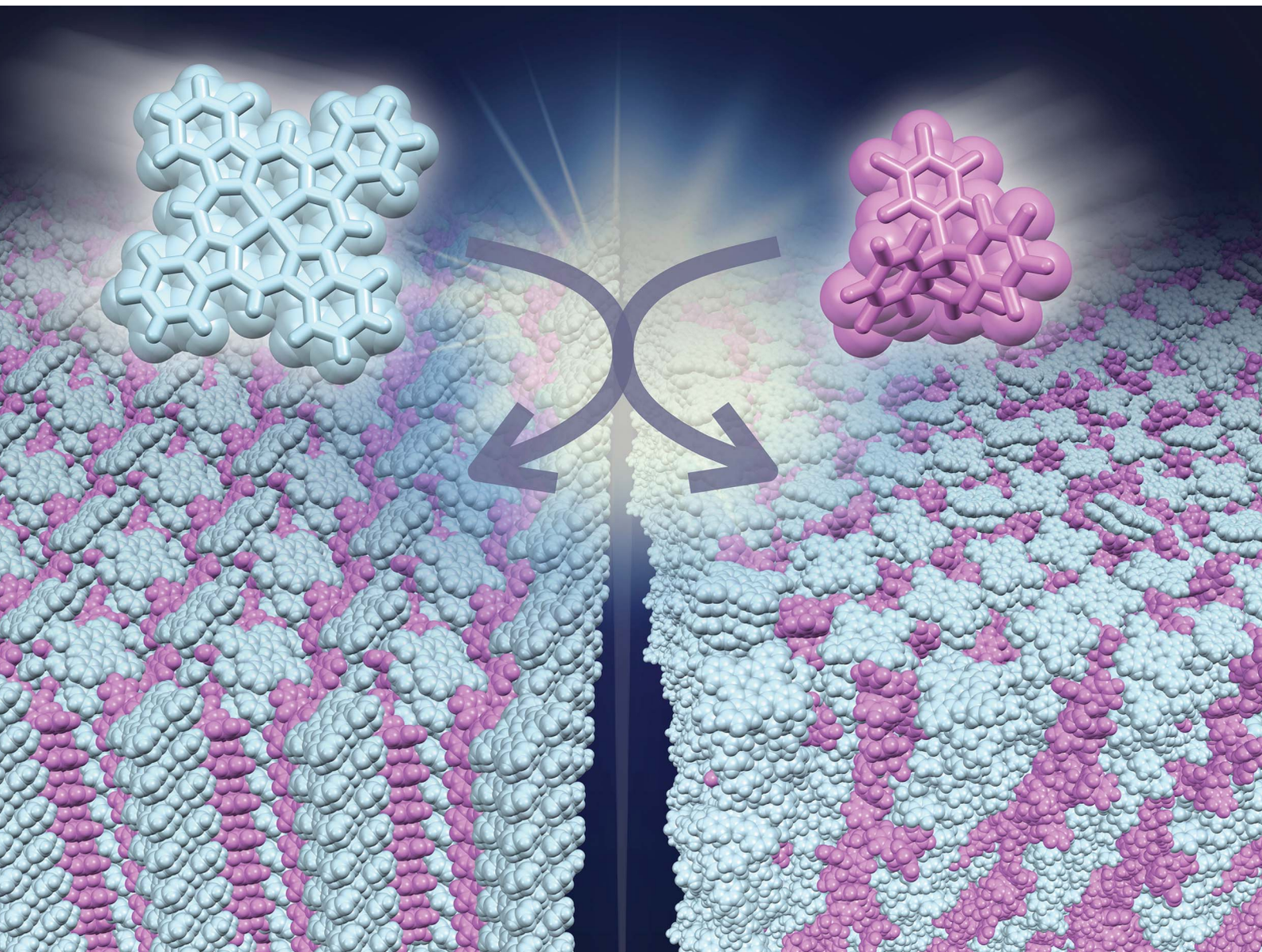


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EDGE ARTICLE

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Electrically conductive charge-segregated pseudo-polymorphs comprising highly planar expanded π -electronic cations†

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Independently stacked positively and negatively charged π -electronic systems in charge-segregated columnar structures are desired for electronic properties derived from their electron-deficient and -rich assembling states, respectively. An expanded π -electronic cation, benzoporphyrin Au^{III} complex, was synthesized as the component of ion pairs in combination with counteranions. In contrast to benzoporphyrin, which is known for its insolubility in organic solvents, the ion pairs with bulky anions in this study are soluble in common organic solvents. The ion pairs formed charge-segregated assemblies as two pseudo-polymorphs of single-crystal and less-crystalline (LeC) states based on the stacking of the benzoporphyrin Au^{III} complex. XRD and solid-state NMR measurements, along with molecular dynamics (MD) simulation, revealed that the LeC states were formed by a less-ordered arrangement of constituting ions induced by bulky counteranions. The electric conductivity properties were observed in the single-crystal and LeC charge-segregated assemblies.

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Introduction

The ordered arrangement of π -electronic systems is crucial for charge-carrier transport properties.¹ Expanded π -planes are adequate for achieving high performance in organic semiconductive materials.^{1g} Since substituents affect the electronic states of molecules and their arrangement, π -electronic systems that have no substituents are in great demand.^{1e} However, such systems have low solubility (high crystallinity), making it difficult to arrange the constituents to form assembled structures (Fig. 1a top left). A promising strategy is the preparation of ion pairs of charged π -electronic systems by combining them with counterions that improve solubility (Fig. 1a top right).² An appropriate combination of charged constituents enables facile handling of π -electronic systems to form counterion-dependent assemblies for applications (Fig. 1a bottom). In particular, independently stacked positively and negatively charged π -electronic systems are fascinating because of their ability to form electron-deficient and -rich assembling states, which can function as n- and p-type semiconductive pathways, respectively, in charge-segregated columnar structures.³ Expanded π -systems contribute to influential dispersion forces that overcome electrostatic repulsion in charged columns. The potential positively charged π -systems are porphyrin Au^{III} complexes, which have been included in various ion-pairing assemblies in the form of crystals, supramolecular gels and liquid crystals

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† Electronic supplementary information (ESI) available: Synthetic procedures, analytical data, computational details and CIF files for the single-crystal X-ray structural analysis. CCDC 2387040–2387044. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d4sc07576e>



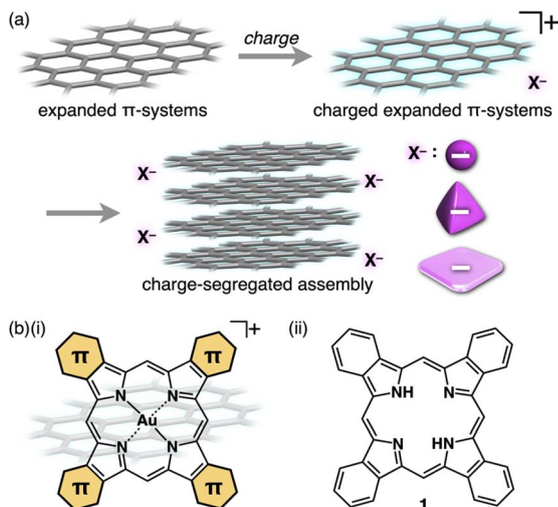


Fig. 1 (a) Expanded π -electronic systems that have no peripheral substituents (top left) and their charged and ion-pairing states (top right, represented as cations), forming a charge-segregated assembly by stacking (bottom) and (b) (i) π -expanded porphyrin Au^{III} complexes as expanded π -electronic cations that can be used in (a) and (ii) benzoporphyrin **1**. The positive signs were omitted in the charge-segregated assembly in (a).

depending on the substituents.^{4,5} Expansion of π -electronic systems can be achieved by modifications at the pyrrole β -positions (Fig. 1b(i)). Numerous porphyrin derivatives that have been synthesized to date offered the choice to use benzoporphyrin⁶ **1** (Fig. 1b(ii)) as a framework to provide highly planar charged expanded π -electronic systems. Large planes of benzoporphyrin-based cations are suitable for stacking to form charge-segregated assemblies, whose packing structures can be controlled by coexisting counteranions. The arrangement of substituent-free planar cations and bulky anions can be modulated by crystallization conditions. This study shows π -expanded cation-based ion-pairing assemblies in single-crystal and pseudo-polymorph less-crystalline (LeC) states and their electric conductivity properties derived from the charge-segregated assemblies.

Results and discussion

Synthesis and characterization of expanded π -electronic cations

Benzoporphyrins, including metal complexes, can be synthesized from bicyclo[2.2.2]octadiene precursors *via* retro-Diels–Alder reactions.^{9f} In this study, Au^{III} complexation was conducted for bicycloporphyrin **2** to afford the Au^{III} complex 2au^+ mainly as a triflate (OTf^-) ion pair by treatment with KAuCl_4 in the presence of AgOTf and NaOAc (Fig. 2 top). The ion pair $2\text{au}^+\text{OTf}^-$ was converted to the Cl^- ion pair $2\text{au}^+\text{Cl}^-$ in 28% yield (two steps) using an ion-exchange resin (Amberlite: IRA402BL Cl). As the selection of the counteranions was crucial in this study, the ion-pair metathesis of $2\text{au}^+\text{Cl}^-$ with AgPF_6 , $\text{LiB}(\text{C}_6\text{F}_5)_4$ (LiFABA), $\text{NaB}(3,5\text{-(CF}_3)_2\text{C}_6\text{H}_3)_4$ (NaBARF) and NaPCCp (PCCp^- : pentacyanocyclopentadienide)⁷ afforded the

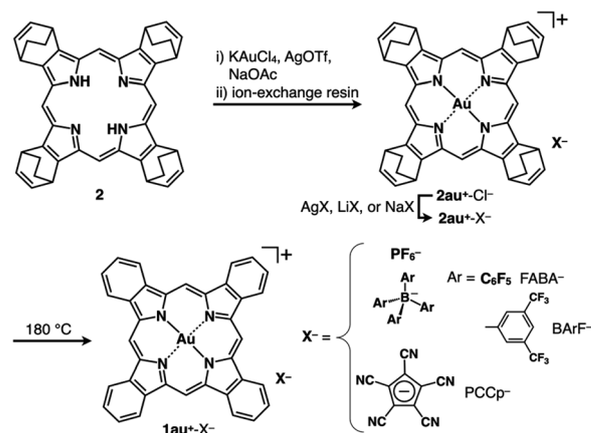


Fig. 2 Synthesis of benzoporphyrin Au^{III} complex ion pairs $1\text{au}^+\text{X}^-$ ($\text{X}^- = \text{PF}_6^-, \text{FABA}^-, \text{BARF}^-$ and PCCp^-) *via* the corresponding bicycloporphyrin Au^{III} complex ion pairs $2\text{au}^+\text{X}^-$.

corresponding ion pairs $2\text{au}^+\text{X}^-$ ($\text{X}^- = \text{PF}_6^-, \text{FABA}^-, \text{BARF}^-$ and PCCp^-) in 66–76% yields. The obtained ion pairs were characterized by ^1H , ^{13}C and ^{19}F NMR and ESI-TOF-MS. DMSO solutions of 2au^+ ion pairs (4 μM) exhibited Soret and Q-bands at 392 and 507/542 nm, respectively, indicating that 2au^+ exists as a monomeric state with a negligible counteranion effect on the electronic properties of 2au^+ (Fig. S12[†]).⁸ $2\text{au}^+\text{FABA}^-$ and $2\text{au}^+\text{BARF}^-$ were also characterized by X-ray analysis for single crystals prepared by vapour diffusion from $\text{CH}_3\text{CN}/\text{water}$ (Fig. 3, S18 and S19[†]). In the solid-state structures, the cation 2au^+ , refined as one of the stereoisomers, formed stacked dimers with stacking/ $\text{Au}\cdots\text{Au}$ distances of 3.56/3.41 and 3.48/3.37 Å, respectively, and a rotation of $\sim 45^\circ$ owing to dispersion forces

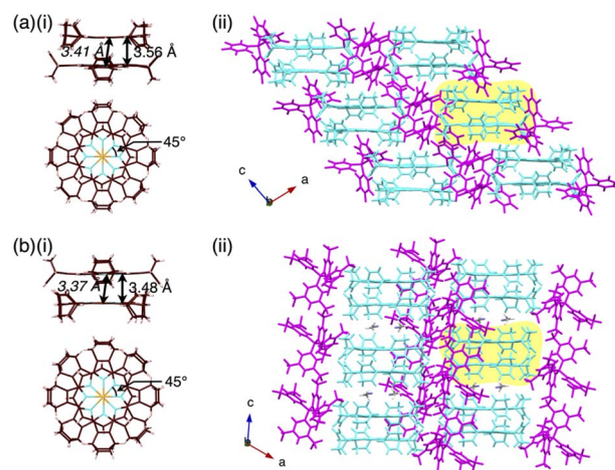


Fig. 3 Single-crystal X-ray structures of (a) $2\text{au}^+\text{FABA}^-$ and (b) $2\text{au}^+\text{BARF}^-$: (i) side and top views of stacked dimers with stacking and $\text{Au}\cdots\text{Au}$ (italic) distances and (ii) packing structures with the yellow parts corresponding to the stacked dimers in (i). Atom colour codes in (i): brown, pink, blue and orange refer to carbon, hydrogen, nitrogen and gold, respectively. Colour codes in (ii): cyan and magenta refer to cation and anion parts, respectively. Solvent molecules are omitted in (a), whereas CH_3CN molecules are shown in (b).



at the core planes and β -bicyclo units (Fig. 3a and b(i)). The stacking of $2\mathbf{au}^+$ is visualized by Hirshfeld surface analysis, which shows a bow-tie arrangement of red and blue triangles in the shape-index property and flat regions in the curvedness property (Fig. S24 and S25[†]).⁹ Mean-plane deviations of 0.052/0.068 and 0.036 Å for stacking $2\mathbf{au}^+$ planes (core 25 atoms), respectively, indicate slightly curved $2\mathbf{au}^+$ planes, owing to the Au...Au distances being less than the stacking distances. Such remarkably close Au...Au distances are also ascribable to the orbital interactions arising from overlap of the $5d_{z^2}$ and $6p_z$ orbitals of the adjacent Au^{III}.¹⁰ The stacked $2\mathbf{au}^+$ dimers in ion pairs are aligned along the *c*-axis with an offset, which is smaller in $2\mathbf{au}^+$ -BARF⁻ with a longer distance between the stacked dimers by including two CH₃CN molecules (Fig. 3a and b(ii)). In either case, counteranions are located at the side of the stacked $2\mathbf{au}^+$ dimers. In particular, BARF⁻ anions are located proximally at the side of the stacked $2\mathbf{au}^+$ dimers, forming a pseudohexagonally arranged packing structure along the *c*-axis (Fig. S19a[†]).

Stacking of $2\mathbf{au}^+$ by overcoming electrostatic repulsion was also observed in the solution state. In CD₃CN, the ¹H NMR of $2\mathbf{au}^+$ -FABA⁻ showed broad signals at 9.96, 8.37/6.94, 5.98 and 3.49–1.30 ppm ascribable to *meso*-CH, bicyclo-*sp*²-CH, bicyclo-bridged methine-CH and bicyclo-*sp*³-CH, respectively, for the stacked dimer.¹¹ Concentration-dependent UV/vis absorption spectra of $2\mathbf{au}^+$ -FABA⁻ in CH₃CN exhibited a blue shift of λ_{\max} from 389 to 374 nm when increasing the concentration from 1.0×10^{-6} to 1.0×10^{-5} M, suggesting the formation of an H-like stacked dimer at the higher concentration (Fig. S71[†]). The transition dipole moments of $2\mathbf{au}^+$ in the optimized structure of the stacked dimer $2\mathbf{au}_2^+$ based on PCM-GD3BJ-B3LYP/6-31+G(d,p) with LanL2DZ for Au (CH₃CN) are arranged at $\sim 45^\circ$, suggesting that the H-like stacking mode induces a blue shift (Fig. S61a[†]).¹² In addition, TD-DFT calculation of the optimized $2\mathbf{au}_2^+$ revealed an absorption at 375 nm, which is blue-shifted by 16 nm compared to the monomer state (Fig. S46[†]).

According to the synthetic procedure for benzoporphyrin 1^{af} , $2\mathbf{au}^+$ -X⁻ (X⁻ = PF₆⁻, FABA⁻, BARF⁻ and PCCp⁻) were quantitatively transformed to the corresponding benzoporphyrin ion pairs $1\mathbf{au}^+$ -X⁻ by heating at 180 °C for 20–60 min in the absence of solvent (Fig. 2 bottom). In contrast to 1^{af} which is insoluble in most organic solvents, the obtained $1\mathbf{au}^+$ -X⁻ showed enhanced solubility. For example, $1\mathbf{au}^+$ -FABA⁻ was soluble in acetone, DMF, CH₃CN and DMSO. In contrast, another ion pair $1\mathbf{au}^+$ -Cl⁻, which was synthesized from $2\mathbf{au}^+$ -Cl⁻, was not soluble in these solvents.¹³ It is noteworthy that $1\mathbf{au}^+$ is soluble with facile handling in the form of the ion pairs with appropriate counteranions, although the optimized structure of $1\mathbf{au}^+$ estimated at B3LYP/6-31+G(d,p) with LanL2DZ for Au¹² exhibits completely planar geometry with a mean-plane deviation of 0.00 Å (Fig. S29[†]). ¹H NMR of $1\mathbf{au}^+$ -FABA⁻ in DMSO-*d*₆ (1.0 mM), as an example, at r.t. exhibited broad signals at 10.21, 9.18 and 8.11 ppm, suggesting soluble but aggregated structures as also indicated by concentration-dependent ¹H NMR (Fig. S74[†]).¹⁴ Such ¹H NMR signals in the downfield region suggested the aromatic ring current effect of $1\mathbf{au}^+$, which was further supported by nucleus-independent chemical shift (NICS) and the anisotropy of the current induced density (ACID) calculations

(Fig. S54 and S55[†]). Interestingly, ¹⁹F NMR in the same solvent shows sharp signals derived from FABA⁻ in the dispersed state. The UV/vis absorption spectrum of $1\mathbf{au}^+$ -FABA⁻ in DMSO (4 μ M), as a monomer state, exhibits Soret and Q bands of 408 and 564/616 nm, respectively (Fig. 4a), which are more red-shifted than those of $2\mathbf{au}^+$ -FABA⁻. The TD-DFT-based UV/vis absorption stick spectrum of $1\mathbf{au}^+$ in DMSO shows that these absorptions are mainly derived from the HOMO–1-to-LUMO+1 and HOMO-to-LUMO+1 transitions, respectively (Fig. 4a inset, S49[†]).

The ¹H NMR of $1\mathbf{au}^+$ -FABA⁻ in CD₃CN (1.0 mM) shows broader signals than those in DMSO-*d*₆, suggesting more aggregated structures in the less polar solvent (Fig. S9[†]). Similar to $2\mathbf{au}^+$ -FABA⁻, $1\mathbf{au}^+$ -FABA⁻ shows a λ_{\max} blue-shift from 402 to 384 nm upon increasing the concentration from 1.8×10^{-7} to 2.0×10^{-5} M in CH₃CN, suggesting the formation of H-like stacked structures (Fig. 4b(i) and S72[†]). Such a blue shift of the λ_{\max} is also observed in variable-temperature (VT)-UV/vis absorption spectra at lower temperatures (Fig. 4b(ii)). TD-DFT calculation of the optimized structure for the stacked dimer $1\mathbf{au}_2^+$ at PCM-B3LYP-GD3BJ/6-31+G(d,p) with LanL2DZ for Au (CH₃CN) suggests a slightly blue-shifted Soret band compared to that of the monomeric state (Fig. S51[†]). The dimerization constant (K_{dim}) of $1\mathbf{au}^+$ for $1\mathbf{au}^+$ -FABA⁻ is estimated to be $5 \times$

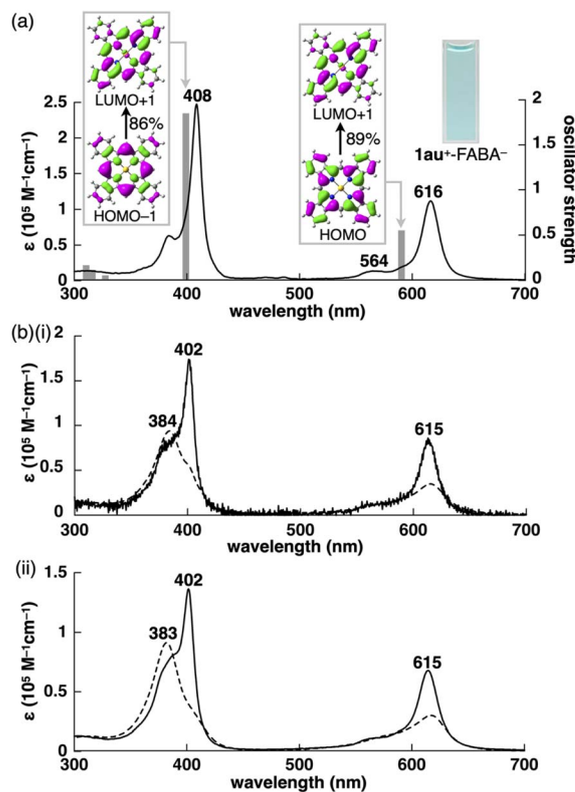


Fig. 4 (a) UV/vis absorption spectrum of $1\mathbf{au}^+$ -FABA⁻ in DMSO (4 μ M) (inset: photograph of the DMSO solution (4 μ M)) and TD-DFT-based UV/vis absorption stick spectrum (grey bar) of $1\mathbf{au}^+$ at PCM-B3LYP-GD3BJ/6-31+G(d,p) with LanL2DZ for Au (DMSO) and (b) UV/vis absorption spectra of $1\mathbf{au}^+$ -FABA⁻ in CH₃CN according to (i) concentrations (solid line: 0.18 μ M and broken line: 20 μ M) and (ii) temperatures (solid line: 70 °C and broken line: –40 °C).



10^6 M^{-1} in CH_3CN at r.t. from concentration-dependent UV/vis absorption spectral changes (Fig. S72†). π -Expansion of positively charged π -electronic systems induces influential dispersion forces, which enable the stacking of identically charged π -electronic systems.

Single-crystal-state charge-segregated assemblies

Prism-shaped single crystals of the ion pairs $1\text{au}^+\text{-X}^-$ ($\text{X}^- = \text{FABA}^-, \text{BARF}^-$ and PCCp^-), obtained from $\text{CH}_3\text{CN}/\text{CHCl}_3$,¹⁵ 1,1,1-trichloroethane/*n*-heptane and DMF/*o*-dichlorobenzene, respectively, were suitable for X-ray analysis, revealing the exact structures of the ion pairs and their assembled structures (Fig. 5). In these structures, the cation 1au^+ , showing planar geometry with mean-plane deviations (core 25 atoms) of 0.016, 0.031/0.011 and 0.023 Å, respectively, forms closely stacked columnar structures with stacking distances of 3.29, 3.36/3.44 and 3.37 Å, respectively, in the charge-segregated mode (Fig. 5a–c(i)). The stacked parts of 1au^+ are clearly shown by Hirshfeld surface analysis, exhibiting a bow-tie arrangement of red and blue triangles in the shape-index property and flat regions in the curvedness property (Fig. S26–S28†).⁹ The Au...Au

distances are 4.76, 3.93/4.59 and 3.81 Å, respectively, suggesting that the offset stacking of the cations is larger for $1\text{au}^+\text{-FABA}^-$. The angles of 44.1°, 48.4°/60.1° and 62.1° are estimated, respectively, for the lines passing through two Au atoms of stacked 1au^+ to the corresponding 41-atom mean planes. In these ion pairs, counteranions FABA^- , BARF^- and PCCp^- are located at the side of the columnar structures (Fig. 5a–c(ii) and (iii)). The ion-pair crystals $1\text{au}^+\text{-FABA}^-$, $1\text{au}^+\text{-BARF}^-$ and $1\text{au}^+\text{-PCCp}^-$ formed orthorhombic, monoclinic and orthorhombic packing, respectively, with the columns of stacked 1au^+ aligned along the *a*-, *c*- and *a*-axes, respectively, which are the long axes of the prism crystals (Fig. S23†). Notably, the stacked 1au^+ in the columns are tilted, with angles of 30.4°, 2.9°/3.0° and 21.2°, respectively, relative to the stacking axis (Fig. 5a–c(ii)). Interestingly, in the crystal packing of $1\text{au}^+\text{-FABA}^-$, an ion-pair framework composed of columnar cation structures and counteranions forms two tubular spaces per unit cell, with a volume of 5.56 nm³, containing disordered solvent molecules. The solvent molecules in the single crystal are not removed after heating at 100 °C under vacuum, as revealed by the X-ray analysis.

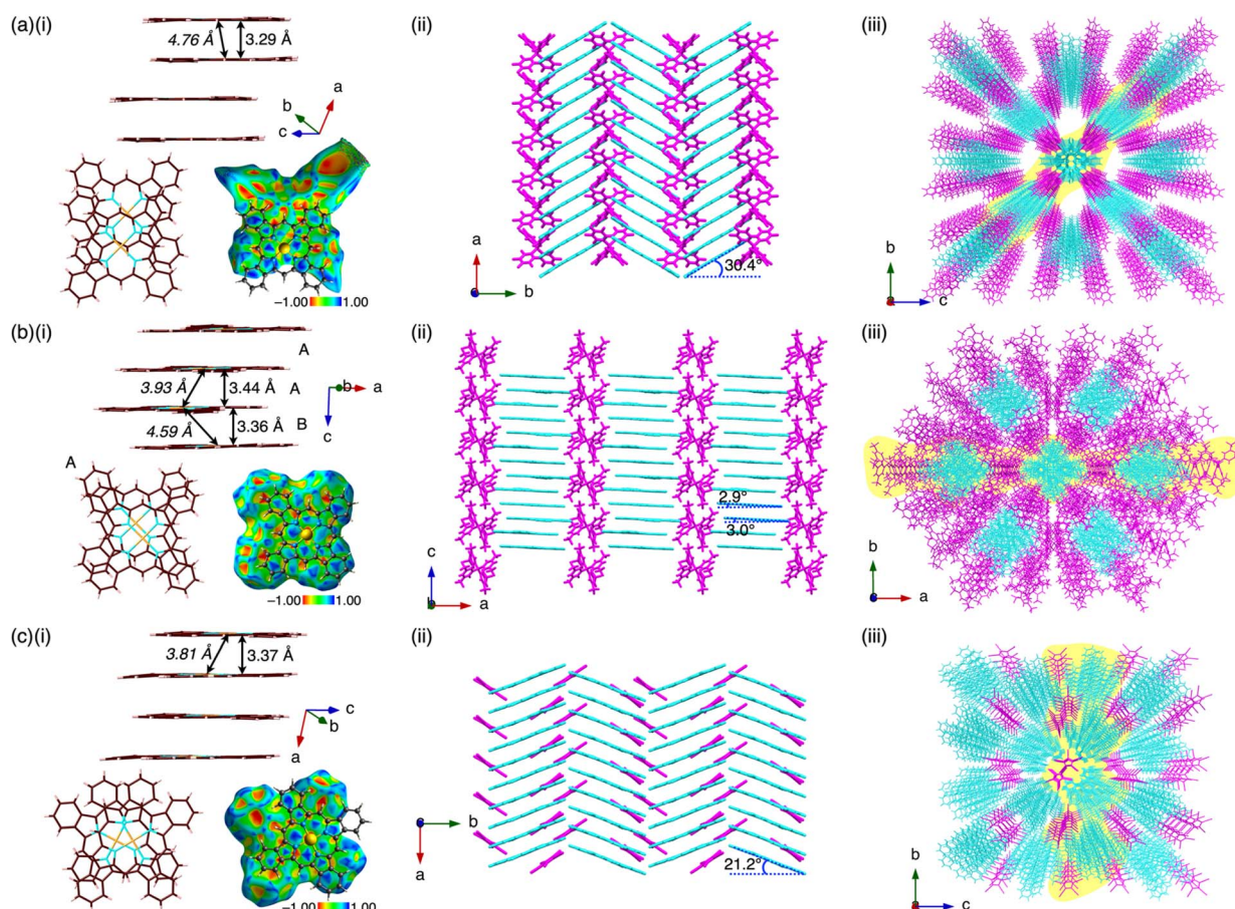


Fig. 5 Single-crystal X-ray structures of (a) $1\text{au}^+\text{-FABA}^-$, (b) $1\text{au}^+\text{-BARF}^-$ and (c) $1\text{au}^+\text{-PCCp}^-$: (i) side views of the columnar structures of stacked 1au^+ with stacking and Au...Au (italic) distances and top views along with Hirshfeld surface mapped over the shape-index property of the stacked dimer, (ii) packing structures of columns of stacked 1au^+ and counteranions and (iii) packing structures with the yellow parts corresponding to the packing structures in (ii). In (b), the different types of stacking arrangements are indicated by A and B (stacking arrangement A is shown as a representative). Atom colour codes in (i): brown, pink, blue and orange refer to carbon, hydrogen, nitrogen and gold, respectively. Colour codes in (ii and iii): cyan and magenta refer to cation and anion parts, respectively.



To evaluate the stacking behaviour of 1au^+ in the crystal structures, energy decomposition analysis (EDA)^{16,17} was performed using the fragment molecular orbital (FMO) method (FMO2-MP2) with mixed basis sets including NOSeC-V-DZP with MCP with NOSeC-V-TZP with MCP for Au.^{18,19} The EDA calculations using FMO yielded E_{es} , E_{disp} , E_{ct} and E_{ex} (energies for electrostatic, dispersion, charge-transfer forces and exchange repulsion, respectively) and E_{tot} (total energy). In the columnar structure of 1au^+ in 1au^+ -FABA⁻, an E_{tot} of -164.2 kcal mol⁻¹ is observed, whereas E_{disp} and E_{es} are -214.0 and 6.5 kcal mol⁻¹, respectively, indicating that E_{disp} is a major force in the stacking structure (Fig. 6 and S58†). EDA calculations for 1au^+ -BARF⁻ and 1au^+ -PCCp⁻ also elucidated similar energy balances for neighbouring π -electronic ions (Fig. S59 and S60†).

Crystal-state absorptions of the ion pairs 1au^+ -FABA⁻, 1au^+ -BARF⁻ and 1au^+ -PCCp⁻ were evaluated *via* optical microscopy for spectroscopic examination (Fig. S75 and S76†). In 1au^+ -FABA⁻, the absorptions at 587 and 623 nm, which are slightly blue- and red-shifted, respectively, compared to those of the monomer in DMSO (616 nm), are ascribable to the exciton coupling in a predominantly J-like arrangement and also very weak coupling for orthogonally arranged transition dipole moments (Fig. 7 and S62†). These behaviours are derived from the D_{4h} geometry of 1au^+ . Similar to 1au^+ -FABA⁻, the crystal-state absorption of 1au^+ -PCCp⁻ shows absorptions at 588 and 623 nm, which are ascribable to the exciton coupling of stacked 1au^+ . On the other hand, 1au^+ -BARF⁻ mainly exhibits a blue-shifted broad absorption at 585 nm with a shoulder at 654 nm. The blue-shifted absorption can be attributed to the larger contribution of the H-like arrangement of transition dipoles in the stacked 1au^+ (Fig. S62†).

Charge-segregated assemblies of π -electronic ion pairs show fascinating electric conductivity properties. The electric conductivity properties of stacked 1au^+ in the ion pairs (1au^+ -FABA⁻, 1au^+ -BARF⁻ and 1au^+ -PCCp⁻) were evaluated by flash-photolysis time-resolved microwave conductivity (FP-TRMC) measurements (Fig. 8a and S91†).²⁰ Electric conductivity ($\phi\Sigma\mu$) values of 4.6×10^{-9} , 9.2×10^{-9} and 2.9×10^{-9} m² V⁻¹ s⁻¹ were

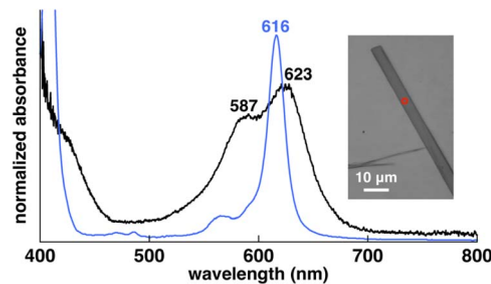


Fig. 7 Solid-state UV/vis absorption spectra of 1au^+ -FABA⁻ in the single crystal (black line) and in DMSO (4 μM) (blue line) as a reference (inset: photograph of the single crystal (red circle indicates the position of measurement)).

observed for the longer axes of the respective single crystals. Clear anisotropic electric conductivity was shown in 1au^+ -BARF⁻ (7.9×10^{-9} m² V⁻¹ s⁻¹ for the shorter axis). These values are comparable to and greater than those of previously reported charge-segregated assemblies.³ The order of the $\phi\Sigma\mu$ values, 1au^+ -BARF⁻ > 1au^+ -FABA⁻ > 1au^+ -PCCp⁻, is consistent with the theoretically estimated transfer integrals t at PW91/TZP for the stacked 1au^+ units in the crystal structures, showing hole transfer integrals $|t|_{\text{h}}$ of 118.5/50.5, 21.3 and 9.8 meV, respectively (Fig. 8b and S63†).²¹ Furthermore, theoretically estimated HOMO band dispersions using the tight-binding approximation²² for the stacked 1au^+ in the single-crystal structures of 1au^+ -FABA⁻, 1au^+ -BARF⁻ and 1au^+ -PCCp⁻ exhibited one-dimensional band structures, which are consistent with the stacking structures of 1au^+ (Fig. S64†). The Fermi levels lie in the middle of the band gaps, suggesting that charge-segregated assemblies comprising stacked 1au^+ exhibit semiconductive behaviours. In the discussed ion pairs, decreased on-site Coulomb repulsion between stacked π -electronic cations with expanded π -electronic systems would induce hole transport rather than electron transport. The conductivity transients

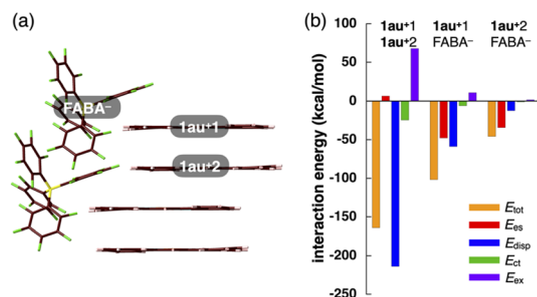


Fig. 6 Decomposition of the total intermolecular interaction energies (E_{tot}) of 1au^+ -FABA⁻ for (a) the single-crystal X-ray structure and (b) estimated interaction energies (kcal mol⁻¹) according to EDA based on the FMO2-MP2 method using a basis set of NOSeC-V-DZP with MCP with NOSeC-V-TZP with MCP for Au (see Table S4† for the complete data list). Colour codes in (i): brown, pink, blue, yellow, green and orange refer to carbon, hydrogen, nitrogen, boron, fluorine and gold, respectively.

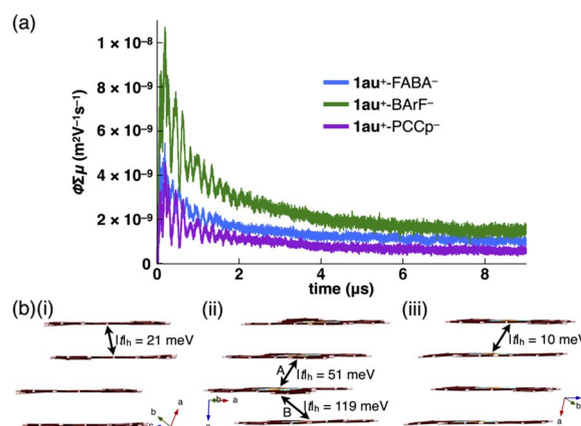


Fig. 8 (a) Photoconductivity transients observed upon excitation at 355 nm, 9.1×10^{15} photons per cm² per pulse, for the long axis of the single crystals of 1au^+ -FABA⁻ (blue), 1au^+ -BARF⁻ (green) and 1au^+ -PCCp⁻ (purple) and (b) stacked 1au^+ in the single crystal structures of (i) 1au^+ -FABA⁻, (ii) 1au^+ -BARF⁻ and (iii) 1au^+ -PCCp⁻ and hole transfer integrals ($|t|_{\text{h}}$) estimated at PW91/TZP. Stacking modes A and B in (ii) correspond to Fig. 5b(i).



recorded under an SF₆ atmosphere with negligible quenching of charge carriers suggest a major contribution from holes photo-injected into the assemblies (Fig. S92†).²³

Charge-segregated assemblies in pseudo-polymorph less-crystalline states

Substituent-free benzoporphyrins in the forms of free base and metal complexes have been known to show highly crystalline states after heating the corresponding bicyclo[2.2.2]octadiene precursors in the film state.^{6h} In contrast, **1au**⁺ as ion pairs can form bulk materials that are not single crystals *via* recrystallization from appropriate solvents. Precipitation of **1au**⁺-FABA⁻ from acetone/*n*-hexane provided a material (labelled as **1au**⁺-FABA⁻_p) that appeared to differ from the single crystals (Fig. 9a inset).²⁴ The synchrotron XRD of **1au**⁺-FABA⁻_p at 25 °C exhibited broad peaks instead of crystalline diffraction, suggesting the formation of an LeC state for the obtained precipitates. The diffraction peaks of 1.81, 1.04, 0.90, 0.68, 0.60, 0.52, 0.50, 0.45, 0.41 and 0.34 nm, showing Debye-Scherrer rings, were assigned to the *hkl* parameters derived from a hexagonal pattern (100, 110, 200, 210, 300, 220, 310, 400 and 320) as *a* = 2.09 nm (Fig. 9a and S81†).²⁵ The intense peak at 0.34 nm was assigned to 001 as the stacking distance of **1au**⁺ in the hexagonal columnar (Col_h) structure (*Z* = 1 for ρ = 1.79 g cm⁻³) (Fig. 9b). Interestingly, heating the powder sample of **2au**⁺-FABA⁻ at 190 °C (labelled as **1au**⁺-FABA⁻_H) also formed a Col_h structure identical to that of **1au**⁺-FABA⁻_p (Fig. S79–S82†). In contrast to the tilted **1au**⁺ plane along the *a*-axis in the single crystal of **1au**⁺-FABA⁻, the **1au**⁺ plane in **1au**⁺-FABA⁻_p should be arranged perpendicularly to the stacking axis, as indicated by the intracolumnar stacking period of 0.34 nm. On the basis of the Col_h structure with *a* = 2.09 nm and the sizes of **1au**⁺ and FABA⁻, columnar structures comprising less-ordered stacking of **1au**⁺, as indicated by cyan circles, are located close to FABA⁻, as indicated by magenta circles (Fig. 9b(ii)).

The proximal location of **1au**⁺ and FABA⁻ is also suggested by the optimized structure of dimeric **1au**⁺-FABA⁻ using B3LYP-GD3BJ/6-31+G(d,p) with LanL2DZ for Au (Fig. 9c). FABA⁻ should be paired with several **1au**⁺ in the proximal location, although the XRD pattern suggests that the location of FABA⁻ is less clear, probably indicating an amorphous-like state. In light of the stoichiometry of the constituents and their contrasting planar and bulky shapes, FABA⁻ would be observed in three sites on average among the **1au**⁺ columns, and, in another cross section according to the **1au**⁺ planes, the anions should be located in the other three sites. As a result, FABA⁻ can be hexagonally arranged around the **1au**⁺ columns. The Col_h structure suggested by the XRD pattern is clearly demonstrated by all-atom molecular dynamics (MD) simulations at 25 °C after 100 ns of the equilibration run (Fig. 9d and S66†). Notably, as the initial structure for the MD simulation, the columns comprising tilted **1au**⁺ units, as observed in the single-crystal structure, are transformed to a structure with barely tilted **1au**⁺. The combination of planar **1au**⁺, suitable for stacking, and bulky FABA⁻, with a less-ordered arrangement *via* noncovalent interactions induces the LeC state.^{27,28} The less-ordered FABA⁻

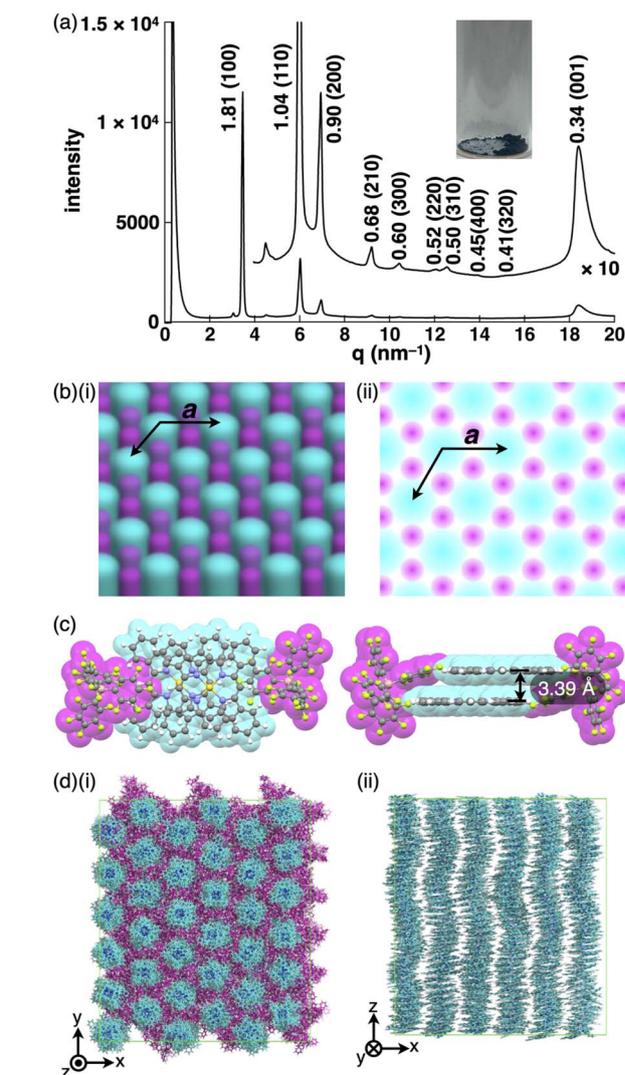


Fig. 9 (a) Synchrotron XRD pattern of **1au**⁺-FABA⁻_p at 25 °C for the sample obtained by precipitation from acetone/*n*-hexane (inset: photograph of the precipitates), (b) schematic representation for the Col_h packing structure: (i) packing diagram and (ii) top view of Col_h (**1au**⁺ and FABA⁻ are represented in cyan and magenta, respectively), (c) top and side views of the optimized structure of **1au**⁺-FABA⁻ as a dimer using B3LYP-GD3BJ/6-31+G(d,p) with LanL2DZ for Au and (d) snapshot of the MD simulation result for **1au**⁺-FABA⁻_p after 100 ns of the equilibration run at 25 °C showing (i) the top view of the packing diagram (**1au**⁺ and FABA⁻ are represented in cyan and magenta, respectively). In (b) (ii), magenta circles suggest the possible locations of FABA⁻ with a 50% occupancy rate on average in a cross section, whereas cyan circles show the diameter of **1au**⁺ columns in slipped stacking.

interferes with the ordered packing of **1au**⁺ for crystallization. **1au**⁺-FABA⁻_p maintains the Col_h structure up to 195 °C and is converted to a complicated crystalline state at higher temperatures, with decomposition at 355 °C. This behaviour is also supported by differential scanning calorimetry (DSC) analysis (Fig. S78†). The appearance of **1au**⁺-FABA⁻_p as a dark blue powder in polarized optical microscopy (POM) was maintained through the heating process. The condition-dependent



assembly (single-crystal and LeC states) as pseudo-polymorphism²⁹ is fascinating for tuneable properties according to the arrangement of building blocks. The Col_h LeC state, in the absence of aliphatic chains, is rare²⁸ but can be achieved by pairing the planar π -electronic cation with a bulky counteranion.³⁰

The details of the structures of 1au^+ -FABA⁻ were investigated by solid-state NMR (SSNMR) spectroscopy. ¹³C NMR was performed using cross-polarization magic-angle spinning (CPMAS) (Fig. 10 and S86†). Dipolar dephasing experiments were conducted to support the signal assignment for the ion pair (Fig. S86 and S88†). The broad signals at 132.2, 120.8, 93.4 and 89.3 ppm were assigned to 1au^+ , whereas those at 149.7, 137.7 and 126.0 ppm were assigned to FABA⁻ (Fig. 10a). ¹³C CPMAS NMR of 1au^+ -FABA⁻_H showed slightly broader signals than 1au^+ -FABA⁻_P, suggesting the formation of a more disordered arrangement of constituting ions (Fig. 10b). In contrast, ¹³C CPMAS NMR for 1au^+ -FABA⁻ as single crystals showed narrower signals than those of 1au^+ -FABA⁻_P and 1au^+ -FABA⁻_H (Fig. 10c), suggesting that the broader signals of 1au^+ -FABA⁻_P and 1au^+ -FABA⁻_H indicated a less-ordered arrangement of ions that form Col_h structures. Similar signal broadening was observed for ¹⁹F and ¹¹B MAS NMR of 1au^+ -FABA⁻_P and 1au^+ -FABA⁻_H (Fig. S89 and S90†). Moreover, the characteristic up-field split signals at 93.4 and 89.3 ppm in 1au^+ -FABA⁻_P are derived from unsubstituted *meso*-carbons,³¹ as also suggested by theoretically estimated NMR signals for 1au^+ using B3LYP/6-311+G(d,p) with SDD for Au (Fig. S65†).³² Such signal splitting is also observed in the ¹³C CPMAS NMR of the single crystals due to slipped stacking of 1au^+ observed in the crystal structure of 1au^+ -FABA⁻ (Fig. 5a(i)).³² These observations, along with the XRD analysis and MD simulation, support a less-ordered slipped stacking structure for 1au^+ in a column of Col_h LeC states, wherein the 1au^+ planes are more perpendicularly arranged to the column compared to those in the single crystal.³³

The formation of such an LeC state was also observed in the precipitate of 1au^+ -BARF⁻_P prepared from acetone/*n*-hexane.³⁴ Similar to 1au^+ -FABA⁻_P, 1au^+ -BARF⁻_P shows a Col_h LeC state (*a*

= 2.28 nm, *c* = 0.34 nm, and *Z* = 1 for ρ = 1.71 g cm⁻³) (Fig. S84 and S85†), also exhibiting condition-dependent pseudo-polymorphs. Col_h LeC states in the precipitates were observed for 1au^+ -FABA⁻ and 1au^+ -BARF⁻, which formed orthorhombic and monoclinic single-crystal packing structures, respectively. Such pseudo-polymorphic phenomena are fascinating because stacking of 1au^+ can be easily controlled by the assembly conditions. The electric conductivity ($\phi\Sigma\mu$) values of LeC-state 1au^+ -FABA⁻_P, 1au^+ -FABA⁻_H and 1au^+ -BARF⁻_P were estimated to be 1.2×10^{-9} , 1.0×10^{-9} and 3.4×10^{-9} m² V⁻¹ s⁻¹, respectively, suggesting that electrically conductive pathways also exist in the columnar structures of LeC materials, although the values are smaller than those of the corresponding single crystals (Fig. S93†).

Conclusions

Ion-pairing assemblies in charge-segregated modes were constructed from a highly planar expanded π -electronic cation in combination with counteranions. Charge-segregated assemblies were formed with both planar and bulky counteranions by means of stable stacked structures of the expanded π -electronic cation. The stacking arrangement and resulting absorption spectra in the single crystals were modulated by coexisting anions. Depending on crystallizing solvent conditions, ion pairs with bulky borate anions also provided LeC states as pseudo-polymorphs of their single crystals. Both single crystals and LeC states exhibited electric conductive properties due to stacking of the π -expanded porphyrin Au^{III} complex.³⁵ It is noteworthy that the discussed planar π -electronic cation, benzoporphyrin Au^{III} complex, is soluble in organic solvents in the ion-pairing states. Ion pairing is an effective strategy to increase the solubility of planar π -electronic systems for their facile handling and the fabrication of assembled structures with ordered arrangements. Further modifications of charged π -electronic systems would lead to ion-pairing assemblies that can be applied for functional electronic materials and devices.

Data availability

Data supporting the work in this publication are available *via* the ESI and associated crystallographic data.

Author contributions

H. M. designed and conducted the project. Y. H., R. N., Y. M. and H. T. carried out the synthesis, characterization and property examinations. W. C. and S. Se. evaluated the electric conductivities. S. Sa., H. B., Y. I. and G. W. conducted the MD calculations. K. B. and K. T. conducted the transfer integral calculations. K. D., S. O., K. H. and T. N. evaluated the SSNMR. Y. I. and T. A. recorded the solid-state absorption spectra. K. O. supported the discussion on the assemblies.

Conflicts of interest

There are no conflicts of interest to declare.

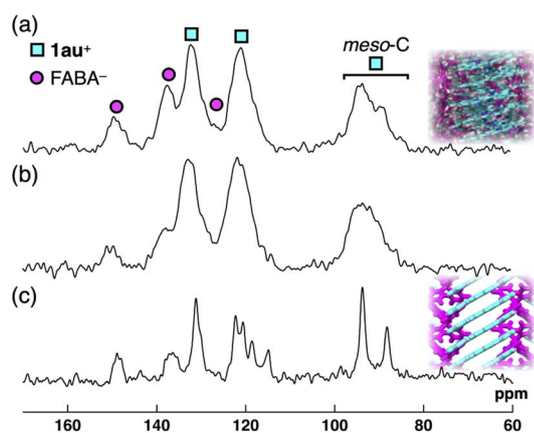


Fig. 10 Solid-state ¹³C CPMAS NMR spectra of (a) 1au^+ -FABA⁻_P, (b) 1au^+ -FABA⁻_H and (c) 1au^+ -FABA⁻ as single crystals recorded at 20 kHz MAS spinning frequency at r.t. with the corresponding packing diagrams.



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