

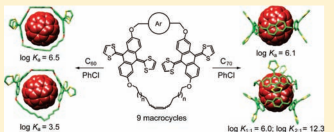
Macrocyclic Hosts for Fullerenes: Extreme Changes in Binding Abilities with Small Structural Variations.

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Supporting Information

ABSTRACT: Exploiting the shape and electronic complementarity of C_{60} and C_{70} with π -extended derivatives of tetrathiafulvalene (exTTF), we have very recently reported a macrocyclic receptor featuring two exTTF recognizing units which forms 1:1 complexes with C_{60} with log $K_a=6.5\pm0.5$ in chlorobenzene at 298 K. This represents one of the highest binding constants toward C_{60} reported to date and a world-record for all-organic receptors. Here, we describe our efforts to fine-tune our macrocyclic bis-exTTF hosts to bind C_{60} and/or C_{70} , through structural variations. On the basis of preliminary



molecular modeling, we have explored p-xylene, m-xylene, and 2,6-dimethylnaphthalene as aromatic spacers between the two exTTF fragments and three alkene-terminated chains of different length to achieve macrocycles of different size through ring closing metathesis. Owing to the structural simplicity of our design, all nine receptors could be accessed in a synthetically straightforward manner. A thorough investigation of the binding abilities of these nine receptors toward C_{60} and C_{70} has been carried out by means of UV—vis titrations. We have found that relatively small variations in the structure of the host lead to very significant changes in affinity toward the fullerene, and in some cases even in the stoichiometry of the associates. Our results highlight the peculiarities of fullerenes as guests in molecular recognition. The extreme stability of these associates in solution and the unique combination of electronic and geometrical reciprocity of exTTF and fullerenes are the main features of this new family of macrocyclic hosts for fullerenes.

1. INTRODUCTION

The search for molecular receptors for fullerenes was initiated soon after their discovery with the pioneering work of the groups of Diederich, Ringsdorf, and Wennerström. The main motivations that have kept it as a very active area of research are the selective association of specific fullerenes to achieve their purification from complex mixtures self-assembly. In both cases the formation of associates with fullerenes of high stability in solution—that is, the construction of hosts that show large binding constants toward the fullerenes—is one of the main goals. To achieve this objective, a broad variety of molecular fragments capable of establishing positive noncovalent interactions with the outer surface of fullerenes have been explored. These include derivatives of calixarenes, 3b,3j,3n,5 cyclotriveratrylenes, 3q,3r,6 corannulenes, 3p,7 cyclic paraphenyleneacetylenes, 8 π -extended tetrathiafulvalenes, 3h,3j,3o,3t and, most frequently, porphyrins. 3c,3s,3v,9

Although the experimental results from different groups are sometimes difficult to compare due to the use of different methods to assess the binding constants and to the different solvents utilized, taking C_{60} as a reference fullerene guest, it is fair to say that obtaining hosts that show binding constants of $\log K_{\rm a} = 5$ or larger remains a major challenge. Aida's bisporphyrin macrocycles—in which porphyrin units are connected via flexible alkyl spacers—constitute perhaps the most successful family of hosts for fullerenes

to date. 3c,3s,3v,9a,9b,9f For example, the Zn(II) metalloporphyrin congener shows a $\log K_{\rm a}=5.8$ toward C₆₀, while the free base shows $\log K_{\rm a}=5.9$, both in benzene at room temperature. 9f The world-record in complex stability is the Ir(III) metalloporphyrin derivative which shows a $\log K_{\rm a}=8.1$ for C₆₀ in 1,2-dichlorobenzene (o-DCB) at room temperature. 3s It is noteworthy that in the latter case, the authors indicate that the supramolecular nature of the host—guest system is arguable, since iridium was found to bind a 6,6 junction of the fullerene in a η^2 fashion. Recently, a macrocycle featuring three porphyrins 10 and a "nanobarrel" formed by four porphyrin units 11 have been shown to bind C₆₀ with $\log K_{\rm a}=6.2$ and 5.7, respectively, both in toluene at room temperature.

2. DESIGN AND SYNTHESIS

We have very recently reported a macrocyclic receptor that associates C_{60} with $\log K_{\rm a} = 6.5 \pm 0.5$ in PhCl at room temperature (*p*-xylmac12 in Chart 1). The design of the macrocyclic host was based on our initial tweezers-like receptors, in which two exTTF units were connected through an isophthalic diester spacer. This simple receptor served as proof-of-principle for the utilization of exTTF as recognizing motif in the construction of hosts for fullerenes, and showed respectable affinities toward C_{60}

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Chart 1. Chemical structures of the nine receptors

Scheme 1. General Scheme for the Synthesis of the Macrocyclic Hosts^a

^a Conditions: (a) K_2CO_3 , NaI (cat.), DMF, reflux; (b) for *p*-xylmac and *m*-xylmac families, α,α' -*p*-dibromoxylene, or α,α' -*m*-dibromoxylene, K_2CO_3 , NaI (cat.), DMF, 60 °C; (c) for naphmac family, 2,6-bis(hydroxymethyl)naphthalene, Ph₃P, diethyl azodicarboxylate, THF, reflux; (d) dimethyl 1,3-dithiol-2-ylphosphonate, BuLi, THF, -78 °C to room temp; (e) Grubb's first generation catalyst, CH₂Cl₂, room temp.

considering its lack of preorganization (log $K_a = 3.5$ in PhCl at room temperature). To increase the degree of preorganization,

we conserved all the basic traits of the exTTF tweezers, but included an alkyl linker with terminal alkenes to achieve macrocyclization through RCM. In the present study, we describe a collection of macrocyclic hosts with systematic structural modification of both the alkyl and aromatic spacers. Considering the precedents and preliminary molecular modeling, we decided to investigate *p*-xylene, *m*-xylene and 2,6-dimethylnaphthalene as aromatic spacers, and hexene, heptene and octene as alkyl spacers and precursors for RCM. The structures of the receptors synthesized are shown in Chart 1. The nomenclature utilized identifies the aromatic spacer and the length of the alkenyl spacers.

The *p*-xylmac and *m*-xylmac families (n = 1-3) were synthesized following a procedure analogous to that reported for pxylmac12 (Scheme 1).¹² The synthesis starts with the Williamson etherification of anthraflavic acid with the corresponding bromoalkene under standard conditions, followed by dialkylation of either para or meta- α , α' -dibromoxylene with the remaining phenolic OH, which yielded linear tetraketone precursors. These were subjected to Horner-Wadsworth-Emmons olefination with dimethyl 1,3dithiol-2-ylphosphonate, to afford the bis-exTTF linear precursors, which were finally treated with Grubb's first generation catalyst to obtain the desired macrocycles. The synthesis of the naphmac family follows the same procedure, but utilizing 2,6bis(hydroxymethyl)naphthalene and a Mitsunobu protocol to obtain linear bis-anthraquinone precursors (Scheme 1). This was followed by olefination to form the exTTF derivatives and RCM to yield the macrocycles as a chromatographically inseparable mixture of E/Z isomers, which was used as such. The identity and purity of the nine macrocyclic hosts and all synthetic intermediates were unambiguously established by standard spectroscopic and analytical techniques (see the Supporting Information for full experimental details and characterization).

3. ASSOCIATION OF C₆₀ AND C₇₀

Binding constants were estimated through UV—vis titrations. In a typical experiment, we prepared stock solutions of the macrocyclic receptor in PhCl at a concentration approximately equal to the reciprocal of the binding constant. Those solutions were utilized as solvents in the solution of fullerene, to ensure working at constant concentration of host. Aliquots of the fullerene solution were

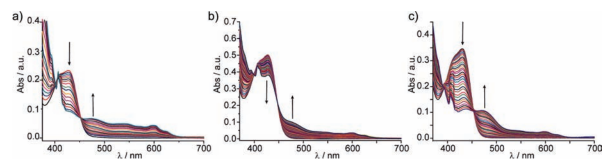


Figure 1. Spectral changes in UV-vis titration experiments of (a) p-xylmac12, (b) m-xylmac12, and (c) naphmac12 against C₆₀ in PhCl at 298 K.

Table 1. Logarithm of the Calculated Binding Constants for the Macrocyclic Hosts towards C_{60} and C_{70} , as Estimated from Three Separate UV—vis Titration Experiments in PhCl at 298 K

	C ₆₀	C ₇₀
p-xylmac10	4.3 ± 0.5	6.0 ± 0.4 ; 12.3 ± 0.6^a
p-xylmac12	6.5 ± 0.5	ь
p-xylmac14	3.5 ± 0.6	5.9 ± 0.3
m-xylmac10	4.0 ± 0.5	4.4 ± 0.4 ; 8.3 ± 0.5^a
m-xylmac12	4.8 ± 0.4	ь
m-xylmac14	4.1 ± 0.4	5.4 ± 0.3
naphmac10	4.0 ± 0.2	4.7 ± 0.3
naphmac12	5.6 ± 0.6	5.6 ± 0.4
naphmac14	3.4 ± 0.1	6.1 ± 0.2

^a First binding constant corresponds to log $K_{1:1}$ and second to log $K_{2:1}$. ^b Titration data did not provide satisfactory fits (see main text).

added to the host solution until reaching 3—4 mol equiv, and UV—vis spectra recorded at room temperature on a Cary UV-50 or on a Shimadzu UV-3600 spectrophotometer equipped with a temperature control unit. Analysis of the UV—vis data to determine binding constants was carried out utilizing Specfit multivariable analysis software. Experiments were repeated three times. The values of the binding constants are reported in Table 1 and were calculated as the mean value of the three experiments, with the error estimated as the standard deviation.

The changes in the absorption spectra during the titrations against C_{60} are very similar for all nine receptors. As representative examples, Figure 1 shows the evolution of the spectra of the best host of each series, p-xylmac12, m-xylmac12, and naphmac12 upon increasing C₆₀ concentration. In all cases, we observe the decrease in absorption of the band at $\lambda_{\rm max} \approx 425$ 430 nm, and the appearance of a charge-transfer band at $\lambda_{\rm max}$ \approx 470—475 nm. An isosbestic point at around 450 nm is also clearly observable. These spectral changes distinctively indicate association of C₆₀ inside the cavity of the macrocycle. 12 The relative intensity of the changes correlates very well with the calculated binding constants (see Figure 1). For p-xylmac12 (log $K_a = 6.5$ \pm 0.5) and **naphmac12** (log K_a = 5.6 \pm 0.6) both the decrease in the absorption of the exTTF and the increase in absorption of the charge-transfer band are significantly more marked than for m-xylmac12 (log $K_a = 4.8 \pm 0.4$).

The titration data versus C_{60} for all nine macrocycles were fitted successfully to a 1:1 binding mode, ¹³ yielding the binding constants summarized in Table 1. From the values of the binding constants, it is clearly apparent that the *m*-xylylene spacer is the least adequate of the aromatics, with *p*-xylylene producing the

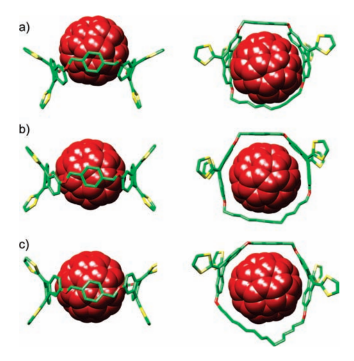


Figure 2. Side and top views of the energy-minimized (AMBER) models of the associates: (a) p-xylmac10·C₆₀; (b) p-xylmac12·C₆₀; (c) p-xylmac14·C₆₀. The macrocycles are depicted in stick representation with carbon atoms in green, sulfurs in yellow, and oxygens in red. Hydrogen atoms are not shown for clarity. [60] Fullerenes are depicted as space-filling models and colored in dark red.

best host for C_{60} , and naphthalene in an intermediate position. With regards to the alkyl spacer, the C_{12} derivative consistently shows the highest binding constant, with a very significant increase compared to C_{10} (too small) and C_{14} (too large). These results are qualitatively well reproduced by molecular mechanics calculations in the gas phase. ¹⁴ Figure 2 shows the geometry optimized models for the *p*-xylmac series to illustrate this point.

In p-xylmac10 · C₆₀ (Figure 2a) the macrocyclic cavity is too small to associate C₆₀, and the host adopts a bowl-type conformation, accommodating the fullerene above the plane of the macrocycle. In contrast, in p-xylmac12 · C₆₀ (Figure 2b) C₆₀ fits perfectly inside the cavity of the macrocycle, with both exTTF units, the aromatic linker, and the alkyl spacer closely wrapping the fullerene unit. Finally, C₆₀ fits loosely in p-xylmac14, so only part of the alkyl chain can approach the [60] fullerene and provide stabilizing van der Waals interactions.

The evolution of the spectra during the titrations against C_{70} is much less uniform across the different families of macrocycles

attending to the aromatic spacer. The spectral changes in the **naphmac** series mirror what we have previously observed for exTTF-cyclotriveratrylene hosts, ^{3h} namely, during the first additions the absorbance at $\lambda_{\text{max}} = 430$ nm decreases, with the concomitant increase of a charge-transfer band centered at $\lambda = 471$ nm. As the concentration of C_{70} increases the spectral changes become difficult to trace due to spectral overlap. Figure 3 depicts the experimental results obtained in the titration of **naphmac14** vs C_{70} (PhCl, 298 K). All three titrations for **naphmac10**, **naphmac12**, and **naphmac14** fitted successfully to a 1:1 binding model (see Figure 3 inset), yielding larger binding constants as the size of the macrocyclic cavity increases (see Table 1), as could be expected. Thus, the most efficient binder for C_{70} was found to be **naphmac14**, with a remarkable binding constant of log $K_a = 6.1 \pm 0.2$.

On the other hand, the *p*-xylmac and *m*-xylmac families showed parallel trends, depending on the length of the alkyl spacer. The larger members of each series, *p*-xylmac14 and *m*-xylmac14 behaved similarly to the naphmac hosts, and analysis of their titration data was performed satisfactorily utilizing a 1:1 binding model, yielding binding constants of $\log K_a = 5.9 \pm 0.3$ and 5.4 ± 0.3 , for *p*-xylmac14 and *m*-xylmac14, respectively.

Unexpectedly, the smaller macrocycles *p*-xylmac10 and *m*-xylmac10 showed singular spectroscopic changes during the titration experiments. Figure 4a shows the variations in the electronic absorption spectrum of *p*-xylmac10 upon additions of 0.1 equiv aliquots of C₇₀ in PhCl at 298 K. Although at first sight the general trends are similar to those we observed for naphmac hosts (see Figure 3), a closer look at the titration data

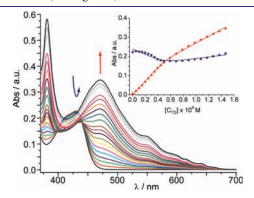


Figure 3. UV—vis spectra as recorded during the titration of naph-mac14 vs C_{70} (PhCl, 298 K). Inset shows the binding isotherms at 430 nm (blue squares) and 471 nm (red triangles); solid lines represent the fits. For this particular experiment a log $K_a = 6.0$ was calculated.

offers major and informative differences. During the first three additions of the titration, where the concentration of the macrocycle is significantly larger than that of the [70] fullerene, two very well-defined isosbestic points at $\lambda = 402$ and 432 nm can be noted (Figure 4b, inset). As the concentration of C₇₀ increases these isosbestic points are lost, indicating the coexistence of more than one type of associates in solution. All these features point to the formation of complexes of both 2:1 and 1:1 host/guest stoichiometry, with the former prevailing in the early stages of the titration, and complexes of both stoichiometries being present in solution when the concentration of C₇₀ increases sufficiently. Indeed, the titration data did not provide satisfactory fits when assuming either a 1:1 or a 2:1 stoichiometry exclusively. On the contrary, the multivariable analysis software converged nicely when we utilized a model where both types of associates are presumed to exist in solution (Figure 4a, inset). The calculated binding constants for *p*-xylmac10 were log $K_{1:1}$ = 6.0 \pm 0.4 and log $K_{2:1} = 12.3 \pm 0.6$, in agreement with both types of associates showing similar stability. Analogously, we estimated $\log K_{1.1} =$ 4.4 ± 0.4 and $\log K_{2:1} = 8.3 \pm 0.5$ for *m*-xylmac10, which also shows smaller binding constants toward C₇₀, as was the case with C₆₀. Conclusive evidence for this model was provided by Job's plot analysis, which showed a broad maximum centered at a molar fraction of 0.3-0.45 (Figure 4c), in an intermediate situation between 2:1 and 1:1, where the maxima would be located at 0.33 and 0.5, respectively.

Finally, *p*-xylmac12 and *m*-xylmac12 are apparently left in an intermediate situation between the C_{10} and C_{14} members, and, although association of C_{70} seems to take place considering the spectral changes, we could not achieve correct fitting of their titration data with any binding model. For instance, for *m*-xylmac12 the best fit was obtained with a 1:1 model, and yielded a clearly overestimated binding constant of log $K_a = 9.9 \pm 2.6$, with excessive error and less than modest fitting for the binding isotherms (Figure 5). Similarly, the best fit for the data corresponding to *p*-xylmac12 vs C_{70} affords log $K_a = 8.4 \pm 2.5$.

As with C_{60} , the experimental results can be rationalized in terms of a simple lock and key model through molecular mechanics calculations. The energy-minimized models of p-xylmac10 \cdot C_{70} and naphmac10 \cdot C_{70} are shown in Figure 6 structures a and c. The cavity of p-xylmac10 is too small to accommodate C_{70} , and leaves a significant part of its ellipsoidal surface exposed, so that it can be bound by another molecule of the host (Figure 6b) without steric congestion. On the other hand, the slight increase in size provided by the naphthalene spacer in naphmac10 is sufficient to allow C_{70} to penetrate deeper into its cavity, stabilizing the 1:1 associate

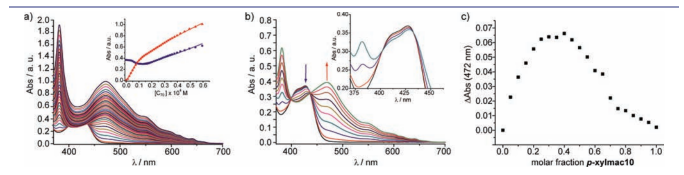


Figure 4. (a) UV—vis spectra as recorded during the titration of p-xylmac10 vs C_{70} (PhCl, 298 K). Inset shows the binding isotherms at 430 nm (blue squares) and 472 nm (red triangles), solid lines represent the fits. (b) First 10 additions, up to 1 equiv of C_{70} . Inset shows the first three additions. (c) Job's plot ([p-xylmac10] + [C_{70}] = 1.0×10^{-4} M).

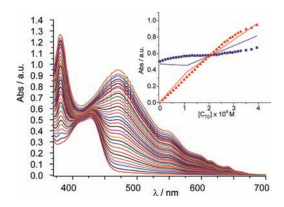


Figure 5. UV—vis spectra as recorded during the titration of *p*-xylmac12 vs C₇₀ (PhCl, 298 K). Inset shows the binding isotherms at 430 nm (blue squares) and 471 nm (red triangles); solid lines represent the fits.

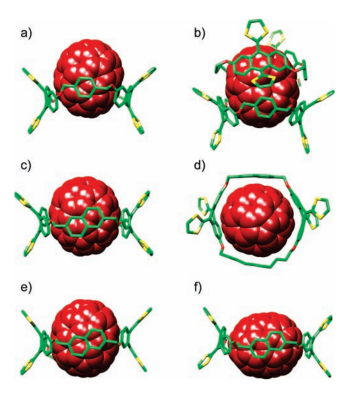


Figure 6. Energy-minimized (AMBER) models of the associates: (a) p-xylmac10· C_{70} ; (b) p-xylmac10· C_{70} ·p-xylmac10; (c) naphmac10· C_{70} , side view and (d) top view; (e) naphmac14· C_{70} with the guest perpendicular to the plane of the macrocycle; (f) naphmac14· C_{70} with the guest parallel to the plane of the macrocycle. The macrocycles are depicted in stick representation with carbon atoms in green, sulfurs in yellow, and oxygens in red. Hydrogen atoms are not shown for clarity. [70] Fullerenes are depicted as space-filling models and colored in dark red.

(Figure 6c,d). Meanwhile, **naphmac14** is the only macrocycle which is large enough to be capable of hosting C_{70} with its longer axis parallel to the plane of the macrocycle (Figure 6f). This provides an alternative to the perpendicular binding we observe for all other macrocycles, and can be responsible for the increase in stability.

To complete our understanding of the recognition process of these macrocyclic systems and gain an insight into its thermodynamic nature, we carried out titrations at different temperatures (288, 298, 308, and 318 K) in PhCl. As model system, we

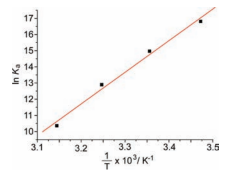


Figure 7. Plot of the natural logarithm of K_a vs 1/T for the association of C_{60} with *p*-xylmac12 in PhCl.

chose the best binder of the family: p-xylmac12 and C_{60} as a host. A plot of the natural logarithm of K_a vs 1/T is shown in Figure 7. The data show a clear linear tendency and was fitted to a straight line ($r^2 = 0.995$) with a slope of 19557 ± 1416 and an intercept of -51 ± 5 . From these data, the standard enthalpy of association was calculated to be $\Delta H^\circ = -163 \pm 12$ kJ mol $^{-1}$ (-38 ± 3 kcal mol $^{-1}$) and $\Delta S^\circ = -424 \pm 42$ J mol $^{-1}$ (-101 ± 10 cal mol $^{-1}$). Thus, the association of fullerenes by our exTTF-based macrocyclic hosts is driven by strong enthalpic interactions. For comparison, the enthalpy value is approximately two to three times higher than that calculated for the association of C_{60} by calix[4]naphthalenes in toluene, which show a log $K_a = 2.8$.

The appearance of a charge-transfer band during the titration suggests that electronic interactions between the fullerenes and our bis-exTTF macrocycles take place. These interactions in the ground state are also noticeable in cyclic voltammetry (CV) experiments. Though adsorption phenomena prevent a reliable measure of the oxidation potential of exTTF units, the cathodic part of the CV allowed us to draw interesting conclusions. Figure 8a shows the CVs of p-xylmac12 (black), C₆₀ (blue), and a mixture of both compounds (red). In the presence of a macrocycle, the half-wave potentials associated with the reduction signals of [60] fullerene are clearly shifted, which definitely confirms an interaction in the ground state between the host and the guest, even though both positive and negative shifts were detected. Similar measurements (Figure 8b), performed with **naphmac14** and C_{70} , indicate a strong electronic communication in the complex with a half-wave potential shift as high as -150mV for the fourth reduction of C_{70} .

To further investigate the influence of the electronic communication on the stability of the complexes, we carried out titrations in solvents of different polarities. In the more polar benzonitrile ($\varepsilon_{\rm r}$ = 25.2), where charge-transfer interactions should be favored, we calculated a binding constant of $\log K_a =$ 7.5 \pm 0.9. The increase in binding constant cannot be ascribed solely to stabilization of the complex through enhanced chargetransfer interactions, since the solubility of C₆₀ in benzonitrile is 0.4 mg/mL. This is conclusively confirmed in the nonpolar toluene ($\varepsilon_{\rm r}$ = 2.4), in which a decrease in binding constant to log $K_{\rm a}$ = 5.4 \pm 0.9 was found. Considering the decrease in solubility of C_{60} when moving from chlorobenzene (6.4 mg/mL) to toluene (2.4 mg/mL)¹⁶ this is evidence of the contribution of the electronic interactions to the stabilization of the *p*-xylmac12. C₆₀ associate, since a decrease in binding constant with increasing solubility is typically found.^{3u} Such effect can be attributed to the solvation penalty associated with the formation of the

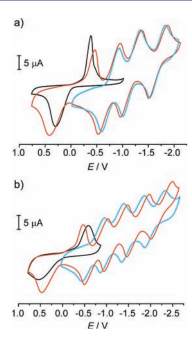


Figure 8. Cyclovoltammograms of (a) *p*-xylmac12 (black), C_{60} (blue), and *p*-xylmac12 $\cdot C_{60}$ (red); (b) naphmac14 (black), C_{70} (blue), and naphmac14 $\cdot C_{70}$ (red), in PhCl, Bu₄NClO₄ (0.1 M), $\nu = 0.1 \text{ V s}^{-1}$, [fullerene] = [macrocycle] = 5×10^{-4} M. Potentials are indicated vs the Ag/AgNO₃ redox couple.

supramolecular assembly. Since a significant portion of the molecular surfaces of both the receptor and the guest becomes unavailable for solvation upon complexation, the solvation penalty is more severe in solvents which solvate the separated entities more efficiently.

4. CONCLUSIONS

We have synthesized a collection of nine macrocycles bearing two units of exTTF as recognition elements for the fullerenes, connected through three different aromatic and alkyl spacers, and investigated their binding abilities toward C_{60} and C_{70} . It was found that *p*-xylmac12 is the best binder for C_{60} with log K_a = 6.5 \pm 0.5 for C_{60} in PhCl and log K_a = 7.5 ± 0.9 in benzonitrile. These values make *p*-xylmac12 one of the most efficient receptors for C₆₀ reported to date. With regards to C₇₀, naphmac14 and p-xylmac14 show log $K_a = 6.1 \pm$ 0.2 and 5.9 \pm 0.3, respectively, in PhCl at 298 K. These are two of the very few examples of hosts capable of associating C_{70} with micromolar affinity. Then the general point of view of the design of hosts for fullerenes, our results show that relatively small structural variations in the receptor lead to large changes in binding affinities, and in, some cases, even stoichiometry. For example, while m-xylmac14 forms 1:1 complexes with C_{70} with a respectable binding constant of log K_a = 5.4 ± 0.3 , the smaller member *m*-xylmac10 forms associates of both 1:1 and 2:1 stoichiometries, with log $K_{1:1}$ = 4.4 \pm 0.4 and $\log K_{2:1}$ 8.3 \pm 0.5. With regards to C₆₀, an impressive change in binding constant of nearly 3 orders of magnitude is found when moving from *p*-xylmac12 (log $K_a = 6.5 \pm 0.5$) to *p*-xylmac14 $(\log K_a = 3.5 \pm 0.6)$. These differences underline the singularities of fullerenes as hosts in molecular recognition and highlight the importance of fine-tuning the structure of the receptors. Remarkably, all our experimental results can be qualitatively explained with

a simple lock and key model through undemanding molecular mechanics calculations.

To shed light on the driving force for binding, we have determined the thermodynamic parameters for the formation of p-xylmac12· C_{60} . To that end, we have carried out titrations at different temperatures (318, 308, 298, and 288 K). The positive slope of the ln K_a vs 1/T indicates that association is driven by enthalpy. Indeed, we calculated a remarkably high $\Delta H^\circ = -163 \pm 12$ kJ mol $^{-1}$ (-38 ± 3 kcal mol $^{-1}$). The spectral features observed during the UV-vis titrations, cyclic voltammetry data, and titrations carried out in solvents of different polarity suggest that electronic interactions between the exTTF and C_{60} contribute significantly to the stabilization of the complex.

In the near future, we plan to exploit the extreme stability of the bis-exTTF macrocycle—fullerene complexes, the lessons extracted from the structure—affinity relationships, and the relatively simple synthetic access to the macrocycles to construct chiral hosts for the enantioselective molecular recognition of the inherently chiral higher fullerenes. ^{3g,3v}

ASSOCIATED CONTENT

Supporting Information. Supplementary figures, characterization and full experimental procedures. This material is available free of charge via the Internet at http://pubs.acs.org.

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