Environmental Stable Chiral-Nematic Liquid-Crystal Elastomers with Mechano-Optical Properties

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Abstract: Chiral-nematic liquid crystal (N* LC) elastomers exhibit mechano-optical responsive behavior. However, practical sensor applications have been limited by the intrinsic sensitivity of N* LC elastomers to environmental conditions, such as temperature. Although densely cross-linked LC network polymers exhibit high thermal stability, they are not proper for the mechanical sensor due to high glass transition temperatures and low flexibility. To overcome these issues, we focused on enhancing thermal stability by introducing noncovalent cross-linking sites via intermolecular interactions between LC molecules bonded to the polymer network. N* LC elastomers with a cyanobiphenyl derivative as a side-chain mesogen exhibited mechano-optical responsive behavior, with a hypsochromic shift of the reflection peak wavelength under an applied tensile strain and quick shape and color recovery owing to high elasticity. Notably, the N* LC elastomers showed high resistance to harsh environments, including high temperatures and various solvents. Interactions, such as π–π stacking and dipole–dipole interactions, between the cyanobiphenyl units can act as weak cross-links, thus improving the thermal stability of the LC phase without affecting the mechano-optical response. Thus, these N* LC elastomers have great potential for the realization of practical mechano-optical sensors.

Keywords: chiral-nematic liquid crystal elastomer; mechanochromism; selective reflection; noncovalent cross-linking; thermal stability

1. Introduction

Stimuli-responsive photonic materials have attracted great interest owing to their tunable optical behavior, which is useful for applications in areas such as flexible electronics and soft robotics [1–3]. In particular, mechano-optical responsive materials change their optical properties under the application of physical stimuli, such as grinding, pressure, and tensile or shear strain [4–9]. Liquid-crystal (LC) elastomers are promising advanced materials in which the orientation of the mesogens is strongly coupled to the polymer network; thus, electro-, mechano-, or thermo-responsive effects on the macroscopic shape and the microscopic molecular orientation can be observed simultaneously [10–18]. Notably, chiral-nematic (N*) LC elastomers have unique optical properties based on the helical structure of the N* LC system [19–21]. N* LC elastomers spontaneously form a helically twisted molecular orientation, with the orientational direction of the mesogens rotating along the helical axis [22–24]. Thus, N* LC elastomers exhibit Bragg reflection and selectively reflect circularly polarized light of a particular wavelength. The reflection wavelength, λ, depends on the helical pitch, P, of the N* LC elastomers, as expressed by the following equation:

$$\lambda = n_{ave} P,$$
where \( n_{\text{ave}} \) is the average of the ordinary and extraordinary refractive indices of the LCs [25–28]. In 2001, Finkelmann et al. reported a dye-doped N* LC elastomer that acted as a mechanically tunable laser [29]. In this system, a mechanical stimulus applied to the elastomer changed the macroscopic shape in accordance with the change in the helical orientation. This pioneering work provided a basis for the development of N* LC elastomers as stimuli-responsive photonic materials [30–34]. The helical orientation of an N* LC elastomer is very sensitive to various external stimuli, such as heat [31], light [32], electric fields [33], and humidity [34]. For the development of mechano-sensing applications, a high sensitivity to multiple stimuli is unfavorable because the external stimuli can interfere with the observation of a pure mechano-optical response. Thus, stability in various environments, except under mechanical stimuli, is highly desirable for the development of N* LC elastomers with highly sensitive mechano-optical response behavior.

In recent decades, N* LC network polymers with high cross-linking densities have attracted practical interest because of their thermally stable optical properties [35–37]. However, dense cross-linking is not appropriate for mechano-optical sensors because of the associated high glass transition temperature and low flexibility [38,39]. Moreover, the cross-linker acts as an impurity that destabilizes the LC phase, leading to a decrease in the isotropic phase transition temperature [40]. Therefore, to utilize an N* LC elastomer for mechano-optical applications, high thermal stability with a low cross-linking density is necessary. Herein, we report a useful design method for thermally stable N* LC networked polymers with low densities of cross-linking sites. Our strategy for enhancing the thermal stability of network polymers involves introducing noncovalent interactions between the mesogenic moieties in the side chains of network polymers. In this study, we employed a cyanobiphenyl derivative as a side-chain mesogen because intermolecular interactions such as \( \pi-\pi \) stacking and dipole–dipole interactions between cyanobiphenyl mesogens are well-known to impart polymeric materials with high resistance to thermal treatment [41–44]. We hypothesized that these interactions between mesogens could act as weak cross-links and thus improve the thermal stability of the LC phase without disturbing the mechanical and optical behavior. The synthesized N* LC elastomers with noncovalent cross-linking sites showed good thermal stability and maintained their mechano-optical response behavior under harsh environmental conditions.

2. Materials and Methods

2.1. Materials

The chemical structures of the materials used to prepare the N* LC elastomers are shown in Scheme 1. The base LC monomers (HAB and CN) were obtained from a commercial supplier and used after recrystallization from methanol. The plasticizer (5CB), cross-linker (HDDA), and photoinitiator (Irgacure 651) were purchased from commercial suppliers and used as received. A chiral monomer (ISBCD) was synthesized and purified as reported previously (Scheme S1, Text S1, and Figures S1–S3) [45,46].

2.2. Preparation of N* LC (x) Elastomers

N* LC (x) elastomers, where \( x \) denotes the molar ratio of ISBCD in the monomer mixture, were prepared by photopolymerization. A schematic of the preparation method is shown in Figure 1, and details are given in Text S2–S4. Monomer mixtures were prepared (Table 1) [47] and then injected into a homemade glass cell at the isotropic temperature (80 °C). In the cell, 110-µm-thick PDMS films were placed on the glass substrate to aid in removing the prepared elastomers. The monomer mixture in the cell was then cooled to the LC temperature (25 °C) and photopolymerized by irradiation with UV light at 365 nm (5.0 mW cm\(^{-2}\)) for 5 min in the N* LC phase. The resultant N* LC (x) elastomer film was peeled off the PDMS layer. The thickness of the obtained N* LC elastomer film was 55 µm. All the resultant films showed bright selective reflection.
2.2. Preparation of N* LC (x) Elastomers

Monomer mixtures for synthesizing N* LC (x) elastomers. The LC monomers (HAB and CN), chiral monomer (ISBCD), cross-linker (HDDA), plasticizer (5CB), and photoinitiator (Irgacure 651) used to synthesize N* LC (x) elastomers.

| x   | HAB (mol%) | CN (mol%) | 5CB (mol%) | Irgacure 651 (mol%) | HDDA (mol%) | ISBCD (mol%) |
|-----|------------|-----------|------------|---------------------|-------------|--------------|
| 2.0 | 2.0        |           |            |                     |             |              |
| 2.2 |            | 2.2       |            |                     |             |              |
| 2.4 |            | 2.4       |            |                     |             |              |
| 2.6 | 40         | 40        | 20         | 1                   | 7           | 2.6          |
| 2.8 |            |           | 2.8        |                     |             |              |
| 3.0 |            |           | 3.0        |                     |             |              |

Scheme 1. Chemical structures of the LC monomers (HAB and CN), chiral monomer (ISBCD), cross-linker (HDDA), plasticizer (5CB), and photoinitiator (Irgacure 651) used to synthesize N* LC (x) elastomers.

2.3. Characterization of N* LC (x) Elastomers

The thermodynamic properties of the N* LC (x) elastomers were determined by differential scanning calorimetry (DSC, X-DSC7000, SEIKO Instruments, 10 °C min⁻¹). The phase sequences and phase transition behavior were determined by polarized optical microscopy (POM, Olympus, EX51TH) using a hot stage (Instec, HCS302 hot stage and mK1000 controller). To evaluate the mechano-optical responsive behavior, the reflectance...
spectra of the elastomers were recorded using a UV-visible diode-array spectrometer (StellarNet BLUE-Wave spectroscope) under static tensile strain. Stress–strain curves were measured using a tensile tester (Instron 5943, Instron) at room temperature with a strain rate of 2 mm min\(^{-1}\). The Young’s modulus of each elastomer film was determined in the tensile deformation range of 0.25–1.0%. Films with dimensions of 20 mm × 5 mm were used for the tensile tests.

3. Results and Discussion

3.1. Phase Transition Behavior

The phase transition behavior of the N\(^*\) LC (\(x\)) elastomer films was investigated using POM (Figure S4) and DSC (Figures S5–S7). In the monomer mixture for N\(^*\) LC (2.0), the isotropic-to-LC phase transition was observed at 28 °C during the cooling process. From the planar texture shown in the POM images (Figure S4), we concluded that the LC phase observed in the monomer mixture was an N\(^*\) LC phase. After polymerization, the N\(^*\) LC phase observed in the monomeric state was fixed in the resultant elastomers. As summarized in Table 2, the LC temperature ranges of the elastomers with different ISBCD concentrations were similar, with an N\(^*\) LC phase observed from ~10 to 90 °C. These results indicate that the LC behavior of the elastomers was not affected by the chiral dopant concentration. ISBCD has been reported to have a very large helical twisting power (HTP = 43 µm\(^{-1}\) mol\(^{-1}\)) [46]; therefore, only a small amount of ISBCD was necessary to induce a helical LC alignment. Because only a low concentration of chiral dopant was used, the thermodynamic properties of the elastomers were not affected.

| x     | Phase Transition Temperature (°C) |
|-------|----------------------------------|
|       | Monomer Mixture \(^1\) | Elastomer \(^1\) |
| 2.0   | N\(^*\) 28 I | G 9 N\(^*\) 93 I |
| 2.2   | N\(^*\) 28 I | G 12 N\(^*\) 90 I |
| 2.4   | N\(^*\) 27 I | G 11 N\(^*\) 90 I |
| 2.6   | N\(^*\) 27 I | G 8 N\(^*\) 91 I |
| 2.8   | N\(^*\) 27 I | G 12 N\(^*\) 95 I |
| 3.0   | N\(^*\) 26 I | G 10 N\(^*\) 92 I |

\(^1\) Abbreviations: G, glassy; N\(^*\), chiral nematic; I, isotropic. The phase transition temperatures were determined by POM for the monomer mixtures and DSC for the elastomers.

The N\(^*\) LC phase of the elastomers was enantiotropic, indicating that the LC phase was reversibly formed during both the heating and cooling processes (Figure 2a). When the N\(^*\) LC (\(x\)) elastomers were cooled from the isotropic phase, the N\(^*\) LC phase exhibited the same optical texture immediately below the isotropic–N\(^*\) LC phase transition temperature. Generally, polymer LCs tend to show large supercooling owing to their viscosity. However, in the current system, most of the mesogens were covalently bonded to the polymer main chains through a methylene spacer, and the mesogenic orientation was fixed by cross-linkers. We consider these structural features to be responsible for the high reversibility of the LC behavior and the quick recovery of the LC phase from the isotropic phase [48,49].

To confirm this hypothesis, we prepared a linear LC polymer (l-LCP) without the cross-linking agent using the same polymerization procedure in the LC cell with polyimide alignment layers. As shown in Figure 2b, l-LCP formed a well-aligned homogeneous N\(^*\) LC phase and exhibited an enantiotropic N\(^*\) LC phase between 0 and 85 °C. As compared to the elastomers, l-PLC had a lower LC-to-isotropic phase transition temperature. In contrast to the N\(^*\) LC (\(x\)) elastomers, which showed complete reversibility of the LC texture, l-LCP became opaque, and a different LC texture was obtained in the N\(^*\) LC phase by cooling from the isotropic phase, as shown in Figure 2b. These observations clearly indicate that l-PLC formed N\(^*\) LC polydomains after cooling from the isotropic phase and that the order
parameter of the LC phase was lower than that just after polymerization. Therefore, we concluded that the mesogenic orientation in the N* LC (x) elastomers is stabilized and memorized by cross-linking.

![Images of phase transition behaviors](image_url)

**Figure 2.** Phase transition behaviors during heating and cooling processes: (a) N* LC (2.0) elastomer and (b) l-LCP.

Next, to investigate the effects of the concentration of the CN monomer on the thermodynamic behavior, we synthesized N* LC (2.0) elastomers with different LC monomer ratios (HAB/CN). Here, the concentration of ISBCD was fixed at 2.0 mol%. The molar ratios of the LC monomers and the phase transition temperatures are summarized in Table 3. All the elastomers with different HAB/CN ratios exhibited the same glass transition temperature (10 °C). Although the LC-to-isotropic phase transition temperature was not affected by the ISBCD concentration, as discussed above, it showed a dependence on the HAB/CN ratio, gradually increasing with an increase in the CN monomer content. CN has been reported to form intermolecular interactions [43,44], and we consider intermolecular interactions between CN units to serve as noncovalent cross-links. As mentioned previously, cross-linking can stabilize the orientation of mesogens in the LC phase. Thus, increasing the CN content in the elastomer enhanced the thermal stability of the N* LC phase. This result supports our working hypothesis that noncovalent cross-linking can be used to improve the thermal stability of the LC phase in the elastomer without increasing the concentration of covalent cross-linkers.
Table 3. Phase transition temperatures of N* LC (2.0) elastomers with different HAB/CN ratios.

| Molar Ratio of HAB/CN | Phase Transition Temperature (°C) |
|-----------------------|----------------------------------|
| 70/10                 | G 10 N* 80 I                     |
| 60/20                 | G 10 N* 83 I                     |
| 50/30                 | G 10 N* 86 I                     |

1 Contents of other materials in the monomer mixture: 5CB, 20 mol%; HDDA, 7 mol%; ISBCD, 2 mol%. 2 Abbreviations: G, glassy; N*, chiral nematic; I, isotropic.

3.2. Thermal Stability of Optical Properties

As a representative example, we investigated the thermal stability of the optical properties of the N* LC (2.0) elastomers. Generally, the helically twisted molecular orientation of an N* LC elastomer is sensitive to temperature [31,50]. The twist of the N* LC helix decreases in the pre-phase transition region upon approaching the phase transition temperature. Therefore, the pitch length increases, and the selective reflection band shifts following thermal treatment. Previous studies have demonstrated that such N* LC elastomers can be utilized as thermochromic materials, as the optical properties show a temperature dependence [50,51]. However, such temperature responsivity degrades the resolution and accuracy of the mechano-optical response, thus limiting the utility of such N* LC elastomers for sensing applications. Although densely cross-linked LC polymers typically exhibit excellent thermal stability because both their macroscopic and microscopic structures are fixed, they have high glass transition temperatures and low flexibility. The resultant low flexibility and stretchability make these materials incompatible with mechano-optical sensing applications. In contrast, our proposed N* LC (x) elastomers have high flexibility and stretchability owing to their low density of covalent cross-linking but high thermal stability derived from the noncovalent cross-linking sites of the CN units.

To confirm the effect of the noncovalent cross-linking sites on the thermo-optical stability of the N* LC (2.0) elastomers, we investigated the thermo-optical properties of N* LC (2.0) elastomers with different HAB/CN ratios by measuring the reflectance spectra at various temperatures. The N* LC (2.0) elastomer with HAB/CN = 40/40 did not exhibit any color change or shift of the reflection peak wavelength with the change in temperature until the isotropic phase transition temperature (Figure 3a,b,e). However, the reflectance intensity decreased spontaneously at the phase transition point (Figure 3d). These results indicate that this N* LC (2.0) elastomer has a high thermo-optical stability. In contrast, an N* LC (2.0) elastomer with a lower concentration of CN units (HAB/CN = 70/10) exhibited a hypsochromic shift in the reflection peak wavelength with increasing temperature (Figure 3c–e). These results clearly suggest that the noncovalent cross-linking sites between CN units in the side chains contribute not only to the thermal stability of the N* LC phase but also to the thermo-optical stability of the N* LC elastomer.
These results indicate that this N* LC (2.0) elastomer has a high thermo-optical stability. In contrast, an N* LC (2.0) elastomer with a lower concentration of CN units (HAB/CN = 70/10) exhibited a hypsochromic shift in the reflection peak wavelength with increasing temperature (Figure 3c–e). These results clearly suggest that the noncovalent cross-linking sites between CN units in the side chains contribute not only to the thermal stability of the N* LC phase but also to the thermo-optical stability of the N* LC elastomer.

3.3. Mechano-Optical Response Behavior and Stability of N* LC (x) Elastomers

Toward the realization of mechano-optical sensors, we investigated the mechano-optical response behavior of the N* LC (x) elastomers under tensile strain. Each of the N* LC (x) elastomers showed selective reflection at a particular peak wavelength (x = 2.0, 835 nm; 2.2, 735 nm; 2.4, 691 nm; 2.6, 635 nm; 2.8, 584 nm; and 3.0, 504 nm; Figure 4a). As the concentration of ISBCD units increased, the peak wavelength hypsochromically shifted in accordance with Equation (1). As a representative example, we elongated the N* LC (2.0) elastomer (Figure 4b,c). The film appeared faint red at 0% strain and changed to an intense orange color at 80% strain. The color change occurred under elongation, and the initial color was recovered after the removal of the applied strain, and this process could be repeated. The color change was evaluated quantitatively using reflectance measurements. The reflection peak wavelength of the N* LC (2.0) elastomer was hypsochromically shifted from the near-IR region (835 nm) in the initial state to the orange region (616 nm) at 80% strain under tensile elongation. The other N* LC (x) elastomers also exhibited similar mechano-optical response behavior, with the reflection peak wavelengths hypsochromically shifted under elongation (Figure S8).
Mechano-optical stability of the N* LC (2.0) elastomer film during compressive/tensile strain cycles from 0% to 50%. As shown in Figure S9, the experimental \( \varepsilon \) values show good agreement with the theoretical \( \varepsilon_{\text{theory}} \) values, which indicates that the N* LC (x) elastomers undergo isotropic deformation. Thus, an applied tensile strain can be evaluated quantitatively by merely measuring the shift of the reflection peak wavelength. Therefore, our proposed N* LC (x) elastomers have great potential as mechano-optical sensors with excellent flexibility and stretchability, unlike densely cross-linked LC polymers.

Subsequently, we conducted a tensile test using the N* LC (2.0) elastomer. The elastomer exhibited a Young’s modulus of 1.02 MPa and a fracture strain of 107%, indicating high stretchability. To further understand the mechano-optical stability, we evaluated the shift of the reflection peak wavelength during compressive/tensile strain cycles. As shown in Figure 4d, the N* LC (2.0) elastomer clearly exhibited the same color change with a slight hysteresis, even after repeated strain and recovery processes. Thus, we conclude that the elastomer maintains high mechano-optical stability during compression and elongation.

The shift in the reflection peak wavelength can be explained by the isotropic deformation of typical rubbery materials. Such materials deform isotropically under incompressible conditions, leading to in-plane uniaxial elongation that causes the compression in both the in-plane perpendicular and thickness directions in the same manner. This phenomenon can be expressed by the following equation:

\[
\varepsilon_{\text{theory}} = 1 - (1 + \varepsilon_x)^{-1/2},
\]

where \( \varepsilon_{\text{theory}} \) is the theoretical strain in the thickness direction under the applied tensile strain \( \varepsilon_x \). In our N* LC elastomers, we experimentally evaluated the thickness strain \( \varepsilon_z \) by transforming Equation (1) into \( \varepsilon_z = \Delta P/P = \Delta \lambda/\lambda_0 \), where \( \Delta P \) is the change in the helical pitch and \( \Delta \lambda \) is the shift of the reflection peak wavelength under the applied tensile strain [52]. As shown in Figure S9, the experimental \( \varepsilon_z \) values show good agreement with the theoretical \( \varepsilon_{\text{theory}} \) values, which indicates that the N* LC (x) elastomers undergo isotropic deformation.

\[\begin{align*}
\text{(a)} & \quad \text{Reflectance spectra of N* LC (x) elastomers with different ISBCD concentrations.} \\
\text{(b)} & \quad \text{Reflectance spectra of N* LC (2.0) elastomer as a function of applied tensile strain (0–80%).} \\
\text{(c)} & \quad \text{Photographs of N* LC (2.0) elastomer under applied tensile strains of 0–80%.} \\
\text{(d)} & \quad \text{Mechano-optical stability of N* LC (2.0) elastomer film during compressive/tensile strain cycles from 0% to 50%}
\end{align*}\]
3.4. Optical Stability in Solvents

For practical mechano-optical sensors, it is desirable that the optical properties show little dependence on the solvent conditions. When a typical N* LC elastomer network is immersed in a solvent, swelling causes network deformation, which drastically affects the optical properties arising from the helically twisted molecular orientation. Although swellable LC polymers have utility as humidity-responsive materials [53,54], this swelling effect interferes with and degrades the resolution of mechano-optical sensing. Therefore, resistance to solvents, especially to common polar solvents such as water and ethanol, is crucial for the broad application of N* LC elastomers as mechano-optical sensors. As shown in Figure S10, the N* LC (2.0) elastomer exhibited hydrophobicity, as water placed on the elastomer surface formed a droplet and was not absorbed into the elastomer. We also evaluated the environmental stability of the N* LC (2.0) elastomer in polar solvents by immersing the elastomer in water or ethanol (Figure 5a). After immersion for 1 week, the N* LC (2.0) elastomer showed no color change in either water or ethanol. To further investigate the solvent resistance, the N* LC (2.0) elastomer was elongated in water and ethanol environments. As shown in Figure 5b, the N* LC (2.0) elastomer exhibited excellent mechano-optical behavior, even in water and ethanol, without any optical degradation. This excellent environmental stability under harsh conditions can be attributed to the wide elastic temperature range and hydrophobicity of the elastomer. These results suggest that our proposed N* LC (x) elastomers provide a pathway for realizing mechano-optical sensors or devices that will function in a wide range of extreme environments.

Figure 5. (a) N* LC (2.0) elastomer immersed in water and ethanol over 1 week, and (b) color change under an applied tensile strain in water and ethanol baths.
4. Conclusions

In conclusion, we proposed a design concept for synthesizing thermally, optically, and environmentally stable N* LC elastomers by employing cyanobiphenyl units in the elastomer as noncovalent cross-linking sites. Because of their low densities of covalent cross-linking sites, the N* LC (x) elastomers exhibited high stretchability and flexibility, which are favorable properties for realizing practical mechano-optical sensors. In addition, the thermal stability of the N* LC phase could be enhanced by increasing the concentration of cyanobiphenyl units that form noncovalent cross-links in the elastomer. Notably, this noncovalent cross-linking had almost no effect on the mechano-optical response behavior, such as the reflection peak wavelength and mechano-optical stability, but significantly enhanced the environmental stability. The N* LC (x) elastomers exhibited excellent mechano-responsive color change properties, with a hypsochromic shift of the reflection peak wavelength under an applied tensile strain and reversible recovery to the original shape and color owing to high elasticity. In addition, the hydrophobicity and large elastic temperature ranges of the N* LC (x) elastomers provided excellent resistance to water and ethanol. Based on the excellent physical properties of the N* LC (x) elastomers, including high stretchability, reversibility, and stability under harsh conditions, our material design concept of using noncovalent cross-linking sites in N* LC elastomers is expected to contribute to various mechano-optical sensor applications in the fields of wearable displays, electric skin, and soft robotics.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/app1115037/s1, Scheme S1: Synthetic route for ISBCD, Text S1: Synthesis of chiral dopant monomer, Text S2: Preparation of PDMS film, Text S3: Cell preparation with rubbing treatment, Text S4: Preparation of N* LC(x) elastomers, Figure S1: 1H-NMR spectrum of ISBCD in CDCl₃, Figure S2: 13C-NMR spectrum of ISBCD in CDCl₃, Figure S3: HMQC spectrum of ISBCD in CDCl₃, Figure S4: POM image of N* LC (2.0) mixtures observed at room temperature before (a) and after (b) polymerization in a rubbing cell, Figure S5: DSC thermogram of I-LCP (2nd scan, scanning rate of 10 °C min⁻¹), Figure S6: DSC thermogram of N* LC (2.0) elastomer (2nd scan, scanning rate of 10 °C min⁻¹), Figure S7: DSC thermogram of N* LC elastomers containing different HAB/CN ratios: (a) 70/10, (b) 60/20, and (c) 50/30 (2nd scan, scanning rate of 10 °C min⁻¹), Figure S8: (a) Color change of N* LC (x) elastomers with different chiral dopant concentrations x (2.0–3.0 mol%) under an applied tensile strain of 0% (left), 40% (middle), and 80% (right), and (b) reflection peak wavelengths of N* LC (x) elastomers as a function of applied tensile strain (0–80%), Figure 59: (a) Schematic illustrations of film deformation by elongation along the x-axis (the thickness direction and helical pitch along the z-axis, and the in-plane direction along the y-axis perpendicular to the elongation direction are compressed as the elongation strain increases) and (b) change in helical pitch εz (red circles and line) experimentally evaluated using the shift in the reflection peak wavelength and the theoretical thickness strain of rubbery elastomers (black line) under the applied tensile strain εz calculated using a Poisson ratio of 0.5, Figure S10: Photographs of an N* LC (2.0) elastomer film: (a) water droplet on the film surface, showing hydrophobicity and (b) contact angle of the droplet on the surface.

Author Contributions: Conceptualization, O.T.; Investigation, K.K., T.S., S.K., K.H., N.A., A.S.; Formal Analysis, K.K., K.H., O.T.; Writing—Original Draft Preparation, K.K., K.H., O.T.; Writing—Review and Editing, K.K., K.H., O.T. All authors have read and agreed to the published version of the manuscript.

Funding: This research was supported by JSPS KAKENHI (18H03764 for OT; 19K21131 and 20K15249 for KH), JST A-STEP (JPMJTM19C9), the Ritsumeikan Global-Innovation Research Organization (R-GIRO), and the Cooperative Research Program of the Network Joint Research Centre for Materials and Devices (Tokyo Institute of Technology).

Data Availability Statement: The data that supports the findings of this study are available in the Supplementary Materials of this article.

Acknowledgments: We thank Osaka Organic Chemical Industry for providing the liquid crystal monomers.
Conflicts of Interest: The authors declare no conflict of interest.

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