Entropy directs the self-assembly of supramolecular palladium coordination macrocycles and cages

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Entropy directs the self-assembly of supramolecular palladium coordination macrocycles and cages†

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The self-assembly of palladium-based cages is frequently rationalized via the cumulative enthalpy (ΔH) of bonds between coordination nodes (M, i.e., Pd) and ligand (L) components. This focus on enthalpic rationale limits the complete understanding of the Gibbs free energy (ΔG) for self-assembly, as entropic (ΔS) contributions are overlooked. Here, we present a study of the M3,L5 intermediate species (M = dinitrato(N,N,N',N'-tetramethylethlenediamine)palladium(n)), in L = 4,4'-bipyridine), formed during the synthesis of triangle-shaped (M3,L3) and square-shaped (M4,L4) coordination macrocycles. Thermochemical analyses by variable temperature (VT) 1H-NMR revealed that the M3,L5 intermediate exhibited an unfavorable (relative) ΔS compared to M3,L3 (triangle, ΔTΔS = +5.22 kcal mol⁻¹) or M4,L4 (square, ΔTΔS = +2.37 kcal mol⁻¹) macrocycles. Further analysis of these constructs with molecular dynamics (MD) identified that the self-assembly process is driven by ΔG losses facilitated by increases in solvation entropy (ΔSolv, i.e., depletion of solvent accessible surface area) that drives the self-assembly from ‘open‘ intermediates toward ‘closed‘ macrocyclic products. Expansion of our computational approach to the analysis of self-assembly in Pd(L),ben,cL8) cages (ben = 4,4’-(5-ethoxy-1,3-phenylene) dipyrindine), demonstrated that ΔSolv contributions drive the self-assembly of both thermodynamic cag products (i.e., Pd3,T,ben,cL6) and kinetically-trapped intermediates (i.e., Pd3,T,ben,cL8).

Introduction

Self-assembled macrocycles,† cage, and oligomers featuring palladium-coordination nodes are a mainstay of supramolecular chemistry due to their unique mechanical, optoelectronic, and catalytic applications. The square-planar coordination geometry of the coordination node permits the synthesis of well-defined constructs invoking simple geometric design principles. These constructs are formed by the spontaneous self-assembly of palladium-based coordination nodes with ditopic ligands, affording highly-ordered assemblies bearing the minimum Gibbs free energy (ΔG). The synthesis of these assemblies from their many constituent components is rationalized by the cumulative enthalpic (ΔH) contributions as a result of the formation ofative bonds between ditopic ligands and Pd-based coordination nodes. However, studies of gas-phase reactivity reveal these ΔH contributions decrease with sequential coordination of additional ligands, favoring the formation of coordinatively unsaturated oligomers. In addition, currently only entropic contributions (ΔS) based on statistical arguments are taken into account, implying that assemblies based on a large number of components are entropically largely unfavorable despite many synthetic examples. Whereas similar enthalpic arguments rationalize the difference in energy between various assemblies, it remains unclear why well-defined structures are favored over oligomers as these coordination bonds in principle can be formed with limited constraints. The current rationale for deducing self-assembly thermodynamics is therefore incomplete, which motivates our current study.

The self-assembly of equimolar amounts of dinitrato(N,N,N',N'-tetramethylethlenediamine)palladium(n) (M) and 4,4'-bipyridine (L) affords a mixture of triangular and square macrocycles in equilibrium (Scheme 1). Previous reports have leveraged this experimentally accessible equilibrium to measure the relative ΔS and ΔH of triangular and square complexes. These studies found that ΔS favors the assembly featuring fewer components (i.e., triangles) while ΔH favors the geometric matching between the square-planar metal center and the ligand geometry (i.e., squares).

The synthesis of (and conversion between) macrocyclic assemblies proceeds via coordinatively unsaturated oligomeric intermediates (Scheme 1, purple).

Interestingly, similar stable oligomer intermediates have been found in the synthesis of polygonal organometallic...
microcyclic assemblies. The equilibrium between these stable oligomeric intermediates and macrocyclic products may be leveraged to quantify $\Delta H$ and $\Delta S$ contributions to self-assembly using the literature described NMR-based approach. Realization of the origin and effect of these thermodynamic contributions enables rational improvement of the self-assembly of highly-ordered constructs used broadly in supramolecular chemistry, including coordination cages, metal–organic frameworks, and dynamic-covalent based constructs.

In this report, we demonstrate that the self-assembly of an equimolar mixture of $\text{M}$ and $\text{linL}$ in deuterated dimethyl sulfoxide (DMSO) affords a mixture of triangular ($\text{M}_3\text{linL}_3$) and square ($\text{M}_4\text{linL}_4$), and oligomeric ($\text{M}_5\text{linL}_{5}$) assemblies as depicted in Scheme 1. We employed variable temperature $^1$H-NMR (VT-NMR) to determine the $\Delta H$ and $\Delta S$ of both oligomeric and macrocyclic assemblies, providing unprecedented thermodynamic insights into the self-assembly process. Importantly, this thermodynamic data-enabled validation of a molecular dynamics (MD) based approach to distinguish respective $\Delta S$ contributions arising from the assembly structure ($\Delta S_{\text{struct}}$, eqn S1$^\dagger$) and its solvation ($\Delta S_{\text{solv}}$, eqn S2$^\dagger$). These individual entropic contributions, alongside calculation of $\Delta H$ (eqn S3$^\dagger$), ultimately provide an accurate $\Delta G$ for self-assembly. We applied our MD-based approach to the study of coordination cages based on 4,4′-bipyridine (ben$\text{L}_4$) and free palladium(II) ions (Pd$^{2+}$), which have been reported in the literature (Scheme 2).

Thermodynamic estimates derived from MD simulations reveal a $\Delta S_{\text{solv}}$-driven, self-assembly process for macrocycles and cages reminiscent of biopolymer folding. The generalization of our MD-based approach may distinguish between kinetically accessible thermodynamic products (i.e., Pd$_{12}$$\text{benL}_{24}$) and undesirable kinetically-trapped intermediate assemblies (e.g., Pd$_8$$\text{benL}_{16}$).

These computational and experimental studies demonstrate that $\Delta S$ drives the self-assembly of supramolecular constructs featuring palladium coordination nodes. As this $\Delta S$ contribution arises from solvation, these findings broadly reflect the thermodynamic drive of self-assembly to form compact supramolecular structures. Furthermore, we demonstrate the utility of MD-based approaches to quantify the thermodynamics of large supramolecular systems, providing a methodology that enables in silico studies of self-assembly processes.

### Results

#### Synthesis and characterization of assemblies based on $\text{M}$ and $\text{linL}$

Previously, we reported that the absence of trace halide impurities during the self-assembly of coordination cages resulted in slower formation kinetics, giving rise to the observation of intermediates. Thus, we developed an alternative preparation for $\text{M}$, using a limiting quantity of palladium dichloride to minimize trace chloride (Scheme S1). With this prepared metal precursor, the self-assembly of stoichiometric quantities of $\text{M}$ and $\text{linL}$ ($\text{M} : \text{linL} = 1.0 : 1.0$, $[\text{M}] = 17.1$ mM) affords a mixture of macrocyclic ($\text{M}_3\text{linL}_3$ and $\text{M}_4\text{linL}_4$) and oligomeric products evidenced by characteristic $^1$H-NMR peaks shown in Fig. 1.

Peaks corresponding to $\text{M}_3\text{linL}_3$ ($\delta = 9.24$ ppm) and $\text{M}_4\text{linL}_4$ ($\delta = 9.40$ ppm) were consistent with the reported values of these macrocyclic species. Two additional peaks ($\delta = 9.32$–9.37 ppm) present, with chemical shifts consistent with reported oligomers, and a single diffusion constant ($D = 1.56 \times 10^{-10}$ m$^2$ s$^{-1}$, Fig. 1 inset). These features indicate the presence of a single coordination assembly with a size larger than free $\text{linL}$ ($D = 1.86 \times 10^{-10}$ m$^2$ s$^{-1}$) but smaller than $\text{M}_3\text{linL}_3$ ($D = 1.20 \times 10^{-10}$ m$^2$ s$^{-1}$). We also observed a near 2 : 1 ratio of $\text{X}$-pyridyl peak areas...
Fig. 1 Macroyclic and oligomeric products observed from the self-assembly of M (17.1 mM) and \(^{15}\text{NtL} (17.1 \text{ mM}) in DMSO characterized by unique \(\alpha\)-pyridyl peaks in \(^1\text{H-NMR} \) (Fig. S1f), with inset DOSY (Fig. S2f) diffusograms. Three distinct assemblies were observed and highlighted for clarity: \(M_2\text{linL}_3\) (slate blue), \(M_4\text{linL}_4\) (pink), and \(M_2\text{linL}_3\) (purple). Numerical data is provided below in Table 1.

**Table 1** \(^1\text{H-NMR} \) characterization of assemblies based on M and \text{linL}\(^a\)

<table>
<thead>
<tr>
<th>Assembly</th>
<th>(\delta_{\alpha\text{-pyridyl}}) (ppm)</th>
<th>(\delta_{\beta\text{-pyridyl}}) (ppm)</th>
<th>% Area</th>
<th>Diffusion, (D) ((\times 10^{-10} \text{ m}^2 \text{ s}^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>(M_2\text{linL}_3)</td>
<td>9.24</td>
<td>8.28</td>
<td>63.9</td>
<td>1.20</td>
</tr>
<tr>
<td>(M_4\text{linL}_4)</td>
<td>9.40</td>
<td>8.21</td>
<td>21.2</td>
<td>1.10</td>
</tr>
<tr>
<td>(M_2\text{linL}_3)</td>
<td>9.37</td>
<td>8.24</td>
<td>9.9</td>
<td>1.58</td>
</tr>
<tr>
<td>(M_4\text{linL}_4)</td>
<td>9.32</td>
<td>8.19</td>
<td>5.0</td>
<td>1.58</td>
</tr>
<tr>
<td>\text{linL} (d)</td>
<td>8.75</td>
<td>7.86</td>
<td>—</td>
<td>1.86</td>
</tr>
</tbody>
</table>

\(^a\) \(^1\text{H-NMR} \) Conditions: DMSO, 300 K, 300 MHz. \(^b\) Outer ligands, i.e., \text{linL}, \text{M}_{\text{linL}}-\text{L-M}_{\text{linL}}-\text{linL}. \(^c\) Inner ligand, i.e., \text{linL}-\text{M}_{\text{linL}}-\text{M}_{\text{linL}}-\text{linL}. \(^d\) Reference values were obtained separately for free \text{linL} (Fig. S3).

Table 2 Experimental thermodynamic differences between \(M_2\text{linL}_3\), \(M_3\text{linL}_3\), and \(M_4\text{linL}_4\) coordination assemblies

<table>
<thead>
<tr>
<th>Assembly</th>
<th>(\Delta H^\circ) ((\text{kcal mol}^{-1}))</th>
<th>(-T\Delta S^\circ) ((\text{kcal mol}^{-1} \text{ K}^{-1}))</th>
<th>(\Delta G^\circ) ((\text{kcal mol}^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>(M_2\text{linL}_3)</td>
<td>0.00 (\pm 0.20)</td>
<td>-2.80 (\pm 0.19)</td>
<td>-2.80 (\pm 0.02)</td>
</tr>
<tr>
<td>(M_4\text{linL}_4)</td>
<td>-1.11 (\pm 0.29)</td>
<td>+0.05 (\pm 0.28)</td>
<td>-1.06 (\pm 0.03)</td>
</tr>
<tr>
<td>(M_4\text{linL}_4)</td>
<td>-3.38 (\pm 0.34)</td>
<td>+2.42 (\pm 0.32)</td>
<td>-0.96 (\pm 0.31)</td>
</tr>
</tbody>
</table>

\(^a\) Values determined for \(T = 298.15 \text{ K}\) by direct fitting a modified van't Hoff model (Scheme S2) to VT-NMR derived \(\Delta G\) values (Fig. 2).

(14.9%, Table 1), assuming the overlap of \(\alpha\)-pyridyl protons adjacent to coordination, we assigned these peaks to an oligomer with the composition \(M_2\text{linL}_3\).

While previous studies rationalized that the self-assembly process is driven to maximize the number of coordination bonds formed, affording coordinatively saturated species that minimize the \(\Delta H\) of the system.\(^1\) However, analysis by Weilandt et al. on mononuclear Pd complexes demonstrated that the formation of successive coordination bonds result in diminishing \(\Delta H\) contributions to the \(\Delta G\) of complex formation, which was partially compensated by \(\Delta S\).\(^26\) Our observation of a significant presence of oligomeric (coordinatively unsaturated) assemblies (14.9%, Table 1), we infer that \(\Delta S\) may play a similarly significant role in macrocycle assembly.

**Thermochemical analysis of macrocycle–oligomer equilibria**

Following the literature, we employed VT-NMR to quantify the relative abundance of assemblies \(\{M_2\text{linL}_3, M_3\text{linL}_3, \text{and } M_4\text{linL}_4\}\) by monitoring the intensity of their unique \(\alpha\)-pyridyl peaks (Table 1) over a wide range of temperatures (297.5–350.0 K, see Fig. S12–S16).\(^6\) To determine the relative \(\Delta G\) of \(M_2\text{linL}_3, M_3\text{linL}_3, \text{and } M_4\text{linL}_4\) we modeled the system as three orthogonal equilibria between each product and a common pool of reactants (Scheme S2f). These relative \(\Delta G\) values (Table S1f) were plotted as a function of temperature (Fig. 2), and directly fit with an expanded van ‘t Hoff equation to compute \(\Delta S\) and \(\Delta H\) (Scheme S2f).\(^38\)

Consistent with literature observations, these experimentally established \(\Delta S\) and \(\Delta H\) values confirm that \(M_2\text{linL}_3\) and \(M_4\text{linL}_4\) are the respective \(\Delta S\)- and \(\Delta H\)-favored macrocyclic products.\(^3\) Interestingly, \(M_2\text{linL}_3\) features a lower \(\Delta H\) than the \(\Delta H\)-favored \(M_4\text{linL}_4\) (Table 2, \(\Delta H = -2.27 \text{ kcal mol}^{-1}\)) and an elevated \(-T\Delta S\) compared to the \(\Delta S\)-favored \(M_2\text{linL}_3\) (Table 2, \(T\Delta S = +5.22 \text{ kcal mol}^{-1}\)). These results are contrary to established geometric and component-number based rationale for \(\Delta H\)- and \(\Delta S\)-favored products.\(^1,4,5\)

We surmise that the \(\Delta H\) of \(M_2\text{linL}_3\) derives from the conformational freedom of the structure, allowing the adoption of an unstrained configuration (\(\angle \text{N-Pd-N} = 90^\circ\)) following the geometric rationale established for \(M_4\text{linL}_4\). However, \(M_4\text{linL}_4\) exhibits an internal strain relative to \(M_2\text{linL}_3\) manifesting as the \(\Delta H\)-difference between the two complexes (\(\Delta \Delta H = -2.27 \text{ kcal mol}^{-1}\)). As both assemblies are presumed to adopt a conformation where \(\angle \text{N-Pd-N} = 90^\circ\), the apparent \(\Delta \Delta H\) is
not accounted for when a simple geometric rationale is invoked. Moreover, ΔS-differences between the 6-component $M_{i}^{\text{lin}L_{3}}$ and 5-component $M_{j}^{\text{lin}L_{3}}$ contrast the typical rationale, which correlates the integration of fewer components to a favorable ΔS. These findings highlight how the current rationale for determining ΔS and ΔH contributions is insufficient to account for oligomeric assemblies, necessitating further computational investigation into the origins of internal strain found in $M_{i}^{\text{lin}L_{4}}$ and the unexpected ΔS penalties associated with $M_{j}^{\text{lin}L_{4}}$ formation.

**MD analysis of experimental assemblies**

Weilandt et al. suggested that two discrete ΔS factors that exert significant influence on metal–organic complex formation. The first is ΔS_struct, which decreases as more molecules (i.e., components) are required to form a complex. The other is ΔS_solv, which decreases as more solvent molecules are required to solvate a complex. Using a previously described methodology, we developed parameters to simulate assemblies comprised of M and $\text{lin}L$ with accurate ΔH contributions (Fig. S4†). Using these parameters, and GBSA model solvation, 50 ns trajectories (Fig. S5–S7†) were propagated by MD for $M_{i}^{\text{lin}L_{3}}$, $M_{i}^{\text{lin}L_{4}}$, and $M_{j}^{\text{lin}L_{4}}$ assemblies (Fig. 3). These trajectories were then used to compute the ΔS_struct, ΔS_solv, and ΔH contributions to ΔG (Table 3).

The thermodynamic contributions to ΔH and ΔS computed from these simulations (Table 3) differ those obtained by VT-NMR (Table 2) in their absolute value. However, the differences (i.e., ΔΔG, ΔΔH, or ΔΔS) between assemblies measured by simulation and experiment are very similar (see below). The differences in absolute value reflect the different reference states in experimental and computational measurements (Fig. S20†). The reproduction of the relative differences in these physical quantities validates our in silico methodology for the thermodynamic values of these and similar assemblies.

The ΔH difference between $M_{i}^{\text{lin}L_{3}}$ and $M_{j}^{\text{lin}L_{4}}$ measured by VT-NMR (ΔΔH_exp = +2.28 kcal mol$^{-1}$, Table 2), is similar to our MD-derived results (ΔΔH_MD = +2.20 kcal mol$^{-1}$, Table 2). As ΔH generally originates from molecular geometry, we infer that the $M_{i}^{\text{lin}L_{4}}$ adopts a geometrically unfavorable (i.e., strained) configuration compared to $M_{i}^{\text{lin}L_{3}}$. Visualization of MD trajectory data for $M_{i}^{\text{lin}L_{4}}$ assemblies reveals that this oligomer prefers a zig-zag conformation with a near-ideal square-planar coordination geometry at the Pd center (∠N–Pd–N = 89°, Fig. 3c). In contrast, visualization of $M_{i}^{\text{lin}L_{4}}$ reveals a folded-square structure that features a hyperbolic geometry (i.e., ∠N–Pd–N = 86°, Fig. 3b) giving rise to an internal strain that is enthalpically unfavorable (i.e., elevates ΔH). Additional simulations of $M_{i}^{\text{lin}L_{4}}$ (performed in vacuo) reinforce that these distortions are a consequence of the solvation incurred to minimize the solvent-accessible surface area (Fig. S8†).

The ΔS difference between $M_{i}^{\text{lin}L_{4}}$ and $M_{j}^{\text{lin}L_{4}}$ measured by VT-NMR (ΔΔS_nmp = +2.37 kcal mol$^{-1}$, Table 2), is similar to our MD-derived results (ΔΔS_MD = +2.37 kcal mol$^{-1}$, Table 2), while other comparisons values have acceptable deviation (Fig. S20†). The calculated −ΔS_struct values from our MD-approach decrease with the number of components (i.e., $M_{i}^{\text{lin}L_{3}} < M_{i}^{\text{lin}L_{4}} < M_{j}^{\text{lin}L_{4}}$), in line with reported trends. 1 While, the computed difference of ΔS_struct for $M_{i}^{\text{lin}L_{3}}$ and $M_{j}^{\text{lin}L_{4}}$ are
similar to experimental values (Table 2), differences between both macrocycles vs. \( M_{\text{lin}L_4} \) deviate significantly (Table 3). The inclusion of \(-\Delta S_{\text{solv}}\) improves the accounting for \( \Delta S \) differences between \( M_{\text{lin}L_3} \), \( M_{\text{lin}L_2} \), and \( M_{\text{lin}L_4} \) assemblies (Table 3). This leads us to infer that experimental \(-\Delta S\) penalties associated with \( M_{\text{lin}L_3} \) formation originate from these \( \Delta S_{\text{solv}} \) contributions, in agreement with thermodynamic studies of mono-nuclear Pd complexes.\(^{26}\)

These thermodynamic parameters demonstrate that \( \Delta S \)-specifically \( \Delta S_{\text{solv}} \)-drives the conversion of oligomeric intermediates (i.e., \( M_{\text{lin}L_3} \)) to their macrocyclic product assemblies (i.e., \( M_{\text{lin}L_3} \) and \( M_{\text{lin}L_4} \)). Moreover, the effect of \( \Delta S_{\text{solv}} \) may overcome \( \Delta H \) contributions, resulting in strained and distorted molecular geometries. While chemists have previously exploited \( \Delta H \) to direct the formation of desired constructs, these findings reveal that \( \Delta S \) ultimately drives the synthetic process of multi-component self-assembly.

**MD modeling of arbitrary \( M_{\text{int}L_3} \) and \( M_{\text{int}L_{n+1}} \) assemblies**

Oligomers similar to (and including) \( M_{\text{lin}L_4} \) are theorized to form as intermediates in the self-assembly process of larger supramolecular structures (Scheme 1). Therefore, we utilized our MD-based approach to compare a range of oligomeric intermediates (\( M_{\text{int}L_{n+1}} \), \( n = 1-28 \)) and the potential macrocyclic products (\( M_{\text{lin}L_{n+1}} \), \( n = 2-28 \)) to elucidate the role of individual thermodynamic parameters (\( \Delta S_{\text{solv}}, \Delta S_{\text{struct}}, \) and \( \Delta H \)) on \( \Delta G \) for the self-assembly of macrocycles (Fig. 4).

The resulting simulations reveal that both macroyclic (Fig. 4a) and oligomeric (Fig. 4b) assemblies exhibit increasing unfavorably \( \Delta G \) with increasing assembly size driven by \( \Delta S_{\text{struct}} \) contributions. The limited range of assemblies observed by NMR measurements (i.e., \( M_{\text{lin}L_2}, M_{\text{lin}L_3} \), and \( M_{\text{lin}L_4} \)) is rationalized by the elevated \( \Delta G \) experienced for other possible structures. This outcome is consistent with ESI-HRMS analysis (Fig. S9) that provides qualitative evidence for the existence of larger assemblies in low abundance (i.e., low signal-to-noise).

The value of \(-\Delta S_{\text{struct}}\) increases with size (Fig. 4, blue trace) for the self-assembly of both oligomeric and macrocyclic products, consistent with the decreased degrees-of-freedom experienced upon aggregation.\(^{27}\) Intriguingly, we find a nonlinear correlation between the size and \( \Delta H \) of oligomeric assemblies that is absent for macrocyclic congeners. Visualization of MD trajectory data reveals that larger oligomer assemblies adopt a compact conformation (Fig. S10), resulting in increased strain (i.e., \( \Delta H \) penalty) on the palladium-aryl bonds compared to the zig-zag conformation found in smaller assemblies such as \( M_{\text{lin}L_3} \) (Fig. 3c). We infer that these compact suprastructures are necessary to realize a more compact assembly, akin to the folded structure observed for \( M_{\text{lin}L_4} \) macrocycles (Fig. 3b). This trade-off between \(-\Delta S_{\text{solv}}\) and \( \Delta H \) originates from solvation and distinguishes oligomeric assemblies from macrocyclic ones. As \(-\Delta S_{\text{solv}}\) favors the formation of compact suprastructures, it is reasonable to deduce that the self-assembly of product macrocycles, in general, is driven by \( \Delta S_{\text{solv}} \) contributions.

**MD modeling of reticular cage self-assembly**

Mixtures of bent ditopic ligands (\( \text{benL} = 4,4'-(5\text{-ethoxy-1,3-phenylene})\text{dipryridine} \)) and free palladium(II) coordination nodes (Pd) self-assemble to afford cuboctahedral cages (Pd\(_{12}\text{benL}_{24}\)) via oligomeric-assembly intermediates. We employed our MD-based approach to gain insight into the self-assembly process,\(^{27}\) as incomplete or partial cages (i.e., intermediate assemblies) are currently inaccessible by experimental methods (e.g., CSI-HRMS, \(^{1}H\)-NMR).\(^{28}\) We developed models for intermediates of self-assembly (Pd\(_n\text{benL}_{mn}\), where \( 2n \leq m \leq 3n + 1 \)) from contiguous sections extracted from Pd\(_{12}\text{benL}_{24}\), Pd\(_8\text{benL}_{16}\) and Pd\(_{15}\text{benL}_{30}\) cages, as well as a polymeric state (Pd\(_n\text{benL}_{mn}\) where \( m = 3n + 1 \)) geometries. Then we elucidated the self-assembly pathway for these partially-formed constructs from their respective \( \Delta S, \Delta H, \) and \( \Delta G \) values from MD-simulation (Fig. 5).

Our simulations reveal the Pd\(_{12}\text{benL}_{24}\) exhibits a lower \( \Delta H \) (i.e., minimal geometric strain) compared to congeneric
assemblies (Table 4). This observation is consistent with the literature and originates from the decreased metal–ligand bond strain experienced by this particular assembly-configuration.⁵⁻¹¹,⁴⁹,⁵⁵ Models of partially-formed assemblies (e.g., Pd₅[^benL₁₄]) bear an elevated ΔH (Fig. 5a, blue trace) as a result of strain originating from deflation or collapse during MD simulations (Fig. S11f). Parallel observations have been made in M[^¹[^linL₄]] macrocycles (Table 2), inferring increases in strain can act to offset penalties from solvation entropy (i.e., ΔS_{solv}), which leads to an overall elevation in ΔH for the system. The sum of entropic contributions (i.e., ΔS_{struct} + ΔS_{solv} = ΔS, Fig. 5a, green trace) suggests that the formation of early intermediates (n < 7) is hindered while the self-assembly of spherical cages (n = 12) is encouraged. These results demonstrate that while ΔH directs the polyhedral geometry of the final assembly,ΔS drives the structure of self-assembly to be spherical.

The comparison of the free energy (ΔG) pathways for the self-assembly of different topologies (Fig. 5b) enables us to distinguish between the thermodynamic product (i.e., Pd₁₂[^benL₂₄]), kinetic traps (e.g., Pd₈[^benL₁₆]), and unrealized topologies (e.g., Pd₁₅[^benL₁₅], red trace). The maximum ΔG of intermediate assemblies of the Pd₁₅[^benL₁₅] pathway (ΔG_{max} = +3.44 kcal mol⁻¹) is greater than that of the Pd₁₂[^benL₂₄] cage (ΔG_{max} = +2.62 kcal mol⁻¹), rationalizing the possibility that intermediates may spontaneously reconfigure towards the latter structure as it is thermodynamically favored. In contrast, the Pd₈[^benL₁₆] pathway features the lowest maximum energy (ΔG_{max} = +2.17 kcal mol⁻¹) of analyzed pathways demonstrating that kinetic traps are readily identified by our MD-based approach.

**Conclusion**

The thermochemical analysis of the self-assembly processes in palladium-based coordination macrocycles revealed unexpected ΔS-contributions that drive the formation of higher-order macrocycle assemblies (M[^¹[^linL₄]] and M[^¹[^linL₄]]) from oligomer intermediates (M[^²[^linL₄]]). Using an MD-based approach, we found that the driving force for self-assembly originates from the solvation entropy (i.e., ΔS_{solv}) of oligomeric intermediates that effects surface-area minimization of the construct. Thermodynamic trends were established by MD analysis of larger assemblies, revealing that both ΔS_{solv} and ΔS_{struct} direct the formation of assemblies that exhibit similar ΔH. Data from MD models of formation pathways for palladium-based coordination cages reveal that ΔS_{solv} is responsible for driving the self-assembly process. Further application of our MD approach enables rationalization of the formation Pd₁₂[^benL₂₄] cage products over kinetically trapped congeners (i.e., Pd₈[^benL₁₆]) directly from the computed thermodynamic quantities (ΔS_{solv}, ΔS_{struct}, and ΔH) of the intermediate assemblies. Overall, these complementary experimental and computational investigations expose ΔS as the driver for the formation of these desirable highly ordered structures that have broad applications across supramolecular chemistry.

**Author contributions**

DAP conceived, designed and performed the experiments. EOB contributed ligand materials, performed HR-MS analyses and provided experimental expertise for self-assembly experiments. JNHR and SM supervised the work. DAP, SM, and JNHR wrote
the manuscript with all authors providing significant contributions to the analysis and interpretation of the work.

Conflicts of interest

There are no conflicts to declare.

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