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Guest recognition enhanced by lateral interactions†

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A hexacationic triangular covalent organic cage, AzaEx²Cage⁶⁺, has been synthesized by means of a tetrabutylammonium iodide-catalyzed S_N2 reaction. The prismatic cage is composed of two triangular 2,4,6-triphenyl-1,3,5-triazine (TPT) platforms bridged face-to-face by three 4,4'-bipyridinium (BIPY²⁺) spacers. The rigidity of these building blocks leads to a shape-persistent cage cavity with an inter-platform distance of approximately 11.0 Å. This distance allows the cage to accommodate two aromatic guests, each of which is able to undergo $\pi - \pi$ interactions with one of the two TPT platform simultaneously, in an A-D-D-A manner. In the previously reported prism-shaped cage, the spacers (pillars) are often considered passive or non-interactive. In the current system, the three BIPY²⁺ spacers are observed to play an important role in guest recognition. Firstly, the BIPY²⁺ spacers are able to interact with the carbonyl group in a pyrene-1-carbaldehyde (PCA) guest, by introducing lateral dipole-cation or dipole-dipole interactions. As a consequence, the binding affinity of the cage towards the PCA quest is significantly larger than that of pyrene as the quest, even although the latter is often considered to be a better π -electron donor. Secondly, in the case of the quest 1,5-bis[2-(2-(2-hydroxyethoxy)ethoxy)ethoxy]naphthalene (BH4EN), the pillars can provide higher binding forces compared to the TPT platform. Hence, peripheral complexation occurs when AzaEx²Cage⁶⁺ accommodates BH4EN in MeCN. Thirdly, when both PCA and BH4EN are added into a solution of AzaEx2Cage6+, inclusion and peripheral complexation occur simultaneously to PCA and BH4EN respectively, even though the accommodation of the former guest seems to attenuate the external binding of the latter. This discovery of the importance of lateral interactions highlights the relationship between the electrostatic properties of a highly charged host and its complexation behavior, and as such, provides insight into the design of more complex hosts that bind guests in multiple locations and modes.

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Introduction

Cyclophanes, such as calix[n]arenes,¹ oxacalixarenes,² resorcinarenes,³ pillar[n]arene,⁴ calixpyrroles,⁵ as well as cationic ones,⁶ including cyclobis(paraquat-p-phenylene) (CBPQT⁴⁺)⁷ and its extended counterparts,⁸ represent one of the major focuses in the field of supramolecular chemistry. In the frameworks of

these cyclophanes, various arenes are connected by a number of aliphatic linkers such as methylene units, resulting in relatively rigid and preorganized cavities where guest encapsulation could occur without significant entropy loss. For example, in the case of CBPQT⁴⁺, π -electron-deficient 4,4'-bipyridinium (BIPY²⁺) are separated by two *p*-xylene spacers in a face-to-face manner by approximately 7 Å, which is two times longer than a typical π - π interaction distance. These geometrical properties imply that CBPQT⁴⁺ could accommodate a π -conjugated guest within its cavity, where π - π interactions⁹ could occur between the guest and both the two BIPY²⁺ platforms in the host.

These kind of host-guest recognition features of CBPQT⁴⁺ and its homologous cyclophanes can be used to fulfill various tasks, including the construction of molecular switches¹⁰ and machines¹¹ in the form of rotaxanes¹² and catenanes,¹³ water purification *via* the extraction of aromatic compounds into the host,¹⁴ and the stabilization of unusual guest conformations.¹⁵ There are still several limitations, however, in the design principles of these cyclophanes.

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Firstly, the spacers that connect the π -electron-deficient moieties often behave as "passive" or non-interactive building blocks in terms of molecular recognition – except for a few examples where these spacers can take part in relatively weak [C–H··· π] interactions. ¹⁶ Secondly, in most cases, host–guest complexation occurs within the host cavity wherein the interactions between the guests and the two platforms occur simultaneously. Furthermore, these interactions are often strengthened by solvophobic effects in polar solvents. As a consequence, most of these cages or rings only have one binding mode or site. There are very few examples ¹⁷ of guests binding externally to these hosts.

In the current work, we report the design and synthesis of a hexacationic triangular prism, $\mathbf{AzaEx^2Cage^{6^+}}$, which is composed of two **TPT** platforms connected face-to-face by three $\mathbf{BIPY^{2^+}}$ pillars. The distance between the two platforms is approximately 11.0 Å, which is approximately three-times larger than a typical π - π interaction distance, implying that the cavity in this prism could encapsulate two aromatic guest molecules such as pyrene. Interestingly, in contrast to previously reported hosts that contain phenyl or biphenyl spacers, the $\mathbf{BIPY^{2^+}}$ spacers in $\mathbf{AzaEx^2Cage^{6^+}}$ play a more important role in host-guest complexation.

The host exhibits significantly better binding affinity towards pyrene-1-carbaldehyde (**PCA**) than towards pyrene in both organic solvents and water because the former guest contains a partially negatively charged carbonyl group that can undergo dipole–cation or dipole–dipole interactions with the positively charged **BIPY**²⁺ pillars. Furthermore, the **BIPY**²⁺ pillars allow the cage to form an external complex with 1,5-bis[2-(2-(2-hydroxyethoxy)ethoxy)ethoxy]naphthalene

(BH4EN), a guest containing two tetraethylene glycol chains grafted onto a 1,5-dioxynaphthalene (DNP) moiety. This external binding is driven by the donor-acceptor interactions between the DNP unit in the guest and the BIPY²⁺ pillar in the host, as well as hydrogen bonding between the glycol oxygen atoms and the relatively acidic protons in the BIPY²⁺ units. The assumption that the DNP unit prefers to interact with the more electron-deficient BIPY²⁺ pillar instead of the triazine-containing platform inside the cage is supported by both single-crystal X-ray diffraction analysis and theoretical calculations. Furthermore, guest encapsulation and external binding to the cage occur simultaneously in the presence of both PCA and BH4EN, even though binding of the latter slightly decreases upon encapsulation of the former.

We envision that our findings will promote the fundamental understanding, and thus design, of hosts that are able to accommodate guests in multiple modes or sites.

Experimental section

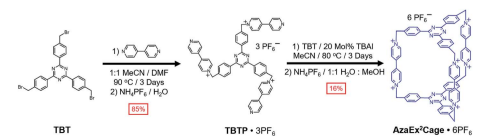
Synthesis and characterization of AzaEx²Cage · 6PF₆

2,4,6-Tris[4-(bromomethyl)phenyl]-1,3,5-triazine a molecule that comprises three benzyl bromide functions grafted onto a TPT moiety - was synthesized in high yield from the acid-catalyzed trimerization of 4-cyanobenzyl bromide.18 TBT was added slowly into a refluxing solution containing a 40fold excess of 4,4'-bipyridine in MeCN and DMF (1:1 v/v) at 90 °C in several aliquots over the course of 6 h, yielding the triscationic compound TBTP·3PF6 after counterion exchange. A 1:1 mixture of TBT and TBTP·3PF6 was then combined and heated at 80 °C in MeCN in the presence of 0.2 equiv. of tetrabutylammonium iodide (TBAI) as a catalyst, leading to the formation (Scheme 1) of a substantial amount of yellow precipitate. The ¹H NMR spectrum of the precipitate, recorded in CD₃SOCD₃, indicates that the cage AzaEx²Cage⁶⁺, the counterions of which could be either Br-, I- or PF₆-, represents the major product in the precipitate, along with a variety of oligomers or polymers as the minor products. Pure AzaEx²Cage · 6PF₆ was obtained in 16% yield by means of silica gel chromatography (1% NH₄PF₆ in MeCN (w/v)), before which counterion exchange was performed. Two more water-soluble counterparts, namely $AzaEx^2Cage \cdot 6Cl$ and $AzaEx^2Cage \cdot 6CF_3CO_2$, were then obtained from AzaEx²Cage · 6PF₆ by counterion exchange.

Results and discussion

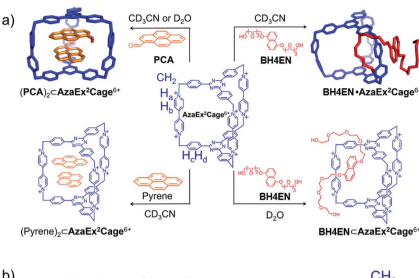
 $AzaEx^2Cage \cdot 6PF_6$ was fully characterized by NMR spectroscopy, high-resolution mass spectrometry (HRMS), and X-ray diffraction analysis. Its 1H NMR spectrum is remarkably simple (Fig. 1b), indicating that the cage has averaged D_{3h} symmetry in solution.

Single crystals of both $AzaEx^2Cage \cdot 6PF_6$ and $AzaEx^2Cage \cdot 6CF_3CO_2$ were obtained by vapor diffusion of ${}^{i}Pr_2O$ into the corresponding MeCN solutions. Crystal twinning was observed, however, in the case of the former crystals (Fig. S40 and S41†). As expected, in the solid-state structure (Fig. 2) of $AzaEx^2Cage \cdot 6CF_3CO_2$, the two TPT platforms in a cage framework are separated by a distance of approximately 11.0 Å, which is three-times larger than a typical π - π interaction distance, 19



Scheme 1 Synthesis of AzaEx²Cage · 6PF₆ from TBT.

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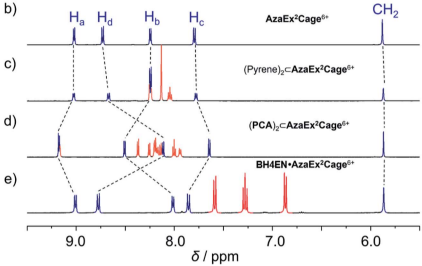


Fig. 1 (a) Schematic representation of the ability of AzaEx 2 Cage $^{6+}$ to recognize a variety of guests, including pyrene, PCA, and BH4EN. In the cases of pyrene and PCA, 2:1 complexes are formed, even though the ability of the cage to host the former guest is much weaker. In the case of BH4EN, the complex BH4EN·AzaEx 2 Cage $^{6+}$ forms in a peripheral and inclusion manner in CD $_3$ CN and D $_2$ O, respectively. 1 H NMR spectra of AzaEx 2 Cage·6PF $_6$ (500 MHz, 1.0 mM in CD $_3$ CN, 298 K) (b) before and after adding (c) pyrene (8.4 equiv.), (d) PCA (10.8 equiv.), and (e) BH4EN (11.4 equiv.). The resonances of the protons in AzaEx 2 Cage $^{6+}$ are labeled in the corresponding spectra. Addition of pyrene and PCA results in upfield shifts of the resonances of the phenylene residues (H $_c$ and H $_d$) in AzaEx 2 Cage $^{6+}$, while the presence of BH4EN leads to upfield shifting of the resonances for the protons on BIPY $^{2+}$ (H $_a$ and H $_b$).

indicating that the cage is able to accommodate two aromatic guests in an A–D–D–A (A = acceptor; D = donor) manner. The distance between two adjacent methylene linkers within each TPT platform is around 11.6 Å. As a result, the cage framework has three relatively large (11.0 Å \times 11.6 Å) rectangular pore windows. These large windows allow potential guests to undergo relatively fast association/dissociation with the cage. The CF $_3$ CO $_2$ counterions are located close to the BIPY $^{2+}$ pillars. Hydrogenbonding interactions occur between the CF $_3$ CO $_2$ counterions and the protons of the cage in the solid state. Furthermore, pairs of adjacent cage frameworks undergo stacking with each other, driven by $\pi-\pi$ interactions between a triazine unit in the platform of one cage and a phenyl moiety of the other.

Cyclic voltammetry (CV) analysis (Fig. S36a†) of $AzaEx^2Cage \cdot 6PF_6$ in degassed DMF reveals two consecutive

reversible redox processes. The reduction processes at -0.264 and -0.662 V can be assigned to two consecutive three-electron reductions, *i.e.*, $\mathbf{BIPY}^{2+}/\mathbf{BIPY}^{+}$ and $\mathbf{BIPY}^{+}/\mathbf{BIPY}^{0}$, by three identical \mathbf{BIPY}^{2+} pillars in a cage. The simultaneous reduction of the three pillars indicates the absence of electron communication between the \mathbf{BIPY}^{2+} units within the cage framework.

The UV-Vis-NIR absorption spectrum (Fig. S36b†) of AzaEx²-Cage³(+•) was obtained by adding Zn dust as a reductant to a solution of AzaEx²-Cage•6PF6 in MeCN. The spectrum is similar to that of 4,4′-dimethylviologen radical cations, further demonstrating that neither radical pairing nor dimerization occurs between any two of the three BIPY* pillars of the cage AzaEx²-Cage³(+•). The absence of intramolecular radical pairing interactions²o is not surprising because the three BIPY* units are separated by approximately 11.6 Å within the rigid cage framework.

a)

Fig. 2 Solid-state (super)structures of AzaEx²Cage·6CF₃CO₂ obtained by single-crystal X-ray crystallography. (a) Stick representation of AzaEx 2 Cage $^{6+}$, CF $_7$ CO $_2^-$ counterions are omitted for the sake of clarity. (b and c) Side-on views of the packing structure revealing that $\pi-\pi$ and hydrogen-bonding interactions play important roles in the crystal packing. The close contacts in the range of 2.22-2.46 Å between $\text{CF}_3\text{CO}_2^{-1}$ counteranions and the protons in the cage indicate the occurrence of Hydrogen-bonding interactions, which are labeled with dashed lines. Hydrogen atoms and disordered solvent molecules are omitted for the sake of clarity. C, gray; N, blue; F, green; O, red.

Host-guest recognition

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We first investigated the ability of AzaEx²Cage⁶⁺ to accommodate two π -electron-rich guests, since the two **TPT** platforms are π -electron acceptors. The ¹H NMR resonances for AzaEx²Cage · 6PF₆ recorded in CD₃CN exhibit only relatively small shifts upon addition of different guests, including pyrene, triphenylene, and perylene. For example, upon addition of 8.37 equiv. of pyrene, the resonances of the H_c and H_d protons in the phenylene residues of AzaEx²Cage·6PF₆ undergo (Fig. 1c) upfield shifts of just -0.02 and -0.06 ppm, respectively. These findings indicate that the ability of AzaEx²Cage⁶⁺ to accommodate pyrene within its cavity is remarkably small. Side-on interactions between the host platforms and the guests in a external manner may also contribute to the upfield shifts of the cage proton resonances in the ¹H NMR spectra, at least partially. For example, after adding 30 equiv. of pyrene into a solution of TBT in CD3CN, the resonances of the corresponding phenylene protons undergo upfield shifts of -0.02and -0.04 ppm, respectively. In fact, the binding constants of $AzaEx^2Cage^{6+}$ to two pyrene molecules in CD₃CN, i.e., K_1 and K_2 , are too low to be determined accurately by means of ¹H NMR titration experiments. A 1:1 binding model was employed to fit the NMR titration data, for which $K_{\rm eq} = 11.1 \pm 0.2 \; {\rm M}^{-1}$ was determined (Fig. S22 and S23†). Several attempts were made to obtain single crystals of the complex (pyrene)₂ \subset AzaEx²Cage⁶⁺ by vapor diffusion of ⁱPr₂O into MeCN solutions containing both pyrene and AzaEx²Cage·6PF₆ at different guest-to-host ratios ranging from 5:1 to 40:1. All of these attempts, however, yielded only a few single crystals corresponding to "empty" AzaEx²Cage · 6PF₆ cages, further supporting our conclusion that the binding affinity of (pyrene)₂ \subset AzaEx²Cage⁶⁺ is at best very weak.

We also envisioned that AzaEx²Cage · 6Cl might be capable of accommodating pyrene in water by taking advantage of the hydrophobic effect. After sonicating suspensions of pyrene in D₂O solutions of AzaEx²Cage·6Cl (1.0 mM) at both room temperature and 80 °C for no less than 5 h, however, no resonances corresponding to pyrene were observed in the ¹H NMR spectra. In addition, the resonances of the cage protons remained almost completely unshifted. These results are in

contrast to our previously reported findings, i.e., that the two smaller cages, namely ExCage⁶⁺ (ref. 14) and AzaExCage⁶⁺,²¹ can encapsulate a variety of polycyclic aromatic hydrocarbons with remarkably high binding constants in both organic solvents and water. For example, the K_a values for pyrene \subseteq **ExCage**⁶⁺ and pyrene \subseteq **AzaExCage**⁶⁺ are 6.77 \times 10⁵ (ref. 14) and 4.93 \times 10⁵ M⁻¹,²¹ respectively, in MeCN. The lower binding affinity in the case of AzaEx2Cage6+ could be explained by the facts that (i) the **TPT** platform of $AzaEx^2Cage^{6+}$ is a weaker π -electron acceptor than the more electron-deficient triscationic counterparts such as 1,3,5-pyridinium-phenyl in ExCage⁶⁺ and 2,4,6-pyridinium-1,3,5-triazine in $AzaExCage^{6+}$, (ii) the formation the 2:1 complex (pyrene)₂ ⊂ AzaEx²Cage⁶⁺ is less entropically favored than formation of the 1:1 complexes pyrene⊂ExCage⁶⁺ and pyrene \subset AzaExCage⁶⁺ (i.e., ExCage⁶⁺ and AzaExCage⁶⁺ have more preorganized cavities for encapsulating pyrene guests) and (iii) the A-D-D-A binding mode in the case of (pyrene)₂ ⊂ AzaEx²Cage⁶⁺ involves a less favored D-D interaction, making its formation more energetically demanding than that of the A-D-A mode that occurs in both pyrene \subseteq ExCage⁶⁺ and pyrene \subseteq AzaExCage⁶⁺.

Pyrene-1-carbaldehyde (PCA) is considered to be a weaker π electron donor than pyrene, since it contains an electronwithdrawing formyl group. Surprisingly, PCA undergoes stronger binding within the cavity of $AzaEx^2Cage \cdot 6PF_6$ compared to that of pyrene. For example, upon addition of 10.8 equiv. of PCA to a CD₃CN solution of $AzaEx^2Cage \cdot 6PF_6$ (Fig. 1d), the resonances of the H_c and H_d protons in the platforms of the cage undergo much more significant upfield shifts, i.e., $\Delta\delta$ for H_c and H_d are -0.154 and -0.618 ppm, respectively, compared with -0.02 and -0.06 ppm, respectively, when 8.37 equiv. of pyrene is added (Fig. 1c). The upfield shifts reveal the presence of π - π interactions between the TPT platform and the pyrene moiety in the guest, providing the phenylene units in the platform with a magnetically shielded environment. Only one set of sharp resonances corresponding to either the host or the guest is observed in the ¹H NMR spectra (Fig. 1d and 3b), regardless of the host-to-guest ratio, indicating that host-guest association/ dissociation occurs relatively rapidly on the ¹H NMR timescale. This observation is consistent with the solid-state

structure of $AzaEx^2Cage^{6+}$ – *i.e.*, each cage contains three large pore windows allowing for rapid guest exchange.

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¹H NMR titration experiments (Fig. 3b and c) revealed that the binding constants for the formation of $(PCA)_2 \subset AzaEx^2Cage^{6+}$ in CD₃CN, *i.e.*, K_1 and K_2 , are 23 \pm 3 and 730 \pm 160 M⁻¹, respectively. The positive cooperativity effect is not surprising, given that the cage cavity with a length of 11 Å is too large and less preorganized to accommodate the first guest, which is able to undergo π - π interactions with only one of the two platforms in the host. After accommodation of the first guest, the cavity becomes smaller in size and therefore more complementary towards binding of the second guest molecule. In addition, the encapsulation of the second guest is driven by an extra guestguest π - π interaction, which does not occur in the case of the first guest. The 2:1 complex $(PCA)_2 \subseteq AzaEx^2Cage^{6+}$ forms exclusively in D₂O upon sonicating PCA solid in a D₂O solution of AzaEx²Cage·6Cl (1.0 mM) at room temperature. Measuring the integration of the resonances for the protons corresponding to both the host and guest in the ¹H NMR spectrum (Fig. 3d) confirms the 1:2 host-to-guest stoichiometry of the complex, which is largely a result of the low solubility of the unbound guest in water. The 2D DOSY spectrum (Fig. 3d) further confirms the formation of the complex $(PCA)_2 \subset AzaEx^2Cage^{6+}$. In addition, in

water, the resonance for the formyl proton in PCA undergoes a remarkable upfield shift, i.e., $\delta = 8.41$ ppm; $\Delta \delta \approx 2$ ppm. This observation indicates that a short contact occurs between the aldehyde and one of the viologen pillars in the host. The latter unit thus provides a magnetically shielded chemical environment for the former group, which supports our assumption that the formyl group can undergo cation-dipole interactions with one of the viologen pillars.

Single crystals of $(PCA)_2 \subset AzaEx^2Cage \cdot 6PF_6$ suitable for X-ray diffraction were obtained by vapor diffusion of ⁱPr₂O into its MeCN solution at 4 °C over 3 days, providing unambiguous evidence for the formation of a 2:1 complex (Fig. 4). As expected, two guest molecules are accommodated within the cavity of the cage framework. π - π Stacking interactions occur between the two platforms in the host and the two guests in an A-D-D-A manner, as inferred from the observation that the distances between triazine/pyrene, pyrene/pyrene, and pyrene/ triazine are 3.42, 3.46, and 3.46 Å, respectively. The closest contacts between the two carbonyl oxygen atoms and the corresponding BIPY²⁺ pillars of the host are 2.78 and 3.01 Å, indicating the occurrence of dipole-cation or dipole-dipole interactions. This secondary interaction is responsible for the enhanced binding affinities of PCA with the cage cavity relative

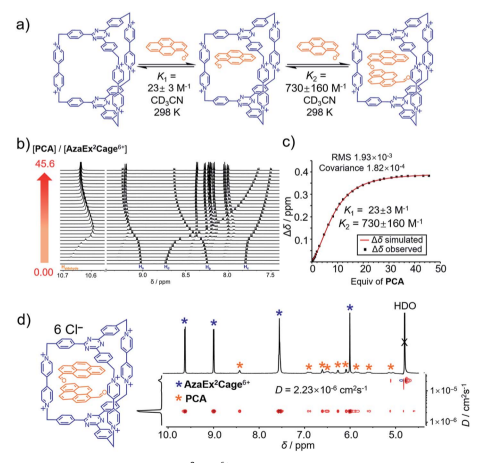


Fig. 3 (a) Schematic representation of the ability of AzaEx²Cage⁶⁺ to accommodate two PCA quests, in which the corresponding binding constants K₁ and K₂ determined by ¹H NMR spectroscopy in CD₃CN at room temperature are shown. (b) Stacked ¹H NMR spectra of AzaEx²Cage 6PF₆ in CD₃CN upon addition of 0 to 45.6 equiv. of PCA relative to AzaEx²Cage⁶⁺. (c) Plot of the downfield resonance shifts of H_b protons in $AzaEx^2Cage^{6+}$ ($\Delta\delta$) versus the amount of PCA guest added relative to $AzaEx^2Cage^{6+}$. [$AzaEx^2Cage \cdot 6PF_6$] = 0.904 mM for all spectra. (d) 2D DOSY spectrum of the $(PCA)_2 \subset AzaEx^2Cage^{6+}$ recorded in D_2O .

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a)
3.42 A
3.46 A
3.46 A

e)

Fig. 4 (a) Side-on stick diagrams view of the crystal structures of $(PCA)_2 \subset AzaEx^2Cage^{6+}$ forms with different chirality including (a) R (blue) and (b) S (red). The axial chirality of the complex results from the different relative rotation directions of the carbonyl groups in the two PCA guests within the cage cavity. The interplane distances between the platforms in the host and the two guests are 3.42, 3.46, and 3.46 Å, indicating the occurrence of $\pi-\pi$ stacking interactions. (c) Top view of the X-ray crystal structure of $(PCA)_2 \subset AzaEx^2Cage^{6+}$. The distances between the two carbonyl oxygen atoms and the corresponding $BIPY^{2+}$ pillars are 2.78 and 3.01 Å, respectively, indicating the occurrence of dipole-cation or dipole-dipole interactions. (d) Side-on stick diagram view overlaid with a space-filling representation of the crystal structure of $(PCA)_2 \subset AzaEx^2Cage^{6+}$. (e) Side view of the crystal lattice of $(PCA)_2 \subset AzaEx^2Cage \cdot 6PF_6$ showing that complexes of different chirality undergo stacking with each other, making the crystal lattice a racemic mixture. PF_6^- counterions are omitted for the sake of clarity.

to those of pyrene. It is noteworthy that, in the solid state, the two carbonyl groups of the two guests interact with two different $BIPY^{2+}$ pillars in the cage in order to minimize the repulsion between the partially anionic carbonyl oxygen atoms. Thus, $(PCA)_2 \subset AzaEx^2Cage \cdot 6PF_6$ exhibits axial chirality in the solid state, even though the two enantiomers co-crystallize in a racemic crystal lattice.

BH4EN, which is another π -electron-rich compound that bears two tetraethylene glycol chains grafted onto a 1,5-dioxynaphthalene (DNP) moiety, was also added into a CD₃CN solution of AzaEx²Cage · 6PF₆ in order to investigate their recognition behavior. Interestingly, instead of forming an inclusion complex in either a 1:1 or 1:2 manner, it seems that the external complex BH4EN·AzaEx²Cage⁶⁺ is formed. This assumption was supported initially by the ¹H NMR spectrum (Fig. 1e). Upon addition of 11.4 equiv. of BH4EN, the resonance of the H_b proton in BIPY²⁺ undergoes a significant upfield shift $(\Delta \delta \approx -0.229 \text{ ppm})$, while those of H_c and H_d in the platforms are barely shifted. This observation indicates that, instead of undergoing π - π interactions with the cage platform, the **DNP** moiety prefers to interact with the BIPY²⁺ pillars in the host, which therefore experiences a magnetically shielded environment. No NOESY cross-peaks between the phenylene units in the host platforms and DNP protons in the guest are observed in the NOESY spectrum (Fig. S27†) of the complex **BH4EN**·**AzaEx**²Cage⁶⁺, indicating that the interactions between **DNP** and $BIPY^{2+}$ occur on the periphery of the cage molecule. The formation of external complex was unambiguously confirmed by single-crystal X-ray diffraction. In the solid state (Fig. 5), the face-to-face distance between **DNP** and **BIPY**²⁺ is

around 3.46 Å, confirming the occurrence of π -electron donoracceptor interactions. The two glycol chains of the guest penetrate into the cavities of two adjacent cage molecules. The shortest contact between an oxygen atom in the glycol chain and one of the H_b protons in BIPY²⁺ is around 2.24 Å, indicating the existence of $[C-H\cdots O]$ interactions. The combination of donor-acceptor and hydrogen-bonding interactions make the formation of the external complex more thermodynamically favorable than formation of the inclusion complex wherein the **DNP** guest undergoes π - π interactions with the triazine residue in the platforms. In water, the recognition mode for BH4EN and AzaEx2Cage6+ is similar, i.e., adding BH4EN to a solution (1.0 mM) of AzaEx²Cage · 6Cl in D₂O also results in an upfield shift of the resonances for the BIPY2+ protons. However, an interesting difference is that strong NOESY cross-peaks between the **DNP** protons $(H_v \text{ and/or } H_z)$ and the phenylene protons (H_c and H_d) in the host platforms are observed (Fig. 6). These observations indicate that, in water, the complex **BH4EN** \subset **AzaEx**²**Cage**⁶⁺ may exist in the form of an inclusion complex instead of the external complex observed in CD₃CN. This observation could be explained by the fact, that in water where the hydrophobic effect becomes predominant, the formation of the inclusion complex is more thermodynamically favored than formation of the external complex. Our attempts to obtain single crystals of the complex **BH4EN⊂AzaEx²Cage**⁶⁺ from water proved unsuccessful, probably because of the relatively high aqueous solubilities of both the host AzaEx² Cage · 6Cl and the guest BH4EN.

When both **PCA** and **BH4EN** are added to a CD₃CN solution of **AzaEx**²Cage · 6PF₆, inclusion and external complexation occur

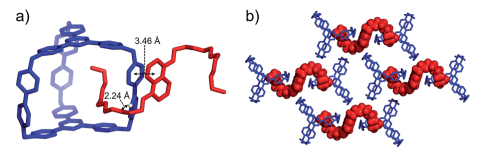


Fig. 5 (a) Side view of the crystal structure of the complex BH4EN·AzaEx 2 Cage·6PF $_6$ in the form of a stick diagram. (b) Top view of the complex BH4EN·AzaEx 2 Cage·6PF $_6$, showing that each BH4EN molecule interacts with two BIPY $^{2+}$ pillars in two adjacent cages. PF $_6$ counterions are omitted for the sake of clarity.

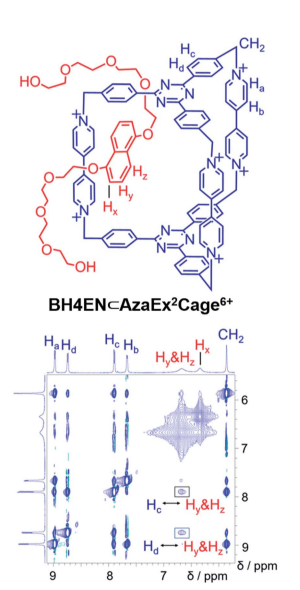


Fig. 6 $^{1}\text{H}-^{1}\text{H}$ NOESY spectrum of a 1 : 6 mixture of AzaEx²Cage·6Cl and BH4EN (500 MHz, D₂O, 298 K). Through-space proton couplings between the H_c and H_d protons in the phenylene groups of the cage and the H_y and/or H_z protons in the guests are labeled in the spectrum, demonstrating that BH4EN·AzaEx²Cage⁶⁺ may exist in the form of an inclusion complex in water.

simultaneously, as inferred from the corresponding 1H NMR spectroscopic analyses (Fig. S30–S33†), in which the proton resonances of both the platforms and pillars undergo shifts upon addition of **PCA** and **BH4EN**, respectively. Diffraction-grade single-crystals of **BH4EN**·(**PCA**) $_2$ \subset **AzaEx** 2 **Cage**·6PF $_6$ were obtained by slow vapor diffusion of i Pr $_2$ O into a three-component 10:4:1 mixture of **PCA**, **BH4EN** and **AzaEx** 2 **Cage**·6PF $_6$ in MeCN (2.0 mM) at 4 $^\circ$ C over a period of 3 days. In the solid state (Fig. 7), the cavity of a cage is occupied by two **PCA** guests, while a **BH4EN** guest resides on the external surface of a **BIPY** $^{2+}$ pillar in the host. The glycol chains of the **BH4EN** guest are excluded from the cage cavity by the two **PCA** guests. This observation implies that hydrogen bonding between the glycol chains and the **BIPY** $^{2+}$ units in the host might be attenuated to some extent in the solid state.

A ¹H NMR titration experiment was performed to evaluate the impact of PCA on the binding affinity of between AzaEx²Cage·6PF₆ and BH4EN. In the absence of PCA, the binding constant (K_a) for the complex BH4EN·AzaEx²Cage⁶⁺ (Fig. S26 and S33†) is approximately around 36.7 \pm 0.4 M⁻¹ in CD_3CN , while upon adding 26.6 equiv. of **PCA**, the K_a value decreases to 4.9 \pm 0.1 M⁻¹. The decrease of the binding constant in solution is consistent with the solid-state observation that encapsulation of PCA guests expels the glycol chains of BH4EN from the cage cavity and helps to decrease the strength of hydrogen bonding. Interestingly, in the presence of 7.8 equiv. of BH4EN, the binding constants between the cage and two PCA guests (Fig. S31†) seem to be barely changed as compared with those in the absence of BH4EN. We interprete this observation to mean that the impact of external binding on the environment inside the cage is of less importance.

Quantum mechanical calculations

In order to gain a better understanding of the recognition characteristics of $AzaEx^2Cage^{6+}$ in terms of both inclusion and peripheral complexation, we have performed density functional theory (DFT) calculations to investigate the electronic properties $AzaEx^2Cage^{6+}$. Geometry optimization of $AzaEx^2Cage^{6+}$, $BH4E-N \subset AzaEx^2Cage^{6+}$, (pyrene)₂ $\subset AzaEx^2Cage^{6+}$, and $(PCA)_2 \subset AzaEx^2Cage^{6+}$ were performed at the M06-2X level of theory²² with the 6-31G(d) basis set.²³ The single-point energies and solvent effects in MeCN were computed at the M06-2X level of theory with the 6-311++G(d,p) basis set for all the atoms, based on the

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a)
3.53 A
3.58 A
3.50 A

Fig. 7 (a) Side view of the crystal structure of the complex BH4EN·(PCA)₂ \subset AzaEx²Cage·6PF₆ in the form of a stick diagram. (b) Top view of the complex BH4EN·(PCA)₂ \subset AzaEx²Cage·6PF₆. PF₆ counterions are omitted for the sake of clarity.

optimized gas-phase structures. Solvation energies were evaluated by a self-consistent reaction field (SCRF) using the SMD model. Natural population analysis of $AzaEx^2Cage^{6+}$ demonstrates that the bipyridinium, methylene and TPT units take +1.301, +0.327 and +0.074 (see the ESI†), respectively, implying that the $BIPY^{2+}$ pillars of the cage, including both pyridinium moieties and the methylene linkers, takes up most of the positive charge on the cage. Expressed another way, the $BIPY^{2+}$ pillars are more electron-deficient than the TPT platforms (Fig. 8a). On one hand, since the DNP unit in a BH4EN guest represents a π -electron-rich donor, it is not surprising that DNP prefers to

interact with BIPY²⁺, forming a BH4EN·AzaEx²Cage⁶⁺ complex (Fig. 8b) whose formation is also strengthened by hydrogen-bonding interactions. On the other hand, the guests pyrene and PCA have larger π -electron surfaces. As a consequence, they prefer to interact with the TPT platform driven by π - π stacking interactions as well as solvophobic effects. To visualize the cation-dipole interactions, we performed reduced density gradient (RDG) analysis²⁶ on the (PCA)₂ \subseteq AzaEx²Cage⁶⁺. The analysis shows clearly the occurrence of noncovalent bonding interactions (Fig. S50†) between the formyl group on PCA and the viologen pillars in AzaEx²Cage⁶⁺.

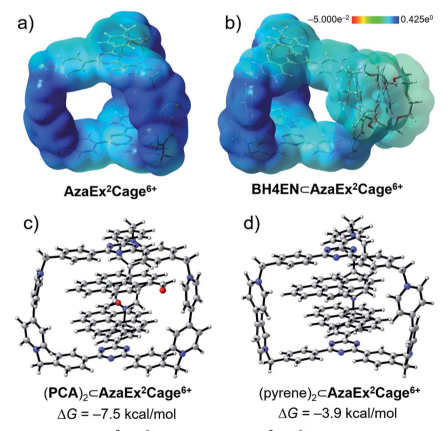


Fig. 8 Electrostatic potential maps for (a) $AzaEx^2Cage^{6+}$ and (b) $BH4EN \subseteq AzaEx^2Cage^{6+}$ obtained by using DFT calculations. Light-yellow and deep-blue colors in the maps represent negative and positive electrostatic potentials, respectively, demonstrating that the $BIPY^{2+}$ units represent better π -electron acceptors on account of their stronger π -electron deficiency. DFT-optimized structures of (c) $(PCA)_2 \subseteq AzaEx^2Cage^{6+}$ and (d) $(pyrene)_2 \subseteq AzaEx^2Cage^{6+}$. The free energy changes (ΔG) for the formation of $(PCA)_2 \subseteq AzaEx^2Cage^{6+}$ and $(pyrene)_2 \subseteq AzaEx^2Cage^{6+}$ complexes are calculated to be -7.5 and -3.9 kcal mol^{-1} , respectively.

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DFT calculations were also performed to evaluate the free energy change (ΔG), which is defined as the free energy difference between the complex and the corresponding unbound guest and host (Fig. 8c and d), for the formation of (pyrene)₂ \subset AzaEx²Cage⁶⁺ and $(PCA)_2 \subset AzaEx^2Cage^{6+}$. The ΔG values for the formation of (pyrene)₂ ⊂ AzaEx²Cage⁶⁺ and (PCA)₂ ⊂ AzaEx²Cage⁶⁺ were calculated to be -3.9 and -7.5 kcal mol⁻¹, respectively. The more negative ΔG for $(PCA)_2 \subset AzaEx^2Cage^{6+}$ indicates that its formation is more thermodynamically favored than that of $(pyrene)_2 \subset AzaEx^2Cage^{6+}$. These results are consistent with the observations that the formation of the former complex is enhanced by dipole-cation or dipole-dipole interactions between the guest carbonyl oxygen atoms, which bear partially negative charges, and the dicationic BIPY²⁺ units in the host.

Conclusion

In summary, we have introduced a hexacationic triangular prismatic cage that is composed of two TPT platforms connected by three BIPY2+ pillar-shaped spacers. Both the platforms and the pillars play important roles in host-guest recognition. The cage can form 2:1 inclusion complexes with polycyclic aromatic hydrocarbons, including pyrene and PCA. The latter guest exhibits remarkably enhanced binding affinities within the cage cavity compared with those of the former, owing to dipole-cation and dipole-dipole interactions between the carbonyl groups and the laterally pillar-shaped spacers in the host. In addition, because the three BIPY²⁺ pillars represent both an efficient π -electron acceptor and hydrogen bonding donor, one of them can interact with a BH4EN guest, which contains a π -electron-donating **DNP** unit and two hydrogenbond-accepting glycol chains. External complexation occurs in organic solvents, while inclusion complexation can occur in water. When both PCA and BH4EN are present, both inclusion and peripheral complexation occur simultaneously, even though accommodation of the former guests seems to suppress the external binding of the latter.

These findings improve our fundamental understanding of the relationship between the electrostatic properties of the building blocks of supramolecular systems and their host-guest recognition properties. Specifically, laterally charged moieties within host molecules could supply effective intermolecular interaction to drive the inclusion and peripheral complexations. Furthermore, this work will inform the development of a design principle for more complex cage molecules that can bind guests in multiple modes and sites.

Conflicts of interest

There are no conflicts of interest to declare.

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References

- 1 A. Ikeda and S. Shinkai, Chem. Rev., 1997, 97, 1713-1734.
- 2 W. Maes and W. Dehaen, Chem. Soc. Rev., 2008, 37, 2393-2402.
- 3 P. Timmerman, W. Verboom and D. N. Reinhoudt, Tetrahedron, 1996, 52, 2663-2704.
- 4 T. Ogoshi, S. Kanai, S. Fujinami, T. A. Yamagishi and Y. Nakamoto, J. Am. Chem. Soc., 2008, 130, 5022-5023.
- 5 P. A. Gale, P. Anzenbacher and J. L. Sessler, Coord. Chem. Rev., 2001, 222, 57-102.
- 6 (a) F. Diederich, Angew. Chem., Int. Ed. Engl., 1988, 27, 362-386; (b) M. Buhner, W. Geuder, W. K. Gries, S. Hunig, M. Koch and T. Poll, Angew. Chem., Int. Ed. Engl., 1988, 27, 1553-1556.
- 7 (a) B. Odell, M. V. Reddington, A. M. Z. Slawin, N. Spencer, J. F. Stoddart and D. J. Williams, Angew. Chem., Int. Ed. Engl., 1988, 27, 1547-1550; (b) P. R. Ashton, B. Odell, M. V. Reddington, A. M. Z. Slawin, J. F. Stoddart and D. J. Williams, Angew. Chem., Int. Ed. Engl., 1988, 27, 1550-
- 8 (a) M. Asakawa, P. R. Ashton, S. Menzer, F. M. Raymo, J. F. Stoddart, A. J. P. White and D. J. Williams, Chem.-Eur. J., 1996, 2, 877-893; (b) P. R. Ashton, S. E. Boyd, A. Brindle, S. J. Langford, S. Menzer, L. Perez-Garcia, J. A. Preece, F. M. Raymo, N. Spencer, J. F. Stoddart, A. J. P. White and D. J. Williams, New J. Chem., 1999, 23, 587-602; (c) H. Y. Gong, B. M. Rambo, E. Karnas, V. M. Lynch and J. L. Sessler, Nat. Chem., 2010, 2, 406-409; (d) E. J. Dale, N. A. Vermeulen, M. Juríček, J. C. Barnes, R. M. Young, M. R. Wasielewski and J. F. Stoddart, Acc. Chem. Res., 2016, 49, 262-273; (e) E. J. Dale, D. P. Ferris, N. A. Vermeulen, J. J. Henkelis, I. Popovs, M. Juríček, J. C. Barnes, S. T. Schneebeli and J. F. Stoddart, J. Am. Chem. Soc., 2016, 138, 3667–3670; (f) X. Gong, R. M. Young, K. J. Hartlieb, C. Miller, Y. Wu, H. Xiao, P. Li, N. Hafezi, J. Zhou, L. Ma, T. Cheng, W. A. Goddard III, O. K. Farha, J. T. Hupp, M. R. Wasielewski and J. F. Stoddart, J. Am. Chem. Soc., 2017, 139, 4107-4116; (g) I. Roy, S. Bobbala, J. Zhou, M. T. Nguyen, S. K. M. Nalluri, Y. Wu, D. P. Ferris, E. A. Scott, M. R. Wasielewski and J. F. Stoddart, J. Am. Chem. Soc., 2018, 140, 7206-7212; (h) Y. Shi, K. Cai, H. Xiao, Z. C. Liu, J. W. Zhou, D. K. Shen, Y. Y. Qiu, Q. H. Guo, C. Stern, M. R. Wasielewski, F. Diederich, W. A. Goddard III and J. F. Stoddart, J. Am. Chem. Soc., 2018, 140, 13835-13842.
- 9 (a) C. A. Hunter and J. K. M. Sanders, J. Am. Chem. Soc., 1990, 112, 5525-5534; (b) L. M. Salonen, M. Ellermann and F. Diederich, Angew. Chem., Int. Ed., 2011, 50, 4808-4842.

Edge Article

10 (a) Molecular Switches, ed. B. L. Feringa, Wiley-VCH, Weinheim, 2001; (b) M. A. Olson, Y. Y. Botros and J. F. Stoddart, Pure Appl. Chem., 2010, 82, 1569–1574; (c) R. Klajn, J. F. Stoddart and B. A. Grzybowski, Chem. Soc. Rev., 2010, 39, 2203–2237; (d) Y. R. Hua and A. H. Flood, J. Am. Chem. Soc., 2010, 132, 12838–12840; (e) A. I. Share, K. Parimal and A. H. Flood, J. Am. Chem. Soc., 2010, 132, 1665–1675.

- 11 (a) B. L. Feringa, R. A. van Delden, N. Koumura and E. M. Geertsema, Chem. Rev., 2000, 100, 1789-1816; (b) R. Ballardini, V. Balzani, A. Credi, M. T. Gandolfi and M. Venturi, Acc. Chem. Res., 2001, 34, 445-455; (c) A. Harada, Acc. Chem. Res., 2001, 34, 456-464; (d) C. A. Schalley, K. Beizai and F. Vögtle, Acc. Chem. Res., 2001, 34, 465-476; (e) J. P. Collin, C. Dietrich-Buchecker, P. Gavina, M. C. Jimenez-Molero and J.-P. Sauvage, Acc. Chem. Res., 2001, 34, 477-487; (f) V. Balzani, A. Credi and M. Venturi, Molecular Devices and Machines: A Journey into the Nanoworld, 2nd edn, Wiley-VCH, Weinheim, 2006; (g) E. R. Kay, D. A. Leigh and F. Zerbetto, Angew. Chem., Int. Ed., 2007, 46, 72-191; (h) J. F. Stoddart, Nat. Chem., 2009, 1, 14–15; (i) M. M. Boyle, R. A. Smaldone, A. C. Whalley, M. W. Ambrogio, Y. Y. Botros and J. F. Stoddart, Chem. Sci., 2011, 2, 204-210; (j) C. Y. Cheng, P. R. McGonigal, S. T. Schneebeli, H. Li, N. A. Vermeulen, C. F. Ke and J. F. Stoddart, Nat. Nanotechnol., 2015, 10, 547-553.
- 12 (a) D. H. Qu, Q. C. Wang and H. Tian, Angew. Chem., Int. Ed., 2005, 44, 5296-5299; (b) H. Zheng, W. Zhou, J. Lv, X. Yin, Y. Li, H. Liu and Y. Li, Chem.-Eur. J., 2009, 15, 13253-13262; (c) Q. Jiang, H. Y. Zhang, M. Han, Z. J. Ding and Y. Liu, Org. Lett., 2010, 12, 1728-1731; (d) C. Romuald, E. Busseron and F. Coutrot, J. Org. Chem., 2010, 75, 6516-6531.
- 13 (a) A. Livoreil, C. O. Dietrich-Buchecker and J.-P. Sauvage, J. Am. Chem. Soc., 1994, 116, 9399-9400; (b) M. Asakawa, R. Ashton, V. Balzani, A. Credi, C. Hamers, G. Mattersteig, M. Montalti, A. N. Shipway, N. Spencer, J. F. Stoddart, M. S. Tolley, M. Venturi, A. J. P. White and D. J. Williams, Angew. Chem., Int. Ed., 1998, 37, 333-337; (c) D. Cao, M. Amelia, L. M. Klivansky, G. Koshkakaryan, S. I. Khan, M. Semeraro, S. Silvi, M. Venturi, A. Credi and Y. Liu, J. Am. Chem. Soc., 2010, 132, 1110-1122; (d) J. C. Barnes, A. C. Fahrenbach, D. Cao, S. M. Dyar, M. Frasconi, M. A. Giesener, D. Benítez, E. Tkatchouk, O. Chernyashevskyy, W. H. Shin, H. Li, S. Sampath, C. L. Stern, A. A. Sarjeant, K. J. Hartlieb, Z. C. Liu, R. Carmieli, Y. Y. Botros, J. W. Choi, A. M. Z. Slawin, J. B. Ketterson, M. R. Wasielewski, W. A. Goddard III and J. F. Stoddart, Science, 2013, 339, 429-433.
- 14 E. J. Dale, N. A. Vermeulen, A. A. Thomas, J. C. Barnes, M. Juríček, A. K. Blackburn, N. L. Strutt, A. A. Sarjeant, C. L. Stern, S. E. Denmark and J. F. Stoddart, *J. Am. Chem. Soc.*, 2014, 136, 10669–10682.

- 15 (a) M. Juríček, N. L. Strutt, J. C. Barnes, A. M. Butterfield, E. J. Dale, K. K. Baldridge, F. Stoddart and J. S. Siegel, *Nat. Chem.*, 2014, 6, 222–228; (b) B. M. Schmidt, T. Osuga, T. Sawada, M. Hoshino and M. Fujita, *Angew. Chem., Int. Ed.*, 2016, 55, 1561–1564.
- 16 (a) P. R. Ashton, C. L. Brown, E. J. T. Chrystal, T. T. Goodnow,
 A. E. Kaifer, K. P. Parry, D. Philp, A. M. Z. Slawin, N. Spencer,
 J. F. Stoddart and D. J. Williams, J. Chem. Soc., Chem. Commun., 1991, 634–639; (b) M. V. Reddington,
 A. M. Z. Slawin, N. Spencer, J. F. Stoddart, C. Vicent and
 D. J. Williams, J. Chem. Soc., Chem. Commun., 1991, 630–634.
- 17 (a) B. E. Tiedemann and K. N. Raymond, Angew. Chem., Int. Ed., 2005, 45, 83–86; (b) C. Sgarlata, J. S. Mugridge, M. D. Pluth, B. E. Tiedemann, V. Zito, G. Arena and K. N. Raymond, J. Am. Chem. Soc., 2010, 132, 1005–1009; (c) T. Sawada and M. Fujita, J. Am. Chem. Soc., 2010, 132, 7194–7201; (d) W. J. Ramsay and J. R. Nitschke, J. Am. Chem. Soc., 2014, 136, 7038–7043; (e) F. J. Rizzuto, W. Y. Wu, T. K. Ronson and J. R. Nitschke, Angew. Chem., Int. Ed., 2016, 55, 7958–7962; (f) D. Preston, J. E. Lewis and J. D. Crowley, J. Am. Chem. Soc., 2017, 139, 2379–2386; (g) X. Bai, C. Jia, Y. Zhao, D. Yang, S. C. Wang, A. Li, Y. T. Chan, Y. Y. Wang, X. J. Yang and B. Wu, Angew. Chem., Int. Ed., 2018, 57, 1851–1855.
- 18 J. Samanta and R. Natarajan, Org. Lett., 2016, 18, 3394-3397.
- (a) M. Yoshizawa, J. Nakagawa, K. Kumazawa, M. Nagao, M. Kawano, T. Ozeki and M. Fujita, *Angew. Chem., Int. Ed.*, 2005, 44, 1810–1813; (b) M. M. Zhang, M. L. Saha, M. Wang, Z. X. Zhou, B. Song, C. J. Lu, X. Z. Yan, X. P. Li, F. H. Huang, S. C. Yin and P. J. Stang, *J. Am. Chem. Soc.*, 2017, 139, 5067–5074.
- 20 A. Trabolsi, N. Khashab, A. C. Fahrenbach, D. C. Friedman, M. T. Colvin, K. K. Coti, D. Benítez, E. Tkatchouk, J. C. Olsen, M. E. Belowich, R. Carmielli, H. A. Khatib, W. A. Goddard III, M. R. Wasielewski and J. F. Stoddart, *Nat. Chem.*, 2010, 2, 42–49.
- 21 N. Hafezi, J. M. Holcroft, K. J. Hartlieb, E. J. Dale, N. A. Vermeulen, C. L. Stern, A. A. Sarjeant and J. F. Stoddart, *Angew. Chem.*, *Int. Ed.*, 2015, 54, 456–461.
- 22 (a) Y. Zhao and D. G. Truhlar, Acc. Chem. Res., 2008, 41, 157–167; (b) Y. Zhao and D. G. Truhlar, Theor. Chem. Acc., 2008, 120, 215–241.
- 23 W. J. Hehre, L. Radom, P. v. R. Schleyer and J. A. Pople, Ab Initio Molecular Orbital Theory, Wiley, New York, 1986.
- 24 A. V. Marenich, C. J. Cramer and D. G. Truhlar, *J. Phys. Chem. B*, 2009, **113**, 6378–6396.
- 25 A. E. Reed, R. B. Weinstock and F. Weinhold, *J. Chem. Phys.*, 1985, **83**, 735–746.
- 26 E. R. Johnson, S. Keinan, P. Mori-Sanchez, J. Contreras-Garcia, A. J. Cohen and W. T. Yang, *J. Am. Chem. Soc.*, 2010, 132, 6498–6506.