Pathway-Dependent Post-assembly Modification of an Anthracene-Edged M\textsuperscript{Ii}\textsubscript{4}L\textsubscript{6} Tetrahedron

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Supporting Information

**ABSTRACT:** Fe\textsuperscript{II}\textsubscript{4}L\textsubscript{6} tetrahedral cage 1 undergoes post-assembly modification (PAM) via a Diels–Alder cycloaddition of the anthracene panels of the cage with tetracyanoethylene (TCNE). The modified cage 2 possesses an enclosed cavity suitable for encapsulation of the fullerene C\textsubscript{60}, whereas original cage 1 forms a unique covalent adduct through a Diels–Alder cycloaddition of three of its anthracene ligands with C\textsubscript{60}. This adduct undergoes further PAM via reaction of the remaining three ligands with TCNE, enabling the isolation of two distinct products depending on the order of addition of C\textsubscript{60} and TCNE. Modified cage 2 was also able to bind an anionic guest, [Co(C\textsubscript{2}B\textsubscript{9}H\textsubscript{11})\textsubscript{2}]\textsuperscript{2−}, which was not encapsulated by the original cage, demonstrating the potential of PAM for tuning the binding properties of supramolecular hosts.

**P**ost-assembly modification (PAM) of discrete supramolecular complexes has recently attracted increasing attention as a means of introducing new functionality, tuning solubility, trapping species out of equilibrium, and constructing interlocked structures and polymers. PAM reactions should proceed quantitatively under mild conditions that do not disrupt the dynamic linkages that hold supramolecular structures together. Although several reaction types, including olefin metathesis, alkylene–azide cycloaddition, imine reduction, Diels–Alder cycloaddition (both normal and inverse electron-demand), Knoevenagel condensation, acylation, and nucleophile–isocyanate coupling have been employed to modify supramolecular species, solution-based PAM remains less explored than the PAM of more robust metal–organic frameworks. To produce new functionalized supramolecular assemblies for recognition, catalysis, sensing, and drug delivery, it is crucial to expand the range of reaction types that can be employed for PAM.

Anthracene is an attractive moiety for incorporation into supramolecular hosts due to its extended π-surface, which may form favorable aromatic interactions with targeted guests, and its panel-like shape, which facilitates effective cavity enclosure and stronger host–guest interactions. Although anthracene has been used as a structural element in metal–organic cages, it has not yet been employed as a reactive element in discrete supramolecular constructs despite its well-known ability to undergo both thermal and photochemical cycloadditions across the 9 and 10 positions with a variety of partners. It has found recent use as a structurally modifiable element of more robust covalent cyclophane and cycloparaphenylene-based hosts, and also as a reagent employed to tether redox functionality onto maleimide-functionalized hexagonal prisms.

Here we show how a new anthracene-edged Fe\textsuperscript{II}\textsubscript{4}L\textsubscript{6} cage 1 undergoes PAM via Diels–Alder reaction with tetracyanoethylene (TCNE) under mild conditions (Scheme 1), and how the resultant conversion from planar anthracene to bent dihydroanthracene panels significantly influences the host–guest properties of the assembly. Remarkably, three of the six anthracene panels of 1 were also observed to undergo regiospecific Diels–Alder reactions with a single C\textsubscript{60} to form a unique covalently trapped adduct; the remaining three anthracene panels of this adduct then underwent further PAM with TCNE. The Fe\textsuperscript{II}\textsubscript{4}L\textsubscript{6} cage is thus converted to two distinct post-assembly-modified products depending on the order of reaction with C\textsubscript{60} and TCNE (Scheme 2). These topologically complex modifications would be impossible to achieve if the isolated components of the system reacted together prior to assembly. The PAM strategy is also crucial due to functional group incompatibilities between the reaction components, as the free diamine precursor undergoes aza-Michael addition with TCNE.

Anthracene-containing subcomponent A was synthesized in a single step from commercially available starting materials via a Pd-catalyzed Suzuki–Miyaura cross-coupling as described in the Supporting Information (SI). Cage 1 was then prepared via subcomponent self-assembly of diamine A (6 equiv) with 2-formylanilide (12 equiv) and iron(II) bis(trifluoromethane)sulfonimide (Fe(NTf\textsubscript{2})\textsubscript{2}, 4 equiv). Electrospray ionization mass spectrometry (ESI-MS) confirmed the Fe\textsuperscript{II}\textsubscript{4}L\textsubscript{6} stoichiometry of 1 (SI, Figure S9). The 1H and 13C NMR spectra of cage 1 display a single set of ligand resonances in solution, consistent with idealized T point symmetry (SI, Figures S3 and S4).

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Scheme 2. Summary of the Distinct Pathways of PAM Involving Cage 1 To Give [C60C2] or [C603], and the One-Pot Formation of [C60C2].

Despite the difficulty of performing meaningful NMR analysis on this mixture of diastereomers, we investigated the ability of cage 2 to act as a host. Fullerene C60 was identified as a potential guest for both 1 and 2 due to a predicted size and shape match with the large internal cavities of the cages and the potential for aromatic stacking with the walls of the cage. Excess C60 (5 equiv) was thus added to a CD3CN solution of 2, and the suspension was stirred at 298 K for 4 days. ESI-MS confirmed formation of the 1:1 host–guest complex [C60C2] with no peaks from the free host detected in the ESI mass spectrum (SI, Figure S32). In contrast to 2, the 1H NMR spectrum of [C60C2] showed a major product with a single set of ligand resonances, consistent with the restoration of T point symmetry (SI, Figure S25). The addition of C60 thus brought about a re-equilibration among the different diastereomers of 2, likely via a deligation–religation process,27 such that the T-diastereomer with all TCNE residues on the exterior of the cage predominated. We infer this diastereomer to maximize cavity size and thus binding affinity for this large guest. The success of the PAM reaction employed to synthesize 2 was verified by a shift in the 1H resonance for the 9,10-anthracenyl protons from 8.16 ppm in 1 to 5.22 ppm in [C60C2] and the appearance of corresponding new aliphatic 13C NMR resonances at 49.7 and 47.0 ppm. The 13C NMR spectrum also showed an intense peak at 141.8 ppm for C60, despite its insolubility in CD3CN,28 providing further evidence for the encapsulation of C60 by 2.

The structure of [C60C2] was confirmed unambiguously by single-crystal X-ray analysis, showing successful PAM of all six cage ligands with TCNE and retention of overall T-symmetry (Figure 2). The Fe–Fe distances of 18.343(3)–18.362(3) Å (average 18.4 Å) are strengthened by an average of 1.3 Å relative to those in 1. The increase in Fe–Fe separation in addition to the convex nature of the modified ligand results in an expanded cavity volume of 752 Å3.

To explore the scope of guest binding within 2, the cage was also investigated as a host for the large anion [Co(C2B9H11)2]−. Addition of Na[Co(C2B9H11)2] (5 equiv) to 2 in CD3CN resulted in formation of a new species with a similar diffusion coefficient to [C60C2] but with four signals per ligand environment in the 1H NMR spectrum, suggesting a C3-symmetric host framework.29 ESI-MS confirmed that the cage remained intact. We infer that [Co(C2B9H11)2]− is encapsulated by 2 as anion metathesis would not be expected to induce diastereomeric reconfiguration. The bound [Co(C2B9H11)2]− was displaced by C60 at 323 K (SI, Figure S40), indicating that 2 has a higher affinity for the fullerenic guest, which we infer to result from a better size and shape match between the host cavity and the spherical guest.

To probe the effect of PAM on guest binding, we also investigated precursor cage 1 as a host for [Co(C2B9H11)2]− and C60. No evidence of encapsulation was observed by NMR.

Figure 1. Two views of the cationic part of the crystal structure of [C60C2]. Counterions, solvents, and disorder are omitted for clarity.

Figure 2. Two views of the cationic part of the crystal structure of [C60C2].
following addition of Na[Co(C5B9H11)2] (5 equiv) to 1 in CD3CN. Binding of [Co(C5B9H11)2]− within 2 but not 1 thus demonstrates the utility of PAM as a means to tailor the host—guest properties of a metal—organic cage.

The reaction between 1 and excess C60 (5 equiv) in CD3CN yielded a new product with a well-resolved but complex 1H NMR spectrum. All peaks between 4.04 and 9.16 ppm displayed a single diffusion constant, nearly identical to that observed for 1, in the diffusion-ordered 1H NMR (DOSY) spectrum (SI, Figure S22), suggesting that they belonged to a single species of similar size to the original cage. One-dimensional pure shift (PSYCHE)30 and two-dimensional NMR spectroscopy allowed the assignment of the major peaks to four magnetically distinct environments of equal intensity per ligand proton, consistent with the formation of a product of C3 symmetry. ESI-MS results were consistent with a 1:1 adduct [C60·1].

The solid-state structure of [C60·1] was elucidated by single-crystal X-ray analysis. The crystal structure revealed a covalent Diels–Alder adduct between 1 and C60 (Figure 3), where the encapsulated C60 had undergone cycloaddition with three anthracene ligands that share an apical vertex of the cage. As is usual with C60 functionalization, exclusively the double bonds between two six-membered rings had reacted. With respect to the first position of reaction, the second position was on the same equator of the fullerene rather than on the same hemisphere (cis) or on the opposite hemisphere (trans).31 The same regiochemical relationship exists between all three functionalization sites giving an overall orthogonal (e,e,e)-tris-adduct. The newly formed long C–C bonds are in the range 1.57(1)–1.61(1) Å, similar to those reported for a related adduct.32 The three modified ligands each link the apical iron(II) center with one of the three basal iron(II) centers (Fe–Fe distances 17.678(2)–18.086(3) Å) while three unmodified basal ligands bridge pairs of iron(II) centers around the bottom face of the tetrahedron with slightly shorter Fe–Fe distances of 17.156(2)–17.307(3) Å. In contrast to homochiral 1, the handedness of the apical Fe6 center of [C60·1] is opposite to those of the basal iron(II) centers; both ∆∆∆∆ and ∆∆∆∆ enantiomers are present in the unit cell. The assembly thus possesses overall non-crystallographic C1 symmetry with a three-fold symmetry axis passing through the center of the basal face (Figure 3). Despite reaction of the C60 with ligands that define one vertex of the cage, the fullerene is located almost exactly at the center of the Fe3 tetrahedron. The presence of attractive aromatic interactions between the fullerene and the uncontracted anthracenes may be stabilizing C60 in this more central position.

The functionalization of fullerenes often leads to multiple products due to different numbers of functionalization events occurring and different regio- and stereoisomers forming.33 Obtaining selectivity for a single adduct is highly challenging and often requires permanent covalent tethering between reactants.34 The orthogonal (e,e,e)-tris-adduct of C60 is rare,32 and this adduct with anthracene units has been reported to be thermally unstable at room temperature.32 Solutions of [C60·1] were stable at 323 K for several days but decomposition was observed at 343 K. The (e,e,e)-tris-adduct is chiral, and it is notable that the stereochemistry of the adduct is determined by the chirality of the cage framework with complete diastereoselectivity.

The solution NMR data for [C60·1] are consistent with the solid-state structure (Figure 4), with the basal metal centers giving rise to three magnetically distinct environments and the apical metal center giving rise to one environment. Remarkably, NMR integration indicated the reaction of C60 to be >95% selective for the formation of tris-adduct [C60·1], despite the presence of six anthracene panels and the potential for multiple adducts involving up to six cycloadditions and multiple regio- and stereoisomers of some adducts.

Noting that adduct [C60·1] still contained three unmodified anthracene moieties, we sought to carry out further PAM through reaction of the remaining anthracene groups with TCNE. The reaction of [C60·1] with excess TCNE (5 equiv per anthracene) in CD3CN at 298 K was followed by 1H NMR. Peaks corresponding to [C60·1] were observed to disappear with appearance of a new set of peaks having the same overall number of signals, indicating that the C3 symmetry of the assembly was maintained in the new product (Figure 4). The presence of four singlets in the range 5.1–5.7 ppm is consistent with complete PAM of the remaining ligands. ESI-MS confirmed formation of mixed ligand adduct [C60·3], where the three basal ligands in [C60·1] had undergone cycloaddition with TCNE (Figure 5).

We have thus shown that anthracene-edged tetrahedral cage 1 can undergo Diels–Alder cycloadditions both with the electron-deficient dienophile TCNE and with the fullerene C60. Two singular adducts can be obtained, depending on the order of reaction with TCNE and C60 (Scheme 2). Treatment with
Communication

TCNE followed by C$_{60}$ gave rise to host–guest complex [C$_{60}$C$_2$], where C$_{60}$ is non-covalently encapsulated inside a TCNE modified cage with bent dihydroanthracene panels. Reversing the order of reactant addition led to the formation of [C$_{60}$]$_2$, where three of the ligands have undergone cycloaddition with C$_{60}$ and the other three have reacted with TCNE, representing an unusual route to mixed-ligand complexes. Addition of an excess of both TCNE and C$_{60}$ concurrently also afforded [C$_{60}$C$_2$] due to the greater rate of reaction between the anthracene panels and TCNE.

Although the insolubility of C$_{60}$ in acetonitrile precluded us from quantifying the affinity of 1 or 3 for this guest, we take the six C–C bonds formed during binding to imply a very high affinity. Future work will seek to adapt the methodology presented here to obtain functionalized fullerene derivatives that would be difficult to obtain selectively by other means.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/jacs.6b06710. Synthetic details, characterization data, and NMR and mass spectra (PDF)

X-ray data for 1 (CCDC 1486377) (CIF)
X-ray data for [C$_{60}$]$_1$ (CCDC 1486378) (CIF)
X-ray data for [C$_{60}$C$_2$] (CCDC 1486379) (CIF)

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Notes

The authors declare no competing financial interest.

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