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Calorimetric Study of Phase Transitions Involving Twist-Grain-Boundary TGB\(_A\) and TGB\(_C\) Phases

L. Navailles \(^{(1,*)}\), C.W. Garland \(^{(1,**)}\) and H.T. Nguyen \(^{(2)}\)

\(^{(1)}\) School of Science and Center for Material Science and Engineering, Massachusetts Institute of Technology, Cambridge, Massachusetts 02139, USA
\(^{(2)}\) Centre de Recherche Paul Pascal, avenue A. Schweitzer, 33600 Pessac, France

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Abstract. — High-resolution calorimetry has been used to determine the heat capacity and latent heat associated with phase transitions in the homologous series of chiral liquid crystals \(n\)F\(_2\)BTFO\(_1\)M\(_T\) [3-fluoro-4(1-methylheptlyoxy)4'-\((4''\text{-alkoxy-2''},3''\text{-difluorobenzoyloxy})\text{tolane}\). These compounds exhibit smectic-C\(^*\) (SmC\(^*\)), twist-grain-boundary (TGB\(_A\) for \(n = 10\), TGB\(_C\) for \(n = 11, 12\)) and cholesteric (N\(^*\)) phases. All the phase transitions are first order with small to moderate latent heats. There is a large rounded excess heat capacity peak in the N\(^*\) phase that is consistent with the predicted appearance of short-range TGB order (chiral line liquid character). This is analogous to the development of an Abrikosov flux vortex liquid in type-II superconductors. Both the \(n = 11\) and \(12\) homologs exhibit two closely spaced transitions in the region where a single TGB\(_C\) - N\(^*\) transition was expected. This suggests the existence of two thermodynamically distinct TGB\(_C\) phases.

Résumé. — Des expériences de calorimétrie haute résolution ont été réalisées pour déterminer les chaleurs spécifiques et les chaleurs latentes associées aux transitions de phase des homologues de la série cristal liquide \(n\)F\(_2\)BTFO\(_1\)M\(_T\): 3-fluoro-4-[1-méthyl-heptyloxyl]-4'-\((4''\text{-alkoxy-2''},3''\text{-difluorobenzoyloxy})\) tolanes. Ces produits présentent la phase smectique C\(^*\) (SmC\(^*\)), les phases à torsion par joint de grain (TGB\(_A\) pour \(n = 10\) et TGB\(_C\) pour \(n = 11, 12\)) et la phase cholestérique (N\(^*\)). Toutes les transitions de phase sont du premier ordre. La chaleur latente associée à ces transitions est faible ou modérée. Nous observons, dans la phase N\(^*\), un grand pic arrondi qui est en accord avec les prédictions de l'apparition d'un ordre TGB à courte distance (liquide de ligne de dislocation). Ce phénomène est l'analogue du liquide de vortex dans les supraconducteurs de type II. Les composés \(n = 11\) et \(12\) présentent, dans la région où nous attendions une transition TGB\(_C\) - N\(^*\) unique, deux transitions sur un très faible domaine de température. Ce résultat suggère l'existence de deux phases TGB\(_C\) thermodynamiquement distinctes.

\(^{(*)}\) Present address: G.D.P.C., Université Montpellier II, c.c. 26, Place E. Bataillon, 34095 Montpellier Cedex 05, France
\(^{(**)}\) Author for correspondence (e-mail: cgarland@mit.edu)

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1. Introduction

The existence of a twist-grain-boundary (TGB) phase in chiral liquid crystals was predicted theoretically by Renn and Lubensky [1] in 1988 and first demonstrated experimentally in 1989 by Goodby et al. [2]. The TGB phase combines a helical twist and smectic-A (SmA) layering. This structure consists of a regular stack of periodic grain boundaries separating regions of SmA order, with the layer normal rotated by \( \Delta \theta \) across each boundary. The grain boundaries themselves are formed by a grid of parallel screw dislocations. Many of the experimental studies characterizing the structure and properties of a TGB phase have been carried out on compounds of the homologous series nP1M7 (methylheptyl-alkoxyphenylpropioloyl-oxybiphenyl carboxylate), which exhibit the phase sequence SmC\(^*\)-TGB\(_A\)-isotropic (I) [2-5].

Recently, there has been a detailed calorimetric study of the phase transitions in the homologous series nFBTFO\(_1\)M\(_7\) [6] with the general formula

\[
\text{H}-\left(\text{CH}_2\right)_n\text{O}\cdots\text{C} = \text{C} \cdots\text{O} \cdots\text{CH}_3
\]

These compounds, whose chemical name is 3-fluoro-4\([(R)\text{ or } (S)]\text{-methylheptoxy}]4'-\(4''\)-alkoxy-3\''-difluorobenzoyloxy) tolan, were synthesized and first characterized by Bouchta et al. [7]. For \( n \leq 11 \), the nFBTFO\(_1\)M\(_7\) compounds exhibit the phase sequence SmA-TGB\(_A\)-N\(^*\)-BP\(_s\)-I, where N\(^*\) is the cholesteric (twisted nematic) phase and BP\(_s\) denotes several blue phases. There is an isomorphism between these chiral liquid crystals and high-\( T_c \) type-II superconductors in which SmA \( \leftrightarrow \) Meissner phase, TGB\(_A\) \( \leftrightarrow \) Abrikosov flux vortex lattice, N\(^*\) \( \leftrightarrow \) normal metal in a field [1]. Indeed, significant short-range TGB structure in the cholesteric phase, corresponding to a liquid of screw dislocations (called a chiral line liquid and denoted here as N\(_L^*\)), has been predicted by Kamien and Lubensky [8] and is strongly supported by the calorimetric results on nFBTFO\(_1\)M\(_7\) [6]. N\(_L^*\) and N\(^*\) are not thermodynamically distinct phases, but they differ appreciably in the extent of short-range smectic layer formation. The N\(_L^*\) region corresponds to the Abrikosov vortex liquid in high-\( T_c \) superconductors.

A chiral extension of the liquid crystal model for the N-SmA-SmC multicritical region has led to the theoretical prediction of tilted twist-grain-boundary (TGB\(_C\)) phases [9]. Experiments by the Bordeaux group [10-12] on the homologous series nF\(_2\)BTFO\(_1\)M\(_7\),

\[
\text{H}-\left(\text{CH}_2\right)_n\text{O} \cdots\text{C} = \text{C} \cdots\text{O} \cdots\text{CH}_3
\]

the 2\''-3\''-difluorobenzoyloxy analog of nFBTFO\(_1\)M\(_7\), have confirmed the existence of a TGB\(_C\) phase. However, the smectic layers in these compounds are tilted relative to the pitch axis [12], in contrast to the structure proposed in reference [9]. In a recent paper, Dozov [13] has proposed the existence of a new phase, called the twist-melted-grain-boundary phase (TMGB). Predicted to be stable close to the N-SmA-SmC\(^*\) triple point in a chiral liquid crystal, this phase consists of SmC slabs with smectic layers inclined with respect to the pitch axis. It is also argued that the TMGB phase has a lower energy than TGB\(_C\); thus the phase sequence should be SmC\(^*\)-TMGB-TGB\(_C\)-N\(^*\) (or N\(_L^*\)). The major difference between the TMGB and the type of TGB\(_C\)
phase reported in reference [12] would be the character of the screw dislocation patterns in the grain boundaries, and this aspect is not yet resolved experimentally.

In the present paper, we report the results of a detailed calorimetric study of \( n \)FBTFO\(_1\)M\(_7\). Three pure compounds (\( n = 10, 11, \) and 12) and one binary mixture (\( n = 10.75 \), corresponding to a mixture of 10 and 11 containing 75 mole percent \( n = 11 \)) were investigated with high-resolution ac calorimetry and nonadiabatic scanning calorimetry. The paper is organized as follows. Experimental procedures are briefly summarized in Section 2, and the results are presented in Section 3. A discussion and comparison with results on \( n \)FBTFO\(_1\)M\(_7\) are given in Section 4.

2. Experimental Methods

A small quantity (typically 50 mg) of each liquid crystal sample was sealed into a thin silver cell containing a helical coil of gold wire to increase the thermal conductivity and eliminate any internal temperature gradients. Details of the sample cell design are given elsewhere [14]. Data were obtained on each sample with a computer-controlled calorimeter capable of both high-resolution ac operation and a nonadiabatic scanning mode [15]. The latter method is a relaxation mode in which the heater power is linearly ramped, and latent heats at first-order transitions can be measured. In both the ac and relaxation mode, the sample temperature is scanned slowly. Heating or cooling rates of 0.02-0.2 K h\(^{-1}\) have been used for ac runs and 2-3.5 K h\(^{-1}\) for relaxation runs. For the ac mode, an oscillating heat input \( P_{ac} \exp(i\omega t) \) is supplied to the sample by a thin resistive heater. The frequency \( \omega \) of the power input is chosen so that \( \omega \tau_{int} \ll 1 \), where \( \tau_{int} \) is the relaxation time for thermal diffusion in the sample cell. The temperature oscillations induced in the sample are slow enough that temperature gradients in the sample are negligibly small and the amplitude \( T_{ac} \) of the sample temperature oscillation is given by

\[
T_{ac} = \frac{P_{ac}}{(R^{-1} + i\omega C)},
\]

where \( C \) represents the heat capacity of the sample cell and \( R \) is the thermal resistance (reciprocal of the conductance) of the link between the sample and computer-controlled thermal bath. In the absence of two-phase coexistence at a first-order transition or complication due to a complex frequency-dependent heat capacity, one finds [15,16]

\[
C = \left( \frac{|P_{ac}|}{\omega |T_{ac}|} \right) \cos \phi,
\]

\[
\phi = \Phi + \frac{\pi}{2} = \arctan \left( \frac{1}{\omega RC} \right).
\]

where \( \Phi \) is the phase shift of \( T_{ac} \) with respect to \( P_{ac} \). For our calorimeter the standard ac operating frequency is \( \omega = 0.196 \) (a 32 s period for \( T_{ac} \)), for which \( \omega \tau_{int} \ll 1 \) and \( \omega RC \gg 1 \), and \( \phi \) is typically 0.1 to 0.2 rad. The specific heat capacity of the liquid crystal is given by \( C_p = [C(\text{filled cell}) - C(\text{empty cell})]/m \), where \( m \) is the mass of the liquid crystal.

For the nonadiabatic scanning mode, an effective heat capacity \( C_{eff} \) is defined by [15]

\[
C_{eff}(T) = \frac{P - (T - T_b)/R}{dT/dt},
\]

where \( P \) is the dc power input, \( T \) is the time-dependent sample temperature, \( T_b \) is the time-independent bath temperature, and \( R \) is the thermal resistance between sample and bath. On
a heating run, \( P = 0 \) for \( t < 0 \), \( P = \dot{P}t \) for \( 0 \leq t \leq t_1 \), and \( P = P_0 = \dot{P}t_1 \) for \( t > t_1 \), where the power ramp rate \( \dot{P} \) is constant; on a cooling run, the power profile is reversed. In a one-phase region, \( C_{\text{eff}} \) is the heat capacity \( C_p \). In a two-phase region, \( C_{\text{eff}} \) has a large value reflecting latent heat effects associated with conversion of one phase into another. In this case, the latent heat \( L \) is given by

\[
L = \int_{T_a}^{T_b} [C_{\text{eff}}(T) - C_p(\text{coex})]dT,
\]

where \( C_p(\text{coex}) \) is the heat capacity of two coexisting phases over the narrow coexistence range from \( T_a \) to \( T_b \) in the absence of phase conversion [6]. The quantity \( C_p(\text{coex}) \) is obtained by a smooth (typically linear) interpolation between \( C_p \) values in the one-phase regions just above and just below the coexistence region. This technique can be used to detect latent heats as small as 0.2 mJ [15].

3. Results

Calorimetric data obtained for \( n\text{F}_2\text{BTFO}_1\text{M}_7 \) compounds indicate the phase sequences

\[
\begin{align*}
\text{n} = 10 & \quad \text{K} - \text{SmC}^* - \text{TGB}_\text{A} - \text{N}^* - \text{BP}_5 - \text{I} \\
\text{n} = 11, 12 & \quad \text{K} - \text{SmC}^* - \text{TGB}_\text{C} - \text{N}^* - \text{BP}_5 - \text{I}
\end{align*}
\]

where \( \text{K} \) denotes a crystalline phase and \( \text{SmC}^* \) is a chiral tilted smectic phase. At high temperatures, there are typically three blue phases with the sequence \( \text{N}^* - \text{BP}_\text{I} - \text{BP}_\text{II} - \text{BP}_\text{III} - \text{I} \), but this region was not studied in detail. It should be noted that the present results support the predicted evolution of the twisted nematic phase from a simple cholesteric \( \text{N}^* \) at high temperatures to a chiral line liquid \( \text{N}_{\text{L}}^* \) with extensive short-range TGB order at low temperatures. The possible presence of a narrow range of TGB\(_\text{A} \) phase in the \( n = 11 \) homolog, reported in reference [10], will be discussed in Section 4.

An overview of the \( C_p \) variation observed with ac calorimetry is given in Figure 1 for the three pure homologs of \( n\text{F}_2\text{BTFO}_1\text{M}_7 \). The high temperature transitions involving isotropic and blue phases \( (T_1 = \text{I} - \text{BP}_{\text{III}}, T_2 = \text{BP}_{\text{III}} - \text{BP}_{\text{II}}, T_3 = \text{BP}_{\text{II}} - \text{BP}_{\text{I}}, \text{and} T_4 = \text{BP}_{\text{I}} - \text{N}^*) \) will not be discussed here, and attention will be focused on the development of short-range order in the twisted nematic region as well as the transitions involving TGB phases.

3.1. Twisted Nematic Phase. — The excess heat capacity \( \Delta C_p = C_p - C_p(\text{background}) \)

associated with the development of \( \text{N}_{\text{L}}^* \) short-range order in this phase is shown in Figure 2a. These \( \Delta C_p \) curves are obviously nonsingular since they exhibit neither a divergent peak nor a sharp cusp. However, they can be characterized by their integrated area

\[
\delta H = \int \Delta C_p dT.
\]

The possibility of very broad two-phase coexistence due to an extremely sluggish first-order transition can be ruled out by a comparison of \( C_p(\text{ac}) \) data with \( C_{\text{eff}} \) values obtained with the nonadiabatic scanning (ramped relaxation) method, as shown in Figure 2b for \( n = 10 \). If a very broad two-phase coexistence were to occur, there should be a larger excess area observed from nonadiabatic scanning since the latent heat will contribute to \( C_{\text{eff}} \) but not to \( C_p(\text{ac}) \). The same sort of match shown in Figure 2b between the broad excess ac heat capacity peak \( \Delta C_p(\text{ac}) \) and that obtained from the nonadiabatic scanning technique was also observed in the \( 9\text{FBTFO}_1\text{M}_7 \) compound investigated in reference [6]. The positions \( T_b \) of this \( C_p \) maximum in the twisted nematic phase and the values of the integrated enthalpy \( \delta H \) are given in Table I.
Fig. 1. — Heat capacity of three $n$F$_2$BTFO$_1$M$_7$ compounds measured with the ac calorimetric mode. The dashed line represents $C_p$ (background), the behavior expected in the $N^*$ phase in the absence of short-range TGB ordering.
Fig. 2. — (a) Excess heat capacity $\Delta C_p = C_p - C_p$ (background) for $n\text{F}_2\text{BTFO}_1\text{M}_7$ obtained from the ac calorimetric data in Figure 1. $N'_L$ and $N^*$ are not distinct phases but represent a cholesteric with and without short-range twisted line liquid structure. (b) $C_{\text{eff}}(T)$ data for $10\text{F}_2\text{BTFO}_1\text{M}_7$ obtained with the nonadiabatic scanning technique. The smooth curve represents $C_p$(ac) data from Figure 1, and the dashed curve is $C_p$(background).

It should be noted that the $C_p$ behavior in the $T_5$ region was reproducible on both heating and cooling runs obtained over a period of a month. There was, however, a small drift of about $-20$ mK/day in the position of the $\Delta C_p$ maximum. This drift in $T_5$ is comparable to that observed for the transitions $T_6$ to $T_9$ also listed in Table I. The drift rates of these transition temperatures ranged from $-7$ to $-19$ mK/day for $n = 10$ and from $-14$ to $-23$ mK/day.
Table I. — Transition temperatures (K) and enthalpies (mJ/g) in three nF₂BTFO₃M₇ chiral liquid crystals. Higher temperature transitions involving isotropic and blue phases (T₁ – T₄) are omitted. In all cases except N₄* – N₄*, the transitions are first order with latent heat L and little if any pretransitional excess heat capacity. If the transition temperature is followed by the symbol h, hysteresis was observed (see text for details). T₅ denotes the position of a very broad rounded Cₚ maximum, but this does not correspond to a phase transition.

<table>
<thead>
<tr>
<th>Transition</th>
<th>Transition</th>
<th>n = 10</th>
<th>n = 11</th>
<th>n = 12</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(N₄* – N₄*)</td>
<td>T₅</td>
<td>380.3 ± 0.5</td>
<td>378.7 ± 0.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>δH</td>
<td>1400 ± 200</td>
<td>700 ± 40</td>
</tr>
<tr>
<td>T₆</td>
<td>TGBₐ – N₄ₐ</td>
<td>T₆</td>
<td>376.1 ± 0.2 h</td>
<td>~ 0</td>
</tr>
<tr>
<td>L₆</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T₇ₐ</td>
<td>TGBₐ* – N₄ₐ</td>
<td>T₇ₐ</td>
<td>375.17 ± 0.02</td>
<td>~ 125(a)</td>
</tr>
<tr>
<td></td>
<td>L₇ₐ</td>
<td></td>
<td>~ 0</td>
<td>~ 255(a)</td>
</tr>
<tr>
<td>T₇₉</td>
<td>TGB₉* – TGB₉*</td>
<td>T₇₉</td>
<td>375.10 ± 0.02</td>
<td>~ 20(a)</td>
</tr>
<tr>
<td></td>
<td>L₇₉</td>
<td></td>
<td>~ 0</td>
<td>~ 40(a)</td>
</tr>
<tr>
<td></td>
<td>Lₗ₇ₐ + Lₗ₇₉</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T₈</td>
<td>SmC* – TGBₐ</td>
<td>T₈</td>
<td>373.03 ± 0.03 h</td>
<td>70 ± 20</td>
</tr>
<tr>
<td>L₈</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T₉</td>
<td>SmC* – TGB₉</td>
<td>T₉</td>
<td>374.35 ± 0.03 h</td>
<td>52 ± 10</td>
</tr>
<tr>
<td>L₉</td>
<td></td>
<td></td>
<td></td>
<td>126 ± 10</td>
</tr>
</tbody>
</table>

(a) It is difficult to separate the overall sum Lₗ₇ₐ + Lₗ₇₉ into individual contributions due to the transitions T₇ₐ and T₇₉ (see text).

Fig. 3. — Layer spacing d as obtained from an X-ray study of nF₂BTFO₃M₇ [10]. The open symbols were obtained from diffuse diffraction peaks, indicating only short-range order, and the solid symbols were obtained from sharp resolution-limited (6.2 × 10⁻⁴ Å⁻¹) peaks.
for \( n = 11 \) and 12. In every case, the temperatures listed in Table I are the values extrapolated back to \( t = 0 \), based on numerous observations of the transition temperature over times \( t \) ranging up to 30 days.

The concept of short-range TGB order evolving in the low-temperature \( N^*_L \) region of the twisted nematic phase is supported by X-ray scattering results on \( nF_{2}BTFO_{1}M_{7} \) compounds [10]. Figure 3 displays the “smectic” layer spacing \( d \) as a function of temperature for \( n = 10, 11 \) and 12. Diffuse scattering peaks corresponding to short-range TGB order were observed over a range of several degrees above the transition into the TGB phase. Indeed, in the case of \( n = 11 \) and 12 there is clear evidence that this short-range order involves tilted smectic character just above the transition temperature, which is reasonable in view of the fact that both these compounds exhibit TGB\(_{C}\) phases. In contrast, the diffuse scattering in \( 10F_{2}BTFO_{1}M_{7} \) is TGB\(_{A}\) like. It should be noted that due to different samples, different thermometers, and different experimental conditions (length of runs, etc.) for the X-ray runs, the transition temperatures inferred from Figure 3 lie 1-2 K lower than those reported in Table I.

The results in Figures 2 and 3 provide strong support for the Kamien-Lubensky [8] theoretical proposal of a chiral line liquid \( N^*_L \) in the low-temperature region of the twisted nematic phase. The sequence SmA-TGB\(_{A}\)-N\(_L^*\)/N* is directly analogous to the Meissner-vortex lattice-vortex liquid/normal metal phase sequence in type-II superconductors with thermal fluctuations and weak quenched disorder [17,18]. There is no thermodynamic transition between \( N^*_L \), a twisted nematic with entangled liquid-like (melted) screw dislocation lines, and N*, a typical cholesteric without short-range TGB order, since these are predicted to have the same macroscopic symmetry [8]. The same argument applies to the Abrikosov vortex liquid and the normal metal in type-II superconductors, and large effects associated with short-range ordering are predicted in that case [17].

3.2. TGB\(_{A}\)-N\(_{L}^*\) Transition. — The TGB\(_{A}\)-N\(_{L}^*\) transition, denoted as \( T_{6} \) in Table I, is very difficult to detect thermally in either \( nFBTFO_{1}M_{7} \) [6,19] or \( nF_{2}BTFO_{1}M_{7} \). Heat capacity \( C_{p} \) and phase shift \( \phi \) data for \( 10F_{2}BTFO_{1}M_{7} \) obtained with the ac mode using a slow scan rate (50 mK/h) are shown in Figure 4. Both heating and cooling runs were made. In both cases, the phase \( \phi \) shows a small anomalous maximum that is consistent with two-phase coexistence at a first-order transition; see Figure 2 in reference [6] for the comparable feature in \( 10FBTFO_{1}M_{7} \) and references [14-16] for a more general discussion. The position of the maximum in \( \phi \) on heating agrees exactly with that of the tiny anomalous “spike” in \( C_{p} \) (heat) shown in Figure 4, whereas the \( \phi \) (cool) maximum lies at a slightly lower temperature. Thus there is a small hysteresis of 0.07 K. Unfortunately, the latent heat at \( T_{6} \) is too small to be detected with our nonadiabatic scanning technique; thus \( L_{6} \) is estimated to be less than 3 mJ/g. In the very similar system \( nFBTFO_{1}M_{7} \), the latent heat for TGB\(_{A}\)-N\(_L^*\) in the \( n = 9 \) homolog is 8.1 ± 1.2 mJ/g [6] and there are qualitative indications that this latent heat is smaller for \( n = 10 \) and 11 [19].

3.3. SmC*—TGB\(_{A}\) Transition. — The heat capacity variations on heating and cooling through the SmC*—TGB\(_{A}\) transition, denoted as \( T_{8} \) in Table I, are shown in Figure 5. The absolute \( C_{p} \) values for these runs are somewhat lower than those observed in earlier runs, and the reasons for this are unclear. In any event the transition features are obvious. A hysteresis of 0.16 K was observed; and phase shift anomalies (not shown), as well as the \( C_{p} \) data, indicate a two-phase coexistence range of \( \sim 0.30 \) K on both heating and cooling. These ac mode indications of a first-order transition are confirmed by a latent heat \( L_{8} = 70±20 \) mJ/g measured with the nonadiabatic scanning mode. The \( C_{p} \) (SmC*—TGB\(_{A}\)) behavior shown in Figure 5 is qualitatively the same as that observed in \( 11FBTFO_{1}M_{7} \) [19].
3.4. TGB\textsubscript{C}-N\textsubscript{L}\textsuperscript{*} TRANSITIONS. — The \( C_p \) and phase shift \( \phi \) variations obtained by ac calorimetry in the vicinity of the transition from N\textsubscript{L} to TGB\textsubscript{C} are shown for the \( n = 11 \) and 12 homologs in Figures 6 and 7. For both compounds, multiple runs were carried out at low scan rates (15-50 mK/h) on heating and on cooling. Although earlier results [10] indicated a single TGB\textsubscript{C}-N\textsubscript{L} transition, Figures 6 and 7 show two major features denoted as \( T_{7A} \) and \( T_{7B} \). In the case of 11F\textsubscript{2}BTFO\textsubscript{1}M\textsubscript{7}, there is even a very tiny third peak in \( C_p \) located 0.09 K above the \( T_{7A} \) peak. This small feature was reproducible on seven scans made over a period of 30 days. No hysteresis was observed for any of these \( C_p \) peaks although there were slow drifts of \(-23\) mK/day for \( T_{7A} \) and \( T_{7B} \) and \(-14 \) mK/day for the small peak 90 mK above \( T_{7A} \).

The ac calorimetry data in Figures 6 and 7 correspond to what one would expect from two closely spaced first-order transitions. It is known from X-ray studies [10–12] that the phase between the N\textsubscript{L} region and the SmC\textsuperscript{*} domain is a tilted TGB structure. Thus we have designated two TGB\textsubscript{C} phases - TGB\textsubscript{C}\textsuperscript{a} over the \( \sim 0.7 \) K range between \( T_6 \) and \( T_{7B} \) and TGB\textsubscript{C}\textsuperscript{b} over the narrow \( \sim 0.06 \) K range from \( T_{7B} \) to \( T_{7A} \). TGB\textsubscript{C}\textsuperscript{a} should correspond to the TGB\textsubscript{C} structure experimentally established in reference [12], whereas the structure of the TGB\textsubscript{C}\textsuperscript{b} phase is unknown.

In an effort to determine the latent heats \( L_{7A} \) and \( L_{7B} \) associated with this pair of closely spaced transitions, nonadiabatic scanning runs were carried out on both the \( n = 11 \) and 12 homologs and the results are given in Figures 8 and 9. Due to the relatively fast scan rates required for nonadiabatic scanning, it was not possible to resolve two transition features in the N\textsubscript{L} to TGB\textsubscript{C} (\( T_7 \)) region. So we could determine only the sum of latent heats \( L_{7A} + L_{7B} \), which was \( 143 \pm 10 \) mJ/g for 11F\textsubscript{2}BTFO\textsubscript{1}M\textsubscript{7} and \( 297 \pm 10 \) mJ/g for 12F\textsubscript{2}BTFO\textsubscript{1}M\textsubscript{7}. As a very
rough approximation, we have assumed that the relative size of $L_{7A}$ and $L_{7B}$ can be estimated from the relative size of the integrated excess $C_p(ac)$ peaks. For both $n = 11$ and 12, the ratio $C_p(7A) \text{area} / C_p(7B) \text{area}$ is about 6.5 and this yields the estimated $L_{7A}$ and $L_{7B}$ values given in Table I.

3.5. SmC$^*$ – TGB$^\alpha$ Transition. — The ac heat capacity and phase shift $\phi$ associated with the SmC$^*$ – TGB$^\alpha$ ($T_8$) transition in the $n = 11$ and 12 homologs are given in Figures 10 and 11. Both $C_p$ and $\phi$ show only steps rather than peaks like those observed at $T_7$ and $T_8$. However, there are two compelling indications of first-order character: the large $C_{\text{eff}}$ peaks at $T_9$ shown in Figures 8 and 9 for nonadiabatic scanning runs and yielding latent heats $L_9 = 52 \pm 10 \text{ mJ/g}$ for the $n = 11$ compound and $L_9 = 126 \pm 10 \text{ mJ/g}$ for the $n = 12$ homolog; a substantial hysteresis for ac heating and cooling runs (0.85 K for $n = 11$ and 0.50 K for $n = 12$). Furthermore, this thermal behavior is exactly the same as that observed at the analogous SmA – TGB$^\alpha$ transition in $n$FBTFO$_1$M$_7$ compounds, where the latent heat is reported to be $40 \pm 1.6 \text{ mJ/g}$ for 9FBTFO$_1$M$_7$ [6]. The relatively large error bounds for the present $L_9$ and $(L_{7A} + L_{7B})$ values reflects the uncertainty on how to apportion the excess $C_{\text{eff}}$ that lies in the temperature range between $T_7$ and $T_9$ in Figures 8 and 9.

3.6. Mixture. — Nonadiabatic scanning and ac calorimetric runs were also carried out on 10.75FBTFO$_1$M$_7$, a binary mixture containing 75 mole percent of the $n = 11$ compound. A broad $C_p$ maximum observed in the $N^*$ phase was completely consistent with those seen for the $n = 10$ and $n = 11$ compounds. This $T_5$ peak was located at 377.9 K and had an integrated area $\delta H$ of 1200 $\pm$ 200 mJ/g. At lower temperatures, a complicated $C_p$ behavior was observed.
Fig. 6. — Heat capacity and phase shift $\phi$ for 11F$_2$BTFO$_1$M$_7$ in the N*$_L$ to TGB$_C$ transition region. These ac calorimetric data were obtained on heating at a scan rate of 22 mK/h.

Fig. 7. — Heat capacity and phase shift $\phi$ for 12F$_2$BTFO$_1$M$_7$ in the N*$_L$ to TGB$_C$ transition region. These ac calorimetric data were obtained on cooling at a scan rate of $-30$ mK/h.
Fig. 8. — $C_{\text{eff}}(T)$ data for 11F$_2$BTFO$_1$M$_7$ obtained with the nonadiabatic scanning technique at a scan rate of $+3.5$ K/h. The dashed line represents $C_{\text{p(coex)}}$; see the text. The area under the peak labelled $T_7$ is 143 mJ/g and corresponds to the sum of latent heats $L_{7A} + L_{7B}$, and the area of 52 mJ/g under peak $T_9$ represents the latent heat $L_9$.

Fig. 9. — $C_{\text{eff}}(T)$ data for 12F$_2$BTFO$_1$M$_7$ obtained with the nonadiabatic scanning technique at a scan rate of $+3.5$ K/h. The $T_7$ area is 297 mJ/g and corresponds to the sum of latent heats $L_{7A} + L_{7B}$, and the $T_9$ area representing $L_9$ is 126 mJ/g.
Fig. 10. — Heat capacity and phase shift $\phi$ for 11F$_2$BTFO$_1$M$_7$ in the SmC$^*$ to TGB$_C$ transition region. These ac calorimetric data were obtained on heating at a scan rate of 22 mK/h.

Fig. 11. — Heat capacity and phase shift $\phi$ for 12F$_2$BTFO$_1$M$_7$ in the SmC$^*$ to TGB$_C$ transition region. These ac calorimetric data were obtained on heating at a scan rate of 15 mK/h.
over the range 372.4–373.4 K. For a nonadiabatic scanning run on heating, four peaks were observed at \( \sim 373.3 \) K, \( \sim 373.1 \) K, \( \sim 372.8 \) K, and \( \sim 372.6 \) K. The total latent heat associated with these four features was \( \sim 70 \) mJ/g. However, only the 373.1 K and 373.3 K peaks were reproducible on nonadiabatic cooling runs and with ac calorimetry. The interpretation of these low temperature \( n = 10.75 \) features is unclear, and more detailed work with mixtures would be helpful.

4. Discussion

A tentative phase diagram based on data for the \( n = 10, 11, \) and 12 compounds from the present study combined with DSC data obtained in Bordeaux [10] for \( n = 9 \) and binary mixtures of 10 + 11 and 11 + 12 is given in Figure 12. Comparison of \( nF_2BTFO_1M_7 \) transition temperatures obtained with high-resolution calorimetric techniques with those obtained from
DSC show that the DSC temperatures are systematically low by ~ 0.7 K, which is consistent with the results from the study of nFBTFO$_1$M$_7$ [19]. To avoid confusion, ten closely spaced and somewhat scattered DSC points for the mixtures are not shown, but these mixture data given in reference [10] confirm the width of the TGB$_C^n$ region and in particular show that a TGB$_C$ phase persists down to $n \simeq 10.5$. The DSC mixture data are also consistent with the dashed $N_l^*/N^*$ line ($T_5$). Since no mixtures of $n = 9$ and $n = 10$ have been studied, the position of the SmA – SmC$^*$ – TGB$_A$ point as well as the SmA – SmC$^*$ and SmA – TGB$_A$ lines is speculative but topologically reasonable in view of theory [9] and data on nFBTFO$_1$M$_7$ [6].

The most important new feature of Figure 12 is the fact that the dashed $N_l^*/N^*$ line does not represent a thermodynamic transition but the evolution of short-range TGB order. The large excess heat capacity peak ($T_5$) in the $N^*$ phase that is consistent with the predicted evolution of short-range chiral line liquid ($N_l^*$) character decreases in magnitude from 1.4 J/g for 10F$_2$BTFO$_1$M$_7$ to 0.28 J/g for 12F$_2$BTFO$_1$M$_7$. The same trend was observed for nFBTFO$_1$M$_7$, but the magnitudes were larger (1.8 J/g for 9FBTFO$_1$M$_7$ to 0.82 J/g for 11FBTFO$_1$M$_7$) [6,19].

The extremely small latent heat associated with the TGB$_A$-$N_l^*$ ($T_5$) transition in 10F$_2$BTFO$_1$M$_7$ is consistent with the small TGB$_A$-$N_l^*$ latent heat of 8.1 mJ/g for 9FBTFO$_1$M$_7$ [6] and indicates that the short-range TGB$_A$ order in the $N_l^*$ phase is very well developed just above this transition. In contrast to this, the enthalpy on going from $N_l^*$ to TGB$_C^n$ is considerably larger (150-300 mJ/g). It would appear that $\delta H$ at $T_5$ is correlated to whether the short-range TGB character in $N_l^*$ is TGB$_A$-like (large $\delta H$) or TGB$_C$-like (small $\delta H$). As noted above, when $\delta H$ ($T_5$) is large the latent heat for $N_l^*$ – TGB$_A$ transition is very small in contrast to the moderately large $N_l^*$ – TGB$_C$ latent heats observed when $\delta H$ ($T_5$) is small. It seems likely that as the alkoy chain grows ($n$ increases) the short-range TGB structure in $N_l^*$ develops a TGB$_C$ character, which is supported by the nature of the diffuse scattering shown in Figure 3. Detailed synchrotron structure studies would be of considerable interest.

All the transitions shown in Figure 12 are first order except for SmC$^*$ – SmA. Although the latter transition was not observed in the present work, it was found to exhibit typical Landau mean-field second-order behavior in 10FBTFO$_1$M$_7$ and 10.5FBTFO$_1$M$_7$ [19], and the same behavior would be expected in nF$_2$BTFO$_1$M$_7$ samples.

The second important new feature of Figure 12 is the fact that both the $n = 11$ and 12 pure homologs exhibit two closely spaced transitions, indicating that there is a different TGB phase, denoted by TGB$_C^n$, lying between $N_l^*$ and TGB$_C^n$. The structural difference between TGB$_C^n$ and TGB$_C^n$ is not known. One might speculate that TGB$_C^n$ is an incommensurate TGB$_C$ phase or a TGB$_C$ phase with a structure like that originally proposed by Renn and Lubensky [9], and X-ray data [12] on 12F$_2$BTFO$_1$M$_7$ shows that TGB$_C^n$ is either a commensurate TGB$_C$ phase or a TMGB phase.

One aspect of the proposed phase diagram that is still unresolved is the detailed behavior of the $N_l^*$ – TGB$_A$ line as it terminates at one of the TGB$_C$ phases. There are many topologically valid possibilities and not yet enough data. Indeed, a narrow range of TGB$_A$ phase was originally reported [10] for $n = 11$ although Figure 3 would suggest that this was due to well-developed short-range TGB$_A$ character in the $N_l^*$ phase just above the $N_l^*$ – TGB$_C$ transition. However, a racemic sample of 11F$_2$BTFO$_1$M$_7$ exhibits the phase sequence N-SmA-SmC [10], which suggests that TGB$_A$ should be stable close to $n = 11$ in the chiral compound. Since the $N_l^*$ – TGB$_A$ latent heat is so small in nF$_2$BTFO$_1$M$_7$, calorimetric measurements are not an ideal way to resolve this issue. Indeed, one might speculate that the tiny $C_p$ feature denoted by the question mark in Figure 6 could correspond to a $N_l^*$ – TGB$_A$ transition. A high-resolution X-ray study that clearly distinguishes diffuse and resolution-limited features is needed to answer such issues.
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References