

Article

Synthesis of an Adaptable Molecular Barrel and Guest Mediated Stabilization of Its Metastable Higher Homologue

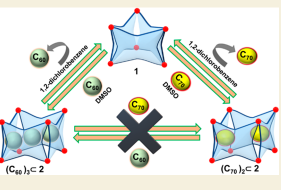
Ranit Banerjee,[‡] Soumalya Bhattacharyya,[‡] and Partha Sarathi Mukherjee*

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ACCESS Metrics & More Article Recommendations Supporting Information ABSTRACT: Structural and functional modulation of three-dimensional artificial macromolecular systems is of immense importance. Designing supramolecular cages that can show stimuli mediated reversible switching Image: Construction of three-dimensional artificial macromolecular systems is of immense importance. Designing supramolecular cages that can show stimuli mediated reversible switching

supramolecular cages that can show stimuli mediated reversible switching between higher-order structures is quite challenging. We report here construction of a Pd₆ trifacial barrel (1) by coordination self-assembly. Surprisingly, barrel 1 was found to exhibit guest-responsive behavior. In presence of fullerenes C_{60} and C_{70} , 1 unprecedentedly transformed to its metastable higher homologue Pd₈ tetrafacial barrel (2), forming stable host-guest complexes $(C_{60})_3 \subset 2$ and $(C_{70})_2 \subset 2$, respectively. Again, encapsulated fullerenes could be extracted from the cavity of 2 using 1,2-dichlorobenzene, leading to its facile conversion to the parent trifacial barrel 1. Such reversible structural interconversion between an adaptable molecular barrel and its guest stabilized higher homologue is an uncommon observation.



KEYWORDS: Self-assembly, Molecular barrels, Fullerene encapsulation, Structural transformation, Coordination chemistry

INTRODUCTION

The flexible and adaptable nature of macromolecular biological receptors is responsible for their structural transformation upon substrate binding. Such flexible and adaptable behavior is mainly due to the noncovalent interactions that drive the biological processes. On the other hand, covalent architectures are generally less prone to structural reorganization to expand their cavity size significantly for better fitting of the substrates. Coordination-driven self-assembly is an efficient technique for designing adaptable large molecular architectures.¹⁻⁹ The dynamic nature of the metal-ligand bonding allows for selfcorrection and conformational changes of the structures.^{10–15} The thermodynamically most stable architecture is formed from a given set of building blocks, whereas the other possible geometries remain inaccessible. However, external stimuli^{16–30} may induce structural changes to access metastable architectures that are not usually formed. Over the years, the internal cavities of such discrete architectures have been used for binding guest molecules for numerous applications such as catalysis, $\frac{31-38}{60}$ sensing, $\frac{39-43}{10}$ light-harvesting, $\frac{44-49}{100}$ biological applications,^{50–52} molecular separations,^{53–56} and modification of the dynamics and chemical reactivity of the trapped guests.⁵⁷⁻⁵⁹ Guest binding within the internal pockets is completely dictated by the size complementarity and favorable noncovalent interactions between the host and the guest. Thus, suitable hosts can be predesigned by incorporating specific functionalities within the building blocks for stabilizing a wide variety of guest molecules within their cavities. Among the

various shapes, architectures resembling barrels are expected to be efficient for encapsulating guest molecules due to their large open windows and parallel orientation of the walls which ensures greater interactions with the incoming guest molecules.^{60,61}

Fullerenes, due to their interesting physicochemical properties, find applications in several fields.⁶²⁻⁶⁷ The major drawback of working with fullerenes is their poor solubility in common organic solvents. Functionalization of fullerenes to generate soluble derivatives has been the common technique employed to overcome this issue. However, such functionalization usually needs multistep reactions and tedious separation steps.⁶⁸ On the other hand, encapsulation of fullerenes within a soluble host is believed to be an easier solution to enhance the solubility of free fullerenes without their chemical modification. Designing suitable receptors that can bind fullerenes in solvents where fullerenes are insoluble has been the prime focus of supramolecular chemists.⁶⁹⁻⁷⁴ Synthesis of hosts that can encapsulate many molecules of fullerene is highly desirable to enhance their solubility. However, it is a challenging task to

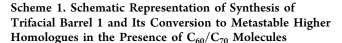
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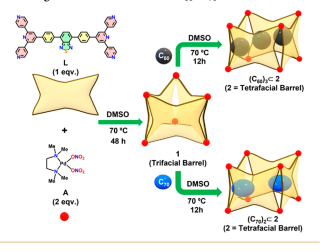




design such receptors with large, confined pockets for accommodating more than one fullerene molecule.

Herein, we report the formation of a $[Pd_6L_3]^{12+}$ trifacial molecular barrel (1) upon coordination self-assembly (Scheme 1) of a tetratopic ligand L with *cis*-[(tmeda)Pd(ONO₂)₂] (A)





as an acceptor in DMSO (tmeda = N, N, N', N'-tetramethylethane-1,2-diamine; DMSO = dimethyl sulfoxide). Trifacial barrel (1) possesses a large internal cavity with open windows suitable for encapsulation of large guest molecules. Interestingly, upon treatment with fullerenes C_{60}/C_{70} , the trifacial barrel undergoes an uncommon structural transformation to its higher homologue $[Pd_8L_4]^{16+}$ tetrafacial barrel (2) with the formation of fullerene encapsulated stable adducts $(C_{60})_3 \subset 2$ and $(C_{70})_2 \subset 2$, respectively. The labile and dynamic nature of the Pd-pyridine bond is ultimately responsible for the transformation of the trifacial barrel to its higher homologue in the presence of external guests to reach its thermodynamically most stable state.^{70,75-79} The addition of fullerene to barrel 1 induced reorganization of the building blocks to generate a metastable tetrafacial barrel (2) by effective hostguest interactions. Extraction of C_{60}/C_{70} from the host-guest complexes led to facile conversion of 2 to its parent trifacial barrel 1 which indicates that the metastable barrel 2 could only be obtained in the presence of C_{60} or C_{70} as the guest. Such guest-induced reversible structural transformation between homologues of self-assembled molecular barrels and stabilization of a metastable homologue by multiple fullerene molecules are unusual observations. Further, these adducts $[(C_{60})_3 \subset 2 \text{ and } (C_{70})_2 \subset 2]$ were found to act as photosensitizers for singlet oxygen generation under visible light irradiation in acetonitrile where free C_{60}/C_{70} is insoluble and shows no singlet oxygen generation.

RESULTS AND DISCUSSION

The ligand L was synthesized by reacting 4,4'-(benzo[c]-[1,2,5]thiadiazole-4,7-diyl)dibenzaldehyde (P) with KOH, 4acetylpyridine, and NH₄OH in ethanol using a standard procedure for the synthesis of terpyridines.^{70,80,81} Precursor P was synthesized via Suzuki coupling between 4,7dibromobenzo[c][1,2,5]thiadiazole and (4-formylphenyl)boronic acid (Scheme S1) following a reported procedure.⁸² Self-assembly of L and *cis*-[(tmeda)Pd(NO₃)₂] (A) was done in a 1:2 molar ratio in DMSO for 48 h at 70 °C which afforded a clear yellow solution (Scheme 1). 1 was isolated as a yellow solid in almost quantitative yield upon addition of excess ethyl acetate to the reaction mixture. ¹H NMR of 1 was recorded in DMSO- d_6 which showed downfield shifts of α -pyridyl protons and a 2-fold splitting of α -pyridyl (H_a and H_a) and β -pyridyl (H_b and H_b) protons of the ligand in comparison to the free L due to rigidification of asymmetric ligand L upon assembly formation (Figures 1a,b, S1, and S4). All the peaks in the ¹H

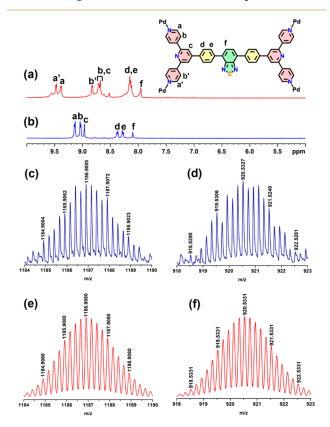


Figure 1. (a) Stacked ¹H NMR spectra of 1 and (b) L in DMSO- d_6 . (c) Experimental (blue) and (e) calculated (red) isotopic patterns of $[1-4PF_6]^{4+}$ fragment. (d) Experimental (blue) and (f) calculated (red) isotopic patterns of $[1-5PF_6]^{5+}$ fragment in ESI-MS analyses.

NMR spectrum were assigned with the help of ${}^{1}\text{H}-{}^{1}\text{H}$ COSY (Figure S6) and NOESY NMR (Figure S7). The same diffusion coefficient ($D = 6.025 \times 10^{-11} \text{ m}^2/\text{s}$) for all the peaks in ${}^{1}\text{H}$ -DOSY NMR confirmed the formation of a single assembly (Figure S5). 1 was converted to the PF₆⁻ analogue by treating the nitrate analogue with excess KPF₆ for better ionization. ESI-MS analyses confirmed the formation of a Pd₆L₃ trifacial barrel 1 by the appearance of peaks at m/z = 1630.8743, 1186.9005, 920.5327, and 742.9542 which correspond to the charged fragments $[1-3PF_6]^{3+}$, $[1-4PF_6]^{4+}$, $[1-5PF_6]^{5+}$, and $[1-6PF_6]^{6+}$, respectively (Figures 1c,d and S9). The isotopic patterns corresponding to these fragments were in good agreement with the theoretically calculated patterns.

Several attempts to grow single crystals of 1 were fruitless. To gain structural insights, the geometry of 1 was optimized using the PM6 method. In the energy minimized structure of barrel 1, three tetrapyridyl ligands (L) are horizontally oriented in a trigonal disposition and the six corners are clipped by the Pd(II) acceptor A (Figure 2). There remains a possibility to obtain an isomeric trifacial tube in which all three ligands are

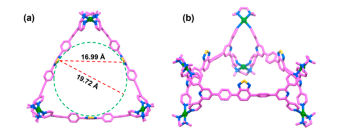


Figure 2. Structure of trifacial barrel 1 obtained by optimization: (a) top view and (b) side view, where ligands are horizontally oriented. Color codes: carbon (pink), nitrogen (blue), sulfur (yellow), and palladium (green). Hydrogen atoms are omitted for clarity.

vertically oriented and clipped to six Pd(II) acceptors. However, the formation of a trifacial tube (Figures S24) is energetically disfavored by about 132 kcal/mol compared to barrel 1 which can be attributed to the greater steric crowding among the vertical panels. From the optimized structure, it is observed that 1 possesses a large internal cavity of diameter 19.72 Å (Figure 2a), π -conjugated aromatic walls, and wideopen windows, suitable for encapsulation and stabilization of large guest molecules.

Fullerenes (C_{60}/C_{70}), on the other hand, are electrondeficient large molecules with negligible solubility in a polar solvent like DMSO. This prompted us to carry out encapsulation of fullerenes C_{60} and C_{70} into barrel 1. We have recently shown structural conversion of a self-assembled trigonal cage to larger analogue upon binding of large guests like fullerenes.⁷⁰ Four equivalents of C_{60} were added to a yellow solution of trifacial barrel 1 in DMSO and stirred at 70 °C for 12 h. The reaction mixture was centrifuged, and a clear brown solution was obtained. The product was isolated as brown solid by treating the solution with excess ethyl acetate. A new set of peaks was seen in the ¹H NMR spectrum with downfield shifts of the α -pyridyl protons of the barrel, indicating the formation of a host–guest complex with C_{60} (Figures 3b and S10).

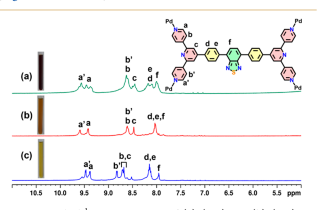
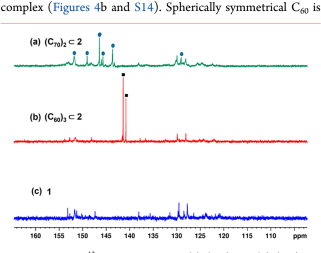


Figure 3. Stacked ¹H NMR spectra of (a) $(C_{70})_2 \subset 2$, (b) $(C_{60})_3 \subset 2$, and (c) trifacial barrel 1 in DMSO- d_6 . Insets show the color of the solutions.

An encapsulation experiment with the ellipsoidal C_{70} was performed in a similar fashion. A deep-red solution was obtained, which upon treatment with excess ethyl acetate yielded a dark-red solid product. The ¹H NMR spectrum of the product was broad due to the presence of unsymmetrical and ellipsoidal C_{70} (Figures 3a and S17). ¹³C NMR spectra of these complexes were recorded to detect the presence of



encapsulated C₆₀ or C₇₀ molecules. Free fullerenes in DMSO

do not show any peak in ¹³C NMR spectra due to poor solubility. Interestingly, two intense peaks of slightly different intensities were found at δ 141.39 and 140.80 ppm for the C₆₀

Figure 4. Stacked ¹³C NMR spectra of (a) $(C_{70})_2 \subset 2$, (b) $(C_{60})_3 \subset 2$, and (c) trifacial barrel 1 in DMSO- d_6 .

expected to show a single intense peak in the 13 C NMR spectrum. The presence of two peaks hinted toward the possibility of the presence of multiple C₆₀ molecules encapsulated within the cavity of the barrel. 13 C NMR spectrum of the C₇₀ complex showed several intense peaks characteristic of C₇₀ which also indicated the possible presence of multiple C₇₀ molecules encapsulated within the barrel (Figures 4a and S21).

To get the exact composition of the host-guest complexes, the nitrate analogues were converted to hexafluorophosphates for easy ionization in ESI-MS experimental conditions. Surprisingly, a composition of three C₆₀ molecules encapsulated in a homologous tetrafacial barrel 2 ($[(C_{60})_3 \subset 2]$) was found for the C₆₀ complex. Peaks were obtained at m/z = 2171.4499, 1708.1593, 1399.3139, 1178.6954, and 1013.2381 which correspond to the charged fragments $[(C_{60})_3 \subset 2-4PF_6]^{4+}$, $[(C_{60})_3 \subset 2-5PF_6]^{5+}$, $[(C_{60})_3 \subset 2-6PF_6]^{6+}$, $[(C_{60})_3 \subset 2-7PF_6]^{7+}$, and $[(C_{60})_3 \subset 2-8PF_6]^{8+}$, respectively (Figures 5b,d, \$15, and \$16). The isotopic patterns were in good agreement with theoretically calculated patterns. On the other hand, a composition of two C70 molecules encapsulated in a similar tetrafacial barrel 2 ([$(C_{70})_2 \subset 2$]) was obtained for the C_{70} complex from ESI-MS analyses. Peaks and the isotopic distribution patterns were obtained at m/z = 2051.1936, 1611.9548, 1319.1342, 1109.9847, and 953.1152 which correspond to the charged fragments $[(C_{70})_2 \subset 2-4PF_6]^{4+}$, $[(C_{70})_2 \subset 2\text{-}5PF_6]^{5+}$, $[(C_{70})_2 \subset 2\text{-}6PF_6]^{6+}$, $[(C_{70})_2 \subset 2\text{-}7PF_6]^{7+}$, and $[(C_{70})_2 \subset 2\text{-}8PF_6]^{8+}$, respectively (Figures 5a,e, S22, and S23). Interestingly, neither the free trifacial barrel (1) nor the tetrafacial barrel (2) was detected from mass analyses. Therefore, trifacial barrel 1 undergoes a complete transformation to a homologous tetrafacial barrel 2 with encapsulated C_{60} or C_{70} . While the free tetrafacial barrel was not formed during the reaction, it was isolated in the form of stable host-guest complexes with three C₆₀ or two C₇₀ molecules.

¹H-DOSY NMR confirmed the presence of a single species for both the complexes $(C_{60})_3 \subset 2$ and $(C_{70})_2 \subset 2$ (Figures S11 and S18). Further, a comparison of the diffusion coefficient

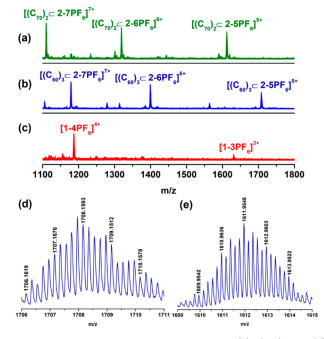


Figure 5. Partial stacked ESI-MS spectra of (a) $(C_{70})_2 \subset 2$, (b) $(C_{60})_3 \subset 2$, and (c) **1.** Experimental isotopic distribution patterns of (d) $[(C_{60})_3 \subset 2\text{-}SPF_6]^{5+}$ and (e) $[(C_{70})_2 \subset 2\text{-}SPF_6]^{5+}$ in ESI-MS.

values for the free trifacial barrel 1 ($D = 6.025 \times 10^{-11} \text{ m}^2/\text{s}$) with host-guest complexes (C_{60})₃ \subset 2 ($D = 5.164 \times 10^{-11} \text{ m}^2/\text{s}$) and (C_{70})₂ \subset 2 ($D = 4.819 \times 10^{-11} \text{ m}^2/\text{s}$) revealed higher hydrodynamic radii for (C_{60})₃ \subset 2 and (C_{70})₂ \subset 2 than the free barrel 1, which also supports the transformation of 1 to the higher homologue (2).

Absorption spectra of 1, $(C_{60})_3 \subset 2$, and $(C_{70})_2 \subset 2$ were recorded (Figure 6) using 10 μ M dimethyl sulfoxide solutions

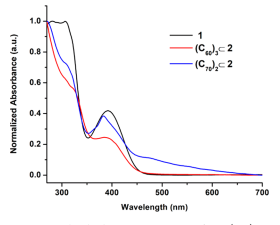


Figure 6. Normalized absorption spectra of 1, $(C_{60})_3 \subset 2$, and $(C_{70})_2 \subset 2$ in DMSO (10 μ M concentration).

of the respective complexes. Trifacial barrel **1** displays two absorptions at 307 and 392 nm due to $\pi - \pi^*$ transitions. $(C_{60})_3 \subset \mathbf{2}$ shows an absorption at 387 nm with a 5 nm blue shift compared to the peak at 392 nm for **1** and a hump at 324 nm (free C_{60} shows an absorption at 321 nm).⁸³ $(C_{70})_2 \subset \mathbf{2}$ shows peaks at 383 and 311 nm and a broad hump at 470 nm (characteristic of C_{70}).⁸⁴ These observations further confirmed the presence of encapsulated C_{60} and C_{70} in these complexes.

The structures of **2**, $(C_{60})_3 \subset 2$, and $(C_{70})_2 \subset 2$ were optimized by the PM6 method. The tetrafacial barrel **2** possesses a cavity large enough to accommodate three C_{60} or two C_{70} molecules (Figure S25). Inside the cavity of **2**, the three C_{60} molecules are linearly oriented along one diagonal (Figure 7a). This

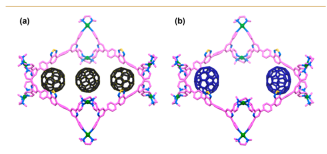


Figure 7. Optimized structures of (a) $(C_{60})_3 \subset 2$ and (b) $(C_{70})_2 \subset 2$. Color codes: carbons of host (pink), nitrogen (blue), sulfur (yellow), palladium (green), carbons of C_{60} guest (black), carbons of C_{70} guest (deep blue). Hydrogen atoms are omitted for clarity.

explains the origin of two peaks corresponding to C_{60} in the ¹³C NMR spectrum of $(C_{60})_3 \subset 2$. The two peripheral C_{60} molecules within the cavity of **2** are in a different chemical environment compared to the central C_{60} molecule, and thus, two time-averaged peaks of slightly different intensities are observed in the ¹³C NMR spectrum (Figure 4b). The two C_{70} molecules are diagonally oriented within the cavity (Figure 7b). Presumably, the $\pi - \pi$ interactions between the cluster of aromatic rings of terpyridine-based building blocks of host **2** and ellipsoidal C_{70} provide the maximum stabilization in two corners of **2**.

Next, competitive binding experiments were performed by treating 1 with excess C_{60} and C_{70} added simultaneously in a 1:1 ratio (4 equiv of each C_{60} and C_{70} was added to 1 equiv of 1 and mixture stirred at 70 °C for 12 h), and the results were monitored by ESI-MS. Interestingly, the peaks obtained correspond to C_{70} encapsulated barrel $(C_{70})_2 \subset 2$ (Figure 8a). Again, when 4 equiv of C_{60} was added to the complex $(C_{70})_2 \subset 2$, no change was observed in mass spectrum, confirming the high stability and selectivity of $(C_{70})_2 \subset 2$ (Figure 8b). However, when 4 equiv of C_{70} was added to the complex (Figure 8b).

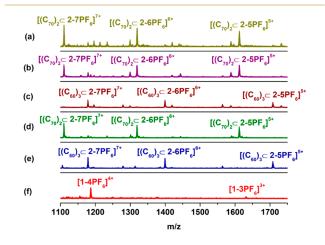


Figure 8. Partial stacked ESI-MS spectra: (a) after treatment of 1 with excess C_{60} and C_{70} (1:1 ratio), (b) after treatment of $(C_{70})_2 \subset 2$ with excess $C_{60'}$ (c) after treatment of $(C_{60})_3 \subset 2$ with excess $C_{70'}$ (d) $(C_{70})_2 \subset 2$, and (e) $(C_{60})_3 \subset 2$ and (f) 1.

complex $(C_{60})_3 \subset 2$, C_{70} could not displace C_{60} molecules from the cavity of 2 (Figure 8c). These results imply that the barrelto-barrel transformation occurs in the presence of both C_{60} and C_{70} . C_{70} has a slightly higher affinity toward the tetrafacial barrel (2); however, once three C_{60} molecules get encapsulated within the cavity of 2, it forms the very stable host-guest complex $(C_{60})_3 \subset 2$, and C_{70} cannot replace the encapsulated C_{60} molecules from the cavity.

It was important to extract the fullerenes from the host– guest complexes to check the fate of the barrel. After screening with different solvents, 1,2-dichlorobenzene was found to be appropriate for the extraction experiments since barrel/host– guest complexes and fullerenes (C_{60}/C_{70}) have orthogonal solubilities in this solvent. C_{60} and C_{70} were successfully extracted from the complexes as was indicated by the pale purple/brown coloration of the 1,2-dichlorobenzene extract in the respective cases. Furthermore, ESI-MS analyses confirmed the existence of only trifacial barrel 1 and not tetrafacial barrel 2 after extraction in both cases (Figure 9a,c).

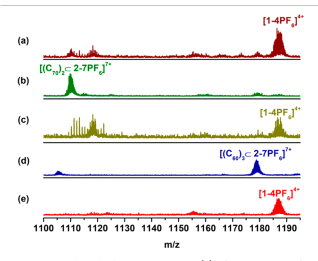
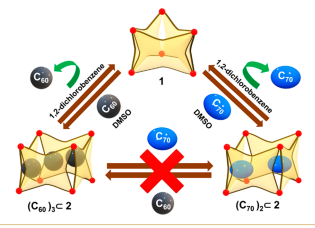


Figure 9. Partial stacked ESI-MS spectra: (a) after extraction of C_{70} from $(C_{70})_2 \subset 2$ using 1,2-dichlorobenzene, (b) $(C_{70})_2 \subset 2$, (c) after extraction of C_{60} from $(C_{60})_3 \subset 2$ using 1,2-dichlorobenzene, (d) $(C_{60})_3 \subset 2$, and (e) 1.

This proves that the structural transformation from trifacial barrel to its higher homologue is feasible only in the presence of fullerenes. The removal of fullerenes from the host-guest complexes readily converted the tetrafacial barrel to the parent trifacial barrel in solution, which indicates that the tetrafacial barrel is a metastable form which can only exist in the presence of fullerenes (Scheme 2).

Singlet oxygen $({}^{1}O_{2})$ generation is important for different uses. Photosensitizers upon light irradiation can transfer its absorbed energy to triplet oxygen to generate singlet oxygen. Fullerenes are well-known photosensitizers that can generate singlet oxygen, but their insolubility in common organic solvents limits their application. Good solubility of the PF₆⁻ analogues of $(C_{60})_3 \subset 2$ and $(C_{70})_2 \subset 2$ in acetonitrile prompted us to investigate the efficiencies of these complexes as photosensitizers for generating singlet oxygen using visible light. To the CD₃CN solutions of anthracene, 2 mol % of PF₆⁻ analogues of these complexes were added separately and irradiated with visible light, and the generation of singlet oxygen was monitored by the formation of anthracene endoperoxide over time using 1 H NMR. The complete Scheme 2. Schematic Representation of Fullerene Induced Barrel to Barrel Transformation, Competitive Binding of C_{60} and C_{70} , and Fullerene Extraction



conversion of anthracene to anthracene-endoperoxide was achieved in 4 and 5 h using $(C_{70})_2 \subset 2$ and $(C_{60})_3 \subset 2$, respectively (Figures S26 and S27). Although, $(C_{70})_2 \subset 2$ has two molecules of fullerene, its slightly higher singlet oxygen generating ability compared to $(C_{60})_3 \subset 2$ can be ascribed to its higher absorption in the visible region (Figure 6). The building block of free barrel 1, $(C_{60})_3 \subset 2$, and $(C_{70})_2 \subset 2$ contains a benzothiadiazole moiety which too is a photosensitizer. So, we also investigated the generation of singlet oxygen using 2 mol % of PF_6^- analogue of free barrel 1 as a photosensitizer in acetonitrile. Even after irradiation with visible light for 20 h, complete conversion of anthracene could not be achieved, thus showing the poor singlet oxygen generating ability of the free barrel 1 (Figures 10 and S28). Encapsulation of multiple fullerenes within the barrel enhanced singlet oxygen generation drastically (Figure 10). Notably, when an excess (20 mol %) of

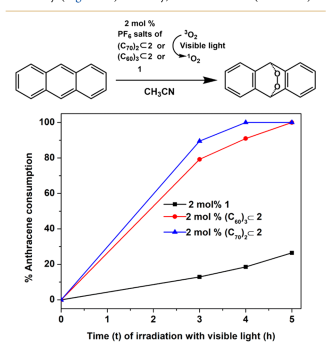


Figure 10. Comparison of singlet oxygen generating ability of $(C_{70})_2 \subset 2$, $(C_{60})_3 \subset 2$, and 1 obtained by monitoring anthracene consumption and formation of its endoperoxide over time upon visible light irradiation using ¹H NMR.

either C₇₀ or C₆₀ was used as a heterogeneous photocatalyst for conversion of anthracene to its endoperoxide in acetonitrile, no conversion was observed (Figures S29 and S30), thus highlighting the importance of solubilizing fullerenes C₆₀/C₇₀ in common organic solvents by encapsulation within a host for singlet oxygen generation.

CONCLUSIONS

In summary, we report here the synthesis of an adaptable trifacial molecular barrel (1) that undergoes structural change to a metastable higher homologue in the presence of suitable guests by efficient host-guest binding. The barrel (1) was obtained via self-assembly of a tetrapyridyl ligand L and a cisblocked Pd(II) acceptor (A). The large cavity, open windows, and π -electron rich aromatic walls of 1 made it a potential candidate for electron deficient large guests like fullerenes C₆₀ and C_{70} . The trifacial barrel (1) showed an uncommon structural transformation⁷⁰ to its metastable higher homologue tetrafacial barrel (2) in the presence of excess C_{60} and C_{70} , stabilizing three C₆₀ and two C₇₀ molecules in its enlarged cavity, respectively. Competitive binding experiments with C₆₀ and C₇₀ confirmed that C₇₀ has a higher affinity for tetrafacial barrel **2** possibly due to greater π -interactions with the walls of the molecular barrel because of the ellipsoidal shape of C₇₀. The host-guest complexes $(C_{60})_3 \subset 2$ and $(C_{70})_2 \subset 2$ were thermodynamically stable and did not show guest exchange upon treatment with C70 and C60, respectively. Trifacial barrel 1 was thus found to be a guest-responsive system which undergoes a structural expansion to a larger homologous molecular container to stabilize multiple fullerene molecules within its cavity. Further, the encapsulated fullerenes were extracted by washing with an appropriate solvent which led to the facile conversion of the metastable tetrafacial barrel to parent trifacial barrel 1. Such a guest mediated reversible structural transformation between a 3D adaptable architecture and its higher homologue is an interesting observation. The PF_6^- analogues of $(C_{60})_3 \subset 2$ and $(C_{70})_2 \subset 2$ generated singlet oxygen under visible light irradiation in acetonitrile. The encapsulation of multiple fullerenes within host 2 was thus pivotal in solubilizing acetonitrile insoluble C_{60}/C_{70} molecules for acting as visible light photosensitizers in acetonitrile.

METHODS

All the chemicals were purchased from commercially available sources and used without further purification. NMR spectra were recorded on Bruker 400 and 500 MHz NMR spectrometers in DMSO- d_6 . Electrospray ionization mass spectra (ESI-MS) were recorded using an Agilent 6538 Ultra-High Definition (UHD) Accurate Mass Q-TOF spectrometer using standard spectroscopic grade solvents. Electronic absorption spectra were recorded on a LAMBDA 750 UV/vis spectrophotometer. White LED (45 W) was used for singlet oxygen generation experiments.

The Gaussian 09 package was used for the computational study.⁸⁵ Trifacial barrel 1, trifacial tube, tetrafacial barrel 2, $(C_{60})_3 \subset 2$, and $(C_{70})_2 \subset 2$ were optimized using the PM6 semiempirical method. No symmetry constraints were used during optimization.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacsau.3c00224.

Experimental section, additional NMR spectra (¹H, COSY, NOESY, DOSY), mass spectra, optimized structures (PDF)

AUTHOR INFORMATION

Corresponding Author

Partha Sarathi Mukherjee – Department of Inorganic and Physical Chemistry, Indian Institute of Science, Bangalore 560012, India; o orcid.org/0000-0001-6891-6697; Email: psm@iisc.ac.in

Authors

- Ranit Banerjee Department of Inorganic and Physical Chemistry, Indian Institute of Science, Bangalore 560012, India
- Soumalya Bhattacharyya Department of Inorganic and Physical Chemistry, Indian Institute of Science, Bangalore 560012, India; orcid.org/0000-0003-4467-6056

Complete contact information is available at: https://pubs.acs.org/10.1021/jacsau.3c00224

Author Contributions

[‡]R.B. and S.B. contributed equally. R.B. carried out the experimental works. S.B. carried out the optimization studies. R.B. and S.B. analyzed the experimental data. P.S.M. supervised the whole project. All authors contributed to writing this manuscript, and they have given approval to the final version of the manuscript. CRediT: **Ranit Banerjee** conceptualization, data curation, formal analysis, investigation, methodology, writing-original draft, writing-review & editing; **Soumalya Bhattacharyya** data curation, formal analysis, investigation, methodology, writing-original draft, writing-review & editing; **Partha Sarathi Mukherjee** conceptualization, funding acquisition, project administration, supervision, writing-original draft, writing-review & editing.

Notes

The authors declare no competing financial interest.

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