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Orthogonal, modular anion—cation and cation—anion self-assembly using pre-programmed anion binding sites†

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Subcomponent self-assembly relies on cation coordination whereas the roles of anions often only emerge during the assembly process. When sites for anions are instead pre-programmed, they have the potential to be used as orthogonal elements to build up structure in a predictable and modular way. We explore this idea by combining cation (M^+) and anion (X^-) binding sites together and show the orthogonal and modular build up of structure in a multi-ion assembly. Cation binding is based on a ligand (L) made by subcomponent metal-imine chemistry ($M^+ = Cu^+$, Au^+) while the site for anion binding ($X^- = BF_a^-$, ClO_a^-) derives from the inner cavity of cyanostar (CS) macrocycles. The two sites are connected by imine condensation between a pyridyl-aldehyde and an aniline-modified cyanostar. The target assembly [LM-CS-X-CS-ML], * generates two terminal metal complexation sites (LM and ML) with one central anion-bridging site (X) defined by cyanostar dimerization. We showcase modular assembly by isolating intermediates when the primary structure-directing ions are paired with weakly coordinating counter ions. Cation-directed (Cu⁺) or anion-bridged (BF₄⁻) intermediates can be isolated along either cation-anion or anion-cation pathways. Different products can also be prepared in a modular way using Au^+ and ClO_4^- . This is also the first use of gold() in subcomponent self-assembly. Pre-programmed cation and anion binding sites combine with judicious selection of spectator ions to provide modular noncovalent syntheses of multicomponent architectures.

Introduction

The role of anions in cation-directed self-assembly is growing in importance. They act as templates¹⁻³ for coordination cages,⁴ helicates⁵ and knots,⁶ triggers for switching between structures,^{1,7,8} they solubilize⁹ the final assembly in desired solvents, and get encapsulated as guests.¹⁰ These many roles have grown on the back of metal-directed subcomponent self-assembly¹¹ involving the elegantly direct and *in situ* Schiff-base condensation of modular ligands. Across all these roles, the anion's

assembly¹²⁻¹⁵ where metal ion coordination remains the primary structure-directing interaction. The anions have only rarely been explored as orthogonal motifs for building up structural features in a manner that matches metal ions. A notable exception¹⁶ from Jansone-Popova is the synergistic one-pot self-assembly of a helicate around divalent metal cations, (Cu^{2+}) and sulfate anions (SO_4^{2-}) all directed by heteroditopic ligands composed of separate cation and anion binding sites. Herein, we investigate the orthogonality between anion coordination chemistry^{17,18} and cation coordination for the modular build-up of ion-driven assemblies (Fig. 1).

binding site is built up dynamically by the cation-directed self-

The realization of structure-directing anion coordination is enabled by advances in receptor design. ^{19–23} Examples include anion helicates²⁴ with ditopic ligands bearing terminal ureas wrapped around phosphate (PO₄^{3–}). ²⁵ Cages, ²⁶ *e.g.*, tetrahedra, have combined urea motifs with highly charged phosphates in the vertices. ²⁷ Polymers have been developed, *e.g.*, the Texassized box electrostatically stabilizes the polymerization of terephthalates, ²⁸ and cyanostar stabilizes polymerization of diphosphonates²⁹ or diphosphates. ³⁰ The counter cations in these cases, once again, play secondary roles.

Examples of multi-ion assemblies with anions and cations serving as equal partners are under-represented.³¹⁻³⁵ Ion-pair

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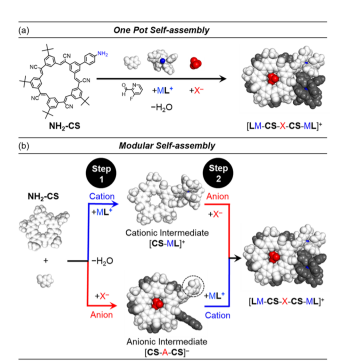


Fig. 1 Schematic summary of the (a) one-pot and (b) two-step self-assembly pathways that use pre-programmed and orthogonal coordination of cations and anions. In the resulting assembly (at right) the top cyanostar is shown in white and the bottom one is shown in dark grey for clarity. The Cu^+ cation is shown in blue, and the BF_4^- anion shown in red. The pyridyl-based aldehyde is highlighted by the dotted circle in the anionic intermediate.

receptors can bind cations, anions and ion pairs but such receptors are not used to build up structure unless both ions³⁶ bind at the same time to pre-formed metal and anion binding sites.³⁷ The most powerful among them afford cooperativity^{38–40} with selective capture of ion pairs over the separate ions.¹⁶ Such systems rely on one-pot processes with mutual binding of anion and cation without the isolation of intermediates for modular assembly.

Orthogonality between metal coordination and anion recognition offers advantages for self-assembly. When cation and anion do not interact either with each other or the other's binding sites then the assembly should be pre-programmable in a predictable way. Orthogonality also lends modularity. Thus, swapping one type of anion for another alters the composition but not the structural features of the final assembly. These characteristics of predictability and modularity can potentially be demonstrated by separating a one-pot multi-component self-assembly (Fig. 1a) into discrete steps (Fig. 1b).

Herein, we explore the orthogonality and modularity of ion-driven assembly by combining cation-directed subcomponent self-assembly between imines and metal moieties 42 (ML; L = POP, PPh₃; Fig. 2b) with pre-programmed anion binding sites (Fig. 2). 41 We use cyanostar macrocycles (CS, Fig. 2c) 22 for strong anion coordination (X $^-$, Fig. 2a). Structure-directing cations (M = Cu $^+$, Au $^+$; Fig. 2b) and anions (X $^-$ = BF $_4$ $^-$, ClO $_4$ $^-$; Fig. 2a) can be added to form target assemblies in one-pot confirming the orthogonality of assembly. Modularity is demonstrated by first isolating supramolecular intermediates of either ion (Fig. 1b). The intermediates are accessed by using non-coordinating counterions (Fig. 2d and e). The cationic intermediate [LCu-CS] $^+$, and the anionic intermediate [NH $_2$ CS-BF $_4$ -NH $_2$ CS] $^-$, can be

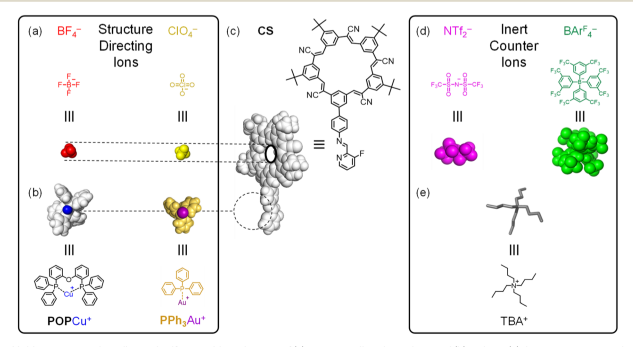


Fig. 2 Multi-component ion-directed self-assembly makes use of (a) structure-directing anions and (b) cations, (c) the pre-programmed anion-binding ligand, cyanostar (CS), as well as (d) inert counter anions and (e) counter cations. The Cu^+ is blue, and the Au^+ cation is dark purple. Their complexes are distinguishable by their ligands where POP is white and PPh₃ is gold. Anions are color-coordinated where BF_4^- is red, ClO_4^- is yellow NTf_2^- is purple, and $BArF_4^-$ is green.

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isolated by trituration and used in subsequent reactions. Exemplary studies with copper(1) and tetrafluoroborate indicate the intermediate can be used as a substrate for a second coordination step to form the final assembly [LCu-CS-BF₄-CS-CuL], (Fig. 1b). Modularity was also demonstrated using Au⁺ and ClO₄⁻ ions. Surprisingly, this is the first example of subcomponent selfassembly⁴³ with Au(1). We used other anions (phosphates, phosphonates, bisulfate) to explore the limits of this strategy. Thus, pre-programmed cation and anion binding sites serve as orthogonal interactions that form supramolecular assemblies and help contribute to the growing exploration of methods^{44,45} capable of orthogonal, modular noncovalent synthesis.

Results and discussions

Design of the components used in cation-anion self-assembly

In order to demonstrate orthogonality, we needed to simplify the number of potential products that form. We used ancillary ligands on the metals to block the possibility of cross linking. We substituted just one of the five cyanostar arms (Fig. 1a) to enable coordination of one metal ion moiety to the macrocycle as opposed to five in the C_5 -symmetric cyanostar. The only drawback to mono-substitution is the presence of overlapping peaks in the NMR spectra. These ligand and receptor features all favor formation of fewer species.

We used knowledge of the recognition patterns of the binding sites to design the various components.42 The monosubstituted cyanostar46 (NH2-CS, Fig. 1) contains a 4-aminophenyl group. We expect this aniline to enable imine formation upon reaction with aldehydes in the presence of metal ions.43 We also anticipate anion coordination with NH₂-CS will match the behavior of parent cyanostar (pCS)22 to direct formation of a 2:1 sandwich complex with a bridging anion, X⁻. For metal complexation, we selected a fluoro-substituted picolinaldehyde, F-PyCHO, as the partner to the NH2-CS aniline for subcomponent assembly.

We identified two metal precursors to explore modularity. The copper(1) complex of bis(2-(diphenylphosphino)phenyl) ether (POP) as the bis-acetonitrile solvato species, {POPCu(MeCN)₂}⁺, was selected as the first. The POP ligand is known to terminate the assembly after imine coordination42 and the acetonitrile is labile. The second is a gold(1) triphenylphosphine moiety, {**PPh**₃Au}⁺. Gold(I) has never been used in subcomponent self-assembly and it has unique coordination geometries.⁴⁷ Gold(1) is typically two coordinate,⁴⁸ but three has also been observed when using σ-donors of comparable strength.49 The imine nitrogen is a relatively strong donor,50 therefore a two-coordinate geometry is possible. Alternatively, the chelating imine is similar to phenanthroline that can favor three-coordinate geometries with gold(1).51 A model gold(1) complex was examined to confirm the preferred mode of coordination prior to the cation-anion self-assembly.

Strategies for orthogonal cation-anion self-assembly

Our strategy for demonstrating the principle of cation and anion self-assembly relies on two design principles: orthogonal

binding sites and weak ion-pair interactions. The cyanostar component has an anion-binding site native to its cavity and one cation binding site added to its exterior. Based on their preferred reactivity,22,42 these sites do not compete with each other. The second criterion requires knowledge of ion coordination and ion pairing. Copper(1) ions are known to coordinate with free imines while BF₄⁻ is complementary to the cyanostar. These ions (Cu⁺ and BF₄⁻) do not display strong interactions with each other in the solvents we use here.52 While BF₄ anions are considered weakly coordinating to metals, they display some of the strongest binding affinities to cyanostar.22 But not all anions that bind tightly to cyanostar are inert to metal coordination (see below). These design principles are anticipated to be transferrable to other pairs of ions. To this end, we investigated gold(I), Au⁺, and perchlorate, ClO₄⁻ to verify the generality of this assembly strategy.

Finally, isolation of intermediates requires selection of nonbinding counter ions. Their use enables structure-directing ions to be delivered separately and sequentially. Bulky anions tetrakis(3,5-bis(trifluoromethyl)phenyl)borate (BAr^F₄, Fig. 2d) and triflimide (NTf2-, Fig. 2d) fulfill the requirements for weak counter anions. They are well-established to coordinate poorly to both metal ions53,54 and cyanostar.55 NTf2 has a low affinity $(250 \pm 30 \text{ M}^{-1})$ for cyanostar.⁵² Similarly, the TBA⁺ counter cation (tetrabutylammonium, Fig. 2e) is not known to coordinate to metal-binding ligands. While the ion pairing of TBA⁺ in dichloromethane with our target anions, e.g. $BF_4^- \log K = 4.6$ (ref. 56) and ClO_4 log K = 4.24 (ref. 57) is not negligible, they are weaker than the coordination with parent cyanostar, e.g., $\log \beta(pCS-ClO_4^--pCS) \sim 12.^{22}$ Thus, any ion pairing does not interfere.

One-pot, multi-component self-assembly using Cu⁺coordination and cyanostar-BF₄ binding

Orthogonality between the two binding sites was verified using a traditional one-pot assembly. The structure-directing Cu⁺ and BF₄ ions can be added as a single precursor (Fig. 3a). The target assembly [POPCu-CS-BF4-CS-CuPOP]+, is a dicopper and anionbridged dimer. The copper binding sites are products of imine condensation between NH2CS and F-PyCHO in the presence of the salt [POPCu(MeCN)₂]·BF₄. Mixtures of the three components proceed near-quantitatively through to the target assembly. The product was isolated as a BF₄⁻ salt after solvent removal. Exploration of the assembly conditions revealed that excess aldehyde enhanced the yield and unreacted aldehyde could be removed with ether washes. Excess NH2CS was not investigated on account of its poorer solubility relative to the aldehyde.

Product identity was confirmed using NMR spectroscopy and electrospray ionization mass spectrometry (ESI-MS). The wellresolved ¹H NMR peaks (Fig. 3b) indicate high-fidelity Cu(1)complexation and BF₄ binding in the target assembly. Diffusion ordered spectroscopy (DOSY, Fig. 3b) verified formation of a single monodisperse species. ESI-MS shows the peak for the parent assembly [POPCu-CS-BF₄-CS-CuPOP]⁺ at 3407.2455 m/z(Fig. 3c). Under ESI-MS conditions, minor multimer peaks are

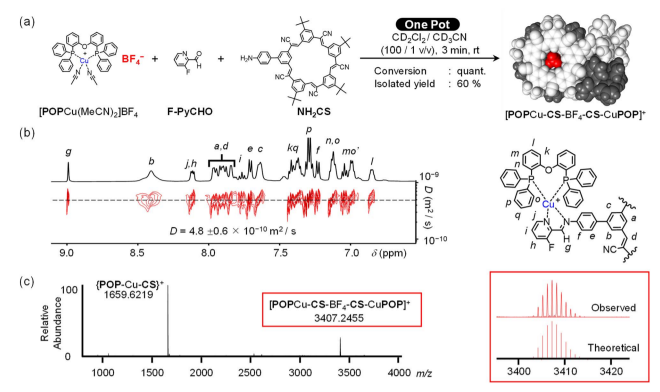


Fig. 3 (a) One-pot reaction to form $[POPCu-CS-BF_4-CS-CuPOP]^+$ and characterization by (b) DOSY NMR $(CD_2Cl_2, 2 \text{ mM})$, and (c) ESI-MS spectrum of assembly $(CH_2Cl_2, 2 \text{ mM})$.

observed from 0.5 Dalton peak-to-peak separation associated with a dimer of the target: [**POPCu-CS-BF**₄-**CS-CuPOP**]₂²⁺. We also see a daughter peak corresponding to the anion-free half of the assembly assigned to the {**CS-CuPOP**}⁺ moiety at 1659.6219 m/z.

Imine formation and metal coordination show characteristic changes in the chemical shifts of key protons upon formation of the target assembly (Fig. S53†). The aldehyde (10.16 ppm) is consumed to form the imine (9.01 ppm). 58 Other proton close to the metal binding site are characteristic of imine bond

(a) (b) (c) -145 -150 -155 Chemical Shift (ppm)

Fig. 4 19 F NMR (a) [POPCu-CS-BF₄-CS-CuPOP]⁺ created in one-pot, (b) [pCS-BF₄-pCS]⁻, and (c) TBABF₄ in CD₂Cl₂ (298 K, 376 MHz).

formation and copper complexation.⁴² The aniline's ring hydrogen (H_f , 6.84 ppm) shifts downfield (7.23 ppm), and the phosphine protons (H_k) shift modestly from 7.02 to 7.09 ppm.

Anion binding in the target assembly is confirmed by ¹H NMR (Fig. S53†). Peak assignments were based on ROESY studies, controls, and assignments of a related cyanostar. ²¹ The cluster of peaks at 8.76 ppm in NH₂CS are assigned to the outer H_b protons of the macrocycle. After one-pot assembly, these shift upfield to 8.41 ppm matching the parent cyanostar. ²² The ¹⁹F NMR signal from BF₄⁻ shows a single broad peak at –150 ppm (Fig. 4a) consistent with exchange averaging between complexed and outer-sphere ions. ⁵⁹ The spectrum of BF₄⁻ as a TBA⁺ salt shows a sharp peak at –152 ppm (Fig. 4c) while the 2:1 complex formed between BF₄⁻ and the parent cyanostar (Fig. 4b) resonates as a sharp peak at –148 ppm.

Cation-directed supramolecular intermediates

Formation of the desired multi-component assembly [POPCu-CS-BF₄-CS-CuPOP]⁺, in one pot (Fig. 1a) with high yields provides a basis for exploring stepwise synthetic strategies (Fig. 1b). We separately investigated generation of cation-directed and anion-directed intermediates by relying on their pre-programmed binding sites. In traditional cation-directed assembly, it is not possible to isolate the anion-directed intermediates because the anion binding site is built *in situ* instead of being pre-programmed.

The cation-directed supramolecular intermediate was prepared under similar conditions as the one pot but using the

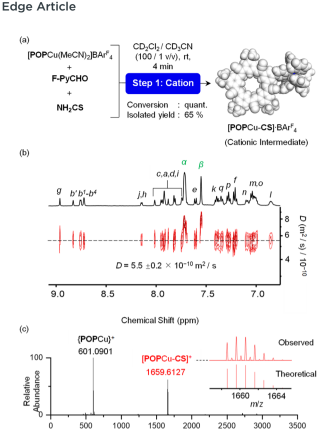


Fig. 5 (a) Reaction to form [POPCu-CS]⁺ and characterization by (b) DOSY NMR (CD₂Cl₂, 2 mM), and (c) ESI-MS spectrum (CH₂Cl₂, 2 mM).

copper precursor [**POP**Cu(MeCN)₂]⁺ as a BAr^F₄⁻ salt (Fig. 5a). Identification of the desired monomeric intermediate [**POP**Cu-CS]⁺ was obtained from the ESI-MS with a peak at 1659.6127 m/z (Fig. 5c). DOSY confirmed formation of a single monodisperse species (Fig. 5b). Characteristic shifts in NMR peaks confirm formation of intermediate [**POP**Cu-CS]⁺ (Fig. S54†). The BAr^F₄⁻ anion peaks are unshifted indicative of its innocence. ROESY data (Fig. 6) on the intermediate confirms imine formation and {**POP**Cu}⁺ complexation.

Modular synthesis using a universal cation-directed supramolecular intermediate with different anions

Preparation of the cationic intermediate [POPCu-CS]⁺ allowed investigation of different anion-driven products with BF₄⁻ or ClO_4^- (Fig. 6 and 7) to illustrate modularity. Addition of TBABF₄ to [POPCu-CS]⁺ led to the cyanostar-mediated dimerization around the bridging BF₄⁻ anion. ¹H NMR titration monitoring addition of TBABF₄ (Fig. 6b) to [POPCu-CS]⁺ indicate formation of a 2:1 complex with saturation at 0.5 equivalents (Fig. 6b). Diffusion NMR confirmed formation of a single monodisperse species (Fig. S62†). The dimeric assembly [POPCu-CS-BF₄-CS-CuPOP],⁺, was verified using ESI-MS (Fig. 7a) matching the one prepared by the one-pot method (Fig. 3c). A peak at 2533.9194 m/z indicates formation of a higher-order trimer [(POPCu-CS)₃BF₄]²⁺ only under the ESI-MS conditions.^{20,29,30,57,60}

Consistent with the modularity of the cation–anion method, addition of TBAClO₄ generated the same types of signatures as

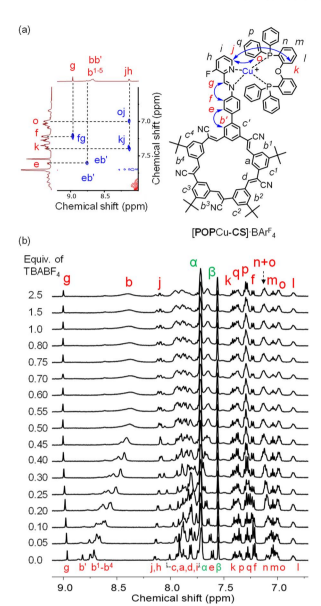


Fig. 6 (a) The aromatic region of ROESY NMR data of [POPCu-CS]- BAr^F_4 . (b) 1H NMR titration of [CS-POPCu]- BAr^F_4 (2 mM) with TBABF $_4$ (CD $_2$ Cl $_2$, 400 MHz, rt).

those seen with $\mathrm{BF_4}^-$. The dimeric assembly was seen by ESI-MS (Fig. 7b) with the same daughter and higher-order peaks. The diffusion coefficient matched the ones seen using the $\mathrm{BF_4}^-$ anion (Fig. S63†). The $^1\mathrm{H}$ NMR titrations also showed saturation at the expected 0.5 equivalence point TBAClO₄ (Fig. 7c).

Stepwise synthesis using anion-linked intermediates

The function of the pre-programmed anion binding site can be demonstrated by inverting the ion addition sequence using an anion–cation pathway. The anion-linked supramolecular intermediate [CS-BF₄-CS]⁻ (Fig. 8) was created by condensation of 2 equivalents each of NH₂-CS and aldehyde F-PyCHO in the presence of 1 equivalent of TBABF₄. The ESI-MS analyses showed the intermediate (Fig. S14†) present as a variety of

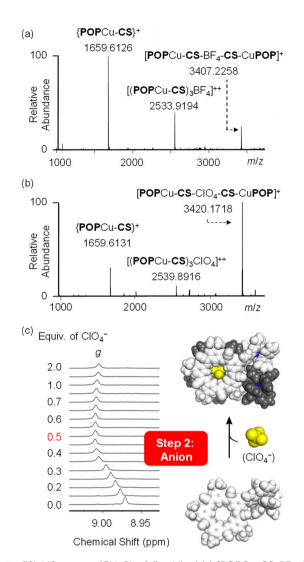


Fig. 7 ESI-MS spectra (CH $_2$ Cl $_2$, 0.5 mM) of (a) [POPCu-CS-BF $_4$ -CS-CuPOP] $^+$ and (b) [POPCu-CS-ClO $_4$ -CS-CuPOP] $^+$ formed by cation—anion pathway. (c) 1 H NMR titrations of intermediate [CS-POPCu]-BAr F_4 (2 mM) with TBAClO $_4$ (CD $_2$ Cl $_2$, 400 MHz, rt).

MeOH adducts under the conditions of the experiment. The 1: 4 ratio seen by NMR between the free aldehyde (**F-PyCHO**) and the imine suggests 80% reaction. Metal-ion chelation is required to drive complete condensation.

Addition of [POPCu(MeCN)₂]⁺ in the second step (Fig. 8) drove complete formation of the imine and the target assembly,

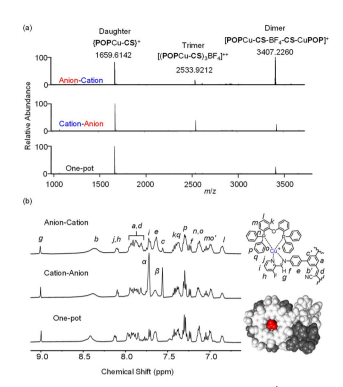


Fig. 9 (a) ESI-MS (CH_2Cl_2 , 0.5 mM) and (b) diagnostic 1H NMR peaks (CD_2Cl_2 , 0.5 mM) for [POPCu-CS-BF₄-CS-CuPOP]⁺ for the anion–cation stepwise, cation–anion stepwise, and one-pot assembly.

[POPCu-CS-BF₄-CS-CuPOP][†]. Notably, the product of the anion-cation assembly pathway shows the same diagnostic signatures in ESI-MS and NMR (Fig. 9) as the products of the one-pot and cation-anion methods confirming the same structure is made along three different synthetic pathways. The nature of the counter anions depends on the pathway used for self-assembly. In the cases where more than one counter anion is present, they are in exchange with the various species present in solution. For example, the ¹⁹F NMR spectrum of the gold assembly obtained *via* the cation-anion pathway shows two peaks. One results from BF₄⁻ in fast exchange between being its cyanostar-bound and unbound states while the other corresponds to the uncomplexed NTf₂⁻ anion (Fig. S30†).

Orientation of the macrocycles in the dimer

Structurally, the copper complexes in the product [POPCu-CS-BF₄-CS-CuPOP]⁺ can be either *syn*, *meta*, or *anti* (Fig. 10).

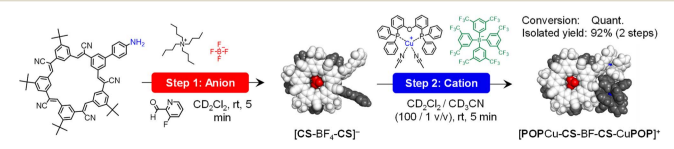


Fig. 8 Stepwise anion—cation synthesis.

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Fig. 10 1 H NMR titration of TBABF₄ into [POPCu-CS]⁺ (4 mM) in CD₂Cl₂ (298 K, 400 MHz).

Chemical Shift (ppm)

Molecular mechanics shows syn to be favored over meta (16 kJ mol⁻¹) and anti (86 kJ mol⁻¹), which is consistent with a related complex.⁴⁶ An NMR titration following addition of BF₄⁻ to [POPCu-CS]⁺ to form the dimerized product [CuCS-BF₄-CSCu]⁺ supports the syn geometry. Therein, the phosphine ligands are close enough to lower their symmetry into two different environments (H_o, Fig. 10). Molecular models on the syn geometry how one of the two phenyls engaging in π stacking. These findings suggest the syn geometry is favored by stabilizing contacts between coordinated POP ligands.

Use of gold(i) showcases modularity of cation-directed supramolecular intermediates

Gold(\mathfrak{i}) was used to investigate the modularity of assembly. To model the first use of gold(\mathfrak{i}) ions in subcomponent self-assembly, we prepared a complex using a simple aniline. We mixed a gold(\mathfrak{i}) precursor as a triflimide salt, [PPh₃Au]·NTf₂ with F-PyCHO and the aniline *p*-toluidine. X-ray diffraction studies showed formation of tricoordinate gold(\mathfrak{i}) cation (Fig. 11) with

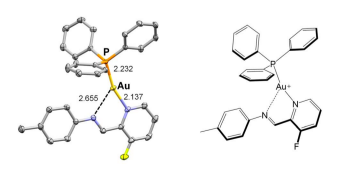


Fig. 11 Molecular structure of the three-coordinate gold(i) model complex as determined by X-ray crystallography (50% ellipsoids). Hydrogen atoms and counter anion omitted for clarity.

a chelating imine. ¹H and ¹⁹F NMR, and ESI-MS spectra (Fig. S36–S38†), are consistent with formation of the gold(t) complex seen in the solid state.

We examined the various synthetic pathways using the Au⁺ and BF₄⁻ ions. The target complex [PPh₃Au-CS-BF₄-CS-AuPPh₃]⁺ was formed along both one-pot and cation–anion pathways (Fig. 12a). Formation was confirmed by ESI-MS with the one-pot producing a peak at 3122.1669 m/z (Fig. S25†) matching the product of cation–anion assembly (Fig. 12b). Consistent with gold(i) complexes being less stable than copper(i), parent ion peaks have lower intensity than daughter ions.

Formation of the target assembly [PPh₃Au-CS-BF₄-CS-AuPPh₃]^{\dagger}, was verified by NMR. All resonances display the same diffusion coefficients (Fig. 12c) for the one-pot and cationanion assembly (Fig. S69†) and the same diagnostic peaks by 1 H-NMR (Fig. 13) and 19 F-NMR spectroscopy (Fig. S27†). These similarities indicate that the same assembly is being produced along different pathways.

The anion–cation stepwise pathway was also investigated. Using the anion-linked intermediate described earlier, addition of the $\mathrm{Au^+}$ precursor produced an $^1\mathrm{H}$ NMR spectrum differing from the one-pot and cation–anion pathways (Fig. S56†). The ESI-MS did not show any peaks consistent with target assembly. Adding excess $\mathrm{BF_4}^-$ to better match the one-pot pathway did not influence the outcome (Fig. S41†). Failure to access the target assembly by the alternative stepwise pathway is surprising and highlights the need to study the impacts of reaction order on multi-component supramolecular assemblies.

Failure modes using competitive anions

Not every anion that binds tightly to cyanostar^{29,57,60,61} is also weakly coordinating to metal cations. We were hoping to leverage this idea to target assemblies that could only be produced by pre-complexation of the anion. To explore this idea, we studied how metal complexes survived in the presence of organophosphates, which are known to ligate metals. These phosphates are also known to form threaded pseudorotaxane complexes with cyanostar.62 This study was undertaken using gold and copper complexes, dibutyl phosphate, dibenzyl phosphate, and the parent cyanostar, pCS, as models. However, precomplexation with cyanostar did not prevent the phosphates from decomposing the copper complexes in 24 hours (Fig. S71-S75†). A gold complex showed similar behavior (Fig. S76 and S77†) as did the cationic intermediate [PPh₃Au-CS]·NTf₂ (Fig. S77†). While we saw some evidence of a threaded pseudorotaxane, the ¹H NMR spectra suggests that the phosphate binds to the gold ions in the complex as opposed to the cyanostar cavity (Fig. S71†). In addition, the bisulfate (HSO₄⁻)^{57,60} and n-hexylphosphonate $(C_6H_{13}PO_3H^-)^{29}$ anions that can drive cyanostar dimerization, behaved poorly and stripped copper ions off the complex (Fig. S71-S75†). An instantaneous loss of color was indicative of decomplexation.

Hence, testing the orthogonality of the ion interactions involved in the noncovalent bond forming reactions and verifying the compatibility of the ions with each other is a key step for successful implementation of ion-by-ion stepwise self-

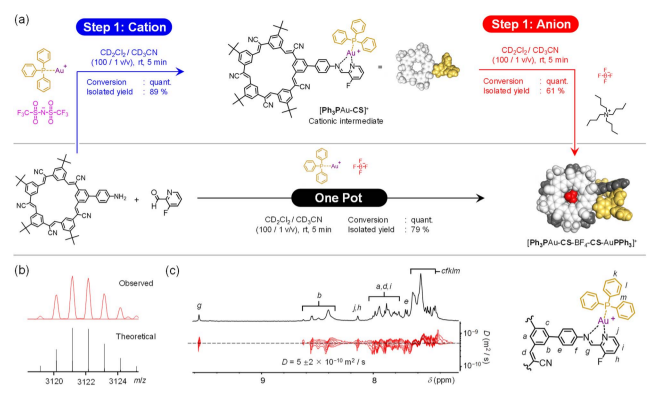


Fig. 12 (a) Reactions to form $[PPh_3Au-CS-BF_4-CS-AuPPh_3]^+$ by one-pot and cation—anion pathways. (b) ESI-MS (CH₂Cl₂, 0.5 mM) of $[PPh_3Au-CS-BF_4-CS-AuPPh_3]^+$ prepared *via* cation—anion pathway. (c) DOSY NMR (CD₂Cl₂, 2 mM) of $[PPh_3Au-CS-BF_4-CS-AuPPh_3]^+$ created in one-pot assembly.

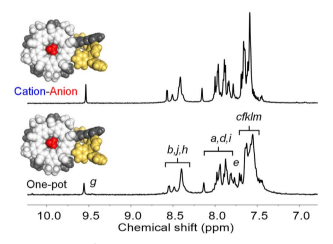


Fig. 13 Diagnostic 1 H NMR peaks (CD₂Cl₂, 0.5 mM) for [PPh₃Au-CS-BF₄-CS-AuPPh₃]⁺ obtained by one-pot assembly and cation–anion assembly.

assembly. Use of cyanostar as a supramolecular element⁴¹ with its strong affinity for anions known to weakly coordinate with metals may be key to this demonstration suggesting other receptors, like bambusuril,^{63–66} may also be used in this way.

Conclusion

We demonstrated that orthogonal cation-anion and anioncation self-assembly can be used to build up the structures of

multi-component assemblies in a modular manner by using pre-programmed binding sites for both cations and anions. This orthogonal ion-by-ion strategy of subcomponent selfassembly enables similar structures to be accessed in a modular manner using common intermediates in both a onepot and stepwise way. The architectures involved Cu(1) and Au(1) complexes situated on the termini of the anion-driven cyanostar dimers form [POPCu-CS-X-CS-CuPOP]⁺ and [PPh₃Au-CS-X-CS-Au**PPh**₃]⁺, where the bridging anion (X) can be either BF₄ or ClO₄⁻. We find that the noncovalent chemistries of anion and cation coordination can be readily paired with the covalent chemistry of sub-component self-assembly based on modular imine-based ligands. The combination of anion receptors with established ligands for metal complexation introduce a new approach to design architectures based on orthogonal cationanion self-assembly.

Data availability

The ESI is available free of charge on the journal website. General methods, NMR titrations, 2D NMR spectroscopy, X-ray diffraction analyses, and ESI-MS analyses.

Author contributions

AD conceived the project under the supervision of AHF; in response to the Covid pandemic, REF assisted AHF with overseeing project completion. AD conducted experiments with assistance from LAK, YC, REF, DVC and VC; AD designed, created and characterized the assemblies with initial input from LAK; YC, REF and LAK made and characterized compounds and collected control data; DVC grew a single crystal and VC collected X-ray diffraction data, solved and refined the crystal structure; AD, REF and AHF analyzed the data, wrote and edited the manuscript with input from all co-authors.

Conflicts of interest

The authors declare no competing financial interests.

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