Viscoelastic Properties of Cholesteric Liquid Crystals from Hydroxypropyl Cellulose Derivatives

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In this study, we investigated the rheological behavior of a cholesteric liquid crystalline polymer prepared by esterification of hydroxyl propyl cellulose (HPC) with propionyl chloride. The HPC ester tethering propionyl side chains (HPC-Pr) showed thermotropic cholesteric liquid crystal (CLC) phase with visible Bragg reflection in the temperature range between 100 °C and 135 °C. Although visible Bragg reflection of HPC-Pr disappeared even after cooling from the temperature above isotropic phase transition temperature, we observed birefringence under crossed Nicols. Therefore, at this state, the molecular structure of HPC-Pr might be partially isotropic. The partially isotropic HPC-Pr had a relatively high viscosity as confirmed by rheological measurements. On the other hand, the viscosity decreased by shearing.

Keywords: Cellulose derivative, Cholesteric liquid crystals, Bragg reflection, Rheology, Viscoelasticity

1. Introduction

Cellulose is the most abundant polymer on the earth, and its derivatives often show liquid crystalline phase [1]. Especially, hydroxypropyl cellulose (HPC) is one of the most frequently investigated cellulose derivatives. For instance, HPC derivatives possessing acyl groups are known to show cholesteric liquid crystal (CLC) phase [2-4]. At the CLC phase, the chiral molecules are self-organized to form the helicoidal molecular structure with a periodic helical pitch. When unpolarized light propagates into the CLC, right-handed or left-handed circularly polarized light of a specific wavelength is reflected depending on the handedness of helical molecular structure. The specific reflection peak wavelength (λref) by CLC is quantitatively determined by the following formula:

\[ \lambda_{\text{ref}} = n \pi \]

where \( n \) is the average refractive index of CLC and \( \pi \) is the helical pitch length [5]. Such a light reflection phenomenon can be regarded as a sort of Bragg reflection.

Previous report on the HPC derivatives with various acyl groups have shown that their average refractive indices are approximately 1.46 regardless of the alkyl chain length of their substitution groups [6]. By considering the experimental result, the reflection peak wavelength can be tuned by controlling the CLC helical pitch length by external stimuli. In the case of thermotropic CLCs, the helical pitch length is changed by temperature due to the thermodynamic motion of CLC molecules. Consequently, the reflection wavelength of CLC can be tuned by controlling the temperature [7].

The rheological behavior of HPC derivatives possessing acyl groups is interesting. Previous studies reported that sticky products are obtained by acylation of HPC [3,4,8-10]. This phenomenon is quite strange in that sticky polymer melts are produced from a powdery raw material of pristine HPC after the simple acylation. In addition, the fundamentals and applications of HPC derivatives with acyl groups are numerous discussed, focusing on their environmental-friendliness and CLC properties [10-15]. As well as the other CLC materials, it is expected to be applied for versatile
photonic devices such as temperature sensors and reflective color displays. In the perspective of practical realization of them, the knowledge of their basic physical and mechanical properties is of prime importance. However, a limited number of studies on rheological behavior of acylated HPC derivatives have been reported, while there are many papers on that of HPC solutions. It is important to comprehend the rheological behavior of polymer melts.

In this report, we describe the optical and rheological properties of a HPC derivative possessing propionyl side chains (HPC-Pr), as shown in Fig. 1. The transmission spectral measurements of HPC-Pr revealed that the reflection peak wavelength is controlled by temperature, arising from the thermotropic CLC feature. Then, the viscoelasticity of HPC-Pr is measured to investigate the rheological behaviors caused by the helical molecular structure of CLC. The present report would pave the way to understand the intrinsic physical properties of HPC derivatives from both scientific and practical viewpoints.

2. Experimental

2.1. Materials

We used a commercially available substance of HPC (Fujifilm Wako Pure Chemical Co.; Viscosity of 2.0 wt% aqueous solution, 2.0~2.9 mPa·s). The number of hydroxypropyl groups per HPC monomer unit, i.e., molar substitution value (MS), was determined to be 4.0 by using the 1H-NMR spectrum [16]. HPC was dried under vacuum before use. Anhydrous pyridine (Kanto Chemical Co., Inc.; 99.5%), anhydrous acetone (Kanto Chemical Co., Inc.; 99.5%), and propionyl chloride (Tokyo Chemical Industry Co. Ltd.; 98.0%) were used as received.

Figure 1 shows the chemical structure of a HPC derivative possessing propionyl side chains (HPC-Pr). According to our previous report [14], HPC-Pr was synthesized by the esterification of HPC with propionyl chloride.

2.2. Measurement of optical properties

To fabricate a CLC cell, HPC-Pr was sandwiched between a pair of two glass plates with a gap of approximately 200 µm. Transmission spectra of the CLC cell were measured using a compact CCD spectrometer (Ocean Optics, USB2000+) equipped with a tungsten halogen light source for the probing white light. The temperature of CLC cell was precisely controlled using a hot stage system (Mettler Toledo, HS82 and HS1).

2.3. Measurement of rheological properties

The viscosity measurements were performed using a stress-controlled rheometer (Anton Paar, MCR102) equipped with a stainless-steel parallel plate with a diameter of 25 mm. HPC-Pr was sandwiched at the gap of approximately 500 µm. In this experiment, the shear rate (γ̇) was increased from 0.01 s⁻¹ to 100 s⁻¹ and then decreased from 100 s⁻¹ to 0.01 s⁻¹ taking 41 sec in each way. By using a forced convection oven, the temperature of HPC-Pr was maintained at 100 ºC during the measurements. As heated at this temperature of 100 ºC, a reflection peak of HPC-Pr emerged around 400 nm, as mentioned below.

Oscillatory shear rheology measurements were also conducted by using the above-mentioned rheometer. The storage modulus (G') and loss modulus (G'') were measured at angular frequencies (ω) ranging from 100 rad/s to 0.1 rad/s at the temperature between 40 ºC to 100 ºC at the intervals of 20 ºC. The strain amplitude was adjusted in the range of 0.2% to 2.0%, which was sufficiently small to measure the linear viscoelasticity. The master curves were created by shifting the G' and G'' results in logarithmic scales according to the time-temperature superposition principle with reference temperature (T_ref) of 100 ºC. When shifting the results, the amounts of horizontal and vertical shifts of the curves at each temperature are called the horizontal shift factor (α_t) and vertical shift factor (β_t), respectively. The values α_t and β_t are dependent on the temperature.

Before each measurement, HPC-Pr was either preheated or presheared to erase any remaining the historical molecular orientation. Hereafter, the samples are noted as Pr-H or Pr-S, respectively. Pr-H was prepared by heating at 155 ºC for 5 min in
the rheometer, which was higher than the isotropic phase transition temperature of HPC-Pr \((T_i = 150 \degree C)\) to ensure the random orientation of cholesteric texture confirmed by polarized optical microscopic observation. Pr-S was prepared by shearing at the rate of 10 s\(^{-1}\) for 200 sec heated at 100 \degree C, which was sufficiently long enough to reach its steady state of molecular alignment.

3. Results and discussion

3.1. Synthesis of HPC-Pr

We confirmed that all hydroxyl groups in the side chains of HPC are completely substituted with propionyl groups to form HPC-Pr. According to the identification method described in our previous report [14], the \(^1\)H-NMR spectrum of HPC-Pr in CDCl\(_3\) indicated that all hydroxyl groups of HPC are esterified by propionyl chloride. Also, the FT-IR spectrum showed the disappearance of a broad peak around 3500 cm\(^{-1}\), assigned to the O-H stretching vibration of pristine HPC, after the esterification of HPC with propionyl chloride.

3.2. Reflection properties of HPC-Pr

The Bragg reflection peak wavelength of HPC-Pr was dependent on the temperature. Figure 2 shows the changes in the transmission spectrum of a cell of HPC-Pr upon stepwise heating from 100 \degree C to 135 \degree C at the intervals of 5 \degree C. The Bragg reflection peak was continuously red-shifted from 400 nm to 635 nm upon heating from 100 \degree C to 135 \degree C. Such a shift of Bragg reflection peak can be attributed to the increase of the helical pitch length of the thermotropic CLC induced by temperature [7,17]. Furthermore, we confirmed that the reflection peak is blue-shifted from 635 nm to 400 nm upon cooling process from 135 \degree C to 100 \degree C. Therefore, the reflection wavelength shift by temperature was found to be fully reversible in the range between 100 \degree C and 135 \degree C.

3.3. Pretreatment of HPC-Pr before rheological measurements [18]

Figure 2 also shows the broadening of reflection band as well as the decrease of reflection intensity by elevating the temperature from 100 \degree C to 135 \degree C, suggesting that the helical axis of CLC gradually becomes the disordered by heating close to the isotropic phase temperature. This is also supported by the fact that the Bragg reflection is not observed at 100 \degree C for at least 3 h after heating at 155 \degree C for 5 min once. This result emphasizes that the preheating process deteriorates the orientational order of helical axis of CLC. On the other hand, as HPC-Pr was heated at 155 \degree C, followed by 100 \degree C, the polarized optical microscopic observation between crossed Nicols confirmed that HPC-Pr shows slight birefringence. However, at this stage, no Bragg reflection color was observed. Taking the overall facts in account, the molecular structure of HPC-Pr after preheating process can be interpreted as follows. Although the CLC molecules form the helicoidal structure at the microscopic scale such as one helical pitch or less, the helical axis is aligned in random directions. As a result, Bragg reflection was not observed due to the randomly-aligned helical axis of CLC structure while the birefringence was observed due to the formation of CLC structure at microscopic scale. This motivated us to investigate the shearing effect of HPC-Pr.

The preshearing time was determined to be 200

![Fig. 2](image2.png)

**Fig. 2.** Transmission spectral changes in the CLC cell of HPC-Pr upon stepwise heating from 100 \degree C to 135 \degree C at the intervals of 5 \degree C. The reflection peak was red-shifted, accompanied by the spectral broadening, upon the heating process.

![Fig. 3](image3.png)

**Fig. 3.** Time dependence of the viscosity \((\eta)\) \((\dot{\gamma} = 10\ s^{-1}, 100\ \degree C)\) of a preheated CLC of HPC-Pr, which reached the steady state by shearing at 200 sec.
sec, which is sufficient to align CLC layers. Figure 3 shows the time dependence of viscosity of HPC-Pr measured at 100 ºC by shearing at a constant rate of 10 s⁻¹. The viscosity gradually decreased with taking the measurement time up to 150 sec. However, the viscosity became constant at 19 Pa·s despite the prolonged measurement time of 150 sec or more, suggesting that the sample reached its steady state. This experimental result also implies that the alignment of CLC layers is completed by shearing at 10 s⁻¹ for 150 sec.

3.4. Rheological properties of HPC-Pr

Although both Pr-H and Pr-S exhibited shear thinning behavior, their flow curves were different when \( \dot{\gamma} \) is increased, as shown in Fig. 4. Shear thinning is a decrease of viscosity with increasing the shear rate, and often occurred by structure deformation of the sample [19]. This indicates that the CLC structure of HPC-Pr is changed by shearing. Thereby, we focus on the difference of viscosity between Pr-H and Pr-S. It is well-known that the viscosity is crucially influenced by the molecular arrangements.

On the other hand, when \( \dot{\gamma} \) was decreased, the viscosity of Pr-H and Pr-S were almost identical regardless of \( \dot{\gamma} \). This is because the CLC alignment is completed in the process of increasing \( \dot{\gamma} \) for both Pr-H and Pr-S.

The flow curve of Pr-H also exhibited shear thickening behavior in the \( \dot{\gamma} \) range from 0.01 s⁻¹ to 0.1 s⁻¹. In general, the shear thickening behavior appears with structure formation by shear. This fact implies the alignment of CLC layers is induced when \( \dot{\gamma} \) is below 0.01 s⁻¹, which is reasonable for considering the randomly-arranged CLC layers. Pr-H seems to be more viscous than Pr-S.

Figures 5 (a) and 5 (b) shows the master curves of \( G' \) and \( G'' \) of Pr-H and Pr-S heated at \( T_{\text{ref}} \) of 100 ºC. The master curve of Pr-H given in Fig. 5 (a) can be divided into two regions by the difference of relationship between \( G' \) and \( G'' \). In the region of \( \omega \) below 4.0 \times 10⁻¹ rad/s, \( G' \) exceeds \( G'' \) and reaches a plateau, which implies that Pr-H is more solid-like in this region. This rigidity of Pr-H can be caused by the randomly oriented CLC layers as described above. At higher region of \( \omega \) over 4.0 \times 10⁻¹ rad/s, \( G'' \) became larger than \( G' \) and the slopes of \( G' \) and \( G'' \) in the range above \( \omega = 1.0 \times 10² \) rad/s were

![Fig. 4. Shear-rate dependences at 100 ºC of the apparent viscosity of Pr-H and Pr-S samples. The value \( \eta \) of Pr-H was higher than that of Pr-S in the increasing \( \dot{\gamma} \) process, whereas they showed the same profiles in the decreasing \( \dot{\gamma} \) process.](image)

For instance, the viscosity of polymers whose chains entangle themselves tends to be higher than that of unentangled one [20]. The viscosity of Pr-H at 0.2 s⁻¹ was 3.0 \times 10² Pa·s, which was up to 7.5 times higher than that of the Pr-S (40 Pa·s). Such high viscosity of Pr-H can be attributed to the randomly oriented CLC layers. When the Pr-H is sheared, the additional force is requisite to generate well alignment of CLC layers from random state [18].

![Fig. 5. Master curves of the storage modulus (\( G' \)) and loss modulus (\( G'' \)) of HPC-Pr heated at \( T_{\text{ref}} \) of 100 ºC (a) The preheated sample (Pr-H). (b) The presheared sample (Pr-S).](image)
estimated to be approximately 1.0 and 0.8, respectively. These values are different from those of the ideal Newtonian fluid, which would be 2.0 and 1.0, respectively. Such smaller \( \omega \) dependences of \( G' \) and \( G'' \) can be attributed to the disturbed flow behavior caused by the molecular orientation of CLCs.

The master curve of Pr-S presented in Fig. 5 (b) is reasonably characterized by two differences when compared with that of Pr-H as follows. First, \( G'' \) exceeded \( G' \) in the entire \( \omega \) range, and no plateau of \( G' \) was observed. This indicates that Pr-S is more liquid-like in lower \( \omega \) regions. This liquid-like behavior might be happened by the aligned CLC structure. Second, both \( G' \) and \( G'' \) of Pr-H were higher than those of Pr-S in the entire \( \omega \) range. On the other hand, the slopes of both \( G' \) and \( G'' \) in the range above \( \omega = 1.0 \times 10^2 \) rad/s were approximately 1.0 and 0.8, respectively, which was the same as those of Pr-H.

Considering the overall results, it turned out that Pr-H is more viscous and rigid rather than Pr-S. This rigidity of Pr-H might stem from the randomly oriented CLC layers, as mentioned above.

4. Conclusion

In this study, the rheological behavior of a CLC polymer derived from HPC was investigated. As the CLC polymer was preheated above the isotropic phase transition temperature, the partially isotropic structure remained even after cooling. This was confirmed by the fact that the color derived from Bragg reflection did not appear but birefringence appears in this state. In this way, the preheated CLC polymers behave more viscously and rigidly than the presheared ones, which might be attributed to their randomly oriented CLC layers. The experimental evidence is expected to be available to apply the general CLC polymers for not only the comprehension of physical properties from a scientific standpoint, but also the fabrication of novel photonic devices with CLC materials from a practical standpoint [21-25].

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