

Formation of a Fluorous/Organic Biphasic Supramolecular Octopus Assembly for Enhanced Porphyrin Phosphorescence in Air**

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Dedicated to Professor Jingzhang Gao and Professor Yuan Wang

Abstract: The trinuclear triangle-shaped system [tris(3,5-bis(heptafluoropropyl)-1,2,4-triazolatosilver(I))] (**1**) and the multi-armed square-shaped metalloporphyrin PtOEP or the free porphyrin base H₂OEP serve as excellent octopus hosts (OEP = 2,3,7,8,12,13,17,18-octaethyl-21H,23H-porphine). Coupling of the fluorous/organic molecular octopi **1** and H₂OEP or PtOEP by strong quadrupole–quadrupole and metal– π interactions affords the supramolecular assemblies [1·PtOEP] or [1·H₂OEP] (**2a**), which feature nanoscopic cavities surrounding the upper triangular and lower square cores. The fluorous/organic biphasic configuration of [1·PtOEP] leads to an increase in the phosphorescence of PtOEP under ambient conditions. Guest molecules can be included in the biphasic double-octopus assembly in three different site-selective modes.

Self-assembly of octopus arms around a guest molecule is a powerful strategy to obtain architectures in which the steric and electronic properties of the shell control the guest environment.^[1] Octopus molecules represent a class of molecular hosts composed of a rigid core with multiple surrounding flexible substituents. A variety of octopus hosts have been explored since the pioneering work of Vögtle and co-workers.^[1] On the other hand, π – π and metal– π interac-

tions between an octopus core and an aromatic guest may act as driving forces towards an ordered one-dimensional (1D) octopus-like supramolecular array. Numerous 1D π -stacked and/or metal-mediated supramolecular polymers have been reported in the last decade.^[2] The use of fluorous octopus arms may induce novel functions akin to those enabled by fluorous metal–organic frameworks (FMOFs)^[3,4] or non-porous analogues thereof that bear fluorous surfaces.^[5] The monovalent Ag^I, Au^I, and Cu^I salts of 1,2,4-triazolate (Tz) readily assemble into discrete trinuclear macrocycles, [M₃(Tz)₃],^[6] which can serve as the octopus core while the six 3,5-substituents serve as flexible octopus arms.^[7,8] The electronic and chemical properties of fluorous [M₃Tz₃]-based octopus complexes are easily tunable using triazole chemistry. This work and parallel prior studies by Dias et al. and our group show that a fluorous octopus such as [Ag₃($\mu^{1,2}$ -3,5-(n-C₃F₇)₂Tz)₃] (**1**) can be constructed and has excellent inclusion and optoelectronic properties.^[9]

Herein, we show that fluorous octopus **1** can be combined with porphyrin octopi to form biphasic double-octopus assemblies with fluorous and non-fluorous arms on opposite sides; the corresponding nanoscopic phases thus display interesting inclusion properties. The metalloporphyrin PtOEP and the free porphyrin base H₂OEP (OEP = 2,3,7,8,12,13,17,18-octaethyl-21H,23H-porphine) were selected because of their wide-ranging applications.^[10–12] DFT calculations showed that the molecular electrostatic potentials (MEPs) of H₂OEP and PtOEP exhibit an electron-rich (π -basic) region at the center of the squares whereas [Ag₃($\mu^{1,2}$ -3,5-(n-C₃F₇)₂Tz)₃] (**1**) is very electron-poor (strongly π -acidic) at the center of the nine-membered ring (Scheme 1b). The electron-rich regions in H₂OEP and PtOEP are susceptible to quadrupole–quadrupole interactions with the strongly electron-deficient region of **1**. The difference between the Q_{zz} values of **1** (+20) and the porphyrin systems (–24 for H₂OEP; –30 for PtOEP) accentuates strong attractive interactions, which assist π – π and/or metal– π forces in the formation of the stacked pairs [1·H₂OEP] (**2a**) and [1·PtOEP] (**2**).

The reaction of **1** with PtOEP or H₂OEP in dichloromethane or benzene afforded double-octopus assembly **2** or **2a** in high yield. Both **2** and **2a** can be recrystallized from benzene to give red [2·(C₆H₆)_{1.5}] or dark-purple [2a·(C₆H₆)] crystals. **2a** crystallizes in the triclinic space group *P* $\bar{1}$ as a discrete [1·H₂OEP] pair (Figure 1) with one benzene molecule per asymmetric unit. The adduct pair consists of

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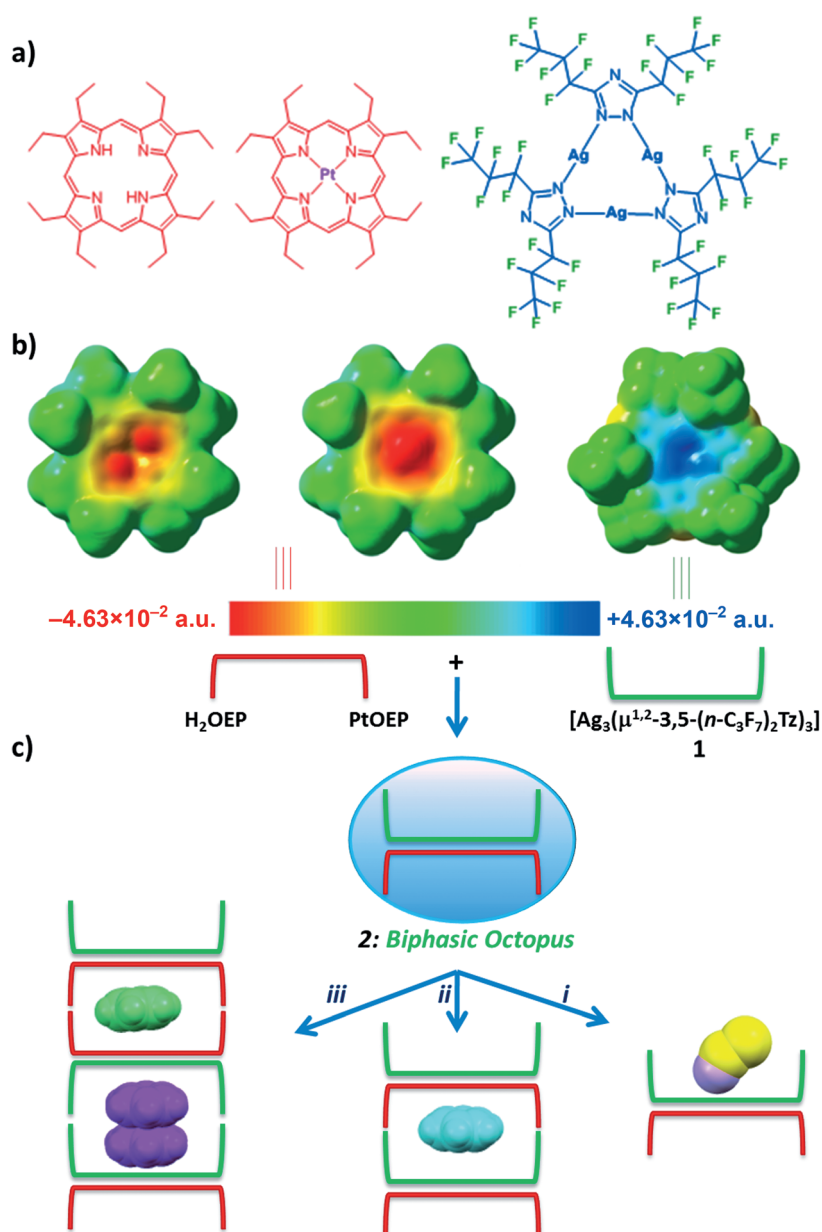
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Scheme 1. a, b) Chemical structure (a) and MEPs (b) of H₂OEP, PtOEP, and **1** mapped on the electron density surfaces. c) Formation of biphasic double-octopus assembly **2** by quadrupole–quadrupole interactions. Three different guest-inclusion modes are possible: i) head-on inclusion of a linear guest within the fluorine-lined cavity; ii) inclusion of an aromatic guest within the nest consisting of a semi-fluorous phase and a semi-organic phase; iii) inclusion of an aromatic π-dimer within the fluorine-lined cage while an aromatic monomer is included within the organic cage. The MEP values are plotted on a scale ranging from -4.63×10^{-2} a.u. (red-most values; a.u. = atomic units) to $+4.63 \times 10^{-2}$ a.u. (blue-most values).

fluorous bowl-shaped **1** and H₂OEP, which are held together by coordination, π -stacking (quadrupole–quadrupole), and van der Waals forces. In this rare type of Ag₃–porphyrin bonding, the imino hydrogen atoms are located on one pair of pyrrole N atoms in the porphyrin. The unprotonated N11/N13 atoms and the Ag₃ cluster were found to strongly interact, with interatomic distances of 2.53 Å (Ag3–N11) and 2.95 Å (Ag1–N13). The shorter Ag3–N11 distance is comparable

with those reported for supramolecular arrays of Ag^I–polypyridines (2.15–2.55 Å),^[2d–f] and shorter than the axial Ag–N(Tz) bond (2.62 Å) found in FMOF-1.^[4] The strong interaction between the porphyrin ring of H₂OEP and Ag₃ cluster **1** is such that the Ag atoms protrude out of the plane of the three Tz rings, with maximum vertical displacements of the silver(I) atoms from the plane of the six N atoms of 0.33 (Ag1), 0.10 (Ag2), and 0.63 Å (Ag3). Remarkably, the two unprotonated N atoms that are bonded to the Ag₃ cluster also protrude out of the porphyrin mean plane, with a vertical distance to the porphyrin plane of 0.12 (N11) and 0.10 Å (N13). DFT calculations gave a binding energy of 62.06 kcal mol⁻¹ for **2a**, indicating that the Ag₃–porphyrin bonding is strengthened by electrostatic quadrupolar interactions between **1** and H₂OEP (Supporting Information, Figure S2). The [**1**·H₂OEP] pair is stabilized by an octopus shell formed by six perfluoroalkyl *n*-C₃F₇ arms oriented upwards to provide a fluorine-lined electron-poor cavity above the stack; eight C₂H₅ arms are oriented downwards to afford a hydrogen-lined electron-rich cavity below the stack, which renders the adduct polar, asymmetric, and biconcave. The [**1**·H₂OEP] units are packed along the *a* axis such that the fluorine-lined bowl and the hydrogen-lined bowl are glued together to form a benzene-encapsulating supramolecular cage (Scheme 1 c, ii).

Biphasic double-octopus **2** crystallizes in the monoclinic space group *P21/c* as two crystallographically independent [**1**·PtOEP] pairs (Figure 1, right) along with three crystallographically independent benzene molecules in the asymmetric unit. The 1:1 cofacial adduct **2** adopts a similar biconcave conformation as **2a**, except that the face-to-face overlap in [**1**·PtOEP] is strengthened by donor–acceptor Pt–Ag metallophilic bonding. The shortest Pt–Ag and Pt–Ag₃ centroid distances in **2** are 3.040 and 2.847 Å, respectively, which fall in the range of Pt–Ag dative-bond distances reported in the literature (2.7–3.2 Å).^[13a,b] The binding energy of **2** was determined to be 67.81 kcal mol⁻¹ by DFT calculations, which indicates that the Pt→Ag metallophilic/dative bond is strengthened by electrostatic quadrupolar interactions between **1** and PtOEP (Figure S2). The quadrupolar interactions in **2** and **2a** are stronger than those in the non-octopus adduct [**1**·Au(bzim)]₃/[[Hg(C₆F₄)₃]₃], which was reported by Burini and co-workers (bzim = 1-benzylimidazole; see the Supporting Information).^[13c]

The asymmetric units in **2** are related by a two-fold axis such that a one-dimensional array of stacked chromophores

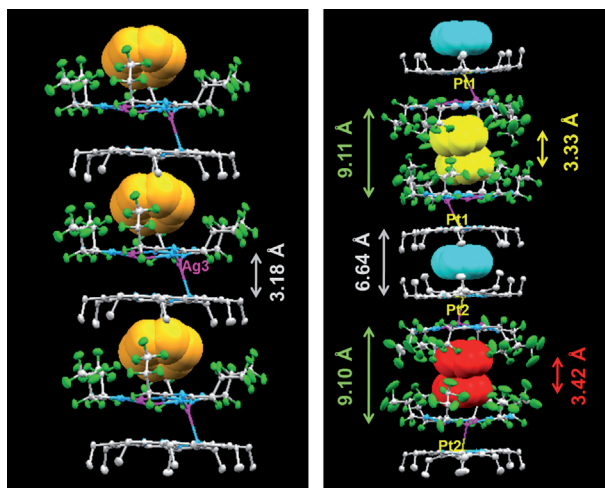


Figure 1. Illustration of infinite π -stacks in crystals of $[2 \cdot \text{C}_6\text{H}_6]$ and $[2 \cdot \text{C}(\text{C}_6\text{H}_6)_{1.5}]$ along the a and c^* axes, respectively. The three independent benzene molecules in $[2 \cdot \text{C}(\text{C}_6\text{H}_6)_{1.5}]$ are indicated as red, blue, and yellow space-filling spheres.

with alternating benzene–1–PtOEP–benzene–PtOEP–1–benzene units is generated along the c^* axis (Figure 1 b). Aside from Pt–Ag metallophilic and electrostatic/quadrupolar forces, the extended stacking is mediated by π – π interactions between adjacent chromophores, including those between benzene–benzene, benzene–1, 1–PtOEP, and PtOEP–benzene units (interplanar angles of 0, 12.7, 1.1, and 3.3°, respectively), resulting in pair-wise separations of 3.33, 6.64, and 9.11 Å for the benzene–benzene, porphyrin–porphyrin and 1–1 pairs, respectively. Such self-assembled aromatic one-dimensional columns that combine metal–metal, π – π , and metal– π interactions favor charge transport along the column chains, favoring applications in organic electronics, for example, in organic photovoltaics (OPVs), light-emitting diodes (OLEDs), and field-effect transistors (OFETs).^[14]

One highlight of the stack is the aromatic (benzene)₂ π -dimer, which is sandwiched between cofacially stacked macrocycles 1 and enclosed by the fluorine-lined hydrophobic cavity formed by twelve fluororous $n\text{-C}_3\text{F}_7$ groups. A single benzene molecule is also sandwiched between two cofacial PtOEP planes and is well-nested within a cavity formed by 16 ethyl groups (Scheme 1 c, iii). To the best of our knowledge, this is the first example of alternating fluororous/non-fluororous cages formed by cofacially stacked octopus cavities. The biphasic cavities act as host cages to encapsulate π -dimer and monomer guest molecules.

We found that formation of the octopus couple in $[2 \cdot (\text{C}_6\text{H}_6)_{1.5}]$ can enhance the PtOEP phosphorescence in air. It is well known that pure PtOEP is a very weak emitter in air owing to its extreme sensitivity to O₂ quenching and concentration quenching through a strong bimolecular annihilation process.^[10–12] This limitation is a critical bottleneck for the application of PtOEP systems in organic light-emitting devices and full-color displays that require high stability.^[15–17] One strategy to circumvent this bottleneck involves the doping of PtOEP into a host matrix of an amorphous organic semiconductor and sealing the device under inert atmos-

phere.^[15] As O₂ quenching of the PtOEP phosphorescence occurs through axial interactions of O₂ with the porphyrin plane and the triplet–triplet annihilation process through intermolecular PtOEP π -interactions, the coupling of PtOEP with a fluororous octopus host effectively protects PtOEP from axial interactions with O₂ and completely prevents the concentration-dependent bimolecular self-annihilation processes.

As seen in Figure 1, the PtOEP molecules in $[2 \cdot (\text{C}_6\text{H}_6)_{1.5}]$ are well-protected by coupling with 1 under the fluororous shell, which captures invasive guests, such as benzene and O₂. The configuration of $[2 \cdot (\text{C}_6\text{H}_6)_{1.5}]$ keeps the PtOEP units apart from each other and prevents their interactions with O₂ (which is “filtered” by the fluororous pore).^[4a] Electronic spectra of 2 under various conditions are shown in Figure 2 a–f. In agreement with its solution absorption spectrum

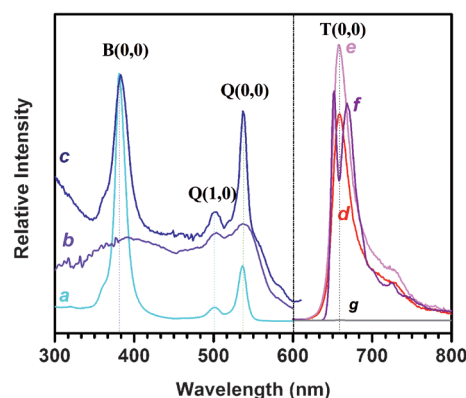


Figure 2. a–c) Solution absorption (a; in CH_2Cl_2), solid-state diffuse reflectance (b), and excitation (c; $\lambda_{\text{em}} = 660 \text{ nm}$) spectra of compound 2. d–f) Luminescence spectra of crystalline 2 ($\lambda_{\text{ex}} = 540 \text{ nm}$) in ambient air (d; $\tau = 55 \mu\text{s}$), in 100% N_2 (e; $\tau = 64 \mu\text{s}$), and at 77 K in air (f; $\tau = 93 \mu\text{s}$). g) The gray trace shows the vanishing luminescence of a solid sample of PtOEP under ambient conditions ($\lambda_{\text{ex}} = 540 \text{ nm}$) for the same instrument settings.

in CH_2Cl_2 , the diffuse reflectance and excitation ($\lambda_{\text{em}} = 660 \text{ nm}$) spectra of a solid sample of 2 are governed by the Q- and Soret-band transitions at 380, 502, and 537 nm, which were classified as B(0,0), Q(1,0), and Q(0,0) transitions, respectively.^[12,18] Although PtOEP exhibits rather weak emission in air (Figure 2 g),^[12] crystals of 2 exhibit bright-red luminescence characterized by a porphyrin-based triplet emission at 658 nm [T(0,0)], with weaker emissions at the vibronic harmonic overtones (690 and 720 nm). The solid-state phosphorescence of 2 under ambient conditions exhibits only small intensity increases under oxygen-free conditions (e.g., in pure N_2 ; Figure 2 e), which indicates a rather insensitive response to the presence of O₂ as a result of the well-protected PtOEP sites (see the crystal structure of $[2 \cdot (\text{C}_6\text{H}_6)_{1.5}]$; Figure 1). The strong Ag₃–Pt interactions and heavy-atom effects result in a strong sensitization of the doublet T(0,0) band. The lifetime of the phosphorescence of the solid sample of 2 also corroborates the strong sensitization of the PtOEP phosphorescence by the fluororous octopus molecules 1, as the τ values that we obtained even in ambient

air and at ambient temperature are similar to those measured for PtOEP-doped thin films in inert environments (Figure 2).

Crystals of $[2 \cdot (C_6H_6)_{1.5}]$ remained intact even after standing in air for extended periods of time, suggesting that the benzene molecules are tightly included in the lattice sites. However, when these crystals were exposed to vapors of polar organic solvents, for example, acetone or acetonitrile, the red emission of **2** was significantly quenched. To reveal whether the guest benzene molecules in $[2 \cdot (C_6H_6)_{1.5}]$ played a role in enhancing PtOEP phosphorescence, we replaced them with other solvent molecules. Slow evaporation of a CH_2Cl_2 solution of **2** in the presence of acetonitrile or acetone resulted in non-luminescent dark-red crystals. X-ray analysis showed that these crystals were formed by linear guest inclusion in the biphasic double-octopus system **2** (Scheme 1 c, i) to form $[2 \cdot MeCN]$ and $[2 \cdot Me_2CO]$, respectively. The MeCN and Me_2CO solvent molecules were well-nested in the hydrophobic fluorine-lined cavity with head-on coordination to the Ag_3 unit (Figure 3). The corresponding $Ag-O$ (acetone)

as O_2 may reach exposed sides of the porphyrin plane. Such interactions likely represent a principal cause for the emission quenching observed for $[2 \cdot MeCN]$ and $[2 \cdot Me_2CO]$ in air, whereas guest inclusion of benzene molecules in $[2 \cdot (C_6H_6)_{1.5}]$ likely plays a sensitizing/protecting role for the PtOEP triplet emission in air. By applying a vacuum to crystals of $[2 \cdot MeCN]$ and $[2 \cdot Me_2CO]$ followed by N_2 exposure, the emission of PtOEP was recovered and gave the same emission profile as $[2 \cdot (C_6H_6)_{1.5}]$, which reinforces the aforementioned hypothesis that acetone and acetonitrile inclusion represents a quenching pathway whereas benzene inclusion represents a sensitizing pathway for porphyrin phosphorescence.

In summary, the crystal structures of **2a** and **2** revealed strong electrostatic quadrupole–quadrupole interactions between the electron-deficient Ag_3 macrocycles and the electron-rich free porphyrin base or the metalloporphyrin cores. These interactions acts as important organizational forces in the formation (“self-recognition”) of stacked pairs of **1** and the porphyrin systems. The van der Waals interactions between the perfluoroalkyl $n-C_3F_7$ groups in **1** and the ethyl groups in the porphyrin systems lead to a conformation consisting of a biconcave biphasic octopus pair. The fluorine-lined cavities formed by the octopus-arm-like $n-C_3F_7$ groups around the Ag_3 cores enhance the chemical, photochemical, and thermal robustness of the immediate metal-cluster environment. In our biphasic octopus assembly, the PtOEP phosphorescence is optimized in two ways as 1) O_2 quenching is effectively prevented and 2) self-quenching and bimolecular triplet–triplet annihilation processes are completely suppressed. Ongoing efforts target “best guest” chromophores for the biphasic octopus couple **2** aside from benzene, such as pyrene, perlene, or coronene, to further optimize the phosphorescence of metalloporphyrins in ambient air.

Keywords: fluorine · heterocycles · luminescent materials · porphyrins · supramolecular chemistry

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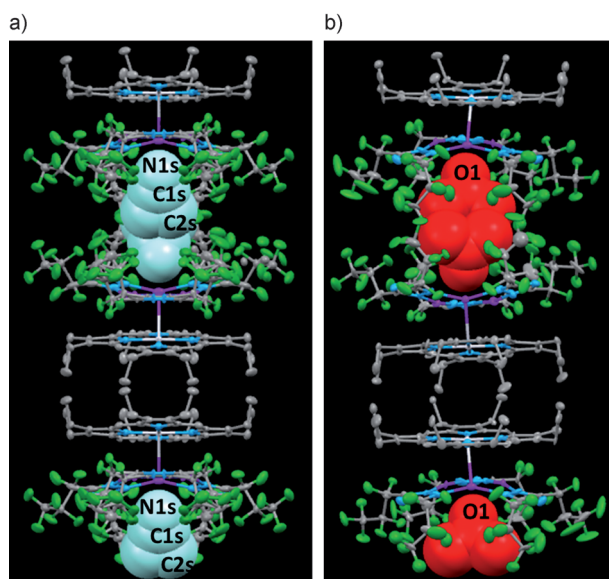


Figure 3. a, b) Crystal packing viewed along the *a* axis in the crystals of $[2 \cdot MeCN]$ (a) and $[2 \cdot Me_2CO]$ (b). Acetonitrile and acetone molecules are indicated as light-blue and red space-filling spheres, respectively.

distances were determined to be 2.746, 2.859, and 3.038 Å, and the $Ag-N(MeCN)$ distances were found to be 2.815, 3.134, 3.148 Å with centroid– Ag_3 separations of 2.097 and 2.304 Å for acetone and acetonitrile, respectively.^[19,20] These data indicate the presence of strong dipole–quadrupole bonding of acetone and MeCN with Ag_3 in $[2 \cdot Me_2CO]$ and $[2 \cdot MeCN]$, respectively, as described in our recent work conducted in collaboration with the Burini and Galassi groups.^[21] The strong interactions between the solvent molecules and **2** suggest that the $[Ag_3(\mu^{1,2}-3,5-(n-C_3F_7)_2Tz)_3]$ macrocycles that bear a fluorine octopus shell are strongly Lewis acidic/ π -superacidic. An analysis of the crystal packing in $[2 \cdot Me_2CO]$ and $[2 \cdot MeCN]$ (Figure 3) revealed small pores between the porphyrin alkyl chains, where gas molecules such

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