Interference color modulation, tunable refractive index, and chiroptical electrochromism in a pi-conjugated polymer with cholesteric liquid crystal order

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Interference color modulation, tunable refractive index, and chiroptical electrochromism in a π-conjugated polymer with cholesteric liquid crystal order

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A π-conjugated polymer film prepared by electrolytic polymerization using a cholesteric liquid crystal (CLC) is demonstrated to have a tunable interference function under electrochemical doping and dedoping. The polymer exhibits a CLC-like periodic structure with a potential-dependent refractive index and optical absorption properties. The interference color of the polymer film can thus be modulated dynamically by appropriate application of a voltage in the ±1 V range. This research involves structural chirality and tunable chiroptical properties, doping-dedoping driven tunable refractive index, electrochromism, and interference color modulation for the present polymer. The phenomena demonstrate the possibility of electrochemical photonics. © 2009 American Institute of Physics. [DOI: 10.1063/1.3126524]

I. INTRODUCTION

Cholesteric liquid crystal (CLC) is a twisted nematic phase of liquid crystal in which the crystal directors are rotated progressively to form a helical structure. The helical orientation of rodlike molecules results in periodicity, and the three-dimensional molecular arrangement produces phase chirality. When CLC molecules are homeotropically anchored to a substrate, a periodic stripe pattern is formed corresponding to the helical half-pitch of the CLC, resulting in a visible spiral domain or fingerprint pattern. This periodic structure can be regarded as a one-dimensional photonic crystal that can function as a diffraction grating by selectively reflecting light according to the half-pitch periodicity of the helical structure. This light-selective property of CLC can be modified by appropriate adjustment of the half-pitch length through external stimuli, such as electrical potential. The helicity-related optical properties of CLCs have been applied in a range of functional optical devices, including lasing devices, optical diodes, and optical reflectors.

CLC has also been employed as a reaction field for the synthesis of optically active polymer films from optically inactive monomers. Such a CLC reaction field has successfully been employed for the preparation of polymer-stabilized CLC materials. Polymer growth by this electrochemical method follows the three-dimensional chiral continuum of the CLC electrolyte solution to form a chiral architecture on the molecular level, effectively imprinting the chiral structure of the CLC field on the product polymer. The optically active polymer thus prepared exhibits intense optical activity, controllable circular dichroism, and optical rotation upon doping. Here, the doping level can be tuned electrochemically by appropriate application of a voltage in an electrolyte solution containing a dopant (supporting salt, perchlorate ion). Counterions are injected into the conductive polymer on the anode side, resulting in p-type doping. However, while the π-conjugated polymer was successfully imprinted with the molecular architecture of the CLC, the surface of the polymer film thus prepared is planar and does not exhibit iridescence.

In the present study, an optically active π-conjugated polymer film with a microperiodic convexo-concave surface is prepared by a modified electrolytic polymerization method in CLC electrolyte solution from an achiral monomer with high polymerization activity. The resultant polymer film acts as a diffraction grating, displaying optical absorption and refractive index modulation that can be reversibly controlled by electrochemical doping-dedoping through the application of a low voltage. This polymer therefore has the functionality of interference color tuning.

II. EXPERIMENTAL

Polymerization was performed by an electrochemical method using a CLC electrolyte solution containing a cholesteric derivative (cholesteryl pelargionate) as a CLC inducer and the 3,4-ethylenedioxythiophene (EDOT)-based starting monomer, 1,4-bis[2-(3,4-ethylenedioxy)thienyl]phenylene (BEDOT-P). Here, the addition of a small amount of a chiral compound having a large helical twisting power (β) to nematic liquid crystal produces CLC with a helical superstructure. Although “chiral dopant” is the generally accepted term, “chiral inducer” is employed in this report to avoid confusion with the term “dopant” (electron accepter or donor) for conductive polymers.

The monomer was prepared in advance by Stille coupling reaction (Grignard metathesis was employed in a previous study). The resulting polymer (poly-BEDOT-P) was prepared on indium tin oxide (ITO) glass substrates using a sandwich cell configuration.

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from that of the cholesteric electrolyte solution. The molecular structure of the polymer, however, differs completely from the macromolecular arrangement of the main chains in the pitch between stripes in the texture corresponds to the helical half-periodic pattern to be transcribed from the CLC structure. The fingerprint pattern is similar to that of typical CLC materials. Scanning electron microscopy (SEM) observations reveal that the polymer prepared in the CLC exhibits a microperiodic convexo-concave surface with a fingerprintlike pattern [Fig. 1(b)]. The infrared (IR) absorption spectrum of the polymer confirms the absence of the CLC inducer (cholesteryl pelargonate, νC=C, 1739 cm\(^{-1}\)) in the final polymer.

The good polymerization activity and excellent crystallinity of the polymer prepared in this study allows a single-pitch periodic pattern to be transcribed from the CLC structure. The fingerprint pattern is similar to that of typical CLC materials, although the polymer has no fluidity. The distance between stripes in the texture corresponds to the helical half-pitch (P/2) of the CLC electrolyte solution and originates from the macromolecular arrangement of the main chains in a helix matching that of the electrolyte solution. The molecular structure of the polymer, however, differs completely from that of the cholesteric electrolyte solution. The molecular imprinting of the product polymer thus appears to occur by topological control of the one-handed helically twisted matrix of the CLC electrolyte solution in the course of the epitaxial electrolytic polymerization process.

The convexo-concave structure of the polymer produces two optical effects: the iridescent reflection of light and the diffraction of light. The polymer film has a rainbowlike appearance reminiscent of a jewel beetle upon irradiation at certain angles and appears monochromatic at other incident angles upon irradiation with white light from a light emitting diode [Figs. 2(a) and 2(b)]. The natural color of the polymer is dark brown in the half-doped state [Fig. 2(c)]. This reflection of light originates from the periodic convexo-concave structure of the polymer surface.

Laser transmission through the polymer film produces a circular Fourier-transformed interference figure with distinct separation of red, green, and blue laser light [Fig. 2(d)] due to the random grating formed by the single-pitch helicity-derived stripes of the polymer. The red (670 nm), green (532 nm), and blue (473 nm) wavelengths are projected to the outer, middle, and inner parts of the circular diffraction pattern, respectively. Upon irradiation with light at 532 nm from a laser diode, the polymer displays \( m=1 \) diffraction, as shown in Figs. 2(e) and 2(g), confirming that the polymer has both transmission and reflection grating functions. The diffraction can be expressed by the grating equation \( mλ = 2d \sin θ \), where \( d=P/2 \). Under irradiation normal to the polymer surface, the light trace draws a conic transmission and reflection pattern [Fig. 2(f)]. The three-dimensional surface of the symmetry cone can be described by a quadric cone, \( x^2/a^2 + y^2/b^2 - z^2/c^2 = 0 \), where \( a, b, \) and \( c \) are constants, and \( a=b \neq 0 \).

Figures 3(a) and 3(b) show the transmission colors in the...
a periodicity process increasing positive voltage in the electrochemical doping
fractive index of the polymer near 600 nm decreases with wavelength.

change in the course of electrochemical doping-dedoping observations indicate that $P_{nm}$ and strengthens the intensities at long wavelengths, as extinction coefficient change in the diffraction wavelength. The variation in the diffraction wavelength; that is, the refractive index drives the crease in
applied voltage reduces absorption and the observed for optical absorption, namely, an increase in the
smetry measurements can be substituted for detection angle
change demonstrates the selectivity of light transmission semicircular trace. The angle-dependent profile of the color change shows in Fig. 4(b). This can be attributed to the same physical property (i.e., $\alpha=4\pi\kappa/\lambda$; where $\alpha$ is the absorption coefficient and $\kappa \neq \alpha$).

Electro-optic mechanisms, such as the Kerr and Pockels effects, by which the refractive index changes with application of a voltage, are known to occur in many materials. The Frantz–Keldysh effect similarly affords a change in the optical absorption and refraction for semiconductors and insulators under an electric field. The modulation of optical properties by these processes, however, typically requires applying very high voltages. In contrast, the optical absorption and refractive index of the polymer prepared in the present study can be controlled by electrochemical doping-dedoping using an applied voltage of less than 1 V. This can be referred to as “doping-dedoping driven change in refractive index.”

The change in visible-near infrared (vis-NIR) optical absorption upon electrochemical doping-dedoping reveals an absorption maximum at 463 nm (2.7 eV), corresponding to shown in Fig. 4(b). This can be attributed to the same physical property (i.e., $\alpha=4\pi\kappa/\lambda$; where $\alpha$ is the absorption coefficient and $\kappa \neq \alpha$).

Ellipsometry measurements reveal that the average refractive index of the polymer near 600 nm decreases with increasing positive voltage in the electrochemical doping process [Fig. 4(a)]. Here, the $n$ values obtained from ellipsometry measurements can be substituted for $n_{av}$. The decrease in $n$ by doping results in a shortening (blueshift) of the diffraction wavelength; that is, the refractive index drives the change in the diffraction wavelength. The variation in the extinction coefficient ($\kappa$) exhibits a similar tendency to that observed for optical absorption, namely, an increase in the applied voltage reduces absorption and the $\kappa$ intensity at 450 nm and strengthens the intensities at long wavelengths, as
the \( \pi-\pi^* \) transition of the main chain (Fig. 5). Figure 5 (inset) displays absorption of the polymer as a function of energy (\( \lambda \) is transformed into energy in Fig. 5). The intensity of this peak decreases with increasing voltage, accompanied by the emergence of a new absorption maximum at 639 nm (1.9 eV) and an absorption band in the red to NIR region (<1 eV) attributable to the generation of polarons and bipolarons on the \( \pi \)-conjugated main chains, respectively.\(^{22-25}\)

The natural color appearance in the CIE chromaticity diagram as calculated from the optical absorption spectra of the polymer at \(-0.6\) and \(0.9\) V indicates that the color of the polymer at each applied voltage shifts toward blue with progressive electrochemical doping. This result corresponds to that of the transmission CIE result.

The transmission intensity at a certain wavelength (\( \lambda_i \)) can be expressed by

\[ I_{\text{tot}} = I_0 - I_{\text{abs}} - I_{\text{ref}}, \]

where \( I_{\text{tot}} \) is the total transmission intensity, \( I_0 \) is the input intensity, \( I_{\text{abs}} \) is the absorption intensity, and \( I_{\text{ref}} \) is the reflection intensity. The

\[ I_0 - I_{\text{abs}} \]

component can be obtained using \( I_{\text{abs}} = I_0 - I_{\text{tot}} - I_{\text{ref}}, \)

where \( \alpha \) is the absorption coefficient according to the Lambert–Beer law. Hence, \( I \) is primarily dependent on the optical absorption of the polymer. The generation of polarons as an electroconduction carrier accompanying doping strengthens optical absorption in the red wavelength region, resulting in overall enhancement of transmission in the blue region.

The modulation of light using materials with tunable optical absorption properties can be explained in terms of the change in the complex index of refraction (\( N \)), which is given by

\[ N = n - ik. \]

Doping causes a reduction in the refractive index \( n \) of the polymer by the electro-optic effect and an increase in \(-ik\) at long wavelengths (corresponding to gain loss) by absorption change. Doping therefore drives the total refractive index.

In the doping process for the polymer, the value of \( n \) in the green-orange region is reduced and the optical absorption intensity in the red region is increased, which enhances the transmission of the blue wavelength region light as a complementary color. Furthermore, the reduction in the absorption intensity in the blue region allows transmission of blue light from the polymer. In other words, the generation of the polarons on the polymer by the doping changes the electronic structure of the polymer. This change affords change in refractive index and absorption at a certain detection angle. The change in the color is reversible with the sequential electrochemical doping-dedoping process. This can be referred to as an interference color modulation electrochromism based on tunable transmission effect for the electroactive polymer with diffraction property.

The change in surface color of the present polymer during redox cycling is displayed in Fig. 6 (−0.6 and +0.9 V). Under obliquely incident white light, the natural color of the polymer changes from dark orange to mazarine according to the oxidation state, while the structural color changes from light blue to dark interference colors.

Optical rotatory dispersion (ORD) for the polymer at three applied voltage in 0.1M tetrabutyl ammonium perchlorate (TBAP)/acetonitrile solution is shown in Fig. 7. Upon doping (increase in the applied voltage), an optical rotation at short wavelengths and a broad optical rotation signal at long wavelengths with an isosbestic point of 490 nm. This result indicates that the polymer is electrochiroptically active and that the optical rotation can be tuned via the electrochemical doping process. This property of the polymer can be regarded as an “electrochemically driven chiroptical effect.” The polymer produces a maximum optical rotation of \(-2.2 \times 10^4\) deg/cm at 445 nm. These values are comparable to the Faraday rotation of magnetic transition-metal materials such as Fe \((+3.8 \times 10^3\) deg/cm),\(^{25}\) and NdFeO\(_3\) \((+4.7 \times 10^4\) deg/cm)\(^{26,27}\) under POM observations. This result indicates that structural chirality produced by cholesterics effectively provides chiral architecture for the polymer. The optical rotations of the polymer during voltage scanning from −0.1 to +0.9 V undergo reversible changes at 440 nm (Fig. 7, inset). This change in optical rotation demonstrates reversible chiroptical electrochromism.
IV. CONCLUSION

The electroactive polymer, poly(1,4-bis[2-(3,4-ethylenedioxy)thienyl]phenylene), prepared by an electrochemical method using a CLC electrolyte solution was demonstrated to have a molecular helical structure transcribed from the CLC. As a result of its helical structure, the polymer film exhibits characteristic diffraction that can be tuned by electrochemical doping and dedoping. The diffraction color is determined by the refractive index and optical absorption, both of which are changed by doping/ dedoping due to the generation/loss of polarons on the polymer main chain and the resultant change in the electronic structure. The optical properties of the polymer are thus dynamically tunable by appropriate application of a voltage in the ±1 V range. This electrochromic effect, a type of interference color modulation determined by potential-dependent diffraction, is demonstrative of the possibility of a new field of “electrochemical photonics.”

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28See EPAPS Document No. E-JAP/IAU-105-153909 for change in δ and ϕ value at the applied voltages obtained with ellipsometry measurement. For more information on EPAPS, see http://www.aip.org/pubservs/epaps.html.

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