



HHS Public Access

Author manuscript

New J Chem. Author manuscript; available in PMC 2023 January 21.

Published in final edited form as:

New J Chem. 2022 January 21; 46(3): 995–1002. doi:10.1039/d1nj05209h.

Thermodynamics of Pillararene•Guest Complexation: Blinded Dataset for the SAMPL9 Challenge

Chun-Lin Deng, Ming Cheng, Peter Y. Zavalij, Lyle Isaacs

Department of Chemistry and Biochemistry, University of Maryland, College Park, Maryland 20742, USA.

Abstract

We report an investigation of the complexation between a water soluble pillararene host (WP6) and a panel of hydrophobic cationic guests (G1 – G20) by a combination of ^1H NMR spectroscopy and isothermal titration calorimetry in phosphate buffered saline. We find that WP6 forms 1:1 complexes with K_a values in the $10^4 - 10^9 \text{ M}^{-1}$ range driven by favorable enthalpic contributions. This thermodynamic dataset serves as blinded data for the SAMPL9 challenge.

Introduction

A crucial step in the drug development process is the discovery and optimization of small molecule ligands that bind to their target proteins in aqueous solution. Experimentally, this process is very expensive and time consuming because it requires an iterative process of chemical synthesis and the measurement of binding affinity.¹ Accordingly, the development of computational methods that successfully rank ligands by relative affinity and deliver binding free energies with errors below 1 kcal mol^{-1} are highly sought by the computational chemistry community and pharmaceutical industry. Validation is an important step in the development of such computational methods. However, testing of new methods on protein•ligand systems can be computationally expensive and time consuming because proteins are large and complex entities which require that extensive conformational sampling to ensure convergence. To address this issue, a group of computational chemists has organized a series of Statistical Assessment of the Modeling of Proteins and Ligands (SAMPL) challenges² to assess and improve the state-of-the-art. Over the years, SAMPL challenges relied upon unpublished blinded datasets including small molecule solvation free energies, HIV integrase inhibitors binding free energy, and pK_a and octanol-water partition coefficient predictions.³ Supramolecular chemists are also deeply involved in the fundamentals and applications of host•guest binding and measurement of the binding free energies.⁴ Given that supramolecular hosts are typically smaller and conformationally more homogenous than proteins and that some supramolecular systems achieve binding affinities

LIsaacs@umd.edu.

Conflicts of Interest.

The authors have no conflicts of interest in relation to the work contained in the paper.

[†]Electronic Supplementary Information (ESI) available: Details of synthesis, NMR, and ITC experiments. See DOI: [10.1039/D1NJ05209H](https://doi.org/10.1039/D1NJ05209H)

and selectivities that rival Nature suggested that host•guest systems (Figure 1) should be included in the SAMPL challenges.⁵

The Isaacs group has a longstanding interest in the cucurbit[n]uril (CB[n]) family of molecular containers⁶ and has been involved in the elucidation of the mechanism of CB[n] formation as a means to create new CB[n]-type receptors and in the delineation of their host•guest recognition properties.⁷ We discovered that CB[n] bind tightly and with high selectivity toward hydrophobic cations in water (K_a typically $10^6 - 10^{12} \text{ M}^{-1}$).⁸ The origin of the tight binding was traced to the presence of intracavity waters that lack a full complement of H-bonds that are released upon complexation.⁹ We, and others, have used CB[n]-type receptors as *in vivo* sequestration agents and for (targeted) drug delivery applications.^{5f, 10} Since SAMPL3, the Isaacs group has provided unpublished K_a values for guests toward various CB[n]-type receptors (e.g. CB[n] (n = 7, 8), acyclic CB[n] (e.g. **MIACid**), and glycoluril derived molecular clips).¹¹ The groups of Bruce Gibb and Michael Gilson have supplied blinded datasets for deep cavity cavitands and cyclodextrin derivatives, respectively.¹² One issue the computational chemists encountered in previous SAMPL challenges with acyclic CB[n]-type receptors was conformational sampling. Recently, we have become interested in the pillararene family¹³ of molecular containers (e.g. **WP6**, **P6AS**) as sequestration agents.^{5f, 14} Pillararenes are macrocyclic and display high affinity toward cationic guests like viologens in water which makes them ideally suited as an alternative scaffold for the SAMPL challenges.^{5c, 15} Herein, we describe the binding of **WP6**¹⁵ – which is a water soluble derivative of pillar[6]arene – toward a series of hydrophobic cations which serves as a blinded dataset for the SAMPL9 challenge.

Results and Discussion

This results and discussion section is organized as follows. First, we present the selection of the host (**WP6**) and guests (**G1 – G20**) used in the study. Subsequently, we present a qualitative investigation of the host•guest complexation by analysis of complexation induced changes in ¹H NMR chemical shift and multiplicity. Thereafter, we present the determination of host•guest binding affinity and enthalpy by isothermal titration calorimetry (ITC). Finally, we discuss the thermodynamic parameters as a function of guest structure and offer some conclusions.

Selection of Host and Guests.

Previous SAMPL challenges have featured macrocyclic CB[7] and CB[8],^{2b, 11b, 11f} glycoluril derived molecular clips and acyclic CB[n] that feature carboxylate or sulfonate groups,^{11a, c, e} deep cavity cavitands,^{12a, b} and cyclodextrins.^{12c} In previous challenges, issues relating to the conformational flexibility of acyclic CB[n] hosts and the degree of deprotonation of ionizable functional groups have arisen. Accordingly, for SAMPL9 challenge we decided to select **WP6** as host because it is more defined conformationally and is known to undergo strong host•guest complexation in water.^{14b, 15} Most studies of host•guest complexation of **WP6** use less competitive media (e.g. unbuffered water or buffered water). To make the SAMPL9 challenge more biologically relevant, we elected to perform our studies in phosphate buffered saline (PBS) at physiological pH (pH 7.4).

Given that **WP6** is anionic at neutral pH, we knew that binding of cationic guests would be favored. Accordingly, we selected guests **G1** – **G20** (Figure 2) which are mono- and diammonium ions which were available from our previous studies of CB[n]•guest complexation events.^{8, 11b, 11d, 16} Guests **G1** – **G20** feature different numbers of cationic residues, different alkylation states (e.g. 1°, 2°, 3°, 4°), and different sized hydrophobic residues. Given that **WP6** is highly negatively charged at neutral pH, we expected that **G1** – **G20** would form **WP6•G** complexes whose K_a values would span several orders of magnitude thereby making it easier for the computationalists to predict changes in binding free energy as a function of guest structure.

Qualitative ¹H NMR Host•Guest Recognition Study

As drawn in Figure 1, **WP6** features a C_6 -axis and overall D_6 -symmetry and is therefore chiral.^{13b} However, because the OCH_2CO_2Na substituents can rotate through the annulus of the macrocycle **WP6** is isolated as a racemic mixture of planar chiral macrocycles (e.g. R_p and S_p).¹⁷ Figure 3a shows the ¹H NMR spectrum recorded for **WP6** which features a single sharp resonance for H_a , H_b , and H_c on the chemical shift timescale. This observation strongly suggests that rotation through the annulus is fast on the chemical shift timescale. Initially, we studied the binding of **WP6** toward the panel of guests (**G1** – **G20**) in D_2O by ¹H NMR stoichiometry at 1:1 and 1:2 **WP6**:guest stoichiometry (Supporting Information). The ¹H NMR spectra recorded for **WP6**, **G1**, and 1:1 and 1:2 mixtures of **WP6** and **G1** (Figure 3) illustrate the spectral changes that are commonly observed. For example, at a 1:1 **WP6**:**G1** stoichiometry (Figure 3c), the resonances for **G1** within the **WP6•G1** complex undergo substantial upfield shifts due to their location in the magnetically shielding environment of the macrocyclic cavity defined by the aromatic walls. Conversely, host resonance H_a undergoes a smaller downfield shift which can be explained by changes in the orientation of the aromatic walls with respect to each other. More interestingly, the H_c resonance of the OCH_2CO_2Na groups with the **WP6•G1** complex shift downfield and split into an AB quartet (H_c , H_c') for the diastereotopic methylene groups. In combination, this indicates that rotation through the annulus is slow on the chemical shift timescale for **WP6•G1** but that exchange of guest **G1** is fast on the chemical shift timescale which renders the top and bottom portals of **WP6** equivalent. Figure 3d shows the ¹H NMR spectrum recorded at a 1:2 **WP6**:**G1** stoichiometry. Compared to Figure 3c, the resonances for **G1** shift back toward their locations for uncomplexed **G1** which further confirms the fast exchange of **G1** on the chemical shift timescale. The D_6 -symmetric conformation of uncomplexed **WP6** is dominant in aqueous solution. However, pillararenes are capable of conformational diastereoisomerism when one or more of the aromatic rings flips. For example, in the case of the **WP6•G13** complex we observe a dramatic increase in complexity in the 5.5 – 7.5 ppm region of the spectrum which is consistent with reduced symmetry of the complexes (Supporting Information, Figure S12).

The ¹H NMR spectra recorded for mixtures of **WP6** and guest **G14** (Figure 4) provide a beautiful example of stereochemistry and chemical exchange in host•guest chemistry. For example, the observation of a sharp singlets for **WP6** (Figure 4a) indicates that the top and bottom portals of **WP6** are equivalent due to the presence of a C_6 -axis and six perpendicular C_2 -axes resulting in D_6 point group symmetry. Similarly, the adamantane residue of guest

G14 has a C_3 -axis and three mirror planes which results in single resonances for H_m and H_n , whereas H_o and H_p are part of the diastereotopic CH_2 -group (Figure 4b). The protons on the $N-CH_2CH_2-N$ group (H_k and H_l) appear as coupled triplets as expected. The situation changes completely within the **WP6•G14** complex (Figure 4c). As can be seen (Figure 4a,c), the aromatic resonance H_a splits into two singlets (H_a, H_a'). Apparently, the **WP6•G14** complex undergoes slow guest exchange which renders the top and bottom portals of the complex chemically distinct with different chemical shifts; complexation maintains the C_6 -axis but eliminates the six perpendicular C_2 -axes. The presence of four doublets for H_c ($H_c - H_c'$) for **WP6•G14** reflects the top-bottom dissymmetry and that this CH_2 -group is diastereotopic within the overall chiral and racemic complex. Figure 5 shows an MMFF minimized molecular model of **WP6•G14** which illustrates these symmetry considerations. Even more interesting is the appearance of the resonances for guest **G14** within the **WP6•G14** complex. For example, H_k and H_l split into four resonances H_l, H_l', H_k, H_k' because the chiral **WP6•G14** complex renders these CH_2 -groups diastereotopic and all four protons are chemically distinct. Protons H_n still appear as a single resonance in **WP6•G14** because the C_3 -axis present in **G14** is maintained in the **WP6•G14** complex. Even more interesting is that the six protons H_m that appear as a single resonance in **G14** split into a pair of coupled doublets H_m and H_m' within **WP6•G14**. The three mirror planes that are present in the adamantane skeleton of **G14** are destroyed upon complexation to form the chiral **WP6•G14** complex which renders these three CH_2 -groups diastereotopic. All of the protons of guest **G14** experience a large upfield shift upon complexation which reflects their complexation inside the hydrophobic magnetically shielding environment of the **WP6** cavity. At a 1:2 **WP6:G14** stoichiometry the guest exchange rate increases which results in averaged NMR where the ethylene diammonium ion tail can point out of either portal which results in a merging of the H_a and H_a' resonances as well as the $H_c - H_c'$ resonances as expected based on symmetry considerations. The guest resonances also merge and shift back toward the chemical shift for uncomplexed **G14** as expected.

Measurement and Discussion of the Thermodynamic Parameters of Complex Formation.

After having qualitatively assessed the binding properties of **WP6** toward the guest panel by 1H NMR spectroscopy we decided to measure the thermodynamic parameters of complexation. Given that **WP6** is known to display tight binding and our desire to use a single analytical method across our measurements we turned to isothermal titration calorimetry (ITC) measurements which allows accurate K_a determination over a wide dynamic range.¹⁸ Figure 6a shows the thermogram measured when **WP6** (200 μM) in the ITC cell was titrated with a solution of **G7** (2.0 mM) in the syringe. All ITC experiments were conducted in duplicate. Figure 6b shows the fitting of the integrated heat values to a 1:1 binding model implemented in the PEAQ ITC data analysis software with $K_a = 1.31 \times 10^5 M^{-1}$ and $\Delta H = -3.18 kcal mol^{-1}$. The K_a and ΔH values for the weaker complexes ($K_a = 5 \times 10^6 M^{-1}$) were determined in an analogous manner by direct ITC titrations and are presented in Table 1. In these direct titrations, the fixed concentration of **WP6** in the cell was manipulated in order to optimize the c -value^{18c} and therefore sample a larger portion of the binding isotherm and therefore deliver more reliable results.

For the tighter binding complexes **WP6•G2** and **WP6•G12** with $K_a > 10^7 \text{ M}^{-1}$ we could not optimize the c -values by reducing the fixed concentration of **WP6** in the cell and therefore turned to competitive ITC titrations.^{18b} In competitive ITC titrations the cell contains a solution of **WP6** and an excess of a weaker binding guest into which a solution of the tighter binding guest is titrated. The integrated heat data from the competitive ITC titration is fitted to the competitive binding model implemented in the PEAQ ITC data analysis software using the known concentrations of host and weak binding guest along with the known K_a and ΔH values for the host•weak guest complexes as inputs to extract the K_a and ΔH values for the host•tight guest complex. Experimentally, it is important that the host•weak guest and host•tight guest complexes have significantly different ΔH values otherwise the titration will not produce sufficient heat to allow a proper fitting of the data. Experimentally, we selected **G7** as the weak binding complex because its K_a toward **WP6** is large enough to make it a reasonable competitor and the ΔH for the **WP6•G7** complex is significantly smaller than those of the other complexes. Figure 7a shows the thermogram recorded during the titration of a solution of **WP6** (100 μM) and **G7** (0.2 mM) in the cell with **G2** (1 mM) in the syringe. Figure 7b shows the fitting of the integrated heat versus **WP6:G7** molar ratio to the competitive binding model that allowed us to determine $K_a = 4.59 \times 10^7 \text{ M}^{-1}$ and $\Delta H = -6.10 \text{ kcal mol}^{-1}$ for the tighter **WP6•G2** complex. Please note that the limiting ΔH value at low molar ratio ($\approx -3.2 \text{ kcal mol}^{-1}$; Figure 7b) corresponds to the difference between the ΔH values for the **WP6•G7** and **WP6•G2** complexes. The K_a and ΔH values for the **WP6•G12**, **WP6•G14**, **WP6•G15** complexes were determined by an analogous competitive ITC titration (Supporting Information).

Measurement of the $\text{p}K_a$ values for **WP6**.

Given the importance of electrostatic interactions on the measured **WP6**•guest K_a values and the complications likely to be encountered by the computationalists in determining the average charge state of **WP6** at neutral pH, we decided to measure the $\text{p}K_a$ values for **WP6**. Previously, the Silveira group reported the $\text{p}K_a$ values for **WP5** obtained by pH metric titrations.¹⁹ The authors assume that each portal acts independently and report a total of five $\text{p}K_a$ values: 4.35, 4.49, 4.89, 5.30 and 6.34. Similar pH metric titrations were performed by a contract research organization (Pion, Supporting Information) in three different THF/water mixtures and the $\text{p}K_a$ values for **WP6** were determined as 3.62 ± 0.01 , 4.16 ± 0.01 , 4.41 ± 0.03 , 4.80 ± 0.07 , and 5.66 ± 0.01 after extrapolation to pure water using the Yasuda-Shedlovsky equation. Accordingly, **WP6** is predominantly present in the dodeca anionic form at pH 7.4

X-ray Crystal Structure of **G2**.

We attempted to grow single crystals of different host•guest complexes of **WP6** but were unsuccessful. In one attempt, we obtained single crystals of **G2•2I⁻** and performed x-ray diffraction measurements and solved the crystal structure of **G2** (Figure 8, CCDC 2114714).²⁰ In brief, guest **G2** adopts a linear geometry in the crystal with both dimethyl piperidine rings in the chair conformation. The dihedral angle of the central HC-CH unit of **G2** is 180° which minimizes unfavorable gauche butane type interactions.

Discussion of the Trends in Binding Affinity.

The binding constants measured for the complexation between **WP6** and **G1** – **G17** differ by over five orders of magnitude from 9010 M^{-1} to $4.69 \times 10^9 \text{ M}^{-1}$ (Table 1). The **WP6•G1** – **WP6•G17** complexes all uniformly driven by favorable changes in enthalpy with ΔH values ranging from $-3.18 \text{ kcal mol}^{-1}$ for **WP6•G7** to $-16.4 \text{ kcal mol}^{-1}$ for **WP6•G14**. Most of the complexes are also driven by energetically favorable entropic changes with $-\Delta T \Delta S$ values (Supporting Information) ranging from $-0.57 \text{ kcal mol}^{-1}$ for **WP6•G11** to $-4.35 \text{ kcal mol}^{-1}$ for **WP6•G2** and **WP6•G10**; the **WP6•G1** ($+1.63 \text{ kcal mol}^{-1}$), **WP6•G8** ($+3.58 \text{ kcal mol}^{-1}$), **WP6•G14** ($+3.25 \text{ kcal mol}^{-1}$), **WP6•G16** ($+1.87 \text{ kcal mol}^{-1}$) complexes are exceptions with positive $-\Delta T \Delta S$ values. These thermodynamic signatures for **WP6**•guest binding are consistent with the non-classical hydrophobic effect that was established in cyclophane chemistry by Diederich²¹ and documented in other systems most notably cucurbiturils.⁹ Some trends are discernible within this limited dataset and are discussed below.

Influence of the Number of Carbons Among Primary Mono Ammonium Ions.—

Guests **G11** (5 C-atoms), **G7** (6 C-atoms), **G3** (7 C-atoms), **G6** (8 C-atoms), and **G1** (12 C-atoms) are all primary mono-ammonium ions that differ in the number of C-atoms in the hydrophobic residue. The K_a values increase as the number of carbon atoms increases from **G11** to **G6** which can be explained by the increasing hydrophobicity of the scaffold as CH_2 units are incrementally added; we have seen related trends previously with **P6AS** and **CB[n]**-type receptors.^{14a, 16d, 22} Cyclododecylammonium ion **G1** binds more weakly ($K_a = 5.29 \times 10^4 \text{ M}^{-1}$) which suggests that **G1** may be too large for the cavity of **WP6**. Alternatively, the $-\Delta T \Delta S$ value for **WP6•G1** is $+1.63 \text{ kcal mol}^{-1}$ which suggests that confinement of the conformationally flexible **G1** imposes a large entropic penalty which reduces K_a . Other primary mono ammoniums whose K_a values were measured include **G4** and **G9**. Guest **G4** ($K_a = 5.08 \times 10^4 \text{ M}^{-1}$) which contains one silicon atom was found to bind somewhat more strongly than **G11** ($K_a = 3.37 \times 10^4 \text{ M}^{-1}$) which can be attributed to the slightly larger volume of **G4** due to the longer C-Si bonds. Adamantane guest **G9** ($K_a = 3.75 \times 10^4 \text{ M}^{-1}$) contains 10 C-atoms but binds even more weakly than **G1** presumably due to the need for the hydrophilic OH functional group of **G9** to remain solvated within the **WP6•G9** complex.

Influence of Guest Charge on Binding Affinity.—

Diammonium ion guests **G13** ($K_a = 1.63 \times 10^6 \text{ M}^{-1}$), **G10** ($K_a = 1.61 \times 10^7 \text{ M}^{-1}$), **G2** ($K_a = 4.59 \times 10^7 \text{ M}^{-1}$), **G12** ($K_a = 9.43 \times 10^7 \text{ M}^{-1}$), and **G14** ($K_a = 4.69 \times 10^9 \text{ M}^{-1}$) are the tighter binders within this dataset. The central hydrophobic cores of **G2**, **G12**, and **G13** each contain 10 carbon atoms which suggests that the lower K_a measured for **G13** is most likely due to the more hydrophilic viologen skeleton. The K_a for **WP6•G13** was previously measured by Huang in less competitive unbuffered water where $K_a = 1.02 \times 10^8 \text{ M}^{-1}$.¹⁵ Guest **G10** which contains only 8 C-atoms in its central hydrophobic core binds somewhat stronger than **G13** but weaker than **G2** and (\pm)-**G12**. The ability of guests **G2**, **G10**, **G12**, and **G13** to engage in favorable ammonium ion•••carboxylate interactions at both portals of **WP6** is likely the source of their high binding affinity. Complex **WP6•G14** which is the tightest complex in the dataset two ammonium•••carboxylate interactions at a single portal (Figure 5).

Cavity size effects.—Interestingly, bis quaternary ammonium ion **G5** binds very poorly to **WP6** ($K_a = 9010 \text{ M}^{-1}$) despite its dimethyl adamantane core and its 2+ charge. Figure 9 shows an MMFF minimized molecular model for **WP6•G5** which shows that **WP6** is too narrow to engulf the hydrophobic core of **G5** and instead simply binds to one of the pendant NMe_3^+ groups which explains the especially poor affinity. Related trends have been observed previously by us with CB[n]-type receptors.^{8b}

Influence of Secondary Electrostatic Interactions.—The diammonium ion guests **G2**, **G10**, and **G12** locate their cationic centers near the anionic portals of **WP6**. We wondered about the influence of pendant charged functionality on the observed K_a values. For example, **G17** is an analogue of **G4** that features a cationic $(\text{CH}_2)_3\text{NH}_3^+$ sidearm that would be expected to engage in attractive secondary electrostatic interactions with anionic **WP6**. We find that the **WP6•G17** complex is 4.5-fold tighter than the **WP6•G4** complex which corresponds to a difference of $-0.89 \text{ kcal mol}^{-1}$. Similarly, **G16** is an analogue of **G10** that features two anionic $(\text{CH}_2)_3\text{SO}_3^-$ sidearms that would be expected to engage in repulsive secondary electrostatic interactions with anionic **WP6**. Complex **WP6•G16** is 1220-fold weaker than **WP6•G10** which corresponds to a difference of $+4.2 \text{ kcal mol}^{-1}$ (or $+2.1 \text{ kcal mol}^{-1}$ per sidearm). Apparently, repulsive secondary electrostatic interactions exert a larger influence on K_a than attractive secondary electrostatic interactions.

Influence of Guest Methylation State.—In our recent study of **P6AS** we found that higher degrees of guest methylation (e.g. $1^\circ < 2^\circ < 3^\circ < 4^\circ$) resulted in significantly higher K_a values.^{14a} In the present dataset, *p*-xylenediamine derived guests **G10** and **G15** differ only in the degree of methylation. We find that the K_a values for **WP6** toward **G10** ($1.61 \times 10^7 \text{ M}^{-1}$) and **G15** ($1.76 \times 10^7 \text{ M}^{-1}$) are quite similar which establishes that methylation state changes do not play a major role in the host•guest trends of **WP6**.

Conclusions.

In summary, we have reported an investigation of the binding of **WP6** toward a panel of cationic hydrophobic guests **G1** – **G20** by a combination of ^1H NMR spectroscopy and ITC. The ^1H NMR measurements establish that the hydrophobic binding domains of the guest are located in the hydrophobic cavity of **WP6** which constitutes an anisotropic shielding region. The ^1H NMR spectra of **WP6•guest** complexes may appear simple when guest exchange is fast (e.g. **WP6•G1**, Figure 3), present a workshop on symmetry considerations when guest exchange is slower (e.g. **WP6•G14**, Figure 4), or be uninterpretable when **WP6** assumes an unsymmetrical conformation (e.g. **WP6•G13**, Supporting Information). The thermodynamic parameters of binding (K_a , ΔH) were measured by direct or competitive ITC and span from a low of 9010 M^{-1} for **WP6•G5** to $4.69 \times 10^9 \text{ M}^{-1}$ for **WP6•G14**. The **WP6•guest** complexes are generally driven by favorable ΔH and less favorable $-\Delta T \Delta S$ values which means that the non-classical hydrophobic effect governs the molecular recognition of **WP6**. The overall guest charge, the number of C-atoms in the hydrophobic binding domain, the presence of secondary electrostatic interactions, and cavity size effects all play a significant role in determining **WP6•guest** binding affinity. Perhaps most significantly, the thermodynamic data presented in Table 1 serves as a blinded dataset for the SAMPL9

challenge to allow to validate and improve their methods to compute binding free energies in aqueous solution. When those methods reach maturity it will significantly advance wide areas of supramolecular and medicinal chemistry.

Experimental.

WP6 was synthesized according to the reported procedure.¹⁵ Guests were available from previous studies.^{8, 11b, 11d, 16} ¹H NMR spectra were measured on Bruker spectrometers operating at 400 or 600 MHz using D₂O as solvent. Chemical shifts (δ) are referenced relative to the residual resonances for HOD (4.80 ppm). ITC experiments were conducted in the 200 μ L working volume of the sample cell of a PEAQ ITC instrument (Malvern) using a 40 μ L injection syringe. Host and guest solutions were prepared in phosphate buffered saline (PBS) at pH 7.4. The sample cell was filled to capacity (200 μ L) with the host solution and the guest solution was titrated in (first injection = 0.4 μ L, subsequent 18 injections = 2 μ L). In select cases, competitive titrations were required where host and an excess of weaker binding guest were included in the cell and the tighter binding guest was titrated into the cell. For direct titrations, the binding data was fitted using the 1:1 binding model implemented in the PEAQ-ITC analysis software whereas for competitive titrations the competition binding model was used.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgements.

We thank the National Institutes of Health (GM-124270) and the National Science Foundation (CHE-2105857) for financial support. We thank Profs. Kata Majerski, Marina Sekutor, and Robert Glaser for the samples of **G5**, **G12**, **G19**, and **G20**.

Notes and References

1. Mohs RC and Greig NH, *Alzheimers Dement*, 2017, 3, 651–657.
2. (a)Isaacs L, SAMPL Challenge, https://en.wikipedia.org/wiki/SAMPL_Challenge, (accessed June 11, 2021);(b)Muddana HS, Daniel Varnado C, Bielawski CW, Urbach AR, Isaacs L, Geballe MT and Gilson MK, *J. Comput.-Aided Mol. Des*, 2012, 26, 475–487; [PubMed: 22366955] (c)Muddana HS, Fenley AT, Mobley DL and Gilson MK, *J. Comput.-Aided Mol. Des*, 2014, 28, 305–317; [PubMed: 24599514] (d)Yin J, Henriksen NM, Slochower DR, Shirts MR, Chiu MW, Mobley DL and Gilson MK, *J. Comput.-Aided Mol. Des*, 2017, 31, 1–19; [PubMed: 27658802] (e)Rizzi A, Murkli S, McNeill JN, Yao W, Sullivan M, Gilson MK, Chiu MW, Isaacs L, Gibb BC, Mobley DL and Chodera JD, *J. Comput.-Aided Mol. Des*, 2018, 32, 937–963. [PubMed: 30415285]
3. (a)Isik M, Levorse D, Rustenburg AS, Ndukwe IE, Wang H, Wang X, Reibarkh M, Martin GE, Makarov AA, Mobley DL, Rhodes T and Chodera JD, *J. Comput.-Aided Mol. Des*, 2018, 32, 1117–1138; [PubMed: 30406372] (b)Isik M, Levorse D, Mobley DL, Rhodes T and Chodera JD, *J. Comput.-Aided Mol. Des*, 2020, 34, 405–420; [PubMed: 31858363] (c)Mobley DL, Liu S, Lim NM, Wymer KL, Perryman AL, Forli S, Deng N, Su J, Branson K and Olson AJ, *J. Comput.-Aided Mol. Des*, 2014, 28, 327–345; [PubMed: 24595873] (d)Guthrie JP, *J. Phys. Chem. B*, 2009, 113, 4501–4507. [PubMed: 19338360]
4. Lowe AJ, Pfeffer FM and Thordarson P, *Supramol. Chem*, 2012, 24, 585–594.
5. (a)Dempsey JM, Zhai C, McGarraugh HH, Schreiber CL, Stoffel SE, Johnson A and Smith BD, *Chem. Commun*, 2019, 55, 12793–12796;(b)Assaf KI and Nau WM, *Chem. Soc. Rev*, 2015, 44,

- 394–418; [PubMed: 25317670] (c)Liu Y, Zhou F, Yang F and Ma D, *Org. Biomol. Chem.*, 2019, 17, 5106–5111; [PubMed: 31070210] (d)Jordan JH and Gibb BC, *Chem. Soc. Rev.*, 2015, 44, 547–585; (e)Bom A, Bradley M, Cameron K, Clark JK, Van Egmond J, Feilden H, MacLean EJ, Muir AW, Palin R, Rees DC and Zhang M-Q, *Angew. Chem., Int. Ed.*, 2002, 41, 265–270; (f)Yin H, Zhang X, Wei J, Lu S, Bardelang D and Wang R, *Theranostics*, 2021, 11, 1513–1526. [PubMed: 33391548]
6. (a)Freeman WA, Mock WL and Shih N-Y, *J. Am. Chem. Soc.*, 1981, 103, 7367–7368; (b)Kim J, Jung I-S, Kim S-Y, Lee E, Kang J-K, Sakamoto S, Yamaguchi K and Kim K, *J. Am. Chem. Soc.*, 2000, 122, 540–541; (c)Day AI, Arnold AP, Blanch RJ and Snushall B, *J. Org. Chem.*, 2001, 66, 8094–8100; [PubMed: 11722210] (d)Day AI, Blanch RJ, Arnold AP, Lorenzo S, Lewis GR and Dance I, *Angew. Chem. Int. Ed.*, 2002, 41, 275–277; (e)Liu S, Zavalij PY and Isaacs L, *J. Am. Chem. Soc.*, 2005, 127, 16798–16799. [PubMed: 16316221]
7. (a)Ganapati S and Isaacs L, *Isr. J. Chem.*, 2018, 58, 250–263; [PubMed: 29805180] (b)Isaacs L, *Chem. Commun.*, 2009, DOI: 10.1039/b814897j, 619–629.
8. (a)Cao L, Sekutor M, Zavalij PY, Mlinaric-Majerski K, Glaser R and Isaacs L, *Angew. Chem. Int. Ed.*, 2014, 53, 988–993; (b)Liu S, Ruspic C, Mukhopadhyay P, Chakrabarti S, Zavalij PY and Isaacs L, *J. Am. Chem. Soc.*, 2005, 127, 15959–15967. [PubMed: 16277540]
9. (a)Nau WM, Florea M and Assaf KI, *Isr. J. Chem.*, 2011, 51, 559–577; (b)Biedermann F, Uzunova VD, Scherman OA, Nau WM and De Simone A, *J. Am. Chem. Soc.*, 2012, 134, 15318–15323; [PubMed: 22881280] (c)Biedermann F, Nau WM and Schneider H-J, *Angew. Chem. Int. Ed.*, 2014, 53, 11158–11171.
10. (a)Deng C-L, Murkli SL and Isaacs LD, *Chem. Soc. Rev.*, 2020, 49, 7516–7532; [PubMed: 33043945] (b)Jung H, Park KM, Yang J-A, Oh EJ, Lee D-W, Park K, Ryu SH, Hahn SK and Kim K, *Biomaterials*, 2011, 32, 7687–7694; [PubMed: 21788071] (c)Sun C, Zhang H, Li S, Zhang X, Cheng Q, Ding Y, Wang L-H and Wang R, *ACS Appl. Mater. Interfaces*, 2018, 10, 25090–25098; [PubMed: 29993234] (d)Wheate NJ and Limantoro C, *Supramol. Chem.*, 2016, 28, 849–856.
11. (a)Ma D, Glassenberg R, Ghosh S, Zavalij PY and Isaacs L, *Supramol. Chem.*, 2012, 24, 325–332; (b)Cao L and Isaacs L, *Supramol. Chem.*, 2014, 26, 251–258; (c)She NF, Moncelet D, Gilbert L, Lu XY, Sindelar V, Briken V and Isaacs L, *Chem. Eur. J.*, 2016, 22, 15270–15279; [PubMed: 27492252] (d)Murkli S, McNeill JN and Isaacs L, *Supramol. Chem.*, 2019, 31, 150–158; (e)Ndendjio SZ, Liu W, Yvanez N, Meng Z, Zavalij PY and Isaacs L, *New J. Chem.*, 2020, 44, 338–345; (f)Murkli S, Klemm J, Brockett AT, Shuster M, Briken V, Roesch MR and Isaacs L, *Chem. - Eur. J.*, 2021, 27, 3098–3105. [PubMed: 33206421]
12. (a)Sullivan MR, Yao W and Gibb BC, *Supramol. Chem.*, 2019, 31, 184–189; [PubMed: 31555042] (b)Suating P, Nguyen TT, Ernst NE, Wang Y, Jordan JH, Gibb CLD, Ashbaugh HS and Gibb BC, *Chem. Sci.*, 2020, 11, 3656–3663; [PubMed: 32864079] (c)Kellek K, Slochower DR, Schauerl M, Duggan BM and Gilson MK, *J. Comput.-Aided Mol. Des.*, 2021, 35, 95–104. [PubMed: 33037548]
13. (a)Xue M, Yang Y, Chi X, Zhang Z and Huang F, *Acc. Chem. Res.*, 2012, 45, 1294–1308; [PubMed: 22551015] (b)Ogoshi T, Yamagishi T-A and Nakamoto Y, *Chem. Rev.*, 2016, 116, 7937–8002; [PubMed: 27337002] (c)Wu J-R and Yang Y-W, *Chem. Commun.*, 2019, 55, 1533–1543; (d)Li Z and Yang Y-W, *Acc. Mater. Res.*, 2021, 2, 292–305; (e)Lou X-Y and Yang Y-W, *Adv. Mater.*, 2020, 32, 2003263; (f)Song N, Lou X-Y, Ma L, Gao H and Yang Y-W, *Theranostics*, 2019, 9, 3075–3093; [PubMed: 31244942] (g)Song N, Kakuta T, Yamagishi T-A, Yang Y-W and Ogoshi T, *Chem.*, 2018, 4, 2029–2053.
14. (a)Xue W, Zavalij PY and Isaacs L, *Angew. Chem., Int. Ed.*, 2020, 59, 13313–13319; (b)Yu G, Xue M, Zhang Z, Li J, Han C and Huang F, *J. Am. Chem. Soc.*, 2012, 134, 13248–13251. [PubMed: 22827832]
15. Yu G, Zhou X, Zhang Z, Han C, Mao Z, Gao C and Huang F, *J. Am. Chem. Soc.*, 2012, 134, 19489–19497. [PubMed: 23130568]
16. (a)Cao L, Hettiarachchi G, Briken V and Isaacs L, *Angew. Chem. Int. Ed.*, 2013, 52, 12033–12037; (b)Cao L, Skalamera D, Zavalij PY, Hostas J, Hobza P, Mlinaric-Majerski K, Glaser R and Isaacs L, *Org. Biomol. Chem.*, 2015, 13, 6249–6254; [PubMed: 25962667] (c)Sigwalt D, Sekutor M, Cao L, Zavalij PY, Hostas J, Ajani H, Hobza P, Mlinaric-Majerski K, Glaser R and Isaacs L, *J. Am. Chem. Soc.*, 2017, 139, 3249–3258; [PubMed: 28182422] (d)Liu W, Lu X, Meng Z and Isaacs L, *Org. Biomol. Chem.*, 2018, 16, 6499–6506. [PubMed: 30155536]

17. Zhu H, Li Q, Gao Z, Wang H, Shi B, Wu Y, Shangguan L, Hong X, Wang F and Huang F, *Angew. Chem., Int. Ed*, 2020, 59, 10868–10872.
18. (a) Wiseman T, Williston S, Brandts JF and Lin L-N, *Anal. Biochem*, 1989, 179, 131–137; [PubMed: 2757186] (b) Velazquez-Campoy A and Freire E, *Nat. Protocols*, 2006, 1, 186–191; [PubMed: 17406231] (c) Broecker J, Vargas C and Keller S, *Anal. Biochem*, 2011, 418, 307–309. [PubMed: 21854755]
19. Silveira EV, Wanderlind EH, Masson AK, Cordeiro PS, Nascimento V, Affeldt RF and Micke GA, *New J. Chem*, 2020, 44, 2701–2704.
20. Crystal Data for C₁₄H₃₀N₂I₂ (M = 480.20 g/mol): monoclinic, space group P2₁/c (no. 14), a = 6.5208(2) Å, b = 14.5116(5) Å, c = 10.1283(3) Å, β = 104.2182(5)°, V = 929.05(5) Å³, Z = 2, T = 296(2) K, μ(MoKα) = 3.375 mm⁻¹, D_{calc} = 1.717 g/cm³, 16743 reflections measured (5.01° 2θ – 62.496°), 3037 unique (R_{int} = 0.0262, R_{sig} = 0.0147) which were used in all calculations. The final R₁ was 0.0240 (I > 2σ(I)) and wR₂ was 0.0553 (all data). Refinement details: H atoms were positioned from the geometric considerations and refined as riding on the attached atoms with U_{iso} constrained to be 20% (50% for methyl group) larger than U_{eq} of the attached group. Orientation of methyl groups was optimized.
21. (a) Meyer EA, Castellano RK and Diederich F, *Angew. Chem., Int. Ed*, 2003, 42, 1210–1250; (b) Persch E, Dumele O and Diederich F, *Angew. Chem., Int. Ed*, 2015, 54, 3290–3327.
22. Murkli S, Klemm J, King D, Zavalij PY and Isaacs L, *Chem. - Eur. J*, 2020, 26, 15249–15258. [PubMed: 32658342]

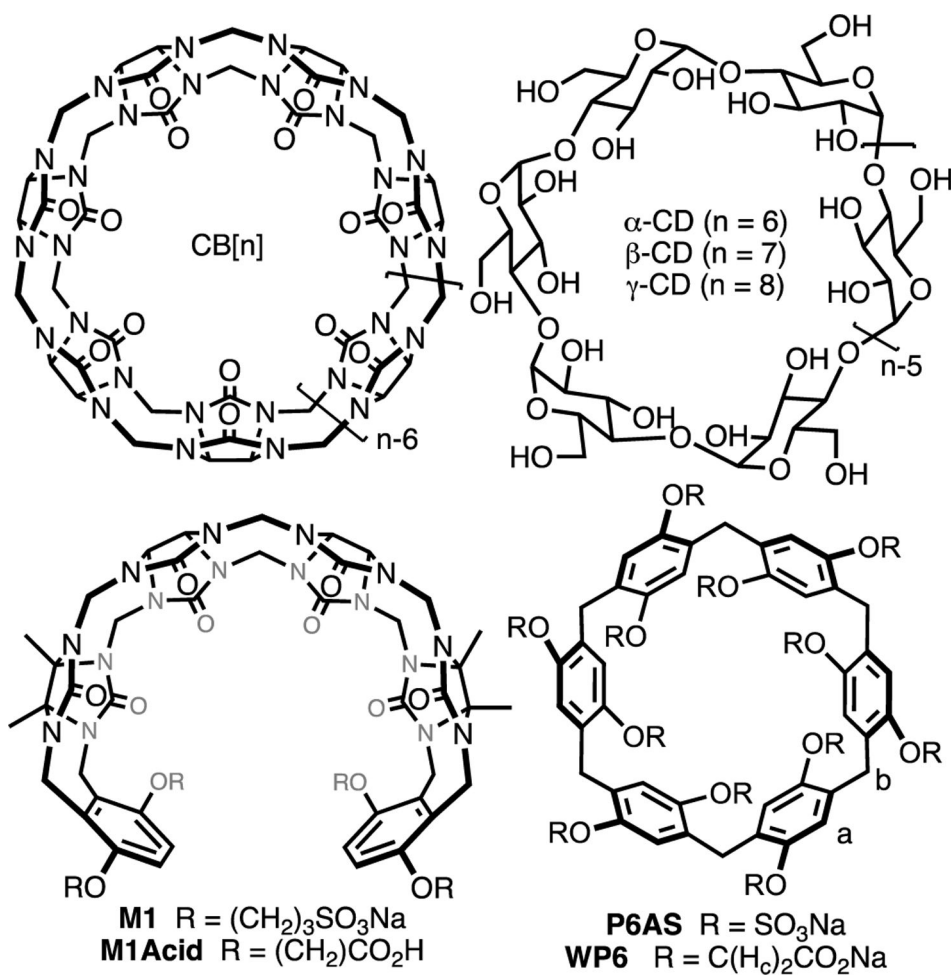


Figure 1.
Structures of (acyclic) CB[n], cyclodextrins, and pillararenes.

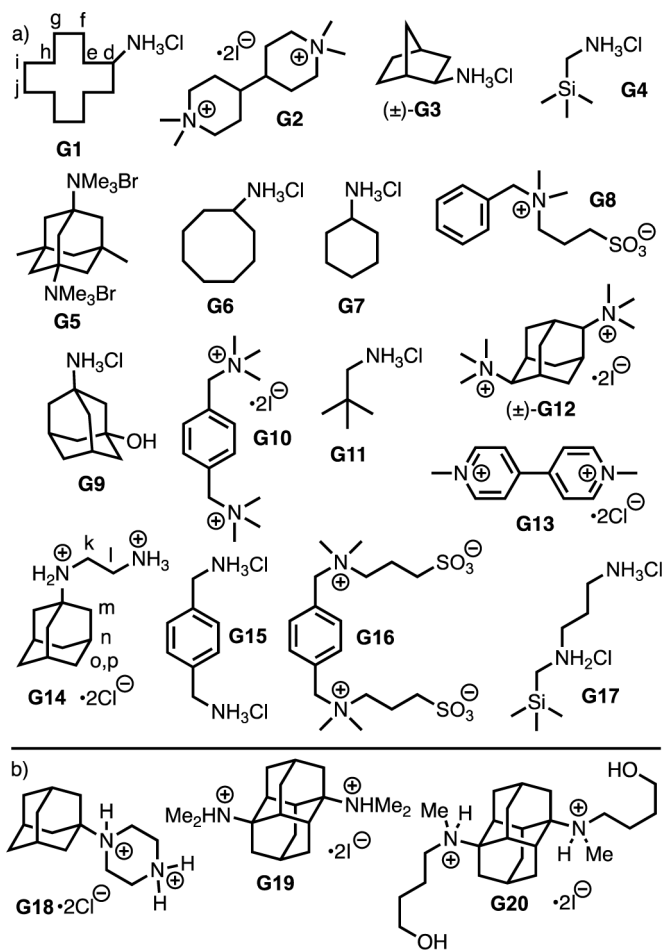


Figure 2. Structures of guests **G1** – **G20** used in this study. Panel a) guests studied by ^1H NMR and ITC, b) guests studied only by ^1H NMR.

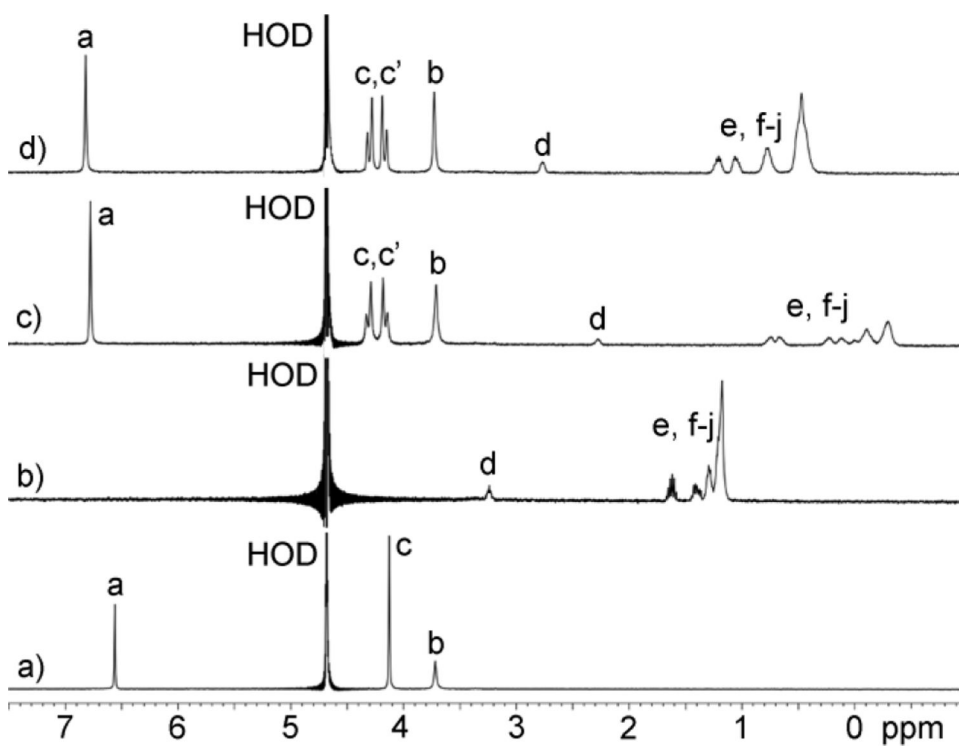


Figure 3. ¹H NMR spectra recorded (400 MHz, RT, D₂O) for: a) **WP6** (1 mM), b) **G1** (1 mM), c) a mixture of **WP6** (1 mM) and **G1** (1 mM), and d) a mixture of host **WP6** (1 mM) and **G1** (2 mM).

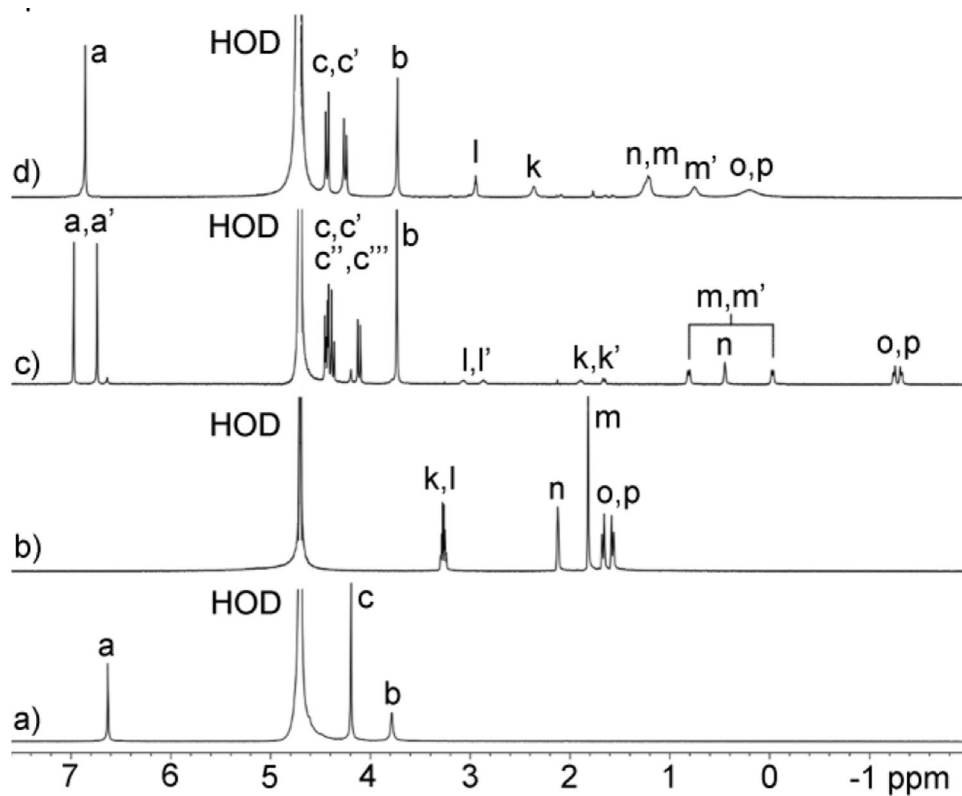


Figure 4. ¹H NMR spectra recorded (600 MHz, RT, D₂O) for: a) **WP6** (1 mM), b) **G14** (1 mM), c) a mixture of **WP6** (1 mM) and **G14** (1 mM), and d) a mixture of host **WP6** (1 mM) and **G14** (2 mM).

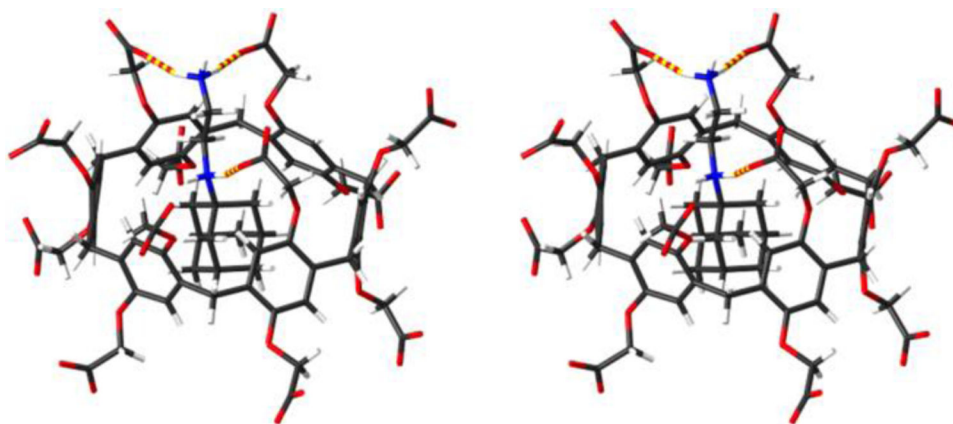


Figure 5. Cross-eyed stereoview of an MMFF minimized model of **WP6•G14**. Color code: C, gray; H, white; N, blue; O, red; H-bonds, red-yellow striped.

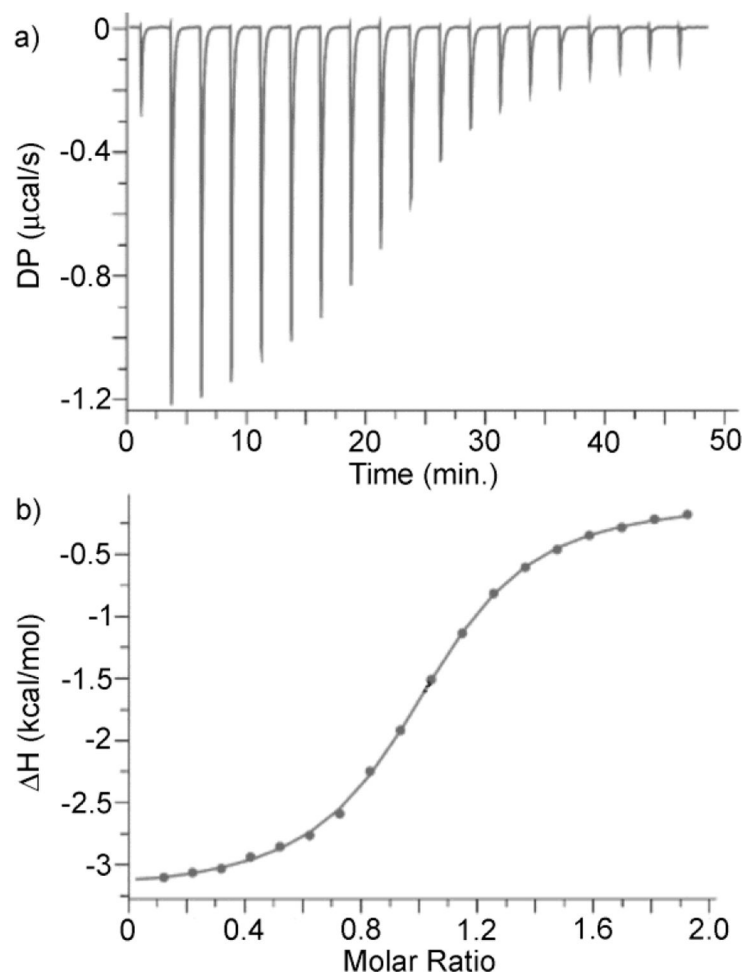


Figure 6.

a) ITC thermogram recorded during the direct titration of **WP6** ($200 \mu\text{M}$) in the cell with **G7** (2.0 mM) in the syringe, b) Fitting of the data to a 1:1 binding model with $K_d = 1.31 \times 10^5 \text{ M}^{-1}$.

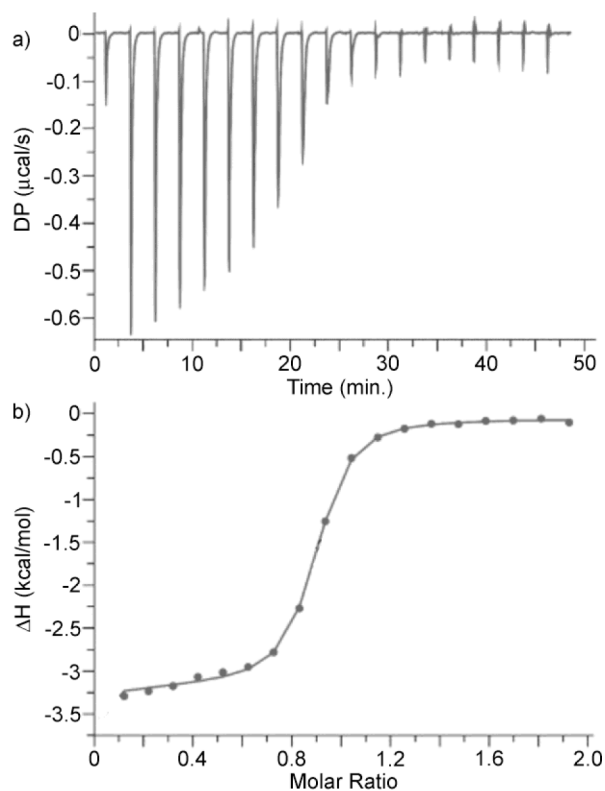


Figure 7.

a) ITC thermogram recorded during the competitive titration of a mixture of **WP6** ($100 \mu\text{M}$) and **G7** (0.2 mM) in the cell with **G2** (1.00 mM) in the syringe, b) Fitting of the data to a competitive binding model with $K_a = 4.59 \times 10^7 \text{ M}^{-1}$ and $H = -6.10 \text{ kcal mol}^{-1}$.

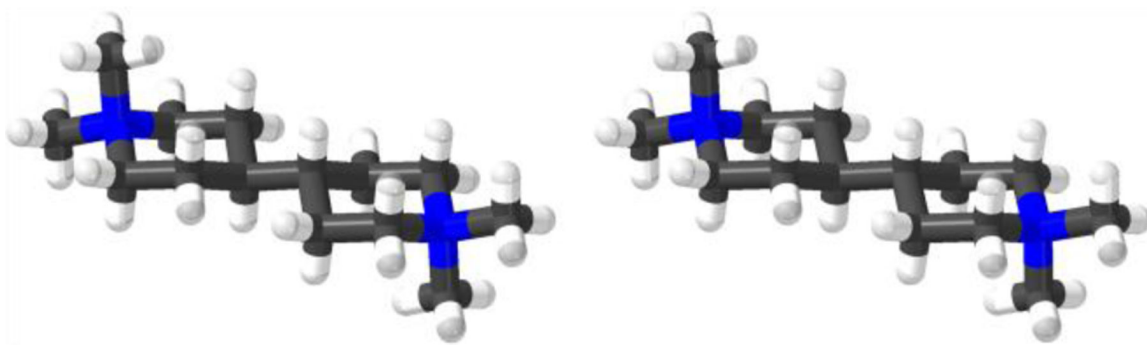


Figure 8.
Cross-eyed stereoview of the x-ray crystal structure of **G2**. Color code: C, gray; H, white; N, blue; I, purple.



Figure 9. Cross-eyed stereoview of an MMFF minimized model of **WP6•G5**. Color code: C, gray; H, white; N, blue; O, red.

Table 1.

Binding constants (K_a , M^{-1}) and enthalpies (ΔH , kcal mol^{-1}) measured for **WP6**•guest complexes. Conditions: 1x PBS buffer, pH 7.4, 298.15 K.

Guest	K_a (M^{-1})	ΔH (kcal mol^{-1})
G1 ^a	$(5.29 \pm 0.07) \times 10^4$	-8.08 ± 0.02
G2 ^b	$(4.59 \pm 0.35) \times 10^7$	-6.10 ± 0.02
(\pm)- G3 ^a	$(6.45 \pm 0.18) \times 10^5$	-4.75 ± 0.02
G4 ^e	$(5.08 \pm 0.11) \times 10^4$	-4.15 ± 0.02
G5 ^f	$(9.01 \pm 0.23) \times 10^3$	-3.95 ± 0.03
G6 ^c	$(7.09 \pm 0.44) \times 10^5$	-6.90 ± 0.07
G7 ^d	$(1.31 \pm 0.05) \times 10^5$	-3.18 ± 0.02
G8 ^e	$(2.35 \pm 0.04) \times 10^4$	-9.55 ± 0.05
G9 ^f	$(3.75 \pm 0.31) \times 10^4$	-5.31 ± 0.08
G10 ^b	$(1.61 \pm 0.08) \times 10^7$	-6.23 ± 0.02
G11 ^f	$(3.37 \pm 0.05) \times 10^4$	-5.61 ± 0.02
(\pm)- G12 ^g	$(9.43 \pm 0.31) \times 10^7$	-7.45 ± 0.02
G13 ^c	$(1.63 \pm 0.11) \times 10^6$	-4.98 ± 0.04
G14 ⁱ	$(4.69 \pm 0.09) \times 10^9$	-16.4 ± 0.02
G15 ^h	$(1.76 \pm 0.06) \times 10^7$	-7.03 ± 0.03
G16 ^e	$(1.32 \pm 0.03) \times 10^4$	-7.49 ± 0.06
G17 ^a	$(2.29 \pm 0.06) \times 10^5$	-4.15 ± 0.02

Measured by direct ITC titration of **WP6** in the cell with guest in the syringe

^a[**WP6**] = 0.1 mM, [guest] = 1.0 mM

^c[**WP6**] = 0.05 mM, [guest] = 0.5 mM

^d[**WP6**] = 0.2 mM, [guest] = 2.0 mM

^e[**WP6**] = 0.5 mM, [guest] = 5.0 mM

^f[**WP6**] = 1.0 mM, [guest] = 10 mM. Measured by competitive ITC titration of a mixture of **WP6** (0.1 mM) and **G7** in the cell with guest (1 mM) in the syringe

^b[**G7**] = 0.2 mM

^g[**G7**] = 0.5 mM

^h[**G7**] = 1.0 mM.

ⁱ Measured by competitive ITC titration of a mixture of **WP6** (0.1 mM) and **G15** (0.5 mM) in the cell with guest (1 mM) in the syringe.