# The Source of Rate Acceleration for Carbocation Cyclization in Biomimetic Supramolecular Cages

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**ABSTRACT:** The results of quantum chemical and molecular dynamics calculations reveal that polyanionic gallium-based cages accelerate cyclization reactions of pentadienyl alcohols as a result of substrate cage interactions, preferential binding of reactive conformations of substrate/ $H_3O^+$  pairs and increased substrate basicity. However, the increase in basicity dominates. Experimental structure-activity relationship studies in which the metal vertices and overall charge of the cage are varied confirm the model arrived at via calculations.

#### INTRODUCTION

Carbocations are involved as intermediates in catalytic reactions accomplished by both synthetic chemists and biological systems. The latter are most prominently found in the realm of terpene biosynthesis, where terpene synthase enzymes generate carbocations in productive conformations, allow them to rearrange while protecting them from premature deprotonation or addition by nucleophiles, and allow them to be deprotonated at specific positions (Scheme 1, top, OPP = diphosphate).¹ Synthetic chemists have developed systems that mimic these characteristics.².³ In both the synthetic and biosynthetic realms, however, the relative importance of each step in the reaction mechanism is still up for debate. Extensive computational work has targeted this issue for terpene synthases,¹c-d,⁴ but much less attention has been afforded to synthetic systems.².3,5

Here we describe molecular dynamics (MD) and quantum chemical computations on a 4-electron electrocyclization of pentadienyl carbocations promoted by the tetrahedral Ga<sub>4</sub>L<sub>6</sub> cage studied by Raymond, Bergman, Toste and co-workers (Scheme 1, bottom).2 The cyclization of dienol 1 (multiple geometric isomers have been examined; only one is shown) typically requires strong Lewis or Bronsted acids in bulk solution to promote protonation of the oxygen and generate a pentadienyl carbocation intermediate (2), which then undergoes a 4-electron electrocyclic ring closure. Under Ga<sub>4</sub>L<sub>6</sub> catalyzed conditions, the hydrophobic substrate 1 is first reversibly bound in the naphthalene-walled host cavity, which enables protonation at the alcohol despite the basic aqueous medium outside the host. The substrate ionizes to release water before undergoing 4-electron electrocyclization (associated with the highest energy transition state on the reaction coordinate), followed by proton loss to generate the cyclopentadiene product **3**. Due to the similar association constants between the product **(3)** and substrate **(1)**, excess maleimide was added to trap the product as the Diels-Alder adduct **4**, which no longer is encapsulated by the host.

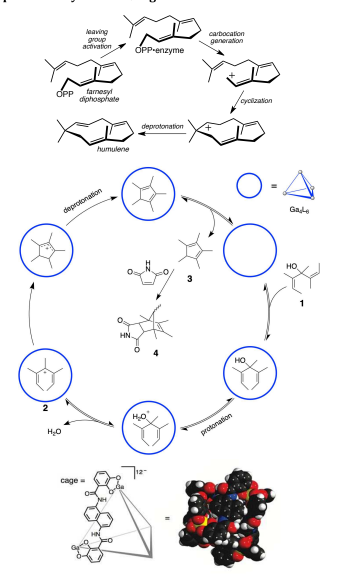
Our results indicate the essential role of the catalyst cage in promoting carbocation formation primarily via leaving group basicity enhancement. We also conducted a comprehensive experimental study to validate the model derived from calculations and determine the contribution of the metal vertices to supramolecular catalytic activity. We found that when the overall charge of the host was held constant, changing the metal center had little effect on the host's catalytic performance. Our results provide evidence that supramolecular hosts can tolerate systematic structural modifications without significant loss of catalytic activity.

## **Computational Methods**

Finding a computational approach for modeling such a large system that strikes the right balance between accuracy and efficiency is a challenge. Inspired by the success of Nitschke and co-workers in computing geometries for related cage structures,<sup>6</sup> we first examined the feasibility of employing various density functionals with relatively small basis sets (see Supporting Information for details). Ultimately, we found that the B3LYP/3-21G level of theory,<sup>7</sup> with the LANL2DZ basis set for Ga,<sup>8</sup> provided reasonable results, suitable for surveying a wide range of structures. Optimizing geometries and computing vibrational frequencies with ostensibly "better" levels of theory proved, in our hands, to be impractical at best and intractable at worst.

However, single point energies on B3LYP/3-21G-LANL2DZ geometries with various other functionals and larger basis sets showed that the results described below were not very sensitive to the method used (see Supporting Information); all led to the same qualitative conclusions. In addition, for substrates in the absence of the cage, we performed optimization and frequency calculations with a variety of methods and larger basis sets; results were again very similar to those obtained with B3LYP/3-21G (see Supporting Information). All calculations were conducted using the CPCM continuum model for water (the solvent used experimentally),9 which allowed the cage, which bears a 12- charge, not to expand significantly beyond its reported crystallographic geometry,<sup>2,10</sup> despite the absence of counterions in these calculations. All reported energies are free energies at 298 K unless stated otherwise.

SCHEME 1. Top: A typical, albeit simple, terpene-synthase promoted reaction. Bottom: Electrocyclization promoted by the  $Ga_4L_6$  cage.



In addition to these static calculations, we carried out molecular dynamics (MD) calculations to address conformational and configurational flexibility. These calculations were set up based on the initial QM optimization results. The organic portions of the cage and the ligand molecules were described by the AMBER general force field (GAFF)<sup>11</sup> and their charges were calculated by the RESP method<sup>12</sup> based on ESP results obtained with HF/6-31G\*. Since there are no parameters for Ga or Si in AMBER,13 we substituted both by Al for our molecular mechanics (MM) calculations (but set the Si charge to 4+). The assumption that any electronic differences aside from charge between Ga and Si are insignificant is validated experimentally (vide infra). All system preparations were done using the tleap program in AMBER16.13 In the MM/MD simulations, a multi-step strategy was used to heat and equilibrate the system gradually. First, energy minimization was carried out, followed by a 100 ps heating process (NVT ensemble). Then, another 100 ps MD simulation was carried out to balance the system (NPT ensemble with the help of the Berendsen barostat). Finally, a 50 ns MD simulation (NVT ensemble) was carried out. During the heating and equilibrium processes, an extra force was first added to the cage to prevent its contraction and the SHAKE<sup>14</sup> algorithm was applied to constrain covalent bond lengths involving hydrogen. The cutoff values for short-range electrostatics and van der Waals interactions were set to 12 Å, and the particle-mesh Ewald method was used for long range summation of electrostatic interactions. 15 All MD calculations were carried out using the OpenMM package.16

In addition, to estimate energy barriers for leaving group activation and departure, combined quantum mechanics (QM)/MM calculations were carried out using the *Q-Chem* and *AMBER* software packages. <sup>17</sup> The QM region was set as the guest molecule, the  $\rm H_3O^+$  and two  $\rm H_2O$  molecules that engage in hydrogen-bond interactions with the  $\rm H_3O^+$ . All atoms in the QM region were treated with B3LYP/6-31G\*, and all the non-QM atoms were described by the force field mentioned above with a 12 Å cutoff for non-bonded interactions. QM/MM free energy calculations made use of umbrella sampling along the defined reaction coordinate (*vide infra*) and the weighted histogram analysis method. <sup>18</sup>

#### **RESULTS AND DISCUSSION**

#### **Effects of Cage Walls on Cyclization**

Raymond, Bergman and co-workers observed rate accelerations of approximately six orders of magnitude for various isomers of the substrate shown in Scheme 1 in the presence of the cage. Here we model the substrate with two E alkenes. First, we address the issue of whether or not direct interactions with the  $Ga_4L_6$  cage lower the barrier for electrocyclization. The transition state structure (TSS) for electrocyclization in the absence of the cage (but in a water continuum) is shown at the top of Figure 1.<sup>19</sup> A barrier of only  $\sim$ 4 kcal/mol was computed for cyclization from a productive, i.e., preorganized, conformer of reactant. Model calculations (Figure 1, middle and bottom) in the presence of a benzene or naphthalene molecule indicate that the aromatic walls of the cage can provide rate acceleration, although the effect is not large (see Supporting Information for

complexes with benzene and an explicit water molecule, which lead to similar results), i.e., while the walls interact with the substrate,  $^{2,3d}$  selective binding of the transition state structure over the pentadienyl cation intermediate is predicted to amount to <2 kcal/mol, *indicating that, at best, the large observed rate acceleration must originate primarily from another source.* 

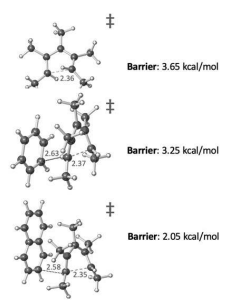


FIGURE 1. Top to bottom: TSS of the model substrate, TSS with a benzene ring nearby, TSS with a naphthalene nearby. Each energy barrier was calculated ( $\omega B97XD/6\text{-}311\text{+}G(d,p)^{19}$ ) relative to the respective reactant with a similar environment, i.e. reactant alone or with a benzene or naphthalene in close proximity.

# **Conformational Preorganization**

Another possible contributor to the observed rate acceleration is reactant preorganization, i.e., adopting the productive conformation in solution is accompanied by an energy penalty that is "paid" by the catalyst upon binding. Our DFT calculations suggest, however, that coiled conformers of the substrate, its O-protonated form and the pentamethylpentadienyl cation, (where the two end carbons of the pentadienyl system are near to each other), are within ~2 kcal/mol of the lowest energy conformer. For the carbocation, this is the lowest energy conformer. Our MD simulations indicate that pentamethylpentadienol in explicit water rapidly explores both extended and compact conformations (Figure 2, top). Upon complexation, it still explores these conformations, although transitions between them are not as smooth (Figure 2, middle). The barrier for transitioning between extended and compact forms inside the cage is estimated to be ~3 kT (roughly 2 kcal/mol at room temperature), while outside the cage, the barrier is close to 0 (energy plot shown in SI). In general, as water molecules flow into the cage, the conformation of the guest changes from extended to compact. Overall, there appears to be no significant preorganization induced by the catalyst for the alcohol. Pentamethylpentadienol along with an explicit hydronium ion next to the hydroxyl group of the substrate was also modeled with MD (Figure 2, bottom). In this case, the substrate mainly adopts a relatively compact conformation, suggesting that the alcohol/ $H_3O^+$  pair is preferentially accommodated in orientations productive for reaction.

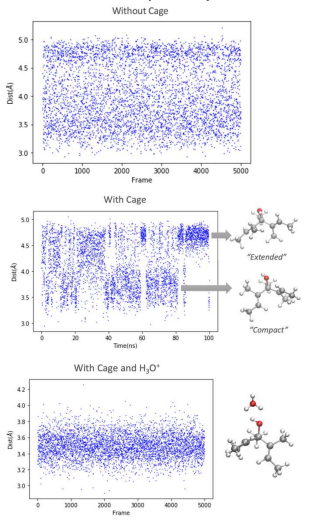


FIGURE 2. Distance between alkene carbons that will form the new C–C bond versus time during MD simulations. Top: pentamethylpentadienol in water. Middle: pentamethylpentadienol in the cage. Bottom: pentamethylpentadienol with hydronium ion in the cage.

# **Leaving Group Activation - Setting the Stage**

That leaves us with leaving group activation. However, an important issue to address in this regard is the number of water molecules contained within the cage, along with any substrates. Warshel and co-workers previously examined the binding of four water molecules plus one H<sub>3</sub>O<sup>+</sup> molecule within the same cage examined here. Based on preliminary DFT calculations, we find that additional water molecules can be encapsulated without major distortions to the cage geometry (see Supporting Information).20 We also addressed this issue with MD simulations of the cage surrounded by water molecules that can enter and leave its interior. Without the substrate present, a range of 0 to 16 water molecules were observed inside the cage over the length of the MD simulation, while 10 to 13 water molecules were the most common numbers (the cage was restrained to prevent shrinking). Recent experimental evidence pointed to 9

± 1 water molecule in the cage in question. <sup>5e</sup> With the pentamethylpentadienol substrate present in the cage, 0 to 4 water molecules were observed in the MD simulation (Figure 3, left, "substrate" bars). When a hydronium ion was included during the MD, 2 to 5 water molecules were present in the cage along with the substrate (water count does not include the hydronium ion; Figure 3, left, "substrate + hydronium" bars). With the *O*-protonated pentamethylpentadienol ligand present, 2 to 6 water molecules were observed inside the cage (Figure 3, left, "cation" bars). A representative snapshot showing the substrate with H<sub>3</sub>O+ and three water molecules in the cage is shown at the right-hand side of Figure 3. The hydronium ion shown is poised to protonate the substrate, while other water molecules inside the cage form a H-bond network.

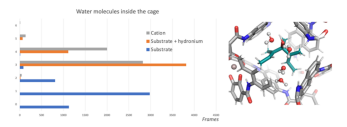


FIGURE 3. Left: water molecules inside the cage in MD. Right: A representative ssnapshot of substrate with  $H_3O^+$  and three water molecules in the cage (water molecules outside the cage are not shown in this picture for clarity).

# **Leaving Group Activation - Protonation**

We begin by discussing DFT (B3LYP/3-21G-LANL2DZ) results on an encapsulated pentamethylpentadienol substrate (Scheme 1), along with various numbers of water molecules and one  $\rm H_3O^+$  molecule. For these calculations, waters were placed manually near to the substrate's hydroxyl group and allowed to relax into the nearest potential energy surface minimum; a consistent solvent configuration was used for all structures. Calculations with different numbers of water molecules and alternative orientations produced qualitatively similar results (see Supporting Information for details).

First, we focused on the degree to which the substrate protonation event is affected by the Ga<sub>4</sub>L<sub>6</sub> cage. To do so, we calculated the free energies associated with protonation of the pentamethylpentadienol reactant inside and outside the cage (both in a water continuum), with various numbers of explicit water molecules present. In all cases, the pentamethylpentadienol reactant was predicted to have a larger (by (≥15 kcal/mol) thermodynamic preference for protonation when encapsulated. We also selected configurations from MD simulations using GROMACS clustering with awareness of symmetry (using RDKit Cookbook) for DFT computations and for these, encapsulated species are again predicted to be ~15 kcal/mol more basic (see Supporting Information for details).21 Warshel and co-workers also argued that the cage interior provides a "remarkable case of a low 'local pH',"5a i.e., encapsulated species are readily protonated. While our computed values here may be overestimates of the  $pK_a$  modulation—previous experiments on amine, phosphine and ester guests indicate that a change in

p $K_a$  of 4-5 units, corresponding to 5-7 kcal/mol can be expected upon encapsulation<sup>10</sup>—it is clear that alcohol basicity is markedly enhanced.

What is the origin of the high acidity of the microenvironment within the Ga<sub>4</sub>L<sub>6</sub> cage? The cage used experimentally has walls made of naphthalene rings and, as shown in Figure 4a, the electrostatic potential within the cage is more negative than that on the cage's outer surface.<sup>22</sup> As a result, it is possible that substrate-cage  $CH-\pi$  interactions<sup>23</sup> strengthen upon protonation, leading to increased substrate basicity. To test the contribution of the cage walls, all naphthalene rings and attached amide groups were removed (and the remaining catechol rings were capped with hydrogen atoms; Figure 4b; note that this does not change the overall charge of the system), and the substrate protonation energy was recomputed (the geometry of the cagesubstrate complex was not allowed to adjust). These changes led to a reduction in predicted protonation energy when no waters are co-encapsulated, i.e., protonation is predicted to be less favorable by 17 kcal/mol, consistent with there being interactions between the  $\pi$ -faces of the cage and the substrate. However, an increase in protonation energy of approximately the same amount is predicted for the complex of substrate and four bound water molecules, highlighting the fact that some bound waters tend to engage in  $OH-\pi$  interactions that, presumably, decrease the strength of their interactions with the protonated alcohol.

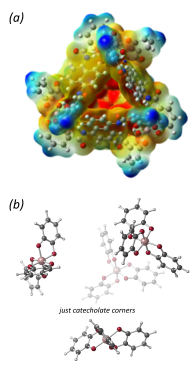


FIGURE 4. (a) Electrostatic potential surface for empty cage (isovalue: 0.005, range: -0.83 to -0.5; red is most negative and blue is least negative). (b) Model of cage with only Ga-cate-cholate corners.

The contributions of specific cage oxygen atoms to basicity modulation were also examined. Keeping the naphthalene walls, but deleting all *ortho* catechol oxygen atoms, all meta catechol oxygen atoms, or all *ortho* and *meta* oxygen

atoms along with all gallium atoms (and capping the remaining aromatic carbons with hydrogen atoms), all changes which lead to a neutral truncated cage, decreased the basicity of bound substrate, in the absence of bound waters, by approximately 15-20 kcal/mol. This result indicates that both cage charge and  $\pi\text{-rich}$  walls contribute roughly equally to the overall effect in the absence of bound water molecules, but are antagonistically coupled. However, in the presence of bound waters, some water molecules will intrude on specific cage-substrate interactions (vide infra).

We also examined the proton transfer process using QM/MM calculations. A representative snapshot from our MD simulations is shown in Figure 5 (cage removed for clarity). Note that this structure showcases the general observation that encapsulated waters cluster on one side of the hydrocarbon group as a hydronium ion hydrogen bonds to the substrate hydroxyl group and to two encapsulated waters. When a QM/MM scan of the proton transfer reaction coordinate is carried out, protonated pentamethylpentadienol results (the dip in the energy plot around r = 0.5), which is predicted to be an endothermic process by  $\sim 2.5$ kcal/mol, associated with a barrier of ~3 kcal/mol. For comparison, we examined the protonation of several amines, reported previously to have  $pK_a$ 's shifted by 4-5 units, with the same approach. 10b However, little to no barrier was observed computationally in each case, likely because these species are more basic compared to the alcohol, precluding meaningful comparisons (see Supporting Information for details).

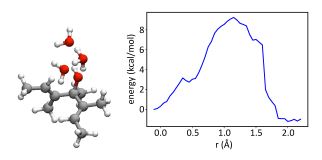


FIGURE 5. MD snapshot (cage and external water molecules removed) and proton transfer scan. r = the difference between the substrate C-O bond and the forming O-H bond. QM region (B3LYP/6-31G\*) includes all atoms shown in figure 4; MM region includes the cage and a water box (not shown in figure).

# Leaving Group Activation - Actually Leaving

What of leaving group loss? Continuing along the scan shown in Figure 5, disconnection of the leaving group inside the cage occurs, resulting in the encapsulated pentamethylpentadienyl cation + 4  $\rm H_2O$ . The barrier for this process is predicted here to be  $\sim\!6.5$  kcal/mol, somewhat higher than expected. In the absence of the cage, once the reactant hydroxyl group is protonated, departure of water is expected to occur with little to no barrier. A TSS was not located for this process (modelling such structures is notoriously difficult at best), $^{25}$  but a constrained calculation with the breaking bond fixed at 1.9 Å led to a predicted barrier of only 2.5 kcal/mol (note also that in the coiled conformer of O-protonated pentamethylpentadienol, the C–O bond is predicted to be approximately 1.7 Å long). This barrier is

similar to the barrier of  $\sim$ 4 kcal/mol estimated previously for this type of reaction.<sup>2b</sup> Nonetheless, the overall barrier from alcohol to pentamethylpentadienyl cation is predicted to be significantly lower (reduced by  $\sim$ 7 kcal/mol) in the cage than in water (Figure 6).

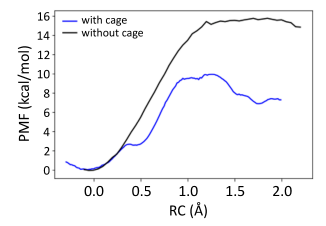


FIGURE 6. Proton transfer scan (RC = reaction coordinate, PMF = potential mean force) in water and in the cage.

# **Changing the Cage Vertices - Predictions and Experimental Validation**

The barrier to form pentamethylpentadienyl cation is predicted to be 2-3 kcal/mol higher for a cage with Si corners than for a cage with Ga corners (compare Figures 6 and 7), which we ascribe to an electrostatic effect that discourages proton transfer to the substrate, since that moves positive charge further from the cage corners (see SI for additional details). This predicted barrier difference is consistent with the experimental observation of a 680-fold reduction in rate associated with changing Ga to Si (for  $\bf 1$  with two  $\bf Z$  alkenes). <sup>26</sup> If our simple electrostatic model is reasonable, a rate difference also should be observed for the In/Ge pair of catalyst cages, but not the Ga/In and Si/Ge pairs.

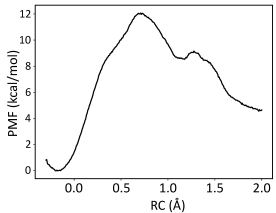


FIGURE 7. Proton transfer scan for cage with Si corners (RC = reaction coordinate, PMF = potential mean force) in water and in the cage.

#### Synthesis of New Cages

To further confirm the calculated contribution of the charged vertices of the cage to leaving group activation, we sought to compare hosts of different overall charge and hosts in which overall charge is maintained but the identity of the metal is changed.

As mentioned above, the metal centers of  $Ga_4L_6$  have previously been replaced to form a catalytically active Si-based host. While it was envisioned that the metal centers could be further diversified to other metals with varied oxidation states, this task presented a major synthetic challenge due to the extreme sensitivity of self-assembly to changes in the structural components. Other catalytically active homogeneous host systems rarely show variability in the metal center. We report here the syntheses of catalytically active  $Si_4L_6$ ,  $In_4L_6$ , and  $Ge_4L_6$  hosts, which were enabled by an optimized and generalizable guest-exchange protocol (Figure 8). We then conducted a comprehensive experimental study to confirm the structural similarity of these assemblies and determine the contribution of their coordinatively saturated metal centers to supramolecular catalytic reactivity.

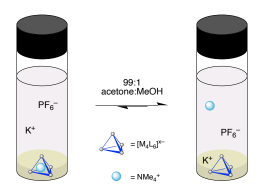


FIGURE 8. Scheme representing guest exchange procedure for replacing NMe<sub>4</sub><sup>+</sup> with K<sup>+</sup>. M = In(III), Ge(IV), or Si(IV). x = 12, 8, or 8, respectively.

We sought to synthesize the In<sub>4</sub>L<sub>6</sub> host to provide an isostructural and homo-anionic comparison to Ga<sub>4</sub>L<sub>6</sub>. Assembly of In<sub>4</sub>L<sub>6</sub> was initially probed by combining six equivalents of ligand L with four equivalents of indium tris-acetylacetonate in the presence of tetraethylammonium bromide, a strongly binding template. The appearance of six aromatic resonances and a diagnostic upfield-shifted guest resonance in the <sup>1</sup>H NMR spectrum confirmed host assembly and encapsulation of the guest (see supporting information). While this was a promising result, a templating guest inhibits host catalysis, and its subsequent removal is hampered by its high association constant. Identifying conditions for a template-free synthesis, however, is often the most challenging step toward accessing these supramolecular catalysts, given the reduced thermodynamic driving force toward host assembly.

Not surprisingly, assembly of  $In_4L_6$  was not observed in the absence of a templating guest, even upon prolonged heating at 50 °C in methanol. We thus sought an alternative approach, where a weakly binding cationic template such as tetramethylammonium could be used to first enable self-assembly and subsequently be displaced by potassium counterions to expose the catalytically active host cavity. The

guest exchange relies on a large excess of potassium ions to drive the binding equilibrium toward K\*-associated host, effectively overcoming the higher association constant of the cationic template. This approach has been previously employed for NMe<sub>4</sub>\* $\subset$ Ga<sub>4</sub>L<sub>6</sub>, which resulted in relatively poor recovery (36–41% yield) due to the similar solubility profiles of the host and excess KI (the K\* source).<sup>27</sup>

Fortunately, assembly of NMe<sub>4</sub>+⊂ In<sub>4</sub>L<sub>6</sub> was observed in the presence of a super-stoichiometric amount of tetramethylammonium bromide. Subjecting NMe<sub>4</sub>+⊂In<sub>4</sub>L<sub>6</sub> to guest exchange conditions identical to those used for Ga<sub>4</sub>L<sub>6</sub>, however, afforded only trace yield (<5%) of K+-associated In<sub>4</sub>L<sub>6</sub>, prompting an extensive screen to identify conditions for a better protocol. Ultimately, stirring NMe<sub>4</sub>+⊂ In<sub>4</sub>L<sub>6</sub> in a saturated solution of KPF6 in acetone and a small amount of methanol overnight yielded optimal exchange and recovery (Figure 8). Differences in solubility between the host and excess KPF<sub>6</sub> under the reaction conditions enabled nearly quantitative recovery of In<sub>4</sub>L<sub>6</sub> by filtration, and after 5 repeated iterations of the exchange protocol less than 0.1 equivalents of tetramethylammonium remained, as confirmed by <sup>1</sup>H NMR spectroscopy (after two rounds of guest exchange, less than 0.2 equivalents of NMe4+ remained, vielding catalytically active host). The stoichiometry and overall charge of In<sub>4</sub>L<sub>6</sub> was further verified by electrospray mass spectrometry (ESI-MS), where it was primarily detected in the 3- and 4- charge states with various counterions. Compared to the Ga<sub>4</sub>L<sub>6</sub> analogue, In<sub>4</sub>L<sub>6</sub> shows an increased sensitivity to oxidation and other decomposition pathways, which was evidenced by rapid attenuation of signal on the ESI-MS.

Next, we turned to the synthesis of Ge<sub>4</sub>L<sub>6</sub>, which we hoped would provide an isostructural and homo-anionic comparison to Si<sub>4</sub>L<sub>6</sub>. While template-free assembly of Ge<sub>4</sub>L<sub>6</sub> was not observed, NMe<sub>4</sub>+⊂Ge<sub>4</sub>L<sub>6</sub> could be formed by heating six equivalents of the ligand and four equivalents of Ge(IV) methoxide in DMF in the presence of five equivalents of NMe<sub>4</sub>Br. NMe<sub>4</sub>+⊂Ge<sub>4</sub>L<sub>6</sub> was subjected to the modified guest exchange conditions, which afforded K+-associated Ge<sub>4</sub>L<sub>6</sub>, albeit in lower yields. This decrease in yield was attributed to differences in solubility between In<sub>4</sub>L<sub>6</sub> and Ge<sub>4</sub>L<sub>6</sub>: the lower 8- charge enhances the solubility of Ge<sub>4</sub>L<sub>6</sub> in organic solvents such as acetone, while slightly decreasing its water solubility. Gratifyingly, addition of a small amount (1 mL) of water to the saturated KPF6 solution in acetone resulted in quantitative recovery of Ge<sub>4</sub>L<sub>6</sub>, which was characterized by <sup>1</sup>H NMR spectroscopy and ESI-MS, detected in the 4- and 5charge states.

While the template-free synthesis of  $Si_4L_6$  has been shown, it required elevated temperatures, long reaction times (60 h), and rigorous purification. We envisioned that applying the guest exchange protocol would not only serve as a proof-of-concept for the general applicability of the method, but could also enable assembly under milder conditions. Indeed,  $NMe_4$ \* $CSi_4L_6$  was quantitatively formed after just 16 h, and guest exchange conditions afforded clean  $Si_4L_6$ , which could be used without further purification.

Structural data obtained by single crystal X-ray diffraction (SCXRD) confirmed that the cavity size and metal-metal distances were very similar between  $Ga_4L_6$ ,  $Si_4L_6$ , and  $Ge_4L_6$ . Crystals of  $NEt_4^+ \subset Ge_4L_6$  were grown from vapor diffusion of

benzene into a solution in DMSO and measured with synchrotron radiation. Like  $Ga_4L_6$  and  $Si_4L_6$ ,  $Ge_4L_6$  is an ideal tetrahedron, with an average edge length of  $12.717(3)\mbox{\normalfont A}$  and a volume of 242.39(4)  $\mbox{\normalfont A}^3$  (Figure 9). Although we could not obtain SCXRD measurements for  $In_4L_6$ , we can expect similarly small deviations in the size and structure of the host, as the difference in metal-oxygen bond lengths between these four elements (Ga, Si, Ge, and In) is small (<0.5\mbox{\normalfont A}) compared to the edge length of the tetrahedron. These data demonstrate that the metal-ligand coordination environment is well-preserved between the hosts, confirming their isostructural relationship in the solid state.

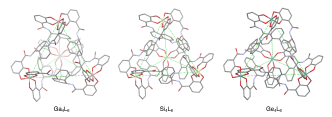


FIGURE 9. Structures obtained by SCXRD of  $Ga_4L_6$ ,  $Si_4L_6$ , and  $Ge_4L_6$ . Metal-metal distances are highlighted in green. For  $Ga_4L_6$ , average Ga-Ga distance is 12.663(3)Å, and volume is 239.3(1)Å $^3$ . For  $Si_4L_6$ , average Si-Si distance is 12.655(2)Å, and volume is 238.85(6)Å $^3$ . For  $Ge_4L_6$ , average Ge-Ge distance is 12.717(3)Å, and volume is 242.39(4)Å $^3$ .

In contrast, total fluorescence yield (TFY) of the O K-edge soft X-ray absorption spectroscopy (sXAS) data of the four hosts are shown in Figure 10. Because all the metal ions have filled d manifolds, the pre-edge features of the O K-edge sXAS (between 530-535 eV) mainly reflect the transitions from O 1s orbitals to the empty orbitals near LUMO with O p characters.<sup>29,30</sup> The differences in energy splitting and relative intensities imply different electronic structures of the LUMOs, although they all have similar geometric structures. The existence of such differences is supported by the varied assembly conditions that were required when the metal vertices were changed.

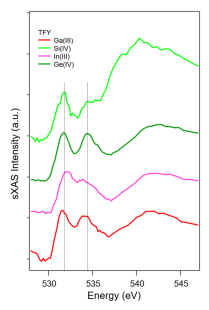


FIGURE 10. Oxygen K-edge sXAS data of Ga<sub>4</sub>L<sub>6</sub>, Si<sub>4</sub>L<sub>6</sub>, In<sub>4</sub>L<sub>6</sub>, and Ge<sub>4</sub>L<sub>6</sub>.

# **Catalytic Rates**

With these four isostructural assemblies consisting of two homo-anionic pairs (12- and 8-) in hand, we sought to compare their catalytic activity and observe the effect of changing the metal vertices on the host-catalyzed Nazarov cyclization. Our aim was to determine whether the structural similarity between the hosts would induce similar catalytic behavior, or if the electronic differences in metal-oxygen binding would cause divergence. Previous comparison of the Ga<sub>4</sub>L<sub>6</sub> and Si<sub>4</sub>L<sub>6</sub> catalyzed aza-Cope rearrangement, which proceeds with retention of cationic charge on the substrate and is driven primarily by constrictive binding within the host, exhibited rates of reaction within error of each other. In<sub>4</sub>L<sub>6</sub> and Ge<sub>4</sub>L<sub>6</sub> were also shown to catalyze the aza-Cope rearrangement at similar rates to Ga<sub>4</sub>L<sub>6</sub> and Si<sub>4</sub>L<sub>6</sub>, further confirming that structurally similar hosts show no difference in reactivity for a reaction that is not sensitive to changes in host charge (see Supporting Information). In contrast, the Nazarov cyclization is initiated by protonation of a neutral dienol substrate, resulting in an overall increase in cationic charge within the host microenvironment. Comparison between the Ga<sub>4</sub>L<sub>6</sub> and Si<sub>4</sub>L<sub>6</sub> catalysts demonstrated a 680-fold difference in rate acceleration, implying that the reaction is indeed sensitive to host charge.<sup>26</sup> Further investigation of the In<sub>4</sub>L<sub>6</sub> and Ge<sub>4</sub>L<sub>6</sub> catalyzed reactions should thus enable us to probe the extent to which the identity of the metal affects charge-based supramolecular catalytic reactivity, and validate whether the substitution of Al for Ga and Si in the computational model is appropriate.

We investigated the Nazarov cyclization of dienol 1 (with two Z alkenes), which previously demonstrated sensitivity to changes in overall host charge (Scheme 1). Due to the large discrepancy in rate between the  $Ga_4L_6$  and  $Si_4L_6$  catalyzed reactions, kinetic data were previously obtained at different temperatures and extrapolated for comparison using an Eyring analysis. Analogously, for this study, we discovered that while the  $In_4L_6$  catalyzed reaction could be monitored over the course of a few hours at  $45\,^{\circ}C$ , the  $Ge_4L_6$ 

catalyzed reaction required heating to 72 °C to measure the reaction rate on a practicable experimental timescale. This initial observation led us to investigate the four hosts as homo-anionic pairs, where the Ga<sub>4</sub>L<sub>6</sub> and In<sub>4</sub>L<sub>6</sub> catalyzed reactions were compared at 45 °C, and Si<sub>4</sub>L<sub>6</sub> and Ge<sub>4</sub>L<sub>6</sub> catalyzed reactions were compared at 72 °C. The rate constant ( $k_{\rm obs}$ ) at 45 °C for the In<sub>4</sub>L<sub>6</sub> catalyzed reaction was found to be 8.3 × 10<sup>-4</sup> s<sup>-1</sup>, compared to 4.2 × 10<sup>-4</sup> s<sup>-1</sup> for Ga<sub>4</sub>L<sub>6</sub>. At 72 °C, the  $k_{obs}$  were found to be 1.7 × 10<sup>-4</sup> s<sup>-1</sup> for Ge<sub>4</sub>L<sub>6</sub>, compared to 2.2 × 10<sup>-4</sup> s<sup>-1</sup> for Si<sub>4</sub>L<sub>6</sub> (Table 1).

Table 1. Rate constants ( $k_{\rm obs}$ ) for the host-catalyzed Nazarov cyclization.

Catalyst	Catalyst Loading (%)	Temperature	<i>k</i> <sub>obs</sub> (s <sup>-1</sup> )
[Ga <sub>4</sub> L <sub>6</sub> ] <sup>12–</sup>	7%	45 °C	4.2(3)×10 <sup>-4</sup>
[ln <sub>4</sub> L <sub>6</sub> ] <sup>12-</sup>	10%	45 °C	8.3(1)×10 <sup>-4</sup>
[Si <sub>4</sub> L <sub>6</sub> ] <sup>8-</sup>	7%	72 °C	2.2(4)×10 <sup>-4</sup>
[Ge <sub>4</sub> L <sub>6</sub> ] <sup>8-</sup>	6%	72 °C	1.7(1)×10 <sup>-4</sup>

Though the host-dienol Michaelis complex is clearly observable by  $^1$ H NMR spectroscopy, encapsulation is not quantitative due to the high dissociation constant ( $K_d$ ) of the neutral alcohol substrate. While substrate dependence studies have not been conducted for the host-catalyzed Nazarov cyclization, it is reasonable to assume that the reaction is not in the saturation regime under these conditions. Thus, the first-order  $k_{obs}$  is dependent upon the value of the Michaelis constant ( $K_m$ ), which can be approximated by  $K_d$ . Because small hydrocarbon molecules are encapsulated by the hydrophobic effect, rather than Coulombic attraction, their association is driven primarily by entropic effects.  $^{31}$  Because of this, we expect negligible difference in  $K_d$  of  $\mathbf{1}$  between the two hosts, as host charge plays a minimal role as a driving force for association.

Evidently, the discrepancies in the reaction rates based on host charge reaffirms the ability of the metal centers to impact catalysis by imparting an overall charge on the host assembly. However, the close agreement in rates between the respective tri- and tetravalent metal centers provide evidence that key electrostatic interactions between the host and the transition state are not significantly perturbed between different homo-cationic metals, consistent with the calculated predictions. These experimental comparisons enable us to pinpoint the specific role of the metal center in catalysis, despite the complexity of structure activity relationships in multi-component, self-assembled systems.

The development of an efficient guest-exchange protocol enabled high-yielding syntheses of catalytically active Si<sub>4</sub>L<sub>6</sub>, In<sub>4</sub>L<sub>6</sub>, and Ge<sub>4</sub>L<sub>6</sub> supramolecular hosts. Together with Ga<sub>4</sub>L<sub>6</sub>, these isostructural catalysts enabled an extensive experimental investigation to isolate the effects of the metal center, an important structural component, on catalytic reactivity. All hosts demonstrated similar catalytic ability for the constrictive-binding driven aza-Cope rearrangement regardless of identity or oxidation state of the metal center. While In<sub>4</sub>L<sub>6</sub> and Ge<sub>4</sub>L<sub>6</sub> hosts demonstrated significant discrepancies in rate ( $k_{obs}$  = 8.3 × 10<sup>-4</sup> s<sup>-1</sup> at 45 °C vs. 1.7 × 10<sup>-4</sup> s

 $^{\rm 1}$  at 72 °C) for the Nazarov cyclization, reactivity was comparable between hosts with homo-cationic metal centers, in agreement with predictions. These results imply that the structural similarity between the hosts determined their similar catalytic behavior, outweighing the differences in electronic structures implied by the LUMOs with 0 p characters indicated by the sXAS data.

#### Overall Model

Thus, we arrive at the following model for rate acceleration. Protonation of the reactant alcohol is greatly enhanced upon complexation, leading to a large net acceleration in water loss. Electrocyclization may be enhanced, but only slightly. Warshel and co-workers similarly concluded that selective transition state stabilization was much less significant than reactant protonation for an orthoformate hydrolysis reaction promoted by  $Ga_4L_6$  cage.  $^{5a,10a}$ 

Evidence continues to mount that the reactivity of carbocations generated in the active sites of terpene synthases reflect their energetics in the gas phase.1d This was predicted on the basis of results of quantum chemical computations, and multiple experiments with enzymatic systems support this model.32 Here we have shown that a very similar scenario occurs for the Ga<sub>4</sub>L<sub>6</sub> cage; the cage facilitates formation of a pentadienyl carbocation, but barrier lowering is not at all necessary for the cyclization reaction step. While Type 1 terpene synthase enzymes activate their substrates' leaving groups (diphosphate groups) primarily by binding to Lewis acidic magnesium ions,1a the Ga<sub>4</sub>L<sub>6</sub> cage activates its substrates' leaving groups via facilitation of protonation. Protonation is also used in other (Type 2) terpene synthases to generate carbocations from epoxides or alkenes.33 In addition to utilizing active site amino acid sidechains with enhanced acidity (resulting from hydrogen-bonding arrays in which they participate), these terpene synthases provide active site cavities that stabilize the carbocations generated upon protonation. Like the Ga<sub>4</sub>L<sub>6</sub> cage, these active sites are lined with aromatic amino acid sidechains that can participate in carbocation- $\pi$  interactions, 3d,22,34 however the cage also appears to bind several water molecules along with the substrate.5b-c While waters do sometimes bind along with substrates for terpene synthases, they tend to be fewer in number.

## CONCLUSION

Supramolecular catalysis systems remain challenging to model, due to the large number of atoms involved and the significant influence of noncovalent and solvent interactions. Our work shows agreement between calculation and experiment on structure-activity relationships, a promising example of the computational modeling of a complex supramolecular system.

# **ASSOCIATED CONTENT**

## **Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website.

Details of computational methods and structures (PDF)

General procedures, synthesis and characterization of compounds, general procedures for kinetics experiments, representative spectra and kinetics data, ESI-MS data, crystallographic analysis of  $Ge_4L_6$ , and  $^1H$  NMR spectra (PDF)

Crystallographic data for Ge<sub>4</sub>L<sub>6</sub> (CIF)

CCDC 2088585 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge via <a href="www.ccdc.cam.ac.uk/data request/cif">www.ccdc.cam.ac.uk/data request/cif</a>, or by emailing <a href="data request@ccdc.cam.ac.uk">data request@ccdc.cam.ac.uk</a>, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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# TOC Graphic

