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# $\pi$ -Electronic ion pairs: building blocks for supramolecular nanoarchitectonics via $i\pi-i\pi$ interactions

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The pairing of charged  $\pi$ -electronic systems and their ordered arrangement have been achieved by  $i\pi-i\pi$  interactions that are derived from synergetically worked electrostatic and dispersion forces. Charged  $\pi$ -electronic systems that provide ion pairs as building blocks for assemblies have been prepared by diverse strategies for introducing charge in the core  $\pi$ -electronic systems. One method to prepare charged  $\pi$ -electronic systems is the use of covalent bonding that makes  $\pi$ -electronic ions and valence-mismatched metal complexes as well as protonated and deprotonated states. Noncovalent ion complexation is another method used to create  $\pi$ -electronic ions, particularly for anion binding, producing negatively charged  $\pi$ -electronic systems. Charged  $\pi$ -electronic systems afford various ion pairs, consisting of both cationic and anionic  $\pi$ -systems, depending on their combinations. Geometries and electronic states of the constituents in  $\pi$ -electronic ion pairs affect the photophysical properties and assembling modes. Recent progress in  $\pi$ -electronic ion pairs has revealed intriguing characteristics, including the transformation into radical pairs through electron transfer and the magnetic properties influenced by the counteranions. Furthermore, the assembly states exhibit diversity as observed in crystals and soft materials including liquid-crystal mesophases. While the chemistry of ion pairs (salts) is well-established, the field of  $\pi$ -electronic ion pairs is relatively new; however, it holds great promise for future applications in novel materials and devices.

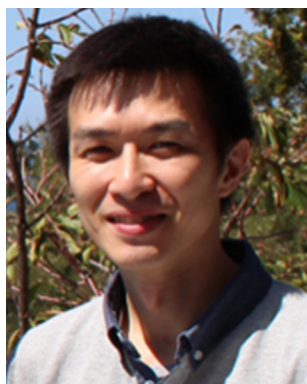
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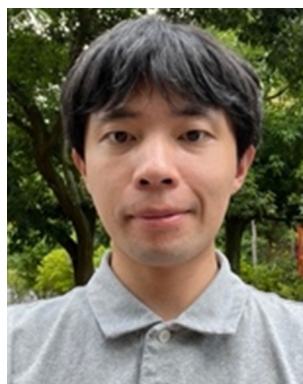
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**Yohei Haketa**

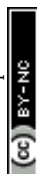
Yohei Haketa received his PhD degree in 2011 from Ritsumeikan University, under the guidance of Prof. Hiromitsu Maeda, focusing on the assemblies of  $\pi$ -conjugated anion receptors. He was selected as a Research Fellow of the Japan Society for the Promotion of Science (JSPS) in 2009–2012. After working in Asahi Kasei Corporation as a researcher (2012–2015), he joined the group of Prof. Hiromitsu Maeda at Ritsumeikan University as a postdoctoral fellow in

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Kazuhisa Yamasumi received his PhD degree in 2020 from Kyushu University under the guidance of Prof. Hiroyuki Furuta, focusing on isomeric porphyrin analogues. His PhD research was focused on the synthesis of infrared dyes based on the expanded or  $\pi$ -extended isomeric porphyrin analogues. He joined the group of Prof. Hiromitsu Maeda at Ritsumeikan University as a postdoctoral fellow in 2020. His current research focuses on  $\pi$ -electronic ion-pairing assemblies.



# 1. Introduction

Various functional materials that are applied in organic electronics can be fabricated by  $\pi$ -electronic systems that have electronic states depending on their core structures and substituents.<sup>1</sup> Electronic properties of such materials are modulated by the arrangement of constituents *via* noncovalent interactions.<sup>2</sup> Understanding and utilization of these interactions offer valuable insights for designing captivating assembled structures and materials. Noncovalent interactions, such as  $\pi$ - $\pi$ , hydrogen-bonding and van der Waals interactions, are facilitated by combining fundamental intermolecular forces, including (i) electrostatic, induction and dispersion forces as long-range forces originating from coulombic forces and (ii) exchange-repulsion and charge-transfer (CT) forces as short-range forces caused by interorbital interaction.<sup>3</sup> Electronically neutral  $\pi$ -electronic systems form stacking assemblies by dispersion forces, the main components of  $\pi$ - $\pi$  interactions.

The introduction of charge into  $\pi$ -electronic systems drastically changes the assembling behaviour based on electrostatic forces. Electrostatic and dispersion forces play a significant role in the arrangement of charged  $\pi$ -electronic systems in their assemblies.<sup>4,5</sup> The assemblies feature stacking structures, with the fundamental units being  $\pi$ -stacked ion pairs ( $\pi$ -sips). These  $\pi$ -sips represent characteristic forms of paired oppositely charged  $\pi$ -electronic systems ( $\pi$ -electronic ion pairs) (Fig. 1a).<sup>6</sup>  $\pi$ -Sips form various ion-pairing assemblies with the modes of charge-by-charge assemblies, wherein oppositely charged species are alternately stacked; moreover, they provide charge-segregated assemblies, in which identically charged species are stacked *via* dissociation and recombination (Fig. 1a). The assembling modes are modulated by the geometries and electronic states of charged  $\pi$ -electronic systems in ion pairs. Various  $\pi$ -electronic ion pairs can be formed by combining  $\pi$ -electronic cations and anions; ten kinds of cations and ten kinds of anions ideally afford 100 kinds of ion pairs. Chemistry of  $\pi$ -electronic ion-pairing assemblies

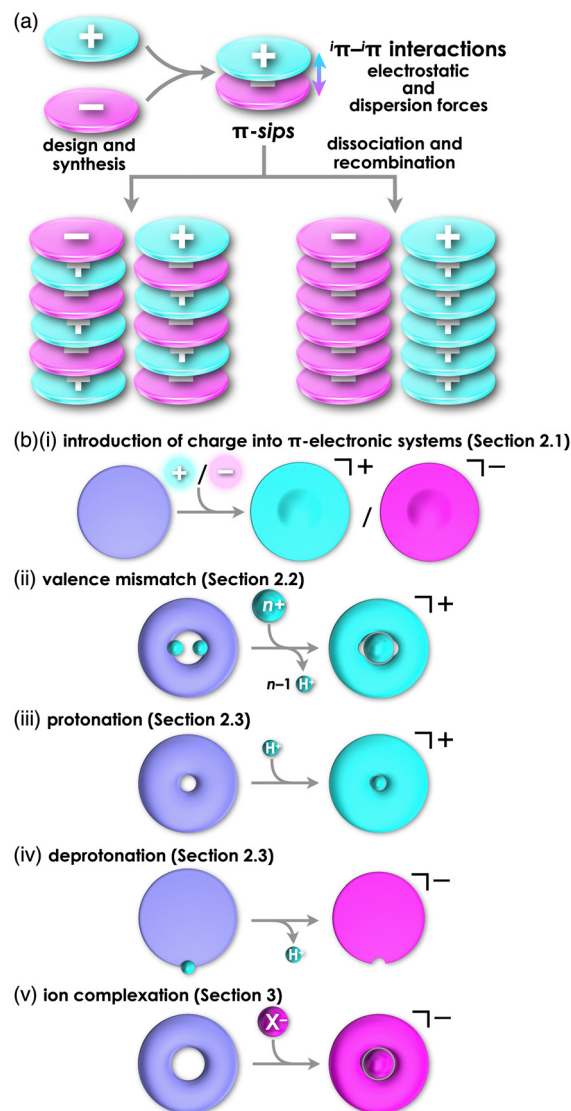


Fig. 1 (a) Conceptual diagrams for the formation of  $\pi$ -sips (top) and charge-by-charge (bottom left) and charge-segregated assemblies (bottom right) comprising charged  $\pi$ -electronic species and (b) formation of charged  $\pi$ -electronic systems *via* "covalent approaches" by (i) introducing charges directly into the core, (ii) valence mismatch by metal complexation, denoted by cation formation as a representative and (iii) protonation and (iv) deprotonation at the basic and acidic sites, respectively, along with (v) ion complexation, as "noncovalent approach" in contrast to (i–iv), denoted by anion binding as a representative.

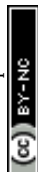


Hiromitsu Maeda

Hiromitsu Maeda received his PhD degree in 2004 from Kyoto University, under the guidance of Prof. Hiroyuki Furuta (Kyushu University) and Prof. Atsuhiko Osuka, after spending three months in the Sessler group, the University of Texas at Austin, in 2001. In 2004, he started an academic career in Department of Bioscience and Biotechnology, Faculty of Science and Engineering, Ritsumeikan University. In 2008, he was transferred to College of Pharmaceutical

Sciences, wherein he was promoted to a professor in 2014. In 2016, he moved to Department of Applied Chemistry, College of Life Sciences. He has been awarded several prizes, including ChemComm Emerging Investigator Lectureship (2012) and Fellow of the RSC (2015).

started in 2010 with the fabrication of charge-by-charge assemblies comprising  $\pi$ -electronic receptor–anion complexes and  $\pi$ -electronic cations in the form of single crystals, supramolecular gel and liquid crystals (*vide infra*, Section 3.1).<sup>7,8</sup> In 2021, cooperative interactions involving electrostatic and dispersion forces were proposed as  $i\pi$ - $j\pi$  interactions *via* detailed experimental and theoretical investigations, including single-crystal X-ray analysis and energy decomposition analysis (EDA) for solid-state ion-pairing assemblies.<sup>9</sup> As previously defined, the  $i\pi$  in " $i\pi$ - $j\pi$  interactions" represents  $\pi$ -electronic ions and charged  $\pi$ -electronic systems.  $i\pi$ - $j\pi$  Interactions induce stacking of oppositely charged



$\pi$ -electronic systems owing to the enhanced contribution of attractive electrostatic and dispersion forces, resulting in the formation of  $\pi$ -sips and their assemblies. For identically charged  $\pi$ -electronic systems,  $i\pi$ - $i\pi$  interactions can be modulated depending on the balance of electrostatic and dispersion forces, facilitating the formation of charge-segregated assemblies. The electronic states of charged  $\pi$ -electronic systems are closely related with their HOMO (highest occupied molecular orbital) and LUMO (lowest unoccupied molecular orbital), which are essential in redox processes, photophysical properties and reactivity. Stacked states of oppositely charged  $\pi$ -electronic systems with relatively high HOMO levels for anions and low LUMO levels for cations can induce charge- and electron-transfer behaviours. Diverse charged  $\pi$ -electronic systems have been reported as potential components of  $\pi$ -electronic ion pairs, including commercially available reagents. This review article summarizes the ion pairs comprising  $\pi$ -electronic cations and anions and their properties for potential applications as functional materials.

Charged  $\pi$ -electronic systems as the components of  $\pi$ -electronic ion pairs can be obtained by diverse methods. A charge can be introduced by covalent and noncovalent approaches (Fig. 1b). In Section 2, several strategies using “covalent approaches” for the preparation of  $\pi$ -electronic charged species are displayed, such as the use of aromaticity and valence mismatch (Fig. 1b(i,ii)). Protonated and deprotonated species of  $\pi$ -electronic systems, using the coordination bond between a proton and a basic site for the former, are also included (Fig. 1b(iii,iv)). Section 3 demonstrates the formation of pseudo- $\pi$ -electronic ions through the complexation of  $\pi$ -electronic systems and inorganic ions by noncovalent interactions (Fig. 1b(v)). This process results in the creation of ion pairs and their assemblies, including the charged  $\pi$ -electronic systems discussed in Section 2. Although various pseudo- $\pi$ -electronic cations and anions are formed using this method, anion-complexation strategies are the main focus. In Section 4, pairs of the charged  $\pi$ -electronic systems shown in Section 2 are summarized as solution-state  $\pi$ -sips and assemblies in crystalline states and mesophases. The photophysical properties of  $\pi$ -electronic ion pairs are also displayed. However, radical ions derived from electronically neutral closed-shell  $\pi$ -systems are not included in this review.

## 2. $\pi$ -Electronic systems with charge introduced by covalent approaches: components of ion pairs with non- $\pi$ -electronic counterions

Diverse strategies, classified as covalent approaches, are available for the production of charged  $\pi$ -electronic systems (Fig. 1b(i–iv)). The introduced charge in  $\pi$ -electronic systems can be delocalized for stabilization. Aromaticity achieved by charge introduction further contributes to the stabilization of charged species. Valence mismatch between  $\pi$ -electronic ligands and metal ions also provides charged  $\pi$ -electronic systems. Furthermore, protonation and deprotonation of electronically neutral  $\pi$ -electronic systems are

facile approaches to forming positively and negatively charged  $\pi$ -electronic systems.

### 2.1. Introduction of charge into $\pi$ -electronic systems

Carbocations and carbanions are generally less stable species, whose reactivity can be reduced by delocalizing the charges through extended  $\pi$ -systems (Fig. 1b(i)). For example, the triphenylmethyl cation (tritylium)  $1^+$  (Fig. 2a(i)), which can be used as a Lewis acid in various organic synthesis, is more stable than the *tert*-butyl cation.<sup>10</sup> The stability of cations is further improved by introducing electron-donating groups, as seen in crystal violet ( $2^+$  as a cation part, Fig. 2a(ii)).<sup>11</sup> Moreover, the  $\pi$ -extension of tritylium can be achieved by reducing the steric repulsion between the *ortho*-CH units to form planar structures. Bridging two aryl groups with an atom results in the formation of planar structures as observed in rhodamine B ( $3a^+$  as a cation part, Fig. 2b(i)), which possesses a planar tricyclic  $\pi$ -unit that is characteristic of xanthene dyes.<sup>12,13</sup> Rhodamine B, represented as  $3a^+Cl^-$ , exhibits strong absorption and fluorescence bands in the visible region at 554 and 627 nm, respectively, in MeOH. These properties make the dye suitable for fluorescence imaging applications.<sup>13a</sup> Furthermore, replacement of the oxygen with other heteroatoms, like phosphorus in  $3b^+$  (Fig. 2b(ii)), alters the electronic states and photophysical properties. The absorption and emission maxima of  $3b^+$  in a phosphate-buffered saline (PBS) buffer (pH = 7.4) are observed at 712 and 740 nm, respectively.<sup>13c</sup> Notably, various positively charged dyes, as seen in tritylium, crystal violet and rhodamine B, are commercially available.

Bridging the *ortho*-CH units of the three aryl groups of tritylium with three N or O linkers affords planar trianguleniums such as triazatriangulenium (TATA)<sup>14</sup> (Fig. 3a) and trioxatriangulenium (TOTA)<sup>15</sup>  $4a^+$  (Fig. 3b(i)). A variety of triangulenium derivatives, including less planar helicene structures,<sup>14a</sup> have been reported thus far for the use in various applications such as phase transition catalysis, bioimaging and DNA intercalation. In particular, the planar charged  $\pi$ -electronic systems are suitable for efficient electron-accepting properties and stacking in various supramolecular assemblies.<sup>16</sup> For example, TATA<sup>+</sup> bearing ether

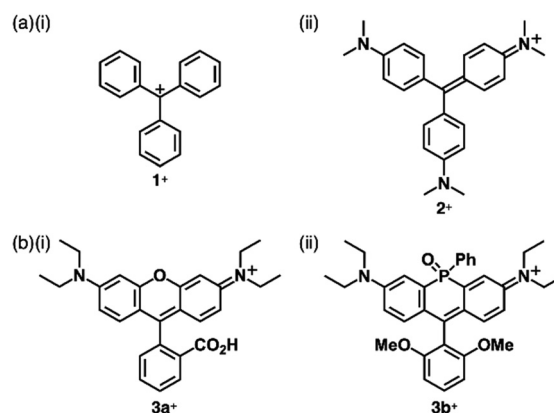
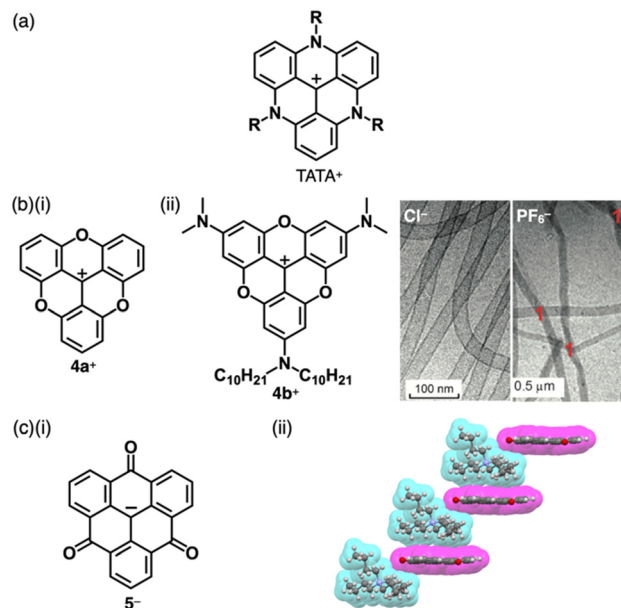


Fig. 2 Representative charged triarylmethane dyes (a)  $1^+$  and  $2^+$  and (b)  $3a^+$ ,  $3b^+$ .

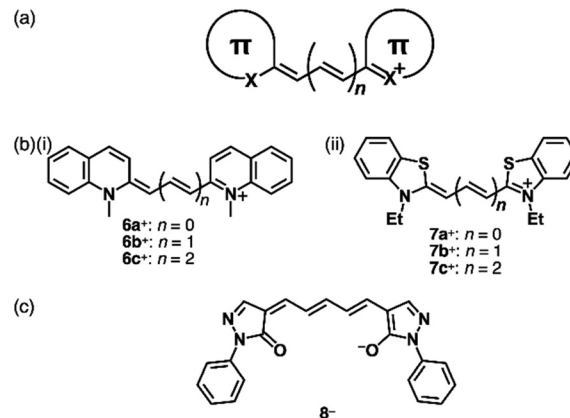




**Fig. 3** (a) General structure of TATA<sup>+</sup>, (b) (i) TOTA<sup>+</sup> **4a**<sup>+</sup> and (ii) alkyl-substituted TOTA<sup>+</sup> **4b**<sup>+</sup> and cryo-TEM images of nanostructures of **4a**<sup>+</sup>-X<sup>-</sup> (X<sup>-</sup> = Cl<sup>-</sup> (left) and PF<sub>6</sub><sup>-</sup> (right)) formed in water (10 vol% CH<sub>3</sub>CN) (redrawn from ref. 15c. Copyright 2014 Wiley) and (c) (i) trioxotriangulene anion **5**<sup>-</sup> and (ii) single-crystal X-ray structure of TBA<sup>+</sup>-**5**<sup>-</sup> (redrawn from a cif file: 1303470). Atom colour code in (c) and the following figures: grey, white, blue and red in the ball-and-stick models refer to carbon, hydrogen, nitrogen and oxygen, respectively, whereas cyan and magenta in the space-filling models refer to cation and anion, respectively.

chains with a terminal hydroxy unit has been used as a visible-light absorbing photosensitizer in metal-catalyzed hydrogen production in water.<sup>16a</sup> Conversely, TOTA<sup>+</sup> with aliphatic alkyl chains at one terminal (**4b**<sup>+</sup>, Fig. 3b(ii)) formed counteranion-depending nanoscale assemblies comprising stacked **4b**<sup>+</sup> (Fig. 3b(ii)).<sup>15c</sup> Planarization can also be applied to anion species: bridging triarylmethyl anion with three electron-withdrawing carbonyl groups afforded the trioxotriangulene anion **5**<sup>−</sup> (Fig. 3c(i)).<sup>17</sup> **5**<sup>−</sup> is thermodynamically stabilized owing to the charge-delocalized closed-shell structure, forming the ion pair with tetrabutylammonium (TBA<sup>+</sup>). Single-crystal X-ray analysis of TBA<sup>+</sup>·**5**<sup>−</sup> revealed that TBA<sup>+</sup> and planar **5**<sup>−</sup> were alternately stacked to form a columnar structure (Fig. 3c(ii)).

Allyl ions and their extended ionic polymethines are stabilized by introducing two aryl groups at both ends (Fig. 4a).<sup>18</sup> Various derivatives were synthesized as polymethine dyes with terminal aryl groups such as pyrylium, quinolinium<sup>18e</sup> (**6a-c<sup>+</sup>**, Fig. 4b(i)) and benzothiazolium<sup>18b</sup> (**7a-c<sup>+</sup>**, Fig. 4b(ii)). As polymethine dyes exhibit sharp absorption bands in the visible-to-near infrared (NIR) region depending on the length of the polymethine chains and the terminal  $\pi$ -electronic systems, cationic dyes are applied as fluorescent probes, laser dyes and photovoltaic materials. Generally, the polymethine dyes exhibit red-shifted absorption bands depending on the length of the linker polymethine units, as seen in the absorption maxima of 425, 558 and 660 nm for **7a-c<sup>+</sup>**, respectively. The photophysical properties are affected by aggregation due to the significant



**Fig. 4** (a) General structure of polymethine dyes, (b) positively charged polymethine dyes based on (i) *N*-methylquinolium (**6a-c<sup>+</sup>**) (ii) benzothiazolium (**7a-c<sup>+</sup>**) and (c) negatively charged polymethine dye based on deprotonated oxonol (**8<sup>-</sup>**).

exciton coupling caused by the large transition dipole moments. Therefore, assemblies of polymethine dyes have attracted great attention to date.<sup>18a,19</sup> Furthermore, negatively charged  $\pi$ -electronic systems (*e.g.*,  $8^-$ , Fig. 4c) are stabilized by the introduction of electron-withdrawing terminal  $\pi$ -units, although applications are limited owing to decreased stability.<sup>20</sup>

Modifications at the centre of the polymethine chains affect the rigidity and modulate the photophysical properties.<sup>18</sup> The positive charge can be compensated by the introduction of negatively charged substituents at the polymethine chains, as seen in zwitterionic squarylium **9** (Fig. 5a) and croconium dyes.<sup>21</sup> These dyes are converted to the corresponding cations by introducing alkyl groups at the anionic site as seen in **10**<sup>+</sup> (Fig. 5a).<sup>22</sup> In the crystal structures of ion pairs **10**<sup>+</sup>X<sup>-</sup> (X<sup>-</sup> = OTf<sup>-</sup> and B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub><sup>-</sup>), the **10**<sup>+</sup> units are stacked by themselves.

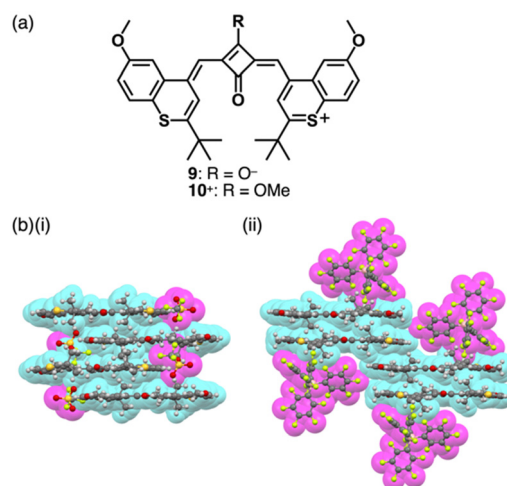


Fig. 5 (a) Zwitterionic squarylium dye **9** and cationic squarylium dye **10<sup>+</sup>** and (b) single-crystal X-ray structures of (i) **10<sup>+</sup>**·OTf<sup>−</sup> and (ii) **10<sup>+</sup>**·B(C<sub>6</sub>F<sub>5</sub>)<sub>4</sub><sup>−</sup> (redrawn from cif files: 2214017 and 2214019). Atom colour code in (b) and the following figures: pink, yellow green and dark yellow in the ball-and-stick models refer to boron, fluorine and sulfur, respectively.

Notably, the stacked  $10^+$  units exhibit antiparallel orientation to cancel their dipoles (Fig. 5b).<sup>23</sup> Furthermore,  $10^+$  shows sharp absorption bands in the NIR region, which are shifted depending on the assembling conditions (*vide infra*, Section 4.1).

Charged units can be stabilized by aromaticity.<sup>24</sup> A smallest class of aromatic compounds is cyclopropenium, a positively charged  $2\pi$  electronic system (Fig. 6a(i)),<sup>25</sup> which is highly reactive and participates in various reactions. The stability of cyclopropenium is increased by the introduction of electron-donating units, as seen in  $11^+$  (Fig. 6a(ii)). Cycloheptatrienyl cation, tropylium ( $Tr^+$ ), a  $6\pi$  electronic system, is a relatively stable  $\pi$ -electronic cation (Fig. 6b).<sup>26</sup> In contrast, cyclopentadienide ( $Cp^-$ ) as a negatively charged  $6\pi$  electronic system is stable enough for various metallocenes (Fig. 6c(i)).<sup>27</sup> Delocalization of negative charges by  $\pi$ -extension can stabilize anions as seen in the tris(7*H*-dibenzo[*c,g*]fluorenylidene)methyl anion (Kuhn's anion)  $12^-$  (Fig. 6c(ii)), a genuine hydrocarbon charged  $\pi$ -electronic anion including  $Cp^-$  units.<sup>28</sup> Kuhn's anion has been synthesized as ion pairs with various cations, including  $\pi$ -electronic cations. Owing to the anion's easily oxidizable nature, the ion pairs must be handled under an inert atmosphere and light shading conditions to prevent unwanted reactions or degradation.

The stability of  $Cp^-$  can be enhanced by the introduction of electron-withdrawing units. For example, various ion pairs have incorporated pentamethoxycarbonylcyclopentadienide  $13^-$  (Fig. 7a(i)).<sup>29</sup> The crystal structure of  $TMA^+-13^-$  ( $TMA^+$  = tetramethylammonium) revealed that dihedral angles between methoxycarbonyl  $CO_2C$  planes and the core cyclopentadienide unit were within a range of  $15.7^\circ$ – $85.9^\circ$ , exhibiting a nonplanar structure (Fig. 7a(ii)).<sup>30</sup> Replacement of the ester groups with cyano groups as less bulky electron-withdrawing groups allows the production of planar anions that are available for more effective stacking with charged  $\pi$ -electronic systems. Since the synthesis of pentacyanocyclopentadienide ( $PCCp^-$ )<sup>31</sup> by Webster in 1965 (Fig. 7b(i)),<sup>31a</sup> the anion has been used as the building block of solid-state coordination networks in combination with metal ions.<sup>32</sup> Exchange of the counteranion with other cations *via* ion-pair metathesis as an ion-exchange

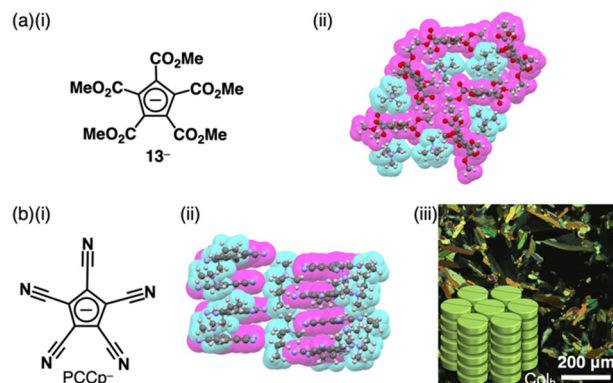


Fig. 7 (a) (i) Pentamethoxycarbonylcyclopentadienide  $13^-$  and (ii) single-crystal X-ray structure of  $TMA^+-13^-$  and (b) (i) pentacyanocyclopentadienide ( $PCCp^-$ ), (ii) single-crystal X-ray structure of  $(C_4H_9)(CH_3)_3N^+-PCCp^-$  and (iii) POM of liquid crystal mesophase of  $(C_{12}H_{25})_3CH_3N^+-PCCp^-$  at  $70^\circ C$  upon cooling (redrawn from ref. 33. Copyright 2016 Wiley). The crystal structures are redrawn from cif files: 1137646 and 1431736.

protocol based on hard and soft acids and bases (HSAB) theory (details are summarized in Section 4) afforded various ion pairs, demonstrating unique assembled structures.<sup>33,34</sup> Single-crystal X-ray analysis of  $(C_4H_9)(CH_3)_3N^+-PCCp^-$  exhibited a columnar structure comprising stacking of  $PCCp^-$  with a distance of  $3.42\text{ \AA}$  (Fig. 7b(ii)). Notably,  $(C_4H_9)(CH_3)_3N^+$  is located proximally to  $PCCp^-$ , constructing a charge-segregated assembly. Furthermore, a charge-segregated assembly in the form of a hexagonal columnar ( $Col_h$ ) structure was observed in the liquid crystal mesophase of  $(C_{12}H_{25})_3CH_3N^+-PCCp^-$  (Fig. 7b(iii)), and its hole mobility was estimated to be  $0.4\text{ cm}^2\text{ V}^{-1}\text{ s}^{-1}$  using field-induced time-resolved microwave conductivity (FI-TRMC) technique.<sup>33</sup>

Replacement of carbons with heteroatoms in aromatic  $\pi$ -electronic systems affords charged aromatic  $\pi$ -electronic systems (Fig. 8). Notably, the replacement of a CH unit in benzene with substituted nitrogen and oxygen/sulfur produces pyridinium, pyrylium and thiopyrylium, respectively ( $14a-c^+$  as triphenyl-substituted derivatives, Fig. 8a).<sup>35</sup> Taking advantage

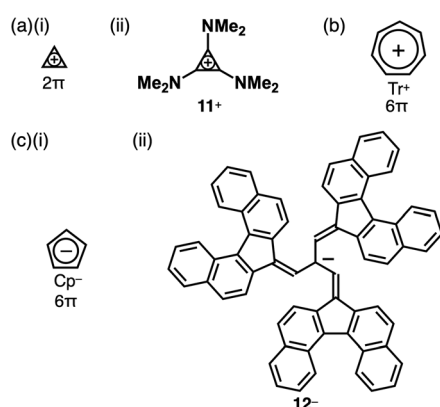


Fig. 6 Representative charged  $\pi$ -electronic systems stabilized by aromaticity: (a) (i) cyclopropenium and (ii) tris(dimethylamino)-substituted derivative  $11^+$ , (b) tropylium and (c) (i) cyclopentadienide and (ii) Kuhn's anion  $12^-$ .

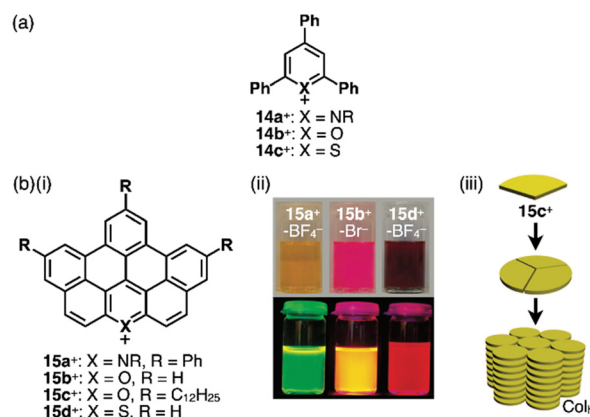


Fig. 8 (a) Triphenylpyridinium and triphenylpyryliums  $14a-c^+$  and (b) (i)  $\pi$ -extended pyridinium and pyryliums  $15a-d^+$ , (ii) photographs of  $CH_2Cl_2$  solutions of  $15a^+-BF_4^-$ ,  $15b^+-Br^-$  and  $15d^+-BF_4^-$  (top: under visible light, bottom: under UV) and (iii) packing diagram of  $15c^+-Br^-$  (redrawn from ref. 36. Copyright 2009 ACS).



of the electron-deficient nature and moderate reactivity, 2,4,6-triarylpyryliums (e.g., **14b**<sup>+</sup>) are used as photocatalysts and building blocks of extended  $\pi$ -electronic systems.<sup>35b</sup> Some charged species, such as pyridinium, are stable under ambient conditions. Other charged  $\pi$ -electronic systems can be stabilized by  $\pi$ -extensions as seen in peripherally heteroatom-containing phenanthro[2,3,4,5-*qgrab*]perylene as  $\pi$ -extended pyridinium, pyrylium and thiopyrylium **15a–d**<sup>+</sup> reported by Feng, Müllen and coworkers (Fig. 8b(i)),<sup>36</sup> whose electronic properties clearly depend on the introduced heteroatoms. Red-shifted UV/vis absorption and emission maxima occur at 512/531, 573/594 and 600/643 nm for **15a,b,d**<sup>+</sup> corresponding to BF<sub>4</sub><sup>−</sup>, Br<sup>−</sup> and BF<sub>4</sub><sup>−</sup> ion pairs, respectively, in CH<sub>2</sub>Cl<sub>2</sub> (Fig. 8b(ii)). Dodecyl-substituted **15c**<sup>+</sup> as a Br<sup>−</sup> ion pair forms liquid crystal mesophases based on the Col<sub>h</sub> structure (Fig. 8b(iii)). Conversely, the internal carbon of naphthalene can be replaced with nitrogen, resulting in quinolizinium as the smallest N-doped polycyclic aromatic hydrocarbon (PAH).<sup>37</sup> Electron-deficient planar charged  $\pi$ -electronic systems with appropriate sizes and substituents can be applied to DNA intercalation. Representative  $\pi$ -extended N-doped PAHs include 9-phenylbenzo[1,2]quinolizino[3,4,5,6-*fed*]-phenanthridinylium (PQP<sup>+</sup>) cations **16a,b**<sup>+</sup> (Fig. 9a), synthesized from a 2,4,6-triphenylpyrylium salt.<sup>38</sup> PQP<sup>+</sup> forms charge-segregated assemblies *via*  $\pi$ - $\pi$  interactions.<sup>38b</sup> In particular, PQP<sup>+</sup> ion pairs **16b**<sup>+</sup>-X<sup>−</sup> (X<sup>−</sup> = Cl<sup>−</sup> and BF<sub>4</sub><sup>−</sup>) form counteranion-dependent ion-pairing assemblies with distinct morphologies (Fig. 9b).<sup>38a</sup> TEM images of the assemblies exhibit nanoribbons and nanotubes for the Cl<sup>−</sup> and BF<sub>4</sub><sup>−</sup> ion pairs, respectively, suggesting that even small counteranions have important roles in the arrangement of charged  $\pi$ -electronic units and resulting organized structures.

Porphyrin, consisting of four pyrrole rings bridged by methine units, is an 18 $\pi$  electronic aromatic macrocycle.<sup>39</sup> Porphyrin and its analogues play important roles in biological systems such as metabolism and photosynthesis. The large  $\pi$ -electronic system of porphyrin contributes to charge delocalization and, thus, stabilization of charged species. Various stacking assemblies can be formed from porphyrin derivatives.<sup>19,40</sup> Therefore, porphyrin is an excellent building block for charged  $\pi$ -electronic systems as components of  $\pi$ -electronic ion-pairing assemblies. In addition, the redox properties of porphyrin can be modulated by peripheral substituents. The introduction of substituted N and O/S at one of the porphyrin *meso* positions affords positively charged heteroatom-containing porphyrins **17a–c**<sup>+</sup> (Fig. 10a(i)).<sup>41</sup> The

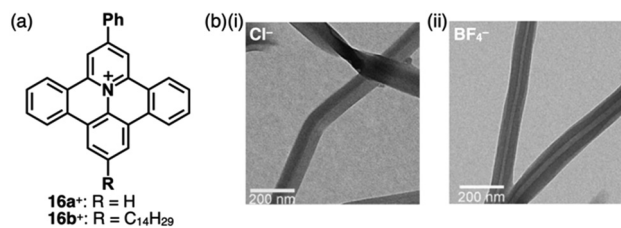


Fig. 9 (a) PQP<sup>+</sup> **16a,b**<sup>+</sup> and (b) TEM images of aggregates in MeOH (1 mM): (i) **16b**<sup>+</sup>-Cl<sup>−</sup> and (ii) **16b**<sup>+</sup>-BF<sub>4</sub><sup>−</sup> (redrawn from ref. 38a. Copyright 2007 Wiley).

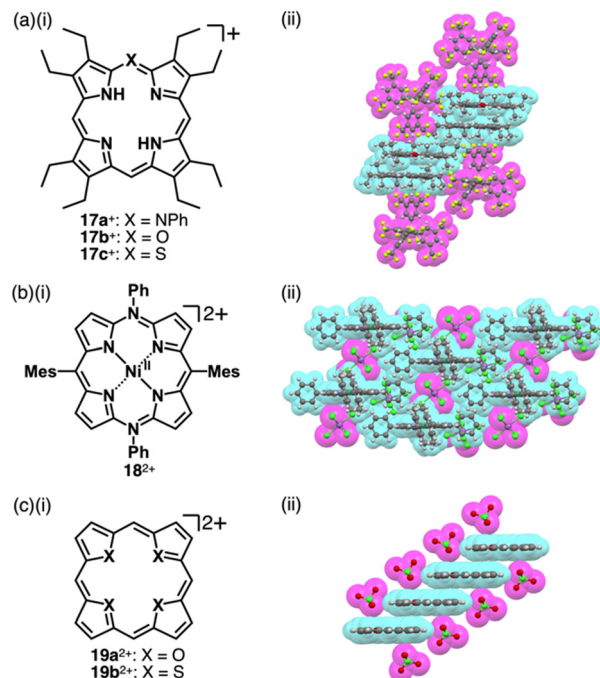


Fig. 10 (a) (i) Heteroatom-containing porphyrins **17a–c**<sup>+</sup> and (ii) single-crystal X-ray structure of **17a**<sup>+</sup>-BARF<sup>−</sup>, (b) (i) tetraaryl-5,15-diazaporphyrin Ni<sup>II</sup> complex **18**<sup>2+</sup> and (ii) single-crystal X-ray structure of **18**<sup>2+</sup>-2SbCl<sub>6</sub><sup>−</sup> and (c) (i) core-modified porphyrins **19a,b**<sup>2+</sup> and (ii) single-crystal X-ray structure of **19a**<sup>2+</sup>-2ClO<sub>4</sub><sup>−</sup>. The crystal structures are redrawn from cif files: 2033992, 1437728 and 1166405. Atom colour codes in (b) (c) (ii) and the following figures: green, blue-green and purple in the ball-and-stick models refer to chlorine, nickel and antimony, respectively.

inner NH protons of **17a,c**<sup>+</sup> are located at the nitrogens in the *trans* positions, whereas those of **17b**<sup>+</sup> are located at those in the *cis* positions, far from the introduced *meso*-oxygen.<sup>41b–d</sup> This arrangement suggests charge delocalization and resulting polarized structures attributed to the introduced atoms. In the crystal structures of **17a–c**<sup>+</sup>-BARF<sup>−</sup> (BARF<sup>−</sup> = B(3,5-(CF<sub>3</sub>)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>)<sub>4</sub><sup>−</sup>), the cations are stacked by themselves on the  $\pi$ -plane, while further stacking is inhibited by BARF<sup>−</sup> located on the  $\pi$ -plane (Fig. 10a(ii)). Furthermore, tetraaryl-5,15-diazaporphyrin Ni<sup>II</sup> complex **18**<sup>2+</sup> as a dication in PF<sub>6</sub><sup>−</sup> and SbCl<sub>6</sub><sup>−</sup> ion pairs can be obtained by oxidizing the corresponding neutral form with AgPF<sub>6</sub> or magic blue (Fig. 10b).<sup>42</sup> Moreover, a similar approach to the one mentioned in the previous paragraph can be applied to heterocyclic systems, where heteroatoms are converted to other heteroatoms with different valences. Heteroporphyrins **19a,b**<sup>2+</sup>, whose inner nitrogens are replaced with chalcogens, form various charged  $\pi$ -electronic systems (Fig. 10c(i)).<sup>43</sup> In the crystal structure of the ClO<sub>4</sub><sup>−</sup> ion pair, **19a**<sup>2+</sup> appears highly planar and forms a columnar structure, slip-stacking by itself on both sides (Fig. 10c(ii)). However, the thiophene units in **19b**<sup>2+</sup> are inclined by 22.8° and 3.7° in the crystal structure.<sup>43c</sup>

## 2.2. Charged $\pi$ -electronic systems derived from valence mismatch

Charges in complexes comprising multiple units can be controlled by the valences of the components. Therefore, if the number of positive charges of metal ions is not matched with



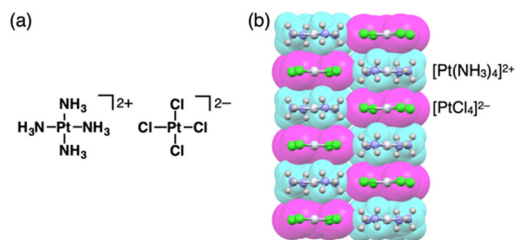


Fig. 11 Magnus' green salt: (a) ion-pairing form and (b) single-crystal X-ray structure (redrawn from a cif file: 2009952). Atom colour code in (b) and the following figures: light grey in the ball-and-stick models refers to platinum.

that of the total negative charges of ligands, the resulting complexes behave as charged species (Fig. 1b(ii)). Metal complexes with such a valence mismatch are used in Magnus' green salt: a salt comprising  $[\text{Pt}(\text{NH}_3)_4]^{2+}$  and  $[\text{PtCl}_4]^{2-}$ , alternately arranged to form an anisotropic quasi 1D linear array in the solid state (Fig. 11).<sup>44,45</sup> The 1D structure of the aligned  $\text{Pt}^{\text{II}}$  complexes in Magnus' green salt is mainly constructed by the electrostatic force between the positively and negatively charged planar  $\text{Pt}^{\text{II}}$  complexes. The fascinating ion-pairing structure has stimulated numerous studies regarding the use of 1D arrays as electrically conductive wires. Importantly, the correlation between the geometries of the constituents and the assembled structure has been clearly demonstrated. This observation has prompted us to design and synthesize charged  $\pi$ -electronic species for fabricating various functional ion-pairing assemblies through  $\pi$ - $\pi$  interactions.

As  $\pi$ -electronic ligands in preparation for charged  $\pi$ -electronic species by valence mismatch, porphyrins serve as dianionic ligands owing to the two inner NH groups. Therefore, complexation with trivalent metal ions provides  $\pi$ -electronic cations accompanied by an anion for charge compensation. Versatile porphyrin complexes with trivalent metal ions such as  $\text{Cr}^{\text{III}}$ ,  $\text{Mn}^{\text{III}}$ ,  $\text{Fe}^{\text{III}}$  and  $\text{Co}^{\text{III}}$  have been prepared thus far.<sup>39</sup> Most of the metal complexes have axial ligands, which compensate for the positive charge and hamper the stacking of the resulting  $\pi$ -electronic systems.  $20^+$ , a porphyrin  $\text{Fe}^{\text{III}}$  complex with bulky triisopropylsilyloxy substituents, can be obtained in the neutral form coordinated by a  $\text{Cl}^-$  ligand.<sup>46</sup> Here, the  $\text{Cl}^-$  is exchanged with other anions by ion-pair metathesis with  $\text{Ag}^+$  salts of weakly coordinating anions ( $\text{ClO}_4^-$ ,  $\text{OTf}^-$  and  $\text{SbF}_6^-$ ) (Fig. 12a). Hexabromocarborane anion ( $\text{CH}_6\text{B}_{11}\text{Br}_6^-$ )<sup>47</sup> is too bulky to approach the axial coordination site owing to the steric

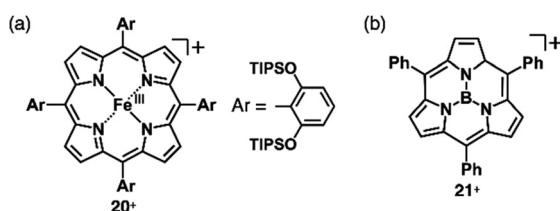


Fig. 12 Positively charged porphyrin and analogue formed by valence mismatch: (a) porphyrin  $\text{Fe}^{\text{III}}$  complex  $20^+$  and (b) subporphyrin boron complex  $21^+$ .

hindrance, resulting in  $\pi$ -electronic cations without axial ligands. Boron complexes of subporphyrins usually exhibit bowl-like structures attributed to the structural distortion caused by axial coordination at the boron centre.<sup>48</sup> Treatment of methoxide-coordinated subporphyrin boron complex with  $\text{Et}_3\text{Si}[\text{CH}_6\text{B}_{11}\text{Br}_6]$  afforded a planar subporphyrin cation  $21^+$  as an ion pair with  $\text{CH}_6\text{B}_{11}\text{Br}_6^-$  (Fig. 12b).<sup>49</sup> Conversely, porphyrin  $\text{Au}^{\text{III}}$  complexes, whose metal centre is in a  $d^8$  electronic state, generally have no axial ligands (e.g.,  $22a-e^+$ , Fig. 13a).<sup>6,50-53</sup> Thus, porphyrin  $\text{Au}^{\text{III}}$  complexes readily form ion pairs in combination with desired anions and can be adopted as building blocks for  $\pi$ -electronic ion-pairing assemblies.<sup>6,52</sup>

Porphyrins can be complexed with  $\text{Au}^{\text{III}}$  using different procedures. One well-known method involves using  $\text{KAuCl}_4$  in refluxing  $\text{AcOH}$ .<sup>51a</sup> Other synthetic approaches, such as the uses of  $\text{HAuCl}_4 \cdot 4\text{H}_2\text{O}/\text{AgOTf}/\text{NaOAc}$ <sup>51d</sup> and  $\text{Au}(\text{tht})_2\text{BF}_4$  (tht = tetrahydrothiophene)/2,6-lutidine,<sup>51c</sup> can be applied depending on the reactivity of the porphyrins. Counteranions of the porphyrin  $\text{Au}^{\text{III}}$  complexes after the  $\text{Au}^{\text{III}}$  complexation are mainly derived from the corresponding  $\text{Au}^{\text{III}}$  reagents. To exchange the counteranions to  $\text{Cl}^-$ , porphyrin  $\text{Au}^{\text{III}}$  complex ion pairs can be passed through an ion-exchange resin (Amberlite). Moreover, the  $\text{Cl}^-$  ions can be exchanged with inorganic salts containing the desired anions through ion-pair metathesis, resulting in various ion pairs.

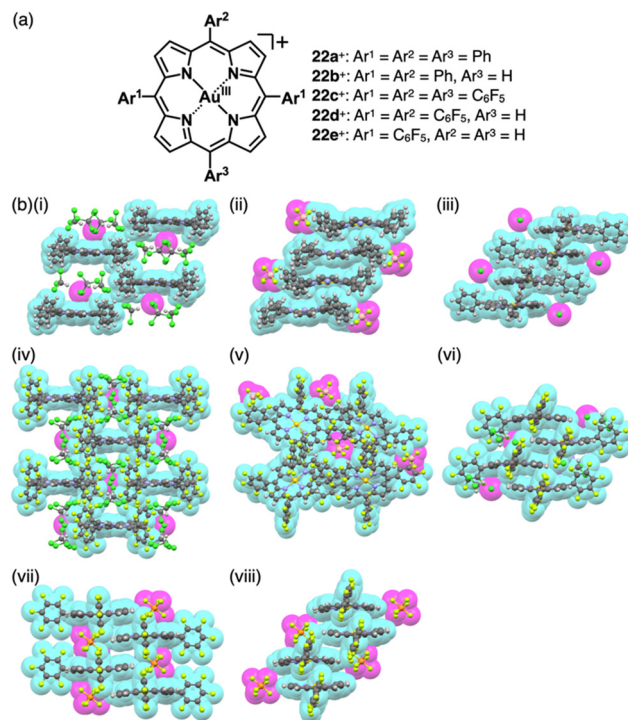
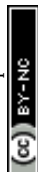


Fig. 13 Positively charged porphyrins formed by valence mismatch: (a) meso-arylporphyrin  $\text{Au}^{\text{III}}$  complexes  $22a-e^+$  and (b) single-crystal X-ray structures of (i)  $22a^+-\text{Cl}^-$ , (ii)  $22a^+-\text{BF}_4^-$ , (iii)  $22b^+-\text{Cl}^-$ , (iv)  $22c^+-\text{Cl}^-$ , (v)  $22c^+-\text{BF}_4^-$ , (vi)  $22d^+-\text{Cl}^-$ , (vii)  $22d^+-\text{PF}_6^-$  and (viii)  $22e^+-\text{PF}_6^-$  (redrawn from cif files: 1877987, 1877988, 2226630, 1904175, 1904176, 1904174, 1904178 and 1904177). Atom colour codes in (b) and the following figures: orange and yellow in the ball-and-stick models refer to phosphorus and gold, respectively.



The high stability of these ion pairs allows for purification using silica gel column chromatography.<sup>52</sup>

Single-crystal X-ray analysis revealed the exact structures of porphyrin Au<sup>III</sup> complexes with no axial ligands. **22a**<sup>+</sup>Cl<sup>−</sup> in a pseudo-polymorph shows a columnar structure based on a charge-by-charge assembly including Cl<sup>−</sup> with four co-crystallized CHCl<sub>3</sub> molecules (Fig. 13b(i)).<sup>51b,52a</sup> The proximal Au...Cl<sup>−</sup> distance was 3.12 Å, comparable to the sum of the ionic radii of Au<sup>3+</sup> and Cl<sup>−</sup>, suggesting the formation of a contact ion pair. In addition, the line passing through both Au and Cl has an angle of 80.2° to the mean plane of **22a**<sup>+</sup>. The non-90° angle indicates that Cl<sup>−</sup> is not directly coordinated to the core Au<sup>III</sup> in proximity to **22a**<sup>+</sup>. Instead, it exists as a π-electronic cation, and the interaction is primarily based on electrostatic interactions. In contrast to the charge-by-charge structure of **22a**<sup>+</sup>Cl<sup>−</sup>, charge-segregated assemblies are formed in **22a**<sup>+</sup>BF<sub>4</sub><sup>−</sup> and **22a**<sup>+</sup>PF<sub>6</sub><sup>−</sup> (Fig. 13b(ii)), exhibiting columnar structures of stacked **22a**<sup>+</sup> with stacking distances of 3.73/3.88 and 3.87/3.89 Å, respectively.<sup>52a</sup> Conversely, ion pairs of *meso*-triphenyl-substituted **22b**<sup>+</sup> with Cl<sup>−</sup> form charge-segregated assemblies with stacking distances of 3.50 and 3.74 Å (Fig. 13b(iii)), whereas **22b**<sup>+</sup>PF<sub>6</sub><sup>−</sup> exhibits a stacked dimer of **22b**<sup>+</sup> with a stacking distance of 3.46 Å. In both cases, counteranions are located at either side of **22b**<sup>+</sup>.<sup>52b</sup>

The assembling behaviours of the ion pairs are affected by the introduction of electron-withdrawing C<sub>6</sub>F<sub>5</sub> groups (**22c**–**e**<sup>+</sup>, Fig. 13a).<sup>52b</sup> The Cl<sup>−</sup> in **22c**<sup>+</sup>Cl<sup>−</sup> is in contact with the β-CH units of two porphyrins, resulting in layered structures separated by the bulky C<sub>6</sub>F<sub>5</sub> groups (Fig. 13b(iv)). Therefore, the Au...Cl distance (7.08 Å) is much longer than those of **22a**<sup>+</sup>Cl<sup>−</sup> (3.00 and 3.12 Å). Meanwhile, in **22c**<sup>+</sup>BF<sub>4</sub><sup>−</sup>, the C<sub>6</sub>F<sub>5</sub> groups are arranged around BF<sub>4</sub><sup>−</sup> with interionic π...F short contacts (Fig. 13b(v)). In **22d**<sup>+</sup>Cl<sup>−</sup>, Cl<sup>−</sup> is separated from the Au<sup>III</sup> centre by CH<sub>2</sub>Cl<sub>2</sub> (Fig. 13b(vi)) and is located proximally to the *meso*-CH of **22d**<sup>+</sup>. The arrangements of the cations and PF<sub>6</sub><sup>−</sup> in **22d**<sup>+</sup>PF<sub>6</sub><sup>−</sup> and **22e**<sup>+</sup>PF<sub>6</sub><sup>−</sup> are similar to that of **22d**<sup>+</sup>Cl<sup>−</sup>, although the interplanar distances are slightly longer (3.50–3.84 Å) than that of **22d**<sup>+</sup>Cl<sup>−</sup> (Fig. 13b(vii,viii)).<sup>52b</sup>

A similar approach can be applied to divalent metal complexes with monoanionic ligands such as thiaporphyrins (tetraphenylthiaporphyrin Ni<sup>II</sup> complex **23**<sup>+</sup> as a representative structure, Fig. 14a).<sup>39,54</sup> In 1989, Latos-Grażyński, Balch and coworkers reported **23**<sup>+</sup> and Cu<sup>II</sup> and Fe<sup>II</sup> complexes as Cl<sup>−</sup> ion pairs.<sup>54</sup> In the crystal structure of **23**<sup>+</sup>Cl<sup>−</sup>, the Ni...Cl<sup>−</sup> distance is 2.28 Å, indicating the axial coordination. The thiophene unit of **23**<sup>+</sup> is inclined by ~25°, leading to a less planar structure. Furthermore, ion pair **23**<sup>+</sup>PF<sub>6</sub><sup>−</sup>, prepared by ion-pair metathesis of **23**<sup>+</sup>Cl<sup>−</sup> with

AgPF<sub>6</sub>, forms the solid-state stacked dimer structure, wherein the S...N distance is 3.77 Å owing to chalcogen bonding (Fig. 14b).<sup>55</sup>

Similarly, anionic species can be obtained by complexing divalent metals with trianionic ligands such as corrole. The introduction of electron-withdrawing substituents at appropriate positions is required for stable π-electronic anions. Gross *et al.* reported *meso*-C<sub>6</sub>F<sub>5</sub>-substituted corrole Ni<sup>II</sup> complex **24**<sup>−</sup> as a monoanionic species in the TBA<sup>+</sup> ion pair (Fig. 15a(i)).<sup>56</sup> Here, the stability of the corrole anion was increased by introducing bromo units at the β-positions. Single-crystal X-ray analysis of TBA<sup>+</sup>·**24**<sup>−</sup> revealed that planar **24**<sup>−</sup> and TBA<sup>+</sup> were alternately arranged, forming a columnar structure (Fig. 15a(ii)). Similarly, the octaethylporphyrin Li<sup>I</sup> complex **25**<sup>−</sup>, consisting of monovalent metal and dianionic ligand, behaves as a π-electronic anion (Fig. 15b(i)).<sup>57</sup> **25**<sup>−</sup> was synthesized as a [Li(thf)<sub>4</sub>]<sup>+</sup> (thf = tetrahydrofuran (THF)) ion pair by Li<sup>I</sup> metallation of octaethylporphyrin in THF. In [Li(thf)<sub>4</sub>]<sup>+</sup>·**25**<sup>−</sup>, the components are alternately arranged, and the **25**<sup>−</sup> units are inclined to each other (Fig. 15b(ii)). Moreover, [Li(thf)<sub>4</sub>]<sup>+</sup>·**25**<sup>−</sup> is sensitive to moisture and undergoes demetallation by hydrolysis.

A variety of charged metal complexes based on neutral π-electronic ligand molecules, such as terpyridine, with another ligand for the partial charge compensation can be used to produce positively charged π-electronic species (*e.g.*, **26**<sup>+</sup>, Fig. 16a).<sup>58</sup> Terpyridine Pt<sup>II</sup> complexes with a variety of ligands at Pt<sup>II</sup> afford planar monocationic species, demonstrating aggregation behaviour using <sup>i</sup>π–<sup>i</sup>π and Pt<sup>II</sup>...Pt<sup>II</sup> interactions.<sup>59</sup> Yam *et al.* reported a terpyridine Pt<sup>II</sup> complex with an aliphatic alkynyl ligand **26a**<sup>+</sup>X<sup>−</sup> with different anions (X<sup>−</sup> = Cl<sup>−</sup>, OTf<sup>−</sup>, BF<sub>4</sub><sup>−</sup> and PF<sub>6</sub><sup>−</sup>), exhibiting luminescence and gelation properties (Fig. 16b(i)).<sup>59b</sup> The OTf<sup>−</sup> ion pair formed coil-shaped helical fibres, whereas other ion pairs formed disordered fibres (Fig. 16b(ii)). Furthermore, in 2012, the groups of Che and Yam independently reported a class of monocationic cyclometallated Au<sup>III</sup> alkynyl complexes **27a**,**b**<sup>+</sup> (Fig. 17a).<sup>60</sup> The positive

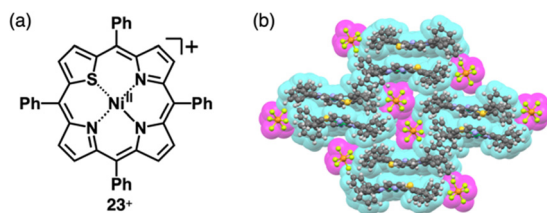


Fig. 14 (a) Thiaporphyrin Ni<sup>II</sup> complex **23**<sup>+</sup> and (b) single-crystal X-ray structure of **23**<sup>+</sup>PF<sub>6</sub><sup>−</sup> (redrawn from a cif file: 2167302).

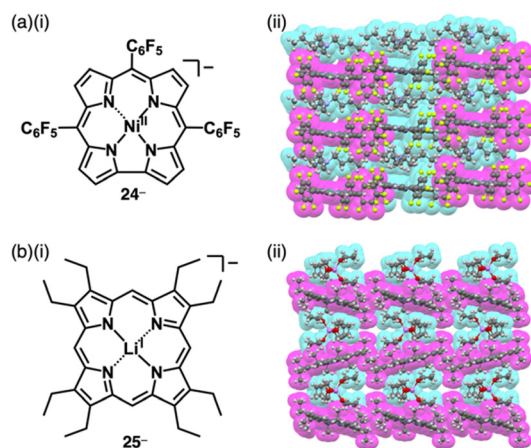


Fig. 15 (a) (i) Corrole Ni<sup>II</sup> complex **24**<sup>−</sup> and (ii) single-crystal X-ray structure of TBA<sup>+</sup>·**24**<sup>−</sup> and (b) (i) porphyrin Li<sup>I</sup> complex **25**<sup>−</sup> and (ii) single-crystal X-ray structure of [Li(thf)<sub>4</sub>]<sup>+</sup>·**25**<sup>−</sup> ([Li(thf)<sub>4</sub>]<sup>+</sup> is highlighted with cyan owing to the coordination of THF to Li<sup>+</sup>). The crystal structures are redrawn from cif files: 2122003 and 1184624. Atom colour code in (b) (ii) and following figures: light purple in the ball-and-stick models refers to lithium.



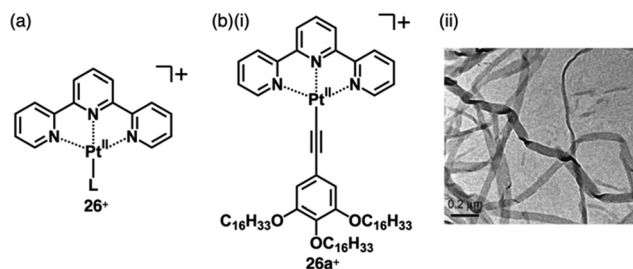


Fig. 16 (a) General structure of positively charged terpyridine Pt<sup>II</sup> complexes **26<sup>+</sup>** and (b) (i) terpyridine Pt<sup>II</sup> complex with aliphatic chains and (ii) TEM image of the DMSO gel of **26a<sup>+</sup>**-OTf<sup>-</sup> (redrawn from ref. 59b. Copyright 2009 Wiley).

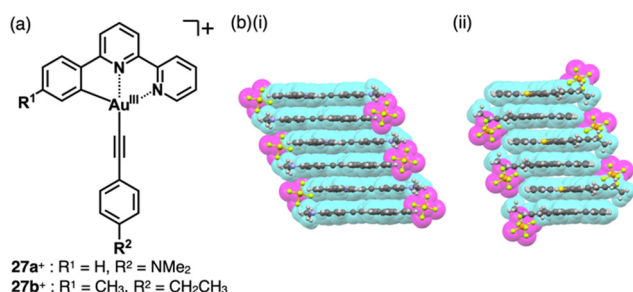


Fig. 17 (a) Positively charged phenylbipyridine Au<sup>III</sup> complexes **27a<sup>+</sup>**, **27b<sup>+</sup>** and (b) single-crystal X-ray structures of (i) **27a<sup>+</sup>**-PF<sub>6</sub><sup>-</sup> and (ii) **27b<sup>+</sup>**-PF<sub>6</sub><sup>-</sup> (redrawn from cif files: 841764 and 897336).

charges of the Au<sup>III</sup> are partially cancelled by phenylbipyridyl and alkynyl ligands, affording charge-segregated assemblies comprising stacked Au<sup>III</sup> complexes (Fig. 17b). The short Au<sup>III</sup>...Au<sup>III</sup> distances of 3.495 and 3.692 Å in **27a<sup>+</sup>**-PF<sub>6</sub><sup>-</sup> and **27b<sup>+</sup>**-PF<sub>6</sub><sup>-</sup> suggest favourable Au<sup>III</sup>...Au<sup>III</sup> interactions as confirmed by theoretical calculations. In **27a<sup>+</sup>**-PF<sub>6</sub><sup>-</sup>, contributions of  $i\pi$ - $\pi$  and Au<sup>III</sup>...Au<sup>III</sup> interactions facilitate the formation of belt-like assemblies.<sup>60a</sup> Tuning of metal centre valences also results in anionic species.<sup>61</sup> Lalinde *et al.* prepared stable anionic Pt<sup>II</sup> complexes with diphenylpyridine and alkynyl ligands (*e.g.*, **28<sup>-</sup>**, Fig. 18).<sup>61a</sup> Here, TBA<sup>+</sup> was selected for charge compensation of anionic Pt<sup>II</sup> complexes. Single-crystal X-ray analysis clearly elucidated the planar geometries of the anionic Pt<sup>II</sup> complexes, which were further used as guest species for cationic Pt<sup>II</sup> tweezers, exhibiting more efficient binding behaviour (log *K* = 13.5 in CH<sub>2</sub>Cl<sub>2</sub>) compared to an uncharged Pt<sup>II</sup> complex.<sup>62</sup>

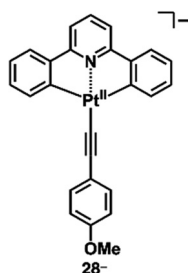


Fig. 18 Negatively charged diphenylpyridine Pt<sup>II</sup> complex **28<sup>-</sup>**.

### 2.3. Charged $\pi$ -electronic systems formed by protonation and deprotonation

Reversible proton attachment and detachment can be considered covalent approach methods *via* the modulation of covalent linkages (Fig. 1b(iii,iv)). Proton (H<sup>+</sup>) is the simplest cation, and protonation of electronically neutral  $\pi$ -electronic systems produces  $\pi$ -electronic cations. The charged  $\pi$ -electronic systems are often deprotonated, reverting neutral  $\pi$ -electronic systems. Thus,  $\pi$ -electronic systems with high proton affinity, achieved by multiple hydrogen bonds, are required to form  $\pi$ -electronic cations suitable for ion pairing.<sup>63,64</sup> Protonated azacalix[3](2,6)pyridine **29a<sup>+</sup>** has a cavity with three imine nitrogen sites, exhibiting high basicity (p*K*<sub>a</sub> = 23.1 in CD<sub>3</sub>CN) (Fig. 19a(i)).<sup>63a,b</sup> A planar structure with a bound proton in the cavity was investigated by single-crystal X-ray analysis of **29a<sup>+</sup>**-PF<sub>6</sub><sup>-</sup> (Fig. 19a(ii)). A columnar structure of slip-stacked **29a<sup>+</sup>** is formed with PF<sub>6</sub><sup>-</sup> located at the side of the columns. The basicity is further enhanced by introducing electron-donating pyrrolidine units at the pyridine 4-positions in **29b<sup>+</sup>** (p*K*<sub>a</sub> = 28.1 in CD<sub>3</sub>CN), facilitating its application as organocatalysis for Michael addition reactions.<sup>63b</sup> Replacement of the pyridyl units with triazine results in a further electron-deficient  $\pi$ -electronic cation azacalix[3]triazine **30<sup>+</sup>** (Fig. 19b(i)), maintaining high basicity (p*K*<sub>a</sub> = 16.7 for the ion pair with **30<sup>+</sup>**-Cl<sup>-</sup> in DMSO).<sup>63d</sup> The protonated species **30<sup>+</sup>** interacts with a counteranion, affecting the p*K*<sub>a</sub> values (17.1 and 16.2 for the NO<sub>3</sub><sup>-</sup> and PF<sub>6</sub><sup>-</sup> ion pairs,

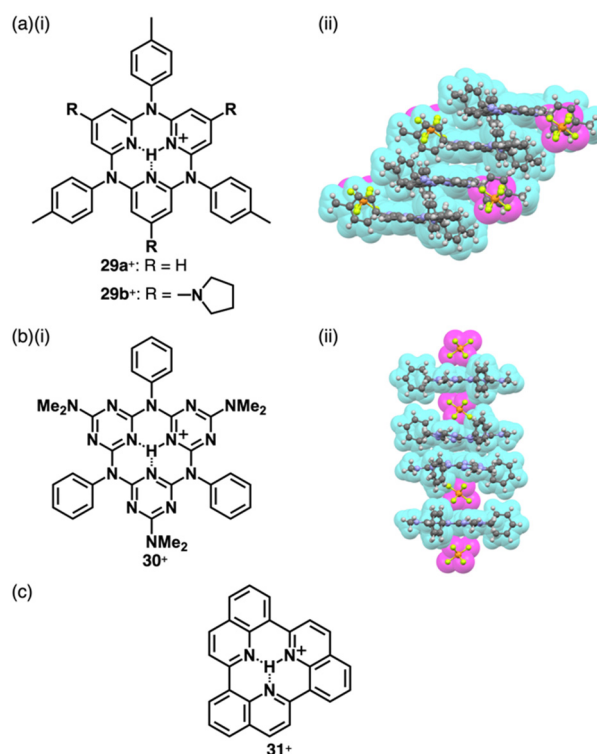


Fig. 19 (a) (i) Protonated azacalix[3](2,6)pyridine **29a<sup>+</sup>**, **29b<sup>+</sup>** and (ii) single-crystal X-ray structure of **29a<sup>+</sup>**-PF<sub>6</sub><sup>-</sup> (redrawn from a cif file: 298963), (b) (i) protonated azacalix[3]triazine **30<sup>+</sup>** and (ii) single-crystal X-ray structure of **30<sup>+</sup>**-PF<sub>6</sub><sup>-</sup> (redrawn from a cif file: 2072959) and (c) protonated triquinoline **31<sup>+</sup>**.

respectively, in  $\text{CD}_3\text{CN}$ ). Single-crystal X-ray analysis of  $30^+\text{-PF}_6^-$  reveals the planar structure of  $30^+$  with anion- $\pi$  interactions between  $\text{PF}_6^-$  and triazine units, resulting in the formation of discrete assemblies in the form of  $\text{PF}_6^-\text{-}30^+\text{-PF}_6^-\text{-}30^+\text{-PF}_6^-\text{-}30^+\text{-PF}_6^-$  (Fig. 19b(ii)). Protonated triquinoline  $31^+$  (Fig. 19c) possesses a planar rigid skeleton with a proton-binding site, as evidenced by the persistence of  $>10\%$  of the inner NH  $^1\text{H}$  NMR signal after one week in  $\text{CD}_3\text{OD}$ .<sup>64</sup> These highly basic macrocycles have been utilized as base catalysts in organic synthesis, and can serve as  $\pi$ -electronic cations in ion-pairing assemblies when paired with suitable anions.

Covalent linkage between charged substituents and  $\pi$ -electronic systems more drastically affects the electronic states of charged  $\pi$ -electronic systems. A ubiquitous negatively charged moiety is  $\text{O}^-$ , formed by the deprotonation of a hydroxy group. A  $\pi$ -electronic anion precursor is phenol, whose  $\text{pK}_a$  is 10.00 in an aqueous solution.<sup>65</sup>  $\pi$ -Extended phenol derivatives, such as fluorescein ( $\text{pK}_a = 6.7$  for the phenolic OH),<sup>66</sup> whose dianion form is uranine  $32^{2-}$  (Fig. 20a), are also commercially available as  $\text{Na}^+$  salts. Fluorescein derivatives that exhibit pH-dependent photophysical properties are used as pH indicators. Phenoxides are further stabilized by introducing electron-withdrawing groups as seen in picric acid ( $\text{pK}_a$  of 0.40),<sup>65</sup> providing  $33^-$  upon deprotonation (Fig. 20b).

4-Nitrophenol ( $\text{pK}_a = 7.14$ ), whose acidity is weaker than picric acid, can be used as a building unit of  $\pi$ -electronic anions. Introduction of pyrrole at the *ortho*-positions of the deprotonated site affords the intramolecular  $\text{N-H}\cdots\text{O}^-$  interactions, which stabilize the  $\pi$ -electronic anions  $34a-c^-$  (Fig. 21a(i)).<sup>67</sup> The  $\text{pK}_a$  value of  $34b$  (7.2) is higher than that of  $34a$  (6.2), comparable to 4-nitrophenol. In the solid state,  $\text{TBA}^+\text{-}34a,b^-$  form planar structures, stabilized by intramolecular  $\text{N-H}\cdots\text{O}$  hydrogen bonding. Planar  $34a,b^-$  are stacked with counter  $\text{TBA}^+$ , providing charge-by-charge columnar assemblies (Fig. 21a(ii)). Introduction of aliphatic alkyl chains at the aryl groups in  $\text{TBA}^+\text{-}34c^-$  induces liquid crystal mesophases with a  $\text{Col}_h$  structure based on charge-by-charge assembly, as observed in the crystal structure of  $\text{TBA}^+\text{-}34b^-$  (Fig. 21a(iii)).<sup>67b</sup> Efforts to introduce electron-withdrawing fluorine at the pyrrole  $\beta$ -positions in search of more stable  $\pi$ -electronic anions yields the desired species. However, handling and purification of this species is challenging owing to its easy deprotonation.<sup>67d</sup> In contrast, introducing thiophene units instead of pyrrole rings affords the  $\pi$ -electronic anion  $35a^-$ , with a  $\text{pK}_a$  of 6.7 (Fig. 21b(i)).<sup>68</sup> Detailed theoretical studies and single-crystal X-ray analysis of  $\text{TBA}^+\text{-}35a^-$  revealed intramolecular  $\text{S}\cdots\text{O}^-$  chalcogen bonding (Fig. 21b(ii)). Similarly,  $\text{TBA}^+\text{-}35b^-$  shows alternate stacking of planar  $35b^-$  and  $\text{TBA}^+$  (Fig. 21b(i,iii)). Furthermore, introducing quinone methide units as electron-withdrawing groups and

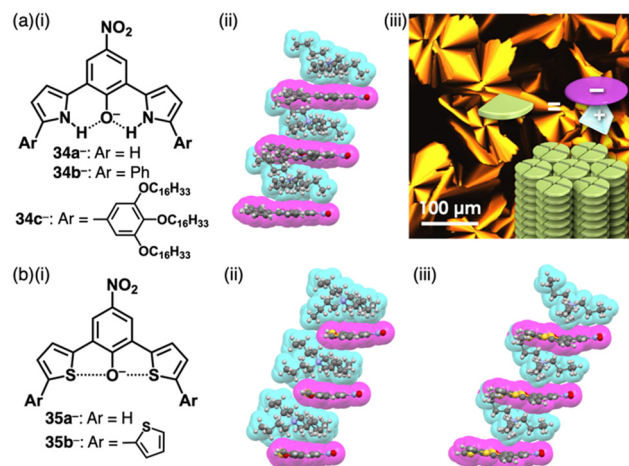


Fig. 21 (a) (i) Deprotonated dipyrrolynitrophenols  $34a-c^-$ , (ii) single-crystal X-ray structure of  $\text{TBA}^+\text{-}34b^-$  and (iii) POM image of  $\text{TBA}^+\text{-}34c^-$  at  $65^\circ\text{C}$  upon cooling (redrawn from ref. 67b. Copyright 2018 Wiley) and (b) (i) deprotonated dithienynitrophenols  $35a,b^-$  and single-crystal X-ray structures of (ii)  $\text{TBA}^+\text{-}35a^-$  and (iii)  $\text{TBA}^+\text{-}35b^-$ . The crystal structures are redrawn from cif files: 1830529, 2253150 and 2253154.

extended  $\pi$ -electronic system effectively stabilizes deprotonated pyrrole units (Fig. 22).<sup>69</sup> Pyrrole-bridged quinone undergoes deprotonation by treatment with tetraalkylammonium hydroxide, inducing a red-shifted absorption band of  $\sim 150\text{ nm}$  for  $36^-$  (Fig. 22a). In  $\text{TPA}^+\text{-}36^-$  ( $\text{TPA}^+$ : tetrapropylammonium),  $\text{TPA}^+$  is located on the  $\pi$ -plane of  $36^-$ , inhibiting the stacking of  $36^-$  (Fig. 22b).

The delocalization of the negative charge is essential to obtain stable  $\pi$ -electronic anions. Deprotonated species of *meso*-hydroxy-substituted porphyrins afford stable  $\pi$ -electronic anionic species  $37^-$  and  $37a-d^-$  ( $\text{Ni}^{\text{II}}$ ,  $\text{Zn}^{\text{II}}$ ,  $\text{Pd}^{\text{II}}$  and  $\text{Pt}^{\text{II}}$  complexes, respectively) (Fig. 23a).<sup>9,70</sup> The  $\text{pK}_a$  value of  $37a$  is estimated to be 7.89, higher than those of dipyrrolynitrophenols  $34a,b$ . The deprotonated structures of  $37a-d^-$  as  $\text{TBA}^+$  ion pairs exhibit the charge-by-charge columnar structures (Fig. 23b(i)). Furthermore, the hydroxy group serving as a directing group enables the selective introduction of phenyl groups at adjacent positions. Treatment of  $37a$  with  $\text{Pd}(\text{OAc})_2$ , *rac*-BINAP and  $\text{Cs}_2\text{CO}_3$  yields the Pd-BINAP-complex; further treatment with  $\text{PhMgBr}$  affords  $\beta$ -phenyl- and diphenyl-substituted  $38a,b$  (Fig. 23a).<sup>70b</sup> The  $\text{pK}_a$  values of 8.18 and 8.25 for  $38a,b$ , respectively, are higher than that of  $37a$ , confirming the electron-donating nature of the phenyl rings. Upon deprotonation, the ion pairs  $\text{TBA}^+\text{-}38a,b^-$

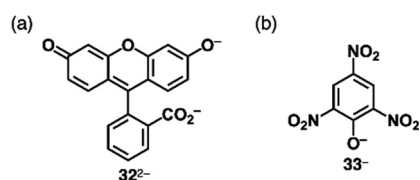


Fig. 20 (a) Uranine  $32^{2-}$  and (b) deprotonated picric acid  $33^-$ .

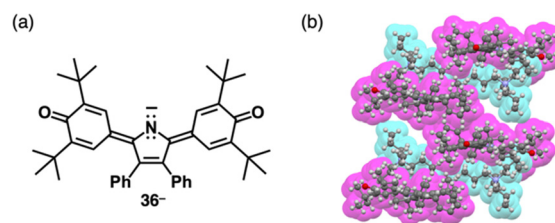
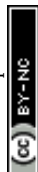


Fig. 22 (a) Deprotonated pyrrole-bridged quinone  $36^-$  and (b) single-crystal X-ray structure of  $\text{TBA}^+\text{-}36^-$  (redrawn from a cif file: 2076443).



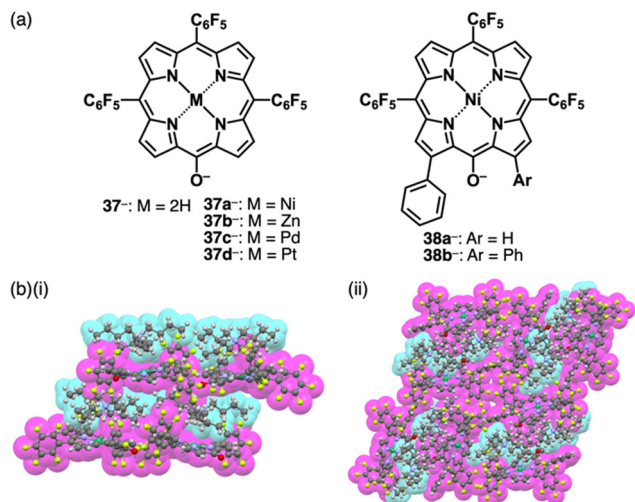


Fig. 23 (a) Deprotonated meso-hydroxyppyrrins **37<sup>-</sup>** and **37a-d<sup>-</sup>** (M = Ni, Zn, Pd and Pt) and  $\beta$ -monophenyl **38a<sup>-</sup>** and diphenyl-substituted **38b<sup>-</sup>** and (b) single-crystal X-ray structures of (i) TBA<sup>+</sup>-**37a<sup>-</sup>** and (ii) TBA<sup>+</sup>-**38b<sup>-</sup>** (redrawn from cif files: 1536339 and 1901215).

form charge-by-charge ion-pairing assemblies in the solid state (Fig. 23b(ii)). The introduction of suitable aryl units allows for the delocalization of the negative charge, resulting in  $\pi$ -extended anionic species with higher stability.

### 3. $\pi$ -Electronic systems with charge introduced by noncovalent interactions: ion complexes that sparked $\pi$ -electronic ion-pairing chemistry

Charge in  $\pi$ -electronic systems can be introduced by noncovalent interactions with inorganic ions, especially, anions (Fig. 1b(v)). Anion binding<sup>71</sup> of anion-responsive  $\pi$ -electronic systems (receptors) leads to the formation of pseudo- $\pi$ -electronic anions, which are incorporated into ion-pairing assemblies along with  $\pi$ -electronic cations, some of which are discussed in Section 2. Anion binding by  $\pi$ -electronic receptors transforms hard inorganic anions to soft “pseudo- $\pi$ -electronic” anions, resulting in charge delocalization within the host  $\pi$ -electronic systems (Fig. 24(i)). These formed anion complexes,

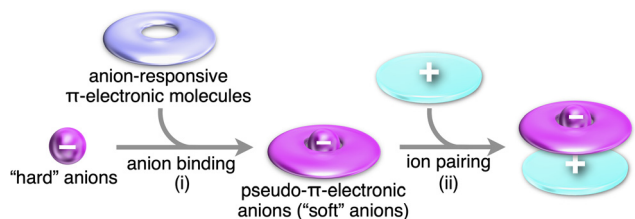


Fig. 24 Conceptual scheme for the formation of  $\pi$ -electronic ion pairs through (i) anion binding and (ii) ion pairing. Pseudo- $\pi$ -electronic anions as soft anions are obtained by the binding of a hard anion by  $\pi$ -electronic receptors. The pseudo- $\pi$ -electronic anions form ion pairs in combination with counter  $\pi$ -electronic cations.

acting as pseudo- $\pi$ -electronic anions, can be stacked with  $\pi$ -electronic cations through  $i\pi-i\pi$  interactions, primarily involving electrostatic and dispersion forces (Fig. 24(ii)). Various pseudo- $\pi$ -electronic anions can be generated based on the guest anions, with their counteranions serving as the counter species of the resulting anion complexes. The stacked pairs of anion complexes and counteranions, termed pseudo- $\pi$ -sips, are fundamental building blocks of ion-pairing assemblies. The focus of this section is on the anion complexes of anion-responsive  $\pi$ -electronic molecules, as the chemistry of ion-pairing assemblies originates from the structures assembled with alternately stacked anion complexes and  $\pi$ -electronic cations.<sup>5</sup>

#### 3.1. Charged $\pi$ -electronic systems formed by ion binding

Various types of anion-responsive molecules have been synthesized as colourimetric selective anion sensors and anion transporters in membranes.<sup>71</sup> However, only a few reports discuss planar receptor-anion complexes. The design of anion receptors using  $\pi$ -electronic systems is essential for the formation of planar receptor-anion complexes. Building subunits such as pyrrole and triazole have polarized structures with an H unit that acts as a hydrogen-bonding donor.

As pyrrole-based anion receptors that show planar structures, dipyrrolyldiketone BF<sub>2</sub> complexes (PBs, **39a-h** as examples) have two pyrrole units attached to the electron-withdrawing 1,3-propanedione unit (Fig. 25).<sup>5a,72</sup> The pyrrole NH units can be directed to the propanedione CH side by rotating the pyrrole rings, and, in the resulting conformation, the two NH and one CH units serve as anion-binding sites with high binding constants for various anions ( $K_a = 15\,000\text{ M}^{-1}$  for **39a** to bind Cl<sup>-</sup> in CH<sub>2</sub>Cl<sub>2</sub>).<sup>72c</sup> Peripheral modifications at the pyrrole and boron units facilitate further functionalization. A previous review article summarizes representative results in anion-binding and assembling behaviours.<sup>5a</sup>  $\alpha$ -Phenyl derivative **39d** (Fig. 25b) shows enhanced anion-binding ability owing to the additional *o*-CH as

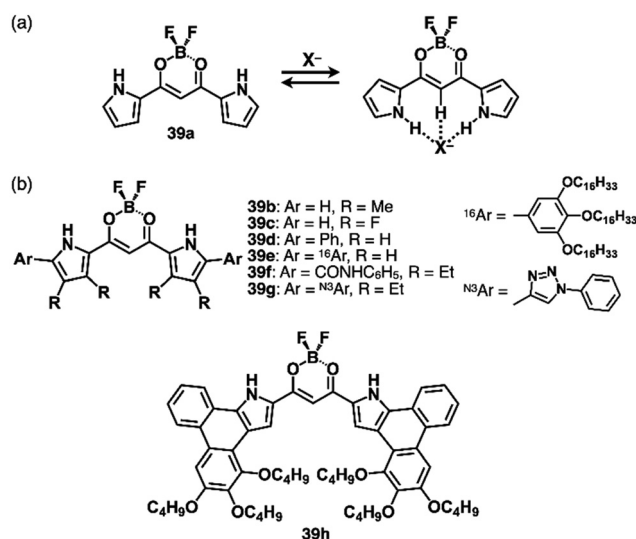


Fig. 25 (a) Anion-binding behaviour of dipyrrolyldiketone BF<sub>2</sub> complex (PB) **39a** and (b) PB derivatives **39b-h**.



supportive interaction sites.<sup>72b</sup> Single-crystal X-ray analysis of  $\text{TPA}^+ \cdot \text{39d} \cdot \text{Cl}^-$  clearly demonstrates the planar geometry of  $\text{39d} \cdot \text{Cl}^-$ . Importantly, the planar  $\text{39d} \cdot \text{Cl}^-$  and counter  $\text{TPA}^+$  are alternately stacked, forming a charge-by-charge columnar structure. The discovery of ion-pairing assemblies with alternately stacked anion complexes and  $\pi$ -electronic cations sparked the idea of incorporating planar counteranions (*vide infra*, Section 3.2).

Efficient anion-binding behaviours of PB have been applied for inducing further functionalities (Fig. 25b). Modifications of the  $\alpha$ -positions such as  $\alpha$ -amide-substituted **39f** exhibit higher anion-binding behaviour ( $K_a \geq 10^8 \text{ M}^{-1}$  for  $\text{Cl}^-$  in  $\text{CH}_2\text{Cl}_2$ ) to form solid-state ion-pairing assemblies.<sup>72g</sup> Moreover, click reactions of an  $\alpha$ -diethynyl-substituted receptor<sup>72f</sup> and arylazides afford triazole-substituted receptors including **39g**.<sup>72i</sup> The triazole-CH functions as an anion-binding site, forming planar receptor-anion complexes in ion-pairing assemblies. Covalently linked PB dimers show high anion-binding affinities by forming helical structures as building blocks of ion-pairing assemblies.<sup>72d,e,73</sup> OFF-ON fluorescence switching by anion binding has been demonstrated in hexaaryl-substituted PB derivatives based on anion-dependent intramolecular photo-induced electron transfer (PET), demonstrating potential logic gate systems.<sup>72j,k</sup> More efficient anion-binding and ion-pairing behaviours can be achieved by including the PB unit in the macrocycles owing to pyrrole-inverted preorganized structures.<sup>72h</sup>

Shape-persistent macrocycles with preorganized anion-binding structures based on  $\pi$ -electronic subunits facilitate the preparation of planar receptor-anion complexes.<sup>74</sup> Flood *et al.* synthesized a triazolophane anion receptor **40** (Fig. 26), containing polarized aryl-CH units as interaction sites, providing an anion-binding cavity.<sup>75</sup> By exclusively employing aryl-CH units, **40** demonstrates a highly efficient anion-binding capability ( $K_a = 4\,700\,000 \text{ M}^{-1}$  for  $\text{Cl}^-$  in  $\text{CH}_2\text{Cl}_2$ ). Moreover, **40** forms stacked [2+1]-type anion complexes with larger anions, such as  $\text{I}^-$ , through the effective  $\pi$ - $\pi$  stacking of the aromatic units present in **40**.

Cyanostar (CS) **41**, serving as another planar  $\pi$ -electronic anion receptor, incorporates five electron-withdrawing cyano groups, making the inner CH units highly efficient hydrogen-bonding donors (Fig. 26).<sup>76</sup> Owing to the presence of an internal anion-binding cavity with five CH units, CS exhibits high binding constants for larger anions, including  $\text{BF}_4^-$ ,  $\text{PF}_6^-$

and  $\text{ClO}_4^-$  ( $\log K_a \sim 5$ ). [2+1]-Type anion complexes for larger anions based on effective  $\pi$ - $\pi$  stacking of CS serve as key structures for fabricating ion-pairing assemblies.

### 3.2. $\pi$ -Electronic ion-pairing assemblies comprising anion receptors

Since the initial discovery of the solid-state charge-by-charge assembly of  $\text{TPA}^+ \cdot \text{39d} \cdot \text{Cl}^-$ , as illustrated in Section 3.1, there has been a trend of introducing more planar counteranions instead of the bulky ammonium cations for the formation of charge-by-charge assemblies. In our preliminary trial for the use of planar counteranions, tris(diethylamino)-substituted  $\text{TOTA}^{+15b}$  was incorporated in the solid-state charge-by-charge assembly with the anion complex of **39d**. Because of the difficulty in excluding  $\text{I}^-$  from the starting ion pair  $\text{TOTA}^+ \cdot \text{Cl}^-$  by anion exchange protocol, other suitable  $\pi$ -electronic cations have been explored to fabricate ion-pairing assemblies. For example,  ${}^3\text{TATA}^+ \cdot \text{Cl}^-$ , which can be synthesized from the  $\text{Cl}^-$  salt a tritylium derivative, was used for an alternative  $\pi$ -electronic cation.<sup>7</sup> Planar  $\text{39d} \cdot \text{Cl}^-$  and  ${}^3\text{TATA}^+$  formed a charge-by-charge assembly *via*  $\pi$ - $\pi$  interactions, with a shorter stacking distance than that of  $\text{TPA}^+ \cdot \text{39d} \cdot \text{Cl}^-$  (Fig. 27a). Furthermore,  ${}^3\text{TATA}^+ \cdot \text{39e} \cdot \text{Cl}^-$  with aliphatic alkyl chains forms a supramolecular gel comprising micrometre-scale fibrous morphologies (Fig. 27b). X-ray diffraction analysis of the xerogel revealed the formation of a  $\text{Col}_h$  structure with a diffraction peak at 0.74 nm, which is nearly twice of  $\pi$ - $\pi$  stacking distances. This distance was assigned as the repeating distance of identically charged species in the charge-by-charge assembly. In addition,  ${}^3\text{TATA}^+ \cdot \text{39e} \cdot \text{Cl}^-$  forms a liquid crystal mesophase based on the  $\text{Col}_h$  packing of charge-by-charge assembly.<sup>7</sup>

Moreover, porphyrin  $\text{Au}^{\text{III}}$  complexes have been used as counteranions of  $\pi$ -electronic ion-pairing assemblies comprising PB-based anion complexes. For example, in the crystal structure of  $22a^+ \cdot 39a \cdot \text{Cl}^-$ , a stacked  $22a^+$  dimer was sandwiched by planar  $39a \cdot \text{Cl}^-$  (Fig. 28a).<sup>77</sup> In contrast, in  $22a^+ \cdot 39b \cdot \text{Cl}^-$ ,  $22a^+$

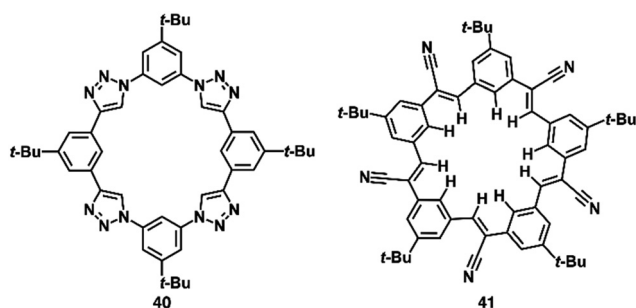


Fig. 26 Macrocyclic anion receptors: triazolophane **40** and cyanostar (CS) **41**.

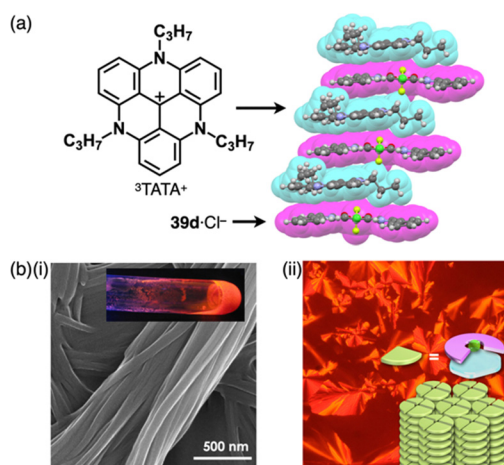
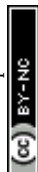


Fig. 27 (a) Single-crystal X-ray structure of  ${}^3\text{TATA}^+ \cdot \text{39d} \cdot \text{Cl}^-$  (redrawn from a cif file: 745780) and (b) (i) photograph and SEM of the supramolecular gel and (ii) POM and packing model of the liquid crystal mesophase of  ${}^3\text{TATA}^+ \cdot \text{39e} \cdot \text{Cl}^-$  (redrawn from ref. 7. Copyright 2010 Wiley).



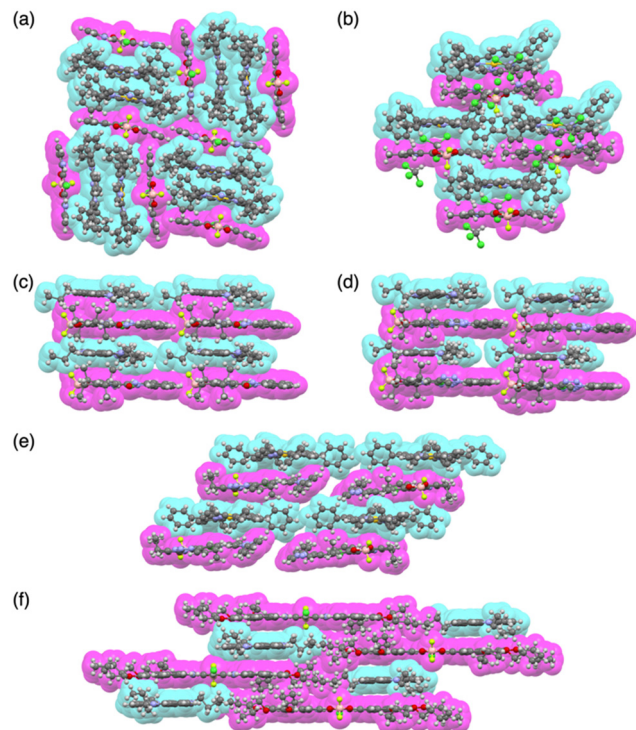


Fig. 28 Single-crystal X-ray structures of (a)  $22a^+-39a\cdot Cl^-$ , (b)  $22a^+-39b\cdot Cl^-$ , (c)  ${}^3TATA^+-39f\cdot Cl^-$ , (d)  ${}^3TATA^+-39g\cdot Cl^-$ , (e)  $22a^+-39g\cdot Cl^-$  and (f)  ${}^3TATA^+-39h\cdot Cl^-$  (redrawn from cif files: 1969168, 1969169, 1546127, 213826, 213827 and 1900650).

and planar  $39b\cdot Cl^-$  are alternately stacked with a  $\pi$ - $\pi$  stacking distance (Fig. 28b).  $22a^+-39c\cdot Cl^-$  forms a similar charge-by-charge assembly. Furthermore, charge-by-charge assemblies of  ${}^3TATA^+-39f\cdot Cl^-$ ,  ${}^3TATA^+-39g\cdot Cl^-$  and  $22a^+-39g\cdot Cl^-$  have been observed (Fig. 28c–e). Effective stacking of the planar receptor-anion complex and counteranion is supported by EDA, suggesting a cooperative contribution of electrostatic and dispersion forces in the stacked ion pairs of  ${}^3TATA^+-39g\cdot Cl^-$  and  $22a^+-39g\cdot Cl^-$ .<sup>72l</sup> Furthermore,  $\pi$ -extended PB  $39h$  exhibits a planar [1+1]-type  $Cl^-$  complex, showing the stacking with  ${}^3TATA^+$  in the solid state (Fig. 28f).<sup>72l</sup>

Recent studies based on the crystal analysis have revealed a variety of charge-by-charge assemblies depending on the constituting receptor-anion complexes and  $\pi$ -electronic counteranions. **42** forms a planar [2+1]-type anion complex  $42\cdot Cl^-$  in the crystal structure of  ${}^3TATA^+-42\cdot Cl^-$ , where the layers of identically charged species are alternately stacked (Fig. 29a).<sup>78</sup> Furthermore, introducing naphthalenediolate units instead of two fluorine moieties at the boron unit affects the assembling modes.<sup>79</sup> In the assembly of the naphthalene-2,3-diolate boron complex as a  ${}^3TATA^+$  ion pair  ${}^3TATA^+-43a\cdot Cl^-$ , the stacked  ${}^3TATA^+$  dimer is sandwiched by the naphthyl unit of  $43a\cdot Cl^-$ . Conversely, in  ${}^3TATA^+-43b\cdot Cl^-$ , alternately stacked  ${}^3TATA^+$  and the receptor- $Cl^-$  complex unit in  $43b\cdot Cl^-$  are observed (Fig. 29b). Square planar dipyrrolyldiketone  $Pt^{II}$  complexes that have phenylpyridine as another ligand part are also suitable for stacking assemblies (Fig. 30a).<sup>80</sup> In the crystals of  $Pt^{II}$  complex ion pairs  $22c^+-44a\cdot Cl^-$

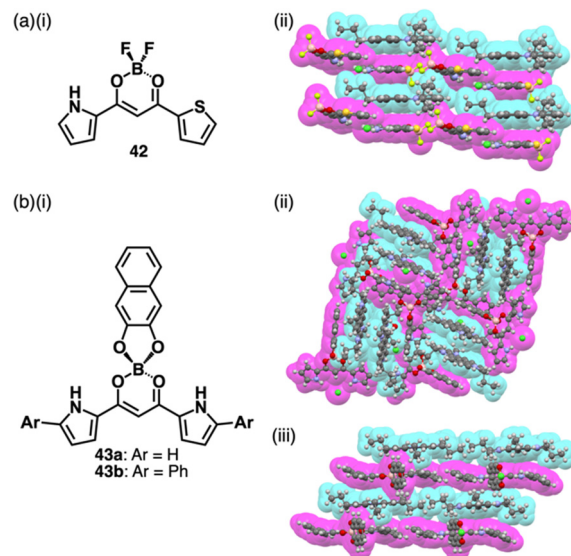


Fig. 29 (a) (i) Pyrrolythienyldiketone  $BF_2$  complex **42** and (ii) single-crystal X-ray structure of  ${}^3TATA^+-42\cdot Cl^-$  and (b) (i) naphthalenediolate boron complexes **43a,b** and (ii) single-crystal X-ray structures of  ${}^3TATA^+-43a\cdot Cl^-$  and (iii)  ${}^3TATA^+-43b\cdot Cl^-$ . The crystal structures are redrawn from cif files: 1840326, 2227398 and 2227399.

and  $22a^+-44b\cdot Cl^-$ ,  $Au^{III}$  complexes and anion complexes are alternately stacked and form columnar structures (Fig. 30b). These solid-state charge-by-charge assemblies, comprising the anion complexes of PB and  $Pt^{II}$  complexes, demonstrate the potential to induce diverse assembling modes and resulting properties based on neutral receptor units.

In the ion-pairing assemblies of CS, the  $4a^+-41_2\cdot BF_4^-$  ion pair is stacked with another ion pair, forming a charge-by-charge assembly, exhibiting a distinct red-shifted emission band derived from  $4a^+$ , in a concentrated solution (0.2 mM) as well as crystalline and film states.<sup>76b</sup> Various cationic dyes can be introduced as the counteranions of  $ClO_4^-$ , and the ion pairs form charge-by-charge assemblies in the crystalline state, as shown in  $45a^+-41_2\cdot ClO_4^-$  (Fig. 31a and b).<sup>76c</sup> Usually, crystal-state photophysical properties, different from those in solution, are affected by exciton coupling depending on the arrangement

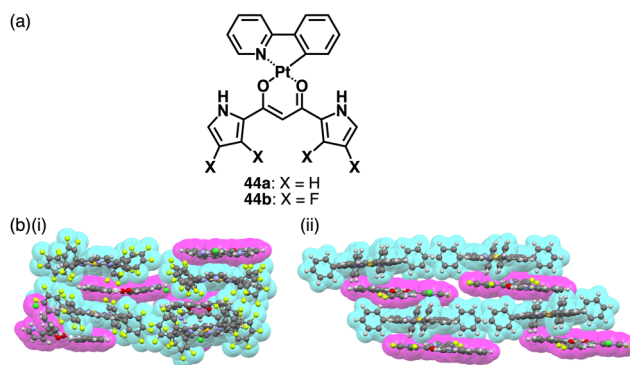
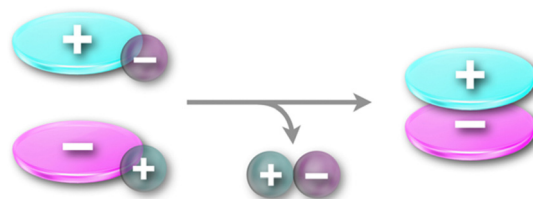


Fig. 30 (a) Dipyrrolyldiketone  $Pt^{II}$  complexes **44a,b** and (b) single-crystal X-ray structures of (i)  $22c^+-44a\cdot Cl^-$  and (ii)  $22a^+-44b\cdot Cl^-$  (redrawn from cif files: 2063433 and 2063434).



of  $\pi$ -electronic molecules. In contrast, the cationic dyes within the charge-by-charge assemblies with CS anion complexes are spatially and electronically isolated, and the exciton coupling is suppressed. Flood *et al.* proposed charge-by-charge assemblies as small-molecule ionic isolation lattices (SMILES). Ensuring that the frontier molecular orbitals of the cationic dyes are located inside those of the anion complex of CS is crucial. Using SMILES, the polymer resin containing  $45a^+ \cdot 41_2 \cdot ClO_4^-$  exhibits an emissive solid-state material (Fig. 31c).<sup>76c</sup> Furthermore,  $45b^+ \cdot 41_2 \cdot PF_6^-$  forms SMILES-based nanoparticles, exhibiting bright emission derived from  $45b^+$  upon excitation of  $41_2 \cdot PF_6^-$  owing to the effective energy transfer ( $\varphi_{ET} = 80\%$ ) between the oppositely charged  $\pi$ -electronic systems.<sup>76d</sup> A similar phenomenon was observed in the nanoparticles of  $46b^+ \cdot 41_2 \cdot PF_6^-$ .<sup>76e</sup> In particular, the energy transfer efficiency of  $46b^+ \cdot 41_2 \cdot PF_6^-$  reaches 100% owing to the larger overlap of emission of  $41_2 \cdot PF_6^-$  and the absorption bands of the cationic dye. However, the fluorescence quantum yield of SMILES is low (12.8%), owing to Förster resonance energy transfer to dark trap states or reabsorption. The deactivation path is suppressed by doping fluorescent dyes into SMILES comprising cationic dyes with higher excitation energies, drastically improving the fluorescent quantum yields (65% and 29% for  $45a^+ \cdot ClO_4^-$ -doped and pristine SMILES comprising  $46a^+ \cdot PF_6^-$ , respectively).<sup>76f</sup>

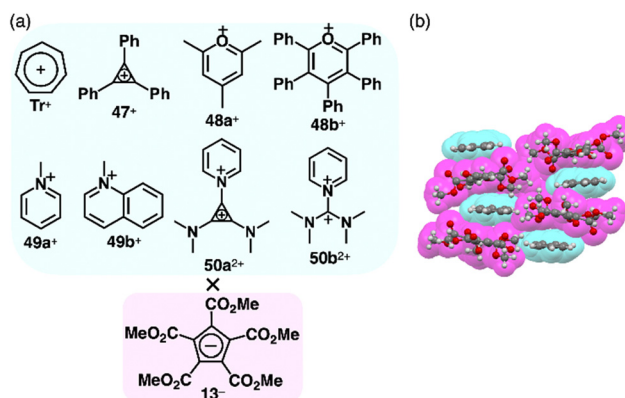
The  $\pi$ -electronic cations described in Section 2 are introduced as the counter species for the anion complexes of anion-responsive  $\pi$ -electronic molecules (Section 3). The combination of  $\pi$ -electronic cations with  $\pi$ -electronic anions, as summarized



**Fig. 32** Ion-pair metathesis for the preparation of  $\pi$ -electronic ion pairs.

#### 4.1. $\pi$ -Electronic ion pairs comprising cyclopentadienide-based $\pi$ -electronic anions

Photophysical properties and ion-pairing assemblies of  $\pi$ -electronic ion pairs were investigated based on cyclopentadienide **13**<sup>-</sup>.<sup>81</sup> In 1963, ion pairs of **13**<sup>-</sup> were prepared by ion-pair metathesis of  $\text{K}^+$ -**13**<sup>-</sup> and various  $\pi$ -electronic cations, such as  $\text{Tr}^+$  and **47**–**49**<sup>+</sup> as  $\text{BF}_4^-$  salts (Fig. 33a).<sup>81a</sup> The ion pairs exhibit countercation-dependent CT bands. The CT band of  $\text{Tr}^+$ -**13**<sup>-</sup> is blue-shifted in polar solvents (474 and 439 nm in  $\text{CH}_2\text{Cl}_2$  and MeOH, respectively) with a smaller solvent dependence than  $\text{Tr}^+$ - $\text{I}^-$  (571 and 403 nm, respectively).<sup>81b</sup> Alternately arranged  $\text{Tr}^+$  and **13**<sup>-</sup> form a charge-by-charge assembly in the crystal structure (Fig. 33b).<sup>81d</sup> The CT absorption band of an *N*-methylpyridinium ion pair **49a**<sup>+</sup>-**13**<sup>-</sup> at  $-170^\circ\text{C}$  in a MeOH/EtOH glassy solution was observed ( $\lambda_{\text{abs}} \sim 340$  nm as a shoulder); compared to other ion pairs with  $\text{Br}^-$ ,  $\text{N}_3^-$ ,  $\text{SCN}^-$  and  $\text{I}^-$ , it was red-shifted, whereas the emission band was blue-shifted.<sup>81c</sup>



**Fig. 33** (a)  $\pi$ -Electronic ion pairs of pentamethoxycarbonyl-substituted cyclopentadienide **13**<sup>−</sup> with  $\pi$ -electronic cations  $\text{Tr}^+$ , **47**–**49**<sup>+</sup> and **50a**,<sup>b,2</sup> and (b) single-crystal X-ray structure of  $\text{Tr}^+$ –**13**<sup>−</sup> (redrawn from a cif file: 1141949).

Therefore, the Stokes shift of  $49a^{+} \cdot 13^{-}$  ( $1.1 \times 10^4 \text{ cm}^{-1}$ ) is smaller than those of the other ion pairs ( $> 1.55 \times 10^4 \text{ cm}^{-1}$ ), ascribed to the smaller structural relaxation in the excited state. Ion pairs of dicationic  $50a^{2+}$  and  $13^{-}$ ,  $50a^{2+} \cdot 13^{-} \cdot X^{-}$  ( $X^{-} = OTf^{-}$  and  $BF_4^{-}$ ) and  $50a^{2+} \cdot 213^{-}$ , were prepared by the ion-pair metathesis of  $50a^{2+} \cdot 2X^{-}$  and  $K^{+} \cdot 13^{-}$ .<sup>81e</sup> Similarly,  $50b^{2+}$  provided ion pairs with  $13^{-}$  in the presence and absence of  $BF_4^{-}$  by the ion-pair metathesis between  $50b^{2+} \cdot 2BF_4^{-}$  and  $K^{+} \cdot 13^{-}$ . The ion pairs of  $50a^{2+}$  exhibit concentration-dependent CT bands ( $\lambda_{\text{abs}} = 406$  and  $424 \text{ nm}$  at  $1.65$  and  $3.30 \text{ M}$ , respectively, in MeCN for  $50a^{2+} \cdot 213^{-}$ ). In addition,  $^1\text{H}$  NMR signals of  $50a^{2+} \cdot 2OTf^{-}$  are shifted upfield in the presence of  $K^{+} \cdot 13^{-}$ , suggesting an interaction between  $13^{-}$  and  $50a^{2+}$ . In the crystal structure of  $50a^{2+} \cdot 13^{-} \cdot BF_4^{-}$ , the dication and  $13^{-}$  are alternately stacked on the cyclopropenyl unit rather than the pyridinium unit, forming a charge-by-charge assembly with  $BF_4^{-}$ , located between the columns.<sup>81e</sup> The  $\text{CO}_2\text{Me}$  groups in  $13^{-}$  are distorted from the cyclopentadienide core owing to their steric repulsion.<sup>81d,e</sup>

The planar structure of  $\text{PCCp}^{-}$  is suitable for more effective stacking of oppositely charged species in combination with  $\pi$ -electronic cations (Fig. 34a). An ion pair of  $\text{PCCp}^{-}$  with trityl cation  $1^{+}$  was prepared as the precursor of triethylsilyl  $\text{PCCp}^{-}$  by the ion-pair metathesis of  $1^{+} \cdot \text{Br}^{-}$  and  $\text{Ag}^{+} \cdot \text{PCCp}^{-}$ .<sup>82</sup>  $1^{+}$  and  $\text{PCCp}^{-}$  afforded a columnar structure comprising identically charged species in a crystalline state (Fig. 34b(i)).  $\text{PCCp}^{-}$  is stacked with another  $\text{PCCp}^{-}$  at an interplanar distance of  $3.42 \text{ \AA}$  to form a pair, which is slip-stacked with other pairs at the cyano groups.<sup>82</sup> In  $2^{+} \cdot \text{PCCp}^{-}$ ,  $2^{+}$  and  $\text{PCCp}^{-}$  are alternately stacked at the two aminophenyl units of  $2^{+}$  (Fig. 34b(ii)).<sup>33</sup> Meanwhile, the other aminophenyl group is stacked with the corresponding part of the neighbouring  $2^{+}$ , resulting in a partially charge-segregated assembly. Meanwhile, the propeller geometries of triarylmethyl cations inhibits effective stacking, resulting in the limited contact with the oppositely charged species. On the other hand,  $\text{PCCp}^{-}$  in  $\pi$ -electronic ion pairs with more planar  $\pi$ -electronic cations such as  $\text{Tr}^{+}$  and  $^3\text{TATA}^{+}$  is effectively stacked with the counteranions. In  $\text{Tr}^{+} \cdot \text{PCCp}^{-}$ ,  $\text{Tr}^{+}$  and  $\text{PCCp}^{-}$  are alternately stacked to afford a charge-by-charge assembly (Fig. 34b(iii)). In contrast, in  $^3\text{TATA}^{+} \cdot \text{PCCp}^{-}$ ,  $^3\text{TATA}^{+}$  and  $\text{PCCp}^{-}$  are stacked with the respective identically charged species to form stacked dimers, resulting in a two-by-two charge-by-charge assembly (Fig. 34b(iv)).<sup>33</sup> The ion pair of  $\text{PCCp}^{-}$  with  $51^{+}$ , an isoelectronic cation of  $\text{TATA}^{+}$  with a B-N unit in place of a central C-C unit, was also prepared by ion-pair metathesis of  $51^{+} \cdot \text{OTf}^{-}$  and  $\text{Na}^{+} \cdot \text{PCCp}^{-}$ .<sup>83</sup> The crystal structure resembles that of  $^3\text{TATA}^{+} \cdot \text{PCCp}^{-}$ , although each component is slip-stacked with the identically charged species (Fig. 34b(v)). Notably,  $^3\text{TATA}^{+}$  and  $51^{+}$  are stacked with  $\text{PCCp}^{-}$  in relatively short contact ( $3.29$  and  $3.25 \text{ \AA}$ , respectively).

Porphyrin  $\text{Au}^{\text{III}}$  complexes, which are discussed in Sections 2.2 and 3.2, have been adopted as the components of  $\pi$ -electronic ion pairs, as seen in  $22a^{+} \cdot \text{PCCp}^{-}$ .<sup>52</sup> The ion pair forms a solid-state charge-by-charge assembly with the interplanar distances of  $3.37$  and  $3.40 \text{ \AA}$  (Fig. 34b(vi)).<sup>52a</sup>  $22e^{+} \cdot \text{PCCp}^{-}$  is also included in a similar charge-by-charge assembly, where the two free *meso*-positions are in contact with  $\text{PCCp}^{-}$  CN units.<sup>52b</sup> Conversely,  $22c^{+} \cdot \text{PCCp}^{-}$  forms a one-by-two charge-

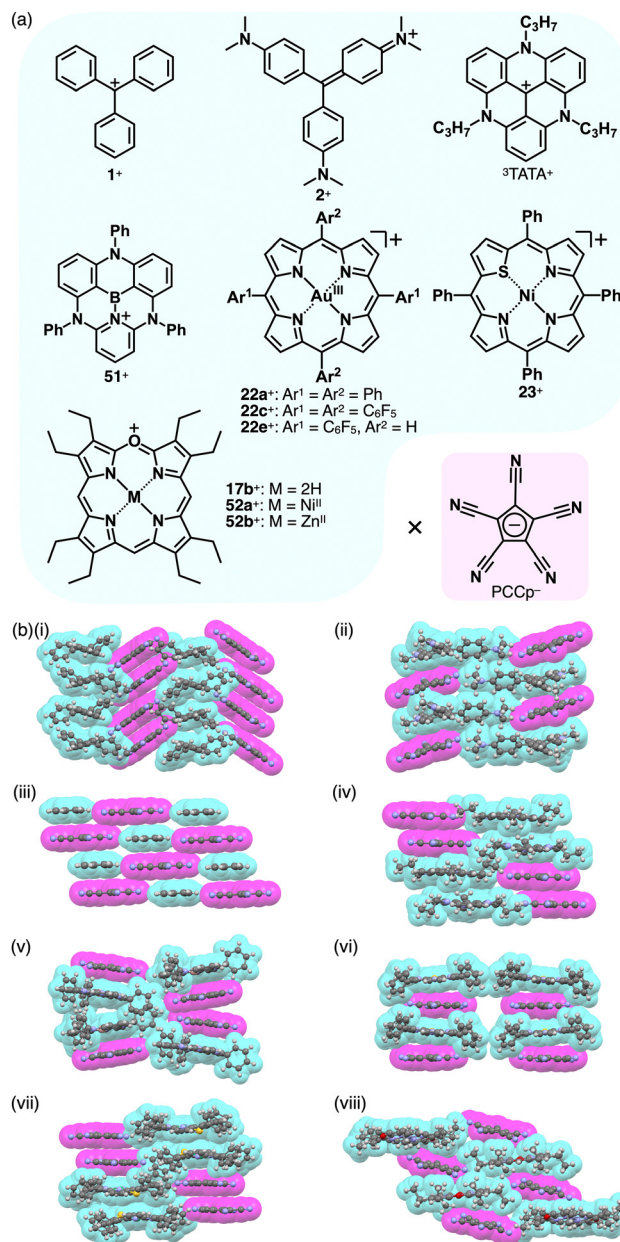


Fig. 34 (a) Combinations of  $\pi$ -electronic ion pairs:  $\text{PCCp}^{-}$  with  $1^{+}$ ,  $2^{+}$ ,  $^3\text{TATA}^{+}$ , BN-TATA  $51^{+}$ ,  $22a, c, e^{+}$ ,  $23^{+}$ ,  $17b^{+}$  and  $52a, b^{+}$  and (b) single-crystal X-ray structures of (i)  $1^{+} \cdot \text{PCCp}^{-}$ , (ii)  $2^{+} \cdot \text{PCCp}^{-}$ , (iii)  $\text{Tr}^{+} \cdot \text{PCCp}^{-}$ , (iv)  $^3\text{TATA}^{+} \cdot \text{PCCp}^{-}$ , (v)  $51^{+} \cdot \text{PCCp}^{-}$ , (vi)  $22a^{+} \cdot \text{PCCp}^{-}$ , (vii)  $23^{+} \cdot \text{PCCp}^{-}$  and (viii)  $52b^{+} \cdot \text{PCCp}^{-}$  (redrawn from cif files: 227124, 1431734, 1431733, 1431735, 2063989, 1877990, 2167304 and 2144318). Atom colour code in (viii): violet in the ball-and-stick models refers to zinc.

by-charge assembly with another  $22c^{+}$  unit located between the columns. Asymmetric heteroatom-containing porphyrin analogues with chalcogen elements has been adopted as the counteranion of  $\text{PCCp}^{-}$ . 21-Thiaporphyrin  $\text{Ni}^{\text{II}}$  complex ion pair  $23^{+} \cdot \text{PCCp}^{-}$  was prepared by ion-pair metathesis of  $23^{+} \cdot \text{Cl}^{-}$  and  $\text{Na}^{+} \cdot \text{PCCp}^{-}$ .<sup>55</sup> In the crystal structure of  $23^{+} \cdot \text{PCCp}^{-}$ , the thiophene unit of  $23^{+}$  is inclined by  $24.5^{\circ}$ . The  $23^{+}$  units are stacked by themselves in antiparallel orientation to form a dimer *via* chalcogen bonding between the thiophene and adjacent pyrrole



units at short S...N distances of 3.26 and 3.33 Å. In addition,  $\text{PCCp}^-$  forms a stacked dimer, providing a two-by-two charge-by-charge assembly (Fig. 34b(vii)).<sup>55</sup> Similarly, in the  $\text{PCCp}^-$  ion pair with oxaporphyrin  $\text{Ni}^{\text{II}}$  complex  $52\text{a}^+$ , the constituent ions form the respective stacked dimers with interplanar distances of 3.31 and 3.28 Å, respectively.<sup>84</sup> Besides,  $\text{PCCp}^-$  ion pairs of free-base  $17\text{b}^+$  and  $\text{Zn}^{\text{II}}$  complex  $52\text{b}^+$  form one-by-one charge-by-charge assemblies. In contrast to the almost parallel arrangement observed in  $17\text{b}^+$  and  $\text{PCCp}^-$ , in the ion pair  $52\text{b}^+\text{-PCCp}^-$ , the components are not arranged in parallel owing to the coordination of  $\text{PCCp}^-$  CN to the  $\text{Zn}^{\text{II}}$  centre (Fig. 34b(viii)).

The introduction of aliphatic alkyl chains into charged  $\pi$ -electronic species induces various soft materials based on ion-pairing assemblies. The porphyrin  $\text{Au}^{\text{III}}$  complex bearing hexadecyloxy chains at the *meso*-aryl moieties  $22\text{f}^+$  provides supramolecular gels and discotic columnar liquid crystals, whose assembled structures depend on the counteranions (Fig. 35).<sup>52a</sup>  $22\text{f}^+\text{-PCCp}^-$  forms a supramolecular octane gel (10 mg  $\text{mL}^{-1}$ ), owing to the entanglement of fibrous morphologies with diameters of 1–3  $\mu\text{m}$  and lengths exceeding 100  $\mu\text{m}$  (Fig. 35b(i)). Conversely,  $22\text{f}^+\text{-PCCp}^-$  exhibits a liquid crystal mesophase (36–293 °C) based on the  $\text{Col}_\text{h}$  structure of the charge-by-charge assembly, as observed in the crystal structure of  $22\text{a}^+\text{-PCCp}^-$  (mentioned earlier) (Fig. 35b(ii)). Meanwhile,  $22\text{f}^+\text{-Cl}^-$  forms a  $\text{Col}_\text{h}$  structure with charge-segregated assembly, excluding the nonplanar anion from the stacked structure of  $22\text{f}^+$ . These findings suggest that the size of  $\text{PCCp}^-$  complements the core part of porphyrin  $\text{Au}^{\text{III}}$  complexes, facilitating the formation of charge-by-charge assemblies. Notably, the charge-by-charge assembly of  $22\text{g}^+\text{-PCCp}^-$ , which possesses icosyloxy chains, shows anisotropic alignment after shearing. The alignment persists after thermal phase transitions, indicating a robust packing state of charge-by-charge assemblies.<sup>52a</sup>

In the previous paragraphs in this section, the  $\pi$ -electronic ion pairs of  $\text{PCCp}^-$  form the charge-by-charge assemblies, some of which comprise the stacked dimers of  $\text{PCCp}^-$  and those of the cations. As seen in the ion pairs with bulky cations, stacking

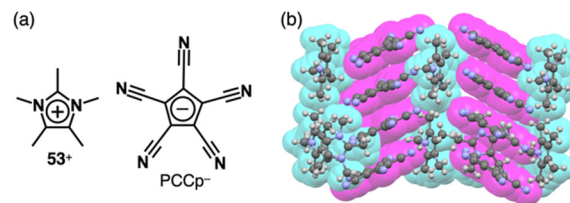


Fig. 36 (a) Ion pair of pentamethylimidazolium  $53^+$  with  $\text{PCCp}^-$  and (b) single-crystal X-ray structure of  $53^+\text{-PCCp}^-$  (redrawn from a cif file: 1868291).

structures of  $\text{PCCp}^-$  can be realized by combining  $\pi$ -electronic cations that aggregate independently. For instance, in the crystal structure of the ion pair comprising  $\text{PCCp}^-$  with pentamethylimidazolium  $53^+$ ,  $\text{PCCp}^-$  is stacked with another  $\text{PCCp}^-$ , leading to a charge-segregated assembly (Fig. 36).<sup>85</sup> In this arrangement,  $53^+$  is located between the  $\text{PCCp}^-$  columns without participating in the stacking. Despite the absence of long alkyl chains, the crystal of  $53^+\text{-PCCp}^-$  undergoes a phase transition and transforms to the mesophase in a temperature range of 234–240 °C. Interestingly, the ion pair, especially when doped with  $\text{Na}^+\text{-PCCp}^-$ , exhibits ionic conductivities.

Charge-segregated assemblies based on stacked structures of  $\text{PCCp}^-$  and those of  $\pi$ -electronic cations can be constructed by introducing the cationic species that induce specific interactions. The dipole of cationic squarylium  $10^+$  can be used for stacking in the  $\text{PCCp}^-$  ion pair (Fig. 37a).<sup>23</sup> The photophysical properties of  $10^+\text{-PCCp}^-$  vary with the stacking modes, forming J- and H-aggregates in 1%  $\text{CH}_2\text{Cl}_2/n$ -hexane and 1% acetone/water, respectively, while a monomeric state is observed in  $\text{CH}_2\text{Cl}_2$  (Fig. 37b). Additionally, J-aggregates are formed in chiral media (0.33% to 1%  $\text{CH}_2\text{Cl}_2/\alpha$ - and  $\beta$ -pinene) and exhibit CD signals, indicating the formation of chiral assemblies. These findings highlight the sensitivity of ion-pairing behaviour to the surrounding environment, offering potential applications in chiral recognition and sensing.  $10^+\text{-PCCp}^-$  gives two crystal pseudo-polymorphs: (i) a charge-segregated assembly ( $10^+\text{-PCCp}^-_{\text{CS}}$ ) (Fig. 37c(i)) and (ii) a partially charge-segregated assembly ( $10^+\text{-PCCp}^-_{\text{SS}}$ ) (Fig. 37c(ii)), where  $10^+$  is partially slip-stacked; hence, the direct contact of identically charged species is smaller than that of  $10^+\text{-PCCp}^-_{\text{CS}}$ . The EDA of the  $10^+\text{-PCCp}^-_{\text{CS}}$  packing structure indicates that electrostatic force in a stacked pair of  $10^+$  (c2–c3 in Fig. 37d), oriented in an antiparallel fashion, is smaller than that in a separated pair of  $10^+$  (c1–c3 in Fig. 37d). The peculiar electrostatic forces contributing to the characteristic assembling behaviour are ascribed to dipole-dipole interactions between identically charged species. Solid-state absorption spectra of the  $10^+$  ion pairs exhibit counteranion-dependent bands derived from the exciton coupling between stacked  $10^+$  units. Flash-photolysis (FP) TRMC measurements of the crystalline states of  $10^+\text{-PCCp}^-_{\text{CS}}$  and  $10^+\text{-PCCp}^-_{\text{SS}}$  reveal that the former exhibits higher electric conductivity ( $\varphi\sum\mu = 3.0 \times 10^{-8} \text{ m}^2 \text{ V}^{-1} \text{ s}^{-1}$ , where  $\varphi$  is the photo-carrier generation yield and  $\sum\mu$  is the sum of hole and electron mobilities) than the latter ( $\varphi\sum\mu = 1.0 \times 10^{-8} \text{ m}^2 \text{ V}^{-1} \text{ s}^{-1}$ ), confirming the effective electric conducting properties in the charge-segregated assembly (Fig. 37e). Moreover,  $10^+\text{-PCCp}^-_{\text{CS}}$

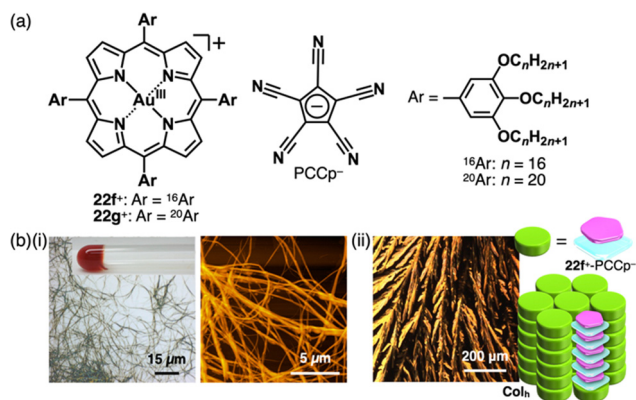
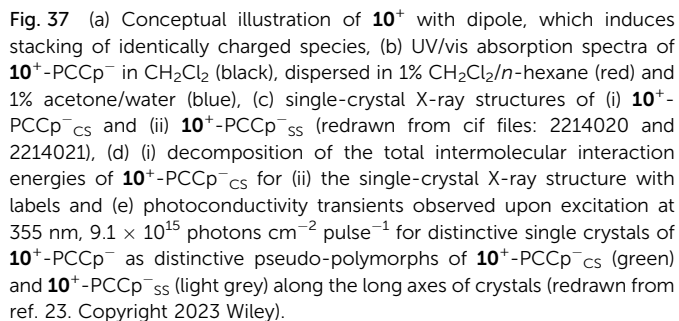


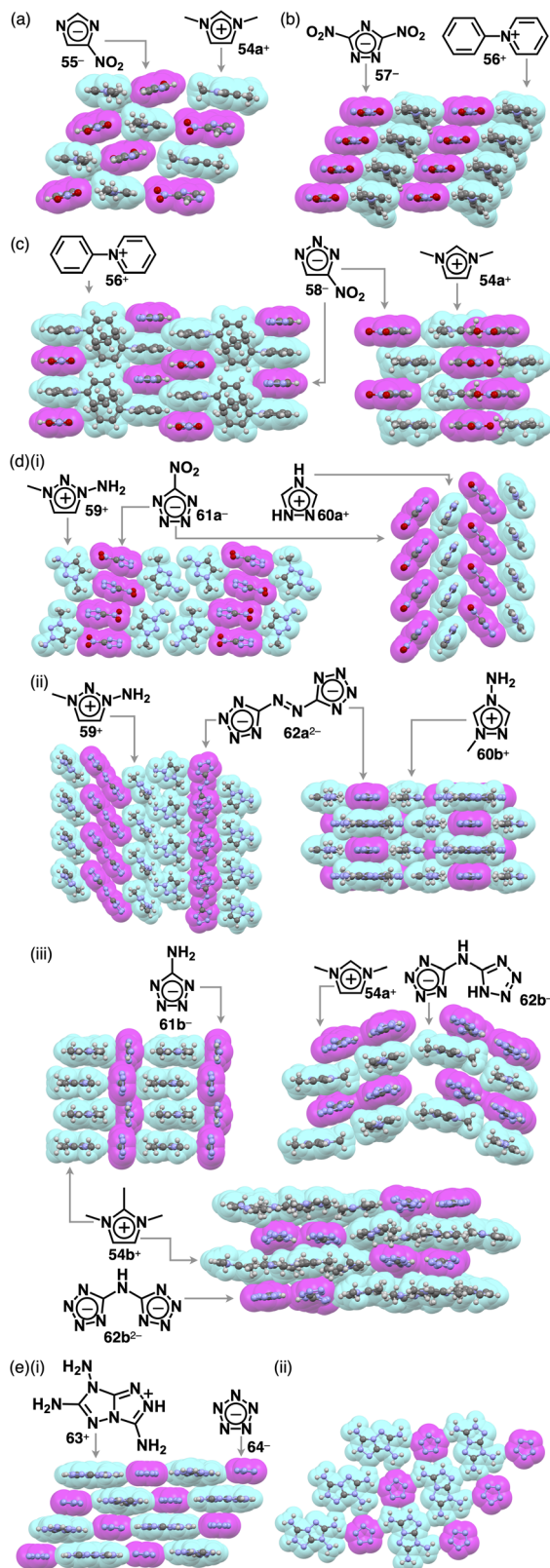
Fig. 35 (a) Ion pair of aliphatic porphyrin  $\text{Au}^{\text{III}}$  complex  $22\text{f}^+$  with  $\text{PCCp}^-$  and (b) (i) optical microscope and atomic force microscope images of  $22\text{f}^+\text{-PCCp}^-$  octane gel (10 mg  $\text{mL}^{-1}$ ) and (ii) mesophase properties of  $22\text{f}^+\text{-PCCp}^-$ : POM and a proposed assembled model at 280 °C upon cooling from the isotropic liquid state (redrawn from ref. 52a. Copyright 2019 Elsevier).





The replacement of CHs with electronegative nitrogens in cyclopentadienides enhances their stability, allowing for the production of various  $\pi$ -electronic anions, even with electron-donating groups. Azolium–azolate ion pairs, investigated by

The ion pairs of a tetrazolate dimer  $62a^{2-}$  were synthesized using ion-pair metathesis of triazolium iodide with  $Ag_2SO_4$ , followed by another metathesis with a  $Ba^{2+}$  salt of a tetrazolate dimer.<sup>87d,g</sup> The resulting  $259^+-62a^{2-}$  ion pair shows  $59^+$  located beside  $62a^{2-}$  owing to the presence of hydrogen bonding between the amino group of  $59^+$  and a tetrazole N of  $62a^{2-}$  (Fig. 38d(ii)). In addition,  $62a^{2-}$  is slip-stacked with adjacent  $62a^{2-}$ , giving a tilted anionic columnar structure.<sup>87g</sup> Conversely, in the crystal structure of  $260b^+-62a^{2-}$ , two  $60b^+$  units are located at both sides of the diazo group of  $62a^{2-}$  (Fig. 38d(ii)).<sup>87d</sup> The ion pair comprising  $60b^+$  and  $62a^{2-}$  forms a columnar structure as they are stacked alternately in a short interplanar distance of 3.04 Å. In the case of  $54b^+-61b^-$ ,  $61b^-$  arranged at the side of adjacent  $61b^-$  through hydrogen-bonding interaction results in a belt-like structure of  $61b^-$  and a stacked structure of  $54b^+$  (Fig. 38d(iii)).<sup>87f</sup> Ion pairs  $54a^+-62b^-$  and  $54b^+-62b^-$  were prepared by treating bis(tetrazolyl)amine, the protonated form of  $62b^-$ , with  $AgNO_3$ , followed by ion-pair metathesis with  $54a,b^+-I^-$ .<sup>87i</sup> A single crystal of  $254b^+-62b^{2-}$  was obtained by recrystallizing  $54b^+-62b^-$ , the corresponding monoanionic ion pair, from the MeOH/EtOAc solution. In the crystal structures of  $54a^+-62b^-$  and  $254b^+-62b^{2-}$ ,  $62b^-$  and  $62b^{2-}$  form hydrogen-bonding dimers, exhibiting alternate

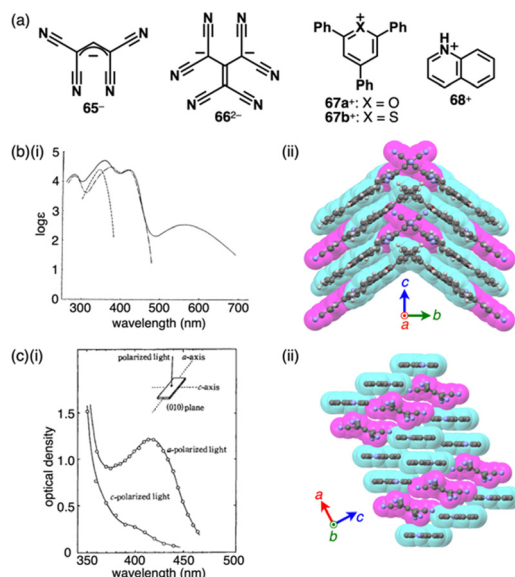


**Fig. 38** Single-crystal X-ray structures of ion pairs comprising (a) imidazolate **55**<sup>−</sup>, (b) 1,2,4-triazolate **57**<sup>−</sup>, (c) 1,2,3-triazolate **58**<sup>−</sup>, (d) tetrazolate (i) **61a**<sup>−</sup>, (ii) **62a**<sup>2−</sup> and (iii) **61b**<sup>−</sup>, **62b**<sup>−</sup> and **62b**<sup>2−</sup> and (e) pentazolate **64**<sup>−</sup> as (i) side and (ii) top views (redrawn from cif files: 902237, 733829, 733828, 902240, 837299, 280499, 837300, 265160, 855087, 913585, 913586 and 1973518).

stacking between the anion dimers and the coexisting two or four cation units, respectively. Additionally, in **54a**<sup>+</sup>-**62b**<sup>-</sup>, the **62b**<sup>-</sup> dimer is partially stacked with another dimer, while in **254b**<sup>+</sup>-**62b**<sup>2-</sup>, the **62b**<sup>2-</sup> dimer is surrounded by **54b**<sup>+</sup>.<sup>87i</sup> Furthermore, pentazolate, cyclopentadienide analogue, with all CHs replaced by nitrogen, was also reported as  $\pi$ -electronic ion pairs. In **63**<sup>+</sup>-**64**<sup>-</sup>, all the amino groups of **63**<sup>+</sup> and nitrogen of **64**<sup>-</sup> participated in hydrogen bonding, providing a two-dimensional hydrogen-bonding network further stacked *via*  $\pi$ - $\pi$  interactions (Fig. 38e).<sup>87j</sup> Hydrogen-bonding interaction often affects the assembling modes of ion-pairing assemblies, resulting in segregated columnar structures consisting of identically charged species.

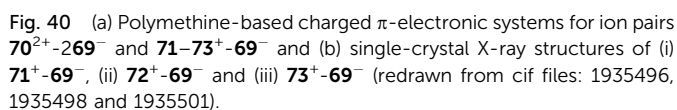
#### 4.3. $\pi$ -Electronic ion pairs comprising polymethine-based anions

Monomethine-based  $\pi$ -electronic anions **65**<sup>−</sup> and **66**<sup>2−</sup> were found to form  $\pi$ -electronic ion-pairing assemblies when combined with  $\pi$ -electronic cations (Fig. 39a).<sup>88–90</sup> The ion pairs of triphenyl(thio)pyryliums **67a,b**<sup>+</sup> with tetracyanoallyl anion **65**<sup>−</sup> were synthesized by ion-pair metathesis of the ClO<sub>4</sub><sup>−</sup> salts of the cations with Na<sup>+</sup>·**65**<sup>−</sup>.<sup>88a</sup> In CHCl<sub>3</sub>, **67a,b**<sup>+</sup>·**65**<sup>−</sup> exhibits CT bands at 570 and 595 nm, respectively (Fig. 39b(i)). The CT bands exhibited negative solvatochromism, which is well-known for zwitterionic systems without electronically neutral resonance structures such as Reichardt's dyes.<sup>91</sup> The difference in the redox potentials of the cations correlates with the



**Fig. 39** Monomethine-based cyanocarbon anions and the counteranions, (b) (i) UV/vis absorption spectra of **67a<sup>+</sup>-65<sup>-</sup>** (solid line), **67a<sup>+</sup>-ClO<sub>4</sub><sup>-</sup>** (chain line) and **Na<sup>+</sup>-65<sup>-</sup>** (dashed line) in CHCl<sub>3</sub> (redrawn from ref. 88a. Copyright 1970 CSJ) and (ii) single-crystal X-ray structure of **67a<sup>+</sup>-65<sup>-</sup>** and (c) (i) polarized absorption spectra of **268<sup>+</sup>-66<sup>2-</sup>** along with the direction of the long and short axes (redrawn from ref. 89a. Copyright 1969 CSJ) and (ii) single-crystal X-ray structure of **268<sup>+</sup>-66<sup>2-</sup>**. The long axes of single crystals of **67a<sup>+</sup>-65<sup>-</sup>** and **268<sup>+</sup>-66<sup>2-</sup>** corresponded to the c and a axes in (ii), respectively. The crystal structures are redrawn from cif files: 1240563 and 1245498.

$\pi$ -Electronic ion pairs of further extended oppositely charged polymethines **74-76<sup>-</sup>** and **77-80<sup>+</sup>** were investigated for the effects on the spectral and assembling features (Fig. 41a).<sup>92-94</sup> Tatikolov *et al.* prepared the ion pairs of oppositely charged polymethine dyes. The absorption spectrum of **77<sup>+</sup>-74a<sup>-</sup>** in



CH<sub>2</sub>Cl<sub>2</sub> is almost identical to the summed spectra of 77<sup>+</sup>-Br<sup>-</sup> and TBA<sup>+</sup>-74a<sup>-</sup>.<sup>92</sup> Similarly, the absorption maxima of 78b<sup>+</sup>-75b<sup>-</sup> are similar to those of the components.<sup>93</sup> In contrast, 78a<sup>+</sup>-75a<sup>-</sup>, whose components show overlapped absorptions in toluene at ~560–580 nm, exhibits a characteristic absorption band at ~520 nm, indicating interchromophoric interactions (Fig. 41b). Theoretical calculations supported the notably blue-shifted absorption bands of 78a<sup>+</sup>-75a<sup>-</sup>, while showing negligible changes in the case of 78b<sup>+</sup>-75b<sup>-</sup>.<sup>93</sup> Photovoltaic cell based on 77<sup>+</sup>-74a<sup>-</sup> shows significantly higher external quantum efficiency compared to that based on Na<sup>+</sup>-74a<sup>-</sup> (Fig. 41c).<sup>92</sup> Focusing on assembled structures, in 79<sup>+</sup>-76<sup>-</sup> forming the charge-segregated assembly, 79<sup>+</sup> and 76<sup>-</sup> are independently stacked in brickwork-like and columnar structures, respectively (Fig. 41d(i)).<sup>94</sup> In contrast, the stacking of identical species is hindered in 80<sup>+</sup>-74b<sup>-</sup> owing to the introduction of a phenyl group at the centre of the polymethine chains, resulting in partial stacking between the oppositely charged species at their terminal  $\pi$ -electronic systems (Fig. 41d(ii)).

In Sections 4.2 and 4.3,  $\pi$ -electronic anions that form different types of  $\pi$ -electronic ion pairs are discussed. These anions are obtained through deprotonation of the corresponding  $\pi$ -electronic systems. Similarly,  $\pi$ -electronic dianions can be prepared by removing protons attached to the  $\pi$ -electronic systems, resulting in the formation of ion pairs and assemblies. By employing ion-pair metathesis between three different deprotonated anions (eosine  $2\text{Na}^+ \mathbf{81a}^{2-}$ , erythrosine  $2\text{Na}^+ \mathbf{81b}^{2-}$  and

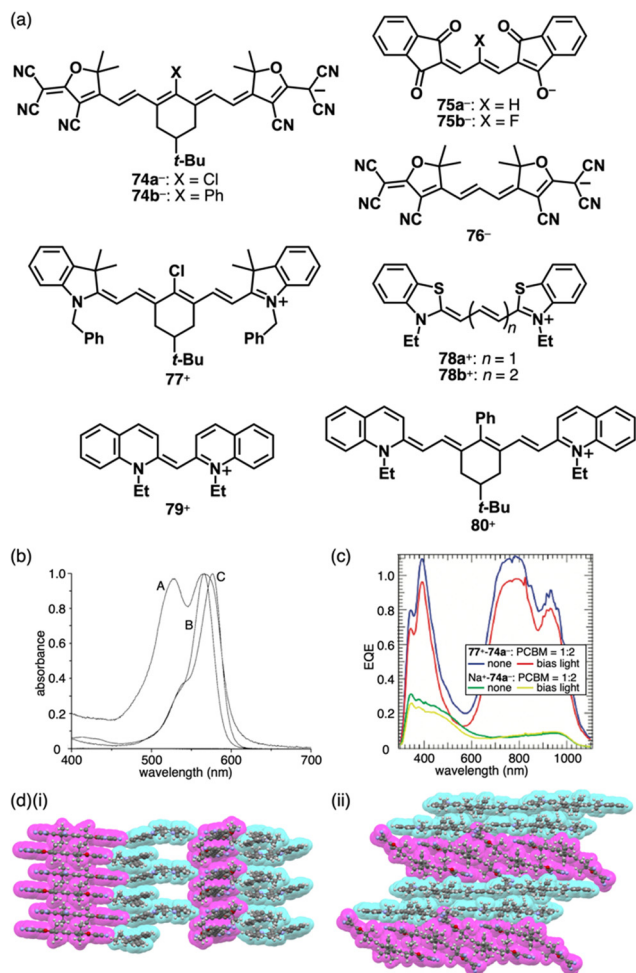


Fig. 41 (a) Extended polymethine-based  $\pi$ -electronic anions **74**<sup>−</sup>–**76**<sup>−</sup> and the counteranions **77**<sup>+</sup>–**80**<sup>+</sup>, (b) UV/vis absorption spectra of **78a**<sup>+</sup>–**75a**<sup>−</sup> (A), 1,3-bis(dimethylamino)propenium-**75a**<sup>−</sup> (B) and **78a**<sup>+</sup>–I<sup>−</sup> (C) in toluene (redrawn from ref. 93. Copyright 2001 ACS), (c) external quantum efficiency (EQE) of bulk heterojunction solar cells fabricated with ion pairs **77**<sup>+</sup>–**74a**<sup>−</sup> (blue, red) and Na<sup>+</sup>–**74a**<sup>−</sup> (green, yellow) blended with [60]PCBM in a 2 : 1 weight ratio (for comparison, the EQE spectra measured with and without white light bias are shown) (redrawn from ref. 92. Copyright 2009 ACS) and (d) single-crystal X-ray structures of (i) **79**<sup>+</sup>–**76**<sup>−</sup> and (ii) **80**<sup>+</sup>–**74b**<sup>−</sup> (redrawn from cif files: 1910541 and 1910529).

rose bengal 2Na<sup>+</sup>–**82**<sup>2−</sup>) and three different cations (crystal violet 2<sup>+</sup>–Cl<sup>−</sup> (Section 2.1), malachite green **83**<sup>+</sup>–Cl<sup>−</sup> and methylene blue **84**<sup>+</sup>–Cl<sup>−</sup>) in aqueous solutions (Fig. 42a), nine types of  $\pi$ -electronic ion pairs (22<sup>+</sup>–**81a**<sup>2−</sup>, 22<sup>+</sup>–**81b**<sup>2−</sup>, 22<sup>+</sup>–**82**<sup>2−</sup>, 283<sup>+</sup>–**81a**<sup>2−</sup>, 283<sup>+</sup>–**81b**<sup>2−</sup>, 283<sup>+</sup>–**82**<sup>2−</sup>, 284<sup>+</sup>–**81a**<sup>2−</sup>, 284<sup>+</sup>–**81b**<sup>2−</sup> and 284<sup>+</sup>–**82**<sup>2−</sup>) were obtained.<sup>95</sup> These ion pairs exhibit distinct electrical resistivity at 20 °C in compressed disk states, with resistivity values ranging from  $2.5 \times 10^9 \Omega \text{ cm}$  for 284<sup>+</sup>–**81a**<sup>2−</sup> to  $3.1 \times 10^{14} \Omega \text{ cm}$  for 283<sup>+</sup>–**82**<sup>2−</sup>.

Ion-pairing behaviours in aqueous solutions were investigated by water-soluble charged  $\pi$ -electronic systems such as **81a**<sup>2−</sup>.<sup>96</sup> Anionic fluorescence dyes in diluted aqueous solutions show fluorescence quenching by adding  $\pi$ -electronic cations.<sup>96a</sup> The absorption spectra of mixed solutions comprising different dyes such as 2Na<sup>+</sup>–**81a**<sup>2−</sup> and 2<sup>+</sup>–Cl<sup>−</sup> are different from the sum

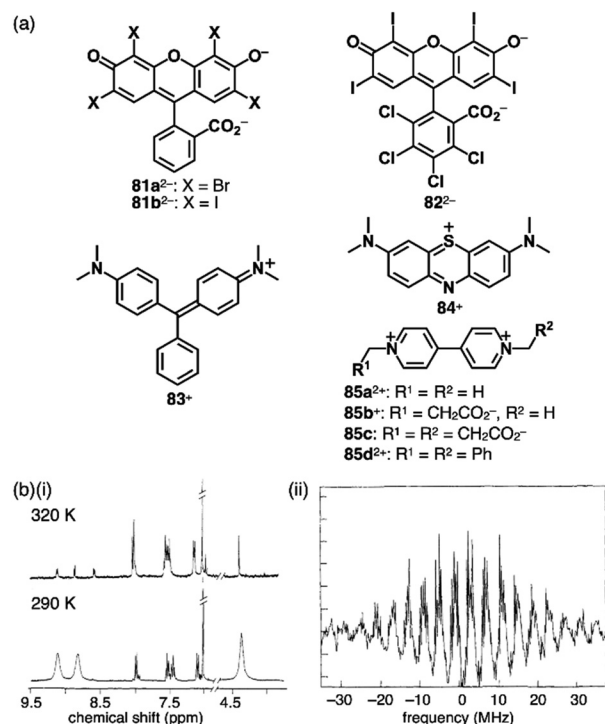


Fig. 42 (a) Xanthene-based  $\pi$ -electronic anions **81a**,**b**<sup>2−</sup> and **82**<sup>2−</sup>, the counteranions **83**,**84**<sup>+</sup>, **85a**,**d**<sup>2+</sup> and **85b**<sup>+</sup> and neutral **85c** and (b) (i) <sup>1</sup>H NMR and (ii) ESR spectra of **85a**<sup>2+</sup>–**81a**<sup>2−</sup> in DMSO-*d*<sub>6</sub> at 320 and 290 K and in DMSO at 295 K, respectively (redrawn from ref. 96c. Copyright 1992 ACS).

of those of the components, suggesting the ion pairing in aqueous solution. Fluorescence quantum yields of the ion pairs in organic solvents are considerably lower than those in aqueous solution.<sup>96a</sup> In particular, a new absorption band appears upon increasing the concentration of 2Na<sup>+</sup>–**82**<sup>2−</sup> in an aqueous solution containing **85a**<sup>2+</sup>–2Cl<sup>−</sup> ( $\sim 10^{-5} \text{ M}$ ) (Fig. 42a).<sup>96b</sup> Variable concentration measurements revealed an association constant of 9000 M<sup>−1</sup> at 298 K. In addition, similar measurements for the methylviologen derivatives whose charges are partially compensated by negatively charged substituents, **85b**<sup>+</sup> and **85c**, suggest that the contributions of electrostatic attractions to the enthalpy changes are  $-1.1$  to  $-1.3 \text{ kcal mol}^{-1}$  in aqueous solutions.

In the crystal structure of **85a**<sup>2+</sup>–**81a**<sup>2−</sup>, **81a**<sup>2−</sup> is stacked with **85a**<sup>2+</sup> in the interplanar distance of 3.43 Å.<sup>96c</sup> The resulting  $\pi$ - $\pi$  forms a slip-stacked arrangement with another  $\pi$ - $\pi$  through contact between the xanthene units of two **81a**<sup>2−</sup> units that are arranged in the antiparallel orientation. The stacked tetrad is slip-stacked with another tetrad using the remaining part of the xanthene unit of **81a**<sup>2−</sup>, resulting in the formation of a columnar structure. In **85d**<sup>2+</sup>–**81a**<sup>2−</sup>, the dianion **81a**<sup>2−</sup> is stacked with another **81a**<sup>2−</sup> in the interplanar distance of 3.76 Å in an antiparallel fashion. The stacked dianion dimer and one dication are alternately stacked with an interplanar distance of 3.58 Å, while another dication is positioned between the columns comprising one dication and two dianion. For **85a**<sup>2+</sup>–**81a**<sup>2−</sup>, partial dissociation is observed in DMF with an

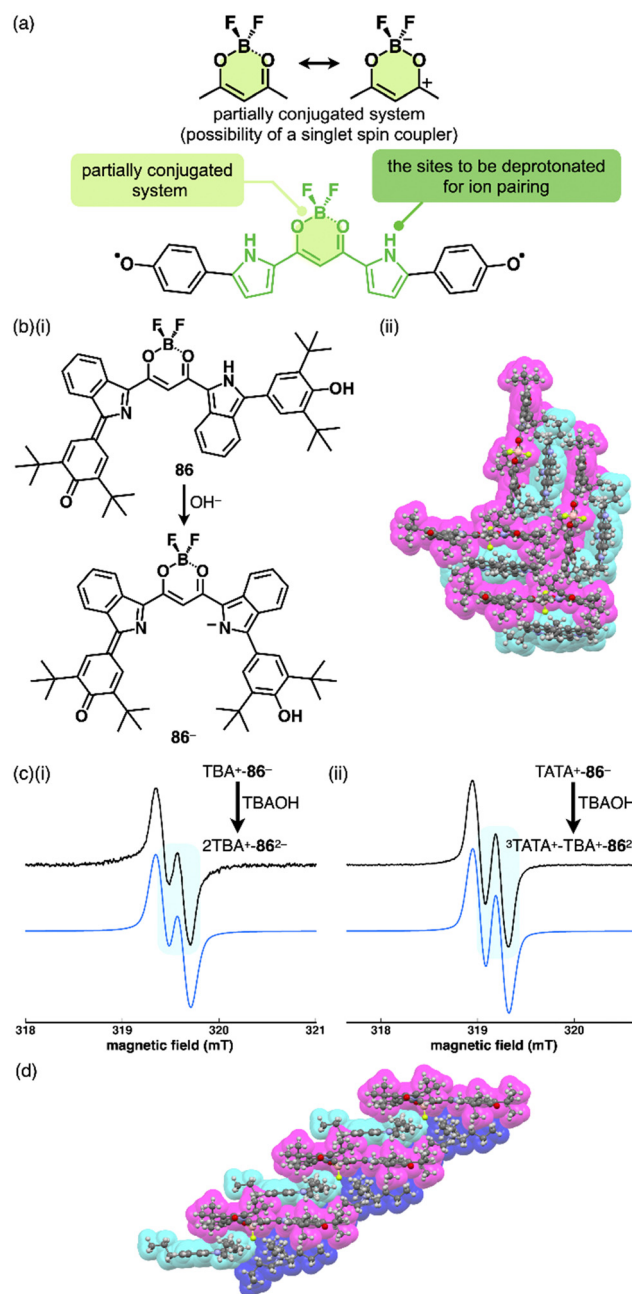


association constant of  $91\,000\text{ M}^{-1}$ . In this solvent, the stacked ion pair is further converted to  $85\text{a}^{2+}\text{--}[\text{81a}^{2-}\text{--}85\text{a}^{2+}\text{--}81\text{a}^{2-}]$ . Intriguingly, the  $^1\text{H}$  NMR signals assigned to  $85\text{a}^{2+}$  of  $85\text{a}^{2+}\text{--}81\text{a}^{2-}$  in  $\text{DMSO-}d_6$  broaden upon increasing temperature and disappear at 320 K, and the ESR is observed in DMSO at 295 K, indicating the presence of a thermally accessible CT state  $85\text{a}^{\bullet+}\text{--}81\text{a}^{\bullet-}$  (Fig. 42b).<sup>96c</sup>

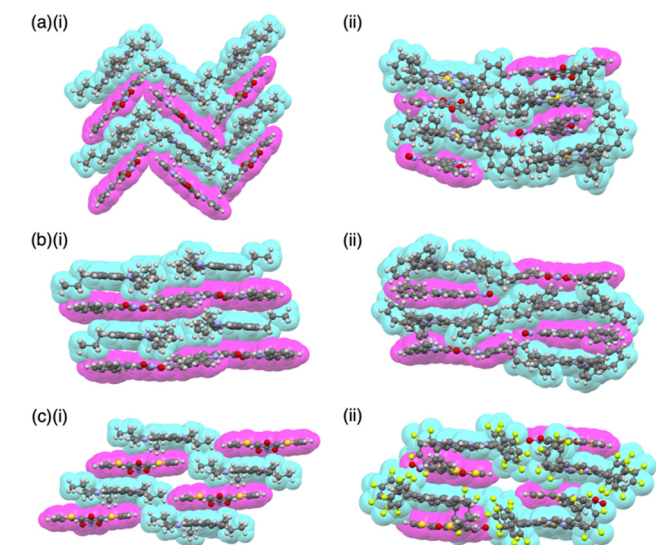
Deprotonated dipyrrolynitrophenols **34a,b**<sup>−</sup> (Fig. 21a) were used for the components of  $\pi$ -electronic ion-pairing assemblies.<sup>67c</sup>  $\pi$ -Electronic cations,  $^3\text{TATA}^+$  and  $22\text{a}^+$ , and  $\pi$ -electronic anions, **34a,b**<sup>−</sup>, are alternately stacked with the stacking distances in the range of 3.27–3.47 Å, forming charge-by-charge assemblies (Fig. 43a and b).<sup>67c</sup> **35a**<sup>−</sup> exhibits charge-by-charge assemblies with  $^3\text{TATA}^+$  and  $22\text{d}^+$  in the solid state (Fig. 43c).<sup>68</sup>

As described in Section 2.3, quinone methide units effectively stabilize  $\pi$ -electronic anions.<sup>69</sup> In addition, as represented by Yang's diradical,<sup>97</sup> combinations of multiple quinone methide units can provide open-shell species, whose spin states depend on the bridging unit as spin coupler; however, the ground state of **36**<sup>−</sup> is closed-shell state.<sup>69</sup> Modifications of the deprotonated pyrrole-based quinoidal systems induce remarkable electronic properties derived from the molecular structures and interactions with coexisting cations. Quinoidal systems exhibit unique diradical properties, with their singlet and triplet states dependent on the linker units acting as spin couplers. The PB core, which has been employed for anion binding (Section 3), possesses a partially cross-conjugated structure that functions as a singlet coupler, resembling a polarized carbonyl unit (Fig. 44a). PB-based quinone methide **86** (Fig. 44b(i)), which is an oxidized form of PB, was synthesized as a benzo-fused derivative to enhance stability.<sup>98</sup> **86** was deprotonated by TBAOH and NaOH to provide **86**<sup>−</sup> in the forms of  $\text{TBA}^+$  and  $\text{Na}^+$  ion pairs, the latter of which was further converted to a  $^3\text{TATA}^+$  ion pair by ion-pair metathesis (Fig. 44b(ii)). In the crystal structure of  $^3\text{TATA}^+\text{--}86^{\bullet-}$ ,  $^3\text{TATA}^+$  and

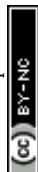
**86**<sup>−</sup> are alternately stacked with their  $\pi$ -systems, creating the charge-by-charge assembly (Fig. 44b(ii)). Upon further deprotonation of the ion pairs of **86**<sup>−</sup> with  $\text{TBA}^+$  and  $^3\text{TATA}^+$  by TBAOH, the dianion **86**<sup>2−</sup> was formed, resulting in the ion pairs  $2\text{TBA}^+\text{--}86^{2-}$  and  $^3\text{TATA}^+\text{--TBA}^+\text{--}86^{2-}$ , respectively (Fig. 44c). Unlike **86** and the



**Fig. 44** (a) Resonance structures of diketone-boron complex (top) and conceptual diagram of PB-based quinoidal molecule represented in the diradical tautomer of the parent form (bottom), (b) (i) formation of quinoidal dipyrrolyldiketone boron complex in the form of monoanion **86**<sup>−</sup> by deprotonation of **86** and (ii) single-crystal X-ray structure of  $^3\text{TATA}^+\text{--}86^{\bullet-}$ , (c) ESR spectra of (i)  $2\text{TBA}^+\text{--}86^{2-}$  and (ii)  $^3\text{TATA}^+\text{--TBA}^+\text{--}86^{2-}$  in mesitylene (1 mM) (black line) and corresponding simulation patterns (blue line) and (d) single-crystal X-ray structure of  $^3\text{TATA}^+\text{--TBA}^+\text{--}86^{2-}$  (redrawn from ref. 98. Copyright 2023 ACS). The crystal structures are redrawn from cif files: 2225435 and 2225436.



**Fig. 43** Single-crystal X-ray structures of diarylnitrophenoxides: (a) (i)  $^3\text{TATA}^+\text{--}34\text{a}^-$  and (ii)  $22\text{a}^+\text{--}34\text{a}^-$ , (b) (i)  $^3\text{TATA}^+\text{--}34\text{b}^-$  and (ii)  $22\text{a}^+\text{--}34\text{b}^-$  and (c) (i)  $^3\text{TATA}^+\text{--}35\text{a}^-$  and (ii)  $22\text{d}^+\text{--}35\text{a}^-$  (redrawn from cif files: 2061520, 2061524, 2061521, 2061525, 2253152 and 2253153).



monoanion  $86^-$ , the dianion  $86^{2-}$  in  $2TBA^+-86^{2-}$  exhibits sharp and broad  $^1\text{H}$  NMR signals at  $-50^\circ\text{C}$  and r.t., respectively, in  $\text{CD}_2\text{Cl}_2$  and displays the ESR signals with a  $g$  value of 2.003 at r.t. in mesitylene (Fig. 44c(i)). These observations indicate a ground-state singlet diradical character. Additionally, the ESR signals of  $^3\text{TATA}^+-\text{TBA}^+-86^{2-}$  differ from those of  $2\text{TBA}^+-86^{2-}$ , displaying counteranion-dependent magnetic properties (Fig. 44c(ii)). The ground-state singlet diradical nature is further confirmed by the enhancement of the variable-temperature ESR for  $^3\text{TATA}^+-\text{TBA}^+-86^{2-}$ , yielding an estimated  $\Delta E_{\text{ST}}$  of  $-6.4\text{ kcal mol}^{-1}$ . The interaction between  $86^{2-}$  and  $^3\text{TATA}^+$  is also observed in single-crystal X-ray analysis of  $^3\text{TATA}^+-\text{TBA}^+-86^{2-}$ , wherein  $\text{TBA}^+$  is situated on the PB unit of  $86^{2-}$  with reduced contact of  $^3\text{TATA}^+$  and  $86^{2-}$  (Fig. 44d).<sup>98</sup>

Moreover, deprotonated hydroxyporphyrins are also adopted in  $\pi$ -electronic ion pairs. Assembling behaviours of  $^3\text{TATA}^+$  ion pairs of anionic porphyrin free base  $37^-$  and  $\text{Ni}^{\text{II}}$  and  $\text{Pd}^{\text{II}}$  complexes  $37\text{a}, \text{c}^-$  were investigated (Fig. 45a).<sup>9</sup> The anions are stacked with  $^3\text{TATA}^+$  to form charge-by-charge assemblies in the crystal structures. In  $^3\text{TATA}^+-37\text{a}^-$ , each component is stacked with identically charged species, resulting in a partially charge-segregated mode (Fig. 45b). The absorption spectrum of  $^3\text{TATA}^+-37\text{a}^-$  in the solid state exhibits blue-shifted and red-shifted bands derived from  $37\text{a}^-$  and  $^3\text{TATA}^+$ , respectively, compared to those for the monomeric states of  $37\text{a}^-$  ( $\text{TBA}^+-37\text{a}^-$ ) and  $^3\text{TATA}^+$  ( $^3\text{TATA}^+-\text{Cl}^-$ ) in  $\text{CH}_2\text{Cl}_2$  (Fig. 45c). This shift is attributed to exciton coupling arising from the stacking of identically charged species in the parallel (H-like) arrangement for  $37\text{a}^-$  and a head-to-tail (J-like) arrangement for  $^3\text{TATA}^+$ . Additionally, solid-state transient absorption spectra of  $^3\text{TATA}^+-37\text{a}^-$  reveal the generation of radical  $37\text{a}^\bullet$  by the PET between the  $\pi$ -electronic systems. Furthermore, EDA of the packing structure of  $^3\text{TATA}^+-37\text{a}^-$  indicates a significant contribution

of electrostatic and dispersion forces in stabilizing the stacked structures of the constituent  $\pi$ -electronic ions.<sup>9</sup>

Remarkable electronic properties emerge from  $\pi$ -electronic ion pairs and their assemblies comprising charged species with large  $\pi$ -systems. Promising  $\pi$ -electronic ion pairs can be produced by introducing charged porphyrins as cations and anions, including  $37\text{a}^-$  for the latter. Modifications of the electronic states and resulting assembled structures can be achieved by introducing substituents, as seen in  $\text{Au}^{\text{III}}$  complexes  $22\text{a}, \text{c}^+$  that possess four  $\text{C}_6\text{H}_5$  and  $\text{C}_6\text{F}_5$  moieties, respectively, at the *meso* positions.<sup>52</sup> Combining oppositely charged porphyrins with distinct electronic states according to introduced substituents provides diverse ion pairs that can be used for functional materials. Thus, in addition to  $37\text{a}^-$ , *meso*-phenyl-substituted  $87^-$  was prepared as an electron-rich “activated” anion (Fig. 46a).<sup>6</sup> The “activated” cation  $22\text{d}^+$ , with three *meso*- $\text{C}_6\text{F}_5$  units for efficient stacking, has been a focal point for creating an activated ion pair with  $87^-$ . Several porphyrin ion pairs, including  $22\text{a}^+-37\text{a}^-$ ,<sup>52a</sup>  $22\text{a}^+-87^-$  and  $22\text{d}^+-37\text{a}^-$  as well as the activated ion pair  $22\text{d}^+-87^-$ , were prepared by ion-pair metathesis from charged porphyrin starting

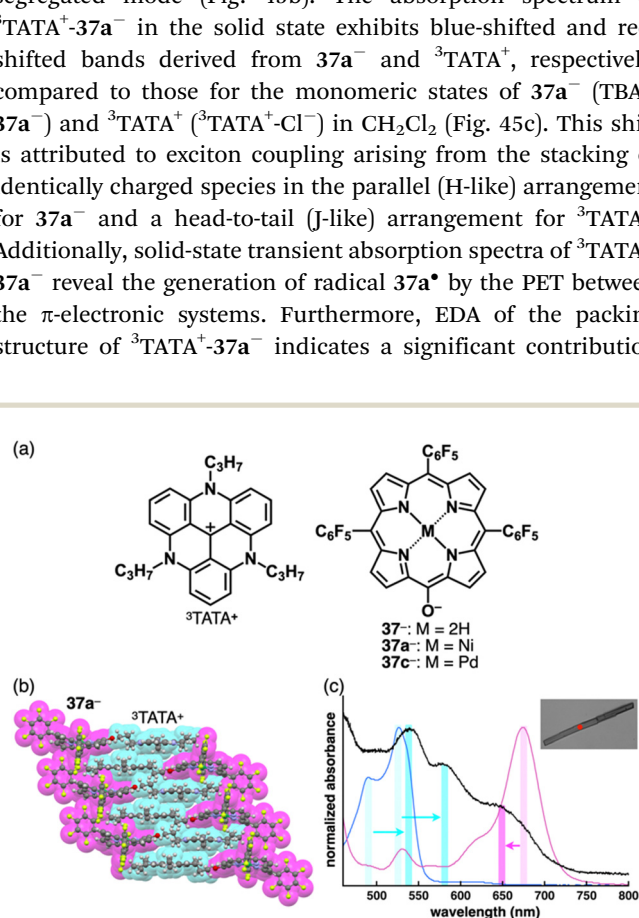


Fig. 45 (a) Ion pairs  $^3\text{TATA}^+-37^-$  and  $^3\text{TATA}^+-37\text{a}, \text{c}^-$ , (b) single-crystal X-ray structure of  $^3\text{TATA}^+-37\text{a}^-$  (redrawn from a cif file: 1906678) and (c) UV/vis absorption spectrum of the single crystal of  $^3\text{TATA}^+-37\text{a}^-$  (black) along with those of the solution states ( $1 \times 10^{-5}\text{ M}$  in  $\text{CH}_2\text{Cl}_2$ ) of  $\text{TBA}^+-37\text{a}^-$  (magenta) and  $^3\text{TATA}^+-\text{Cl}^-$  (cyan) (inset: optical microscopy image of a single crystal of  $^3\text{TATA}^+-37\text{a}^-$ ) (redrawn from ref. 9. Copyright 2021 RSC).

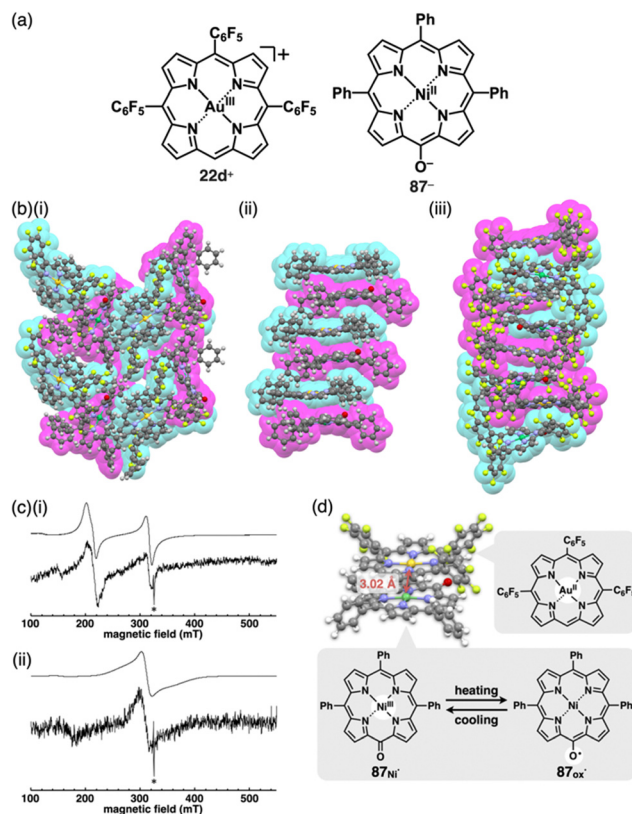


Fig. 46 (a) Ion pair of charged porphyrins,  $22\text{d}^+-87^-$ , as an activated ion pair, (b) single-crystal X-ray structures of (i)  $22\text{d}^+-87^-$ , (ii)  $22\text{a}^+-87^-$  and (iii)  $22\text{d}^+-37\text{a}^-$  (redrawn from cif files: 2201074, 2201075 and 2201076), (c) simulated (top) and observed (bottom) ESR spectra of  $22\text{d}^+-87^-$  in toluene at (i) 10 and (ii) 140 K and (d) closely stacked structure ( $\pi$ -srp) of  $22\text{d}^+$  and  $87^-$  with the spin-spin distance and the change in spin distribution of  $87^\bullet$  ( $87\text{Ni}^\bullet$  and  $87\text{Ox}^\bullet$ ) as a result of temperature change (redrawn from ref. 6. Copyright 2022 ACS). Signals marked with \* in (c) are ascribable to a trace amount of dissociated  $87^\bullet$ , whereas white circles in (d) represent spin distributions.



materials, resulting in solid-state ion-pairing assemblies. In the case of  $22a^+-37a^-$ , stacking arrangement exhibits a zigzag pattern owing to the steric effect of *meso*-phenyl groups.<sup>52a</sup> In contrast, the assembly of  $22d^+-87^-$  involves a charge-segregated mode, with four benzene molecules included for each ion pair (Fig. 46b(i)). Nevertheless,  $\pi$ - $\pi$ -based charge-by-charge assembly modes in  $22a^+-87^-$  results in  $\pi$ - $\pi$  interaction between porphyrin ions, as the steric hindrance of the porphyrin anion is relaxed (Fig. 46b(ii)). The packing diagram in  $22d^+-37a^-$  demonstrates a 1D-chain structure of  $\pi$ - $\pi$  assembled in a four-fold helical symmetry, with  $22d^+$  and  $37a^-$  arranged in a stacking distance of 3.57 Å (Fig. 46b(iii)).<sup>6,99</sup>

Charged porphyrins form stacked dimers in solution, as detected by  $^1H$  NMR chemical shifts derived from shielding and deshielding effects. Concentration-dependent  $^1H$  NMR of  $22d^+-37a^-$  in  $CD_2Cl_2$  provides the heterodimerization constant ( $K_{dim} = 2.8 \times 10^5 M^{-1}$ ) at 20 °C. On the other hand, the UV/vis absorption spectrum of  $22d^+-37a^-$  in the  $\pi$ - $\pi$  is almost the sum of the spectra of constituent  $22d^+$  and  $37a^-$ . In contrast, the activated ion pair  $22d^+-87^-$  exhibits a UV/vis absorption spectrum that differs from the corresponding summed spectra in less polar solvents. The spectrum observed in toluene is consistent with the combined spectra of the reduced  $Au^{II}$  complex  $22d^\bullet$  and the oxidized  $Ni^{II}$  complex oxyradical  $87_{ox}^\bullet$ , indicating an electron transfer from  $87^-$  to  $22d^+$  to generate a  $\pi$ -stacked radical pair ( $\pi$ - $srp$ ). Notably, the ESR spectrum in frozen toluene at 10 K (Fig. 46c(i)) displays the  $\pi$ - $srp$  as a pair of  $22d^\bullet$  and  $Ni^{III}$  complex  $87^\bullet$  with a short spin-spin distance of 3.02 Å (Fig. 46d top). VT ESR spectra suggest spin transfer in  $87^\bullet$  and antiferromagnetic interaction in  $\pi$ - $srp$  (Fig. 46c(ii) and d bottom).<sup>6</sup> The less activated porphyrin ion pairs exhibit  $\pi$ - $\pi$ s that correlate with the redox potentials of their constituent components based on the substituents. Further investigations on the PET for ion pairs, including triaryl-substituted  $22b^+$  and  $22d^+$ , revealed varying electron-transfer rates, leading to the formation of excited-state radical species. These findings suggest application potential of these ion pairs in photocatalysis owing to their tuneable electronic state.<sup>6,53</sup>

## 5. Conclusions

$\pi$ -Electronic ion pairs, comprising countercharged  $\pi$ -electronic systems, hold immense potential in generating structured assembled arrangements and displaying unique properties. Assembling modes of charged components are modulated by their geometries and electronic states. Thus, the design and synthesis of charged  $\pi$ -electronic systems are crucial for the fabrication of ion-pairing assemblies (Fig. 47). The presence of charges within the core  $\pi$ -electronic units makes their preparation challenging owing to the inherent reactivity arising from electron-deficient and -rich states. Furthermore, the specific combinations of  $\pi$ -electronic ions are crucial for determining the nature of the assemblies and the resulting properties. Less stable  $\pi$ -electronic ions can be stabilized by combining specific counterions *via* the formation of  $\pi$ - $\pi$ s based on  $\pi$ - $\pi$

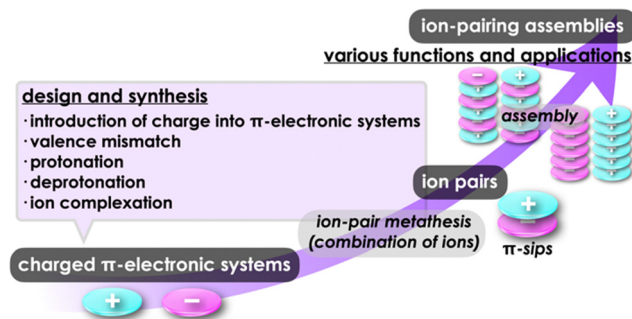


Fig. 47 Formation of ion-pairing assemblies through design and synthesis of charged  $\pi$ -electronic systems, ion-pair metathesis and assembly.

interactions. Hence, diversity is a central aspect in ion-pairing chemistry.

The assemblies of ion pairs reveal their distinct characteristics not only in solution but also in bulk states like crystals and soft materials. The ability to control assembling modes, whether charge-by-charge or charge-segregated, is crucial for exhibiting specific electronic properties based on the arrangements. The electronic properties of  $\pi$ -electronic ion pairs and their assemblies are distinct from those of electronically neutral  $\pi$ -systems. The proximally located oppositely charged  $\pi$ -electronic systems induce the polarized columnar structures, most polarizations of which are cancelled by those of neighbouring columns. Spontaneous polarization with uniform alignment of polarized columns facilitates the development of ferroelectric materials.<sup>100</sup> The manipulation of redox potentials between  $\pi$ -electronic cations and anions enables electron transfer in both excited and ground states. The resulting radical pairs, especially under photoirradiation, can serve as catalysts, with cation- and anion-derived radicals acting as reducing and oxidizing species, respectively.<sup>101</sup> Furthermore, columnar structures comprising identically charged species offer potential applications in efficient semiconductive materials. The electrostatic repulsion between identically charged species can be overcome by dispersion forces originating from  $\pi$ -electronic systems and dipole-dipole interactions in polarized charged species. These strategies are easily attainable through simple modifications for  $\pi$ -electronic systems, in contrast to inorganic ionic species. This review article showcases the production of diverse  $\pi$ -electronic ion pairs, demonstrating possibilities for future property investigations and applications. The authors have made significant strides in understanding their intriguing behaviours of these ion pairs through thorough comprehensive studies in synthesis, combination, assembly and electronic and photophysical properties in recent years.<sup>102</sup> A significant portion of ion-pairing chemistry remains unexplored and awaits exploration through ongoing research endeavours.

## Conflicts of interest

There are no conflicts of interest to declare.

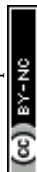


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## Notes and references

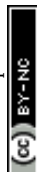
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