Tubular and lamellar hydrogen-bonding molecular assemblies of isophthalic acid derivatives bearing a $-\text{CONHC}_n\text{H}_{2n+1}$ chain†

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Isophthalic acid derivatives (CnIP), bearing alkylamide chains ($-\text{CONHC}_n\text{H}_{2n+1}$; $n = 6, 10, 14, \text{and} 18$) at the 5-position that can participate in hydrogen bonding, were prepared and evaluated for their hydrogen-bonding molecular assembly structures for organogelation and liquid crystal formation. The hydrogen-bonding carboxylic acid ($-\text{COOH}$) groups form a ring-shaped [CnIP]∞ hexamer or a one-dimensional (1D) zig-zag ([CnIP]m) chain. Although neither organogelation nor liquid crystal formation was observed in the isophthalic acid derivative bearing an alkoxy ($-\text{OC}_{14}\text{H}_{29}$) chain, C14IP and C18IP derivatives could form both organogel and liquid crystal states through intermolecular N–H⋅⋅⋅O = amide-type hydrogen-bonding interactions. A discotic hexagonal columnar liquid crystal (Colh) phase was observed in hydrated [C14IP]6–(H2O)2+n and [C18IP]6–(H2O)2+n, whereas a lamella-type liquid crystal (Lα) phase was confirmed in the unhydrated C18IP. In the Colh phase, O–H⋅⋅⋅O hydrogen-bonding ring-shaped ([C14IP]6) and [C18IP]6 hexamers assembled to form the tubular molecular assembly stabilized by intermolecular N–H⋅⋅⋅O = hydrogen-bonding interactions along the tube growth direction, where H2O molecules were contained within the hydrophilic space. On the other hand, the N–H⋅⋅⋅O = hydrogen-bonding interactions between the 1D zig-zag ([CnIP]m) chains formed a layer-type molecular assembly of the Lα-phase in the absence of water molecules.

Introduction

Amide-type N–H⋅⋅⋅O = hydrogen-bonding in polypeptide chains plays an important role in forming the secondary protein structures of $\alpha$-helices and $\beta$-sheets, which are further assembled to form ternary protein structures through weak van der Waals interactions.1 The bonding energy of the amide-type hydrogen-bonding interaction is approximately 10 kJ mol$^{-1}$, a magnitude favourable for achieving association/dissociation of secondary protein structures at approximately room temperature ($k_B T = 2.5$ kJ mol$^{-1}$).2,3 Therefore, amide-type N–H⋅⋅⋅O = hydrogen-bonding interaction is a key intermolecular interaction that controls hierarchical molecular assembly structures. In the field of supramolecular chemistry, directional hydrogen bonding has been used for the construction of various anisotropic molecular assemblies from one-dimensional (1D) chains of phthalic or isophthalic acids,4,8 two-dimensional (2D) layers of squaric or trimesic acids,9,10 and three-dimensional (3D) diamond-like networks of tetracarboxylic acid derivatives.11–13 Among a variety of directional hydrogen-bonding interactions such as O–H⋅⋅⋅O, N–H⋅⋅⋅O, and N–H⋅⋅⋅N, amide-type N–H⋅⋅⋅O = hydrogen-bonding has been used to design single crystal organic ferroelectrics14–15 and discotic hexagonal columnar liquid crystal (Colh) phases of benzene derivatives bearing multiple alkyamide ($-\text{CONHC}_n\text{H}_{2n+1}$) chains.16–22 For instance, ferroelectric polarization ($P$)–electric field ($E$) hysteresis curves have been observed for $N_N',N_N''$-trialkyl-1,3,5-benzenetricarboxamide (3BC) derivatives; a response is achieved by collective inversion of the 1D N–H⋅⋅⋅O = hydrogen-bonding interactions along the $\pi$-stacking column.16–18 The application of the opposite $E$-value along the hydrogen-bonding column inverts the direction of the macro dipole moment arising from the N–H⋅⋅⋅O = chains via rotation of the $-\text{CONHC}_n\text{H}_{2n+1}$ groups in Colh phase. In addition, derivatives of $N_N',N_N''$-dinitredecyl-1,4-benzenedicarboxamide (2BC) and $N_N',N_N'',N''_N'''$-pentatetradecyl-1,2,3,4,5-benzenepentacarboxamide (5BC) were used to prepare ferroelectrics.22 Interestingly, a Colh liquid crystalline material composed of $N_N',N_N'',N''_N'''$-tetratetradecyl-1,3,6,8-pyrenetetracarboxamide also formed a ferroelectric material with fluorescent and current switching properties.22,24 The 1D hydrogen-bonding molecular assembly and its dynamic behaviour are essential to the ferroelectric behaviour of the $\pi$-molecular system bearing multiple $-\text{CONHC}_n\text{H}_{2n+1}$ chains. Improved design of amide-type hydrogen-bonding has the potential to form low-
dimensional hierarchical molecular assemblies such as rings, tubes, chains, and layer structures.

Additional hydrogen-bonding sites such as –OH, –NH, and –COOH are usually introduced into molecular structures to form the low-dimensional molecular assemblies.\(^{25-29}\) Well-known hydrogen-bonding simple benzene carboxylic acid derivatives of trimesic acid can form two types of O–H⋯O = hydrogen-bonding molecular assemblies: a six-fold 2D hexagonal layer and an infinite 1D zig-zag chain, which can be controlled by crystallization conditions. On the other hand, terephthalic and isophthalic acids typically form 1D linear and 1D zig-zag type O–H⋯O = hydrogen-bonding molecular assemblies, respectively.\(^{26-29}\) It can be reasonably expected that these benzene carboxylic acid derivatives can act as effective platforms to construct various molecular assemblies in a flexible and rational way by introducing additional hydrogen-bonding group. For instance, an interesting ring-shaped molecular assembly was reported using an isophthalic acid (IP) derivative bearing a hydrophobic –OC\(_{n}\)H\(_{2n+1}\) chain at the 5-position, which formed a hydrogen-bonding O–H⋯O = hexamer structure at \(n<12\), and each hexamer ring was isolated in the absence of interactions between the hexamers.\(^{26-29}\) In contrast, 1D zig-zag hydrogen-bonding structures have been observed in long alkyl chain compounds with \(n>12\), where the much longer alkyl chains enhanced hydrophobic interactions and stabilized the interdigitate lamellar-type molecular assembly structure. Although interesting ring-shaped hydrogen-bonding hexamer assemblies have been obtained by introducing alkoxy group into IP derivative, additional introduction of amide-type N–H⋯O = hydrogen-bonding interaction at –CONHC\(_{n}\)H\(_{2n+1}\) chain has a potential to form various kinds of low-dimensional hierarchical molecular assembly structures.

Herein, we designed a hydrogen-bonding IP derivative bearing an additional hydrogen-bonding –CONHC\(_{n}\)H\(_{2n+1}\) chain to achieve low-dimensional molecular assemblies, which will fabricate different types of molecular assemblies from –OC\(_{n}\)H\(_{2n+1}\) substituted IP derivative. Four kinds of amphiphilic IP derivatives bearing a different alkyl chain length of –CONHC\(_{n}\)H\(_{2n+1}\), C\(_{6}\)IP \((n=6)\), C\(_{10}\)IP \((n=10)\), C\(_{14}\)IP \((n=14)\), and C\(_{18}\)IP \((n=18)\), were synthesized and corresponding molecular assembly behaviours were studied systematically. Different from the IP derivative with an –OC\(_{n}\)H\(_{2n+1}\) chain, the hydrogen-bonding –CONHC\(_{n}\)H\(_{2n+1}\) chain in CnIP can provide additional N–H⋯O = hydrogen-bonding interactions to form high-order molecular assembly structures (Scheme 1). The O–H⋯O = hydrogen-bonding ring-hexamers are connected by six amide-type N–H⋯O = hydrogen-bonding interactions along the \(\pi\)-stacking direction of the hexamer, resulting in a tubular hierarchical molecular assembly with an inner pore diameter of \(~1.1\) nm. Since the outer surface and inner pore of the tubular assembly are hydrophobic and hydrophilic, respectively, the inner hydrophilic pore can capture hydrophilic species such as H\(_2\)O and various ions. Another possible molecular assembly is a zig-zag type O–H⋯O hydrogen-bonding 1D assembly, which are connected by amide-type N–H⋯O = hydrogen-bonding interactions along the direction normal to the 1D chain. These chains are formed in a similar manner as the lamellar-type 2D molecular assembly structure with alternating arrangement of the hydrogen-bonding layer and hydrophobic alkyl chains. Accordingly, both organogel and liquid crystal states were formed in the assembly system via additional intermolecular N–H⋯O = amide-type hydrogen-bonding interactions in a prime IP molecular core. Moreover, the liquid crystalline phase was modulated by the gelation process of C18IP. Pure unhydrated state formed the lamellar type liquid crystalline phase, whereas the xerogel state of (C18IP\(_{n}\)–(H\(_2\)O)\(_{x}\)) from C\(_{2}\)H\(_{3}\)OH–H\(_2\)O formed the tubular structure and hexagonal columnar liquid crystalline phase. The gelation ability was directly associated with the formation of lamellar and/or hexagonal columnar phases. The thermal stability, organogelation ability, liquid crystal formation, phase transition behaviour, and ion inclusion properties of the prepared assemblies were systematically examined.

**Experimental section**

**Preparation of CnIP**

Commercially available reagents were used without further purification, and dry triethylamine was obtained by distillation from KOH. The preparation of CnIP \((n=6, 10, 14,\) and 18) was performed by a 4-step procedure starting from 1,3,5-benzenetricarboxylic acid trimethyl ester (Scheme 2). Benzene-1,3,5-tricarboxylic acid dimethyl ester was prepared following the literature.\(^{33}\) Benzene-1,3,5-tricarboxylic acid trimethyl ester (5.00 g, 7.93 mmol) and NaOH (790 mg, 19.8 mmol) were dissolved in CH\(_3\)OH (175 mL) and refluxed for 12 h. The reaction mixture was diluted with water (500 mL), and washed with...
diethyl ether (3 × 250 mL). The aqueous phase was acidified to pH ~ 1 with 5% hydrochloric acid and extracted with diethyl ether (3 × 250 mL). The extracted organic layers were dried over MgSO₄ and concentrated under vacuum to afford benzene-1,3,5-tricarboxylic acid dimethyl ester (3.83 g) with a yield of 81%. ¹H NMR (400 MHz, DMSO-d₆): δ 3.93 (s, 6H, C₆H₁₂O₂), 6.21 (br s, 1H, NH), 8.61 (d, J = 1.6 Hz, 2H, ArH), 8.79 (t, J = 1.6 Hz, 1H, ArH).

C6IP

Benzene-1,3,5-tricarboxylic acid dimethyl ester (2.10 g, 8.81 mmol) in SOCl₂ (80 mL, 1.10 mol) was refluxed for 20 h. After cooling the reaction mixture to room temperature, the excess SOCl₂ was removed under vacuum. The resultant acid chloride was dissolved in dry CH₂Cl₂ (40 mL), and hexylamine (1.32 mL, 10.11 mmol) was slowly added dropwise over 10 min followed by the addition of triethylamine (1.28 mL, 9.24 mmol). The mixture was subsequently stirred for 18 h at room temperature. The resultant solution was washed 3 times with aqueous 1 M HCl and brine, and then dried over Na₂SO₄. Evaporation of CH₂Cl₂ yielded a white precipitate, which was recrystallized from hexane: AcOEt (2:5). A total of 1.52 g of 1 was obtained with a yield of 56%. ¹H NMR (400 MHz, CDCl₃): δ 0.90 (t, J = 7.1 Hz, 3H, CH₃), 1.30–1.45 (m, 6H, alkyl), 1.62–1.69 (m, 2H, –NHCH₂CH₂), 3.45–3.52 (m, 2H, –NHCH₂CH₂), 3.98 (s, 6H, 2Ar–COOCH₃), 6.26 (s, 1H, NH), 8.61 (d, J = 1.6 Hz, 2H, ArH), 8.80 (t, J = 1.6 Hz, 1H, ArH).

A solution of C6IP (1.52 g, 4.73 mmol) in CH₂OH (25 mL) was added to a solution of KOH (2.91 g, 51.8 mmol) in water (20 mL), and the resulting solution was refluxed for 2 h. After evaporation of CH₂OH under vacuum, the reaction mixture was diluted with H₂O (300 mL) and aqueous HCl (1 M) until the pH reached 1. The resulting suspension was extracted by AcOEt and dried over Na₂SO₄. Removal of solvent under vacuum afforded a crude product, which was recrystallized twice from CH₃OH: H₂O (4:1) to give C6IP (880 mg) with a yield of 63%. Mp = 279–280 °C. ¹H NMR (400 MHz, DMSO-d₆): δ 0.87 (t, J = 6.8 Hz, 3H, –CH₃), 1.23–1.36 (m, 6H, alkyl), 1.48–1.58 (m, 2H, –NHCH₂CH₂), 3.23–3.31 (m, 2H, –NHCH₂), 8.56 (t, J = 1.6 Hz, 1H, ArH), 8.63 (d, J = 1.6 Hz, 2H, ArH), 8.88 (t, J = 5.6 Hz, 1H, –NH), 13.50 (s, 2H, –(C=O)=OH). IR: ν (cm⁻¹): 3287 (N–H stretch), 1688 (C=O of COOH), 1634 (C=O of CONH), 1544 (C=N). Elemental analysis: calculated for C₁₉H₂₈NO₅: C, 65.31; H, 8.70; N, 3.45, found: C, 65.28; H, 7.91; N, 4.00. HRMS (FAB): calculated for C₁₉H₂₈NO₅: m/z 350.1967 [M + H]+, found m/z 350.1972.

C14IP

Yield 646 mg, 64%. Mp: decomposed at above 241 °C. ¹H NMR (400 MHz, DMSO-d₆): δ 0.85 (t, J = 6.8 Hz, 3H, CH₃), 1.18–1.35 (m, 22H, alkyl), 1.48–1.58 (m, 2H, CONHCH₂CH₂), 3.29 (m, 2H, CONHCH₂), 8.56 (t, J = 1.6 Hz, 1H, ArH), 8.62 (s, 2H, –(C=O)=OH), 8.85 (br t, J = 4.8 Hz, 1H, NH), 13.47 (br s, 2H, –(C=O)=OH). IR: ν (cm⁻¹): 3300 (N–H stretch), 1720 and 1695 (C=O of COOH, C=O stretching vibration of the free, non-hydrogen bonded and laterally hydrogen-bonded COOH groups, respectively), 1640 (C=O of CONH), 1544 (C=N). Elemental analysis: calculated for C₁₉H₂₈NO₅: C, 65.31; H, 7.79; N, 4.01. Found: C, 65.28; H, 7.91; N, 4.00. HRMS (FAB): calculated for C₁₉H₂₈NO₅: m/z 350.1967 [M + H]+, found m/z 350.2594.
Preparation of organogels

Temperature and solvent dependent gelation behaviors were examined following in the literature. A series of commonly used solvents such as CH₂OH, C₆H₅OH, acetone, THF, CH₃CN, AcOEt, toluene, and H₂O was screened for the organogel formation of CnIP. The gelation behavior was not confirmed in each pure solvent. After the screening in the mixed solvent system, the organogels of CnIP were observed in C₆H₅OH–H₂O. CnIP in hot C₆H₅OH–H₂O with concentration of 10 mM was gradually cooled down to room temperature, and the minimum volume percentage of H₂O to induce the organogels at 10 mM was summarized in Table 1.

Physical measurements

Infrared spectroscopy (IR; Thermo Fisher Scientific Nicolet 6700, 400–4000 cm⁻¹) measurements were conducted with a resolution of 4 cm⁻¹ using KBr pellets. Thermogravimetry-differential thermal analyses (TG-DTA) were conducted using a thermal analysis station (Rigaku Thermo plus TGS8120) with Al₂O₃ as a reference from 293 to 600 K with a heating rate of 5 K min⁻¹ under a nitrogen atmosphere. Temperature-dependent powder pattern X-ray diffraction (XRD) data were collected using a diffractometer (Rigaku Rint-Ultima III) with Cu Kα (λ = 1.54187 Å) radiation. Scanning electron microscopy (SEM; JEOL JSEM-5400F) and atomic force microscopy (AFM; JEOL JSPM-5200) were performed on highly ordered pyrolytic graphite (HOPG) and micro substrates, respectively. Acceleration voltages of 5 or 10 kV under a vacuum of less than 10⁻⁵ Pa was used for the SEM measurements. Commercially available Si cantilevers with a force constant of 4.5 N m⁻¹ were used for the AFM measurements.

Table 1 Formation of OG and LC states of CnIP derivatives at n = 6, 10, 14, and 18

<table>
<thead>
<tr>
<th>C6IP</th>
<th>C10IP</th>
<th>C14IP</th>
<th>C18IP</th>
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<tr>
<td>Formation of OG**</td>
<td>—</td>
<td>O*</td>
<td>O*</td>
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<tr>
<td>H₂O (v/v%) in 10 mM solutionb</td>
<td>—</td>
<td>50</td>
<td>40</td>
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<td>Xerogelsc</td>
<td>—</td>
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<td>—</td>
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<tr>
<td>H₂O (wt%) of xerogel</td>
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<td>—</td>
<td>3.7–7.2%</td>
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<tr>
<td>H₂O (wt%) of crystal</td>
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<td>0.0</td>
<td>0.0</td>
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<tr>
<td>Type of LC phase</td>
<td>—</td>
<td>C₀₁₀</td>
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**OG formation was evaluated in a 10 mM solution of CnIP in C₆H₅OH–H₂O (v/v = volume percentage of H₂O) at 300 K. bVolume percentage of H₂O for OG formation of CnIP at ca. fixed concentration of 10 mM in C₆H₅OH. cXerogels were obtained by vacuum evaporation of the OG and the weight percentage of H₂O was determined by TG analyses. dLC phases of C₀₁₀ and Lₙ were discotic hexagonal columnar and lamella phases, respectively. Solvent loss of the C₁₀IP xerogel was not obtained in the vacuum drying process at room temperature, and OG state was transformed to a crystalline solid. eThe notation of O* represented the formation of OG or xerogel states at a fixed concentration of 10 mM in C₆H₅OH–H₂O. fOG states were not observed in C₆H₅OH–H₂O at v/v range from 100/0 to 0/100.

Results and discussion

Formation of organogels

The CnIP molecules (n = 6, 10, 14, and 18) possess two kinds of hydrogen-bonding sites: two hydrophilic –COOH groups and one hydrophobic –CONH(CH₂)ₓCOOH chain, resulting in amphiphilic properties. The hydrophobic character of the CnIP derivatives was enhanced with increasing alkyl chain length (n) from C₆IP, C₁₀IP, C₁₄IP, to C₁₈IP, which affected the molecular assembly structures of the organogel (OG) and liquid crystal (LC). The CnIP molecules were soluble in CH₂OH, C₆H₅OH, acetone, and THF, whereas only slightly soluble in H₂O, CH₃CN, AcOEt, and toluene. Among the four CnIP derivatives, C₁₀IP, C₁₄IP, and C₁₈IP can form OGs in the mixed solvent system of C₆H₅OH (CH₃OH)–H₂O (Fig. 1a) at a fixed concentration of 10 mM. Table 1 summarizes the formation of OGs in C₆H₅OH–H₂O and LCs of the CnIP derivatives. C₆IP with the shortest alkyl chain did not form the OG or LC state due to insufficient hydrophobicity. The formation of an OG is closely related to the formation of 1D fibrous molecular assemblies and their 3D entanglements.

Therefore, the formation of 1D hydrogen-bonding tubular assembly was consistent with the formation of an OG. Among the poor solvents, H₂O was essential for the formation of the OG of C₁₀IP, C₁₄IP, and C₁₈IP, and C₆H₅OH and/or CH₃OH were also needed. Although we tried to prepare OG state from a variety of solvent such as THF, CH₃CN, AcOEt, and toluene, the addition of H₂O in C₆H₅OH achieved the corresponding organogel states. The thermal stability of the OG decreased in the order of C₁₈IP, C₁₄IP, to C₁₀IP, and crystalline powder coexisted with the OG in the shorter chain C₁₀IP molecule. Therefore, crystallinity was increased by decreasing the alkyl chain length. The solubility of CnIP in C₂H₅OH decreased in the order of C₆IP, C₁₀IP, C₁₄IP, to C₁₈IP, and the minimum H₂O content required for the formation of the respective OGs also decreased in the same order (Table 1 and Fig. S1†). The volume percentages of H₂O to form stable OGs at 20, 10, 5, and 2.5 mM solutions of C₁₄IP in C₂H₅OH were approximately 20, 30, 40, and 50%, respectively.

Hydrated xerogels and unhydrated crystals

The formation of an OG was not observed in the shortest alkyl chain C₆IP derivative, whereas the OG state of C₁₀IP was unstable at 300 K due to the coexistence of OG and crystalline...
solid (CS) states. Both C14IP and C18IP derivatives can form stable and uniform OG states in C2H5OH–H2O mixed solvents in the absence of CS at approximately 300 K. Two kinds of molecular assembly states of the xerogel (XG) and CS were obtained for C14IP and C18IP (Fig. S2†) by changing the solvent system. The XG state was obtained by vacuum drying from the OG state in C2H5OH–H2O, whereas the CS was obtained by recrystallization from CH3OH, C2H5OH, or aceton in the absence of H2O. The presence of H2O was a necessary condition to form both the OG and XG states. Fig. 1a shows a cloudy OG state of C14IP in C2H5OH–H2O at a fixed concentration of 10 mM. The microscale morphologies of the CS and XG states of C18IP on silicon differed significantly in the SEM images. Fig. 1b and c show the SEM images of the CS and XG states of C18IP. The morphology of the CS state has a clear edge with a flat crystal surface (Fig. 1b), whereas the 3D entanglement network of each microfiber was observed in the XG state of C18IP (Fig. 1c). The latter 3D entangled microstructure was a prerequisite for the formation of the OG state. Differences in the microscale morphologies of the XG and CS are associated with microscale hydrogen-bonding interactions.

The results of the TGA of the XG and CS states of C14IP and C18IP differed to a significant degree (Fig. 2a). Although the CS state exhibited high thermal stability up to 520 K, as indicated by the absence of weight-loss, the XG state showed weight-loss at approximately 400 K corresponding to the desorption of H2O molecules. The magnitude of the weight-losses for the XG states of C14IP and C18IP at 400 K were 3.7–7.2 and 3.3–3.7%, respectively, corresponding to 5 ≤ n ≤ 12 (Fig. S3†). It should be noted that the amount of H2O molecules may be influenced by environmental humidity during the vacuum drying process, accounting for the deviation in H2O content. The presence of H2O is a requirement for the formation of the OG state, where the hydrophilic H2O molecules stabilize the 1D fibrous molecular assemblies. Since the microfibrous morphology of the XG state was maintained even under vacuum during the SEM measurements, it can be concluded that the 1D fibrous molecular assemblies of C14IP and C18IP were stabilized by intermolecular hydrogen-bonding interactions.

**Liquid crystalline property**

The LC phases were not observed in the short alkyl chain compounds of C6IP and C10IP with direct decomposition at approximately 550 K (Fig. S4†). In contrast, both fluidic and birefringent behaviours of C14IP and C18IP were observed in the polarized optical microscopy (POM) images under the cross-Nicol optical arrangement by increasing the temperature to 420 K.

The XG and CS states of C14IP and C18IP showed different phase transition behaviours in the DSC diagrams. The endothermic peaks of the XG state of C14IP and C18IP were observed at approximately 270 and 300 K during the heating process, which corresponded to H2O melting within the molecular assembly and were also observed in the exothermic peaks during cooling (Fig. 2b). However, the phase transition from the LC to the isotropic liquid (IL) phase was not observed in the DSC of C14IP and C18IP due to decomposition at approximately 540 K. Similar phase transition behaviour was observed in the XG state of C14IP and C18IP, where a reversible S–LC phase transition peak was observed at approximately 410 K (red DSC chart in Fig. 2b). Although the phase transition behaviour of unhydrated C14IP was fundamentally similar to that of the hydrated C14IP, the CS state of unhydrated C18IP exhibited a reversible S–LC phase transition peak at approximately 410 K (blue DSC chart in Fig. 2b). The XG and CS states of C18IP and C18IP formed different LC phases from different molecular assembly structures. Fig. 2c shows the POM images of the two mesophases derived from the XG and CS states of C18IP and C18IP at 450 K. Although both POM textures resembled each other, a striped pattern was observed in the texture of the CS state. The molecular assembly structure within the LC phase is affected by the alkyl chain length (n) of the –CONHC6H2n+1 group.

To identify the molecular assembly structures of the LC phases, the XRD patterns of the two types of LC phases of C14IP and C18IP were compared. Fig. 3 summarizes the XRD patterns of the LC phases of C14IP and C18IP. Typical XRD patterns of C14IP and C18IP at 450 K were consistent with the diffraction pattern of the discotic hexagonal columnar (Colh) LC phase with a d100 = 3.62 nm, where the diffraction peaks at 2θ = 24.4, 42.7, 49.1, and 56° were consistent with the index values of $d_{100}$, $d_{110}$, $d_{200}$, and $d_{210}$, respectively. In the large angle region, two broad diffraction peaks around 2θ = 20 and 25° could be assigned to the melting state of the alkyl chains and...
average interplanar distance \(d_{001}\) along the \(\pi\)-stacking direction within a column. Almost identical XRD patterns were observed in LC phase of \((C18IP)_6(H_2O)_n\) at 480 K, which was also consistent with the \(Col_h\) phase of \((C14IP)_6(H_2O)_n\). A periodicity of \(d_{100} = 3.84\) nm for \((C18IP)_6(H_2O)_n\), was approximately 0.22 nm longer than that of \((C14IP)_6(H_2O)_n\) (\(d_{100} = 3.62\) nm). The \(d_{100}\) spacing for the \(Col_h\) phases of \((C14IP)_6(H_2O)_n\) and \((C18IP)_6(H_2O)_n\) was approximately 40 and 45% smaller than the maximum molecular lengths of 5.98 and 6.98 nm for the \((C14IP)_6\) and \((C18IP)_6\) hexamers, respectively, assuming an all-trans \(-CONHC_2H_{2n+1}\) conformation, consistent with the melting state of the alkyl chains in the \(Col_h\) phase. It is worth to mention that the three-dimensional molecular assemblies of \(CnIP\) is observed even in fluid high temperature LC phase. Similar molecular assemblies of alkoxysapphylic acid derivative in the absence of additional hydrogen-bonding \(-CONHC_2H_{2n+1}\) unit did not form the mesophase by a direct solid-isotropic liquid phase transition,\(^{31}\) which were completely different from the phase transition behaviors of \(C14IP\) and \(C18IP\). This difference is clearly accounted for the role of amide-type hydrogen-bonding interaction, which significantly connects each two-dimension hexameric ring structures to form tubular assembly through the strong amide-type hydrogen-bonding interaction of \(C14IP\) and \(C18IP\). As a result, the corresponding tubular molecular assemblies were stabilized to exhibit LC phase and organogel state. Interestingly, the XRD pattern of the LC phase of unhydrated \(C18IP\) was not consistent with the formation of the \(Col_h\) phase. Highly ordered diffraction peaks at \(2\theta = 4.31, 8.68, 17.45\), and \(21.91\)° at \(T = 460\) K were consistent with the index values of \(d_{100}, d_{200}, d_{300}\), and \(d_{400}\), suggesting a layered molecular assembly structure with an interlayer spacing of \(d_{001} = 2.05\) nm. Therefore, the lamella (\(L_a\)) type LC phase is formed during the heating of unhydrated \(C18IP\).\(^{35-37}\) Infinite \(O-H-O = \) hydrogen-bonding chains were assembled to form a hydrogen-bonding 2D sheet through the interchain \(N-H-O = \) hydrogen-bonding interactions. The interlayer spacing of \(d_{100} = 2.05\) nm in the \(L_a\) phase was 0.74 nm shorter than the maximum length of \(C18IP\) assuming an all-trans conformation of \(-CONHC_4H_{37}\).

The \(H_2O\) molecules play a key role in determining the type of LC phase (\(Col_h\) and/or \(L_a\)). It should be noted that the hydrated \((C14IP)_6(H_2O)_n\) and \((C18IP)_6(H_2O)_n\) primarily formed the \(Col_h\) phase and unhydrated \(C18IP\) adopted the \(L_a\) phase. In the vibrational spectra (Fig. S6 and S7†), both the \(-OH\) and \(-NH\) stretching energies of \(v_0^\text{a\_h}\) of the \(-COOH\) groups and of \(v_0^\text{a\_h}\) of \(-CONH-\) for different molecular assemblies of the \(XG\) and \(CS\) states were observed at 3080 and 3302 \(cm^{-1}\), respectively. This suggests the formation of intermolecular \(O-H-O = \) and \(N-H-O = \) hydrogen-bonding interactions. In hydrated \((C18IP)_6(H_2O)_n\) amide-I and -II bands were observed at 1637 and 1599 \(cm^{-1}\), respectively, indicating the formation of intermolecular \(N-H-O = \) hydrogen-bonding interactions.\(^{39}\) On the other hand, the amide-I and -II bands of unhydrated \(C18IP\) were observed at 1616 and 1582 \(cm^{-1}\), respectively. Since the formation of strong hydrogen-bonding interactions usually results in a red-shift of the vibrational band, it can be concluded that the strength of hydrogen-bonding interaction of unhydrated \(C18IP\) was stronger than that of the hydrated \((C18IP)_6(H_2O)_n\). This corresponds to a much denser packing structure of the former state due to its high crystallinity. There was no significant difference in the vibrational spectra of the unhydrated \(C14IP\) and hydrated \((C14IP)_6(H_2O)_n\) (Fig. S7†), which formed a similar packing structure in both LC phases.

The most widely accepted molecular assembly structure of \((C18IP)_6(H_2O)_n\) in the \(Col_h\) phase is a tubular ring-type structure, where the \(O-H-O = \) hydrogen-bonding hexamer-rings of \((C18IP)_6\) are assembled to form a \(\pi\)-stacking tubular structure through intermolecular \(N-H-O = \) hydrogen-bonding interactions along the tube. The hydrophilic pore with a diameter of \(~1.1\) nm was observed on the inner side of the ring-shaped \((C18IP)_6\) hexamer, which was filled by 5–12 hydrophilic \(H_2O\) molecules in the \(XG\) state. Assuming the \(\pi\)-stacking distance of \((C18IP)_6\) is \(d_{001} = 3.8\) \(Å\) in the \(Col_h\) phase and \(H_2O\) molecular volume of \(~30\) \(Å^3\), the hydrophilic volume available inside the pore is \(361\) \(Å^3\) per \((C18IP)_6\) unit, consistent with the occupied volume of \(360\) \(Å^3\) for the \(12\) \(H_2O\) molecules. The presence of \(H_2O\) inside the tubular pore was also consistent with the TG and DSC analyses, formation of the \(Col_h\) phase, and XRD patterns. On the other hand, the outer surface of the tubular molecular assembly was covered by hydrophobic \(-CONHC_4H_{37}\) chains (Scheme 3).

Two types of LC phases (\(Col_h\) and \(L_a\)) were only observed in hydrated \(C18IP\). In the \(L_a\) phase, each \(C18IP\) molecule formed one zig-zag \(O-H-O = \) hydrogen-bonding chain as a primary assembly structure. Secondary intermolecular \(N-H-O = \) hydrogen-bonding interactions generated the 2D molecular assembly, where the hydrophobic \(-CONHC_4H_{37}\) chains were elongated along the direction normal to the hydrogen-bonding sheet to form a lamella-type molecular assembly in the absence of \(H_2O\).

Dielectric responses and ion inclusion

Molecular motion in the \(Col_h\) and \(L_a\) phases were evaluated by the temperature- and frequency-dependent dielectric constants. The motional freedom of polar \(H_2O\) molecules can be easily detected by dielectric spectroscopy.\(^{39}\) Fig. 4a shows the
Hydrophilic ion pairs such as MX = LiCl, NaCl, KCl, and H$_2$SO$_4$ were introduced into the inner channel of tubular molecular assembly of (C14IP)$_n$. Such channel formation based on tubular molecular assembly enables to achieve the transport property for various kinds of ions along the π-stacking and amide-type hydrogen-bonding direction.$^{36-41}$ A mixing ratio of MX : (C14IP)$_n$ = 1 : 1 can form the OG state in H$_2$O–C$_2$H$_5$OH, which was subsequently dried under vacuum to form the XG state of (C14IP)$_n$–(MX). From the elemental analyses (Table S1†), the formulæ observed were (C14IP)$_n$–(LiCl–H$_2$O)$_6$, (C14IP)$_n$–(NaCl–H$_2$O)$_6$, (C14IP)$_n$–(KCl–H$_2$O)$_6$, and (C14IP)$_n$–(H$_2$SO$_4$–H$_2$O)$_n$. The formation of hexagonal columnar structure was confirmed by XRD showing typical hydrogen-bonding tubular molecular assembly structures. Therefore, the inner hydrophilic channel of the tubular molecular assembly of (C14IP)$_n$ was filled with the corresponding hydrophilic MX species. The dielectric responses of (C14IP)$_n$–(MX) differ according to MX species. Fig. 4b summarizes the imaginary part of the dielectric constant, $\varepsilon_2$, of (C14IP)$_n$–(MX) at a constant frequency of $f = 1$ kHz in the second heating process. Since the H$_2$O molecules in the second heating process were completely removed from the tubular molecular assemblies, the intrinsic dielectric responses of the unhydrated (C14IP)$_n$–(MX) were observed in the temperature dependent $\varepsilon_2$. The $\varepsilon_2$ values corresponded to the dielectric-loss and/or conductive component in a parallel RC-circuit. The $\varepsilon_2$ values below 350 K were almost zero due to a lack of contribution from the conducting component. However, large $\varepsilon_2$ differences were observed above 400 K according to the MX species present in the tubular channel. The $\varepsilon_2$ values of (C14IP)$_n$–(LiCl), (C14IP)$_n$–(NaCl), (C14IP)$_n$–(KCl), and (C14IP)$_n$–(H$_2$SO$_4$) at 450 K were 7.5, 13, 60, and 100, respectively. The relatively high $\varepsilon_2$ value of (C14IP)$_n$–(H$_2$SO$_4$) originated from the protonic conductivity, and the magnitude of the $\varepsilon_2$ values decreased in the order of H$_2$SO$_4$, KCl, NaCl, to LiCl. Large MX salts have a tendency to increase the $\varepsilon_2$ values and the compatibility of pore diameter ($\sim$1.1 nm) and size of the MX pair is essential in determining the $\varepsilon_2$ values.

**Conclusions**

Hydrophobic alkylation chains (–CONHC$_n$H$_{2n+1}$) were introduced as an effective intermolecular N–H····O = hydrogen-bonding unit into a O–H····O hydrogen-bonding isophthalic acid derivative (CnIP). The molecular aggregation behaviours of the prepared derivatives were examined by changing the alkyl chain length from $n$ = 6, 10, 14, to 18. The O–H····O hydrogen-bonding interactions at the two –COOH sites of the isophthalic acid derivatives could form two kinds of hydrogen-bonding structures; a 1D zig-zag chain and ring-shaped (CnIP)$_n$ hexamer according to the parameter $n$. Ring-shaped O–H····O hydrogen-bonding hexamers were obtained in the OG state of (C14IP)$_n$–(H$_2$O)$_n$ and (C18IP)$_n$–(H$_2$O)$_n$, which were further assembled to a tubular molecular assembly via interhexamer N–H····O = hydrogen-bonding. The formation of the OG in H$_2$O–C$_2$H$_5$OH was consistent with the fibrous 1D molecular assembly with a tubular structure. The hydrophilic inner pores of the 1D tubular assemblies of the XG states of

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**Scheme 3** Formation of the hierarchical molecular assembly structures of C18IP to form the Colh (upper) and La phases (lower).
(C14IP)$_n$·(H$_2$O)$_n$ and (C18IP)$_n$·(H$_2$O)$_n$ were occupied by H$_2$O molecules, forming a thermotropic Col$_h$ phase upon heating. In contrast, enhanced hydrophobic interactions in C18IP formed a lamella-type molecular assembly structure in the absence of H$_2$O, where the 1D zig-zag O·H···O hydrogen-bonding chains interacted through additional N·H···O = hydrogen-bonding to form a 2D sheet assembly. No void space in the lamella phase was observed in the absence of H$_2$O. On the other hand, the 1D hydrophilic pore with a diameter of 1.1 nm in the Col$_h$ phase was suitable for H$_2$O occupation and formation of an ion pair (MX = LiCl, NaCl, KCl, and H$_2$SO$_4$) inclusion environment. Slight modification of the hydrophobic interactions in the −CONHC$_2$H$_{2n+1}$ chains was essential for the formation of the tubular or lamella type hydrogen-bonding molecular assembly structures in simple alkylamide-substituted isophthalic acid derivatives. The hydrophilic inner channel of (CnIP)$_n$ tubular assemblies can be used for molecular and ionic transport.

Conflicts of interest
There are no conflicts to declare.

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Notes and references


