Pd(II) Coordination Sphere Engineering: Pyridine Cages, Quinoline Bowls, and Heteroleptic Pills Binding One or Two Fullerenes

Bin Chen,† Julian J. Holstein,‡ Shinnosuke Horiuchi,‡,† Wolf G. Hiller,† and Guido H. Clever*†‡

†Faculty of Chemistry and Chemical Biology, TU Dortmund University, Otto-Hahn Straße 6, 44227 Dortmund, Germany
‡Division of Chemistry and Materials Science, Graduate School of Engineering, Nagasaki University, Bunkyo-machi, Nagasaki 852-8521, Japan

ABSTRACT: Fullerenes and their derivatives are of tremendous technological relevance. Synthetic access and application are still hampered by tedious purification protocols, peculiar solubility, and limited control over regioselective derivatization. We present a modular self-assembly system based on a new low-molecular-weight binding motif, appended by two palladium(II)-coordinating units of different steric demands, to either form a [Pd_L2 ]4+ cage or an unprecedented [Pd_L3(MeCN)]4+ bowl (with L1 = pyridyl, L2 = quinolinyl donors). The former was used as a selective induced-fit receptor for C60. The latter, owing to its more open structure, also allows binding of C70 and fullerene derivatives. By exposing only a fraction of the bound guests’ surface, the bowl acts as fullerene protecting group to control functionalization, as demonstrated by exclusive monoaddition of anthracene. In a hierarchical manner, sterically low-demanding dicarboxylates were found to bridge pairs of bowls into pill-shaped dimers, able to host two fullerenes. The hosts allow transferring bound fullerenes into a variety of organic solvents, extending the scope of possible derivatization and processing methodologies.

INTRODUCTION

As stable carbon allotropes, fullerenes feature curved, fully \( \pi \)-conjugated surfaces with unique electronic properties that render them highly versatile for application in functional materials.† Established techniques for the separation of industrial fullerene mixtures are based on sublimation, crystallization, extraction, and chromatographic protocols.‡ The scope of solution processing methods, e.g., for controlled derivatization, preparation of composite materials, surface deposition, and device fabrication, is still limited due to the small choice of suitable aromatic and halogenated solvents.

Within the field of supramolecular chemistry, large efforts have been devoted to the construction of fullerene receptors aimed at facilitating selective purification and derivatization methods.§ In this regard, covalent organic tweezers and macrocycles based on extended aromatic panels, such as calixarenes, porphyrins and extended tetrathiafulvalenes, fully conjugated belts, as well as curved architectures such as triptycenes, have been intensively studied.¶ As most of these host compounds are the result of lengthy syntheses, self-assembled fullerene binders composed of much simpler building blocks have moved into focus, recently. Among those, metal-mediated rings and cages are notably versatile, owing to their modular composition and tunable cavity. Examples include Yoshizawa’s anthracene-lined \([Mn_L3 ]4+\) cages,¶ Nitschke’s tetrahedral aromatic-paneled arrangement,§ cubic porphyrin boxes,¶ and Ribas’ heteroleptic prismatic cage.¶

Common to most previously reported metal-mediated fullerene receptors is their rather high molecular weight, as a result of maximizing the offered \( \pi \)-surface area. We were therefore interested in designing a new metallo-supramolecular receptor based on the well-studied \([Pd_L2 ]4+\) coordination cage motif,§ that is (1) of lower molecular weight than existing hosts, (2) straightforward to synthesize and derivatize, and (3) capable of discriminating different fullerenes and dissolving them in a range of organic solvents. On the basis of computer-aided modeling, we further aimed at finding a perfect structural match between a curved binding motif, the connective metal complex, and the spherical \( C_{60} \) fullerene. Our design is based on a curved dibenzo-2.2.2-bicyclo-octane backbone reminiscent of triptycene but lacking the third benzene ring that is not in touch with the fullerene guest (Figure 1a and Figure S117).

We introduced phthalimide-based joints to pyridines, that coordinate to Pd(II) cations, thereby bringing all four ligands into perfect relative distances to be able to encapsulate \( C_{60} \) fullerene. Indeed, the as-planned host performed excellently as a fullerene host, and initial concerns that a receptor containing eight phthalimide units would become too electron-deficient to bind \( C_{60} \) did not materialize. In addition, ligand derivatization with sterically more demanding quinoline donors yielded an unprecedented \([Pd_L2 X_2 ]4+\) bowl structure (\( X = CH_2, CN \) or Cl), which allowed us to extend the guest scope to \( C_{70} \) and

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use it as a fullerene-protecting group in cycloaddition reactions with anthracene. The bowl’s sterically congested coordination environment allows clean conversion into a heteroleptic assembly via hierarchical reaction with bridging carboxylates, thus yielding a pill-shaped dimer, capable of binding one ligand, both empty hosts, and three di-atomic complexes to cage [Pd(L^4)^4+] capable of selectively binding C_{60}.

**RESULTS AND DISCUSSION**

**Ligand Synthesis, Cage Assembly, Fullerene Binding.** Backbone dianhydride, depicted in Figure 1a, was prepared in four steps starting from 4-bromo-1,2-dimethylaniline and 2,5-hexane-dione according to a reported procedure. Straightforward conversion into ligands L^1 and L^2 was achieved by reaction with 3-amino-pyridine and 6-aminoquinoline, respectively. Likewise to other banana-shaped, bis-monodentate pyridyl ligands reported previously, heating a 2:1 mixture of L^1 and [Pd(MeCN)_4](BF_4)_2 in deuterated acetonitrile at 70 °C for 1 d resulted in the quantitative formation of cage [Pd(L^4)^4+], unambiguously characterized by ^1H NMR spectroscopy (Figure 1b and c), mass spectrometry (Figure S8), and a single crystal X-ray structure (Figure 4a). The ^1H NMR spectrum reveals that the proton signals of the pyridine moieties undergo a downfield shift associated with metal complexation. Encapsulation of C_{60} and C_{70} were tested by stirring acetonitrile solutions of the cage over the solid fullerenes (usually adding an excess of finely ground material, while about 1.2 equiv were found to be sufficient). Intriguingly, [Pd(L^4)^4+] is only able to encapsulate C_{60} but not C_{70}, which we attribute to the good match in terms of shape (spherical C_{60} vs ellipsoidal C_{70}) and size (572 Å^3 void space; Figure S114; vs 547 Å^3 of C_{70})

Diffusion of isopropyl ether into the acetonitrile solution of [C_{60}@Pd(L^4)^4+] allowed us to grow single crystals in the form of red plates, suitable for synchrotron X-ray diffraction analysis. Three crystallographically independent host–guest complexes were found in the asymmetric unit, all showing the same C_{60}-occupied cage [C_{60}@Pd(L^4)^4+] with the C_{60} guest disordered over two positions, but featuring slightly different Pd–Pd distances (14.66, 14.61, and 14.55 Å, respectively) due to a certain degree of backbone flexibility and crystal packing effects (see Figure 4b and S108). The average distance from the ligand benzene ring centroids to the center of C_{60} is 6.72 Å (3.56–3.88 Å to the six/five-membered rings of C_{60}, see Table S8), verifying that the precisely designed concave inner surface can serve as fullerene receptor through strong π–π interactions. Similar distances were reported for C_{60} receptors based on other aromatic systems. The average distance from pyridine hydrogen H_2 to the C_{60} centroid is 6.11 Å (2.76–3.29 Å to the six/five-membered rings of C_{60} see Table S8), further indicating significant contribution from CH–π interactions. Colorless, block-shaped crystals could be grown when tetrahydrofuran (THF) was diffused into the acetonitrile solution of [C_{60}@Pd(L^4)^4+] X-ray analysis revealed a [Pd(L^4)^4+] cage, not containing any fullerenes, but two BF_4^- counteranions (Figure 4a and S107). The D_{4h}-symmetric cage shows a Pd–Pd distance of 15.94 Å. When comparing the geometries of the BF_4^- containing cage [Pd(L^4)^4+] the guest–cage geometry [C_{60}@Pd(L^4)^4+] it turns out that fullerene binding leads to an averaged Pd–Pd distance shortening of 1.3 Å. This goes along with a twist of all four ligands in a helical manner around the encapsulated C_{60}. As a result, the dihedral angle between two pyridine arms of the same ligand is 62.3° (Figure S108), whereas the corresponding dihedral angle in the free cage [Pd(L^4)^4+] is only 1.0°. Hence, uptake of C_{60} leads to a conformational change of the cage geometry indicating an induced-fit structural adaptation to perfectly accommodate C_{60} within the cavity.

**Self-Assembly of Bowls and Guest Uptake.** The initial aim of synthesizing quinoline-modified ligand L^2, featuring a larger N–N distance than found in L^1, was to enlarge the cavity size, thus allowing to accommodate larger guests inside the cage. To our surprise, prolonged heating of a 2:1 mixture of ligand L^2 and [Pd(MeCN)_4](BF_4)_2 in deuterated acetonitrile
yielded the expected [Pd\(L^2\)]\(^{4+}\) cage only as a minor product, accompanied by a 4-fold excess of a new species whose peculiar \(^1\)H NMR signal pattern as well as unambiguous mass spectral features allowed us to assign it to bowl-shaped compound [Pd\(L^2\)(MeCN)]\(^{4+}\), explainable as a [Pd\(L^2\)] cage lacking the fourth ligand (Figure S19 and S22). Previous work on Pd-mediated assembly with banana-shaped bis-monodentate pyridyl ligands never reported such bowl species (however, few reports exist for sterically more demanding ligand systems). Closer inspection of the steric situation around the metal coordination sites suggested that the quinolines’ hydrogen atoms (H\(_I\)) adjacent to the coordinating nitrogen atoms seem to cause significant steric crowding. This hypothesis was indeed supported by the observation that reacting ligand \(L^2\) with [Pd(MeCN)]\(^{4+}\)(BF\(_4\))\(_2\) in a 3:2 ratio at room temperature yielded [Pd\(L^2\)(MeCN)]\(^{4+}\) as a single product after 2 days. Further proof for the suggested bowl geometry came from characteristic cross peaks in the sample’s \(^1\)H–\(^1\)H NOESY NMR spectrum (Figure S15), the quinoline moieties’ \(^1\)H signal splitting into two sets with 2:1 integral ratio and the observation of prominent peaks in the ESI mass spectrum consistent with the formula [Pd\(L^2\)(MeCN)]\(^{4+}\), alongside further adducts with various anions [Pd\(L^2\)(MeCN)]\(^{−}\)C\(_I\)l, explainable as a [Pd\(L^2\)] cage. Surprisingly, slow vapor diffusion of isopropyl ether into a MeCN solution of bowl [Pd\(L^2\)(MeCN)]\(^{4+}\) produced single crystals for which a synchrotron-based diffraction experiment revealed the structure of cage [Pd\(L^2\)]\(^{4+}\) (Figure 4d), indicating the fine energetic balance between these two species.

Owing to the bowl’s open geometry, we found [Pd\(L^2\)(MeCN)]\(^{4+}\) to be able to bind both C\(_60\) to give [C\(_60@PdL^2\)(MeCN)]\(^{4+}\) and C\(_70\) to yield [C\(_70@PdL^2\)(MeCN)]\(^{4+}\) in a quantitative manner, both accompanied by characteristic color changes. In a competitive binding experiment, exposing bowl [Pd\(L^2\)(MeCN)]\(^{4+}\) to an equimolar mixture of powdered C\(_60\)/C\(_70\) (5 equiv/5 equiv), preferred binding of C\(_70\) was observed (ratio of [C\(_60@PdL^2\)(MeCN)]\(^{4+}\) to [C\(_70@PdL^2\)(MeCN)]\(^{4+}\) ≈ 4:1; Figure S78). The UV–vis spectrum of [C\(_60@PdL^2\)(MeCN)]\(^{4+}\) in acetonitrile showed a shoulder ranging from 330 to 420 nm compared to empty bowl [Pd\(L^2\)(MeCN)]\(^{4+}\), whereas [C\(_70@PdL^2\)(MeCN)]\(^{4+}\) displayed enhanced absorption in the longer wavelength region with one band around \(\lambda_{max} = 383\) and a broad band around 473 nm (Figure S105). Interestingly, the herein observed spectral features of fullerences bound to a rather electron-deficient host differ significantly from Yosizawa’s recent report on binding the same fullerences to electron-rich anthracene-based hosts in the same solvent.\(^{5b}\) In the \(^1\)H NMR spectra of the host–guest complexes, protons H\(_a\), H\(_b\), H\(_d\), and H\(_e\) located inward from the coordination sites, show distinct shifts upon binding of the fullerences within the bowl (Figure 2b). ESI mass spectrometry unambiguously supported the formation of the 1:1 host–guest complexes, while partial substitution of the acetonitrile ligands by trace amounts of anions was observed (Figure 2c). Consequently, we titrated 2 equiv of a solution of tetrabutylylammonium chloride into a solution of bowl [Pd\(L^2\)(MeCN)]\(^{4+}\), which afforded a quantitative conversion into compound [Pd\(L^2\)Cl\(_2\)]\(^{4+}\). Likewise, the fullerene carrying complexes could be converted into chloride-coordinated species, thus giving rise to very clean NMR and mass spectral results (Figure S42 and S45). We evaluated the thermal stability of the different bowl species in solution and found that

Figure 2. Self-assembly and characterization of bowl compounds. (a) L\(^2\), comprising sterically demanding quinoline donors, reacts with Pd\(^{II}\) to the bowl-shaped host, which binds both C\(_60\) and C\(_70\). (b) \(^1\)H NMR spectra of ligand L\(^2\) (600 MHz, 298 K, CD\(_3\)CN), bowl [Pd\(L^2\)(MeCN)]\(^{4+}\), [C\(_60@PdL^2\)(MeCN)]\(^{4+}\), [C\(_70@PdL^2\)(MeCN)]\(^{4+}\), [C\(_70@PdL^2\)(MeCN)]\(^{4+}\) all 0.64 mM, 298 K, CD\(_3\)CN) and photos of solutions. Red and blue marked proton signals are assigned to edge and central ligands, respectively. (c) High resolution ESI mass spectrum of [C\(_60@PdL^2\)(MeCN)]\(^{4+}\), prepared in pure CH\(_3\)CN.

[Pd\(L^2\)(MeCN)]\(^{4+}\) and [C\(_60@PdL^2\)(MeCN)]\(^{4+}\) partly converted into [Pd\(L^2\)]\(^{4+}\) and [C\(_70@PdL^2\)]\(^{4+}\), respectively, upon prolonged heating at 70 °C, whereas [C\(_60@PdL^2\)(MeCN)]\(^{4+}\) remained unchanged under these conditions (Figure S18, S29 and S36).

We were able to obtain the single crystal X-ray structures of ligand L\(^2\), cage [Pd\(L^2\)]\(^{4+}\), and host–guest complex [C\(_60@PdL^2\)(MeCN)]\(^{4+}\) (Figure 4c–e), thus allowing us to compare structural features between L\(^2\) in its free form, as part of the cage as well as the bowl geometry. In case of the free ligand and cage [Pd\(L^2\)]\(^{4+}\), angles between the backbone’s benzene planes are 119.8° and 120.2°, respectively, while slight widening (123.9°) was observed in case of the bowl. Likewise, the ligands’ N–N distance of 19.11 Å for free L\(^2\) and Pd–Pd distance of 18.80 Å for cage [Pd\(L^2\)]\(^{4+}\) compared to 20.22 Å for bowl [C\(_60@PdL^2\)(MeCN)]\(^{4+}\) indicate a slight opening of the cavity when the fourth ligand is missing. Of particular interest is comparing the immediate ligand environment around the Pd\(^{II}\) cations in the cage and bowl structures: as expected, the quinoline \(H_I\) hydrogen substituents lead to
significant steric crowding in case of cage [Pd,L3]4+, where four protons have to squeeze in the small space under the coordinated metal cation, leading to an average H–H distances of 2.31 Å (less than double the van-der-Waals radius of hydrogen, 1.2 Å) and a small deviation of the Pd3-coordination geometry from planarity with N–Pd–N angles of 176°.19 Even more compelling is the situation in the quinoline-based bowl [C60@Pd,L3(MeCN)]4+: here, the H–H hydrogen substituents belonging to the central ligand pushes both H–H hydrogens of adjacent quinolines aside in direction of the lean acetanilide ligand, giving rise to H–H distances of 2.56, 2.50, and 2.47 Å. In this bowl structure, the average distance from the ligand benzene ring centroids to the center of C60 is 6.79 Å (3.67–3.93 Å to the six/five membered rings of C60, see Table S14), and the average distance from quinoline H6 hydrogens to the C60 centroid is 6.39 Å (2.92–3.53 Å to the six/five membered rings of C60, Table S14), which are all slightly longer than the corresponding distances in [C60@Pd,L4]4+, still indicating both π–π and CH–π interactions between C60 and the bowl.

**Bowl-Protected Diels–Alder Reaction of C60 with Anthracene (Ac).** As observed in the crystal structure of [C60@Pd,L3(MeCN)](BF4)4 (Figure 4c), most of the C60 surface is covered by the bowl geometry, while exposing only a patch measuring about 25% of the total surface area to the acetonitrile solution environment. In order to elucidate whether the bowl-shaped host can modulate the fullerene’s chemical reactivity, we subjected complex [C60@Pd,L3(Cl2)]2+ (featuring chloride instead of acetanilide ligands) to a Diels–Alder reaction with anthracene. The same reaction had been studied before within a cubic coordination cage and a metal–organic-framework, both leading to formation of the bisadduct.3a,20 With pure C60, this reaction requires the use of problematic solvents such as benzene, chlorinated aromatics or CS2 and is known to deliver mixtures of mono-, di-, or even triadducts.21 Our system, on the other hand, allows the smooth conversion into the anthracene monoadduct (90% yield) when a MeCN solution of [C60@Pd,L3(Cl2)]2+ (0.56 mM, 1 equiv) is treated with up to 10 equiv of anthracene at 50 °C in the dark under a nitrogen atmosphere for about 12 h (equilibrium constant K12 = 2210 L mol⁻¹) (Figure S99 and Table S1).

The 1H NMR spectrum of the reaction product shows upfield shifts of bowl protons H4, H3, H2, H1. DOSY NMR analysis reveals that both the guest’s and host’s 1H NMR signals belong to a single species with a hydrodynamic radius of about 10.0 Å. The ESI mass spectra shows a single peak for the expected [C60Ac@Pd,L3(Cl2)]2+ ion (Figure 3b and 3c). We were further able to obtain single crystals of [C60Ac@Pd,L3(Cl2)](BF4)3, thereby delivering the first X-ray crystallographic report of the C60-anthracene monomonoadduct structure, which is a light- and oxygen-sensitive compound in solution. The structure reveals that no second anthracene molecule would be able to add to the encapsulated guest for sterical reasons. Interestingly, unlike the disordered C60 guests in the other structures, substituted fullerene C60Ac could be refined without geometrical restraints and did not require modeling of a second conformer. The fused substituent is not positioned in the middle of the window of the bowl structure, but rather tilted to one side (presumably for maximizing a stabilizing CH−O contact measuring 2.2 Å between one of the guest’s bridgehead protons and a ligand oxygen). Since no signal splitting is observed in the 1H NMR spectrum of the host–guest complex that would point to such a fixed, unsymmetrical conformation existing in solution, we expect the system to show a dynamic behavior comparable to a ball- and-socket joint.

**Dimerization of Bowls to Give Difullerene Complexes.** With the X-ray structures of bowls [C60@Pd,L3(Cl)2](MeCN)2]4+ and [C60Ac@Pd,L3(Cl2)]2+ in hand, carrying acetanilide and chloride ligands, respectively, to complement the Pd(II) square-planar coordination environments, the question arose whether other, sterically low-demanding donors could be installed in this position as well. Comparable to Stang’s methodology of mixing N-donor with carboxylate ligands,12 we succeeded in substituting both of the bowl’s acetanilides with carboxylate anions. Most interestingly, this allowed us to cleanly dimerize two bowls into pill-shaped assemblies using two terephthalate bridges (BDC, Figure 5). Dimerization could be shown for the empty bowl and its C60 as well as C70 complexes, as unambiguously demonstrated by NMR, UV–vis spectra, and ESI MS results.

The 1H NMR spectrum of the guest-free dimer shows the 2:1 signal splitting for the quinoline moieties retained and a single signal for the four protons of the phenylene bridge, indicating the formation of a single product containing the terephthalates symmetrically joining two identical halves. The dimer binds two fullerenes (C60 or C70) without experiencing any further symmetry-related signal splitting effects. Encapsulation-induced chemical shift changes are analogous to what was observed for the monomeric bowls (compare Figures 2b and 5b). Further, taking C60-occupied dimer [2C60@Pd,L3(BDC)2]4+ as an example, the NOE NMR spectrum
reveals a contact between signals Hc of the flanking quinolines (but not Hc′ of the central one) and the terephthalate proton HA, in accordance with the relatively short distance between these atoms observed in the PM6-optimized structure (Figure 5d and Figure S60). Furthermore, the increase in size from monomeric bowl to the pill-shaped dimer was confirmed by DOSY experiments, giving diffusion coefficients of \(5.20 \times 10^{-10}\) m\(^2\) s\(^{-1}\) for the monomer (Figure S27) and \(4.08 \times 10^{-10}\) m\(^2\) s\(^{-1}\) for the dimer (Figure S62). Comparison of UV−vis spectra of empty and fullerene-filled dimers shows very similar absorption bands compared to the monomeric bowl system, further confirming the binding of fullerenes within the inner cavities (Figure S106). The high-resolution ESI mass spectrum of \([2\text{C}_{60}@\text{Pd}_{4}\text{L}_{2}\text{(BDC)}_{2}]^{4+}\) revealed prominent signals at \(m/z 1488.7\) and \(2014.0\), consistent with the simulated isotopic pattern of \([2\text{C}_{60}@\text{Pd}_{4}\text{L}_{2}\text{(BDC)}_{2}]^{4+}\). Analogous NMR and MS results were also obtained for the guest-free (Figure S53−S57) and (C\(_{70}\))\(_2\)-containing (Figure S64−S69) dimers, showing that guest encapsulation is orthogonal to the dimerization process.

### CONCLUSIONS

On the basis of a rational design approach, we synthesized new self-assembled, low-weight fullerene receptors that allow the solution handling and facile crystallization of fullerenes as well as their adducts. The first receptor, consisting of pyridyl-terminated, bent ligands assembling with Pd\(^{II}\) cations to a \([\text{Pd}_{4}\text{L}_{2}\text{(BDC)}_{2}]^{4+}\) cage, is highly selective for C\(_{60}\). The second receptor was assembled using quinoline donors, which—owing to their steric demand—led to an unprecedented \([\text{Pd}_{2}\text{L}_{2}\text{(MeCN)}_{2}]^{4+}\) bowl geometry. This bowl not only was found to display a wider guest encapsulation scope (including C\(_{70}\)), but also is capable of serving as a supramolecular protecting group, allowing selective monofunctionalization of its fullerene guest. We further show that these bowls, both with and without bound fullerenes, can be cleanly dimerized by exchanging the acetonitrile ligands for terephthalate bridges, giving large, pill-shaped architectures of heteroleptic nature. The herein introduced receptors have the potential to serve as new tools for handling fullerenes and their derivatives in a wider range of organic solvents, and further allow their selective uptake and regioselective modification. Applications related to advanced fullerene derivatization, purification, structure elucidation, and device fabrication are sought to benefit from our findings.
Experimental details and further NMR, MS, UV–vis and crystal data (PDF)
Crystal data for \([C_{60}\text{Ac}@Pd_2L_1\text{Cl}_2]_4\), CCDC 1858158 (CIF)
Crystal data for \([C_{60}@Pd_2L_2(\text{MeCN})]_2\), CCDC 1850362 (CIF)
Crystal data for \([\text{Pd}_2L_4]^2_4\), CCDC 1850361 (CIF)
Crystal data for \(L_2^1\), CCDC 1850360 (CIF)
Crystal data for \([C_{60}@Pd_2L_1\text{Cl}_2]^4\), CCDC 1850359 (CIF)
Crystal data for \([\text{Pd}_2L_4]^4\), CCDC 1850358 (CIF)

**AUTHOR INFORMATION**

**Corresponding Author**
Guido H. Clever: 0000-0001-8458-3060
Shinnosuke Horiuchi: 0000-0002-5692-6377
Julian J. Holstein: 0000-0002-8385-7805

*Notes*
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