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Molecular baskets form inclusion complexes with phenethylamine drugs in water†

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Molecular basket 1^{6-} comprising a nonpolar cavity and an anionic nest of six carboxylates at its rim was found to form inclusion complexes with (1*R*, 2*S*)-ephedrine, (1*R*, 2*R*)-pseudoephedrine, and (1*S*, 2*R*)-tranilcypromine. Experimental results (NMR) and theory (MM/DFT) suggest the basket encapsulates phenethylamines in unique and predictable fashion.

Substituted phenethylamines (Fig. 1A) are organic molecules with the 2-phenylethan-1-amine framework and a broad range of biological activities. To mention a few, phenethylamines include stimulants (*e.g.*, methamphetamine), hallucinogens (*e.g.*, mescaline), entactogens (*e.g.*, MDMA), vasopressors (*e.g.*, ephedrine), decongestants (*e.g.*, pseudoephedrine), bronchodilators (*e.g.*, albuterol), and antidepressants (*e.g.*, tranilcypromine).¹ With facile access to a great number of phenethylamines,² there has been a considerable interest toward derivatizing their framework and examining potentially novel pharmacological effects.³ In this vein, development of more potent but also cheap hallucinogens and stimulants⁴ is having a negative impact on society by expanding the drug epidemic and illicit activities.⁵ Accordingly, supramolecular chemists have used macrocyclic hosts (*i.e.*, cavitands) for examining facile detection,⁶ resolution,^{1e,7} and sequestration⁸ of phenethylamine drugs. The rationale behind these studies is to use functionalized but also conformationally restricted cavitands for complementing ammonium and/or phenyl units within targeted compound. In the cases of resorcin[4]arenes^{1e} and cucurbit[6]urils,^{1d,6c,9} the formation of perching complexes ensued in polar solvents and/or solid state with cavitands' polar rim holding onto the ammonium site of phenethylamines *via* hydrogen bonding (Fig. 1B). With cucurbit[7]urils,^{6a} calix[4]arenes^{8b} β , γ -cyclodextrins,¹⁰ pillar[5]arenes,¹¹ and pillar[6]arenes,^{8a} the phenyl group was found to, in

aqueous media, insert into each host's nonpolar cavity thereby placing the ammonium at the polar rim to form hydrogen bonds (Fig. 1B). We wondered, would molecular basket 1^{6-} (Fig. 1C), comprising a nonpolar cavity and an anionic nest of six carboxylates at the rim (*i.e.*, three glutamic acid residues),¹² form an inclusion complex with positively charged phenethylamines in water? In this regard, easily accessible and enantiopure (1*R*, 2*S*)-ephedrine **2**, (1*R*, 2*R*)-pseudoephedrine **3**, and (1*S*, 2*R*)-tranilcypromine **4** (Fig. 1A) seemed suitable for the study. While these drugs appeared to have structural complementarity to 1^{6-} (Fig. 1C), we were curious about the exact mode of interaction (*i.e.*, stoichiometry and docking geometry)¹³ as well as the stability of these complexes. In fact, host 1^{6-} was previously shown to form inclusion complexes with organophosphorus agents (OPs)¹⁴ and α,ω -diammonium alkanes^{13,15} with millimolar (or even greater) stabilities. While nonpolar P-CH₃/P-OCH₃ groups from OPs occupy the inner space of 1^{6-} , polar P=O units reside at the rim of binary [OPs $\subset$ 1]⁶⁻. In the case of diammonium alkanes, two RNH₃⁺ groups would form salt bridges with α - and γ -carboxylates from two molecules of 1^{6-} surrounding the guest to give a ternary complex.¹⁵

Molecular basket **1** was obtained by following an earlier described synthetic protocol.¹² This molecule can be dissolved in aqueous 30 mM phosphate buffer solution (PBS) at pH = 7.0 to give hexaanionic 1^{6-} . At concentrations of 5.0 mM or lower, amphiphilic 1^{6-} was shown to be monomeric with results from DLS, UV-Vis, MS, and NMR experiments all suggesting the absence of aggregation.¹² Importantly, ¹H NMR spectrum of 1^{6-} (Fig. 3A) comprises a set of sharp signals corresponding to, on average, a C₃ symmetric compound. A Monte Carlo conformational search (OPLS3) in implicit water solvent showed the presence of two conformers **1_{A/B}**⁶⁻ within 1 kcal mol⁻¹ (Fig. 2; Table S3, ESI†); each conformer was additionally optimized at a higher level of theory (DFT:B3LYP/6-31+G*). While **1_A**⁶⁻ has all three α -carboxylates from glutamic acid residues pointing to the concave side of the host, **1_B**⁶⁻ has one of these carboxylates at the convex face. Along with computational results, we reasoned that a rapid interconversion of **1_A**⁶⁻ to **1_B**⁶⁻ (Fig. 2), in

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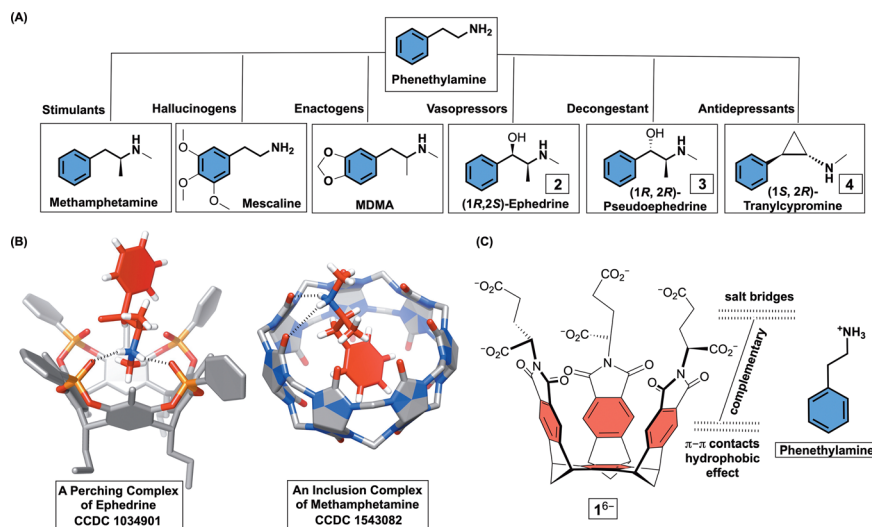


Fig. 1 (A) Chemical structures of selected phenethylamine drugs and their biological functions. (B) Line representations of X-ray structures of a derivative of resorcin[4]arene holding ephedrine at its top (CCDC 1034901) and cucurbit[7]uril forming an inclusion complex with methamphetamine (CCDC 1543082). (C) Chemical structures of basket 16^- and phenethylammonium cation showing their structural and electronic complementarity.

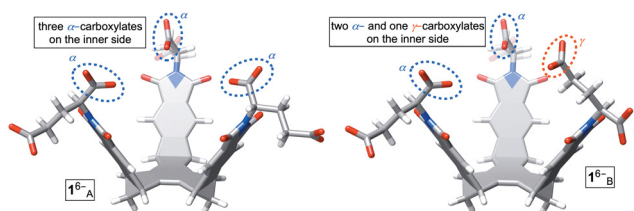


Fig. 2 Stick representations of two most stable conformers (Monte Carlo, OPLS3) of basket 16^- . Conformer $1A6^-$ (top) and $1B6^-$ (bottom) are energy optimized using density functional theory (B3LYP/6-31+G*).

addition to each molecule's conformational dynamics, contributed to the observed ^1H NMR spectrum of 16^- .

An incremental addition of a standard solution of phenethylamines 2^+ – 4^+ to 16^- was monitored (30 mM PBS at pH = 7.0) with ^1H NMR spectroscopy (Fig. 3 and Fig. S1–S3, ESI †); each titration was repeated three times. Importantly, additions prompted a steady change (*i.e.*, magnetic shielding) of the basket's resonances (Fig. 3A). We reasoned that noncovalent complexation ought to be taking place with the host residing in the shielding region of aromatic guests (Fig. 3B). To determine the stoichiometry of the complexation, the change in chemical shift of seven resonances from 16^- as a function of the concentration of 2^+ – 4^+ was subjected to nonlinear regression analyses using 1:1, 1:2 and 2:1 binding models (Fig. S4–S6, ESI †).¹⁶ In this regard, it is important to note that ^1H NMR

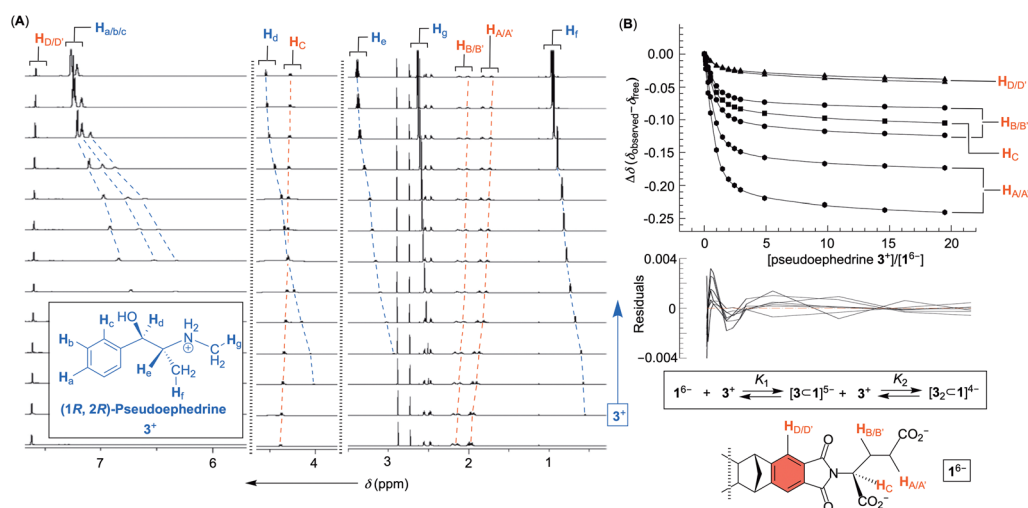


Fig. 3 (A) ^1H NMR spectra of basket 16^- (30 mM PBS at pH = 7.0) obtained after incremental addition of a standard solution of (1R, 2R)-pseudoephedrine 3^+ (up to 20 molar equivalents). (B) A plot showing normalized chemical shifts of ^1H NMR resonances from basket 16^- ($\Delta\delta$) as a function of the quantity of (1R, 2R)-pseudoephedrine 3^+ added to its aqueous solution. The data were fit to 1:2 binding model (supramolecular.org) with a distribution of residuals shown below.



resonances of drugs 2^+-4^+ remained constant as a function of their variable concentrations (Fig. S7–S9, ESI†) so no competing dimerization or aggregation of drug molecules took place during supramolecular titrations. The curve fitting result for the titration of pseudoephedrine 3^+ to 1^{6-} using 1:2 binding model (*i.e.*, $[3\subset 1]^{5-}$ and $[3_2\subset 1]^{4-}$ complexes) is shown in Fig. 3B. The model is accepted given a somewhat random distribution of residuals¹⁷ and the small covariance of the fit (*i.e.*, $cov_{fit} = 3 \times 10^{-4}$; Fig. S5, ESI†).^{18,19} When the same criteria were used for evaluating the goodness of fit (GOF) for 1:1 and 2:1 binding models, we could disqualify both 1:1 ($cov_{fit} = 3 \times 10^{-3}$) and 2:1 complexations ($cov_{fit} = 6 \times 10^{-4}$).¹⁹ For all three drugs, the covariance of the fit slightly favoured 1:2 over 2:1 models of association (Fig. S4–S6, ESI†). Additionally, we decided to examine Akaike's information criterion (AIC).²⁰ This information-theoretic approach was recently introduced²⁰ to the field of host-guest chemistry as a superior alternative for evaluating the GOF. To calculate AIC, one needs to use the formula $AIC = N \ln(SSR/N) + 2k$ in which N is the number of fitted points, SSR is the sum of the squared residuals and k is the number of fitted parameters. The smaller the AIC the better the GOF. After computing AIC values for three independent titrations of each drug to 1^{6-} (Table 1 and Tables S1 and S2, ESI†), in addition to Akaike weights w_i , we compared the data. First, 1:1 complexations are uniformly showing the least favorable AIC scores (*i.e.*, the most positive numbers). Between the other two models, 1:2 gave lower AIC values than 2:1 in two out of three titrations for each drug. We conclude that 1:2 association ought to be dominating under the experimental conditions. To corroborate statistical considerations of the fit, ESI† mass spectrometry measurements of a mixture of phenethylamine drugs 2^+-4^+ with basket 1^{6-} were taken in 1:10, 1:1, and 10:1 ratios (Fig. S10–S12, ESI†). At all ratios, only the presence of 1:1 and 1:2 complexes were observed. While the electrostatic repulsion between two anionic 1^{6-} disfavors 2:1 complexation, we hypothesize that 1:2 complex comprises the first guest occupying the cavity of 1^{6-} while the second one undergoes an exo-complexation.

After elucidating the binding stoichiometry, we went on to analyse the thermodynamic stability of binary $[drug\subset 1]^{5-}$ and ternary $[drug_2\subset 1]^{4-}$ complexes (Table 2; see also Fig. S4–S6, ESI†). Evidently, the formation of binary complexes dominates each equilibria with $K_1 \gg K_2$. This bodes well with the earlier notion that $[drug\subset 1]^{5-}$ is an inclusion and more stable

Table 2 Experimental binding constants (K_1 and K_2) for the complexation of phenethylamine drugs $2-4^+$ to 1^{6-} and describing the formation of binary $[2-4\subset 1]^{5-}$ and ternary $[(2-4)_2\subset 1]^{4-}$ structures. From three independent titrations, each number is reported as mean $\pm$ standard deviation (Fig. S4–S6)

Phenethylamine drug	K_1 (M^{-1})	K_2 (M^{-1})
(1 <i>R</i> , 2 <i>S</i>)-ephedrine 2^+	$7 \pm 1 \times 10^3$	54 ± 11
(1 <i>R</i> , 2 <i>R</i>)-pseudoephedrine 3^+	$6 \pm 2 \times 10^3$	58 ± 11
(1 <i>S</i> , 2 <i>R</i>)-tranilcypromine 4^+	$2 \pm 1 \times 10^4$	39 ± 14

complex than $[drug_2\subset 1]^{4-}$ in which the association takes place on the basket's outer side. The stability of 1:1 complexes are in the millimolar range, but interestingly the affinity of tranilcypromine 4^+ toward 1^{6-} is an order of magnitude greater than ephedrine 2^+ and pseudoephedrine 3^+ .

After quantifying the thermodynamic parameters characterizing the binding, we sought out to learn more about the mode of complexation and intermolecular interactions driving the association. First, the magnetic perturbation of resonances from each drug undergoing complexation show phenyls' $H_{a/b/c}$ experiencing the largest magnetic shielding ($\Delta\delta = \delta_{bound} - \delta_{free}$, Fig. 4A; Fig. S1–S3, ESI†). Inclusion of the phenyl ring into the basket's cavity with H_a reaching for the cavity's bottom would explain the observation.¹³ Along with such docking of the aromatic, the ammonium site from $2-4^+$ ought to reside in the anionic nest (Fig. 1C). If so, what is the exact role of α - and γ -carboxylates in the complexation and what is the dominant pose of each guest with respect to the basket host? To answer these questions, we developed a computational protocol to estimate a pose which best represents the observed NMR shielding effect. First, the magnetic environments of 1_A^{6-} and 1_B^{6-} were mapped by computing nucleus independent chemical shift values (NICS) for a grid around the molecules (Fig. S14 and S15, ESI†).²¹ Next, we ran Monte-Carlo conformational searches (OPLS3) for inclusion complexes $[2-4\subset 1_A]^{5-}$ and $[2-4\subset 1_B]^{5-}$. During MC searches, we froze conformational motions of $1_{A/B}^{6-}$ but allowed guests to change their position and conformation, akin to docking protocols. Finally, we algorithmically assigned shielding values ($\Delta\delta_{comp}$) for proton nuclei of all MC conformers of $[2-4\subset 1_A]^{5-}$ and $[2-4\subset 1_B]^{5-}$ using the NICS maps from 1_A^{6-} and 1_B^{6-} . Having computed ($\Delta\delta_{comp}$) and experimental ($\Delta\delta_{exp}$) values of proton shifts, we analysed the data in two different ways. For the first approach, we

Table 1 Akaike information criterion (AIC) scores and Akaike weights (w_i) obtained for three independent titrations in which (1*R*, 2*R*)-pseudoephedrine 3^+ was added to basket 1^{6-} (Fig. 3A and Fig. S5). The data were fit to three models (1:1, 1:2, and 2:1) with AIC and w_i obtained using formulas from ref. 20

		Model 1:1	Model 1:2	Model 2:1
Titration I	AIC	−1039.7	−1192.6	−1113.3
	w_i	4×10^{-67}	1	2×10^{-26}
Titration II	AIC	−1108.8	−1238.1	−1232.5
	w_i	7×10^{-57}	1	4×10^{-3}
Titration III	AIC	−1541.3	−1619.2	−1978.5
	w_i	1×10^{-190}	9×10^{-157}	1

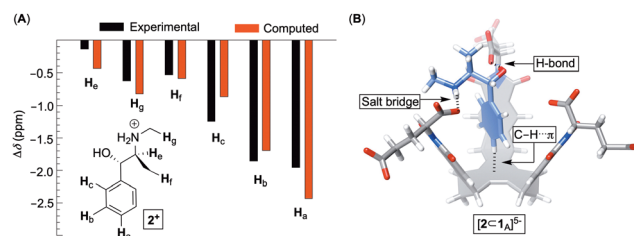


Fig. 4 (A) Experimental (black) and computed (red, for an ensemble of structures) chemical shift perturbations ($\Delta\delta$) of ephedrine resonances within $[2\subset 1_A]^{5-}$. (B) A stick representation of single conformer of $[2\subset 1_A]^{5-}$ having the lowest RMSE.



Table 3 Root mean squared error (RMSE) data for ensemble and single poses of $[2-4 \subset 1_A]^{5-}$ and $[2-4 \subset 1_B]^{5-}$ complexes

Phenethylamine drug	Type	1_A^{6-} (RMSE)	1_B^{6-} (RMSE)
(1R, 2S)-ephedrine 2^+	Ensemble	0.52	0.43
	Single	0.79	0.89
(1R, 2R)-pseudoephedrine 3^+	Ensemble	0.86	0.88
	Single	0.65	0.70
(1S, 2R)-tranilcypromine 4^+	Ensemble	0.50	0.78
	Single	0.95	1.0

determined root mean squared error

$$\left(\text{RMSE} = \sqrt{1/N \sum (\Delta\delta_{\text{comp}} - \Delta\delta_{\text{exp}})^2} \right) \text{ for each computed}$$

pose and the pose with the lowest RMSE was assumed to be the best representation of the experimental result (Table 3, Fig. 4B; see also Fig. S19–S21, ESI†). As for the second approach, we Boltzmann weighted $\Delta\delta_{\text{comp}}$ values for each proton from all computed poses and summed each poses contribution to the ensemble average $\langle\Delta\delta_{\text{comp}}\rangle$. Next, RMSEs for $[2-4 \subset 1_A]^{5-}$ and $[2-4 \subset 1_B]^{5-}$ were determined using the $\langle\Delta\delta_{\text{comp}}\rangle$ values (Table 3). From Table 3, one notes that the ensemble of poses is, in more cases than not, giving lower RMSEs than single structures. It follows that the ensemble is more effective than a single pose at describing the experimental result. Furthermore, the values for RMSEs from complexes of 1_A^{6-} and 1_B^{6-} are comparable to indicate that both conformers likely participate in the binding; however, we noted that 1_A^{6-} showed a closer agreement to experimental NMR shielding values (Fig. S16–S18, ESI†). For (1R, 2S)-ephedrine 2^+ occupying 1_A^{6-} , RMSE values for the ensemble of poses (0.52) and a single pose (0.79) are similar (Table 3).

With an assumption that this single pose presents an average picture of the ensemble, the structure of $[2 \subset 1_A]^{5-}$ in Fig. 4B shows ephedrine anchoring its benzene in the basket's cavity (via C–H $\cdots \pi$ contacts) while using hydroxyl and ammonium groups to form a hydrogen bond and salt bridge with α -carboxylates. Importantly, all of the best fit single poses of $2-4^+$ bound to $1_{A/B}^{6-}$ show the benzene from phenethylamines occupying the basket's cavity to contribute to non-classical hydrophobic effect²² with $\Delta H^\circ < 0$ (Fig. S13, ESI†) while OH and NH^+ groups hydrogen bond/salt bridge α - and to a smaller degree γ -carboxylates (Fig. S19–S21, ESI†).

In conclusion, molecular basket 1^{6-} forms inclusion complexes with phenethylamines in water. The results of both experimental and computational studies suggest that the probed drug molecules anchor their phenyl ring in the basket's cavity as driven by C–H $\cdots \pi$ contacts and the hydrophobic effect while forming hydrogen bonds and/or salt bridges with primarily α -carboxylates at the rim. With a well characterized mode of binding, the opportunity to install alternative amino acids or peptides atop our basket scaffold is clear. In addition to the investigation of structure function relationships, further

derivatization may indeed furnish a better agent for the preparation of chemosensors or sequestrers of phenethylamines.

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Conflicts of interest

There are no conflicts to declare.

Notes and references

- (a) M. D. Anglin, C. Burke, B. Perrochet, E. Stamper and S. Dawud-Noursi, *J. Psychoact. Drugs*, 2000, **32**, 137–141; (b) G. K. Aghajanian and G. J. Marek, *Neuropsychopharmacology*, 1999, **21**, 16S–23S; (c) D. J. Heal, J. Gosden, S. L. Smith and C. K. Atterwill, *Neuropharmacology*, 2023, **225**, 109375; (d) O. Danylyuk, *CrystEngComm*, 2018, **20**, 7642–7647; (e) E. Biavardi, F. Ugozzoli and C. Massera, *Chem. Commun.*, 2015, **51**, 3426–3429.
- S. Freeman and J. F. Alder, *Eur. J. Med. Chem.*, 2002, **37**, 527–539.
- D. E. Nichols and W. E. Fantegrossi, *Emerging designer drugs*, 2014, pp. 575–596.
- M. E. Nelson, S. M. Bryant and S. E. Aks, *Emerg. Med. Clin. North Am.*, 2014, **32**, 1–28.
- R. Gonzales, L. Mooney and R. A. Rawson, *Ann. Rev. Public Health*, 2010, **31**, 385–398.
- (a) Y. Jang, M. Jang, H. Kim, S. J. Lee, E. Jin, J. Y. Koo, I.-C. Hwang, Y. Kim, Y. H. Ko, I. Hwang, J. H. Oh and K. Kim, *Chemistry*, 2017, **3**, 641–651; (b) D. King, C.-L. Deng and L. Isaacs, *Tetrahedron*, 2023, **145**, 133607; (c) E. Biavardi, S. Federici, C. Tudisco, D. Menozzi, C. Massera, A. Sottini, G. G. Condorelli, P. Bergese and E. Dalcanele, *Angew. Chem., Int. Ed.*, 2014, **53**, 9183–9188.
- J. Vachon, S. Harthong, E. Jeanneau, C. Aronica, N. Vanthuyne, C. Roussel and J.-P. Dutasta, *Org. Biomol. Chem.*, 2011, **9**, 5086–5091.
- (a) A. T. Brockett, W. Xue, D. King, C.-L. Deng, C. Zhai, M. Shuster, S. Rastogi, V. Briken, M. R. Roesch and L. Isaacs, *Chemistry*, 2023, **9**, 881–900; (b) S. Ganapati, S. D. Grabitz, S. Murkli, F. Scheffenbichler, M. I. Rudolph, P. Y. Zavaliy, M. Eikermann and L. Isaacs, *ChemBioChem*, 2017, **18**, 1583–1588.
- O. Danylyuk, V. P. Fedin and V. Sashuk, *Chem. Commun.*, 2013, **49**, 1859–1861.
- M. T. Garcia-Valverde, M. L. Soriano, R. Lucena and S. Cardenas, *Anal. Chim. Acta*, 2020, **1126**, 133–143.
- R. Zhang, Y. Ren, Q. Zhang, W. Huang, H. Bai and X. Zeng, *New J. Chem.*, 2022, **46**, 20909–20917.
- S. E. Border, R. Z. Pavlovic, L. Zhiquan, M. J. Gunther, H. Wang, H. Cui and J. D. Badjic, *Chem. – Eur. J.*, 2019, **25**, 273–279.
- R. Z. Pavlovic, S. E. Border, T. J. Finnegan, L. Zhiquan, M. J. Gunther, E. Munoz, C. E. Moore, C. M. Hadad and J. D. Badjic, *J. Am. Chem. Soc.*, 2019, **141**, 16600–16604.
- S. E. Border, R. Z. Pavlovic, Z. Lei, M. J. Gunther, H. Wang, H. Cui and J. D. Badjic, *Chem. Commun.*, 2019, **55**, 1987.
- R. Z. Pavlovic, S. E. Border, Y. Li, X. Li and J. D. Badjic, *Chem. Commun.*, 2020, **56**, 2987–2990.
- P. Thordarson, *Supramolecular Chemistry: From Molecules to Nanomaterials*, 2012, vol. 2, pp. 239–274.
- (a) F. Ulatowski, K. Dabrowa, T. Balakier and J. Jurczak, *J. Org. Chem.*, 2016, **81**, 1746–1756; (b) D. Brynn Hibbert and P. Thordarson, *Chem. Commun.*, 2016, **52**, 12792–12805.
- J. E. A. Webb, M. J. Crossley, P. Turner and P. Thordarson, *J. Am. Chem. Soc.*, 2007, **129**, 7155–7162.
- P. Thordarson, *Chem. Soc. Rev.*, 2011, **40**, 1305–1323.
- K. Ikemoto, K. Takahashi, T. Ozawa and H. Isobe, *Angew. Chem., Int. Ed.*, 2023, **62**, e202219059.
- Z. Chen, C. S. Wannere, C. Corminboeuf, R. Puchta and P. V. R. Schleyer, *Chem. Rev.*, 2005, **105**, 3842–3888.
- L. Zhiquan, S. M. Polen, C. M. Hadad, T. V. RajanBabu and J. D. Badjic, *Org. Lett.*, 2017, **19**, 4932–4935.

