PHYSICAL REVIEW A

VOLUME 46, NUMBER 2

Phase diagram exhibiting a smectic-A-smectic-C-smectic-F meeting point

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We report an observation of a smectic-A-smectic-C-smectic-F(A-C-F) meeting point in a binary liquid-crystalline system. Detailed high-precision x-ray measurements enabled us to characterize the meeting point as a *triple point*, a point at which the three phases coexist. A direct consequence of this result is the existence of a *tricritical point* (TCP) on the A-C boundary. Contrary to the known results, the TCP appears despite a large temperature range for the A phase and the occurrence of a nematic phase at a higher temperature. Bearing in mind that F is a tilted hexatic phase, it is suggested that a possible reason for this behavior is the molecular-tilt field arising from the coupling between the tilt and the bond-orientational-order parameters.

PACS number(s): 61.30.-v, 64.70.Md

Smectic-A(A) and smectic-C(C) liquid crystals are characterized as orientationally ordered fluids with a onedimensional mass-density wave. In the former the wave vector of this wave is along the director, while in the latter it is tilted. Experimentally, the A-C transition is generally found to be second order. As the transition is governed by a two-component order parameter it was proposed initially that the transition might exhibit heliumlike critical behavior [1]. However, subsequent studies [2-7] showed that, owing to a large bare correlation length [8], the transition is almost always mean-field-like with a relatively large sixth-order term in the Landau free-energy expansion. Based on heat-capacity results Lien and Huang [9] postulated that this transition can be driven in a firstorder manner by fluctuations seen as a consequence of the nearby isotropic-to-A phase transition. Recent experiments [10] have shown that the temperature range of the A phase has a pronounced effect on the order of the A-Cphase transition.

Extension of the two-dimensional melting theory to liquid crystals led to the prediction [11] and subsequent experimental observation [12] of bulk phases of matter referred to as hexatic phases which exhibit long-range bond orientational order (BOO) as in a solid but a short-range positional order like in a fluid. The tilted versions of this phase, viz. smectic F(F) and smectic I(I) are described by two order parameters, namely, sixfold-BOO and the molecular-tilt order parameter. Studies on these have been made all the more interesting since in the C phase the existence of finite molecular tilt gives rise to an induced BOO [13] although of a very small amplitude. Thus both C and F phases have the same symmetry. Consequently C phase transforms into F phase either through a first-order transition or evolves continuously without a transition. There are many theoretical studies [13,14] to map out different phase diagrams involving hexatic phases. However, no phase diagram with A, C, and the tilted hexatic phases has been studied so far. In this Rapid Communication we present phase diagram and x-ray diffraction studies on a binary liquid-crystal system of terephthal-bis-butylaniline (TBBA) and $n - (4 - n - n on y \log x)$ benzylidene)-4-n-butylaniline (90.4) which shows an A-

C-F meeting point. High-precision layer-spacing measurements performed on either side of the meeting point enabled us to characterize it as a triple point, at which all the three phases coexist. An interesting offshoot of these studies is the observation of a tricritical point on the A-C boundary (and the concomitant first-order-transition line). A notable feature is that for all the mixtures exhibiting the A-C transition, the A phase has a relatively large temperature range and in addition is followed by a nematic phase. This is puzzling since the A-C transition is expected [10] to become first order, in the absence of a strong lateral dipole moment of the molecule, only when the temperature range of the A phase is quite small.

The partial temperature-concentration (T-X) phase diagram, obtained by optical microscopy and x-ray diffraction studies, is shown in Fig. 1. It can be seen that for X < 0.28 (where X is the weight fraction of 90.4 in TBBA) C phase goes to smectic-G (G) phase directly and for X > 0.28 the F phase intervenes between the C and G phases and with increasing X the range of the F phase grows at the expense of the C phase. Finally, for X > 0.585 the C phase ceases to exist resulting in an A-C-F meeting point.

The x-ray experiments have been conducted on aligned samples obtained by slowly cooling the sample from the nematic phase in the presence of a 2.4-T magnetic field. The setup is essentially identical to the one described earlier [15]. The precision in the determination of the wave vector is 2×10^{-4} Å⁻¹ while the resolution in the equatorial direction is 1×10^{-3} Å⁻¹ half-width at half maximum. The temperature was maintained to a constancy of better than 10 mK during each measurement. The temperature variation of the layer spacing for X = 0.45 in the neighborhood of the A-C transition is shown in Fig. 2. The most striking feature of this diagram is the existence of the two-phase coexistence region wherein the modulations corresponding to both the phases are present clearly signifying the first-order A-C transition. This feature is exactly as expected and indeed observed earlier for a first order A-C transition [10]. Figure 3 shows the tilt angle (Φ) variation for different concentrations in the temperature range very close to A-C transition. [Φ was evaluated



FIG. 1. Partial temperature-concentration phase diagram for varying weight fraction (X) of n-(4-n-nonyloxy benzylidene)-4-n-butylaniline (90.4) in terephthal-bis-butylaniline (TBBA). The solid lines are guides to the eye. The A-C-F point is denoted by the solid circle.

by using the expression $\Phi = \cos^{-1}(d_c/d_A)$ where d_c and d_A being the layer spacing corresponding to the C and A phases, respectively.] It is observed that for $X \ge 0.45$, the tilt angle jumps abruptly to zero indicating a first-order transition while for $X \le 0.40$ it goes continuously to zero as for a second-order transition. Evidently there is a tricritical point in the concentration range $0.4 \le X \le 0.45$.

The temperature variation of layer spacing near the C-F transition is shown in Fig. 4. The abrupt jump in the



FIG. 2. Temperature variation of the smectic-layer spacing (d) in the vicinity of the A-C transition for X = 0.45. The vertical dashed lines indicate the two-phase region where the data are represented by the solid circles.



FIG. 3. Smectic-C tilt-angle variation vs the reduced temperature for X=0.54 (\blacktriangle), 0.51 (\blacksquare), 0.45 (\bullet), 0.4 (\triangle), 0.36 (\square), and 0.32 (\bigcirc).

layer spacing and the presence of the two-phase region unambiguously show that this transition is first order for all the concentrations studied. The inset shows the jump in layer spacing across C-F transition for four different mixtures. The jump in the layer spacing is observed to increase with the increase in the temperature range of the Fphase. It may be noted that a similar feature [16] was observed for the transition between C and I phases. Beyond the triple point the A phase directly goes to the F phase. The temperature variation of the layer spacing across the A-F transition for X = 0.6, a concentration which lies very close to the A-C-F meeting point is shown in Fig. 5. Clearly, the transition is first order in nature.



FIG. 4. Variation of the smectic-layer spacing (d) as a function of the reduced temperature in the vicinity of the C-F transition for x = 0.56 (\oplus), 0.44 (\odot), 0.36 (\blacktriangle), and 0.32 (\Box). For the sake of clarity, the data in the two-phase region corresponding to the C phase are not shown. The jump in the layer spacing across the C-F transition for these mixtures is shown in the inset.



FIG. 5. Temperature variation of the smectic-layer spacing (d) in the vicinity of the *A*-*F* transition for X = 0.60. The solid circles stand for the data in the two-phase region which is marked by the vertical dashed lines.

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These results lead us to infer that A-C-F meeting point is a meeting point of three first-order phase boundaries and hence is a triple point. To our knowledge this kind of a phase diagram has not been envisaged by theory so far. Furthermore, as mentioned earlier, in the vicinity of the triple-point concentration, the A-C transition changes from second order to first order even though the A-phase range is quite large. This is contrary to the known experimental results that the A-C transition becomes a firstorder one only when the strength of the transverse dipole moment of the constituent molecules is large [17] and/or the temperature range [10] of the A phase is very small. But in the present system neither of these two conditions is met and still the transition becomes a first-order one. One possible cause for this behavior is the molecular-tilt arising due to the coupling between the BOO and the molecular-tilt order parameter. Such a possibility still remains to be investigated theoretically.

The authors are indebted to Professor S. Chandrasekhar for many useful suggestions and valuable discussions.

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