Study of the mesophase behavior of the possible dimers formed through mixing of two different parasubstituted benzoic acid derivatives

Refaat I. Nessim¹ and Imad Ed. Albalaa²

¹ Dept. of Chemistry, Faculty of Science, Cairo University, Giza- Egypt
² Dept. of Science, Faculty of Basic Education, PAAET
refaat nessim@hotmail.com

Abstract: Binary mixtures of two types of 4-substituted benzoic acids, covering the whole composition range, were prepared by melting both components together, stirring to give an intimate blend, and then cooling to room temperature. The mixtures prepared were characterized for their mesophase behavior by differential scanning calorimetry (DSC) and polarized-optical microscopy (POM). The first type of the acid is substituted with a small compact polar group that cover a wide range of polarity. The other type of the acid is 4-alkoxy benzoic acid bearing an alkoxy (OCnH2n+1) group with varying chain length (n = 6−16). Binary phase diagrams were constructed whereby all phases observed were identified by POM. The smectic C. (SmC) mesophase was observed in all mixtures investigated, while the nematic phase (N) was observed, together with the SmC phase, only in mixtures possessing the acid homologues with n≤12 carbons. Complex formation was also supported by FT-IR spectroscopic measurements.

[Refaat I. Nessim and Imad Ed. Albalaa. Study of the mesophase behavior of the possible dimers formed through mixing of two different para-substituted benzoic acid derivatives. *Nat Sci* 2014;12(11):108-113]. (ISSN: 1545-0740). http://www.sciencepub.net/nature. 17

Keywords: Alkoxy benzoic acid, Smectic C, Nematic phase

Introduction

Usually, at least two rings are required to enable the generation of liquid crystal phases; however, 4-alkoxy (or alkyl) benzoic acids are the only examples of one-ring mesogen. In a single molecule, the actual molecular species is certainly not long or lathlike; however, dimerization through hydrogen-bonding creates a long lath-like structure (1) with a three-ring core and two flexible terminal chains [1].

Dimmer A-A (1)

Benzoic acid derivatives were the most frequently employed components for the formation of LC materials through such hydrogen-bonding interaction ^[2]. Wing polar groups are usually used in the modification of a liquid crystal material. Since calamitic (rod-like) molecule has two ends, the wing groups may be similar or different. Many terminal units have been employed in the generation of liquid crystals, but the most successful route ^[3] is to use either a fairly long, straight hydrocarbon (usually alkyl or alkoxy), or small polar substituent (e.g. CH₃, CH₃O, CN, NO₂, Cl, etc.).

The role of these groups is to act either as a flexible extension to the core or as a dipolar moiety to introduce anisotropy in the physical properties of the molecule. The option to choose dissimilar wing groups in the case of 4-substituted benzoic acids is not allowed, since we come with either an **acid B**, (I_{a-f}) , substituted with a small compact polar terminal group (X) to give the non-mesomorphic dimmer (2), or the mesomorphic benzoic acid (In) substituted

with 4-alkoxy group (acid A) to give the mesomorphic dimmer (1).

$$C_nH_{2n+1}O$$

ln

Acid A...I6, n=6; I8, n=8; I10, n=10; I12, n=12; I16, n=16

 $\begin{aligned} &\textbf{I}_{a:f} \\ \textbf{Acid B} ...I_{a^{\prime}} &\textbf{X=OCH}_{3}; I_{b^{\prime}} &\textbf{X=CH}_{3}; I_{c^{\prime}} &\textbf{X=CI}; I_{d^{\prime}} &\textbf{X=Br}; I_{a^{\prime}} &\textbf{X=CN}; I_{f^{\prime}} &\textbf{X=NO}_{2} \end{aligned}$

Dimmer B-B (2)

The goal of the present investigation is to allow hydrogen bond formation between two dissimilar acids (In and I_{a-f}) aiming to form a new type of mesogen (3) that bears two dissimilar wing groups (X and CnH2n+1O).

$$C_nH_{2n+1}O$$
 O OH OH OH

Dimmer A-B (3)

The idea can be realized simply by investigating, first, the binary phase behavior through binary phase diagrams of two differently substituted benzoic acids; the first bears a small compact group of different polarity and the other bears a terminal alkoxy group of varying chain length. The study aims further to investigate the mesomorphic characteristics of the all possible supramolecular complexes that could be formed alternatively between each of the acids I_{a-f} and I_{n-f} .

Experimental

Chemicals were of pure grades and purchased from the following Companies: Fluka, Buchs, Switzerland; MP Biomedicals, Inc., Illkirch, France; BDH, Poole, England; Aldrich, Wisconsin, USA; and E. Merck, Darmstadt, Germany. Infrared spectroscopic measurements were recorded at room temperature in a Nicolet, model Magna 560 FT-IR spectrometer at a resolution of 2 cm⁻¹.

Calorimetric measurements were carried out using a PL-DSC of Polymer Laboratories, England. The instrument was calibrated for temperature, heat and heat flow according to the method recommended by Cammenga et al. [4]. DSC measurements were carried out for small samples (2–3 mg) placed in sealed aluminum pans. All of the thermograms have been achieved at a heating rate of 10 °C/min in an inert atmosphere of nitrogen gas (10 ml/min).

Transition temperatures were checked and type of mesophase identified for the mixed derivatives (In), and the binary associates of the acids (In and I_{a-f}), using a standard polarized-optical microscope POM (Wild, Germany) attached to a home- made hot-stage where the temperature is measured by a thermocouple attached to Brookfield temperature controller, England.

2.1. Preparation of supramolecular complexes

For the preparation of the supramolecular acid complexes (In/I_{a-f}), binary mixtures of any two complimentary acids, were prepared in a 1:1 molar ratio of samples, (acid A:acid B equal 1:1), by melting the appropriate amounts of each component, stirring to give an intimate blend, and then cooling with stirring to room temperature. For the construction of binary phase diagrams, the mixtures of the two components were prepared to cover the whole range of composition. Transition temperatures

obtained for all prepared blends, as measured by both DSC and POM, agreed within 2–3 °C. In the phase diagrams constructed by plotting the transition temperatures versus mixture composition, the symbol "○" denotes solid-mesophase, "□" mesophase-isotropic transitions, "●" mesophase-another mesophase, and, "▲" eutectic temperature.

Results And Discussion

3.1. Characterization of hydrogen-bonded complexes.

Hydrogen-bonded complex formation was confirmed by Table 1, which shows the acid carbonyl stretching frequencies (cm⁻¹) in their homo and hetero substituted states for some selected examples of the complexes.

Table 1

Complex formed	Acid carbonyl absorption frequency Cm ⁻¹
Homo complex I12	1682
Homocomplex Ie	1682
Homocomplex Ia	1720
Hetero complex I12/Ia	1682
Hetero complex I12/Ie	1702

3.1.1. Infrared spectra

Complex formation was supported by FT-IR measurements, according to which the initial dimeric acid, A–A or B–B, is replaced by the hetero complex dimmer, A–B [2]. Certainly, the relative acid strength of the two complementary components of the complex has a crucial effect upon the strength of newly formed hydrogen-bond, and consequently upon mesomorphism. This in turn, is influenced by the nature of the terminal substituents on both acids. Thus, for instance, electron-withdrawing substituents are expected to enhance acidity of the acid-proton, and consequently add extra stability of the hydrogenbonding. Generally, the hydroxyl stretching vibration band appears in the region 3400–3600cm⁻¹, however, the resolution is so poor that it is not possible to discern between free and hydrogen-bonded ones. Therefore, only the carbonyl stretching region is used to confirm hydrogen-bond formation [5]

Table 1 summarizes the acid carbonyl frequencies in their similar and dissimilar states, for some selected examples of the hetero complexes that show extremes in polarities.

As can be seen from Table 1, Carbonyl groups of the homo acids of the type A–A, that bearing either the alkoxy group e.g. $C_{12}H_{25}O$ in I12 or the methoxy group in I_a of the homo acid, both absorb at the same frequency (1682cm^{-1}) revealing that the length of the alkoxy group does not affect the polarizability of the carbonyl group. On the other hand, the electron-withdrawing cyano group, located in the para position with respect to the carbonyl

group in I_e increases its double bond character and a red-shift is observed in the carbonyl-absorption frequency that appeared at 1720cm^{-1} .

With respect to the complex $I12/I_a$, that is substituted on both sides with two different alkoxy groups (CH₃O and C₁₂H₂₅O), it absorbs at the same frequency as that of the pure components, namely 1682cm^{-1} . Alternatively, the complex $I12/I_e$ that is terminally substituted with the electron-donating C₁₂H₂₅O on one side and the electron-withdrawing CN group on the other side, proved to exhibit one absorption band at intermediate frequency, 1702cm^{-1} . 3.1.2. Constructing the binary phase diagrams covering the whole range of composition for any two dissimilar acid components

It is well known that 4-substituted benzoic acids form dimmer via hydrogen-bonding as given by the formulae 1 (A–A) and 2 (B–B). Upon mixing two different acids, one from each group (I_{a-f} and In), there are two possibilities to arrange the molecules of one component within the molecular arrangement of the other.

The first possibility is that long-lath dimmers (A-A and B-B) will be arranged side-by-side without being ruptured during mixing. In such case, a solid solution is expected either to from both components with melting temperature changes regularly from that of the first component to that of the other, or at least pass through one eutectic composition. Also in such case, the mesophase stability of the alkoxy component (1) will be gradually disrupted by the gradual addition of the non-mesomorphic component (2), and consequently, linear composition dependence is expected for mixed mesophase stability.

The other possibility for molecular arrangement in mixed systems may take place by rupturing the hydrogen bonds in both dimmers resulting in the formation of a new unsymmetrical dimmer (3).

To investigate which type of molecular arrangement actually takes place in our system, let us first construct the binary phase diagrams for selected representative examples composed from both types of acids. Two homologous of extreme chain length (I8 and I16) were chosen from the acid 1 that bears 8 and 16 carbons, respectively. The longer homologue is smectogenic, possessing the smectic C (SmC) phase as the only mesophase, while the shorter one is dimorphic, possessing both the SmC and nematic (N)mesophases.

On the other hand, two extremes of the acid (2) were selected where one (I_a) bears the strong electron-donating CH_3O group and the other (I_e) is substituted with the strong electron withdrawing CN group

3.2. Binary phase behavior

Figure (1) represents the two binary phase diagrams of the electron-donating substituted acid (I_a) with the two homologues of 4-alkoxy benzoic acid (I_a) with the two homologues of 4-alkoxy benzoic acid (I_a) and I_a 16). As can be seen from Fig. 1, independent of the length of the alkoxy chain (8 or 16 carbon atoms), supramolecular complexes are produced in both systems. Complex formation in the solid phase is evidenced by the two eutectic points, one preceding and the other following the 1:1 molar ratio (50 mol%). The hydrogen-bonded complex is retained in the mesophase as confirmed by the enhanced stability of the mesophase. These results revealed that the second possibility of molecular rearrangement, as given by formula (I_a 1), most probably prevails.

To test if complex formation is restricted by the polarity of the substituent X, we are going to investigate the binary phase behavior of the other extreme in polarity, where X is the strong electron-withdrawing CN group, in I_e

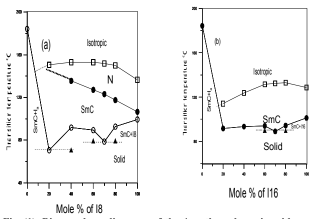


Fig. (1). Binary phase diagrams of the 4-methoxy benzoic acid with: (a) octyloxy benzoic acid and (b) hexadecyloxy benzoic acid.

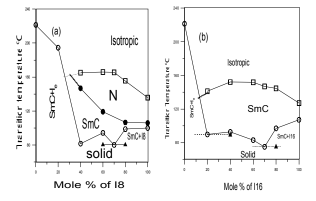


Fig. (2). Binary phase diagrams of the 4-cyano benzoic acid with: (a) octyloxy benzoic acid and (b) hexadecyloxy benzoic acid

Figure (2) describes the binary phase behavior of I_e when mixed with the same homologues of acid I8 and I16. As can be seen from Fig. 2, hydrogen bonded complexes are again observed in either the solid or mesophases, independent of the alkoxy-chain length. These results revealed that the affinity of individual acids towards hydrogen bond formation with each other is much greater than that occurring within molecules of the same acid. This may be ascribed to the dual electronic effect of the polar substituent on the individual members of the group I_{a-f} . That is, electron-donating group will increase the electron density on the carbonyl group of the acid, thus enhances the strength of the hydrogen bond. Alternatively, electron-withdrawing group lead to an increase of the acidity of hydrogen, leading to increased probability for hydrogen bond formation.

Being confirmed, all possible supramolecular complexes (3) were prepared using each of the 4-substituted benzoic acids $I_{\rm a-f}$, alternatively with each

of the homologues In, in a 1:1 molar ratio and their mesophase behavior investigated by DSC and phases detected by POM. The results are collected in Table 2

3.2.1. Effect of alkoxy-chain length on complex formation

In order to investigate the effect of the alkoxychain length on complex 3, on the liquid crystalline properties of the supramolecular complexes, the dependences, on the acid alkoxy-chain length (n), of transition temperatures of the supramolecular complexes are depicted in Fig. 3. As can be seen from Fig. 3, polar substituent, whether it is electron-donating or electron-withdrawing, are effective in promoting the smectic C mesophase. Irrespective of the substituent, X, the stability of the SmC phases decreases slightly with the increase of the length of the alkoxy chain.

Table 2. Phase transition Table temperatures (°C) of the 1:1 complexes of the systems In/Ia-f.

System	n	X	$T_{\text{Cr-C}}$	T _{C-N}	T _{C-I}	T _{N-I}	$T_{\rm C}^{-112}$
I6/Ia	6	СН3О	95.2	144.3	-	155.3	20.7
I6/Ib	6	СН3О	96.5	120.7	-	143.1	20.4
I6/Ic	6	Cl	100.6	132.2	-	147.1	20.5
I6/Id	6	Br	93.5	132.5	-	151.2	20.6
I6/Ie	6	CN	99.5	144.0	-	163.7	20.9
I6/If	6	NO2	107.7	122.0	-	172.1	21.1
I8/Ia	8	СН3О	94.2	140.1	-	155.7	20.7
I8/Ib	8	CH3O	106.2	123.2	-	143.1	20.4
I8/Ic	8	Cl	95.1	136.2	-	151.2	20.6
I8/Id	8	Br	96.5	131.4	-	147.1	20.5
I8/Ie	8	CN	80.0	144.8	-	172.2	21.1
I8/If	8	NO2	89.6	140.1	-	176.3	21.2
I10/Ia	10	CH3O	94.4	-	147.1	-	20.5
I10/Ib	10	CH3O	89.5	130.3	-	138.9	20.3
I10/Ic	10	Cl	87.7	-	143	-	20.4
I10/Id	10	Br	86.5	-	147.1	-	20.5
I10/Ie	10	CN	108.2	-	159.5	-	20.8
I10/If	10	NO2	88.9	-	-	167.9	21.0
I12/Ia	12	CH3O	108.7	-	140.8	-	20.4
I12/Ib	12	CH3O	97.0	-	133.9	-	20.2
I12/Ic	12	Cl	96.6	-	140.6	-	20.3
I12/Id	12	Br	100.0	-	142.6	-	20.4
I12/Ie	12	CN	95.7	-	160.5	-	20.8
I12/If	12	NO2	99.1	-	164.0	-	20.9
I16/Ia	16	CH3O	95.8	-	129.7	-	20.1
I16/Ib	16	CH3O	93.5	-	126.0	-	19.9
I616/Ic	16	Cl	86.5	-	130.9	-	20.1
I16/Id	16	Br	87.5	-	129.7	-	20.2
I16/Ie	16	CN	86.2	-	147.8	-	20.5
I16/If	16	NO2	95.3	-	142.9	-	20.4

The nematic phase appears only with the shorter homologues (I6 and I8) and remains up to the homologue I10 with 4-methyl benzoic acid (I_b). The greatest nematic temperature range is observed with

the 4-nitro analogue, while the smallest range is exhibited by the 4-methoxy substituted complex. As usual the melting point of the complexes does not change regularly with chain length.

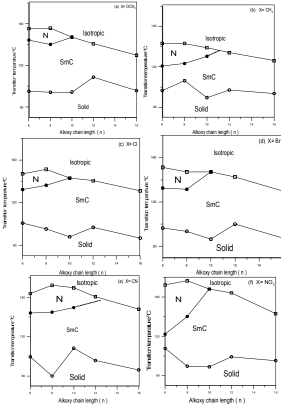


Fig. (3). Effect of the alkoxy chain length on the mesophase behavior of the supramolecular complexes (In / Ia-f).

3.2.2. Transition temperatures and polarizability anisotropy of the Car-X

The relationship between the stability of the mesophase, expressed as the clearing temperature, T_C , and the anisotropy of polarizability ($\Delta\alpha X$) of bonds to the small compact terminal substituent (Car–X), was studied by van der Veen ^[6]. The relationship

is expressed as:

 $T_{C} \alpha (\Delta \alpha_{M} + \Delta \alpha_{X})^{2}$ (1) where T_{C} is measured in Kelvin. Eq. (1) can be put in the form [7]:

$$T_C^{1/2} \alpha \left(\Delta \alpha_M + \Delta \alpha_X \right) = a \Delta \alpha_M + a \Delta \alpha_X \tag{2}$$

The term $\Delta\alpha M$ is the polarizability anisotropy of the whole molecular structure except the terminal substituent, X, and "a" is the proportionality constant. The values of $\Delta\alpha X$ were calculated $^{[4]}$ from the data given by Le Fevre and co-workers $^{[8-10]}$. Thus, if, $T_C^{\ 1/2}$ of any of the investigated hydrogen-bonded dimmer bearing one and the same acid alkoxy substituent "C_nH_2n+1" but of varying polar substituent "X", is plotted against $\Delta\alpha_X$, a straight line is expected with a slope "a" and intercept equals "a $\Delta\alpha_M$ " where

" $\Delta \alpha_{\rm M}$ " is the polarizability anisotropy of the whole of the molecular structure except the substituent (X).

Figure (4) illustrates the dependency of the $T_C^{1/2}$ values on the polarizability anisotropy of the bonds to X in the polar substituted benzoic acid component of the supramolecular complex, individually, with different substituted acids (I_a – I_f).

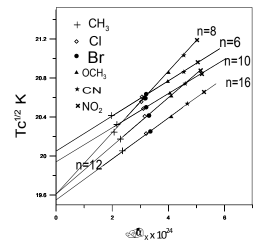


Fig. (4). Dependence of the mesophase stability $(Tc^{1/2})$ of the investigated supramolecular complexes on the polarizability anisotropy $(\Delta\alpha_X)$ of the acid substituent (X).

As can be seen from Fig. 4, fairly linear dependencies were observed in nearly all series investigated. This indicates that polarity and/or polarizability on one side of the complex play a main role in stabilizing the mesophase of the resulting unsymmetrical dimmer. It would be worth mentioning here that the chloro and bromo substituted acid derivatives are more effective, than the unsubstituted analogue, in stabilizing the hydrogen-bonded associates.

The electromeric releasing power of the halogen is completely counterbalanced by its electron-withdrawing inductive character, resulting in an increased mesophase stability of the resulting associate, as evidenced by their relatively high $T_{\rm C}$ values. Unsubstituted benzoic acid was tested but the mixtures do not show a mesophase.

Figure (4) was used to calculate the slope and intercept of each of their regression lines from which $\Delta\alpha_M$ are calculated. The results of the computation are given in Table 3 which shows that the polarizability anisotropy $\Delta\alpha_M$ of the whole complex, except the substituent X, decreases according to n in the order: 10>6>16>12>8. This order reveals that the polarizability anisotropy of the whole molecular structure, expect the substituent X, is independent on the length of the molecule.

Table 3. Regression analysis data for the Van der Veen correlations of Supramolecular complexes investigated.

System	n	X	Slop2x10 ⁻²³	Intercept	$\Delta \alpha_{\rm M} x 10^{23}$
I6/I _{a-f}	6	varies	1.8	20.05	11.13
I8/I _{a-f}	8	varies	3.15	19.61	6.23
$I10/I_{a-f}$	10	varies	1.75	19.94	11.39
I12/I _{a-f}	12	varies	2.47	19.61	7.94
I16/I _{a-f}	16	varies	2.1	19.55	9.31

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10/25/2014