## GHAPTER 7

## ORIENTATIOMAL ORDER AND TRICRITICAL BEHAVIOUR IN SOME HOMOLOGUES OF THE n-OMCPO SERIES OF LIQUID CRYSTALS

### 7.1 Introduction

Many physical properties of liquid crystals exhibit marked changes within a homologous series as the length of the end alkyl chains of the molecules is varied, The even-odd effects in the nematicisotropic transition temperature ${ }^{1,2}$ ( $T_{N I}$ ) as well as the entropy ${ }^{1}$ and orientational order parameter ${ }^{3}$ ( $S$ ) at $T_{N I}$ are some well known examples of such a behaviour. Following the recent theoretical studies of Marcel ja ${ }^{4}$ end Pink, ${ }^{5}$ who showed that the even-odd effect originates primarily from the various conformations adopted by the hydrocarbon end ohains of the molecules, there is now a renewed interest in investigating the orientational order and phase transitions in different homologous aeries, both in the pure compounds and their binary mixtures. ${ }^{6-8}$ In comparison to the pure compounds, binary mixtures have the added advantage that their average alkyl chain length may be varied in a continuous manner over any desired range. ${ }^{9}$ This property of mixtures has al so proved to be a convenient method of studying
the trioritical behaviour of the smectic A-nematic (A-K) transition in a few cases. 7,8

In this chapter, we report the results of an infrared study of the even-odd effect in the orientational order of some homologues of the trans-p-n-alkoxy- $\alpha$-methyl-p'-cyanophenyl cinnamate ( $n$-OMOPC) series of liquid crystals. These compounds possess stable, colourless mesophases at relatively low temperatures. ${ }^{10}$ The birefringence and elastic constants of some members of this series have been studied recentiy. ${ }^{11}$ We have determined the temperature dependence of the $S$ values in six homologues, the number of carbon atoms, $n$, in their alkyl ohain varying from 7 to 12 . As will be seen later, our results on the orientational order of these pure compounds as also their phase diagram prompted us to study the enthalpy of the $A-N$ transition in several binary mixtures of these homologues. Baaed an these calorimetric measurements, we have obtained evidence For tricritical behaviour of the $\mathrm{A}-\mathrm{N}$ transition in the mixtures. These results are also presented and discussed here.

### 7.2 Experimental

The liquid crystals used in these experiments were synthesized and purified in our chemistry laboratory. 10 The transition temperatures of the compounds studied are
listed in Table 7.1. Pox infrared measurements, the eagles were homoeotropically aligned between MaCl wind ows and the alignment was preserved In the smectic phase as well for compounds 10,11 and 12 . We observed that the smectic phases of these three compounds were optically uniaxial. This coupled with the fact that all these smectic phases show a simple fan shaped texture ${ }^{10}$ shower that they are all of the smectic A type. As the samples were homoeotropically aligned, unpolarized radiation could be used in all the, measur ementa. Other advantages of this technique as well as our experimental procedure have been discussed in chapters 2 and 6. As the mesophases could be supercooled consider:ably, the S values were obtained from spectra taken while cooling the samples.

The technique of randomly polishing NaCl surfaces on chamois leather did not yield well aligned samples of the lower homologues of this series with $n$ values below 7. Our attempts at inducing uniform homoeotropic alignment by treating the NaCl surfaces with a dilute solution of cetyl trimethyl ammonium bromide in chloroform were also unsuccessiful. It is for this reason that our order parameter: measurements had to be restricted to the higher homologues of this aeries,

DSC measurementa were performed using a Perkin-
Table 7.1
Pransition temperatures (in ${ }^{\circ}$ ) of the n-OMCPC homologues studied. Monotropic transitions are indicated by asterisk. given homologue are left blank. may be.

| Alkyl chain | Short notation for the homologue | Crystal to |  |  | A- | ${ }^{2}$ |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | A | N | I |  | A-I | N-I |
| $\mathrm{C}_{7} \mathrm{H}_{15}$ | 7 | - | - | 344.2 | - | - | 340.5 ${ }^{\text {m }}$ |
| $\mathrm{C}_{8} \mathrm{H}_{17}$ | 8 | - | 331.0 | - | - | - | 345.0 |
| $\mathrm{C}_{9} \mathrm{H}_{19}$ | 9 | - | 329.0 | - | - | - | 343.3 |
| $\mathrm{C}_{10} \mathrm{H}_{21}$ | 10 | - | 335.8 | - | 330.6 \% | - | 346.6 |
| $\mathrm{C}_{11} \mathrm{H}_{23}$ | 11 | - | 343.0 | - | 343.1 ${ }^{\text {\# }}$ | - | 346.7 |
| $\mathrm{C}_{12} \mathrm{H}_{25}$ | 12 | 338.0 | - | - | - | 9.5 |  |

Blmer oalorimeter (Model DSC-2). The transition temperatures were determined from the onset of the endotherma during heating cycles and are nominally accurate to $+0.5^{\circ} \mathrm{C}$; in the ease of very weak $A-N$ transition the acouracy is $\pm^{\circ} 0$. An indium calibration standard was used for calculating the enthalpies of the A-I transitions. As this transition does not superoool perceptibly, enthalpy data were obtained during both heating and oooling cycles and the average values are quoted here. A full scale range of 0.3 or $0.2 \mathrm{mcal} / \mathrm{sec}$ was used during these measurements in order to facilitate more acourate eatimates of the usually low enthaipy of the a-iil transition.

### 7.3 Results and Discussion

a) Orientational Order: For deducing the $S$ values, we utilized the strong, isolated and well defined $-C \equiv \mathbb{I}$ absorption band exhibited by all the compounds end evaluated the integrated absorbance of this band, corrected for the effects of finite spectral slit width. ${ }^{12}$ If $R$ denotes the ratio of the absorbance in the homoeotropically alignod mesophase to that in the isotropic phase, it is anown in Appendix $C$ that the orientational order parameter is given by

$$
\begin{equation*}
S=\frac{1-R}{1-\frac{3}{2} \sin ^{2} \alpha} \tag{7.1}
\end{equation*}
$$

where $\alpha$ is the angle between the transition dipole moment of the band and the long axis of the molecule. The $n-O M C P C$ molecule may adopt either of two aifferent conformations ${ }^{10}$ as shown in Fig. 7.1. Assuming that the long axis of the molecule lies along the line joining the centers of the two benzene rings, it is seen that $\alpha$ 1s $\sim 18.4^{\circ}$ for conformer I and $4^{\circ}$ for conformer II. Although the presence of conformer II cannot be ontirely ruled out, Sadashiva has indicated ${ }^{10}$ that conformer I is gterfoally more favoured and that free rotation around the $=0-\mathrm{C}$ - is not too likely since the single bond has

some double bond character due to resonance. For this reason, we have taken $a$ to be $18.4^{\circ}$ and calculated the $S$ values assuming that the population of conformer TI is negligible in the mesophases. This assumption, of course, remains $t o$ be verified by other methods such as X-ray difiraction which can determine the average breadth of the molecules in the mesophases and hence the preferred conformation. In any case, this question is important only with regard to the magnitude of the correction factor due to $a$, as seen from ohe denominator of eqn.7.1. Thus, should subsequent $X$-ray studies indicate that our assumption is not justified, then given a knowledge of the relative abundances of the two conformers, it is a simple matter to accordingly scale down the $S$ values

(1)

(II)

## Figure 7.1

Molecular structure and two possible conformations of the rigid gart of trans-p-n-alkoxy- $\alpha$-methyl p'-cyanophenyl cinnamates ( $n$-OMCPC). The long axis of the molecules is taken to be the line joining the centres of the two benzene rings. The alkyl chain $\left(\mathrm{C}_{n} \mathrm{H}_{2 n+1}\right)$ is denoted by $R$.
determined bj us. But for this consideration, our experimental geometry leads to the absolute values of $S$ as other corrections due to the anisotropy in the refractive indices and the polarizability of the ilquid crystalline medium are known to be well below our normal experimental error. The oalculated $S$ valuea ers estimated to be accurate to well within $5 \%$.

Figures 7.2 and 7.3 show the temperature dependence of $S$ for compounds 7-9 and 10-12, reapectively. In the nematic phases of compounds $7-9$, it is seen that as usual the $S$ values decrease monotonically as $T_{c}$ is approached. 10 OMOPC exhibits a monotropic A-18 transition $16^{\circ} \mathrm{K}$ below $\mathrm{T}_{\mathrm{NI}}$ It is seen from Fig. 7.3 that in this case the change in 8 st It $_{\text {AH }}$ is hardy discernible. This is in accord with our calorimetric measurements discussed below, which gives a rather low value of $\sim 1.0 \mathrm{cal} / \mathrm{mole}$ for the enthalpy of this transition.

On the other hand, there is a fairly large and abrupt change in $S$ at the monotropic $A-N$ transition of 11 OMOPC, characteristic of a distinctly first order transition; the enthalpy of this transition is also found to be nearly 40 times larger when compared to 10 OMCPC.

12 OMCPC exhibits only a smeotic A mesophase. The observed $S$ values in this case, even close to the is otropic transition, are accordingly much higher than


Figure 7.2

Variation of $S$ in the nematic phase of compounds 7-9.


Figure 7.3

Variation of $S$ in the smectic $A$ and nematic phases of 10 and 1 and in the smectic A phase of 12. The arrows show the $A-N$ transition points.
> those obtained near $\mathrm{T}_{\mathrm{mI}}$ in the other compounds studied here.

$X t$ is of interest to compare the $S$ values of all the se mesogens at a common relative temperature in the vicinity of the isotropic transition. Figure 7.4 shows these data $2^{\circ}$ below the respective isotropic transition temperature in each case. Here each $S$ value corresponds to the average of three independent measur ements. Accordingly, the error limit on these values is within $\sim 3 \%$. The isotropic transition temperatures of the compounds are also shown for oomparison. For compounds 8-10, Karat ${ }^{11}$ has determined the relative $S$ values from birefringence measurements. We have compared his data with ours, after normalizing the two sets of data for 9 OMOPG. Within the accuracy of the present measurements, a reasonable agreement exists between both sets of data for compounds 8-10.

For compounds 7-10, it is seen that the trend of even-odd effect in $S$ values is at best very week. Their respective $\mathrm{I}_{\text {MI }}$ values also show a similar behaviour. As this series possesses an alkoxy end group, it is waderstandable that among these four compounds, the odd members may have slightly lower $S$ values than the even members.4,5 Even so, the observed variation in $S$ is admittedly comparable to the maximum possible error limit on these
data. This is not entirely surprising considering that the end chains are already quite long fax the homologues considered hers and that the even-odd offect progressively diminishes for higher homologues within any series. Furthermore, the even-odd effect in $\mathrm{T}_{\mathrm{HI}}$ itself is seen to be quite mall far compounds 7-11. A more pronounced even-odd effect in $T_{\text {MI }}$ and relative S values has been observed in the lower homologues of this seriea. ${ }^{10,11}$

In Fig. 7.4, the $S$ value is seen $t o$ increase on going from 10 to 11. This behaviour is at variance from the pattern exhibited by the lower odd homologues. At $T_{c}-2,11$ OMCPO is barely $1.6{ }^{\circ} \mathrm{K}$ above $T_{A N}$. The presence of near neighbour correlations resulting in short range smectic like order within the nematic phase is sexy probably the explanation for the anomalous increase in this case. Evidently, prediotions based on mean-field theories ${ }^{4,5}$ cannot account for the effects arising irom such interactions.
b) Tricritical behaviour: Considerable interest has been evinced recently in tricritical $A-N$ transitions. By varying a parameter related to the molecular packing or intermolecular interactions, the character of the, $A-N$ transition can be made to evolve in a continuous manner from first to second order. For this purpose, concentration


## Figure 7.4

Variation of $S$ (circles) and the isotropic transition temperature, $\mathbb{T}_{c}$ (triangles), with the number of carbon atoms in the alkyl chain. The ordinate marks corresponding $t o \mathbb{T}_{c}$ are shown on the right vertical axis. She $i$ values are referred to a common relative temperature of $\left(\mathbb{T}_{c}-2\right)^{\circ} \mathrm{K}$. The variation from 11 to 12 is shown by broken lines as 12 exhibits only the smectic A phase.
serves as a convenient variable in binary mixtures, ${ }^{9}$ whereas pure compounds require 4hs application of high pressures. 13-16 at the tricritical point, where the Ines of first and second order transitions meet, the enthalpy of the transition vanishes and the first order transition yields place to one of critical behaviour. Our results on the variation of $S$ values at $\Psi^{A N}$ as also the phase diagram of compounds 9-12 led us to investigate the thexmodynamic behaviour of binary mixtures of these homologues. Based on the theories proposed by McMillan ${ }^{17}$ and Lee et al. ${ }^{18}$, a tricritical point in the $A-N$ traneition is expected to occur at the reduced temperature $T_{R}=T_{A N} / T_{N I} \sim 0.88$. From Table 7.1 it is seen that $T_{R}$ decreases on going from 11 to 10, their respective values being 0.99 and 0.95 . It was also noted earlier that the enthalpy of the A-N transition decreases drastically while going from 11 to 10, i.e., as the alkyl chain length. decreases. These factors suggest that one may be observing here a tricritical behaviour which can be studied in further detail by varying the average alkyl chain length in a continuous manner. Thus binary mixtures of 10 or 11 with lower homologues may reveal, at atmospheric pressure, the presence of a trioritical point at an appropriate ratio In the phase diagram of the mixtures. In fact, some homologues in this series with $n$ values below 10 might
conceivably have second order 816 transitions which are not observed presumably because of the intervention of the solid phase. Another noteworthy feature from Table 7.1 is that the nematic range decreases on going from 10 to 11 and disappears altogether for 12 . Here again, the phase diagram of binary mixtures of 11 and 12 may reveal, at atmospheric pressure, the existence of a triple point between the smectic $A$, nematic and isotropic phases.

Motivated by these considerations, we performed calorimetrio measurements on several binary mixtures wherein 11 was one component while the other was 12, 10 or 8 as the case may be. If $A$ and $B$ denote the two components and $X_{A}$ and $X_{B}$ their respective mole par cent concentration in the mixture, the average number of carbon atoms in that alkyl chain is given by

$$
n_{a}=\frac{X_{A} n_{A}+X_{B} n_{B}}{100},
$$

where $n_{A}$ and $n_{B}$ are the respective number of carbon atoms in the alkyl chain of the pure homologues. The different mixtures studied and their $n_{2}$ and $T_{R}$ values are summarized in Table 7.2. Figure 7.5 shows their transition temperatures $T_{A N}$ and $T_{N I}$ as a function of $n_{a}$. The transition temperatures of the relevant pure compounds

## TABLE 7.2

Mixtures investigated and their corresponding $n_{a}$ and $\mathbf{R}_{\mathbf{R}}$ values. $A$ and $B$ denote the two components and $x_{A}$ and $x_{B}$ their reapective mole par cent concentration in the mixture

| $A-B$ | $X_{A}$ | $X_{B}$ | $n_{\mathbf{a}}$ | $\mathbf{T}_{\mathbf{R}}$ |
| :---: | ---: | ---: | ---: | :--- |
|  | 26.5 | 73.5 | 11.74 | 1.0 |
|  | 53.3 | 46.7 | 11.47 | 0.996 |
|  | 75.1 | 24.9 | 11.25 | 0.992 |
|  | 0.0 | 100.0 | 11.00 | 0.990 |
|  | 28.6 | 71.4 | 10.71 | 0.983 |
|  | 52.4 | 47.6 | 10.48 | 0.977 |
|  | 70.3 | 29.7 | 10.30 | 0.971 |
|  | 100.0 | 0.0 | 10.0 | 0.954 |
|  | 5.2 | 94.8 | 10.84 | 0.987 |
|  | 7.2 | 92.8 | 10.78 | 0.985 |
|  | 9.0 | 91.0 | 10.73 | 0.983 |
|  | 13.2 | 86.8 | 10.60 | 0.980 |
|  | 19.0 | 81.0 | 10.43 | 0.972 |
|  | 22.4 | 77.6 | 10.33 | 0.970 |
|  | 35.3 | 64.7 | 9.94 | 0.947 |
|  | 37.4 | 62.6 | 9.88 | 0.933 |



Figure 7.5

Variation of $\mathrm{T}_{A N}$ and $\mathrm{I}_{\mathrm{NI}}$ in three fanilies of mixtures as a function of the average number of carbon atoms, na' in the alkyl chain. The transition temperatures of the pure compounds are also shown for comparison. Circles: 8-11 mixtures; triangles: 10-11 mixtures; squares: 11-12 mixtures; crosses: pure compounds.
are also shown for comparison. It is interesting to note that although data corresponding to three separate families of mixtures are being compared, they all point to a common trend when the average chain leng th or $n_{a}$ is considered to be the variable, i.e., the nematio range increases as $n_{2}$ deoreases. Also, as expected, $T_{R}$ is seen to diminish as $n_{a}$ is lowered in the mixtures and this indicates an approaching tricritioal point in the $A-\mathbb{N}$ transition. The antioipated triple point between the smectic A, nematic and isotropic phases is seen to $000 u \leq$ in a $11-12$ mixture at the mole per cent ratio of $\sim 27: 73$.

When compared to $\boldsymbol{T}_{\text {AII }}$, the variation of $\mathbb{T}_{\text {MI }}$ in all three families of mixtures is inuoh smaller. This is not surprising considering that the $\mathbb{T}_{\mathbb{N I}}$ values of the relevant pure compounds are themselves quite olose to each other.

It is of interest to examine the specific manner in which the enthalpy of the $A-N$ transition $\left(\Delta H_{A N}\right)$ varies against $\mathbf{T}_{\mathbf{R}}$ as the tricritical point is approached. In Fig. 7.6 the $\Delta H_{A N}$ values obtained in 8-11 and $10-11$ mixtures are plotted as a function of $\mathbf{T}_{\mathbf{R}^{\prime}}$. In the case of the mixtures, the $\Delta H_{A H}$ values pertain to the appropriate average molar weight for any ven composition of the mixture. To the extant that $\Delta H_{A N}$ varies against $\mathbb{T}_{R}$ in a non-linear fashion. the present results are in


Figure 7.6

Variation of $\Delta H_{A N}$ with $R_{R}$ in $8-11$ and $10-11$ mixtures, Experimental uneertainties in the three lowest enthal py values are also indicated.
qualitative agreement with the theory proposed by Lee et al. ${ }^{18}$ Our extrapolated data show that the A-M transition would become second order for $\mathrm{m}_{\mathrm{R}}<0.94 \pm 0.005$. In order to ascertain this, we investigated an 8-11 mixture with a $T_{R}$ value of 0.933 . The $A-N$ transition in this case was detected only as a leeble discontinuity in the baseline of the DSC trace, thus indicating that the transition is of second order. The corresponding $\triangle H_{A N}$, if any, was indeed too small to be measured within the accuracy of the DSC-2. The T AN and TNI values of this mixture were also confirmed through microscopic observation using a hot stage.

Our $T_{R}$ value at the tricritical point is considerably higher than that predicted by theory. Similar deviations have been reported earlier in a few other cases as well. For example, Johnson et al. ${ }^{8}$ obtained a value of 0.96 from studies on binary mixtures of another series. Using hydrostatic pressure rather than the concentration of mixture as the variable, $T_{R}$ values of 0.92 and 0.90 were reported in two instances. 13,14 In certain other cases, however, pressure as well as concentration dependent studies have yielded $T_{R}$ values in very close agreement with theory. 7,15,16 Such differences amongst different homologous series on the one hand and between theory and experiment on the other,
remain to be explained satisfactorily. En this respect, differences between the rigid core sizes of mesogenic molecules may be a relevant factor in determining the exact tricritical behaviour observed in each series.

### 7.4 Conclusions

The results of this study emphasize the olose link that exists between the molecular ordering and thermodynamic behaviour of mesophases. The even-odd effect in both $S$ and $T_{N I}$ is not very pronounced in the higher homologues we have investigated, as they all possess relatively long alkyl end chains. Nevertheless, the observed disoontinuities in $S$ values at the $A-N$ transition clearly reflect the enthalpy of this transition in the two homologues 10 and 11 . Che calorimetric results on binary mixtures show the presence of a concentration dependent triple point between $A, N$ and I phases. In addition, a tricritical point is shown to oocur in the $A-\mathbb{N}$ transition at a reduced temperature of $\mathbf{- 0 . 9 4}$. Further studies of the critical exponent as well as the fluctuations in concentration and order parameter in the vicinity of this tricritical point would be of considerable interest.

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