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Hydrogen-bonded liquid crystals built from hydrogen-bonding donors and acceptors

Infrared study on the stability of the hydrogen bond between carboxylic acid and pyridyl moieties

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The stability of a hydrogen-bonded complex built through inter-molecular hydrogen bonding between carboxylic acid and pyridine fragments has been examined using infrared spectroscopy. Infrared spectra as a function of temperature have been recorded for the 1:1 complex of 4-hexyloxybenzoic acid and *trans*-4-propoxy-4'-stilbazole from the crystalline state to the isotropic state. A dependence of the stability of the hydrogen bond on molecular orientation is observed clearly in the infrared spectra. The spectra also suggest that the hydrogen bond is an unionized type with a double minimum potential energy.

1. Introduction

In 1988, we discovered [1-3] that mesogenic structures could be obtained by selfassembly of pyridine and carboxylic acid fragments through the formation of intermolecular hydrogen bonds. For example, complexation of 4-butoxybenzoic acid 4OBA and *trans*-4-[(ethoxybenzoyl)oxy]-4'-stilbazole 2OPhSz through hydrogen bonding results in the formation of an extended mesogen that exhibits a more stable mesophase than either of the single components [1,3]. Mesogens with well-defined structures are formed from independent and different molecules. The linear and rigid structure of the complex is achieved through the connecting part of the single hydrogen bond as shown in the scheme.



The concept of building mesogenic structures through the hydrogen bond between the H-bond donor and acceptor moieties has been extended to a variety of mesogenic

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structures that form a novel family of liquid crystals [1-10]. A non-mesogenic molecule can function as a part of a core unit through the hydrogen bonding [5, 6]. For example, a bifunctional H-bonding acceptor, 4,4'-bipyridine, which is capable of recognizing and binding H-bonding donor molecules at each of its pyridyl ends, forms a mesogen with two benzoic acid molecules. The H-bonded mesogen shows stable mesomorphic behaviour. Hydrogen-bonded polymer systems based on a variety of backbones such as polyacrylates or polysiloxanes exhibit mesogenicity over wide temperature ranges [2, 3, 7-10]. Hydrogen bonding is one of the key interactions used by nature to assemble molecules. This process of molecular self-assembly using the hydrogen bond enables us to design and build new types of functional molecular complexes through non-covalent bonding.

Significant thermal stability is observed for the hydrogen-bonded mesogenic complex obtained from the carboxylic acid and the pyridyl group. For example, the mesophase of a 1:1 complex of 4OBA and 2OPhSz is stable up to 238°C [1]. The 1:2 complex from isophthalic or terephthalic acid and *trans*-4-octyloxy-4'-stilbazole exhibits a smectic phase up to 250°C [6]. Molecular ordering of the mesomorphic state may contribute to the stabilization of the hydrogen bonding of the molecular complex.

The objective of the present study is to examine the stability of the hydrogenbonded mesogenic complex within various phases by infrared spectroscopy. Infrared spectroscopy provides useful information on hydrogen bonding. The mesogenic 1:1 complex from 4-hexyloxybenzoic acid 6OBA and *trans*-4-propoxy-4'-stilbazole 3OSz has been examined by infrared spectroscopy from the crystalline state to the isotropic state. A hydrogen-bonded dimer of 4-hexyloxybenzoic acid, $(6OBA)_2$ which is a conventional liquid crystalline compound has also been studied by infrared spectroscopy to compare its spectral features with those of the 1:1 complex from 6OBA and 3OSz.

2. Experimental

2.1. Preparation of the hydrogen-bonded complex

The preparation and characterization of *trans*-4-propoxy-4'-stilbazole 3OSz, that functions as a H-bond acceptor have been reported elsewhere [10, 11].

The hydrogen-bonded complex was prepared by an evaporation technique from pyridine solution containing an equimolar amount of H-bonding donor and acceptor moieties followed by drying *in vacuo* at 60° C.

2.2. Characterization

DSC measurements were conducted on a Mettler DSC 30. A heating rate of 10° C min⁻¹ was used to determine phase transition temperatures. Onset points of the endothermic peaks were taken as the transition temperatures. A polarizing microscope (Olympus BH 2) equipped with a Mettler FP 82 HT hot stage was used for visual observation. Once melted samples were used for the measurements.

2.3. FT-IR measurements

Infrared spectra at various temperatures were obtained by using a JASCO Micro FT/IR-200 spectrometer equipped with a Mettler FP 84 hot stage at a heating rate of $10^{\circ}C \text{ min}^{-1}$ from 70 to $168^{\circ}C$. The resolution was 4 cm^{-1} . The sample sandwiched by two KBr crystal plates was pressed to prepare the disk for the measurement. The thickness of the KBr disk was less than 1 mm. Samples heated to $20^{\circ}C$ higher than the melting point and then cooled to room temperature were used for the measurements.

3. Results and discussion

The hydrogen-bonded 1:1 complex prepared from an equimolar mixture of 4-hexyloxybenzoic acid 6OBA and *trans*-4-propoxy-4'-stilbazole 3OSz was examined by infrared spectroscopy at various temperatures. The equimolar complex of 6OBA and 3OSz will be denoted as 6OBA–3OSz. The structure of complex 6OBA–3OSz is shown in figure 1. In this structure, 6OBA acting as the H-bond donor and 3OSz functioning as the H-bond acceptor form a well-defined structure of the mesogen through the single hydrogen bond. Table 1 summarizes the transition temperatures of 6OBA–3OSz and each of the H-bond donor and acceptor moieties. The complex, 6OBA–3OSz, behaves as a single component and shows clear phase transitions and homogeneous mesophases. After the sharp melting transition at 102° C, a smectic phase is observed in contrast to $(6OBA)_2$ which exhibits only a nematic phase, and 3OSz, which is non-mesogenic. A homeotropic and focal-conic for texture characteristic of a smectic A phase is observed under microscopic observation. The nematic–isotropic transition is clearly seen at 155°C. These results confirm the formation of the complex shown in figure 1.

Infrared spectra of 6OBA-3OSz and $(6OBA)_2$ in the range of $1500-3400 \text{ cm}^{-1}$ at room temperature are shown in figures 2(*a*) and (*b*), respectively. The O-H bands at 2500 and 1920 cm⁻¹ indicative of strong hydrogen bonding between the carboxylic acid and the pyridyl unit [1-10, 12-14] are observed in figure 2(*a*). The broad O-H band centred at 3000 cm^{-1} is seen in the spectrum of $(6OBA)_2$ (see figure 2(*b*)). The bands at 2560 and 2670 cm⁻¹ for $(6OBA)_2$ are considered to be Fermi resonances [15, 16]. The carbonyl band at 1682 cm^{-1} due to the dimer formation of the carboxylic acids (see figure 2(*b*) is replaced by the C=O band at 1697 cm^{-1} when 6OBA-3OSz is formed (see figure 2(*a*)). The 1697 cm^{-1} band is attributable to the 'free' carbonyl group due to the formation of the H-bonded complex between the O-H group of the carboxylic acid and the pyridyl group. It should be noted that the intensity of the C=O bands increases after the samples are melted.



Figure 1. The structure of the hydrogen-bonded mesogenic complex of 6OBA and 3OSz.

 Table 1. Transition temperatures of the H-bonded complex 6OBA-3OSz and H-bonding donor and acceptor molecules on heating.

	Phase transition behaviour						
(6OBA) ₂ 6OBA–3OSz 3OSz	C C C	106°C 102°C 110°C	N S _A I	153°C 130°C	I N	155°C	I



Figure 2. Infrared spectra of 6OBA-3OSz(a) and $(6OBA)_2(b)$ in the range of 1400-3400 cm⁻¹.

Infrared spectra have been recorded for the complex of 6OBA-3OSz while increasing the temperature from the crystalline phase to the isotropic phase to examine the stability of the hydrogen bond of the mesogenic complex. Figure 3 shows the infrared spectra of 6OBA-3OSz in the range of $1500-1800 \text{ cm}^{-1}$ from 70 to 168° C.

The carbonyl band of 6OBA-3OSz shows significant changes at the melting and isotropization transitions. In particular, the carbonyl band becomes broader with a shift to higher frequencies once the isotropization temperature at 155°C is reached. Curve fitting has been conducted at 92°C (crystalline), 140°C (nematic), and 165°C (isotropic). The results are shown in figure 4 and table 2. In the crystalline state, the carbonyl band is separated into two bands at 1693 and 1705 cm⁻¹, suggesting the existence of two states of the carbonyl group involving the hydrogen bond between the carboxylic acid and pyridyl group. In the nematic state, the fraction of the area of the 1705 cm^{-1} band increases. It was reported that for a 1:1 complex of pyridine with a benzoic acid with pKa \sim 4, the hydrogen bond is an un-ionized type which has double minimum energy potential [13]. The present results may show that an un-ionized hydrogen bond with double minimum potential is formed between 6OBA and 3OSz. In the isotropic state (see figure 4(c)), the carbonyl absorption is separated into three bands. The bands at 1687 and 1703 cm^{-1} are due to the hydrogen-bonded complex of 6OBA-3OSz. The band at 1722 cm^{-1} indicates the existence of monomeric 6OBA. The C=O band of monomeric carboxylic acid is observed at a higher frequency than those of



Figure 3. Infrared spectra of 6OBA-3OSz in the range of $1500-1800 \text{ cm}^{-1}$ from 70 to 168°C .



Figure 4. Curve fitting results of the carbonyl bands of 6OBA–3OSz at $92^{\circ}C(a)$, $140^{\circ}C(b)$, and $165^{\circ}C(c)$.

	C=O bands of 6OBA-3OSz		C=O band of monomeric 6OBA		
	λ/cm^{-1}	Fraction of area	λ/cm^{-1}	Fraction of area	
Crystalline (92°C)	1705 1693	0·13 0·87			
Nematic (140°C)	1705 1687	0·50 0·50			
Isotropic (162°C)	1702 1687	0·59 0·23	1723	0.18	

Table 2. Curve fitting results of the carbonyl peak of 6OBA-3OSz for various phases.



Figure 5. Infrared spectra of $(6OBA)_2$ in the range of $1500-1800 \text{ cm}^{-1}$ from 70 to 168°C .

hydrogen-bonded carboxylic acids [15, 16]. These results show that the stability of the hydrogen bond decreases greatly once the temperature reaches the isotropization temperature, while the amount of free 6OBA increases.

Infrared spectra have also been recorded for the dimer of 4-hexyloxybenzoic acid, $(6OBA)_2$ on heating, as shown in figure 5. A significant effect of molecular ordering on the stability of the H-bond is also observed for the dimer of the benzoic acid. The carbonyl band at 1630 cm^{-1} due to monomeric 6OBA appears after the crystal-nematic transition at 106° C, whereas no peak due to monomeric 6OBA is observed for 6OBA-3OSz after the crystal-mesophase transition occurs (see figure 4(c)). This difference suggests that the double H-bond between two benzoic acid moieties is weaker than the single H-bond between the carboxylic acid and pyridine fragments

disussed earlier in this study. The abrupt increase in the intensity of the C=O band at $1630 \,\mathrm{cm}^{-1}$ is observed at the nematic-isotropic transition at $153^{\circ}\mathrm{C}$, showing the stability of the hydrogen bonds decreases significantly in the disordered phase.

The present results may provide useful information to design functional molecular complexes through intermolecular hydrogen bonding. Further infrared studies on a variety of H-bonded mesogenic complexes are now in progress.

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