Supramolecular polymers with tunable topologies via hierarchical coordination-driven self-assembly and hydrogen bonding interfaces

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A powerful strategy to obtain complex supramolecular materials is the bottom-up construction of noncovalently bound materials by hierarchical self-assembly. This assembly process involves stepwise, uniform increases to the architectural complexity of a substrate, starting from discrete precursors and growing in dimensionality through controlled reactivity to a final product. Herein, two orthogonal processes are exploited: coordination-driven self-assembly and hydrogen bonding. The former relies on the predictable formation of metal–ligand bonds wherein the directionalities of the rigid precursors used determine the structural outcome. The latter uses 2-ureido-4-pyrimidinone interfaces that are structurally robust by virtue of the quadruple hydrogen bonding that can occur between subunits. By combining these two processes into a single system, it is possible to generate hierarchical materials that preserve the attractive tunability associated with discrete supramolecular coordination complexes. For instance, the synthesis of a one-dimensional chain comprising linked metallrahomoids is readily adapted to a 2D cross-linked hexagonal network by simply selecting a different metal acceptor precursor as an assembly component. The specific interactions between subunits, in this case platinum(II)-pyridyl bonds and the quadruple H-bonding of ureidopyrimidinone, are unchanged, establishing a unique strategy to obtain supramolecular polymers with marked topological differences with minimal synthetic redesign. In addition, the structural rigidity imposed by the inclusion of the platinum metallacycles serves to minimize the formation of cyclic oligomers, increasing the efficacy of formation and improving the properties of the resultant materials. Furthermore, this study taps the potential of organoplatinum(II) metallacycles in materials science.

Nature provides many examples of structurally complex and functional architectures by integrating multiple simple interactions that act in concert to produce the final macro- or supramolecular compound. For example, the primary structure of a protein comprises covalent amide bonds. Secondary structure is concerned with the formation of local segments of peptide chains as defined by simple hydrogen bonding. The next level of complexity, tertiary structure, is largely determined by hydrophobic interactions with contributions from disulfide bonds. Taken separately, these interactions are basic and fundamental in chemistry, yet when combined as components of hierarchical synthesis, impressive examples of complex, functional proteins may be found (1). Natural systems remain an ongoing motivation for scientists seeking new methods to obtain elaborate molecular assemblies using simple interactions as the impetus for formation. Hierarchical polymerization, in which growing complexity is achieved through well-defined steps that iteratively increase the dimensionality of a material, is particularly attractive in that each unique interaction can be carefully designed. As with proteins, the local interactions between the smallest subunits are often simple, yet taken as a whole, these interactions lead to complex architectures over various length scales (2, 3). In some cases, the most effective means to emulate natural systems is simply to exploit the moieties found therein. As such, the nucleobases found in DNA have been incorporated into numerous polymerization strategies that take advantage of their proclivity to hydrogen bond and provide favorable electrostatic interactions (4). Related examples can be found in the form of peptide amphiphiles that, on self-assembly and mineralization, can form nanofibers (5), as well as abiological organic polymers that form helical rods that exhibit host–guest chemistry (6), all with relatively simple interactions as their impetus for formation.

Organizing a single type of interaction between subunits is itself useful, and as new methods for controlling H-bonding, hydrophobic interactions, π–π stacking, Van der Waals forces, and metal-ligand bonding emerge, impressive examples of novel supramolecular polymeric materials tend to follow (7). For instance, organic precursors with hydrogen bonding interfaces can self-assemble into columnar structures (8), a pioneering example in a field that has steadily grown to include a number of materials (9–16), which have helped establish supramolecular polymerization as a critical subset of materials science (7, 17–20).

Metal–ligand bonding has proven to be a particularly valuable method to guide the organization of molecular precursors through strategies such as coordination-driven self-assembly (21–26). By carefully combining selected building blocks, discrete supramolecular coordination complexes (SCCs) can be obtained (27–31), based on the number and orientation of the labile coordination sites (for metal acceptors) and Lewis basic moieties (for organic donors). Because these metal–ligand interactions are general for a wide number of transition metals and organic linkers, a sizable library of 2D and 3D structures spanning multiple synthetic approaches has been developed over the last two decades. These synthetic strategies are unified under the theme of directional bonding, which reduces a target architecture to its constituent edges, faces, and vertices and seeks to replicate these on a molecular level (27). The substantial body of synthetic research has paved the way for multifunctional applications of SCCs ranging from molecular topology (32–34) and catalysis (35, 36) to light harvesting and gas capture (37, 38). Herein, we unify the themes of coordination-driven self-assembly and supramolecular polymerization through a hierarchical design strategy. Two SCC platforms are used, a $^{2 + 2}$ rhomboid constructed from a 60° acceptor and a 120° donor and hexagonal

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metallacycles that result from the [3 + 3] assembly of 120° acceptors and the same 120° donor. The self-assembly that occurs spontaneously on mixing these building blocks represents one component of the polymerization. To link these discrete metallacycles together, a second orthogonal interaction was designed to take advantage of hydrogen bonding. Specifically, the well-defined quadruple hydrogen bonding motif, 2-ureido-4-pyrimidinone (UPy), as developed by Meijer and coworkers, was selected (39). The UPy motif complements coordination-driven self-assembly by offering an attractive combination of high thermodynamic stability ($\Delta G \approx 10$ kcal/mol; $k_{\text{dimer}} > 10^7$ M$^{-1}$ s$^{-1}$ in chloroform) and rapid kinetic reversibility ($k_{\text{off}} \approx 8$ s$^{-1}$) (39). These favorable parameters have been previously exploited to link UPy-containing subunits, furnishing supramolecular polymers with high synthetic efficiencies and possessing dynamic reversibility (10, 17, 40). Moreover, the presence of self-selective UPy moieties was not expected to disrupt the platinum (Pt)-pyridyl bonding used for metallacycle formation, meaning the two interactions could independently occur within a single reaction flask. Coordination-driven self-assembly of the UPy-functionalized ligand with metal acceptors resulted in metallacycles decorated with a number of pendant hydrogen bonding sites. Hydrogen bonding could be mediated by selecting an appropriate solvent, thus linking the metallacycles into supramolecular polymers. This work establishes (1) the UPy functionalization of a 120° pyridyl ligand through ary-ether and amide linkers, which is readily used in (2) the formation of bis (phosphine) Pt(II) rhomboidal and hexagonal metallacycles decorated with H-bonding moieties that facilitate (3) the formation of linear chains (rhomboid) or cross-linked supramolecular polymers (hexagons; Fig. 1). The resulting materials highlight the advantages of SCC-based supramolecular polymers, in particular by attenuating the formation of cyclic oligomers due to the rigidity of the metallacyclic subunits. These polymers exhibit solvent swelling effects and the adaptive deformation of the cross-linked hexagonal networks via external forces results in macroscopic fibers that are robust enough to arrange into a knot. The strength, stretchability, and flexibility of these fibers is attributed to the ability of the cross-linked hexagons to reestablish Pt-pyridyl bonds that may be severed on mechanical stress, a property that is absent in supramolecular polymers that lack the SCC component described here.

**Results and Discussion**

The UPy-functionalized 120° dipyrindyl donor (4) was synthesized in a six-step pathway starting from commercially available 3,5-dibromophenol (Scheme 1A). After nucleophilic substitution of the phenolic hydroxyl, palladium-catalyzed introduction of the TMS-protected acetylene (10, 73% isolated yield) and subsequent desilylation in 86% isolated yield, compound 11 was reacted with 4-bromopyridine hydrochloride via a Sonogashira coupling to give 120° dipyrindyl ligand 12 in 95% isolated yield. This stable indoline-1,3-dione precursor is particularly useful in that it can be readily reduced to amino intermediate 13 in the presence of hydrazine (97% isolated yield). Given the well-established coupling reactions of primary amines, this compound provides a generalized route to furnish 120° donors functionalized with any moiety that is compatible with amine couplings. Herein, we linked amine-terminated compound 13 with a 1,1’-carbonyldimimidazol-activated alkyl-substituted pyrimidine (14), delivering the 120° UPy-functionalized dipyrindyl ligand (4) with an 80% isolated yield.

Stirring a mixture of 4 and 60° organoplatinum(II) acceptor 3,6-bis[trans-Pt(PES)2(NO3)2]phenanthrene (5) in a 1:1 ratio in CD2OD at 50 °C for 8 h induced the self-assembly of [2 + 2] rhomboid 1 decorated with UPy at its vertices (Scheme 1B). Self-assembled 120° [5 + 3] hexagons were prepared in a similar fashion and [3 + 3] hexagons were prepared in a six-step pathway starting from commercially available 3,5-dibromophenol (Scheme 1A). After nucleophilic substitution of the phenolic hydroxyl, palladium-catalyzed introduction of the TMS-protected acetylene (10, 73% isolated yield) and subsequent desilylation in 86% isolated yield, compound 11 was reacted with 4-bromopyridine hydrochloride via a Sonogashira coupling to give 120° dipyrindyl ligand 12 in 95% isolated yield. This stable indoline-1,3-dione precursor is particularly useful in that it can be readily reduced to amino intermediate 13 in the presence of hydrazine (97% isolated yield). Given the well-established coupling reactions of primary amines, this compound provides a generalized route to furnish 120° donors functionalized with any moiety that is compatible with amine couplings. Herein, we linked amine-terminated compound 13 with a 1,1’-carbonyldimimidazol-activated alkyl-substituted pyrimidine (14), delivering the 120° UPy-functionalized dipyrindyl ligand (4) with an 80% isolated yield.

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extend above and below the cores (SI Appendix, Fig. S28). As such, these UPy groups are well situated to undergo intermolecular hydrogen bonding.

In the $^1$H NMR spectrum of rhomboid 1, downfield shifts of the $\alpha$- and $\beta$-pyridyl protons relative to those of ligand 4 were observed, consistent with the loss of electron density that occurs on Pt–pyridyl bond formation. As shown in Fig. 3 (spectrum H), the $\alpha$- and $\beta$-pyridyl protons are split into two sets of two doublets on coordination. The $H_\alpha$ protons (8.62 ppm in 4; Fig. 3, spectrum I) are split into two doublets at 9.31 and 8.74 ppm (Fig. 3, spectrum H). Similarly, the $H_\beta$ protons (7.41 ppm in 4; Fig. 3, spectrum I) are also split into two doublets, at 7.94 and 7.81 ppm (Fig. 3, spectrum H).

The orthogonal relationship between the metal–ligand bonding responsible for SCC formation and the H-bonding expected of the UPy moieties manifests in two routes toward polymer formation. If a solvent such as methanol or DMSO is used, SCC formation occurs under conditions in which UPy dimerization is blocked due to hydrogen bonding with solvent molecules. The discrete metallacycles thus formed can then be polymerized by dissolution in a solvent that facilitates UPy–UPy interactions, such as CH$_2$Cl$_2$. Alternatively, the dissolution of 4 in CH$_2$Cl$_2$ will result in the formation of dimeric ligands that will self-assemble into polymeric materials directly on the introduction of metal acceptor 5 (SI Appendix, Fig. S29).

The formation of linear chains of 1 (Fig. 4A) was indicated by $^1$H NMR spectra in CD$_2$Cl$_2$ at concentrations ranging from 1.00 to 110 mM (Fig. 3, spectra A–H). The UPy N–H signals displayed large downfield shifts (observed between 10.0 and 13.5 ppm) and a lower intensity relative to the one in DMSO, giving direct evidence for the formation of H-bonds between UPy groups. Furthermore, the chemical shifts of the UPy protons showed little change on increasing monomer concentration. The single set of signals of the UPy protons indicates that there is no significant

formation of low-molecular-weight cyclic oligomers. The lack of cyclic oligomers was also supported by 2D diffusion-ordered $^1$H NMR spectroscopy (DOSY) NMR (vide infra). The circumvention of oligomer formation is attributed to the rigidity of the metallacyclic cores, which limits the formation of cyclic species and thereby improves the efficiency of polymerization in SCC–UPy systems. As the concentration of 1 approaches 40.0 mM, the $H_\gamma$ signal associated with acceptor 5 shifts upfield, ultimately overlapping with one set of the $H_\alpha$ protons resonances (Fig. 3, spectrum D). Above 20.0 mM, the peaks undergo broadening that obscures the well-resolved splitting associated with the free ligand and indicates the formation of high-molecular-weight rhombic aggregates (14, 19). Despite these spectral changes in the $^1$H NMR, the $^{13}$P ($^1$H) NMR spectra of 1 have no significant concentration dependencies, with the exception of some broadening that occurs at high concentrations (SI Appendix, Fig. S30).

DOSY was used to probe the dimensions of the polydispersed supramolecular aggregates (14, 19). As the concentration of rhomboid 1 was increased between 10.0 and 90.0 mM, the measured weight average diffusion coefficient decreased from $1.07 \times 10^{-10}$ to $5.83 \times 10^{-12}$ m$^2$s$^{-1}$ ($D_{10.0 \text{mM}}/D_{90.0 \text{mM}} \sim 19$; Fig. 4A, b–g), indicating a concentration dependence on the polymerization of 1 to give high-molecular-weight structures. On the DOSY NMR time scale, no evidence for low-molecular-weight cyclic oligomer formation was observed (SI Appendix, Figs. S32–S37), further supporting the utility of rigid metallacyclic cores in producing high fidelity polymers. The supramolecular entities have narrow dispersities at the concentration ranges of 1 used here. The DOSY NMR of unfunctionalized rhomboid 15 was also performed using a 20.0-mM solution to yield an average diffusion coefficient of $3.94 \times 10^{-10}$ m$^2$s$^{-1}$ (Fig. 4A, a; SI Appendix, Fig. S31), further supporting that the UPy moieties are responsible for aggregate formation in CD$_2$Cl$_2$. Moreover, dynamic light scattering (DLS) measurements were conducted to

![Fig. 1. Cartoon representations for the formation of a linear supramolecular polyrhomboid (A) and cross-linked three-dimensional supramolecular polymeric networks (B) from self-assembly of rhomboid 1 and hexagon 2 or 3, respectively.](image)
study the size distributions of the polymeric material with respect to the concentration of 1 (Fig. 4B). At the concentration of 10.0 mM, polymerized 1 showed an average hydrodynamic diameter ($D_h$) of 100.2 nm. The value of $D_h$ was concentration dependent, increasing at higher molarities, indicating that larger aggregates are formed when more 1 is present. Given the ~5-nm size of rhomboid 1 as predicted by the force-field calculations, the measured average $D_h$ obtained by DLS suggest that there are ~41 rhombic units in the linear polymeric chain at 10.0 mM. Although the polymeric material comprised of rhomboid 1 was sufficiently soluble to allow for the solution-based characterization methods described above, rod-like fibers could be mechanically drawn from high viscosity CH$_2$Cl$_2$ solutions. These fibers were observed using SEM (SI Appendix, Fig. S44), providing evidence for the formation of a polymeric material formed via linkage of the rhomboids into a linear chain. Furthermore, transmission electron microscopy (TEM) was used to provide further insight into the hierarchical assembly of the polymer material. As shown in Fig. 5C, samples of 1 ultimately organize into long fibers with a diameter of ~300 nm. These fibers represent bundles of single strands of linear rhomboid chains that rapidly form in solution as the UPy moieties undergo hydrogen bonding. Once formed, these single polymeric chains aggregate in solution, entangling together to form associated clusters which ultimately self-assemble into tightly packed nanoscale fibers (Fig. 5D).

Subsequently, we used UPy-functionalized hexagons 2 and 3 to prepare cross-linked supramolecular hexagonal networks (Fig. 1B) with the aim of exploiting their material properties benefiting from the introduction of organoplatinum(II) metallacycles. However, solution-based characterization techniques could not be used to investigate the polymerization of UPy-functionalized hexagons 2 and 3 due to the insoluble materials that formed on introduction of solvents that facilitated hydrogen bonding. The anticipated cross-linked polymer networks (Fig. 1B) expected for these hexagonal metallacycles could instead be studied by FTIR methods, which indicated bands associated with N-H hydrogen bond formation and characteristic signatures of the ethynyl moieties of ligand 4 (SI Appendix, Figs. S39–S41). Despite the lack of solubility of polymeric 2 and 3 in dichloromethane, the introduction of this solvent resulted in the formation of gel-like soft matter due to swelling (Fig. 5B). The swelling properties of cross-linked polymeric materials are well established, and the observation of such a phenomenon with 2 and 3 supported the formation of extended hexagonal networks. Swelling tests based on the sample weight before and after swelling indicated that the network resulting from 3 exhibited a larger effect than that of hexagon 2 (SI Appendix, Figs. S42 and S43).

Attempts to draw out macroscopic fibers from the swelled samples resulted in the formation of long, flexible fibers (Movie S1). SEM imaging established that fibers obtained from gels of 2 had diameters of ~8 µm (Fig. 5D). As a testament to the stretchability, flexibility, and strength of the resulting material, one such fiber, obtained from a gel of 3, was readily fashioned into an overhand knot. The knot structure persisted over the course of 24 h, with no cracking or agglomeration. The combination of stretchability, strength, and flexibility associated with the formation of such a structure deriving from multicomponent self-assembly based on low-molecular-weight building blocks is noteworthy and illustrates the promising characteristics of SCC-based supramolecular polymers, which are reminiscent of the skeletal muscle protein titin, which possesses a remarkable combination of strength, toughness, and elasticity (41). The observed properties of the cross-linked hexagonal networks are attributed in part to the efficient polymerization that can occur when rigid metallacyclic cores are present. These cores preorient their pendant UPy groups for the facile formation of hexagonal networks. On mechanical stress, which can potentially rupture both H-bonding and the SCC cores, both the metal–ligand coordination and UPy dimerization can readily reform due to the reversibility of both processes. These features allow the material to adapt to physical deformations by stabilizing new orientations of the precursors through intermolecular bond formation, ultimately increasing the robustness of the materials. Even if mechanical stress distorts the cores such that hexagons cannot reform, the multtopic nature of the precursors ensures that some cross-linking can persist, thereby increasing the cross-link density and strengthening the new 3D network.

**Conclusion**

In summary, by combining the directional-bonding approach of coordination-driven self-assembly and the quadruple hydrogen bonding of UPy moieties, we established a method for hierarchical supramolecular polymerization. The tunability associated with SCC formation allowed for the formation of both a rhomboid and two hexagons from a single UPy-functionalized dipyr- idyl ligand by selecting specific metal acceptors. The hierarchical supramolecular polymerization of this ligand furnished discrete SCCs decorated with UPy groups as a first level of complexity. The complexity was then amplified on hydrogen bonding of the UPy groups to link the metallacycles into either a linear chain (rhomboid) or cross-linked networks (hexagons). Characterization

![Figure 2](image-url)  
*Fig. 2.* $^{31}$P ($^1$H) NMR spectra (room temperature, 121.4 MHz) of (A) 60° acceptor 5 in CD$_2$Cl$_2$, (B) supramolecular polyrhomboid in CD$_2$Cl$_2$ at a concentration of 40.0 mM, (C) rhomboid 1 in CD$_2$OD, (D) hexagon 2 in DMSO-d$_6$, and (E) hexagon 3 in DMF-d$_7$.

![Figure 3](image-url)  
*Fig. 3.* Partial $^1$H NMR spectra of rhomboid 1 (CD$_2$Cl$_2$, room temperature, 300 MHz) at different concentrations: (A) 110 mM; (B) 78.0 mM; (C) 60.0 mM; (D) 40.0 mM; (E) 20.0 mM; (F) 10.0 mM; (G) 5.00 mM; (H) 1.00 mM; and (I) 120° UPy-functionalized ligand 4 in CD$_2$Cl$_2$. Peaks of linear polymer are designated by lin.
of the resulting materials revealed that the polymerizations were particularly effective in avoiding the formation of low-molecular-weight cyclic oligomers, a property attributed to the presence of the rigid metallocyclic cores. In addition, the solvent swelling formation of gels observed for the hexagonal networks lent themselves to the formation of long, macroscopic fibers that possessed enough strength and flexibility to permit the construction of stable knots. The robustness of these fibers is consistent with heightened cross-link densities that occur due to the reversibility of the metal–ligand and hydrogen bonding that form the material. Deformations that can permanently sever polymer strands in other materials can reform in these gel-like soft matters, restoring intermolecular cross-linking and allowing the fibers to adapt to stresses. Given the vast library of readily functionalized SCCs and favorable properties of supramolecular polymers established here, the hierarchical polymerization of metallocycles is a promising new method to obtain novel functional materials.

Materials and Methods

All reagents were commercially available and used as supplied without further purification. Deuterated solvents were purchased from Cambridge Isotope Laboratory. Compounds 5 (42), 7 (32), 8 (43), 14 (39), and parent rhomboid 15 (44) were prepared according to published procedures. NMR spectra were recorded on a Varian Unity 300 MHz spectrometer. 1H and 13C NMR chemical shifts are reported relative to residual solvent signals, and 31P (1H) NMR chemical shifts are referenced to an external unlocked sample of 85% H3PO4 (δ 0.0). The 2D diffusion-ordered (2D DOSY) NMR spectra were recorded on a Bruker DRX500 spectrometer. DLS was carried out on a Malvern Nanosizer S instrument at room temperature. FTIR spectra were recorded on a NEXUS-47 FTIR spectrometer. SEM investigations were carried out on a JEO L6390VL instrument. TEM images were obtained using a Philips TEM/TECNAI-12 instrument with an accelerating voltage of 120 kV. Mass spectra were recorded on a Micromass Quattro II triple-quadrupole mass spectrometer using electrospray ionization with a MassLynx operating system. The melting points were collected on a SHPSIC WRS-2 automatic Laboratory. Compounds 5a, 5b, 5c, 5d, 5e, and 5f were added to the ligand 1 in a 2:1 molar ratio. The solids were dissolved in methanol (for 1) or DMSO (for 2 and 3) and allowed to stir at 50 °C for 8 h. To the resulting homogeneous solution, diethyl ether was added to precipitate the product, which was then isolated and dried under reduced pressure for 4 h and redisolved in methanol or DMSO for characterization.

Rhomboid 1. 1H NMR (CD2OD, room temperature, 300 MHz) δ (ppm): 9.02 (d, J = 5.7 Hz, 4H), 8.93 (d, J = 6.0 Hz, 4H), 8.67 (s, 4H), 7.94–8.00 (m, 4H), 7.89–9.00 (m, 4H), 7.70 (s, 4H), 7.65 (s, 2H), 7.62–7.64 (m, 4H), 7.58–7.61 (m, 4H), 7.33 (d, J = 1.2 Hz, 4H), 5.85 (s, 2H), 4.12 (t, J = 6.3 Hz, 4H), 2.26–2.42 (m, 2H), 1.80–1.96 (m, 4H), 1.53–1.73 (m, 3ZH), 1.35–1.52 (m, 4ZH), 1.10–1.30 (m, 7ZH), 0.76–0.93 (m, 12ZH). 31P (1H) NMR (CD2OD, room temperature, 121.4 MHz) δ (ppm): 11.77 ppm (δ 2H, Pt satellites, 2JPH = 2.67,6.4 Hz). ESI-TOF-Ms: m/z 834.85 [M − 4NO3]+, 1,112.81 [M − HNO3 − 3NO3]2+, 1,124.79 [M − 2HNO3 − 2NO3 + K]3+, 1,133.80 [M − 3NO3]3+, 1,731.68 [M − 2NO3]3+.

Hexagon 2. 1H NMR (DMSO-d6, room temperature, 300 MHz) δ (ppm): 11.41 (s, 4H), 9.55 (s, 3H), 8.85 (d, J = 5.7 Hz, 12H), 7.81 (d, J = 5.4 Hz, 12H), 7.74 (s, 3H), 7.52 (s, 3H), 7.39 (d, J = 5.1 Hz, 6H), 6.97–7.12 (m, 12H), 5.71 (s, 3H), 3.97–4.14 (m, 6H), 3.09–3.19 (m, 6H), 1.65–1.90 (m, 72H), 1.29–1.56 (m, 51H), 0.95–1.18 (m, 108H), 0.67–0.85 (m, 18H). 31P (1H) NMR (DMSO-d6, room temperature, 121.4 MHz) δ (ppm): 13.15 ppm (δ 19Pt satellites, 3JPH = 2.309.5 Hz). ESI-TOF-Ms: m/z 808.94 [M − 6OTf]1+, 818.34 [M − 2OTf− + 4OTf− + Na + K]2+, 970.41 [M − 2OTf− − 3OTf− + 2NaH]2+, 1,000.40 [M − 2OTf−]2+, 1,212.50 [M − 2OTf− − 4OTf−]3+, 1,288.23 [M − 4OTf−], 1,628.65 [M − 4OTf− + 2OTf + K]3+.

Hexagon 3. 1H NMR (DMSO-d6, room temperature, 300 MHz) δ (ppm): 11.40 (s, 4H), 9.55 (s, 3H), 8.85 (d, J = 5.7 Hz, 12H), 7.84 (d, J = 5.4 Hz, 12H), 7.49–7.59 (m, 18H), 7.36–7.48 (m, 18H), 7.50 (s, 3H), 3.97–4.14 (m, 6H), 3.09–3.19 (m, 6H), 1.64–1.81 (m, 6H), 1.39–1.56 (m, 45H), 1.21–1.38 (m, 72H), 0.94–1.13 (m, 108H), 0.68–0.83 (m, 18H). 31P (1H) NMR (DMSO-d6, room temperature, 121.4 MHz) δ (ppm): 10.01 ppm (δ 19Pt satellites, 3JPH = 2.63,6,9 Hz). ESI-TOF-Ms: m/z 836.85 [M − 6OTf]2+, 846.53 [M − 2HNO3 − 4NO3 + Na + K]3+, 1,004.03 [M − HNO3 − 3NO3]2+, 1,011.42 [M − 2HNO3 − 4NO3 + K]3+, 1,016.43 [M − 5NO3]3+, 1,254.53 [M − 2HNO3 − 4NO3]2+, 1,263.53 [M − 3HNO3 − 3NO3 + K]3+, 1,270.54 [M − 4HNO3 − 2NO3 + Na + K]3+.

In a 1:1 stoichiometric fashion, 60% or 120% bis(phosphine) organoplatinum (II) acceptor 5, 6, or 7 was added to the ligand 1 in a 2:1 ratio of vinyl. The solids were dissolved in methanol (for 1) or DMSO (for 2 and 3) and allowed to stir at 50 °C for 8 h. To the resulting homogeneous solution, diethyl ether was added to precipitate the product, which was then isolated and dried under reduced pressure for 4 h and redisolved in methanol or DMSO for characterization.
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