure has been explained in detail in the literature
4 (Yamaguchi et al. 2012). After stretching to a draw ratio of 1.3, some of the films were
5 used to measure the relaxation behavior of both stress and retardation by holding the
6 stretched films at a constant temperature. The films obtained by blowing air were also
7 used to measure the optical retardation at room temperature as a function of wavelength
8 by changing the color filters using a KOBRA-WPR optical birefringence analyzer (Oji
9 Scientific Instruments). Three sample films were prepared as follows: (1) cooled
10 immediately after pre-heating prior to stretching; (2) cooled immediately after stretching
11 to a draw ratio of 1.3; and (3) cooled over 60 s after cessation of stretching. The films
12 were stored at 25°C and 50% RH for 1 day prior to conducting the retardation
13 measurements.

14 Two-dimensional wide-angle X-ray diffraction (2D-XRD) patterns were
15 collected with a graphite monochromatized $\text{CuK}\alpha$ radiation beam using an R-AXIS IIC
16 flat imaging plate detector (Rigaku). The films were exposed for 6 min per shot in a
17 geometrical manner by directing the X-ray beam normal to the film plane.

18 Stress-optical coefficients in the glassy state were evaluated as a function of
19 wavelength at room temperature using the optical birefringence analyzer. Measurements
20 were taken after applying various loads to the sample films.

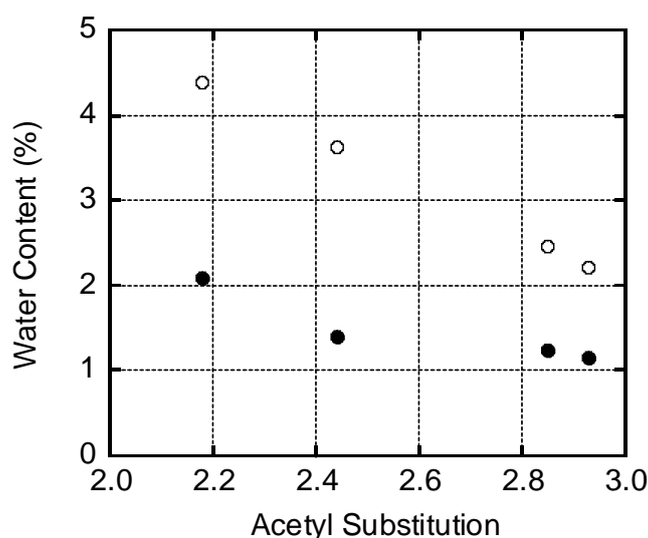
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22 **Results and Discussion**

23 The optical anisotropy of cellulose derivatives is affected by their water content.
24 In general, moisture enhances the optical anisotropy of CA films (Mohd Edeerozey et al.

1 2011b). Therefore, the moisture content of the films was evaluated first. Figure 1 shows
2 the water content of the films used for the optical and mechanical measurements, i.e.,
3 the samples that had been kept in a chamber at 25°C and 50% RH for 1 day. Dried films
4 that had been kept in a vacuum oven at room temperature for 24 h were used as a
5 reference. As expected for CA films, the moisture content, which was high in the films
6 stored at 50% RH, decreased rapidly with the degree of acetyl substitution. Regardless
7 of the storage conditions, the water content of the CA2.2 films was approximately twice
8 that of the CA2.9 films.

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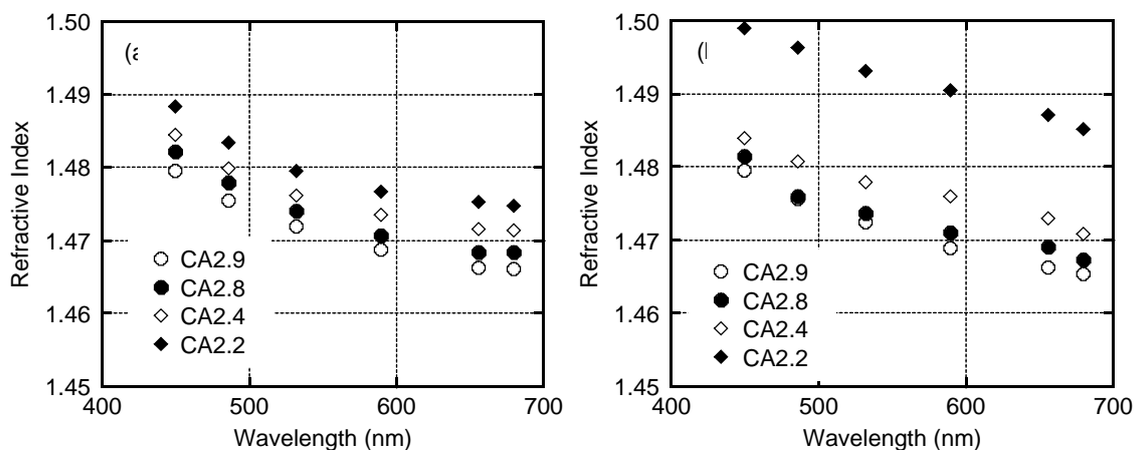
11 Figure 1 Water content in the CA films: (closed circles) after vacuum drying at 25°C for
12 24 hours and (open circles) after keeping in the temperature-and-humidity controlled
13 chamber at 25°C and 50% RH.

14

15 The wavelength dispersion of the refractive index is shown in Figure 2 for the
16 vacuum-dried and moisture-controlled (25°C, 50% RH) samples. The refractive index
17 decreased with increasing acetyl substitution, whereas the wavelength dispersion was

1 barely affected. Figure 3 represents the relationship between the degree of acetyl
 2 substitution and the refractive index at 589 nm (D-line). The refractive index decreased
 3 linearly with the degree of acetyl substitution in the dried samples. In contrast, the
 4 refractive index increased in the moisture-controlled samples, and the increase was most
 5 pronounced in CA2.2. The increase in the refractive index can be attributed to the
 6 absorbed water, which is an interesting phenomenon. Considering that the refractive
 7 index of water is 1.333, the simple addition of the refractive indices of the individual
 8 components—i.e., CA and water—cannot account for the refractive index observed. The
 9 polarizability of CA would be enhanced by the strong hydrogen bonding with water
 10 molecules.

11

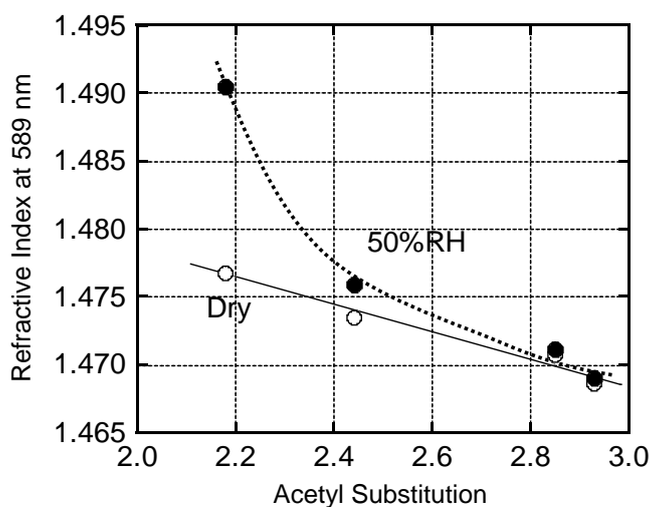


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13 Figure 2 Wavelength dispersion of refractive index for (a) vacuum-dried films and (b)

14 moisture-controlled films at 25°C and 50% RH.

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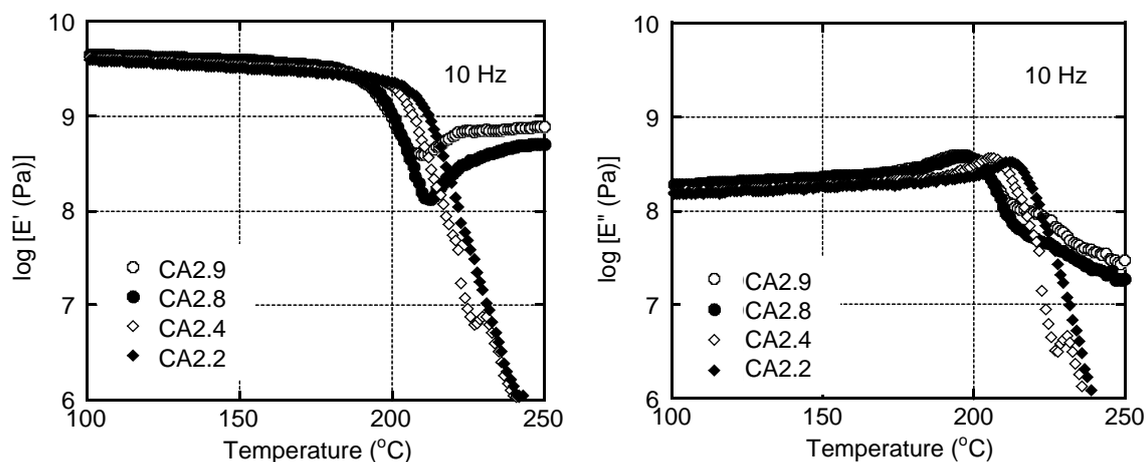
2 Figure 3 Relation between acetyl substitution and refractive index at 589 nm for (open
3 circles) vacuum-dried films and (closed circles) moisture-controlled films at 25°C and
4 50% RH.

5

6 Figure 4 shows the temperature dependence of the tensile storage modulus E'
7 and the loss modulus E'' . Both CA2.2 and CA2.4—i.e., the samples with low acetyl
8 substitutions—exhibited a rapid decrease in E' , which can be attributed to the
9 glass-to-rubber transition. In contrast, the E' curve for the other samples—i.e., CA2.8
10 and CA2.9—featured a plateau, even beyond the glass transition temperature T_g . This
11 was expected because the crystallites act as crosslink points. In other words, a high
12 degree of acetyl substitution increased the crystallinity of the CA films, as reported
13 previously (Edgar et al. 2001; Sata et al. 2004), although the films were transparent. The
14 E' curve for CA2.8 exhibited a slight increase beyond 210°C. This can be attributed to
15 cold crystallization. Because the solution-cast film had a low degree of crystallinity,
16 further crystallization occurred beyond T_g . A similar behavior was detected in CA2.4 at
17 approximately 230°C, indicating that CA2.4 was crystalline, as explained in detail later.
18 The peak temperature in the E'' curve, defined as T_g in the present study, decreased

1 slightly with increasing degree of acetyl substitution: 216°C for CA2.2, 207°C for
 2 CA2.4, 199°C for CA2.8, and 198°C for CA2.9.

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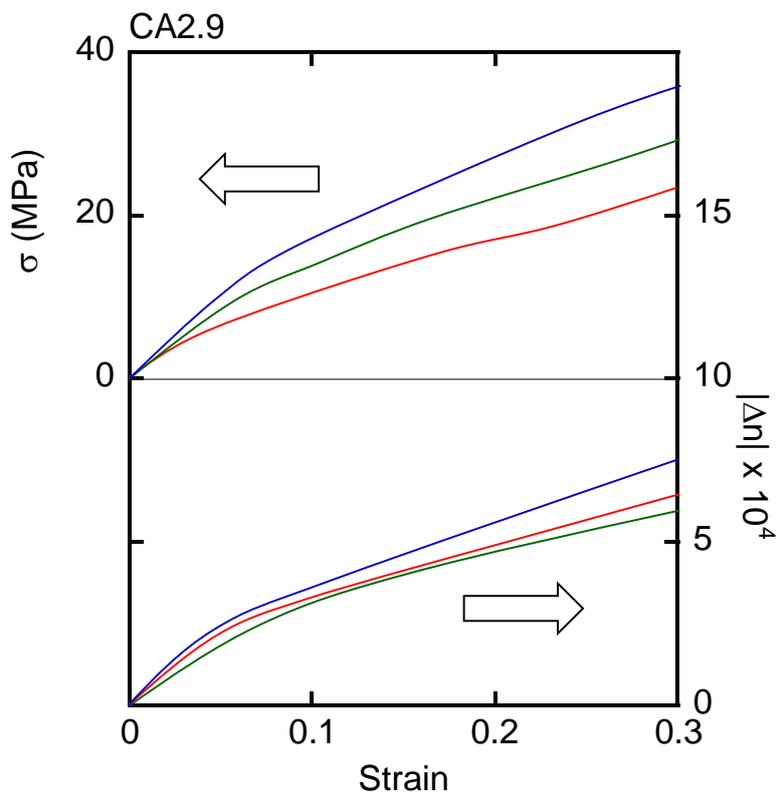


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5 Figure 4 Temperature dependence of dynamic tensile moduli (a) storage modulus E'
 6 and (b) loss modulus E'' .

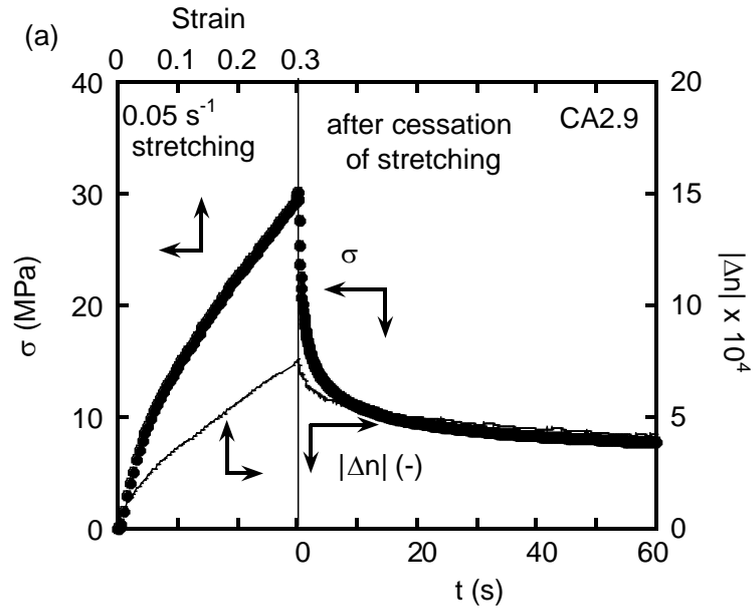
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8 The engineering tensile stress and orientation birefringence growth curves for
 9 CA2.9 during hot-stretching are shown in Figure 5. The birefringence was negative, as
 10 reported previously (El-Diasty et al. 2007; Yamaguchi et al. 2009). The stress level
 11 increased with increasing strain rate, which is the typical mechanical behavior of a
 12 viscoelastic body. Because the degree of main-chain orientation dictates the stress in the
 13 rubbery region, films stretched at the highest strain rate (0.1 s^{-1}) had the highest
 14 orientation birefringence. It should be noted that the orientation birefringence, which is
 15 in general determined by the product of the intrinsic birefringence and the orientation
 16 function, was hardly affected by the strain rate, i.e., the stress level. The result
 17 demonstrated that the stress-optical law in the rubbery region (Kuhn & Gr \ddot{u} n 1942) was
 18 not applicable to sample CA2.9.

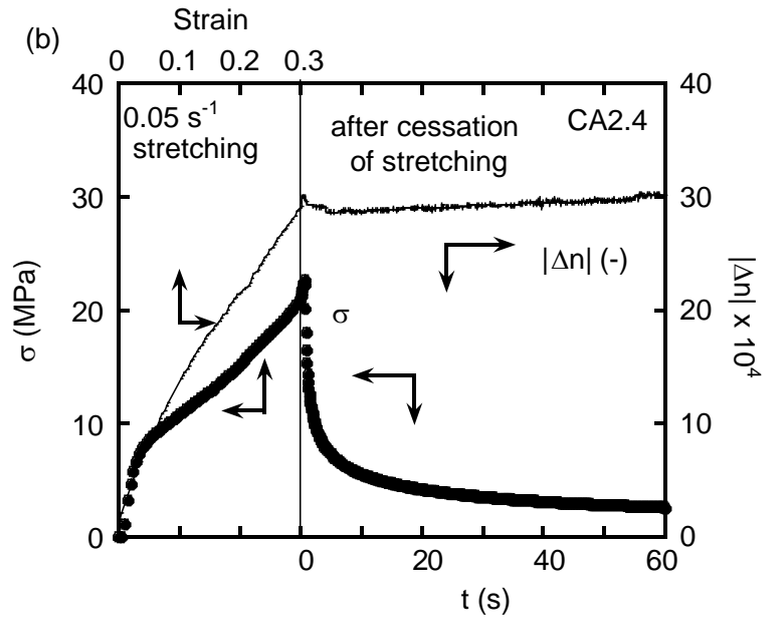


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 2 Figure 5 Growth curves of the tensile stress σ and orientation birefringence Δn during
 3 stretching for CA2.9 at various strain rates. The sign of birefringence was negative.

4
 5 The stress and birefringence decay curves during relaxation, and following the
 6 cessation of stretching at a draw ratio of 1.3 (applied at 0.05 s^{-1}) were evaluated for
 7 CA2.9 and CA2.4. As shown in Figure 6, the birefringence ($t > 0$) did not decrease
 8 much in either sample, although the stress dropped off rapidly. The results also
 9 demonstrated that the stress-optical rule was not applicable to the CA samples. This
 10 anomalous and interesting behavior indicates that the crystalline regions of CA, which
 11 barely exhibit orientation relaxation, are responsible for the orientation birefringence, as
 12 discussed later. From an industrial perspective, this is a great advantage, because it is
 13 easy to provide appropriate retardation by hot-stretching CA films, owing to the slow
 14 relaxation of birefringence after stretching ceases.



1



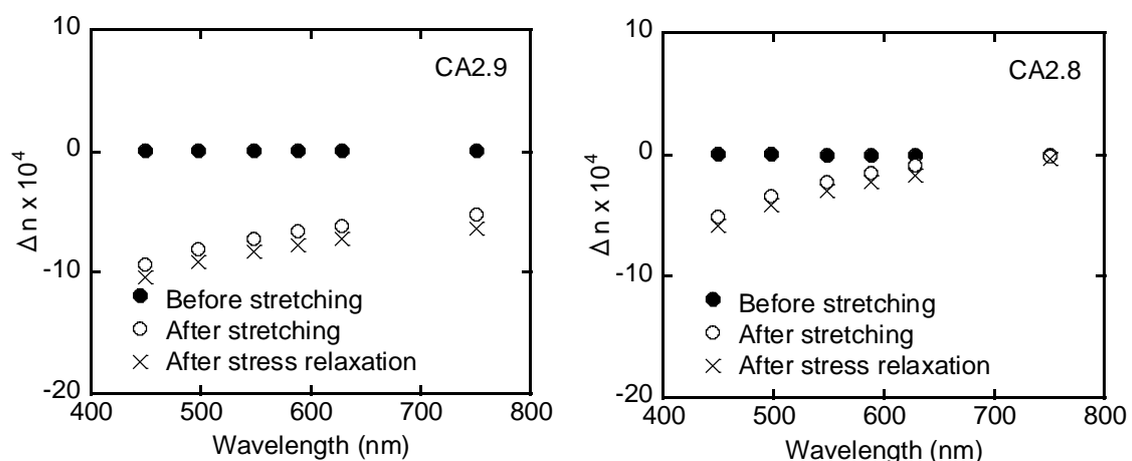
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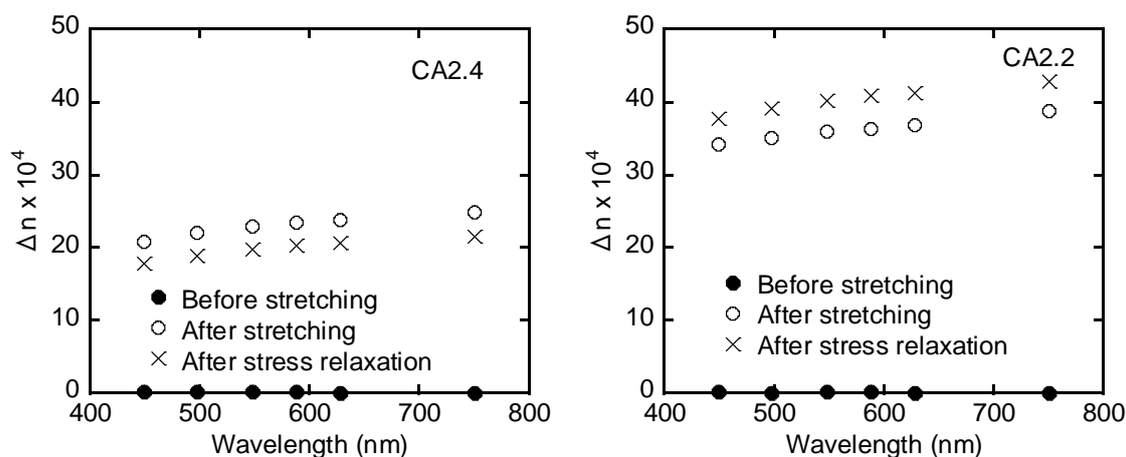
3 Figure 6 Growth curves of the tensile stress σ and orientation birefringence Δn during
 4 stretching at 0.05 s⁻¹ and after cessation of stretching at a draw ratio of 1.3 for (a) CA2.9
 5 and (b) CA2.4. The sign of the birefringence was denoted in the parenthesis.

6

7 The films that had been stretched at 0.05 s⁻¹ were cooled by blowing air after

1 the relaxation process (60 s after the cessation of stretching), and the chucks were
 2 removed from the tensile machine to evaluate the wavelength dispersion of
 3 birefringence. The same measurements were performed using the stretched films cooled
 4 immediately after the cessation of stretching without relaxation. The results are shown
 5 in Figure 7. The birefringence of the films pre-heated at the stretching temperature prior
 6 to stretching is also shown as "before stretching" in Figure 7 (no birefringence).
 7 Because the measurements were performed at room temperature after cooling, the
 8 birefringence of the stretched films shown in Figure 7 is slightly different from the data
 9 in Figures 5 and 6. As seen in Figure 7, we confirmed that there was hardly any
 10 relaxation of orientation birefringence in any of the samples. CA2.8 exhibited negative
 11 and markedly low orientation birefringence with strong wavelength dispersion. In
 12 contrast, CA2.4 and CA2.2 exhibited large and positive orientation birefringence values
 13 with weak wavelength dispersion. Furthermore, CA2.4 and CA2.2 exhibited unusual
 14 wavelength dispersion; i.e., the birefringence increased with the wavelength. This
 15 phenomenon can be explained by the sum contributions of the hydroxyl and acetyl
 16 groups (Yamaguchi et al. 2009).



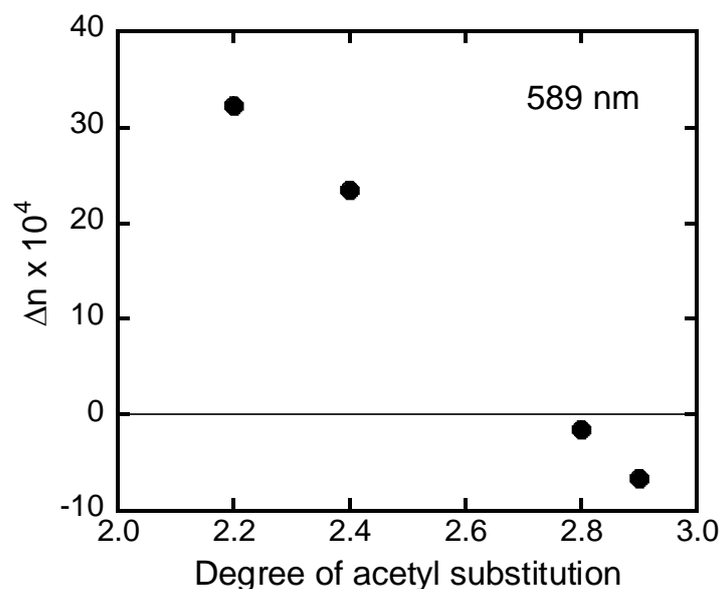


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2 Figure 7 Wavelength dispersion of the orientation birefringence for the films: (closed
 3 circles) after pre-heating prior to stretching, (open circles) immediately after stretching
 4 without relaxation process, and (x) after the relaxation process (60 sec after cessation of
 5 stretching). (a) CA2.9, (b) CA2.8, (c) CA2.4, and (d) CA2.2.

6

7 The orientation birefringence at 589 nm is plotted as a function of the degree of
 8 acetyl substitution in Figure 8. The values were measured at room temperature using the
 9 unrelaxed stretched films. The orientation birefringence increased monotonically with
 10 an increasing number of hydroxyl groups, i.e., low acetyl substitution. This is due to the
 11 strongly anisotropic polarizability arising from the hydroxyl groups in a stretched film.
 12 The direction of the polarizability anisotropy for the samples having a high degree of
 13 acetyl substitution was perpendicular to that of the polymer chain; i.e., the acetyl groups
 14 contribute negative birefringence. Figure 8 indicates that a CA sample with an
 15 appropriate acetyl substitution that is close to 2.8 is free from orientation birefringence.



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2 Figure 8 Orientation birefringence at 589 nm as a function of the acetyl substitution.

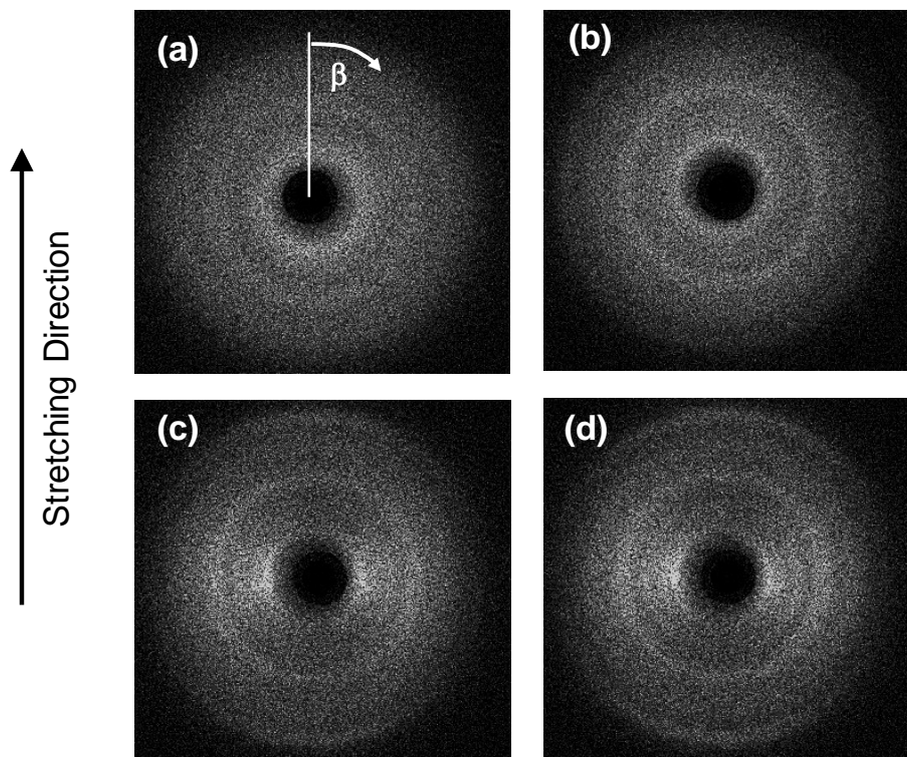
3 The values were measured at room temperature using the stretched films without
4 relaxation process.

5

6 Figure 9 shows the 2D-XRD images of the through views of CA2.4 films: (a) a
7 solution-cast film; (b) a film after pre-heating prior to stretching; (c) a film after
8 stretching at 0.05 s^{-1} without relaxation; and (d) a film after relaxation for 60 s. There
9 were two noticeable rings in the 2D-XRD images of the solution-cast film. These
10 diffraction peaks are attributable to the (010) and (020) planes, respectively (Roche et al.
11 1978; Siroski et al. 2013). The peak intensities increased slightly following pre-heating
12 because of crystallization. After stretching, the peaks were clearly visible on the equator
13 because the polymer chains orient to the stretching direction. Moreover, the peaks did
14 not diminish even after relaxation, demonstrating that the orientation of the crystalline
15 phase remained. The result suggests that the orientation of amorphous chains does not
16 play an important role in birefringence, although the stress in the rubbery state is

1 governed by the orientation of amorphous chains (Read 1975).

2



3

4

5 Figure 9 2D-XRD profiles of CA2.4: (a) solution-cast film, (b) after pre-heating prior
6 to stretching, (c) immediately after stretching without relaxation process, and (d) after
7 the stress relaxation process (60 sec after cessation of stretching).

8

9 Figure 10 shows the azimuthal intensity distribution at $2\theta = 7.8^\circ$, i.e., the (010)
10 plane, for all CA samples. The strongest peaks were detected on the equator, i.e., $\beta = 90$
11 and 270° , revealing the chain orientation in the crystals. The figures demonstrate that
12 there was hardly any orientation relaxation of the crystalline chains in any of the
13 samples including CA2.2, i.e., the sample with low crystallinity.

14

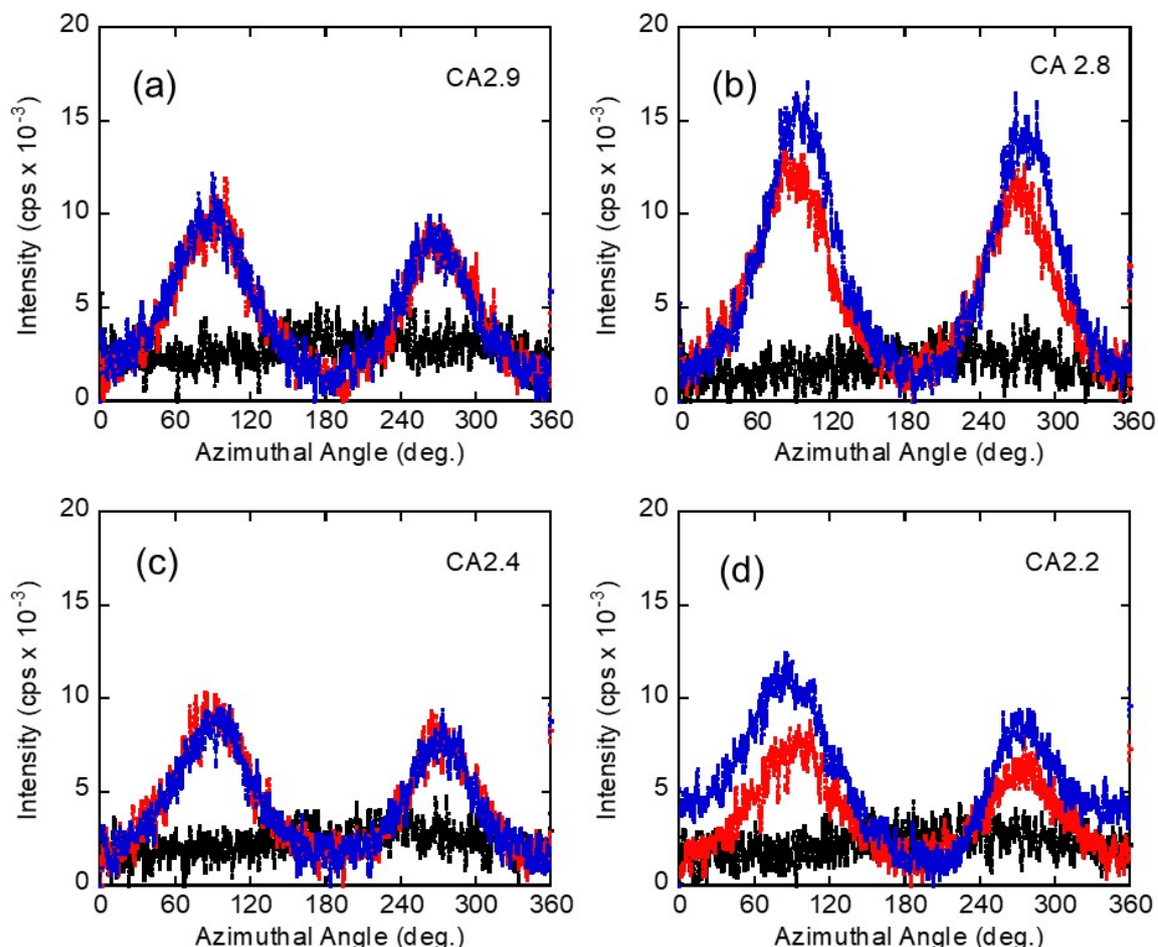
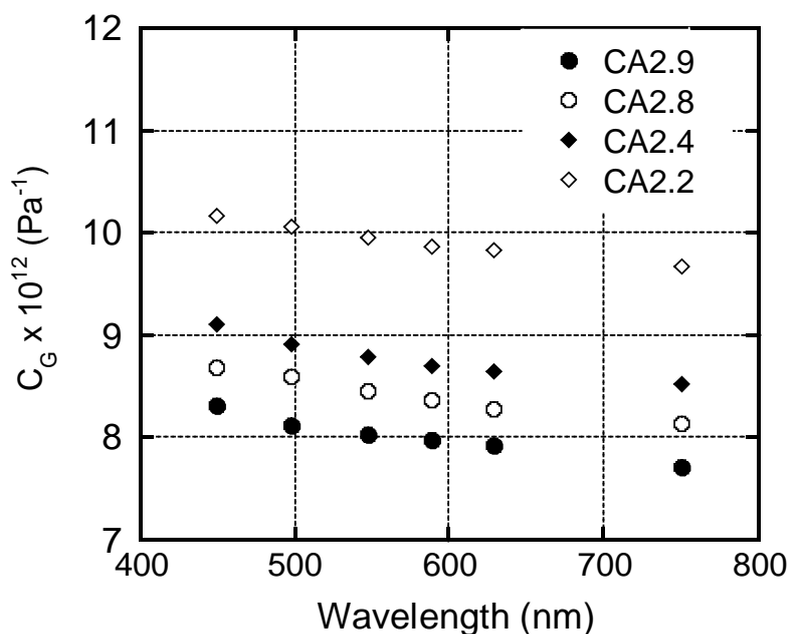


Figure 10 Azimuthal intensity distributions at $\angle\sigma = 1.6$, i.e., the (010) plane of (a) CA2.9, (b) CA2.8, (c) CA2.4, and (d) CA2.2; (black) after pre-heating prior to stretching, (red) immediately after stretching without relaxation process, and (blue) after the relaxation process (60 sec after cessation of stretching).

The stress-optical coefficient in the glassy state C_G was shown in Figure 11, which was evaluated by the relationship between the applied load and the retardation at room temperature. It was confirmed that the birefringence was proportional to the applied stress; i.e., the stress-optical law in the glassy region was applicable to all CA

1 films. The stress-optical coefficient, i.e., the slope of the birefringence – stress relation,
 2 decreased with increasing the acetyl substitution. Furthermore, all films exhibited
 3 weak and ordinary wavelength dispersion. The absolute value of CA2.9 was almost
 4 similar to those of poly(methyl methacrylate) and polystyrene, and much lower than
 5 that of polycarbonate (Liu et al. 1996; Okita et al. 2005; Szczurowski et al. 2010).



6
 7 Figure 11 Wavelength dispersion of the stress-optical coefficient C_G in the glassy
 8 region.

9

10 Conclusion

11 The optical anisotropy was studied using various CA samples with different
 12 degrees of acetyl substitution. As increasing the acetyl substitution, the orientation
 13 birefringence decreased. Furthermore, the orientation birefringence was found to be
 14 determined mainly by the orientation of crystalline region, which was revealed by

1 2D-XRD measurements. As a result, the birefringence showed slow relaxation after the
2 cessation of hot-stretching. This anomalous behavior will be a great benefit for CA
3 because the retardation is not sensitive to the cooling condition at hot-stretching process.
4 The stress-optical coefficient in the glassy region was found to decrease with the acetyl
5 substitution. The value of CA with a large amount of acetyl substitution is similar to
6 those of poly(methyl methacrylate) and polystyrene.

7

8 **Acknowledgement**

9 A part of this work was supported by JSPS Grant-in-Aid for Scientific
10 Research (B) Grant Number 16H04201. The authors would like to express their sincere
11 gratitude to Daicel Corporation for their valuable suggestions and the kind supply of the
12 samples employed in this study.

13

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