Chiral Amide Directed Assembly of a Diastereo- and Enantiopure Supramolecular Host and its Application to Enantioselective Catalysis of Neutral Substrates

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Supporting Information

ABSTRACT: The synthesis of a novel supramolecular tetrahedral assembly of $K_{12}Ga_4L_6$ stoichiometry is reported. The newly designed chiral ligand exhibits high diastereoselective control during cluster formation, leading exclusively to a single diastereomer of the desired host. This new assembly also exhibits high stability toward oxidation or a low pH environment and is a more robust and efficient catalyst for asymmetric organic transformations of neutral substrates.

Inspired by nature, recent work in supramolecular chemistry has focused on the design and construction of assemblies that imitate the properties of enzymes.1 Many such synthetic nanovessels can function in aqueous environments at physiological pH,2 contain well-defined cavities for selective guest encapsulation and recognition,3 and have been shown to stabilize otherwise reactive and unstable species.4 Furthermore, many supramolecular hosts have proven to be efficient catalysts that increase both the rate and selectivity of a variety of chemical reactions.5 Raymond et al. have developed tetrahedral supramolecular assembly 1 of $K_{12}Ga_4L_6$ stoichiometry, where $L = N,N$-bis(2,3-dihydroxybenzoyl)-1,5-diaminonaphthalene.6 The highly charged anionic host 1 has been shown to encapsulate a variety of cationic and neutral guests;7 however, to date, its use in enantioselective catalysis has been limited to the charged substrates of the Aza-Cope rearrangement.8 While Fujita et al. have reported the [2 + 2] cycloaddition of neutral guests in stoichiometric chiral hosts,9 the use of nanoscale molecular flasks possessing chiral cavities as catalysts for asymmetric transformations of neutral guests remains elusive.8−10

Complex 1 is a chiral species because the three catecholates coordinate to a given gallium atom and can form either a right ($\Delta$)- or a left ($\Lambda$)-handed helicity at each metal center. Enforced by mechanical coupling that leads to chirality transfer between the four vertices,11 complex 1 is formed as a racemic mixture of two homochiral enantiomeric forms, namely $\Lambda\Lambda\Lambda\Lambda$-1 and $\Delta\Delta\Delta\Delta$-1. Resolution of the racemate was realized using (−)-N′-methylnicotinium iodide, giving access to enantiopure $\Lambda\Lambda\Lambda\Lambda$-(S-nic $\subset$ 1) and $\Delta\Delta\Delta\Delta$-(S-nic $\subset$ 1) stereoisomers.12 Sequential ion exchange chromatography with large excess amounts of tetramethylammonium and potassium iodide salts then afforded “empty” and enantiopure clusters. However, the instability of the isolated cationic guest-free or K'-filled $\Lambda\Lambda\Lambda\Lambda$-1 and $\Delta\Delta\Delta\Delta$-1 clusters warrants improvement.12 We describe herein the design and synthesis of a new enantiopure supramolecular $Ga_4L_6$ cluster that spontaneously self-assembles. In addition to circumventing the need for resolution, these new assemblies provide enhanced stability and catalytic reactivity required for asymmetric organic transformations of neutral guests.

Our strategy for achieving an enantiopure supramolecular $M_4L_6$ assembly without resolution involves the addition of an amide-containing chiral directing group at the vertex of ligand 2, as shown in Figure 1. We envisioned that this chiral source would control the helical configuration of the proximal metal center during cluster formation and direct a highly diastereoselective process in which the desired $M_4L_6$ supramolecular assemblies would be formed enantioenriched rather than as a racemate. We also suspected that this additional amide functional group would stabilize the resulting assembly via hydrogen bonding with the catecholates and could prevent ligand oxidation and decomposition due to its electron withdrawing nature.

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Ligand (R)-S was prepared as shown in Scheme 1. The terephthalate sodium salt was converted to the corresponding acyl chloride. This was followed by amide bond formation with commercially available chiral amine (R)-(−)-3,3-dimethyl-2-butyramine and subsequent saponification with KOH in methanol to afford the desired intermediate (R)-S. Reaction between (R)-S and 1.2 equiv of O-(7-azabenzotriazol-1-yl)-N,N,N′,N′-tetramethyluronium hexafluorophosphate in THF for 1 h at room temperature, followed by addition of 1,5-diaminonaphthalene gave the desired methyl-protected chiral ligand (R)-4. Methyl group deprotection of (R)-4 was achieved by treatment with BBr3 and hydrolysis of the resulting borate to produce the desired terephthalamide-based chiral ligand (R)-S in 52% yield over 5 steps. The enantiomer (S)-S was also synthesized according to the procedures shown in Scheme 1.

We next investigated whether ligand (R)-S would form the desired tetrahedral supramolecular assembly. The initial reaction between 4 equiv of Ga(acac)3, 6 equiv of ligand (R)-S, and 12 equiv of KOH in methanol at room temperature, in the absence of any cationic species as a template, gave a mixture of products as analyzed by 1H NMR spectroscopy (see SI). However when the reaction was repeated at 50 °C for 1 h, highly symmetric complex 6, as suggested by the simplicity of its 1H NMR spectrum (see SI), was isolated as a yellow solid in 78% yield. Analysis of 6 by ESI mass spectrometry confirmed its stoichiometry as K12Ga4S5. Furthermore, when 5 equiv of PEt4I was added to a D2O solution of 6, encapsulation of PEt4+ was observed as indicated by the proton resonances at δ = −1.45 and −1.78 ppm (see SI). This observation can also be taken as an indication of the successful formation of the desired tetrahedral assembly 6.6 Furthermore, 6-K12Ga4(R)-S6 was synthesized without the use of any cationic species as a template, whereas enantiopure 1 could only be obtained as a stable species after treatment with excess amount of NMe4+. The enantiomer of 6-K12Ga4(S)-S6 was also synthesized by using ligand (S)-S, Ga(acac)3, and KOH following a procedure directly analogous to that outlined in Scheme 2.

Complex 6 was also found to be benchtop stable in both the solid and solution states at elevated temperature, whereas complex 1 was sensitive to oxidation and relatively less stable at 40 °C in the absence of a strong binding guest in solution over time. More importantly, complex 6 proved to be stable in aerobic D2O at pD 5 and readily encapsulates PEt4+ even after heating at 70 °C for 6 h, while complex 1 and (NEt4)121

Scheme 1. Synthesis of Ligand (R)-S

Scheme 2. Synthesis of Supramolecular Assembly 6 and Its Encapsulation of PEt4+ Cation

It was reported previously that the UV x−π* transitions of the catechol moiety of assembly 1 produced a strong and distinct exciton couplet.14 This property enabled the determination of absolute configuration of the resolved enantiopure parent assembly 1 by circular dichroism (CD) spectroscopy.12 When assemblies 6-K12Ga4(R)-S6 and 6-K12Ga4(S)-S6 were examined by CD spectroscopy, the spectra of the two enantiomers proved to be perfect mirror images of each other and to contain a shape and sign of the Cotton effect similar to those of ΔΔΔΔ-1 and ΔΔΔΔ-1 (see SI).15 Thus, we infer by comparison and assign complex 6-K12Ga4(R)-S6 as the ΔΔΔΔ stereoisomer and 6-K12Ga4(S)-S6 as the ΔΔΔΔ stereoisomer.

The absolute stereochemical assignment of ΔΔΔΔ-6 was further supported by X-ray crystallographic analysis. Single crystals were obtained by slow diffusion of THF vapor into a water solution of ΔΔΔΔ-6 without any strong binding and cationic guest molecules under aerobic conditions. The structure conforms to the chiral space group R3 with three molecules of the enantiopure complex in the unit cell, each with crystallographic three-fold symmetry. As shown in Figure 2, all four gallium centers adopt the Δ configuration, with an average Ga–Ga distance of 12.6 Å, similar to that found in the resolved parent assembly 1.8,12 The chiral directing groups bury the metal vertices of the cage with additional intramolecular hydrogen bonds between the amide proton and the catecholate oxygen, which could be responsible for the observed stability of this new cluster. By crystal packing, each cage is part of a larger network of 12 neighboring cages, forming a 3-dimensional

Figure 2. X-ray structure of ΔΔΔΔ-6.
molecular organic framework. A huge solvent accessible void of 25,000 Å³ is calculated for the unit cell (65% of total unit cell volume), as a result of the large channels found along both the a and b axes of the crystal.

As a further probe of the stereochemistry of ΔΔΔΔ-6 and ΔΛΛΛ-6, we investigated their host-guest chemistries individually with both enantiomers of ammonium salt 8. As illustrated in Figure 3, host-guest complex 9, or ΔΔΔΔ-[((S)-8]-contentsubject_reference{345x522}ΔΔΔΔ-6, should have different and distinguishable properties from complex 10, ΔΔΔΔ-[((R)-8]⊂6, due to their diastereomeric relationship. 1H NMR spectroscopy (Figure 3) reveals that the two complexes are indeed different, most notably in the encapsulation region of the spectra. On the other hand, complex 11, ΔΛΛΛ-[(R)-8]⊂6, and complex 9 are enantiomers and exhibit exactly the same spectroscopic behaviors when analyzed by 1H NMR; the same result was also observed for complexes 10 and 12. This evidence, combined with results from X-ray crystallography and CD spectroscopy, demonstrates that complex 6 is highly enantioenriched. The chiral group of ligand 5 exhibits strong control during cluster formation to give the desired supramolecular Κ12Ga4L6 cluster as a single diastereomer. 15

One challenge to the development of asymmetric organic reactions catalyzed by enantiopure host ΔΔΔΔ-1 is the requirement for cationic starting material or substrates that are more tightly bound than is NMe4+ to the cavity of ΔΔΔΔ-1. Since ΔΔΔΔ-6 was synthesized without the use of any templates or cationic species, this new supramolecular host makes possible the enantioselective transformations of neutral compounds.

We recently reported the chemoselective carbonyl-ene cyclization of compounds 13a and 13b catalyzed by complex 1 to give exclusively products 14a,b and 15a,b respectively, as compared to a reaction performed in bulk solution. 16 When the reaction was repeated with 10 mol % of (NMe4)12H8 at 60 °C in D2O buffered at pD 8 for 14h, no desired products were observed. On the other hand, when compound 13a was treated with 2.5 mol % of ΔΔΔΔ-6 in a solvent mixture of CD2OD and D2O buffered at pD 8 at room temperature, the desired products 14a and 15a were obtained in 92% NMR yield with a trans:cis ratio of 8:1 and 61% ee for 14a over two days (Table 1, entry 1). Compared to reaction with complex 1 as the catalyst at the same pD, cyclization of 13a in the presence of a catalytic amount of ΔΔΔΔ-6 proved to be faster by 7-fold (see SI).

| Table 1. Enantioselective and Chemoselective Monoterpene-Like Cyclization of Neutral Substrates Catalyzed by ΔΔΔΔ-6 |

<table>
<thead>
<tr>
<th>entry</th>
<th>R</th>
<th>pD</th>
<th>temp (°C)</th>
<th>time (h)</th>
<th>yield (trans:cis)</th>
<th>ee of 14</th>
</tr>
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<tbody>
<tr>
<td>1</td>
<td>Me</td>
<td>8</td>
<td>25</td>
<td>50</td>
<td>92% (8:1)</td>
<td>61%</td>
</tr>
<tr>
<td>2</td>
<td>Me</td>
<td>5</td>
<td>25</td>
<td>16</td>
<td>94% (7:5:1)</td>
<td>58%</td>
</tr>
<tr>
<td>3</td>
<td>Me</td>
<td>5</td>
<td>−25</td>
<td>16</td>
<td>70% (8:1)</td>
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<td>H</td>
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<td>6</td>
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<td>60</td>
<td>16</td>
<td>92% (8:1)</td>
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The stability and turnover capability of catalyst ΔΔΔΔ-6 was further illustrated as only 0.3 mol % of the complex is required to achieve 33% yield of 14a and 15a with no loss in enantiomeric excess of 14a (Table 1, entry 4), representing 99 TON of the catalyst. Interestingly, carbonyl-ene cyclization of 13b proceeded with complex 6 at pD 8 over 16 h at 60 °C to give the desired products in only 12% yield (Table 1, entry 5), whereas reaction at pD 5 led to much better conversion over the same reaction time to give the desired product mixture in 92% yield and 65% ee of 14b. 17,18

In conclusion, a new enantiopure supramolecular Κ12Ga4L6 assembly has been synthesized, fully characterized, and applied as a rare example of chiral host-catalyzed enantioselective transformations of neutral guests. The chiral amide in the terephthalamide-based ligands (R)-5 and (S)-5 directs cluster formation to afford highly diastereo- and enantiomerically enriched complexes. Remarkably, cationic guest-free variants of complexes ΔΔΔΔ-6 and ΔΛΛΛ-6, which in comparison to 1 vary only in modification to the exterior of the assembly, show increased stability toward air oxidation in both the solid and solution states and to low pH in solution. These features allow complexes ΔΔΔΔ-6 and ΔΛΛΛ-6 to serve as efficient catalysts for chemo-, diastereo-, and enantioselective carbonyl-ene cyclization.

ASSOCIATED CONTENT

Supporting Information

Experimental procedures and spectroscopic data. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
The authors declare no competing financial interest.

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(15) Molecular mechanics calculations of \( \Delta \Delta \Delta \Delta [Ga_4(R)-5] \) and \( \Delta \Delta \Delta \Delta [Ga_4(R)-5_a] \) reveal that the ground-state energy difference between the two diastereomers is \( \sim 19 \text{ kcal/mol} \), favoring the experimentally observed isomer \( \Delta \Delta \Delta \Delta [Ga_4(R)-5] \) (see SI for details).

The observed chemoselectivity, yield, and enantiomeric excess of \( 15b \) are improved as compared to Lewis acid-catalyzed transformation of the same substrate reported in literature: (a) Mikami, K.; Sawa, E.; Terada, M. Tetrahedron: Asymmetry 1991, 12, 1403.
(18) For an additional example of an enantioselective transformation of neutral substrate catalyzed by complex 6, see SI.