Crystallographic Evidence for the Host-Guest Interaction of Metallamacromolecules (A Highlight Review)

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ABSTRACT

Metallamacromolecules offer opportunities to create structural diversity and interesting properties based on their unique frameworks and host-guest chemistry. Various types of assemblies can be created by the appropriate choice of the predesigned organic ligands containing various backbones and connectivity information and metal centers. These macrocyclic hosts contain a large and tunable hydrophobic inner cavity, which can be able selectively to recognize guest molecules. The goal of this highlight review is to describe the synthetic routes for the preparation of metallamacromolecules including stepwise and self-assembly strategies as well as their molecular recognition properties.

Key words: Host-Guest Chemistry, Metallamacromolecules.

INTRODUCTION

Macrocyclic host molecules such as calixarenes, cucurbiturils, and cyclodextrins act as catalytical nanoreactors,1 drug delivery agents,2 or molecular detectors.3 The successful application of these molecules often requires functionalization or derivatization with active functional groups. However, their preparation remains very challenging and often requires multistep syntheses that lead to overall low yields. Metallamacro molecules are a class of unique supramolecules, which have space inside to encapsulate cationic, neutral, or anionic guests.4,5 Generally, they do not expose them to the outside environment but bind with them via covalent or non-covalent bond interactions.6 The ability of these metal-organic hosts with the capability of binding guests strongly and selectively is difficult and still remains a challenge for chemists.7 The size and shape of the host structures could be fine-tuned by adjusting or modifying the ligand structure and the preferred geometry of the metal ions. The compounds of palladium(II) and platinum(II) with multidentate pyridine-based ligands were of the first reported cyclic hosts because their square planar geometry provides corners for the formation of different sizes and shapes.8 The simplicity of self-assembly has resulted in a plethora of self-assembled complexes having structures containing molecular triangles, squares, higher-order cyclic species, and closed structures with nanometer-sized cavities. Many impressive examples of
metallasupramolecular hosts can be found in studies reported by Lehn,9 Fujita,10 Sauvage,11 Stang,12 Mirkin,13 Hupp,14 Lu,15 Jin16 and other groups.17 Numerous types of metallacyclic hosts have been constructed in a step-wise or one-step synthetic route by using metallo-corner building blocks with bidentate rigid or semirigid linkers and difunctional pyridyl linkers. The preparation of metallacyclic hosts and the investigation of their complexations have produced many insights into significant noncovalent binding mechanisms. The formation and stability of complexation depend on the magnitude of the covalent bonding interaction.

Therefore, the preparation of compounds 1-6 has produced many insights into significant noncovalent interactions, including the nature of relevant noncovalent interactions, including those occurring in biological systems. Herein, for the sake of brevity, we highlight some specific examples of synthetic approaches and host behavior of discrete metallacyclic compounds in the solid state.

**Preparation and Host-Guest Studies**

The use of shape-specific designed ligands such as 1,3-bis(benzimidazol-1-ylmethyl)-4,6-dimethylbenzene L1, 1,3-bis(benzimidazol-1-ylmethyl)-2,4,6-trimethylbenzene L2, or 1,4-bis(benzimidazol-1-ylmethyl)-2,3,5,6-tetramethylbenzene L3 with different metal salts to form a series of metallacyclic structures [AgL1] (BF4)2 (1), [AgL2][CF3SO3]2 (2) and [CF3SO3]3[AgL3]BF4 (3), [CF3SO3]3[AgL4][BF4] (4), [ClO4]3[AgL5][BF4] (5) and [4H2O·NiL6]Cl2 (6), respectively (Scheme 1).20

X-ray diffraction analysis reveals that in proceeding from 1 to 6 the molecules display an increasingly regular shape, especially with respect to the inner cavity. The cavity in 1-3, however, is crucial for thermal stability and kinetics of formation and decomposition of such materials.18 In addition, computational studies are helpful for steering host-guest assembly into prescribed crystal architectures based on well-defined structure directing noncovalent bonding interactions.19 Therefore, the quantitative knowledge of host-guest assembly is of considerable importance for understanding the nature of relevant noncovalent interactions, including those occurring in biological systems. Herein, for the sake of brevity, we highlight some specific examples of synthetic approaches and host behavior of discrete metallacyclic compounds in the solid state.

![Scheme 1: Synthetic route for the preparation of compounds 1-6](image-url)
Fig. 1: Ball and stick representation of the crystal structure of (a) 2 showing two dangling CF$_3$SO$_3^-$ anions, and (b) 3 including the disordered CF$_3$SO$_3^-$ guest

Fig. 2: Ball and stick representation of the (a) [CF$_3$SO$_3^-\subset$Ag$_2$L$_3^-$]$^+$ cation in 4, (b) [ClO$_4^-$\subset$Cu$_2$L$_3$]$^{3+}$ cation in 5 and (c) [4H$_2$O\subset$Ni$_2$L$_3$] cage in 6
β-diketonate ligand (L4H₂ = 1,1’-(4,4’-biphenylene)bis-3,3-dimethylpentane-1,3-dione) in warm pyridine solution with cobalt(II) acetate tetrahydrate in pyridine afforded the neutral trinuclear Co³⁺ compound, [Co₃L₄₃(py)₆]. 5.55py· 0.6H₂O (7, py = pyridine) (Scheme 2).²²

Single-crystal structural analysis revealed that the β-diketonate ligands in 7 were not only bridged by three Co(III) ions but also formed a neutral equilateral triangle (Fig. 50). In addition, the pyridine coligands were axially coordinated to pseudooctahedral mode of Co³⁺ ions. In the solid state structure of 7, a triangular void space was found about 118 Å² in which the pyridine guest molecule was disordered in this void (Fig. 3). Compound 7 was claimed as the largest neutral M₃L₃ triangle so far characterized structurally.²³

The characteristics associated with the recognition of molecular rectangles with respect to the planar aromatic molecules and the Ag ion was reported. The alkoxy- or thiolato-bridged molecular rectangles \[ \{(CO)₃Re(m-ER)₂Re(CO)₃\}₂(m-L₅)₂ \] (8-11) (8, -OC₈H₁₇; 9, -OC₁₂H₂₅; 10, ER = -SC₄H₉; 11, ER = -SC₈H₁₇) were prepared by the reaction of Re₂(CO)₁₀ with the 4,4’-bipyridine (L₅, bpy) in the presence of higher aliphatic alcohols or a mercaptan under solvothermal reaction (Scheme 3).²⁴ The more hydrophobic nature of rectangles containing a dodecyl group enhanced their solubility in less polar solvents compared to those carrying an octyl and a butyl groups.

Fig. 3: Crystal structure of 7 along with the pyridine guest occupies in the void

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\text{Scheme 3: Self-assembly of neutral Re(I)-based rectangles 8-11}
\]
Using X-ray crystallographic studies, the rectangular architecture of 10 was confirmed by the distances between Re1···Re2 (3.787 Å) and Re1···Re2A (11.560 Å). The two bpy ligands in 10 are arranged in a face-to-face mode with a weak p-p stacking interaction (both centroid···centroid distances are 3.730 Å), which significantly stabilize the structure of 10. Single crystals of [10·pyrene] suitable for X-ray crystallographic analysis were grown up by the slow evaporation of solvent from an acetone solution of 10 in the presence of pyrene at 25 °C. It was found that the bpy ligands in 10 interact with pyrene via CH···p interactions. The face of the pyrene guest sits over the edges of the bpy linkers, nearly orthogonal with H(pyridyl)···C (pyrene) distances of 2.769-3.295 Å and a dihedral angle of 95° (Fig. 4). Crystal packing studies showed that the host-guest pairs, 10-pyrene, are arranged in a stair-like fashion, in which the guest pyrene molecules are not located within the molecular cavity of 10. This is an example of a CH···p interaction that is rarely designed into a host-guest pair.

Fig. 4: Crystallographic drawing of [10·pyrene] showing CH-p interactions in the solid state

Single crystals of [{10·(Ag⁺)₂(NO₃⁻)₂(C₃H₆O)₂}(C₃H₆O)] suitable for X-ray crystallographic study were obtained by dissolving the host 10 with the guest AgNO₃ in acetone, followed by slow evaporation at room temperature. The thiolato groups of 10 were firstly coordinated to a Ag atom [Ag1-S1, 2.4405 Å], which was further linked by two NO₃⁻ ions with Ag-O distances of 2.308-2.542 Å in addition to the coordination of one acetone molecule [Ag1-O10, 2.432 Å]. The intrinsic affinity of the sulphur atom towards the Ag(I) ion together with the Ag-ONO₂ interactions preceded to the formation of a one-dimensional supramolecular array via S···Ag···O connections. Furthermore, two adjacent Ag salts were connected together through NO₃⁻ ions with Ag···Ag distances of 4.05 Å, indicating weak argentophilic interactions (Fig. 5).

Fig. 5: Crystallographic drawing indicating the inclusion of AgNO₃ moieties by host 10 through Ag-S side-arm interactions and formation of a linear supramolecular array
The highly specific shape, the structural design or modification of the framework for the inclusion phenomena is difficult. Fujita and co-workers have utilized an ethylenediamine (en) derivative or pyridyl “capping” ligands to enforce the 90° cis geometry around square-planar coordinated Pd(II) or Pt(II) ions. The reaction of six (en)Pd²⁺ and four exo-tridentate ligands such as 1,3,5-tris(4-pyridyl)triazine (L6) or its derivative led to the formation of \( M_6L_4 \)-type coordination cages (12-14) (Scheme 4).

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\[ \text{Scheme 4: The reaction scheme for the self-assembly of compounds 12-14} \]

X-ray crystallographic study revealed that the four triazine panels in 12-14 occupy alternate faces of the octahedron and generate a very large hydrophobic cavity with a diameter of approximately 2 nm. The cationic cage (overall charge: 12⁺) is remarkably stable, encapsulates a variety of organic molecules. The unique and efficient binding of large guests by cage 12a has been well reported previously. Unlike the 12a-(o-carborane)₄ complex, which has been never crystallized, 12c-(o-carborane)₄, 12c-(1-adamantanol)₄, 12c-(diphenylmethane)₄, 12c-(tri-tert-butylbenzene), 12c-(1,2-bis(4-methoxyphenyl)-1,2-ethanedione)₂, and 12c-(tetrabenzylsilane) were easily crystallized by allowing its aqueous solution or organic solvents to stand at room temperature for a few days. The crystal structure of 12c-(o-carborane)₄ revealed that four guest molecules generate (o-carborane)₄ aggregates in the cavity of 12c (Fig. 6a). In the solid-state structure of 12c-(o-carborane)₄, it was difficult to distinguish carbon from boron atoms since the guest molecules were believed to spin at

\[ \text{Fig. 6: Crystal structure of (a) 12c-(o-carborane)₄ and (b) 12c-(diphenylmethane)₂} \]
low temperature. However, \(^1\)H NMR study elucidates that guest molecules are oriented in such a way that positive CH is outward while negative (BH)\(_n\) part is pointed inward.

The guest molecules in 12c·(1-adamantanol)\(_4\) are located at a fixed geometry without disordered. The hydrophobic adamantyl groups are pointed inside, whereas hydrophilic hydroxyl groups are pointed outside. In 12c·(diphenylmethane)\(_2\), two diphenylmethane guest molecules are located orthogonally inside the cavity of 12c through CH-p and p-p interactions (3.3-3.6 Å) between two aromatic rings of guests (Fig. 6b). The guest molecule 12c·(trityl-butylnbenzene) was disordered and triply located with 0.33 occupancy for each. The benzene ring of the guest was located on the triazine ring of one ligand in 12c with a distance of 4.5 Å. In the crystal structure of 12c·(tetrabenzylsilane), each portal accommodated one phenyl group of the guests. In addition, the guest geometries determined by X-ray analyses were consistent with the spectroscopic observation of the solution structure. Thus, the use of 2,2’-bipy ancillary ligand in these macrocycles was found to improve the crystallity of host-guest complexes.

The key to success with the covalently bonded dimetal units is that some coordination sites could be blocked by using non-labile bridging ligands such as \(N,N’\)-di\((p\)-anisyl)formamidinate (L7, DAniF), giving a building block [M\(_2\)(DAniF)\(_n\)]\(^{4-}\). With such a building block and linkers, a variety of discrete dimetal-containing macrocycles have been synthesized. For example, when cis\(-\)Mo\(_2\)(DAniF)\(_2\)^{2+} as a vertex building block and was treated with \(p\)- or \(m\)-trifluoromethyl substituted terephthaloyl-diamidate aromatic unit (L8a and L8b) as the linker, two neutral molecular squares 15a and 15b, respectively, were obtained and were structurally characterized. X-ray studies of 15a showed that the central square area, approximately 10 \(\times\) 10 Å\(^2\), is fenced by eight \(p\)-trifluoromethyl substituted terephthaloyl-diamidate aromatic unit (L8a and L8b) as the linker, two neutral molecular squares 15a and 15b, respectively, were obtained and were structurally characterized. X-ray studies of 15a showed that the central square area, approximately 10 \(\times\) 10 Å\(^2\), is fenced by eight \(p\)-trifluoromethyl substituted terephthaloyl-diamidate aromatic unit (L8a and L8b) as the linker, two neutral molecular squares 15a and 15b, respectively, were obtained and were structurally characterized. X-ray studies of 15a showed that the central square area, approximately 10 \(\times\) 10 Å\(^2\), is fenced by eight \(p\)-trifluoromethyl substituted terephthaloyl-diamidate aromatic unit (L8a and L8b) as the linker, two neutral molecular squares 15a and 15b, respectively, were obtained and were structurally characterized. X-ray studies of 15a showed that the central square area, approximately 10 \(\times\) 10 Å\(^2\), is fenced by eight \(p\)-trifluoromethyl substituted terephthaloyl-diamidate aromatic unit (L8a and L8b) as the linker, two neutral molecular squares 15a and 15b, respectively, were obtained and were structurally characterized. X-ray studies of 15a showed that the central square area, approximately 10 \(\times\) 10 Å\(^2\), is fenced by eight \(p\)-trifluoromethyl substituted terephthaloyl-diamidate aromatic unit (L8a and L8b) as the linker, two neutral molecular squares 15a and 15b, respectively, were obtained and were structurally characterized. X-ray studies of 15a showed that the central square area, approximately 10 \(\times\) 10 Å\(^2\), is fenced by eight \(p\)-trifluoromethyl substituted terephthaloyl-diamidate aromatic unit (L8a and L8b) as the linker, two neutral molecular squares 15a and 15b, respectively, were obtained and were structurally characterized.

Macromolecule 15b was considered as a class of bowl-shaped molecule in which four aryl groups on the bottom of the square were oriented inward, whereas the four on the top of each molecule were oriented outward (Fig. 7). When the two bowls were approached each other, they formed a cavity where two THF molecules could be encapsulated selectively although a mixture of tetrahydrofuran, toluene, and hexane was chosen as guest molecules in addition to intermolecular F\(\times\)\(\times\)H interactions.

Self-assembly of giant metallasacrocycles from metal ions and bridging ligands is one of the intriguing topics in current chemistry. The way of the CS\(_2\) loss from the metal complexes could be employed in the construction of high nuclearity metallasacrocycles. The reaction of ZnCl\(_2\) with 3,5-dimethylpyrazolate (L9, K[7mpzdtc]) in a mixture of MeOH and H\(_2\)O afforded [Zn(L9)]\(_2\), which was further dissolved in a mixture of DMF and water followed by the elimination of CS\(_2\) to give [Zn\(_4\)(m-dmpz)\(_6\)(m-OH)\(_2\)]\(_4\) (16). An X-ray structural analysis revealed that compound 16 was consisting of four [Zn\(_4\)(m-dmpz)\(_6\)(m-OH)\(_2\)]\(_4\) units coordinated by sharing pairs of m-dmpz and m-OH\(_2\) anions, forming another...
cyclic structure with a crystallographic four-fold axis running through the centre of the structure (Fig. 8). The distances of the edge and diagonal of 16 were found in the range of 15.7 and 18.0 Å, respectively, indicating a nanosized molecular crown structure.

The solvent DMF molecules were located inside and outside of 16 through O2–H···O3 and O1–H···O4 hydrogen bonding interactions, respectively. A very large voids in 16·4DMF with some $10^{16}$ Å$^3$ centered at (0, 0, 0.132) and (0.5, 0.5, 0.632) were estimated.

Conclusions and Outlook

In this review, the developments of metallamacromolecules from bi-, tri-, tetra- and hexanuclear through octanuclear to hexakaidecanuclear assemblies and their host-guest properties were highlighted. The versatility and efficient synthetic methodologies enable straightforward access to metallamacromolecules, thus allowing the properties to be fine-tuned through a precise chemical alteration of the architectures, opening the way to their use in host-guest chemistry. Studies of synthetic macromolecules with guests have provided significant insight into the nature of relevant non-covalent interactions, including those occurring in biological systems. We demonstrated through some examples that coordination assemblies with well-defined cavities could be used as molecular hosts to encapsulate a variety of guests. According to their solid-state structural studies, the π–π, CH–π interactions, hydrogen bonding and metal–ligand coordination are involved in these systems.

In order to understand the effects on the shape and size of metallamacromolecules with guest molecules, solution-state methods such as NMR spectroscopy and mass spectrometry and solid-state method (X-ray structure analysis) should be integrated for understanding of the host–guest behavior. When the binding interactions are relatively weak and labile, the host–guest complex found in solution and in the crystal may be different. Although the nature of their binding from $^1$H NMR spectral study in solution is informative, X-ray crystallography provides the clearest structural evidence of the subtle intermolecular interactions. The several mechanisms including guest inclusion, guest exchange, and host–guest interactions are really tedious to follow up because of dynamic and flexible nature of metallahosts. The another challenge to this field is to engineer a system that produces significant changes in selectivity, taking into account the many factors or interactions that may be reinforcing each other or may compete. Thus the functionalization and application of host–guest chemistry of metallamacromolecules still in its fancy and have to be carried out in the coming years. We believe that this highlight review will increase awareness of metallamacromolecules and accelerate the development of host-guest functional materials.

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