

NIH Public Access

Author Manuscript

Tetrahedron. Author manuscript; available in PMC 2009 December 8.

Published in final edited form as:

Tetrahedron. 2008 December 8; 64(50): 11495–11503. doi:10.1016/j.tet.2008.08.062.

Carbon-rich supramolecular metallacycles and metallacages

Brian H. Northrop* , **Dennis Chercka**, and **Peter J. Stang**

University of Utah, Department of Chemistry, 315 So. 1400 E. Salt Lake City, UT 84112

Abstract

Coordination-driven self-assembly via the directional-bonding approach utilizes rigid transition metal acceptors and electron-rich donors to allow for complex, nanoscale 2D polygons and 3D polyhedra to be prepared under mild conditions and in high yields. To ensure proper rigidity and directionality, many acceptor and donor precursors contain largely carbon-rich aromatic and/or acetylenic moieties. This article introduces self-assembly as an alternative means of synthesizing carbon-rich materials and discusses the development, design, synthesis, and applications of carbonrich supramolecular metallacycles and metallacages as well as the self-assembly of new diastereomeric carbon-rich supramolecular triangles.

1. Introduction

Carbon-rich materials¹ have attracted the interest and attention of chemists ever since the extraction and identification of polycycles during petroleum analysis began in the late 1800's. In the intervening years, seminal studies, led initially by Glaser, Scholl, and Clar, have transformed the science of carbon-rich materials from one primarily of analysis to one of planned preparation.^{2,3} Whereas it was once a distinct challenge to synthesize even a relatively simple polycyclic aromatic hydrocarbon (PAHs) such as naphthalene – a challenge originally overcome by Radziszewski⁴ in 1878 – it is now possible to carry out a chemical synthesis of such highly complex PAHs as C_{60} , as was done by Scott⁵ in 2001. Even so, as the field of carbon-rich molecules expands further, continually branching out into new areas with new applications, there is a persistent need for new and innovative ways of preparing these important compounds. Recent advances in coordination-driven self-assembly have provided such a means and have enabled a wide variety of carbon-rich metal-organic supramolecules to be prepared.

Over the past 150 years a variety of methods have been devised for preparing carbon-rich compounds. Originally, most techniques relied upon fairly harsh conditions typically requiring high temperatures and/or strong oxidizing agents and often resulted in low yields. Milder, higher yielding methods have since been developed such as Clar's Zn-dust reductions,² oxidative couplings of terminal alkynes,⁶ Scholl-type intramolecular cyclodehydrogenation, 7 *hv*-induced ring closures, 8 cyclodehydration, 9 and Diels-Alder cycloaddition, 10 just to name a few. High temperature flash vacuum pyrolysis (FVP), however, remains a very popular means of preparing carbon-rich molecules despite its often-poor yields. Self-assembly, on the other hand, is a synthetic technique that is typified by mild reaction conditions and high yields.

^{© 2008} Elsevier Ltd. All rights reserved.

^{*}B. H. Northrop. Tel.: 801-581-8011; fax: 801-581-8433; e-mail: b.northrop@utah.edu.

Publisher's Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

1.1. Coordination-Driven Self-Assembly of Carbon-Rich Supramolecules

Molecular self-assembly¹¹ is a process through which properly designed, complementary molecular subunits spontaneously assemble according to the specific noncovalent "information" encoded within their structures. The use of noncovalent interactions (e.g. hydrogen-bonding, π-π, donor-acceptor, charge-charge, etc.) to direct self-assembly enables product formation to be thermodynamically controlled. Individual building-blocks are able to continually and reversibly assemble in a variety of ways until the most thermodynamically stable product is obtained. More often than not, the result of such dynamic¹² self-assembly processes is one highly symmetric, thermodynamically stable product in high and often quantitative yield. With appropriate planning, individual precursor building-blocks may be designed such that they lead specifically to a single desired supramolecule possessing unique architectural or topological¹³ characteristics. While many examples of self-assembled supramolecules involve flexible, significantly saturated carbon backbones11c,d,f and, thus, may not fit the typical definition of being carbon-rich compounds, there are also myriad ways of using the virtues of self-assembly (high yield, mild conditions, complex architectures) to prepare carbon-rich molecules.

As a design and synthesis methodology, coordination-driven¹⁴ self-assembly is especially useful for preparing carbon-rich molecules. Coordination-driven self-assembly relies upon the use of dative metal-organic interactions between electron-poor transition metal acceptors and electron-rich organic donors and is often used in conjunction with the directional-bonding¹⁵ approach to self-assembly. According to the directional-bonding protocol, discrete supramolecular assemblied will be formed in accordance to the specific geometric information – the "directionality" – stored within the pre-defined angles and symmetry of rigid precursors and the stoichiometric ratio in which those precursors are mixed. Rigid acceptor and donor precursors are necessary so that they retain their directionality, increasing the likelihood of a successful self-assembly, i.e. one that results in a high yield of a single discrete product as opposed to a mixture of polymeric, oligomeric, or other side products. With these design criteria in mind, the vast majority of examples that follow the directional-bonding approach to coordination-driven self-assembly utilize largely aromatic and alkynyl building-blocks. The resulting self-assembled metal-organic supramolecules are often quite carbon-rich.

1.2. Carbon-Rich Polygons and Polyhedra

The use of coordination-driven self-assembly in the synthesis of carbon-rich compounds has many advantages. When precursors are carefully designed it is possible to self-assemble 2D metallacyclic polygons and 3D metallacage polyhedra while exerting precise control over the often highly-complex structures of the supramolecular products. Representative collections of 2D and 3D metallacycles that can be prepared by the directional-bonding approach are shown in Figure 1 and Figure 2, respectively. It would likely prove exceedingly difficult, if not altogether impossible, to prepare analogous structures in high yields by traditional synthetic methods. Dynamic self-assembly, on the other hand, as a thermodynamically controlled process, enables such complex structures to be prepared in significantly fewer synthetic steps and often in near quantitative yields. Preparation of these highly symmetric polygons and polyhedra is very modular, enabling both their architecture and properties to be easily manipulated simply by changing the size, shape, coordination number, and/or functionality of individual building units.

2. Metallacyclic Polygons

While earlier examples of metal-organic self-assembled metallacages exist, most notably from Maverick and coworkers¹⁶ in 1984, it wasn't until about 1990 that systematic approaches to metallacycles began to appear.17 Most initial examples of self-assembled metallacycles took

the form of supramolecular squares. One of the most commonly utilized protocols for designing a supramolecular square calls for a combination of four linear ditopic units $(L²)$ and four complementary angular ditopic units (A^2) , generating a square of type $A^2_4L^2_4$. Square planar $Pt(II)$ and Pd(II) metals have provided the basis for 90 $^{\circ}$ angular corner units of myriad carbonrich supramolecular squares in conjunction with rigid pyridyl or nitryl substituted linear donors. $18-22$ Example carbon-rich angular and linear components are shown in Figure 3, while Scheme 1 summarizes the self-assembly of a number of $A^2_4L^2_4$ supramolecular squares.

It is also possible to form related squares of type $\rm A^2_2A^2_2$ wherein two sets of ditopic 90° angular units, one donor and one acceptor, are combined in a 1:1 ratio. 23

By employing ditopic donor and/or acceptor building-blocks with turning angles that deviate from 90°, a variety of other carbon-rich 2D metallacycles can be prepared (Figure 1). Representative examples of ditopic acceptors and donors used in the construction of carbonrich supramolecular rectangles, $24-26$ rhomboids, $26-28$ triangles, 29 pentagons, 30 and hexagons27–28,31–32 are shown in Figure 4. As can be seen in Scheme 2, the variety of complementary angular and linear ditopic building units provides access to a series of metallacyclic polygons.

Rigid supramolecular rectangles can be prepared upon mixing the 0° molecular "clip" with linear donor ligands such as **4**, **10**, **11**, and **13**. Given the close proximity of the two Pt(II) acceptors in **14**, care must be taken when designing a linear donor ligand because of steric crowding. Given that a rhomboid can be defined as any four sided parallelogram that is neither a square nor a rectangle, a variety of donor and acceptor combinations may be used to prepare metallacyclic rhomboids. Carbon-rich supramolecular rhomboids have thus been prepared using 180° donor bis(pyridyl)-substituted perylene diimide **13** as well as 120° donors **21–23**. It is possible to form a rhomboid structure using the linear donor **13** because its length, \sim 3.5 nm, confers enough overall flexibility to self-assemble when combined with a 90° metal acceptor.

Combining three ditopic 60° phenanthrene-based Pt(II) acceptors (**15**) with three ditopic 180° 4,4'-bipyridyl linear donors (**4**) results in a metallacyclic triangle.29 Changing the acceptor unit from a 60° phenanthrene-based **15** to a 120° benzophenone-based Pt(II) acceptor (**17**) results in a carbon-rich supramolecular hexagon when mixed with the same linear donor (**4**) in a 1:1 molar ratio.³¹ The modularity of the coordination-driven directional-bonding approach is easily seen when comparing some of the many carbon-rich metallacyclic hexagons that have been self-assembled. Hexagons of type $A^2 A^2$, devoid of linear building blocks, have been prepared²⁷ as have large hexagons of type $A^2{}_6L^2{}_6$ wherein complementary pairs of 120° acceptors and 180° donors and analogous 120° donors and 180° acceptors have both been used. Pentagonal metallacycles may be formed by using a slightly more flexible 109° acceptor, however, they are often in competition with related hexagonal metallacycles.

3. Metallacage Polyhedra

More recently, considerable attention has been paid to expanding these coordination-driven self-assembly techniques to the 3rd dimension. The protocol for preparing 3D metallacages only requires the preparation of donor and/or acceptor precursors with greater than two binding sites. As shown in Figure 2, combining carbon-rich tritopic and ditopic building units allows for the construction of metallacages with truncated tetrahedral, distorted and non-distorted trigonal prism, cuboctahedral, adamantoid, and dodecahedral geometries (Scheme 3).14b,c,e Combining carbon-rich, 3-fold symmetric planar tritopic donors or acceptors (Figure 5) with complementary 90° units in a 3:2 ratio allows the self-assembly of truncated tetrahedral assemblies.^{33–35} In addition to these carbon-rich examples, Fujita and coworkers have also prepared related nitrogen-rich analogues based on tritopic trispyridyl donors with a

triazabenzene core.³⁶ If the self-assembly involving donors 25 or 26 is performed with a 0° molecular "clip" rather than a 90° acceptor, also in a 3:2 ratio, the result becomes a trigonal prism.37,38 An alternative method for preparing a carbon-rich trigonal prism has been achieved by Fujita using a combination of two tritopic trigonal planar donors, three ditopic linear donors, and six 90° metal acceptors.³⁹

Other trigonal prism geometries may be formed using three carbon-rich tetratopic donors that function as planar "walls" that are linked using six 90° metal acceptors as "corners" as shown in Scheme 4. Tetrapyridyl-substituted porphyrins⁴⁰ as well as tetrapyridyl-substituted benzene and cyclobutadiene-cyclopentadieneyl cobalts⁴¹ have been used as the carbon-rich walls of such trigonal prism geometries. In the case of tetratopic donor **34**, restricted rotation of pyridyl ligands in a trigonal prism renders the assembly helically chiral with MMM and PPP enantiomers.

An entirely different geometry, that of a cuboctahedron, is obtained when planar tritopic units such as carbon-rich donor **25** or acceptor **28** are combined in an 8:12 ratio with complementary 120° ditopic units (Scheme 5).⁴² Robson and coworkers⁴³ have self-assembled a metallacage with cuboctahedral geometry by combining a 2,4,6-triazophenyl-1,3,5-trioxybenzene trianion and Cu(NO₃)₂ in a 2:3 molar ratio. By using "naked" Pd metals in conjunction with 120° dipyridyl donors, Fujita and coworkers⁴⁴ have prepared a variety of cuboctahedral complexes of the form $M_{12}L_{24}$. The large variety of carbon-rich 3D architectures available simply using planar triropic of tetratopic units with 0°, 90°, or 120° units – resulting in assembly geometries that range from truncated tetrahedral, to trigonal prisms, to cuboctahedra, respectively – demonstrates the power and modularity of the directional-bonding, coordination-driven approach.

Using non-planar tritopic donors (Figure 6) opens the door to an even greater variety of 3D metallacages. Combining trigonal pyramidal tritopic donors with 0° or 90° acceptors in a 2:3 molar ratio results in the formation of distorted trigonal prisms^{45–46} or, more precisely, trigonal bipyramids (Scheme 6). The extent of distortion from a "planar" trigonal prism geometry is controlled by the size and geometry of the tritopic donor and the angle of the acceptor, with a greater acceptor angle resulting in greater distortion from a trigonal prism geometry.

Fujita has self-assembled a "double-square" metallacage⁴⁷ by combining four equivalents of a tritopic trispyridyl methyl donor derivative with six equivalents of a 90° Pd(II) acceptor. Whereas a 90° Pd(II) acceptor in the previous case is used to prepare a double-square, a 109– 120° Pt(II) can be combined with a related tritopic trispyridyl methyl derivative (**35**) to selfassemble an adamantoid structure (Scheme 6).⁴⁸ We have also prepared a variety of carbonrich dodecahedral assemblies using a trispyridylmethanol tritopic donor and different sized 1,4-substituted benzene and 4,4'-biphenyl substituted Pt(II) linear (180°) ditopic acceptors.⁴⁹ Combining the tritopic donors and ditopic acceptors in a 30:20 ratio results in carbon-rich supramolecular metallacages with dodecahedral geometries. These dodecahedra, with diameters ranging from 5–7 nm, represent some of the largest discrete supramolecular assemblies prepared to date.

4. Materials Applications

As with many other carbon-rich molecules, which have found applications in materials such as organic field-effect transistors (OFETs), artificial photosynthetic systems, and luminescent devices, the potential materials applications of carbon-rich supramolecular assemblies have also been explored. Our phenanthroline-containing supramolecular rectangle, 25 for example, has been shown to function as an optical sensor for $Ni(II)$, $Cd(II)$, and $Cr(III)$. Fujita and coworkers have demonstrated the ability of a carbon-rich supramolecular tetrahedron cage to

selectively recognize and bind sequence-specific peptides⁵⁰ and, in the case of a supramolecular trigonal prism, capable of folding an Ala-Ala-Ala tripeptide in to a β-turn on account of hydrophobic encapsulation.⁵¹ The use of largely aromatic moieties in various precursors enables many supramolecular cages to act as hosts for other aromatic guests through a combination of π -π and hydrophobic interactions.^{39,52} Both Hupp⁵³ and Mirkin⁵⁴ have recently used porphyrin-based supramolecular rectangles as catalysts for epoxidation and acyl transfer reactions, respectively.

4.1. Microporous Molecular Materials

In addition to aforementioned host-guest applications, rigid supramolecular 2D polygons offer unique opportunities to be used as highly-selective nanofilters. The size and shape of nanoscale cavities present in metallacyclic polygons can be easily modified in order to allow for sizespecific discrimination of various molecular permeants. Hupp and coworkers⁵⁵ have extensively explored the use of metal-organic supramolecules as selective nanofilters. Small and large supramolecular squares, for example, composed of $Re(I)Cl(CO)$ ₃ corners and either pyrazine or dipyridyl porphyrin edges, respectively, were shown capable of filtering planar and spherical permeants based upon their size.⁵⁶ Further studies utilizing thin-films of $Mn(I)$ and Re(I) carbon-rich rectangles demonstrated that only those molecular permeants that were narrow in at least one dimension, similar to the metallacyclic rectangles, were able to pass through the intramolecular rectangular cavities whereas permeants lacking a narrow dimension were blocked.⁵⁷ In addition to applications as highly-modular nanofilters, the gas adsorption properties of carbon-rich supramolecular metal-organic assemblies are currently being explored.⁵⁸

4.2. Higher-Order Self-Assembly on Surfaces

Recently, in collaboration with Wan and coworkers, we have explored the self-assembly of supramolecular assemblies on $Au(111)$ and HOPG surfaces. Both polygonal (rectangular⁵⁹) and square⁶⁰) and polyhedral (trigonal bipyramid⁶⁰ and chiral trigonal prism⁶¹) structures have been assembled onto surfaces and their structures characterized. Results of these studies have shed considerable light on the factors that influence the orientation of these supramolecules on various surfaces. For example, one carbon-rich supramolecular rectangle investigated adopts an edge-down orientation on HOPG while the same rectangle adopts a face-down orientation on Au(111).⁵⁹ In the case of chiral trigonal prisms, deposition of a racemic mixture of supramolecular assemblies resulted in spontaneous symmetry-breaking and formation of enantiomerically-segregated chiral domains.⁶¹ Furthermore, we have demonstrated the ability of using a molecular template⁶² to promote the formation of a well-ordered, monodisperse monolayer of molecular rectangles on Au(111) whereas without the template only a disordered adlayer is observed. Having the ability to prepare well-ordered arrays of these carbon-rich supramolecules and control their orientation on surfaces may lead to a variety of applications as sensors, electronic, photonic, or magnetic materials, or catalysis, especially in the case of chiral 2D assemblies which may have a range of heterogeneous stereoselective synthetic applications.

4.3. Functionalized Metal-Organic Assemblies

Another development in the use of carbon-rich supramolecular metallacycles and metallacages has been in their functionalization. The rigid structures and precise, unique geometries of many supramolecular assemblies makes them very effective "scaffolds" for a variety of functionalities simply by preparing derivatized donors and/or acceptors. Given the rigid structural characteristics of precursor building-blocks and the fact that donors and acceptors are mixed in specific ratios, the location and stoichiometry of covalently linked functional moieties attached to carbon-rich aupramolecules can be precisely controlled. We have recently

prepared 120° donors functionalized with Frechét-type dendritic, 63 crown-ether, 64 and ferrocenyl65 moieties attached. Combining these donors with complementary 90°, 120°, or 180° Pt(II) acceptors allows for the formation of bisfunctionalized rhomboids as well as trisand hexakisfunctionalized hexagons, respectively. We have further demonstrated the ability of crown-ether derivatized assemblies to act as hosts for multiple dibenzylammonium cations64 and have explored the electrochemistry of ferrocenyl metallacycles, which may have applications as multiple electron redox catalysts or as electron sponges.65 Fujita and coworkers66 have also successfully prepared carbon-rich 3D cuboctahedra that have been functionalized endohedrally with azobenzene,66a,c oligo(ethyleneglycol),^{66a} polyfluoro,^{66b} and polymerizable methyl methacrylate units^{66d} and exohedrally with saccharide moieties. 66e These 3D assemblies have found uses in a variety of host-guest applications.

5. Novel Stereoisomeric Supramolecular Triangles

Chirality is a ubiquitous and important phenomenon throughout nature in general and chemistry in particular. The general lack of tetrahedral carbon atoms throughout most carbon-rich compounds usually rules out the possibility of synthesizing carbon-rich molecules with point chirality. There are, however, ways to introduce chirality into carbon-rich molecules by means of helicity. For example, beginning in 1967, Martin and coworkers⁶⁷ prepared helically chiral hepta-, octa-, and nonahelicenes. Building upon these purely aromatic helically chiral structures, in 1972 Staab and coworkers⁶⁸ prepared a carbon-rich helically chiral benzannulene structure composed solely of benzene and acetylene moieties. Since that time a rich chemistry of chiral benzannulene compounds has been developed.⁶⁹ Similarly, chirality can be introduced into carbon-rich systems through the incorporation of chiral moieties, such as binaphthyl units.

Coordination-driven self-assembly techniques provide a means of synthesizing stereoisomers of carbon-rich supramolecules in high yield under thermodynamic control. In the mid-1990's, Stang and coworkers used a chiral binaphthyl-substituted 90° Pt(II) acceptor to prepare optically active carbon-rich 2D metallacyclic squares²¹ and 3D metallacage truncated tetrahedra.33 It is also possible to use self-assembly to synthesize carbon-rich supramolecules from completely achiral building blocks. To demonstrate the utility of coordination-driven selfassembly we report herein the design and synthesis of stereoisomeric carbon-rich metallacyclic triangles.

The combination of 2,6-diazaanthracene (**6**) and the 60° Pt(II) acceptor **15** in a 1:1 ratio results, theoretically, in the formation of two possible $A^2 {}_3L^2 {}_3$ metallacyclic triangle diastereomers (Scheme 7), which are both meso compounds. The diastereomers differ in the relative orientations of their donor components as indicated by arrows in the Scheme. In the symmetric diastereomer, all three donor moieties are oriented in the same direction while in the asymmetric diastereomer, one donor moiety is oriented opposite the other two. Rotation around the Pt-N bonds in each assembly is restricted, $\frac{70}{10}$ preventing direct interconversion between stereoisomers through dynamic bond rotations.

Stirring 6 and 15 in dry CH_3NO_2 for 2 h results in a homogeneous yellow solution. Following the exchange of the NO₃^{$-$} counterions for weakly coordinating PF₆^{$-$} counterions the formation of the new metallacyclic triangles was initially investigated by ${}^{31}P$ and ${}^{1}H$ NMR spectroscopies. Analysis of the carbon-rich triangles with ${}^{31}P[{^1}H]$ NMR revealed the presence of a sharp signal around 13.7 ppm, indicating the formation of discrete supramolecules as opposed to oligomeric species. Close examination of the sharp signal, however, revealed that it is in fact composed of two peaks: one high intensity signal at 13.69 ppm and another lower intensity signal at 13.59 ppm (Figure 7). The observation of two peaks indicates that both carbon-rich diastereomers are present in solution, though more of one stereoisomer is formed than the other. Both peaks are upfield shifted by about 5.3 ppm as compared to the starting 60°

acceptor **15** due to back donation from the platinum atoms. Further evidence of back donation and Pt-N coordination was also observed by the decrease in coupling of the ¹⁹⁵Pt satellite peaks $(\Delta J = 109 \text{ Hz})$. Examination of the ¹H NMR spectrum also revealed the presence of highly symmetric species, though proton signals were broadened slightly as a result of the presence of two different stereoisomers. It was further observed that hindered Pt-N rotation gives rise to in equivalent signals for donor **6**, the four protons that point inward with respect to the core of the supramolecular triangle display are shifted slightly relative to the four that point outward (see Supporting Information). Further confirmation of the formation of discrete supramolecular triangles was provided by Electro-Spray Ionization mass spectrometry (ESI-MS). A peak was found at $m/z = 986.8$, corresponding to the $[M - 4PF_6]^{4+}$ peak of the carbon-rich $A^2 {}_3L^2 {}_3$ triangle, and the distribution of peaks matched their theoretical values (see Supporting Information) establishing the molecularity of the assembly.

The observation of two peaks of different intensity in the $31P$ NMR spectrum of the carbonrich supramolecular triangle can be rationalized through molecular modeling studies. The supramolecular structures of both diastereomers were built within the input mode of the program Maestro v8.01.10 and were subjected to a 1.0 ns molecular dynamics simulation (MMFF force field, 300 K, gas phase) in order to equilibrate their structures, followed by energy minimization to full convergence (Figure 8). The fully equilibrated and minimized structure of the "symmetric" supramolecular triangle, having all three donor moieties oriented in the same direction, was found to be 6.9 kcal/mol more stable than the asymmetric supramolecular triangle. This difference in stability likely arises from differences in strain energy induced by the relative orientations of the donor moieties in the two diastereomers. In the asymmetric diastereomer two of the three donors are oriented opposite to each other, increasing the structural strain in the supramolecule and causing the asymmetric triangle to deviate from a perfectly triangular structure as shown in Figure 8b. The aligned arrangement of the three donor moieties in the symmetric triangle distributes and equalizes any strain, resulting in a more stable, and more triangular, structure (Figure 8a). The differences in structural strain can be seen, in part, in comparing the distances between neighboring phenanthrene units as measured from molecular modeling: all three distances are 1.46 nm in the symmetric diastereomer while in the asymmetric diastereomer they range from 1.43–1.49 nm. Due to the energetic difference between the two diastereomeric carbon-rich triangles it is likely that the symmetric triangles are formed preferentially, giving rise to the ^{31}P NMR signal of higher intensity, while the less stable asymmetric triangles can be accounted for by the ^{31}P NMR signal of lower intensity.

These results demonstrate how self-assembly may be used to prepare different stereoisomers of carbon-rich supramolecular assemblies from achiral precursors. Though not the first examples of chiral carbon-rich molecules or supramolecules, the preferential formation of one stereoisomer at the expense of another indicates that it may be possible to prepare enantiomerically pure carbon-rich supramolecular assemblies by coordination-driven selfassembly. In addition to structural strain it may be possible to use other driving forces such as electrostatics to bias the formation of one desired stereoisomer. Having the ability to prepare novel, enantiomerically pure carbon-rich compounds in high yield under the mild reaction conditions of self-assembly is very desirable. Experiments aimed at such chiral discrimination during self-organizing self-assembly are currently underway.

6. Conclusions

In conclusion, these examples demonstrate many of the ways in which coordination-driven self-assembly can be used to prepare carbon-rich supramolecular 2D and 3D assemblies. The key characteristics of metal-ligand self-assembly – mild reaction conditions, high yields, thermodynamic control, and a high degree of modulatiry – combine to make the directional-

bonding approach to self-assembly a very valuable addition to the many other synthetic protocols available for the synthesis of carbon-rich materials. While carbon-rich molecular polygons and polyhedra have also been prepared via traditional covalent means – from triangles⁷¹ and rectangles⁷² composed purely of aromatic and acetylenic moieties, to molecules such as Kohnkene,⁷³ to Diederich's synthesis of expanded cubane⁷⁴ – the overall number of synthetic steps and yields of such synthesis often limit their practicality. Self-assembly allows access to related supramolecular systems at significantly lower synthetic cost. It has also been demonstrated that coordination-driven self-assembly can be used to prepare diastereomeric supramolecular triangles and that the structural characteristics of these metallacyclic systems are able to bias the formation of one stereoisomer over another. Such observations may lead to the development of methods that utilize self-assembly to form enantiomerically pure carbonrich supramolecules. Furthermore, the unique geometries and physical, electronic, and magnetic properties of these metal-organic structures opens a wide range of potential applications. It is likely that as their solution, solid-state, and surface-confined properties continue to be studied, and as new functionalized examples of this already important class of compounds continue to be investigated, researchers will find an ever widening number of applications in materials and nanosciences. They are certainly a welcome recent addition to the diverse world of carbon-rich materials.

7. Experimental

Self assembly of diastereomeric supramolecular triangles

The 2,6-diazaanthracene donor **6** and 60° Pt(II) acceptor **15** were prepared according to literature procedures.21,29 In a glass vial equipped with a magnetic stir bar was placed **15** (32.2 mg, 0.0277 mmol). In another glass vial was placed solid **6** (5.0 mg, 0.0277 mmol). Donor **6** was then dissolved in 0.5 mL dry CH_3NO_2 , transferred to the vial with acceptor 15, and then washed with additional $CH_3NO_2 (3 \times 0.5 \text{ mL})$ to ensure quantitative transfer. The suspension was stirred for 5 h at room temperature, resulting in a homogeneous clear yellow solution. The solvent was removed under reduced pressure, the NO_3^- counterions were exchanged for PF_6^- using a concentrated aqueous solution of KPF_6 , and the resulting precipitate was dried to completeness. Yield 39 mg (yellow solid), 93%. ¹H NMR (CD₃COCD₃, 300 MHz): δ 10.43 $(s, 3H, H_e)$, 10.42 $(s, 3H, H_e)$, 9.58 $(s, 3H, H_f)$, 9.54 $(s, 3H, H_f)$, 9.20-9.13 $(m, 6H, H_H)$, 9.03-8.93 (m, 6H, H_g), 8.61 (br, 6H, H_a), 7.86 (d, 6H, H_b, *J* = 7.7 Hz), 7.75 (d, 6H, H_c, *J* = 7.7 Hz), 7.71 (s, 6H, H_d), 1.42 (m, 72H, PCH₂CH₃), 1.19 (m, 108H, PCH₂CH₃). ³¹P{¹H} NMR $(CD_3COCD_3, 121.4 MHz)$:δ 13.7 (major), 13.6 (minor) ($^1J_{\text{Pt-P}} = 2662.7 \text{ Hz}$). MS (ESI) calcd for $[M - 4 \text{ PF}_6]^{4+} m/z$ 986.8, found 986.8.

The majority of synthetic procedures outlined in Scheme 1–Schemes 6 involve the selfassembly of pyridyl or nitryl donors and Pt(II) metal acceptors as either their OTf[−] or NO₃[−] salts. The following two synthetic self-assembly protocols, one for a Pt(II) triflate salt and one for a Pt(II) nitrate salt, serve as representative examples of the procedures used to prepare 2D and 3D supramolecules.

Self assembly of supramolecular rectangles.²⁴

In a glass vial equipped with a magnetic stir bar were placed solid 1,8-bis(*trans*-Pt (PEt3)2(NO3))anthracene (**14**) (20.0 mg, 0.0172 mmol) and an equimolar amount of the appropriate linear bridging ligand (e.g. 4 , 10 , 11 , 13). Next, 0.5 mL of CD_3COCD_3 and 0.5 mL of D₂O were added to the vial, which was then sealed with Teflon tape and heated in an oil bath at $50-55^{\circ}$ C, with stirring. After 10–15 h, the initial pale yellow suspension gradually turned bright orange and became homogeneous. The orange solution was then transferred to an NMR tube for analysis. The product was precipitated with KPF_6 , collected on a frit, washed with excess water, and dried in vacuo.

Self assembly of an A² 6L 2 ⁶ supramolecular hexagon.³⁰

In a glass vial equipped with a magnetic stir bar were placed solid bis(4-*trans*-Pt (PEt3)2(OTf)2)methanone (**17**) (20.0 mg, 0.0149 mmol) and 4,4'-bipyridyl (**4**) (2.3 mg, 0.0149 mmol). Next, 1.0 mL of CD_2Cl_2 was added to the vial, which was then stirred at 298 K for 30 min, resulting in quantitative formation (as observed by ¹H and ³¹P NMR spectroscopy) of the desired supramolecular hexagon.

For details of the synthetic procedures used in the preparation of **1–37** please see the original texts.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

The authors thank our co-workers and collaborators who have made this work possible. We are also grateful to Dr. Rik Tykwinski for guidance and materials. P.J.S. thanks the NIH (Grant GM-057052) and the NSF (Grant CHE-0306720) for financial support. B.H.N. thanks the NIH (Grant GM-080820) for financial support.

References

- 1. Haley, MM.; Tykwinski, RR., editors. Carbon-Rich Compounds. Weinheim: Wiley-VCH; 2006.
- 2. Clar, E. Polycyclic Hydrocarbons. Vol. Vol. I–II. London: Academic Press; 1964.
- 3. (a) Harvey, RG. Polycyclic Aromatic Hydrocarbons. New York: Wiley-VCH; 1997. (b) Modern Arene Chemistry. Weinheim: Wiley-VCH; 2002.
- 4. Radaziszewski B. Ber. Dtsh. Chem. Ges 1876;9:260–262.
- 5. (a) Boorum MM, Vasil'ev YV, Drewello T, Scott LT. Science 2001;294:828–831. [PubMed: 11679663] (b) Scott LT, Boorum MM, McMahon B, Hagen S, Mack J, Blank J, Wegner H, de Meijere A. Science 2002;295:1500–1503. [PubMed: 11859187]
- 6. (a) Glaser C. Ber. Dtsh. Chem. Ges 1869;2:422–424. (b) Baeyer A, Landsberg L. Ber. Dtsh. Chem. Ges 1882;15:50–56. 57–61. (c) Sondheimer F. Acc. Chem. Res 1972;5:81–91. (d) Siemsen P, Livingston RC, Dietrich F. Angew. Chem. Int. Ed 2000;39:2632–2657.
- 7. Kovacic P, Jones MB. Chem. Rev 1987;87:357–379.
- 8. Muszkat KA. Top. Curr. Chem 1981;88:89–143.
- 9. See, for example, Harvey RG, Pataki J, Cortez C, DiRaddo P, Yang C. J. Org. Chem 1991;56:1210– 1217.
- 10. See, for example, (a) Zander M. Chem. Ber 1959;92:2740–2743. (b) LeHoullier CS, Gribble GW. J. Org. Chem 1983;48:1682–1685.
- 11. (a) Lindesy JS. New J. Chem 1991;15:153–180.(b) Lehn, J-M. Supramolecular Chemistry. Weinheim: Wiley-VCH; 1995. (c) Rebek J Jr. Acc. Chem. Rec 1999;32:278–286. (d) Philip D, Stoddart JF. Angew. Chem. Int. Ed 2003;42:1210–1250. (e) Brancato G, Coutrot F, Leigh DA, Murphy A, Wong JKY, Zerbetto F. Proc. Natl. Acad. Sci. USA 2002;99:4967–4971. [PubMed: 11959948] (f) Yaghi OM, O'Keeffe M, Ockwig NW, Chae HK, Eddaoudi M, Kim J. Nature 2003;423:705–714. [PubMed: 12802325]
- 12. Huc I, Lehn J-M. Proc. Natl. Acad. Sci. USA 1997;94:2106–2110. [PubMed: 9122156]
- 13. Lukin O, Godt A, Vötgle F. Chem. Eur. J 2004;10:1878–1883.
- 14. (a) Stang PJ, Olenyuk B. Acc. Chem. Res 1997;30:502–518. (b) Leininger S, Olenyuk B, Stang PJ. Chem. Rev 2000;100:853–907. [PubMed: 11749254] (c) Seidel SR, Stang PJ. Acc. Chem. Res 2002;35:972–983. [PubMed: 12437322] (d) Schwab PFH, Levin MD, Michl J. Chem. Rev 1999;99:1863–1933. [PubMed: 11849014] (e) Gianneschi NC, Masar MS III, Mirkin CA. Acc. Chem. Res 2005;38:825–837. [PubMed: 16285706] (f) Cotton FA, Lin C, Murillo CA. Acc. Chem. Res 2001;34:759–771. [PubMed: 11601960] (g) Fujita M, Tominaga M, Hori A, Therrien B. Acc. Chem. Res 2005;38:369–378. [PubMed: 15835883] (h) Fiedler D, Leung DH, Bergman RG,

Raymond KN. Acc. Chem. Res 2005;38:349–358. [PubMed: 15835881] (i) Steel PJ. Acc. Chem. Res 2005;38:243–250. [PubMed: 15835871]

- 15. (a) Lehn, J-M. Supramolecular Chemistry Concepts and Perspectives. Weinheim: VCH; 1995. p. 139-160.(b) Chambron, J-C.; Dietrich-Buchner, C.; Sauvage, J-P. Comprehensive Supramolecular Chemistry. Lehn, J-M.; Atwood, JL.; Davies, JED.; MacNicol, DD.; Vötgle, F., editors. Vol. Vo. 9. Oxford: Pergamon; 1996. p. 43-83. (c) Holliday BJ, Mirkin CA. Angew. Chem. Int. Ed 2001;40:2022–2043. (d) Cotton FA, Lin C, Murillo CA. Acc. Chem. Res 2001;34:759–771. [PubMed: 11601960]
- 16. Maverick AW, Klavetter FE. Inorg. Chem 1984;23:4129–4130.
- 17. Fujita M, Yazaki J, Ogura K. J. Am. Chem. Soc 1990;112:5645–5647.
- 18. Stang PJ, Cao DH. J. Am. Chem. Soc 1994;116:4981–4982.
- 19. Stang PJ, Cao DH, Saito S, Arif AM. J. Am. Chem. Soc 1995;117:6273–6283.
- 20. Stang PJ, Olenyuk B, Fan J, Arif AM. Organometallics 1996;15:904–908.
- 21. Stang PJ, Olenyuk B. Angew. Chem. Int. Ed. Engl 1996;35:732–736.
- 22. Stang PJ, Fan J, Olenyuk B. Chem. Commun 1997:1453–1454.
- 23. For examples of $A^2{}_{2}A^2{}_{2}$ squares utilizing mixed Pt and Pd metals see: (a) Stang PJ, Whiteford JA. Organometallics 1994;13:3776–3777. (b) Whiteford JA, Lu CV, Stang PJ. J. Am. Chem. Soc 1997;119:2524–2533. For examples of $A^2{}_{2}A^2{}_{2}$ squares utilizing hypervalent iodonium corners see: (a) Stang PJ, Chen K. J. Am. Chem. Soc 1995;117:1667–1668. (b) Stang PJ, Chen K, Arif AM. J. Am. Chem. Soc 1995;117:8793–8797.. For examples of $A^2{}_{2}A^2{}_{2}$ squares utilizing 5,10-dipyridyl substituted porphyrins as 90° units see: (a) Drain CM, Lehn JM. J. Chem. Soc., Chem. Commun 1994:2313–2315. (b) Fan J, Whiteford JA, Olenyuk B, Levin MD, Stang PJ, Fleischer EB. J. Am. Chem. Soc 1999;121:2741–2752.
- 24. Kuehl CJ, Huang SD, Stang PJ. J. Am. Chem. Soc 2001;123:9634–9641. [PubMed: 11572685]
- 25. Resendiz MJE, Noveron JC, Disteldorf H, Fischer S, Stang PJ. Org. Lett 2004;6:651–653. [PubMed: 14986941]
- 26. Addicott C, Oesterling I, Yamamoto T, Mullen K, Stang PJ. J. Org. Chem 2005;70:797–802. [PubMed: 15675834]
- 27. Chi K-W, Addicott C, Arif AM, Das N, Stang PJ. J. Org. Chem 2003;68:9798–9803. [PubMed: 14656109]
- 28. Yamamoto T, Arif AM, Stang PJ. J. Am. Chem. Soc 2003;125:12309–12317. [PubMed: 14519016]
- 29. Kryschenko YK, Seidel SR, Arif AM, Stang PJ. J. Am. Chem. Soc 2003;125:5193–5198. [PubMed: 12708871]
- 30. For a discussion of the equilibrium between metallacyclic pentagons and hexagons that exists when complementary 109° and 180° precursors are mixed in a 1:1 ratio, see 14b 882.
- 31. Stang PJ, Perskyc NE, Manna J. J. Am. Chem. Soc 1997;119:4777–4778.
- 32. Leininger S, Schmitz M, Stang PJ. Org. Lett 1999;1:1921–1923. [PubMed: 10836049]
- 33. Stang PJ, Olenyuk B, Muddiman DC, Smith RD. Organometallics 1997;16:3094–3096.
- 34. Leininger S, Fan J, Schmitz M, Stang PJ. Proc. Natl. Acad. Sci. USA 2000;97:1380–1384. [PubMed: 10677470]
- 35. Schweiger M, Yamamoto T, Stang PJ, Bläser D, Boese R. J. Org. Chem 2005;70:4861–4864. [PubMed: 15932331]
- 36. Fujita M, Yazaki J, Oka H, Yamaguchi K, Ogura K. Nature 1995;378:469–471.
- 37. Kuehl CJ, Yamamoto T, Seidel SR, Stang PJ. Org. Lett 2002;4:913–915. [PubMed: 11893184]
- 38. For a more recent example of carbon-rich, flexible trigonal prisms see: Yang H-B, Ghosh K, Das N, Stang PJ. Org. Lett 2006;8:3991–3994. [PubMed: 16928056]
- 39. Kumazawa K, Biradha K, Kusukawa T, Okano T, Fujita M. Angew. Chem. Int. Ed 2003;42:3909– 3913.
- 40. Kumazawa K, Biradha K, Fujita M, Sakamoto S, Yamaguchi KA. Angew. Chem. Int. Ed 2001;40:1718–1721.
- 41. Caskey DC, Yamamoto T, Addicott C, Shoemaker RK, Vacek J, Hawkridge AM, Muddimann DC, Kottas GS, Michl J, Stang PJ. J. Am. Chem. Soc 2008;130:7620–7628. [PubMed: 18491898]
- 42. Olenyuk B, Whiteford JA, Fechtenkötter A, Stang PJ. Nature 1999;398:796–799. [PubMed: 10235260]
- 43. Abrahams BF, Egan SJ, Robson R. J. Am. Chem. Soc 1999;121:3535–3536.
- 44. Tominaga M, Suzuki K, Kawano M, Kusukawa T, Ozeki T, Sakamoto S, Yamaguchi K, Fujita M. Angew. Chem. Int. Ed 2004;43:5621–5625.
- 45. Radhakrishan U, Schweiger M, Stang PJ. Org. Lett 2001;3:3141–3143. [PubMed: 11574015]
- 46. Kuehl C, Kryschenko YK, Radhakrishnan U, Seidel SR, Huang SD, Stang PJ. Proc. Natl. Acad. Sci. USA 2002;99:4932–4936. [PubMed: 11880599]
- 47. Fujita M, Yu S-Y, Kusukawa T, Funaki H, Ogura K, Yamaguchi K. Angew. Chem. Int. Ed 1998;37:2082–2085.
- 48. Schweiger M, Seidel SR, Schmitz M, Stang PJ. Org. Lett 2000;2:1255–1257. [PubMed: 10810721]
- 49. (a) Olenyuk B, Levin MD, Whiteford JA, Shield JE, Stang PJ. J. Am. Chem. Soc 1999;121:10434– 10435. (b) Levinc MD, Stang PJ. J. Am. Chem. Soc 2000;122:7428–7429.
- 50. Tashiro S, Tominaga M, Kawano M, Therrien B, Ozeki T, Fujita M. J. Am. Chem. Soc 2005;127:4546–4547. [PubMed: 15796499]
- 51. Tashiro S, Kobayashi M, Fujita M. J. Am. Chem. Soc 2006;128:9280–9281. [PubMed: 16848433]
- 52. (a) Kumazawa K, Yamanoi Y, Yoshizawa M, Kusukawa T, Fujita M. Angew. Chem. Int. Ed 2004;43:5936–5940. (b) Yoshizawa M, Nakagawa J, Kumazawa K, Nagao M, Kawano M, Ozeki T, Fujita M. Angew. Chem. Int. Ed 2005;44:1810–1813.
- 53. Merlau ML, del Pilar Mejia M, Nguyen ST, Hupp JT. Angew. Chem. Int. Ed 2001;40:4239–4242.
- 54. Oliveri CG, Gianneschi NC, Nguyen ST, Mirkin CA, Stern CL, Wawrzak Z, Pink M. J. Am. Chem. Soc 2006;128:16286–16296. [PubMed: 17165783]
- 55. Dinolfo PH, Hupp JT. Chem. Mater 2001;13:3113–3125. and references therein.
- 56. Czaplewski KF, Hupp JT, Snurr RQ. Adv. Mater 2001;13:1895–1897.
- 57. Williams ME, Benkstein KD, Abel C, Dinolfo PH, Hupp JT. Proc. Natl. Acad. Sci. USA 2002;99:5171–5177. [PubMed: 11959966]
- 58. Mulfort KL, Hupp JT. J. Am. Chem. Soc 2007;129:9604–9605. [PubMed: 17636927]
- 59. Gong J-R, Wan L-J, Yuan Q-H, Bai C-L, Jude H, Stang PJ. Proc. Natl. Acad. Sci. USA 2005;102:971– 974. [PubMed: 15657148]
- 60. Yuan Q-H, Wan L-J, Jude H, Stang PJ. J. Am. Chem. Soc 2005;127:16279–16286. [PubMed: 16287321]
- 61. Yuan Q-H, Yan C-J, Yan H-J, Wan L-J, Northrop BH, Jude H, Stang PJ. J. Am. Chem. Soc 2008;130:8878–8879. [PubMed: 18549212]
- 62. Li S-S, Yan H-J, Wan L-J, Yang H-B, Northrop BH, Stang PJ. J. Am. Chem. Soc 2007;129:9268– 9269. [PubMed: 17622151]
- 63. (a) Yang H-B, Das N, Huang F, Hawkridge AM, Muddiman DC, Stang PJ. J. Am. Chem. Soc 2006;128:10014–10015. [PubMed: 16881621] (b) Yang H-B, Hawkridge AM, Huang SD, Das N, Bunge SD, Muddiman DC, Stang PJ. J. Am. Chem. Soc 2007;129:2120–2129. [PubMed: 17256935]
- 64. (a) Yang H-B, Ghosh K, Northrop BH, Zheng Y-R, Lyndon MM, Muddiman DC, Stang PJ. J. Am. Chem. Soc 2007;129:14187–14189. [PubMed: 17963382] (b) Ghosh K, Yang H-B, Northrop BH, Lyndon MM, Zheng Y-R, Muddiman DC, Stang PJ. J. Am. Chem. Soc 2008;130:5320–5334. [PubMed: 18341280]
- 65. Yang H-B, Ghosh K, Zhao Y, Northrop BH, Lyndon MM, Muddiman DC, White HS, Stang PJ. J. Am. Chem. Soc 2008;130:839–841. [PubMed: 18166061]
- 66. (a) Tominaga M, Suzuki K, Murase T, Fujita M. J. Am. Chem. Soc 2005;127:11950–11951. [PubMed: 16117528] (b) Sato S, Iida J, Suzuki K, Kawano M, Ozeki T, Fujita M. Science 2006;313:1273– 1276. [PubMed: 16946067] (c) Murase T, Sato S, Fujita M. Angew. Chem. Int. Ed 2007;46:5133– 5136. (d) Murase T, Sato S, Fujita M. Angew. Chem. Int. Ed 2007;46:1083–1085. (e) Kamiya N, Tominaga M, Sato S, Fujita M. J. Am. Chem. Soc 2007;129:3816–3817. [PubMed: 17343384]
- 67. (a) Flammang-Barbieux M, Nasielski J, Martin RH. Tetrahedron Lett 1967:743–744. (b) Martin RH, Flammang-Barbieux M, Cosyn JP, Gelbcke M. Tetrahedron Lett 1968:3507–3510.
- 68. Staab HA, Wehinger E, Thorwart W. Chem. Ber 1972;105:2290–2309.

- 69. Campbell, K.; Tykwinski, RR. Chiral Carbon-Rich Macrocycles and Cyclophanes. In: Haley, MM.; Tykwinski, RR., editors. Carbon-Rich Compounds. Weinheim: Wiley-VCH; 2006.
- 70. (a) Fuss M, Siehl H-U, Olenyuk B, Stang PJ. Organometallics 1999;18:758–769. (b) Vacek J, Caskey DC, Horinek D, Shoemaker RK, Stang PJ, Michl J. J. Am. Chem. Soc 2008;130:7629–7638. [PubMed: 18491897]
- 71. For an example of a purely covalent, carbon rich triagular dehydrobenzo[12]annulene see: (a) Campbell ID, Eglinton G, Henderson W, Raphael RA. Chem. Commun 1966:87–89. (b) Staab HA, Graf F. Tetrahedron Lett 1966:751–757.
- 72. For an example of a purely covalent, carbon-rich 1,8-diethynylanthracene rectangle see: Akiyama S, Nakagawa M. Chem. Ind 1960:346–347.
- 73. (a) Kohnke FH, Slawin AMZ, Stoddart JF, Williams DJ. Angew. Chem. Int. Ed. Engl 1987;26:892– 894. (b) Kohnke FH , Mathias JP, Stoddart JF. Angew. Chem. Int. Ed. Engl 1989;28:1103–1110.
- 74. Manini P, Amrein W, Gramlich V, Diederich F. Angew. Chem. Int. Ed 2002;41:4339–4343.

Figure 1.

The directional-bonding approach to the self-assembly of 2D metallacyclic polygons.

Figure 2.

The directional-bonding approach to the self-assembly of select 3D metallacages.

Carbon-rich ditopic 90° metal acceptors (**A**) and ditopic 180° donors (**B**).

Northrop et al. Page 16

Figure 4.

Carbon-rich Pt(II) acceptors with turning angles of 0°, 60°, 109°, 120°, and 180° (**A**) as well as non-linear ditopic donors (**B**).

Northrop et al. Page 17

Figure 5. Trigonal planar tritopic donors and acceptors.

Figure 7.

 $31P$ NMR spectrum (121.5 MHz, CD₃COCD₃, 298 K) of the self-assembled carbon-rich supramolecular diastereomers. The inset shows that what appears to be a single sharp peak is actually composed of two peaks of differing intensity, indicating one diastereomer is formed preferentially.

Figure 8.

Structures of symmetric and asymmetric diastereomers obtained from molecular modeling. Measurements of the distance between neighboring phenanthrene units partially demonstrates differences in structural strain between the diastereomers. Color scheme: $C = grey, N = blue,$ $P =$ purple, $Pt =$ yellow.

Scheme 1.

The self-assembly of carbon-rich metallacyclic squares. In all cases the acceptor and donor are mixed in a 1:1 molar ratio. $M = Pt$, Pd.

Scheme 2.

Summary of protocols for self-assembling a variety of carbon-rich 2D polyhedra. In each case the donor and acceptor precursors are mixed in a 1:1 molar ratio. $a = Pt(PMe₃)₂(OTf)₂$.

Scheme 3.

Use of trigonal planar tritopic donors to self-assemble carbon-rich truncated tetrahedra and trigonal prisms. $a = Pt(PMe₃)₂(OTf)₂$.

Scheme 4.

The self-assembly of "walled" trigonal prism geometries using carbon-rich tetratopic donors and 90° metal acceptors in a 3:6 molar ratio.

Scheme 5.

The self-assembly of cuboctahedra from a 12 ditopic 120° precursors and 8 complementary tritopic trigonal planar precursors.

Scheme 6.

Self-assembly of trigonal bipyrimidal, adamantoid, and dodecahedral polygons. Component acceptor and donor precursors are mixed in 3:2, 6:4, and 30:20 molar ratios, respectively.

Scheme 7.

Self-assembly of "symmetric" and "asymmetric" diastereomers of a carbon-rich supramolecular triangle from achiral building blocks.