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Pitch and Handedness of the Cholesteric Order in Films of a Chiral Alternating Fluorene Copolymer

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ABSTRACT: The molecular organization in thermally annealed films of poly(9,9-bis((S)-3,7-dimethyloctyl)-2,7-fluorene-alt-benzothiadiazole) is investigated using polarized light. Measurement of linear polarization in transmission and reflection as a function of layer thickness and orientation directly show a left handed cholesteric organization with a pitch length of 600 nm. Results are corroborated by measurements of circularly polarized reflection and generalized ellipsometry and are compared to calculations of the optical properties based on the Maugin–Oseen–DeVries model. For wavelengths near the lowest allowed optical transition, light with the same handedness as the cholesteric arrangement (left) is found to be reflected and transmitted with a probability higher than right circularly polarized light. The high transmission for left polarized light is interpreted as an optical manifestation of the Bormann effect.

INTRODUCTION

π-conjugated molecular materials find application in optoelectronic devices, such as light emitting diodes and photovoltaic cells. The probability for optical transitions in these materials, which is an important factor controlling the efficiency in light emission and photovoltaic diodes, can in principle be influenced by the mesoscopic structure of the material. A periodic variation in the polarizability of the material with a typical length comparable to the wavelength of the optical transition of interest, changes the density of photon states and will affect the probability of spectroscopic transitions in a limited frequency range. In the extreme case, a photonic bandgap is established, i.e., in a certain frequency range practically all photon states are eliminated. The exclusion of photon states prohibits the absorption and/or emission of photons by the molecule in the material in the forbidden frequency range. A well-established route to create such a forbidden energy gap for photons in a particular direction is the introduction of chiral nematic order. For films of polymer with thicknesses that are relevant for practical application in light emitting diodes or photovoltaic diodes, few methods exist to determine the pitch length of the cholesteric arrangement. The well-known Grandjean–Cano or wedge method does not work because of the high viscosity of the material. Second, because the films are relatively thin, no clear selective reflection band can be observed. Also optical rotation has been used to determine pitch length, because according to the De Vries theory, optical rotation should diverge for wavelengths approaching the selective reflection band. Since often the reflection band cannot be discerned, the viability of the De Vries description for optical rotation seems questionable. In this contribution we develop a novel method to determine pitch length in thin films based on linear dichroism in transmission and reflection, relying on the strong optical anisotropy of the individual polymer chains. This work extends the seminal contribution of Chen and co-workers, who showed that polarized transmission measurements on a thin film of chiral fluorine nonamers provide information on handedness and pitch of the cholesteric molecular arrangement. Making use of this linear dichroism method, we determine the handedness and pitch of the cholesteric arrangement in a layer of chiral conjugated polymer. Complementary measurements on the circular polarization in reflection and optical rotation confirm the handedness. For aligned, thermally annealed films of 1 we find a pitch length of around 600 nm which is close to the wavelength of the lowest allowed absorption band in this material with an onset around 500 nm. For π-conjugated polymers, the contrast in...
dielectric contrast in the direction parallel and perpendicular to the main chain is large for wavelengths near an optical resonance. Because the wavelength of the lowest allowed optical transition in the polymer 1 and its pitch length are comparable, many of the intriguing optical properties of cholesteric liquid crystals which are usually only observable in very thick films, now become apparent already in thin films.

### EXPERIMENTAL SECTION

**Synthesis.** Synthesis of 1 was carried out according to a previously reported procedure. However, the crude product was purified according to reported procedures to remove palladium(0) catalyst from the polymer.

Crude polymer was dissolved in chloroform (150 mL) and aqueous ammonia solution (28–30%, 150 mL) was added and refluxed for 3 h followed by stirring at room temperature overnight. The organic layer was separated using a separating funnel and to this was added ethylendiaminetetraacetic acid disodium salt dihydrate (2.1 g) and the solution was stirred overnight. The organic layer was extracted with water, washed with brine, dried over sodium sulfate, and evaporated under reduced pressure to about 10 mL. To this concentrated organic layer, a palladium scavenger, diethyldithiocarbamic acid diethylammonium salt (60 mg) was added and stirred under dark and argon atmosphere for 6 h. Then the solution was poured onto cold methanol (300 mL) and the precipitate was collected by suction filtration and dried under vacuum at 60 °C overnight. The precipitate was further subjected to Soxhlet extraction with acetone to remove small oligomers. The insoluble precipitate was dissolved in minimum amount of THF and precipitated into cold methanol. The precipitate was filtered and dried under vacuum at 60 °C overnight to obtain a yellow powder.

**Gel Permeation Chromatography.** PS standard, THF; Apparent $M_n$: 5.54 kg mol$^{-1}$, $M_w$: 12.24 kg mol$^{-1}$, PDI ($M_w/M_n$) = 2.20.

**Nuclear Magnetic Resonance.** $^1$H- and $^{13}$C spectra were in accordance with the previous reports.

**Film Preparation.** A solution of 1 in 9:1 (v/v) chloroform-chlorobenzene was heated in an oil bath at 70 °C for 2 h and then within an hour was spin-coated (2000 rpm, 60 s, maximum acceleration) on clean glass slides and rubbed polyimide coated glass slides. The glass slides were annealed at 150 °C for 15 min under nitrogen atmosphere (glovebox). To obtain films of different thickness, concentration of 1 was varied from 5 mg/mL to 50 mg/mL. Film thickness at each concentration (5, 7.5, 15, 30, and 50 mg/mL) of 1 was determined for films on clean glass slides using Dektak 150 Surface Profiler and a similar thickness was assumed on rubbed polyimide coated glass slides as well.

**Polyimide Alignment Layer.** 2.5 cm × 2.5 cm glass slides were cleaned by sonication for 10 min each with acetone and isopropyl alcohol. The glass slides were then etched in a UV-ozone photoreactor (PR-100) for 30 min. Polyimide (AP 1051) was spin-coated on etched glass-slides (5000 rpm, 40 s, 500 rpm acceleration) and then annealed at 180 °C for 90 min. The polyimide facing side was rubbed on a velvet cloth to induce planar alignment.

**Polarized Reflection and Transmission Measurement.** Polarized reflection and transmission measurement were performed using a Woollam WVASE ellipsometer using the general Mueller matrix measurement protocol. Reflection measurement under normal incidence with selection of the reflected intensity, were done used a home-built setup described earlier.

### RESULTS AND DISCUSSION

In Figure 1 we show polarized transmission spectra for a thermally annealed film of 1 with a thickness of $d = 24$ nm on a rubbed polyimide substrate. Transmission spectra for two different orientations of the polymer film are shown: one with the rubbing direction of the underlying alignment layer in the vertical direction and the other with rubbing direction rotated clockwise over +45° with the vertical as seen from the light source directly facing the polymer layer. The minimum in transmission near 460 nm corresponds to electronic excitation of the polymer chain via the allowed $S_1 \leftrightarrow S_0$ electronic transition. This transition is of π-π* orbital nature and polarized in a direction along the polymer backbone. As expected for the vertically oriented polymer film, the transmission of light resonant with the electronic transition, shows strong linear dichroism with vertically polarized (V) incident light being...
transmitted with far lower probability that horizontally polarized (H) incident light. The dichroism is consistent with alignment of the polymer chains in the direction of rubbing in the polyimide orientation layer. Also the reflection of the vertically oriented film is strongly polarized in the vertical direction. If we rotate the film over +45°, the dichroism for V and H polarized light in reflection and transmission is strongly reduced. Yet we notice that for the +45° orientation, H polarized light is transmitted with lower probability than V. In reflection, H is reflected more strongly than V for light at resonance with the S₁ ← S₀ transition. This shows that in the film oriented +45°, the orientation of the chains is biased toward the H direction. This bias can be accounted for by assuming a left-handed cholesteric arrangement of polymer chains.

In Figure 2, we summarize linearly polarized transmission and reflection measurements as shown in Figure 1 for films of I with various thickness. First we calculate the maximum extinction (\( E = -10 \log T \)) for the \( S₁ ← S₀ \) transition. This shows that in the film oriented +45°, the orientation of the chains is biased toward the H direction. This bias can be accounted for by assuming a left-handed cholesteric arrangement of polymer chains.

The degree of polarization for the films with vertical and with +45° orientation are shown in the top part of Figure 2. Clearly for the thinnest films the degree of polarization is the largest. As the film thickness increases a damped oscillatory pattern emerges. In order to account for the measurements we assume simplistically that the extinction in, say, the V direction is proportional to the square of the component of the transition dipole moment integrated over the entire thickness of the film:

\[
P_E^{VH} = \frac{E_V - E_H}{E_V + E_H}
\]  (1)

The degree of polarization for the V and H polarizations according to

\[
E_V(d, \chi) \propto \int_0^{2\pi/d/pitch} \cos^2(\phi + \chi)d\phi
\]

\[
E_H(d, \chi) \propto \int_0^{2\pi/d/pitch} \sin^2(\phi + \chi)d\phi
\]  (2)

with \( d \) the thickness of the film, \( \phi \) the angle between the vertical direction and the local director of the polymer chains film and \( \chi \) the angle describing the orientation of the film with \( \chi = 0° \) referring the rubbing direction parallel to V. In eq 2 the integration ranges from zero to \( \phi_{top} \), i.e., the angle between the director of the cholesteric at the top surface of the film and the direction of rubbing of the underlying substrate:

\[
\phi_{top} = 2\pi \frac{d}{\text{pitch}}
\]  (3)

Good agreement between experimental data and predictions from eq 2 can be obtained assuming a left-handed cholesteric order with a pitch length of 600 nm. To illustrate this we note that for films with 300 nm thickness the director of the cholesteric arrangement, makes exactly half a turn when going from the bottom to the top surface of the film. As a consequence of this half turn, the dichroism in extinction vanishes for all orientations of the film, consistent with the experiment.

For the linear polarization in reflection \( R \) by the films of I, an analysis similar to the one described above applies. We first introduce the degree of polarization in reflection

\[
P_R^{VH} = \frac{R_V - R_H}{R_V + R_H}
\]  (4)

Next, we assume that the linear polarization in reflection is determined by the orientation of the polymer chains at the top surface of the polymer film facing the light source.

\[
R_V(d, \chi) \propto \cos^2\left(\frac{2\pi d}{\text{pitch}} + \chi\right)
\]

\[
R_H(d, \chi) \propto \sin^2\left(\frac{2\pi d}{\text{pitch}} + \chi\right)
\]  (5)

Assuming a pitch length of 600 nm and a left-handed arrangement, the dependence of the degree of polarization in reflection on film thickness can quantitatively be accounted for (see Figure 2 middle).

An objection that may be raised to the above treatment of especially the linearly polarized reflection data, is that due the
left-handed helical arrangement with pitch length close to the wavelength of observation, the reflection should be mainly circularly polarized. In order to investigate the light reflection in more detail, we have resorted to reflection measurements under exactly normal incidence in which not only the polarization of the incoming light but also the polarization of the reflected light can be selected, see Figure 3.

\[
P_{R}^{\text{VHHH}}(\chi) \propto \left( \cos^2 \left( \frac{2 \pi d}{\text{pitch}} + \chi \right) - \sin^2 \left( \frac{2 \pi d}{\text{pitch}} + \chi \right) \right)
\]

(7)

The relationship (eq 7) is illustrated in the left part of Figure 3 for pitch length \( d = 600 \text{ nm} \).

Interestingly, the experimental setup illustrated in Figure 3 can also be used to measure the reflection of left circularly (L) polarized light reflected as L and right circularly polarized light reflected as R. This type of reflection is characteristic for cholesteric liquid crystals. In contrast, reflection of circularly polarized light at dielectric interface occurs with inversion of circular polarization. We define a degree of polarization for this type of cholesteric reflection:

\[
P_{R}^{\text{RLL}} = \frac{R_{RR} - R_{LL}}{R_{RR} + R_{LL}}
\]

(8)

Results for the cholesteric circularly polarized reflection are shown in the left part of Figure 3. We find that cholesteric reflection of left circularly polarized is more efficient that the corresponding reflection of right circular polarized. Because cholesteric liquid crystals selectively reflect light with the same helicity as their cholesteric ordering, these measurements support the conclusion of a left-handed organization in the films of 1. The variation of the degree of circular polarization in the reflection with the orientation of the film which can be observed for the film with 300 nm thickness (Figure 3c) is currently not understood and may indicate that the reflected light may actually be polarized elliptically rather than circularly. In summary, by controlling also the polarization of the light detected in the reflection, we find that both linear and circular polarization measurements confirm a left-handed molecular organization in films of 1.

Having now established a firm basis for the assignment of a left-handed cholesteric arrangement, the question arises naturally whether any circular selective reflection bands arise. It is well-known that in thick films of cholesteric liquid crystals of at least several microns, spectrally quite narrow selective reflection bands occur. In thinner films, however, the width of these reflection bands broaden. In Figure 4 we present spectrally resolved circular selective reflection measurements on aligned films of 1. These selectivities can be extracted out of the so-called Mueller matrix obtained from generalized ellipsometric measurements. In these experiments the angle of incidence is 15° and the polarization of the incident light is controlled and the total (unpolarized) reflected intensity is recorded.

In the left part of Figure 4 we show the degree of circular polarization in reflection obtained as the 1,4 element of the Mueller matrix, defined according to

\[
P_{R}^{\text{RL}} = \frac{R_{R} - R_{L}}{R_{R} + R_{L}}
\]

(9)

As can be seen, even the thinnest films show preferential reflection of left circularly polarized light for wavelengths exceeding the wavelength of maximum absorption for the \( S_1 \leftarrow S_0 \) transition. The degree of polarization in these thin films is however quite low. For the two thickest films, circular selective reflection is more pronounced and the typical undulatory pattern...
of the circular polarization expected for thin cholesteric films is indeed observed.

The handedness and pitch lengths for cholesteric materials have previously also been estimated from study of the optical rotation and circular dichroism.24−26 We now also apply these methods to the films of 1. In Figure 4b, the rotation of plane of linearly polarized light upon transmission through films of 1 under normal incidence is shown. The optical rotation shown is an average over two orientation of the polymer films (0° and 45°). Curiously, the optical rotation spectra show no specific feature in the wavelength region matching the pitch of the cholesteric arrangement.

The optical rotation \( \phi \) of cholesteric liquid crystal is usually interpreted in terms of the De Vries equation. The De Vries equation for the specific rotation reads:

\[
\phi = \frac{d}{P} \frac{k_1^4}{8 q_0 (k_0^2 - q_0^2)}
\]

with

\[
k_0 = \left( \frac{2\pi}{\lambda_{vac}} \right)^2 \left( \frac{\epsilon_\parallel + \epsilon_\perp}{2} \right)
\]

\[
k_1 = \left( \frac{2\pi}{\lambda_{vac}} \right)^2 \left( \frac{\epsilon_\parallel - \epsilon_\perp}{2} \right)
\]
and \( q_0 = 2\pi/\text{pitch} \). Eq 10 is an approximate solution to the wave equation for electromagnetic waves traveling along the helical axis in the Maugin–Oseen–DeVries model for a cholesteric. \(^{27-29}\) Interestingly the DeVries equation predicts a discontinuity at the wavelength of the electromagnetic waves inside the medium that matches the pitch of the chiral nematic. This discontinuity is illustrated in Figure 5b by the red line.

The discontinuity in the optical rotation predicted by eq 10 is not observed in our experiment. In fact this discontinuity also does not occur in the exact solution of wave equation for electromagnetic waves as given by Belyakov \(^{30}\) and Good and Karali. \(^{31}\) In Figure 5 we illustrate these exact solutions computed using an adaptation of the method described by Good and Karali. In the calculations we use complex dielectric functions determined experimentally from ellipsometry on thin annealed and unaligned films of 1. Comparing Figures 5 and 4, we notice that the calculated reflection and optical rotation are in qualitative agreement. The calculations correctly predict strong thickness dependence of the selective reflectivity and the optical rotation. The optical rotation according to the exact solution matches very closely the prediction from the DeVries eq 10, except for the discontinuity at the wavelength matching the pitch of the medium. Calculations using the Good and Karali formalism for films with thickness in the range of several microns, show features that start to resemble the discontinuity in optical rotation predicted by eq 10. We conclude that the DeVries equation does not correctly predict the optical rotation for thin films in the wavelength range corresponding to the pitch length.\(^{32}\)

Finally we focus on the circular differential transmission of light through aligned films of 1. In Figure 4c the degree of circular polarization in the transmission of light is shown, defined according:

\[
\begin{align*}
 p_T^{RL} &= \frac{T_R - T_L}{T_R + T_L} \\
 \phi &\equiv \frac{2(\Delta_L - \Delta_R)}{(\Delta_L + \Delta_R)}
\end{align*}
\]

(fig 13)

Reasons for studying transmission rather than the more commonly used circular differential absorption also known as circular dichroism are the following. Because of the selective reflection and scattering of light by the aligned cholesteric material, determining true absorption and the associated circular differential is complicated and involves combination of several spectroscopic observables. Circular differential transmission is well-defined and can be measured directly. The second reason is a practical one. Commercial circular dichroism spectrometers are designed, optimized and calibrated to detect very small circular dichroism signals. The differentials for the materials under study can, however, be extremely large and partially fall outside the usual measurement range of the commercial spectrometers. The degree of polarization in eq (13) equals the 1,4 element of the normalized Mueller matrix in general ellipsometry and the sign of eq (13) is consistent with the sign of the dissymmetry ratio used in circular dichroism spectroscopy (\( g_{ab} = 2(\Delta_L - \Delta_R)/(\Delta_L + \Delta_R) \) with \( A \) the absorbance).

As can be seen in Figure 4c, the aligned films of 1 all show a negative degree of polarization for wavelengths near the maximum in the \( S_1 \leftarrow S_0 \) transitions. The degree of polarization increases rapidly with film thickness. For the thickest film (300 nm), the value for \( p_T^{RL} = -0.8 \) is close to the theoretical limit for fully selective transmission of left polarized light, \( p = -1 \). The \( p \) value for wavelengths just below the lowest allowed transition is positive, which can be attributed to selective reflection.

Up to now, we have interpreted chiroptical properties of films of 1 in terms of their mesoscopic chiral nematic molecular organization. \(^{33-35}\) In principle, however, also the helicity of individual polymer chains could contribute to the circular differential signals. \(^{56}\) In the latter case, because the chiroptical effects arise from the intramolecular organization, one expects the degree of circular polarization and the specific optical rotation are independent of the thickness of this film. Indeed from molecular chiroptical spectroscopy on dilute solutions, the dissymmetry ratio and specific optical rotation are known to be intrinsic properties of the individual molecules. In Figure 6 we have summarized the maxima of the various degrees of circular polarization as well as the specific optical rotation for films with different thickness. As can be seen, all these chiroptical properties show a strong variation with the thickness of the film. These dependencies, which are in qualitative agreement with the predictions for the Maugin–Oseen–DeVries model seen Figure 5, indicate that contributions to the chiroptical properties under study arising from the intrinsic chiral molecular structure are negligibly small.

For wavelengths near the allowed \( S_1 \leftarrow S_0 \) transition we find that, left circularly polarized light is both reflected and transmitted with a higher probability than right circularly polarized light. This implies that the right circularly polarized light must be absorbed more strongly that left. In contrast, linearly polarized light with polarization parallel to the direction of the main chain of the conjugated polymers is reflected with higher probability than the orthogonal polarization but transmitted with lower probability by thin films, see Figure 1. The peculiar anomalous behavior of the circular polarizations may be interpreted as an optical manifestation of the Borrmann effect. \(^{37,38}\) The latter effect is known from X-ray spectroscopy and describes an anomalously low absorption of X-rays in thick crystals when the crystal is set for a Bragg reflection. In absorbing cholesteric liquid crystals a related optical effect is known, where upon addition of an infrared absorbing dopant, the circular
polarization which is reflected with the highest probability in the absence of dopant acquires the highest transmission coefficient after addition of the absorbing species.\textsuperscript{39,40}

\section*{CONCLUSION}

The optical properties of thin films of the alternating fluorene copolymer 1 with a vitrified chiral nematic order have been investigated covering linear dichroism in transmission and reflection, circular differential reflection and transmission as well as optical rotation. All optical signatures are consistent with a left-handed cholesteric organization with a pitch length around 600 nm. In contrast to common belief, we find that even films with thickness an order of magnitude smaller than the pitch length of the cholesteric arrangement show preferential reflection of one of the circular polarizations of light.

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Notes

The authors declare no competing financial interest.

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