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How much do coulombic interactions stabilize a mesophase? Ion pair and non-ionic binary isosteric derivatives of monocarbaborates and carboranes

Aleksandra Jankowiak, Ajan Sivaramamoorthy, Damian Pociecha, and Piotr Kaszyński

Replacement of the N⁺ atom in the pyridinium cation [Pyr] and the B⁻ atom in the monocarbaborate anion, 1[10] or 1[12], of a liquid crystalline ion pair with C atoms leads to an isoelectronic and isosteric non-ionic binary liquid crystalline mixture of carborane (2[10] or 2[12]) and benzene ([Ph]) derivatives lacking coulombic interactions. A comparison of mesogenic properties of ion pairs, 1[10]c–[Pyr]c and 1[12]c–[Pyr]c, with their analogous non-ionic mixtures, 2[10]c–[Ph]c and 2[12]c–[Ph]c, shows a 181 K higher clearing temperature, Tc, for the ion pair. This corresponds to a DFT-calculated difference in association energy ∆∆H = 24.5 kcal/mol in a typical dielectric medium (ε = 2.5). Pure compounds and binary mixtures were characterized using thermal, optical, and XRD methods.

Introduction

Ionic liquid crystals (ILC) are capable of anisotropic ion transport,1-4 and for this reason they are becoming increasingly attractive electrolytes for ion batteries,5 photovoltaic6-8 and other applications.9,10 In contrast to non-ionic mesogens, typical ILC are binary systems, comprising of a cation and an anion. Consequently, mesogenic properties of ILC are controlled not only by the anisometry of the two components, but also by electrostatic interactions.11 While the first factor is well understood,12 the effect of coulombic interactions on phase stability is poorly characterized and has only recently become the subject of theoretical considerations.13,14 Despite the large number of examples of ILC,10 there exists no single report of a molecular system that permits experimental evaluation of the coulombic component of the phase stability in the absence of other factors. We now report such a molecular system and for the first time assess the effect of coulombic interactions, as the only variable, on mesophase stability in ILC.

During the past several years, we have developed new ILC materials in which anisometric anions derived from monocarbaborates, [closo-1-CB₉H₉]⁻ ([10]) and [closo-1-CB₁₁H₁₁]⁻ ([12]), are the driving force to induce mesogenic behaviour.15-18 Due to the high delocalization of the charge and the resulting weak nucleophilicity of the boron cluster anions,19 the ion interactions in such ILC, e.g. 1a–[Pyr]a (Fig. 1),16 are non-specific and mainly electrostatic. Such ion pairs typically form SmA and soft crystalline phases, although some exhibit a nematic phase, e.g. azo derivative 1[12]b–[Pyr]a (Fig. 1).16

![Fig. 1. The structure of ions (1 and [Pyr]) and non-ions (2 and [Ph]). In the cage structures, each vertex represents a BH fragment, and the sphere is a carbon atom.](image-url)

Here, we report the synthesis and detailed thermal, optical and structural (XRD) analysis of ion pairs 1[10]–[Pyr] and 1[12]–[Pyr] as well as their electrically neutral analogues, 2 and [Ph]. We emphasize the comparison of the mesogenic behaviour of ion pairs 1–[Pyr] with the isosteric, non-ionic equimolar mixtures 2–[Ph]. The experimental results are supported with DFT modelling of the association of ions and non-ionic analogues in a dielectric medium.

Results and Discussion

Synthesis

Esters 1c and 2c were obtained by reaction of phenol 3c with appropriate acid chlorides which were prepared from carboxylic acids 4[10]–15, 4[12]–16, 5[10]–20 and 5[12] (Method A, Scheme 1). Without isolation, ion pairs 1c–[NEt4] were obtained from acid 4[10]–[NEt4]15 and 4[12]–[NEt4]16 were converted to 1c–[Pyr] by cation exchange with [Pyr]Br as described before for esters 1a–[Pyr]a and 1b–[Pyr]a.15,16 Esters 2a and 2b were prepared directly from phenols 3a and 3b, respectively, and carboxylic acid 5 in the presence of DCC (Method B).

Scheme 1. Synthesis of esters 1 and 2.1

\[ \begin{align*}
4, X & = B' \\
5, X & = C \\
6, X & = B \\
2, X & = C \\
\end{align*} \]

Reagents and conditions: i) Method A: (COCl)2, cat. DMF, CH2Cl2; 2. Et3N, CH2Cl2; or Method B: DCC, DMAP, CH2Cl2, rt, 12 hrs.

Synthesis of carboxylic acid 5[12].

\[ \text{H} \xrightarrow{i} C_6H_{13}\text{O}_2\text{COOH} \]

Reagents and conditions: i) 1. BuLi 1 eq. THF -80 °C → rt → 80 °C; 2. C6H5I 1 eq. 0 °C; 3. BuLi 1 eq. -80 °C → 0 °C; 4. CO2; 5. HCl.

12-Hexyl-p-carborane-1-carboxylic acid (5[12]) was obtained by alkylation followed by carboxylation of p-carborane (Scheme 2) as described before for the pentyl analogue.21

Pyridinium bromides [Pyr]b–Br and [Pyr]c–Br were prepared by N-alkylation of 4-(trans-4-pentylcyclohexylmethoxy)-pyridine (6) with appropriate alkyl bromide in MeCN following the procedure described for [Pyr]a–Br (Scheme 3).15 The substituted pyridine 6 was obtained from 4-chloropyridine and trans-4-pentylcyclohexylmethanol22 (7) in the presence of NaH in DMSO following a general method.23


Reagents and conditions: i) NaH, DMSO, rt; ii) C6H5MgBr, MeCN, rflx.

Scheme 4. Synthesis of phenyl ethers [Ph].

Reagents and conditions: i) K2CO3, MeCN, cat Aliquat, reflx.

The cyclohexylmethyl phenyl ethers [Ph] were obtained by alkylation of appropriate 4-alkylphenol (8[n]) with heptyl tosylate (for [Ph]a) or trans-4-pentylcyclohexylmethyl tosylate22 9 for [Ph]b and [Ph]c (Scheme 4). Phenol 8[11] was obtained by alkylation of 4-bromophenol with C11H23MgBr following a general procedure.21

Thermal analysis

Transition temperatures and enthalpies of the newly prepared compounds were determined by differential scanning calorimetry (DSC) and results are shown in Tables 1 and 2.

Single component materials. All six non-ionic carborane derivatives 2 form a nematic phase (Table 1), which is consistent with nematogenic behavior of their benzene analogues. The three-ring compounds 2a and 2b exhibit a wide temperature range enantiotropic phase with clearing temperatures around 200 °C. A comparison of the carborane derivatives with their benzene analogues demonstrates that the phase stability follows the order [10] < [12] < [Ph], which is in agreement with our previous findings.16,25 The two-ring mesogens, 2c, exhibit monotropic phases with significantly lower N–I transition temperatures.
The phenyl ether [Ph]c is a new member of a known homologous series and it exhibits a mesophase identified by XRD analysis as a soft crystalline B phase (Cr 32 SmB crystal, N-nematic, I-isotropic). Temperatures obtained on heating at 5 K min⁻¹. Enthalpies are listed in the ESI.

DSC, POM, and XRD analyses revealed that pyridinium bromide [Pyr]c–Br exhibits a SmA phase and another more ordered unidentified phase. Unfortunately, the salt decomposes at about 165 °C. The shorter analogue, [Pyr]b–Br, is not mesogenic.

Table 1. Transition temperatures for 2.

<table>
<thead>
<tr>
<th>R¹</th>
<th>A</th>
<th>Cr</th>
<th>N</th>
<th>I</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>[10]</td>
<td>114*</td>
<td>175</td>
<td>*</td>
</tr>
<tr>
<td>[Ph]²</td>
<td>89.5</td>
<td>200</td>
<td>*</td>
<td></td>
</tr>
<tr>
<td>b</td>
<td>[10]</td>
<td>67</td>
<td>197</td>
<td>*</td>
</tr>
<tr>
<td>[Ph]³</td>
<td>91</td>
<td>206</td>
<td>*</td>
<td></td>
</tr>
<tr>
<td>c</td>
<td>[10]</td>
<td>44*</td>
<td>36</td>
<td>*</td>
</tr>
</tbody>
</table>

Cr-crystal, N-nematic, I-isotropic. Temperatures obtained on heating at 5 K min⁻¹. Enthalpies are listed in the ESI.

Two-component materials. Ion pairs 1a–[Pyr]a and 1b–[Pyr]a were reported before, and they display SmA and nematic phases. Previous results demonstrated that ions pairs 1c–[Pyr]a, in which the total number of rings is 3, melt about 110 °C and do not form liquid crystalline phases. Inclusion of the cyclohexyl ring in the structure of the pyridinium cation increased the melting temperature of ion pair 1[12]c–[Pyr]b and induced an enantiotropic soft crystalline phase E (Table 2). Extension of the butyl chain in the [Pyr]b cation to undecyl in [Pyr]c resulted in the appearance of an enantiotropic SmA phase above the E phase in both ion pairs, 1[10]c–[Pyr]c and 1[12]c–[Pyr]c, with a clearing temperature of about 210 °C (e.g. Fig. 3 and 4). DSC analysis of 1[12]c–[Pyr]c also revealed another soft-crystalline phase X below the E phase.

Previous results demonstrated that ions pairs 1a–[Pyr]a and 1b–[Pyr]a do exhibit enantiotropic SmA and N phases.

For comparison purposes, equimolar mixtures of non-ionic analogues of 1–[Pyr] were investigated. Initial studies of 2a–[Ph]a and 2b–[Ph]a demonstrated phase separation upon cooling from the isotropic phase. It was considered that the significant difference in the molecular length and mesogenic behaviour of the two components leads to their incompatibility. Therefore, carboranes 2c and phenyl ether [Ph]c were considered, since the total length of the alkyl chains in each compound is 16 atoms.

Table 2. Transition temperatures for binary systems 1-[Pyr] and 2-[Ph].

<table>
<thead>
<tr>
<th>X</th>
<th>Y</th>
<th>Cr</th>
<th>E</th>
<th>SmA</th>
<th>N</th>
<th>I</th>
</tr>
</thead>
<tbody>
<tr>
<td>1[10]c–[Pyr]c</td>
<td>B N°</td>
<td>126</td>
<td>163</td>
<td>207</td>
<td>*</td>
<td></td>
</tr>
<tr>
<td>2[10]c–[Ph]c</td>
<td>C C</td>
<td>27</td>
<td>*</td>
<td>26</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1[12]c–[Pyr]c</td>
<td>B N°</td>
<td>148*</td>
<td>200</td>
<td>210</td>
<td>*</td>
<td></td>
</tr>
<tr>
<td>2[12]c–[Ph]e</td>
<td>C C</td>
<td>30</td>
<td>*</td>
<td>29</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1[10]c–[Pyr]e</td>
<td>B N°</td>
<td>107*</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1[12]c–[Pyr]b</td>
<td>B N°</td>
<td>132</td>
<td>172</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Cr-crystal, SmA-smectic A, E-soft crystalline phase, N-nematic, I-isotropic. Temperatures obtained on heating at 5 K min⁻¹. Enthalpies are listed in the ESI. For definition of substituents see Fig. 1. Cr 85 Cr 41 Cr 41 Cr 101 X 107.
26 °C and 29 °C, respectively (Table 2). A comparison of these transition temperatures with those obtained for the analogous ion pair 1c-[Pyr]c shows a difference in the clearing temperature of 181 K. Since the ion-pair and the non-ionic binary mixture differ only by a molecular charge, the increase in the clearing temperature can be attributed solely to the strong coulombic interactions present in the ion pair and absent in the non-ionic binary mixture.

![Fig. 4](image1)

**Fig. 4.** The optical textures of a) SmA phase (205 °C) and b) E phase (175 °C) obtained for 1c-[Pyr]c on cooling from the isotropic phase in the same region of the sample.

**XRD data**

The formation of the SmA phase by ion pairs 1[10]c-[Pyr]c and 1[12]c-[Pyr]c was confirmed by XRD measurements. Diffractograms obtained for the high temperature mesophase in each material showed a series of sharp commensurate reflections consistent with a lamellar structure (Fig. 5). Considering the calculated molecular length of about 33.8 Å for both anions 1[10]c and 1[12]c (including Van der Waals radius for the terminal H atoms) the measured layer spacing of about 27 Å indicates approximately 20% of interdigitation and/or imperfect orientational order of molecular long axes and molecular folding in both ion-pairs.

The wide-angle region of the diffractograms shows an unsymmetric broad halo, which can be deconvoluted into two signals with the maxima about 4.7 Å and 5.1 Å for 1[10]c-[Pyr]c and 4.8 and 5.3 for 1[12]c-[Pyr]c. The diffused signals correlate with the mean distance between the alkyl chains (former) and the separation between boron clusters (latter).

**Table 3.** Unit cell parameters for E phase in 1c-[Pyr].

<table>
<thead>
<tr>
<th>Temperature / °C</th>
<th>a / Å</th>
<th>b / Å</th>
<th>c / Å</th>
</tr>
</thead>
<tbody>
<tr>
<td>1[12]c-[Pyr]c</td>
<td>180</td>
<td>9.16</td>
<td>6.50</td>
</tr>
</tbody>
</table>

![Fig. 5](image2)

**Fig. 5.** Integrated XRD patterns taken in SmA phase of 1[10]c-[Pyr]c (T = 175 °C, upper red curve) and 1[12]c-[Pyr]c (T = 205 °C, lower black curve). Upper curve is vertically shifted for clarity of presentation.

![Fig. 6](image3)

**Fig. 6.** 2D XRD patterns for E phase of (a) 1[10]c-[Pyr]c at 150 °C and (b) 1[12]c-[Pyr]c at 180 °C. Note the presence of equatorial signals (100) and (010) in a) and their absence in b) indicating regular ABA layer stacking for 1[12]c-[Pyr]c.

XRD analysis of the mesophase formed by phenyl ether [Ph]c revealed a soft crystalline B phase (SmBcry), which is consistent with a relatively large enthalpy of the isotropic transition (12.7 kJ mol⁻¹) and the polarized optical texture (Fig. 2). The measured layer spacing is 31.2 Å and correlates well with the calculated molecular length of 32.8 Å (including Van der Waals radius for H atoms).

**Molecular modelling**

The energy of coulombic interactions in ion pairs 1c-[Pyr]c were assessed computationally by comparison of the association energy for 1[12]c-[Pyr]c and 2[12]c-[Ph]c for an arbitrary chosen relative orientation of the molecules in the pair. The calculations were conducted using the M06-2x functional, which reasonably well reproduces non-covalent interactions.29
Analysis of fully optimized individual structures revealed a nearly linear anion $\text{I}[\text{12}]\text{c}$ and a bent cation $\text{[Pyr]}\text{c}$ (Fig. 7). The hexyl chain and the carbonyl group in the former are staggered relative to the \textit{closo-1-CB$_{11}$} cage. The two carboxy groups are nearly co-planar with the benzene ring rendering the “organic” half of the molecule planar (Fig. 7a). The angle between the planes of the terminal alkyl chains is 15°, and the molecular length ($L$, as the $\text{H-H}$ distance) is 31.6 Å. The carborane analogue $\text{2}[\text{12}]\text{c}$ has a nearly identical molecular length ($L = 31.3$ Å). The main difference between $\text{1}[\text{12}]\text{c}$ and $\text{2}[\text{12}]\text{c}$ is that the carbonyl group nearly eclipses the B–H bond in the latter, which result in the wider interplanar angle of 27°.

The undecyl chain in the pyridinium cation $\text{[Pyr]}\text{c}$ is orthogonal to the ring plane, which results in a bent molecular structure as shown in Fig. 7b. The interplanar angle is 78° and the $\text{H-H}$ separation $L$ of 30.3 Å, which is close to the length of anion $\text{1}[\text{12}]\text{c}$. Molecular dimensions of the benzene analogue $\text{[Ph]}\text{c}$ are nearly the same with the interplanar angle of 81° and $L = 30.2$ Å.

Full geometry optimization of $\text{2}[\text{12}]\text{c}$ and $\text{[Ph]}\text{c}$ in vacuum gave a tight molecular pair $\text{2}[\text{12}]\text{c}–\text{[Ph]}\text{c}$, which served as the starting point for geometry optimization of ion pair $\text{1}[\text{12}]\text{c}–\text{[Pyr]}\text{c}$. The resulting close-contact structure is shown in Figure 7c. A comparison of energies of the individual components and that of the binary system gave the enthalpy of association, $\Delta H_a$, after BSSE correction. For the non-ionic pair the association is calculated to be mildly exothermic ($\Delta H_a = -4.3$ kcal mol$^{-1}$), while for the ion pair, $\text{1}[\text{12}]\text{c}–\text{[Pyr]}\text{c}$, the association enthalpy is dramatically larger $\Delta H_a = -64.3$ kcal mol$^{-1}$ in vacuum. Placing molecules at low strength dielectric medium ($\varepsilon = 2.5$), typical for weakly polar liquid crystals, had little effect on the association enthalpy of $\text{2}[\text{12}]\text{c}–\text{[Ph]}\text{c}$ ($\Delta H_a = -3.4$ kcal mol$^{-1}$). In contrast, the dielectric medium significantly reduced the exotherm of formation of ion pair $\text{1}[\text{12}]\text{c}–\text{[Pyr]}\text{c}$ ($\Delta H_a = -27.9$ kcal mol$^{-1}$). As a result, the difference in association energy, $\Delta\Delta H_a$, of the non-polar pair and ion pair is 24.5 kcal mol$^{-1}$ in a weakly dielectric medium. Increasing $\varepsilon$ to 10.0, which is reasonable for an ionic liquid, the $\Delta\Delta H_a$ value for the two pairs falls to 7.0 kcal mol$^{-1}$. Thus, the calculated significant difference in association enthalpy in the two model binary systems is consistent with the observed 181 K higher thermal stability of the mesophase in ILC than in the non-ionic binary mixture. Also, the observed tendency for the formation of high-temperature lamellar phases (SmA and E) by the ion pair instead of a nematic phase is a consequence of the tight molecular arrangement dictated by significant coulombic interactions.

**Conclusions**

For the first time, we have assessed experimentally the impact of coulombic interactions on mesophase stability by comparison of ionic ($\text{1c}–\text{[Pyr]}\text{c}$) and isosteric non-ionic ($\text{2c}–\text{[Pyr]}\text{c}$) systems in the absence of other factors. Thus, replacement of appropriate carbon atoms in an equimolar binary mixture with B$^+$ and N$^+$ fragments leads to an isosteric and isoelectronic ion pair. Results show that the difference in the clearing temperature between the two binary systems is 181 K, which corresponds to a difference in association enthalpy of 7 kcal in appropriate dielectric media. The boron cluster size in the ionic material appears to have little effect on the magnitude of this phenomenon. It is possible that the size of the ions is the determining factor. A comparison was demonstrated for components of approximately equal length due to their best compatibility of the electrically neutral compounds (i.e. formation of a homogeneous nematic phase).

The presented concept of isosteric and isoelectronic replacement of C,C with B,N is applicable to other molecular systems and provides unique experimental models to support theoretical considerations of the liquid crystalline state.

**Computational Details**

Quantum-mechanical calculations were carried out using a Gaussian 09 suite of programs. Geometry optimizations for unconstrained conformers of appropriate molecules and ions with the most extended molecular shapes were performed at the M06-2x/3-21G* level of theory using default convergence limits. The alkoxy group was set in all-trans conformation maintaining co-planarity with the aromatic ring in the input structure. The aromatic ring and the carboxyl group were staggered with respect to the carborane cage as found experimentally and computationally in related structures. The orientation of the alkyl and carboxyloxy substituents on the alicyclic ring was set according to conformational analysis of the corresponding 1-ethyl and 1-acetoxy derivatives of cyclohexane. Optimized structures of non-polar molecules served as starting points for optimization of the ionic analogues after replacing the carbon atoms with B or N.

For calculations of binary system $\text{2}[\text{12}]\text{c}–\text{[Ph]}\text{c}$, molecules of $\text{2}[\text{12}]\text{c}$ and $\text{[Ph]}\text{c}$ at equilibrium geometry (M06-2x/3-21G*) were set parallel to each other at a 4 Å distance, and the geometry of the pair was minimized. The resulting energy was corrected for basis set superposition error (BSSE) by running
single point calculations (M06-2x/3-21G*/M06-2x/3-21G*) at the equilibrium geometry with the keyword COUNTERPOISE2 = 2.33 and compared to that of isolated molecular components. The optimized geometry of non-polar pair 2[12]c–[Pyr]c was served as the starting point for calculations involving ion pair 1[12]c–[Pyr]c, after replacement of the two carbon atoms with B and N. Conformational search was not attempted. For each pair, the association energy was calculated for isolated molecules (ΔH LAP = Hpair – (Hcat + Ha)).

Thermodynamic parameters were requested with the FREQ keyword. The PCM model14 was implemented using the SCRF(Solvent=Generic, Read) keyword and specified “epsinf=2.25” and “eps=2.5” or “eps=10” and the total energies were obtained in single point calculations with the same method.

**Experimental Part**

**General.** NMR spectra were obtained at 128 MHz (1H) and 400 MHz (13C) in CDCl3 or CD3CN. Chemical shifts were referenced to the solvent (1H) or to an external sample of B(OH)3 in MeOH (13C, δ = 18.1 ppm). Optical microscopy and phase identification were performed using polarized optical microscopy (POM) using a hot stage. Thermal analysis was obtained using a DSC with samples of about 0.5-1.0 mg and a heating rate of 5 K/min under a flow of nitrogen gas.

**General Procedure for the Preparation of 1–[Pyr].** To a solution of carboxylic acid 4[12]–[NEt4] or 4[10]–[NMe4] (0.1 mmol) in dry CH2Cl2 (2 mL) was added oxalyl chloride (5 eq) and a catalytic amount of DMF. The mixture was stirred at rt for 30 min, and the mixture was evaporated to complete dryness. Dry CH2Cl2 was added (3 mL) followed by appropriate phenol 3 (0.1 mmol) and Et3N (0.2 mmol), and the reaction was stirred overnight. Organic products were extracted (CH2Cl2), the organic layer was dried (Na2SO4) and the solvent was evaporated. The residue was passed through a silica gel plug (hexane/CH2Cl2, 4:1) giving crude ester 2, which was purified further by recrystallization.

**Method A.** A solution of carboxylic acid 5 (0.1 mmol) in dry CH2Cl2 (2 mL) was added oxalyl chloride (5 eq) and a catalytic amount of DMF. The mixture was stirred at rt for 30 min, and the mixture was evaporated to complete dryness. Dry CH2Cl2 was added (3 mL) followed by appropriate phenol 3 (0.1 mmol) and Et3N (0.2 mmol), and the mixture was stirred overnight. Organic products were extracted (CH2Cl2), the organic layer was dried (Na2SO4) and a solvent was evaporated. The residue was passed through a silica gel plug (hexane/CH2Cl2, 4:1) giving crude ester 2, which was purified further by recrystallization.

**Method B.** A solution of carboxylic acid 5 (0.1 mmol) and appropriate phenol 3 (0.11 mmol) was treated with N,N'-dicyclohexylcarbodiimide (DCC, 31 mg, 0.15 mmol) and a catalytic amount of 4-dimethylaminopyridine. The reaction mixture was stirred overnight. The reaction mixture was then washed with 10% HCl, the CH2Cl2 layer separated, dried (Na2SO4), and evaporated. The crude material was passed through a silica gel plug (CH2Cl2) giving crude ester 2, which was purified further by recrystallization.

**Method A.** Obtained as a white solid from acid 5[10] according to Method B and recrystallized from EtOH: 1H NMR (400 MHz, CDCl3) δ 0.88 (t, J = 6.9 Hz, 3H), 0.94 (t, J = 7.0 Hz, 3H), 1.03–1.10 (m, 2H), 1.12–1.31 (m, 6H), 1.32–1.46 (m, 20H), 1.49–1.58 (m, 2H), 1.5–3.5 (m, 8H), 1.95 (quint, J = 7.9 Hz, 2H), 2.49–2.57 (m, 2H), 2.32 (t, J = 8.3 Hz, 2H), 7.18 (d, J = 8.6 Hz, 2H), 7.23 (d, J = 8.6 Hz, 2H); 11B NMR δ = –10.3 (d, J = 160 Hz, 8B). Anal. Calcd. for C13H53B3O2: C, 68.29; H, 9.98. Found: C, 68.56; H, 9.95%.

**Method A.** Obtained as a white solid from acid 5[12] according to Method B and recrystallized from EtOH: 1H NMR (400 MHz, CDCl3) δ 0.85 (t, J = 7.1 Hz, 3H), 0.87 (t, J = 6.9 Hz, 3H), 1.00–1.07 (m, 2H), 1.08–1.42 (m, 30H), 1.5–3.5 (m, 10H), 1.60–1.64 (m, 2H), 2.42–2.49 (m, 2H), 6.83 (d, J = 8.5 Hz, 2H), 7.09 (d, J = 8.5 Hz, 2H); 11B NMR δ = –13.9 (5B), –12.7 (5B). Anal. Calcd. for C13H53B3O2: C, 65.45; H, 9.92. Found: C, 65.74; H, 9.84%.
8.79. Found: C, 57.36; H, 8.70%.

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4-(trans-4-Pentaylcyclohexylmethyl)pyridine (6). Following a general literature procedure,23,35 a suspension of NaH (755 mg, 19.7 mmol, 60% in oil) in dry DMSO (50 mL) was treated with trans-4-pentaylcyclohexylmethanol23 (7.30 g, 16.4 mmol) under an Ar atmosphere. The mixture was stirred for 2 h at rt, and then freshly prepared 4-chloropyridine23,35 was added. The reaction mixture was stirred at rt overnight. Water was added, and the mixture was extracted (hexane). The hexane layer was separated, dried (Na2SO4), and evaporated. The crude product was passed through a silica gel plug (CH2Cl2/EtOAc, 4:1) giving 2.25 g (53% yield) of pyridine 6 as a white solid: mp 72-72 °C; 1H NMR (400 MHz, CDCl3) δ 0.89 (t, J = 7.0 Hz, 3H), 0.90-1.11 (m, 3H), 1.12-1.36 (m, 10H), 1.70-1.78 (m, 18H), 1.81 (br d, J = 13.8 Hz, 2H), 1.88 (br d, J = 13.8 Hz, 2H), 3.79 (d, J = 6.4 Hz, 2H), 6.78 (d, J = 6.4 Hz, 2H), 8.40 (d, J = 6.4 Hz, 2H). Anal. Calcd. for C12H2NO: C, 78.11; H, 10.41; N, 5.36. Found: C, 78.15; H, 10.24; N, 5.41%.

4-(trans-4-Pentaylcyclohexylmethyl)-1-butylyl-pyridinium bromide ([Pyr]b–Br). A solution of pyridine 6 (300 mg, 1.15 mmol) and 1-bromobutane (1.02 mL, 9.4 mmol) in dry MeCN (10 mL) was refluxed for 15 h. The solvent was evaporated, and the residue was passed through a silica gel plug (MeCN/CH2Cl2, 4:1) yielding 620 mg (62% yield) of pyridinium salt [Pyr]b–Br which was recrystallized from EtOH to give a white solid: DSC Cr1 53 Cr4 80 107 I; 1H NMR (400 MHz, CDCl3) δ 0.88 (t, J = 7.0 Hz, 3H), 0.96 (t, J = 7.3 Hz, 3H), 0.95-1.14 (m, 3H), 1.15-1.34 (m, 10H), 1.40 (sext, J = 6.0 Hz, 2H), 1.75-1.89 (m, 5H), 1.95 (quint, J = 7.6 Hz, 2H), 4.06 (d, J = 6.0 Hz, 2H), 4.73 (t, J = 7.4 Hz, 2H), 7.39 (d, J = 7.2 Hz, 2H), 9.16 (d, J = 7.2 Hz, 2H). Anal. Calcd. for C12H8BrNO: C, 63.31; H, 9.11; N, 3.52. Anal. Calcd. for C12H8BrNO·0.5H2O: C, 61.91; H, 9.15; N, 3.44. Found: C, 62.14; H, 9.02; N, 3.50%.

4-(trans-4-Pentaylcyclohexylmethyl)-1-undecyl-pyridinium bromide ([Pyr]c–Br). It was prepared in an analogous way to the synthesis of [Pyr]b–Br using 6 and undecyl bromide and isolated as a white solid. The salt was further purified by crystallization (3x) from EtOH containing a few drops of hexane giving a white solid: DSC Cr75 Sm145 Sma163 dec. 1; 1H NMR (400 MHz, CDCl3) δ 0.87 (t, J = 6.8 Hz, 2H), 0.89 (t, J = 6.9 Hz, 3H), 0.95-1.15 (m, 3H), 1.16-1.39 (m, 26H), 1.76-1.89 (m, 5H), 1.96 (quint, J = 7.1 Hz, 2H), 4.07 (d, J = 6.0 Hz, 2H), 4.70 (t, J = 7.4, 2H), 7.40 (d, J = 7.4 Hz, 2H), 9.09 (d, J = 7.4 Hz, 2H). Anal. Calcd. for C25H30BrNO·H2O: C, 67.72; H, 10.15; N, 2.82. Anal. Calcd. for C25H30BrNO·H2O:
C. 65.35; H. 10.18; N. 2.72. Found: C. 65.62; H. 10.18; N. 2.75%.

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**Notes and references**

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Graphical Abstract

A comparison of two isosteric, ionic and non-ionic binary systems, provides a measure of coulombic interactions impact on mesophase stability in the absence of other factors.