Flexo-electric behavior of bimesogenic liquid crystals in the nematic phase - observation of a new self-assembly pattern at their twist-bend nematic and nematic interface.

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Flexoelectric properties - the effective flexoelectric coefficient and flexoelectric polarisation are investigated for a bimesogenic liquid crystal CBC11CB with a twist bend nematic phase (N_{tb}). Results show that the effective flexoelectric coefficient is at least two times higher than for bimesogens that do not exhibit the N_{tb} phase. The flexoelectric polarization for the N_{tb} phase is also two times greater than in its nematic phase. In a temperature gradient cell, we observe surprisingly a new phenomenon, in the bimesogens CBC7CB and CBC9CB and their mixtures with SCB, of periodic self-assembly in the N phase close to the N - N_{tb} phase transition temperature. This phenomenon is reminiscent of the self-deformation and spontaneous chirality that appears in the system.

I. INTRODUCTION

Two mesogens, connected by an odd number of methylene units are shown to exhibit a second nematic phase (N_2) at temperatures below those observed for the characteristic uniaxial nematic (N) phase. This low temperature phase is now characterized as a twist bend nematic phase (N_{tb}) with ~8 nm helical pitch, determined independently by two groups using freeze-fracture transmission electron microscopy. This is in amazing coincidence with the pitch found from the measurements of the electro-clinical coefficient, on assuming the effective flexoelectric coefficient, ε, to be 5 pC/m. The complicated internal structure of the phase results in a number of fascinating macroscopic self-assembly properties. When the LC in the N_{tb} phase is confined to a planar homogeneously rubbed cell of thicknesses ranging from 2 to 10 μm, the cell shows spontaneously formed self-assembled periodic patterns appearing under a polarising optical microscope as uniform stripes parallel to the rubbing direction with a periodicity two times the sample thickness. The phase also exhibits a linear optical response similar to the electroclinic effect with the switching time of the order of a few microseconds. In the achiral materials this is possible due to the formation of macroscopic chiral domains of opposite handedness separated by the domain walls. When a large electric field of a certain frequency range is applied, these domains turn into another type of periodic striped texture, with handedness of the neighboring domains alternating from one to the other. The walls of these domains are normal to the rubbing direction. A range of interesting properties exhibited, so far, by bimesogens in N_{tb} motivates us to investigate additional macroscopic properties with the final objective as to how the macroscopic structure evolves from the nanoscale one. In this paper, we investigate the bulk flexoelectric coefficient ε = [ε_1 - ε_3] in the conventional nematic phase, and the flexoelectric polarisation, P_f, in both N and N_{tb} phases of the bimesogen CBC11CB in order to gain better understanding of the material properties of such systems. In addition, we also report an unusual pre-transitional phenomenon observed in the bimesogens CBC7CB and CBC9CB, and their mixtures with SCB. The chemical formulae of the liquid crystalline materials studied are given in Fig 1. To the best of our knowledge, such results for this class of materials have not been reported in the literature before. The ultimate question as to what drives this nanoscale helical pitch to the formation of beautiful stripes under cell confinement of thickness of a few μm needs to be explored. Some of the other intriguing issues that remain to be addressed are (i) how do the competing elastic forces of splay, bend and twist play a role in LC cells under dimensions of a few μm confinement? (ii) do the bimesogens display large flexoelectric coefficient that drives the electroclinic coefficient? (iii) what happens at the phase transition from a conventional nematic to the N_{tb} phase?

II. RESULTS AND DISCUSSION

The flexoelectric polarization arising from a coupling between the elastic deformations and the electric polarizability is expressed in terms of the splay and bend flexoelectric coefficients by Meyer as follows:

\[ P_f = ε_1 n (\nabla \cdot n) - ε_3 [n \times (\nabla \times n)] \]

In equation (1), ε_1 and ε_3 are the flexoelectric coefficients that correspond to the splay and bend deformations, respectively. Flexoelectric coefficients in bimesogens, measured using indirect methods, are found to

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be larger than conventional calamitics\textsuperscript{15,16}. The indirect/converse methodology for obtaining the flexoelectric coefficient is based on the gradient stress produced by the external electric field through exploiting the uniform lying helical (ULH) geometry obtained in a cholesteric nematic phase\textsuperscript{17}. The latter is obtained by adding a small concentration of a large helical twisting power dopant to the liquid crystal sample. Flexoelectricity in such systems manifests itself in terms of the rotation of optic axis on the application of an electric field perpendicular to the helical axis\textsuperscript{17}. For small $E$, tilt angle is linearly related to the applied electric field by

$$
tan(\phi) = \frac{e'p}{2\pi K}E$$

(2)

$e'$ and $K$ are the effective flexoelectric and the elastic coefficients defined as: $e' = \frac{\epsilon_1 - \epsilon_2}{2}$, $K = \frac{\epsilon_1 + 2\epsilon_2}{2}$, $p$ is the helicoidal pitch of the ULH. Hence, one can calculate the effective flexoelectric coefficient if the splay, and bend elastic constants are known.

Planar aligned cells are fabricated using indium tin oxide (ITO) glass plates coated with polymer alignment layer RN175 (Japan) and cured at 250°C for ~1.5 hours. The glass plates are assembled together with antiparallel rubbing directions on the top and bottom substrates. The typical thickness of the cell used for the flexoelectric experiment is 5-6 μm. The cholesteric mixture is prepared by adding a small concentration (~3%) of a large helical twisting power chiral agent R5011 (right handed dopant supplied by E. Merck Korea, helical twisting power (HTP) ~100 μm$^{-1}$) to the original sample. The low concentration of the additive ensures that the properties of the host sample are not modified by the chiral dopant. The optical pitch is obtained by measuring the wavelength of selective reflection of light incident normally on a planar cell by a UV/VIS Spectrometer (Avantes-2048). The pitch is found to be nearly invariant for higher temperatures, however it increases significantly from ~420 to 535 nm at temperatures closer to the N-N$\text{ch}$ phase transition temperature as shown in Fig. 3.

The uniform lying helical structure (ULH) is formed only under special conditions\textsuperscript{17} as the minimum free energy of a chiral nematic leads to a Grandjean configuration where the helical axis lies perpendicular to the substrates. In order to induce ULH, the cell is cooled from the isotropic to the cholesteric nematic state at a rate of 0.1°C/min under the influence of a moderate electric field ($E \sim 2$V/μm, $f = 100$ Hz) applied across the cell. Once the ULH evidenced by its characteristic texture is obtained, the tilt angle is calculated using the procedure given before\textsuperscript{18}. The applied fields are lower than required for complete helical unwinding in order to satisfy the conditions used in deriving Eqn(2). Data for the elastic constants and the dielectric anisotropy for C811CB are taken from our results on the same material\textsuperscript{19}. A linear relationship between $tan\phi$ and $E$ (Fig. 2) is obtained for several reduced temperatures. As seen from this figure, flexo-electrically induced rotation angle $\phi$ shows noticeable increase at temperatures closer to the N-N$\text{ch}$ transition temperature. This is possibly due to increase in the helical pitch as temperature decreases (see Fig. 3).

The convention introduced by Coles et al.\textsuperscript{20} about the sign of $(\epsilon_1 - \epsilon_2)$ when applied, sets the sign positive. It is noted that $|e_1 - e_3|$ increases from 21 pC/m at $T-T_{N^\text{ch}} = -1^\circ$C to 35 pC/m for $T-T_{N^\text{ch}} = -15^\circ$C. It would have been interesting to obtain data at temperatures closer to the N-N$\text{ch}$ phase transition. However, the resulting flexoelectric signal degrades at such temperatures significantly; hence reliable flexoelectric coefficient data close to N-N$\text{ch}$ phase transition could not be obtained.

The effective value of the flexoelectric coefficient is higher by a factor 1.5 to 2.0 than the highest values de-
determined for the bent core systems C4 to C7 recently studied by us.\textsuperscript{19} These are also greater by a factor of 2 than ether-linked bimesogens\textsuperscript{15} that do not exhibit the N\textsubscript{th} phase. Values are rather similar to those measured for ester-linked fluorine substituted asymmetric bimesogens\textsuperscript{21}. Note that Atkinson et al.\textsuperscript{15,21} plot |e\textsubscript{1} - e\textsubscript{3}|/2 instead of |e\textsubscript{1} - e\textsubscript{3}| being plotted here in Fig. 3.

Figure 4 gives the temperature dependence of the pyroelectric signal. The flexoelectric polarization $P_f$ is obtained by integrating the pyroelectric signal measured from a 7.7 \( \mu \)m hybrid cell incorporating CBC11CB. The hybrid cell was specially constructed for making pyroelectric measurements. The alignment on one of the substrates was planar and on the second was homeotropic. The peaks that appear at \( T = 126.6^\circ C \) and \( 107^\circ C \) (on cooling the liquid crystal) refer to the Iso-N and N-N\textsubscript{th} phase transitions, respectively. The pyroelectric signal in the isotropic phase is not zero and differs in phase presumably due to the presence of a strong surface polarization in the sample\textsuperscript{32}. This phasor vector was subtracted when calculating $P_f$. The measurements were made in the absence of an electric field, hence the potential effects such as the ionic screening and coupling of dielectric anisotropy to the electric field were neglected. The absolute values of $P_f$ can be calculated by determining the temperature increment i.e. the time rate of change of temperature of the sample as a result of heating from the light source. Alternatively the system can be calibrated by determining the scaling factor required to obtain the absolute value of the pyroelectric coefficient. The absolute calibration of the amplitude of the pyroelectric response, necessary for determining the pyroelectric coefficient, $\gamma$, was carried out by using the spontaneous polarization $P_S$ value of a well known ferroelectric liquid crystal 12OF1M7. This was measured using the standard repolarization current technique\textsuperscript{23}. The pyroelectric signal for 12OF1M7 was recorded using the same measurement conditions as for CBC11CB. The integrated pyroelectric signal of 12OF1M7 was then scaled to $P_f$ value obtained using field reversal method. This yielded the necessary calibration factor. Fig. 4 shows the magnitude of $|P_f|$ determined for CBC11CB. Its value increases on transition from the isotropic phase and is seen to grow marginally from 0.08 to 0.10 nC cm\textsuperscript{-2} in the nematic phase, as the temperature is further reduced in the N phase. It is observed to increase steadily on cooling at the transition into the N\textsubscript{th} phase. It reaches almost 0.17 nC cm\textsuperscript{-2} at the lowest temperatures for which measurements were carried out. Results show for the first time that N\textsubscript{th} phase has a larger flexoelectric polarization by a factor of almost 2, compared to the classical nematic phase. This possibly is the source of a large electroclinic effect observed in this phase\textsuperscript{9,12}. In the literature, flexoelectric polarization has been determined in the nematic phase of a classical calamatic system 5CB\textsuperscript{24}. This was found to be 0.04 nC cm\textsuperscript{-2} at 25 C and its value within the experimental error is being confirmed using our experimental set-up. Furthermore these results on CBC11CB may also indicate that splay deformations may significantly influence the macroscopic effects exhibited by twist-bend nematic phase especially close to the transition temperature. This implies that as the macroscopic structure evolves from the nanostructure, splay deformations can also have a role especially in exhibiting macroscopic effects.

We observe unusual effects in bimesogens CBC7CB and CBC9CB and their mixtures with the monomer (5CB) in a temperature gradient cell. In a narrow temperature range close to the N-N\textsubscript{th} phase transition, a new type of stripe pattern is observed under a polarizing microscope (Fig. 5). This is accompanied by the appearance of rainbow colors and is present only in a narrow
FIG. 5. (color online) POM textures obtained in a 25µm cell filled with the sample (CB-C7-CB + 30% 5CB) when viewed under cross-polarizers on slowly cooling the cell from the nematic phase. (a) the observed striped region between the N and N anarchists phase using a 10X lens magnification observed at a temperature 52.75°C (length of the magenta bar is 24 µm). The stripes are uniform and have a periodicity of 2.94 µm. The red arrow depicts the rubbing direction. (b) the effect of an electric field (10 V, frequency 10 kHz, temperature 52.82°C) applied across the cell in this narrow temperature range. With the field applied, the stripes are not uniform anymore but the region is distinct from the other two phases, with a wider, non-uniform striped pattern. N anarchists is still planar, while nematic is almost homeotropic at this field. In (c) and (d) colors obtained when the rubbing direction rotated left (c) and right (d) from the axis of the crossed polarizers. The circle diameter is 80µm.

The periodicity of the new stripes varies as a function of both concentration and the cell gap. In the pure materials, the stripes are not readily visible but these can be detected as the concentration of 5CB is increased. However, beyond a certain concentration of 5CB, the phenomenon disappears completely. The periodicity of the stripes together with a temperature range of the rainbow colors for the various cell thicknesses and concentrations have been studied for both sets of bimesogens CBC7CB and CB9CB, and their mixtures with 5CB. The temperature range of this transition phenomena is estimated to be less than 0.1°C and appears to be primarily a function of the temperature gradient in the particular experiment, but also as a function of cell gap and to a certain extent depends on concentration of 5CB in the mixture. When a second heater was introduced on top of the sample cell, the temperature range of the observed phenomena decreases significantly and in some cases vanishes altogether, highlighting its association with the temperature gradient across the cell. The periodicity of the striped pattern, however, remains unaffected, suggesting that the rainbow colors are related to the temperature gradient, while the stripes are independent of it. Table 1 shows temperature ranges found in different cells and mixtures under similar temperature gradient conditions in the absence of the second heater. To explain the observed phenomenon, we refer to the geometry shown in Fig 6. The rainbow colors and their behavior as a function of the cell gap can then be explained by the interference on the wedge made with the materials of different birefringence and refractive indices (i.e. N and N anarchists phases). However, the interchange of the colors on rotating the sample between the crossed polarizers (Figure 5(c, d)) implies that not only the birefringence, but also an optical rotation is involved. This indicates presence of self-deformation and spontaneous chirality in the system. The fluctuations seen in the transition region appear to originate from a thin layer of the nematic phase present at the bottom (warmer) glass substrate, while the N anarchists phase is present closer to the top substrate (colder surface) [Fig. 6]. The periodic pattern can be attributed to the interface present between the two phases. The electric field switches the nematic phase by Fredericksz transition whereas the applied field causes the stripes to become non-uniform and wider, seen in Fig. 5(b). It is likely

TABLE I. The periodicity of the striped patterns (bold) and the temperature ranges for the various mixtures and cell gaps.

<table>
<thead>
<tr>
<th>% of 5CB</th>
<th>0%</th>
<th>20%</th>
<th>30%</th>
<th>40%</th>
<th>50%</th>
</tr>
</thead>
<tbody>
<tr>
<td>CB-C7-CB</td>
<td>0°C</td>
<td>0.03°C</td>
<td>0.06°C</td>
<td>0.06°C</td>
<td>0.03°C</td>
</tr>
<tr>
<td>15µm</td>
<td>no</td>
<td>1.93µm</td>
<td>2.74µm</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>CB-C7-CB</td>
<td>0.16°C</td>
<td>0.08°C</td>
<td>0.25°C</td>
<td>0.12°C</td>
<td>0.06°C</td>
</tr>
<tr>
<td>25µm</td>
<td>no</td>
<td>2.2±µm</td>
<td>2.9±µm</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>CB-C9-CB</td>
<td>0.08°C</td>
<td>0.1°C</td>
<td>0.17°C</td>
<td>0.08°C</td>
<td>0.03°C</td>
</tr>
<tr>
<td>15µm</td>
<td>no</td>
<td>2.9±µm</td>
<td>3.0±µm</td>
<td>no</td>
<td>no</td>
</tr>
<tr>
<td>CB-C9-CB</td>
<td>0.1°C</td>
<td>0.2°C</td>
<td>0.13°C</td>
<td>0.08°C</td>
<td>0.04°C</td>
</tr>
<tr>
<td>25µm</td>
<td>no</td>
<td>3.27µm</td>
<td>3.0µm</td>
<td>no</td>
<td>no</td>
</tr>
</tbody>
</table>

FIG. 6. (color online) The temperature gradient in a heating stage causes the interface between N anarchists and nematic phases to form wedge-like confinement for each phase.
that the presence of a thin film of aligned nematic phase improves the visibility of an intrinsic pattern of the N_{th} phase. We however note that the observed phenomenon does not arise from the Grandjean texture, likely to be seen on confining a material with a short helical pitch in a wedge geometry. In our case the stripes appear normal to the rubbing direction and these are independent of the magnitude and direction of the temperature gradient (visualized by the rainbow colors), i.e. these are not caused by the wedge shape (Fig. 6) of the N_{th} (or N) sections of the sample.

III. CONCLUSION

We have obtained the difference in the bulk flexoelectric coefficient, |\epsilon_1 - \epsilon_3| in CBC11CB in the N phase for a bimesogenic system which exhibits the N_{th} phase below the ordinary nematic phase by studying the converse flexoelectric effect in a ULH system. The optical pitch is nearly temperature independent at higher temperatures, but increases steadily at lower temperatures, closer to the N_{th} phase. \tan \phi follows a linear relationship at lower fields in accordance with Eqn(2). The tilt angle at temperatures close to the N_{th} phase was seen to increase substantially in comparison to the higher temperatures, where \phi was observed to be nearly independent of it. The value of |\epsilon_1 - \epsilon_3| was seen to be nearly independent of temperature after an initial increase below the transition from the isotropic phase with values ranging from ~21-35 pC/m. The e/K ratio in this compound is almost greater by a factor of 1.7 and |\epsilon_1 - \epsilon_3| is almost twice higher for CBC11CB than reported for the bimesogen studied in ref.15 and similar to that estimated for ester-linked fluorine substituted asymmetric bimesogens21. These bimesogens show relatively large flexoelectric polarization which undoubtedly gives rise to the large electroclinic effect already reported by the authors12. In some of these bimesogens and their mixtures with 5CB, we have observed micrometer scale stripes at the N-N_{th} transition temperature, indicating yet another surprising chiral self-assembly phenomenon observed in this class of materials. The change in the elastic constants with temperature closer to N-N_{th} transition may be one of the reasons for such observations.

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IV. REFERENCES