Host-Enhanced Phenyl-Perfluorophenyl Polar−π Interactions
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ABSTRACT: Phenyl-perfluorophenyl polar−π interactions have been revisited for the design and fabrication of functional supramolecular systems. The relatively weak associative interactions (ΔG ≈ −1.0 kcal/mol) have limited their use in aqueous self-assembly to date. Herein, we propose a strategy to strengthen phenyl-perfluorophenyl polar−π interactions by encapsulation within a synthetic host, thus increasing the binding affinity to ΔG = −15.5 kcal/mol upon formation of heteroternary complexes through self-sorting. These heteroternary complexes were used as dynamic, yet strong, cross-linkers in the fabrication of supramolecular gels, which exhibited excellent viscoelasticity, stretchability, self-recovery, self-healing, and energy dissipation. This work unveils a general approach to exploit host-enhanced polar−π interactions in the design of robust aqueous supramolecular systems.

Fluorinated small molecules and macromolecules have been widely utilized in organic synthesis, polymer science, medicinal chemistry, drug delivery, liquid crystals, and supramolecular chemistry. Noncovalent interactions between aromatic molecules are important in both natural and artificial supramolecular systems. The introduction of electron-withdrawing fluorine substituents onto aromatic molecules serves to polarize their π-conjugated systems, enhancing π−π stacking through electrostatic interactions. Phenyl-perfluorophenyl polar−π interactions are an elegant example of electrostatic reinforcement. This interaction emerged from the first discovery of a 1:1 cocrystal of hexafluorobenzene and benzene in 1960 (Figure 1a) followed by structural elucidation in 1992 by William et al. Coates and co-workers further developed phenyl-perfluorophenyl polar−π interactions to align diyne molecules for photopolymerization in the crystalline state.

While the perfluorophenyl group maintains preferential stacking selectivity with electron-rich arene groups, it also offers a position for functionalization. Thus, this interaction has been extensively adopted as a facile binding motif for crystal engineering and protein/peptide folding. Although significant progress has been made, the relatively low binding affinity of phenyl-perfluorophenyl polar−π interactions in water (ΔG ≈ −1.0 kcal/mol, K ≈ 1.5 M−1) limits its widespread application. Herein, we propose a host–guest strategy to mediate phenyl-perfluorophenyl polar−π interactions, leading to a substantial increase in overall binding strength. There are several different host molecules capable of encapsulating multiple guests simultaneously. Among them, cucurbit[8]uril (CB[8]) is one of the ideal options on account of its high-energy water release from its nanocavity upon guest incorporation, which can provide tremendous enthalpic contributions for stabilizing ternary host–guest complexes. CB[8]-mediated ternary interactions often exhibit high binding strengths ≥ 15 kcal/mol, which can be further enhanced in aqueous solution.

Figure 1. Schematic representation of (a) the first 1:1 cocrystal of hexafluorobenzene and benzene; (b) design concept of CB[8]-mediated polar−π interaction between perfluorophenyl (5FBVI) and phenyl (BVI) guests; (c) overview of thermodynamic pathways toward ternary complexation: social and narcissistic self-sorting. Br−counterions omitted for clarity.
both homoternary and heteroternary interactions have been widely used as the driving force for fabrication of various aqueous supramolecular systems including supramolecular oligomers and polymers, supramolecular hydrogels, supramolecular organic frameworks, protein/peptide self-assemblies, and functional interfaces. Therefore, we posited that CB[8] could enhance phenyl-π interactions. Heteroternary complexation of singly charged guest molecules within CB[8] mediated by polar-π interactions would represent a new class of CB[8]-mediated heteroternary interaction, generating a promising and powerful noncovalent driving force for supramolecular self-assembly.

Two N-vinyl imidazolium guests were designed (Figure 1b), consisting of perfluorophenyl (5FBVI) and phenyl (BVI) groups. We envisioned that an equimolar mixture of 5FBVI, BVI, and CB[8] should result exclusively in a 1:1:1 heteroternary complex. Perfluorophenyl moieties should not form homoternary complexes on account of electrostatic repulsion, instead favoring secondary binding with an electron-rich phenyl group. To study the influence of electrostatic repulsion, a series of guest molecules varying in fluorine substituents and cationic side groups were synthesized (Chart S1). We utilized the 5FBVI-BVI-CB[8] complex as a supramolecular cross-linker for fabricating robust transparent polyacrylamide networks exhibiting excellent mechanical properties and ionic conductivity.

Extensive 1H and 19F NMR titrations were performed to study formation of the 5FBVI-BVI-CB[8] complex. Titration of BVI into an equimolar mixture of 5FBVI and CB[8] resulted in the gradual appearance of the BVI phenyl protons at 6.39, 6.49, and 6.59 ppm, respectively (Figure 2a (left), S4). The phenyl proton peaks had similar chemical shifts to the BVI and CB[8] should result exclusively in a 1:1:1 heteroternary complex. Perfluorophenyl moieties should not form homoternary complexes on account of electrostatic repulsion, instead favoring secondary binding with an electron-rich phenyl group. To study the influence of electrostatic repulsion, a series of guest molecules varying in fluorine substituents and cationic side groups were synthesized (Chart S1). We utilized the 5FBVI-BVI-CB[8] complex as a supramolecular cross-linker for fabricating robust transparent polyacrylamide networks exhibiting excellent mechanical properties and ionic conductivity.

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homoternary complex (2-BVI-CB[8]) at 6.23, 6.46, and 6.54 ppm, suggesting the phenyl group in the SFBVI-BVI-CB[8] complex is in a different chemical environment compared to 2-BVI-CB[8]. 19F NMR (Figure 2a (right), S5) showed three peaks in the binary mixture of SFBVI-CB[8] (−146.2, −153.0, and −162.8 ppm), which gradually disappeared while three new peaks at −141.2, −152.2, and −160.9 ppm appeared. Both titrations indicated formation of a new host–guest complex, SFBVI-BVI-CB[8]. Additionally, a series of MS titration experiments were conducted. Titrations of BVI into a 1:1 solution of SFBVI-CB[8] yielded the heteroternary complex with an intense ion peak at m/z = 894.2794, Figure 2b, which gradually increased with a concomitant decrease of the binary complex at m/z = 802.2300, confirming the CB[8]-mediated polar–π interaction.

Isothermal titration calorimetry (ITC) was employed to study the thermodynamic mechanism behind this new complexation. Titrations of BVI and SFBVI into CB[8] led to two different cooperative binding curves. A clear transition at a 2:1 molar ratio was observed upon titration of BVI into CB[8] (Figure 3a), indicating formation of a homoternary complex. This binding curve is indicative of positive cooperativity, as reported previously. In contrast, titration of SFBVI into CB[8] showed a sharp transition at 1 equiv (Figure 3b), indicating favored formation of a binary complex SFBVI-CB[8] (K_a = 6.83 × 10^4 M^-1, Table 1). Incorporation of a second SFBVI guest into the binary complex is significantly suppressed, exhibiting negative cooperativity. Free BVI was then titrated into the 1:1 binary complex of SFBVI-CB[8] resulting in a secondary binding curve, K_2 = 3.36 × 10^4 M^-1 (Figure 3c). The calculated overall binding constant of the heteroternary SFBVI-BVI-CB[8] complex is 2.29 × 10^11 M^-2. Compared to the homoternary complex, 2-BVI-CB[8], this heteroternary complex exhibits higher binding strength on account of the additional electrostatic interactions between the phenyl and perfluorophenyl units.

Binding cooperativity plays a critical role in stabilizing the heteroternary SFBVI-BVI-CB[8] complex; unexpectedly, a strong negative cooperativity with an interaction parameter α = 0.0004 was observed between SFBVI and CB[8]. To gain insight into the importance of cooperativity, three N-vinyl imidazolium guests (4FBVI, 3FBVI, 2FBVI) containing 2,3,5,6-tetrafluorophenyl, 3,4,5-trifluorophenyl, and 3,5-difluorophenyl groups, respectively, were synthesized. ITC indicated that 2:1 complexation with CB[8] was dramatically enhanced compared to SFBVI, Figure 4.

Table 1 reports key thermodynamic parameters; α values of 4FBVI, 3FBVI, and 2FBVI showed a steady increase, indicating a change from negative cooperativity for 2FBVI and 4FBVI to positive cooperativity for 3FBVI and 2FBVI. These data indicate that weakening electrostatic repulsion between guests by decreasing the number of fluorine substituents leads to favorable 2:1 homoternary complexation, thus changing the binding cooperativity from negative to positive. A previous report of a heteroternary complex between tetrafluorophenyl, naphthalenyl, and CB[8] could support this effect; however, in this case additional long-range Coulombic and CH–π interactions are required to achieve binding as the complexation lacks strong negative cooperativity. In our study the SFBVI-CB[8] complex is the only dominant 1:1 complex observed, Figure 3b.

To validate this point, and further extend the structural diversity for CB[8]-mediated polar–π interactions, three additional perfluorophenyl guests bearing N-methyl imidazolium, pyridinium, and trimethylammonium moieties were evaluated. The heteroternary complexations of these three guests with CB[8] all exhibited strong negative cooperativity as expected (Figure S7), unambiguously demonstrating that the perfluorophenyl motif displays strong electrostatic repulsion within the CB[8] cavity, key to favoring the formation of a 1:1 complex.

The mechanism behind the formation of the heteroternary SFBVI-BVI-CB[8] complex is attributed to social self-sorting, Figure 1c. Two complexation pathways exist for an equimolar mixture of SFBVI, BVI, and CB[8]: social and narcissistic self-sorting. Thermodynamic data showed a substantial preference for social self-sorting with a difference in Gibbs free energy of 3.1 kcal/mol (Figure 1c), and a relative ratio between heteroternary and homoternary complex formation of nearly 200:1. Recently it was reported that a mixture of both social and narcissistic self-sorting is obtained with difluorophenyl guest motifs. In contrast, we are able to control the outcome of the self-sorting process, which results solely in the formation of the CB[8]-mediated phenyl-perfluorophenyl complex through polar–π interactions.

To demonstrate the utility of quantitative social self-sorting in CB[8]-mediated polar–π interactions, we employed the heteroternary complex SFBVI-BVI-CB[8] as a supramolecular cross-linker to fabricate robust viscoelastic polyacrylamide networks. A mixture of 2.5 mol % of SFBVI-BVI-CB[8] cross-linker and 95 mol % of acrylamide monomer with 0.003 mol % photoinitiator was photopolymerized for 6 h (λ = 350 nm, Figure 5a). The resultant transparent gel could be readily stretched to 8-fold its original length demonstrating complete self-recovery within 3 min. By reducing the ratio of supramolecular cross-linker, the storage and loss moduli of

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**Table 1. Overview of Thermodynamic Data for Homoternary Complexations (H_2O, 298 K)**

|          | K_1 (10^4 M^-1) | K_2 (10^4 M^-1) | α       |
|----------|-----------------|-----------------|---------|
| SFBVI    | 683 ± 187       | 0.07 ± 0.01     | 0.0004 ± 0.0001 |
| 4FBVI    | 230 ± 36        | 33 ± 6          | 0.57 ± 0.14 |
| 3FBVI    | 95 ± 28         | 102 ± 54        | 4.27 ± 2.61 |
| 2FBVI    | 61 ± 25         | 69 ± 43         | 4.52 ± 3.35 |
| BVI      | 44 ± 5          | 14 ± 1          | 1.29 ± 0.16 |

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Figure 4. ITC plots (H_2O, 298 K) of CB[8] homoternary complexation with SFBVI, 4FBVI, 3FBVI, and 2FBVI. [Guest] = 1.0 mM, [CB[8]] = 0.05 mM.
the obtained gels are readily tuned over a wide range, $10^5$–$10^7$ Pa, Figures 5b and S12.

The supramolecular gels (Figure 5b; green, blue, and red data sets) showed higher viscoelasticity with both storage and loss moduli $1$–$2$ orders of magnitude above the functionalized copolymer gel without CB[8] present (yellow data set). Furthermore, when the 2.5 mol % cross-linked gel was subjected to multiple step−strain cycles (5% and 500% strain at 1 rad/s), a reversible sol−gel transition was observed, Figure 5c. This reversible behavior was replicated over a substantially longer time period (>1 h) without loss of material integrity, Figure S13. The resilience of the SFBV1-BVI-CB[8] cross-linked gel toward step-strain measurements demonstrates the reversible and robust nature of the network and highlights its rapid self-healing properties.

Owing to the dynamic nature of CB[8]-mediated cross-links, the transient polymeric networks dissipate energy upon application of a force, giving rise to excellent mechanical properties. To obtain the activation energy for local chain motion, a series of time−temperature superposition (TTS) experiments were carried out, quantifying the energy-dissipation capability of the SFBV1-BVI-CB[8] cross-linked gel (Figure S14). Fitting the shift parameters ($a_T$, $b_T$) revealed the activation energy to be 18.0 kcal/mol, comparable to the unfolding barrier of the I27 domain of the human muscle protein titin, which has an activation energy of 17.0 kcal/mol.

On account of the charged SFBV1 and BVI guest moieties, we probed the ionic conductivity of the obtained material through incorporation within a series circuit connected to two LEDs, Figure S15. The high brightness of the LEDs connected to the gel in its original state illustrates its considerable ionic conductivity. Stretching the gel reduced its cross-sectional area, leading to an increase in resistance and an observed decrease in brightness of the LEDs. Removal of the stretching force returned the gel to its original length with concomitant increase in brightness over a 2 min period, demonstrating the gel’s self-recovery properties.

In conclusion, we have successfully developed a new type of CB[8]-mediated polar−π interaction in aqueous media. Through host−guest complexation within CB[8], the binding affinity of phenyl-perfluorophenyl polar−π interactions is significantly enhanced with an increase in ΔG from $-1.0$ kcal/mol to $-15.5$ kcal/mol, assuring sole formation of the heteroternary complex upon social self-sorting. To demonstrate the utility of CB[8]-mediated polar−π interactions, we applied a model complex (SFBV1-BVI-CB[8]) as a supramolecular cross-linker in the fabrication of completely transparent polyacrylamide gel networks, which exhibited excellent viscoelasticity, stretchability, self-recovery, self-healing, and ionic conductivity. These materials are promising candidates for research in areas including structural biomaterials, flexible electronics, artificial muscles, soft robots, etc. We highly anticipate that this work will inspire renewed interest and research into the exploitation of polar−π interactions as robust building blocks in supramolecular chemistry and materials science.

■ ASSOCIATED CONTENT

* Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c02275. Materials and methods; molecular design and synthetic protocols; procedures for preparation of supramolecular polyacrylamide networks; study on CB[8]-mediated polar−π interactions by $^1$H and $^{19}$F NMR, ESI-MS, and ITC; rheological characterization of supramolecular polyacrylamide networks; photographs for demonstration of ionic conductivity (PDF)

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Notes
The authors declare no competing financial interest.

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