

Transient equilibrium mapping: A new tool for measuring the thermodynamics of slowly assembling supramolecular systems

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Supramolecular chemistry involves the non-covalent assembly of monomers into materials with unique properties and wide-ranging applications. Thermal analysis is one of the key analytical tools in this field, as it provides quantitative thermodynamic information on both the structural stability and nature of the underlying molecular interactions. However there exist many supramolecular systems whose kinetics are so slow under conditions approaching chemical equilibrium that thermodynamic data are inaccessible. We have developed a simple, general, and rapid spectroscopic method for extracting thermodynamic parameters from such slowly assembling systems. It is based on repeatedly raising and lowering the temperature during assembly and identifying the points of transient equilibrium as they are passed on the up- and down-scans. In a proof-of-principle application to the slow co-assembly of polydeoxyadenosine containing 15 adenosines (polyA₁₅) and cyanuric acid (CA), we found that roughly 30% of the CA binding sites on the polyA chains were unoccupied, with implications for the assembly of high-valence systems.

Introduction

Supramolecular chemistry is emerging as a rich source of diverse materials with novel and valuable properties. Potential applications range from drug-delivery and tissue regeneration to optical sensors and organic electronics.¹ This approach involves the non-covalent self-assembly of tens to thousands of monomeric units into larger structures with emergent physical properties that derive from both the structures of the individual components and their interactions and arrangement with respect to one another.² Reversible assembly has some distinct advantages compared to traditional covalent synthesis. The dynamic nature of supramolecular interactions allows bonds to break and reform leading to materials with self-healing properties. Furthermore, many supramolecular systems have the ability to generate multiple morphologies and sets of physical properties from a single set of building blocks with only small modifications of the assembly conditions.³ Nevertheless, there are unique challenges associated with this approach. Chief among these is characterizing the products of a non-covalent assembly reaction. Much of the excitement surrounding supramolecular chemistry comes from the fact that there remains much to be understood regarding the relationships between the chemical structures of the monomeric units, the supramolecular architectures, and the emerging physical properties, and there is wide possibility for new and unexpected discoveries. However, this implies that the nature of supramolecular products is difficult to predict and that rigorous structural and thermodynamic analyses are critical to advancing the field.

A variety of tools have been used to elucidate the structures produced by assembly, including atomic force, electron, and super-resolution microscopies, and solid-state NMR spectroscopy.⁴⁻⁶ The stabilities of the assemblies are most commonly measured by thermal analysis. Most supramolecular structures dissociate when they are heated and reassemble when the monomer mixtures are cooled. This process can be quantified either by calorimetry⁷ or by spectroscopically-detected melting and annealing.⁸ Detailed analyses of melting curves provide substantial insight into the forces holding together the supramolecular structures and yield additional thermodynamic parameters such as the enthalpies, ΔH , entropies ΔS , and free energies, ΔG , of critical steps in the assembly process.⁹ However, there exists a subset of supramolecular systems for which thermal analysis fails to give reliable thermodynamic data due to extremely slow assembly kinetics. Such systems only assemble or disassemble at useful rates when they are pushed far from equilibrium, i.e. under very highly stabilizing or destabilizing conditions. Common examples include amyloid fibrils, viral capsids, and a variety of self-assembling non-biological small molecules.¹⁰⁻¹⁶ Thermodynamic analyses rely on assembled and disassembled monomers existing at their equilibrium distributions over the range of temperature measured during the experiment. Slowly assembling systems require so much time to equilibrate that this is not achievable on practical timeframes using standard melting and annealing experiments. For this reason, rigorous thermodynamic information is largely missing for slowly assembling systems, hampering progress and discouraging their study.

We have developed a new experimental approach we term transient equilibrium (TE) mapping that gives rigorous thermodynamic data on slowly assembling systems and can be performed using a standard temperature-controlled spectrophotometer. Rather than waiting for the system to equilibrate (which can take days or weeks), the temperature is repeatedly raised and lowered, driving cyclic, non-equilibrium disassembly and assembly. Since the T_m lies between the starting and ending temperatures of the scans,

the system briefly passes through a moment of transient equilibrium on each up-scan and down-scan. The transient equilibria (temperature and concentration values) can be clearly identified from the spectroscopic trace, allowing the full equilibrium melting curve to be mapped out in just a few hours.

For example, we applied TE mapping to better understand the recently discovered co-assembly of polydeoxyadenosine (polyA) and the small molecule cyanuric acid (CA) into fibres whose biocompatibility and low cost make them promising candidates for tissue engineering and drug delivery.¹⁷ A cross-section of the proposed structure (*Figure 1*) shows the adenosine of three different DNA strands hydrogen bonding to CA molecules in a continuous supramolecular helicene.¹⁸ We note that the ideal helicene structure has a 1:1 ratio of dA residues and CA molecules. We recently characterized the kinetics of polyA-CA fibre assembly using non-equilibrium melting methods.¹⁴ Equilibration of the fibres near the melting point could take up to a month of constant instrument use. Using TE mapping, we determined the ΔG , ΔH , and ΔS values for adding a polyA chain to the end of a growing fibre in a single 10-hour experiment. By repeating these measurements at different concentrations of CA, we determined the minimum polyA:CA ratio necessary for assembly and made the surprising discovery that about 30% of the available CA binding sites are unfilled under our conditions. These results have implications for the future development of asymmetric systems involving components of very different valences, such as polyA and CA, and demonstrate the potential of the TE approach for learning about slowly assembling systems.

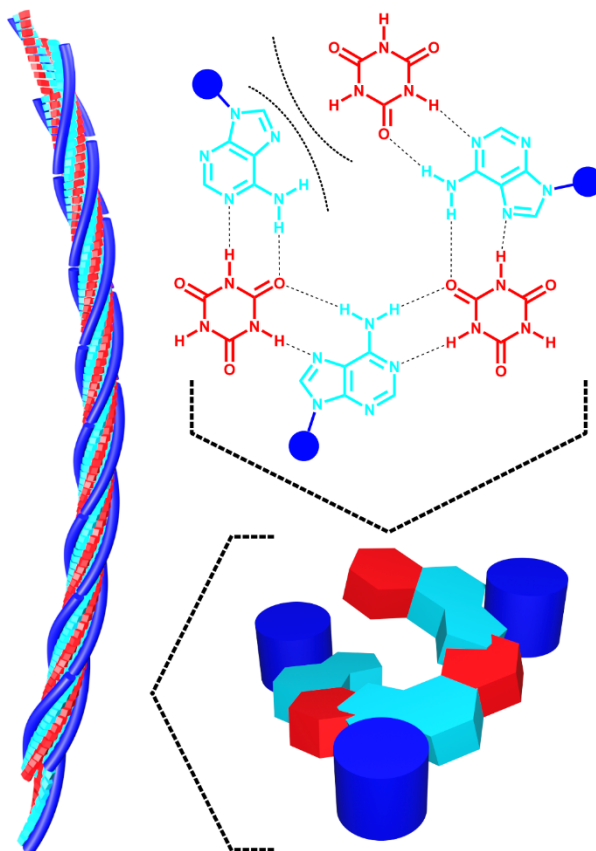


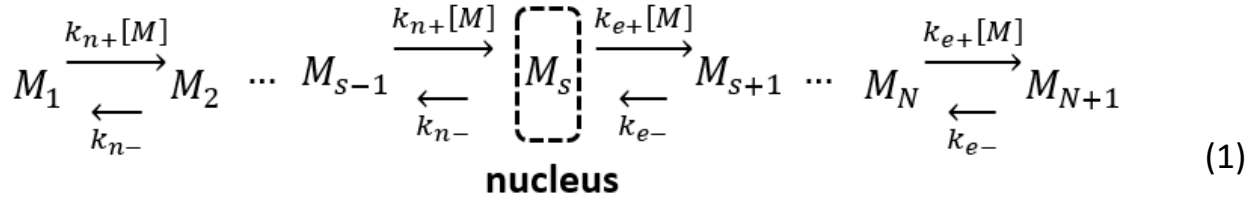
Figure 1: Proposed structure of supramolecular fibers formed from the co-assembly of polyadenosine strands and cyanuric acid (left). A cross section of a single hexameric helicene (right).

Results and Discussion

Theory

We previously characterized the kinetics of polyA-CA fibre assembly using thermal hysteresis melting and annealing experiments in which the temperature is varied and the concentration of free monomers is detected spectroscopically. For slowly assembling systems, such as polyA-CA, the rate at which the system relaxes to equilibrium is far slower than the temperature scan rate, thus both folding (cooling) and unfolding (heating) occur out of equilibrium. The populations effectively lag behind the changing temperature such that the cooling and heating scans are offset, with the apparent T_m reached at a higher temperature than the true T_m on the up-scan and at a lower temperature than the true T_m on the down-scan, as illustrated in *Figure 2a*. The resulting hysteresis loops can be analyzed to give folding and

unfolding rates and activation energies in the case of 2-state folding reactions.⁹ For fibres, as is the case here, hysteresis data can be fit to more complicated assembly schemes,^{14, 17} such as the Goldstein-Stryer (GS) cooperative kinetic model, where M_N describes a fibre with N polyA chains:¹⁹



Association and dissociation of polyA monomers (i.e. dA₁₅ oligodeoxynucleotides, M_1) and short oligomers less than the critical nucleus size, s , are described by the nucleation rate constants k_{n+} and k_{n-} respectively, while oligomers larger than s are described with the elongation rate constants k_{e+} and k_{e-} . All four rate constants are typically assumed to obey Arrhenius kinetics with activation energies E_{n+}^a , E_{n-}^a , E_{e+}^a , and E_{e-}^a . Fits to the hysteresis data give the four rate constants and four activation enthalpies. In principle, these values contain thermodynamic information. For example, the ratio $k_{e-}/k_{e+} = K_e$ is the equilibrium dissociation constant for adding a polyA chain to the end of an elongating fibre. The differences $\Delta H_e = E_{e-}^a - E_{e+}^a$ and $\Delta S_e = \Delta H_e/T + R \ln(K_e)$ are the enthalpy and entropy changes for the same process, respectively. In practice, however, thermodynamic parameters are very poorly defined by the hysteresis data. To illustrate, simulated GS fibres with equilibrium melting temperatures spanning over 10 degrees can produce virtually superimposable hysteresis data (Figure 2a). Similarly, experimental hysteresis data for polyA-CA fibres give visually very similar fits, even when ΔH_e and ΔS_e values are varied by a factor of 1.6

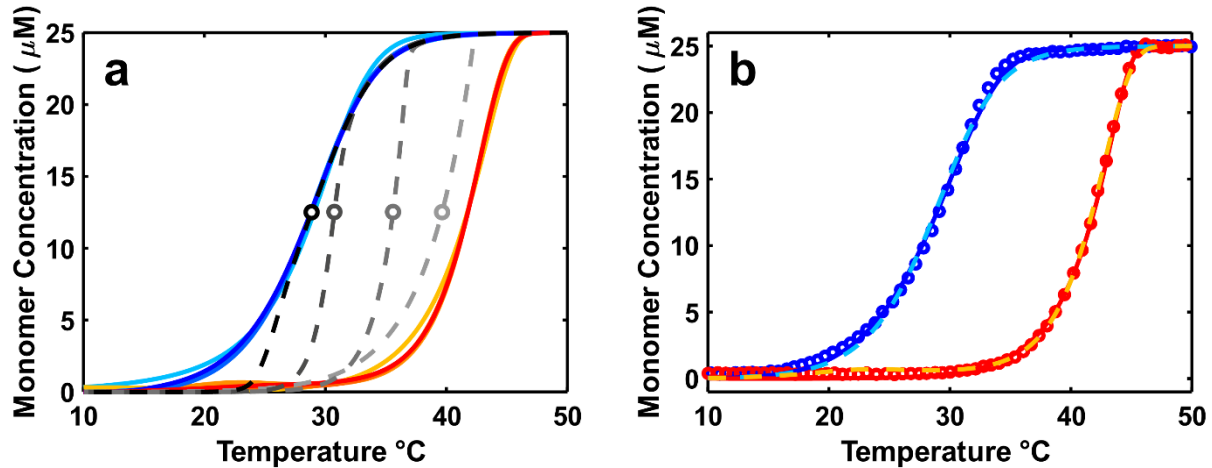


Figure 2: **a)** Fiber assembly/disassembly simulated using the Goldstein-Stryer model and kinetic parameters that give similar melting and annealing curves (solid lines) with drastically different equilibrium curves (dashed lines). Heating curves are shown in red/orange and cooling curves are shown in blue/cyan. The offset between heating and cooling data is due to thermal hysteresis (TH). Simulation parameters are listed in Supplementary Table 1 **b)** Fits of experimental TH curves with ΔH_e extracted from TH data alone (blue and red solid lines) and ΔH_e constrained to be equal to the value extracted from TE measurements (dashed cyan and orange lines). Simulation parameters are listed in Supplementary Table 2

(Figure 2b, Supplementary Table 1). Thus new approaches are needed to accurately determine thermodynamic parameters for slowly-equilibrating systems.

Recent work from the Yamaguchi lab¹⁶ has explored how the spectra of slowly equilibrating, self assembling systems respond to repeated heating and cooling cycles.²⁰ Depending on the starting and ending temperatures and ramp rates, a rich diversity of shapes (thermal hysteresis loops) have been observed, providing qualitative information on the underlying assembly reactions. However, to date there has not been a straightforward way to extract quantitative thermodynamic information from these data. Our new TE mapping approach uniquely fills this gap. In order to illustrate the fundamental principles, we performed kinetic simulations using the GS assembly model and parameters for polyA-CA fibres (Figure 3a, see Supplementary Information). The dashed black line indicates the concentration of monomers that would be present after extensive equilibration, i.e. the critical monomer concentration, $[M]_c$, as a function of temperature. This curve is rich in thermodynamic information and can be analyzed to obtain ΔH_e , ΔS_e ,

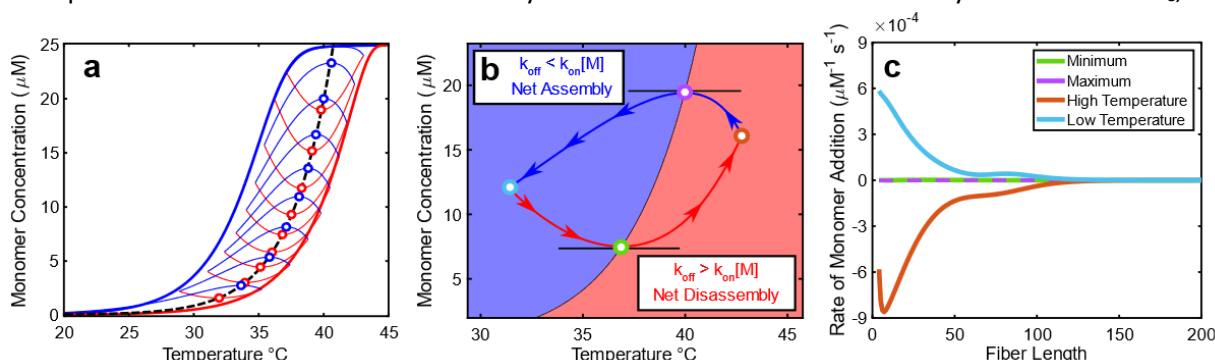


Figure 3: a) Kinetic simulations of a typical hysteresis experiment (bold lines) and TE mapping experiment (narrow lines). Cooling traces are shown in blue, heating traces are shown in red. The experiment begins by cooling from 45°C to 36°C, this is followed by the first up-scan (36°C to 44°C), a second down-scan (44°C to 35°C), a second up-scan (35°C to 43°C), a third down-scan (43°C to 34°C), etc. The equilibrium profile is shown as the dashed black line, with the extrema of each TE cycle shown as points. **b)** An isolated TE mapping cycle: assembly occurs only in the blue shaded region; disassembly only occurs in the red shaded region. The interface of these two regions represents a system at equilibrium. Coloured points represent the position of calculated monomer flux in panel c. **c)** Calculated monomer flux of fibers for points shown in panel b, the horizontal extrema of the TE cycle have 100-fold less flux than either the high or low temperature values.

and K_e , as well as the corresponding parameters for fibre nucleation.²¹ For a rapidly-equilibrating system, the $[M]_c$ curve can be measured directly by traditional melting or reannealing experiments.²¹ In contrast, for the slowly assembling polyA-CA system, heating and cooling scans give the left- and right-shifted blue and red curves, respectively, and the location of the $[M]_c$ curve is obscured by the thermal hysteresis, as illustrated in Figure 3a.

The TE mapping method is based on our discovery that repeatedly raising and lowering the temperature such that it repeatedly traverses the true T_m reveals the precise locations of the hidden equilibria. Simulating TE data for polyA-CA assembly gives a series of concave-up and concave-down arcs on the heating and cooling scans, respectively (narrow red and blue curves in Figure 3a). Strikingly, the $[M]_c$ values (black line) pass directly through the extrema (concentration maxima and minima) of the cooling and heating arcs. Thus experimentally-determined extrema can be interpreted as a set of $[M]_c(T)$ values.

The physical process underlying this behaviour can be understood as follows: for cooperatively assembled fibres, such as polyA-CA, equilibrium is reached when the rate of monomer addition to the end of a fibre ($k_{e+}[M]_c$) is exactly equal to the rate of monomer dissociation from the end of a fibre (k_{e-}), such that the net rate of fibre growth is zero.²¹ When $[M_1] < [M]_c$ there is net dissociation and $[M_1]$ increases with time, corresponding to the red region below the $[M]_c$ curve in *Figure 3b*. When $[M_1] > [M]_c$ there is net association and $[M_1]$ decreases with time, corresponding to the blue region above the $[M]_c$ curve. Every cooling scan starts in the red region with net dissociation (increasing $[M_1]$) and ends in the blue region with net association (decreasing $[M_1]$). As the temperature crosses the boundary where $[M_1]=[M]_c$, net fibre growth is zero, the arc is exactly horizontal, and the maximum is reached. Conversely, every heating scan starts in the blue region with decreasing $[M_1]$ and ends in the red region with increasing $[M_1]$. As the temperature crosses the $[M_1]=[M]_c$ boundary, the free monomer concentration is at a minimum. To validate this interpretation, we calculated the net rate of monomer addition to each length of fibre in the simulation. At the lower and upper limiting scan temperatures (orange and cyan), the rates of monomer addition and release are at least 100-fold greater than at the horizontal extrema of the heating and cooling arcs (green and purple) (*Figure 3c*). Furthermore, since $k_{e+}[M]_c=k_{e-}$ at all of the extrema, this implies that the experimental monomer concentrations at these points are equal to the equilibrium dissociation constant $K_e=k_{e-}/k_{e+}$. In other words, thermodynamic parameters can be read directly from TE data.

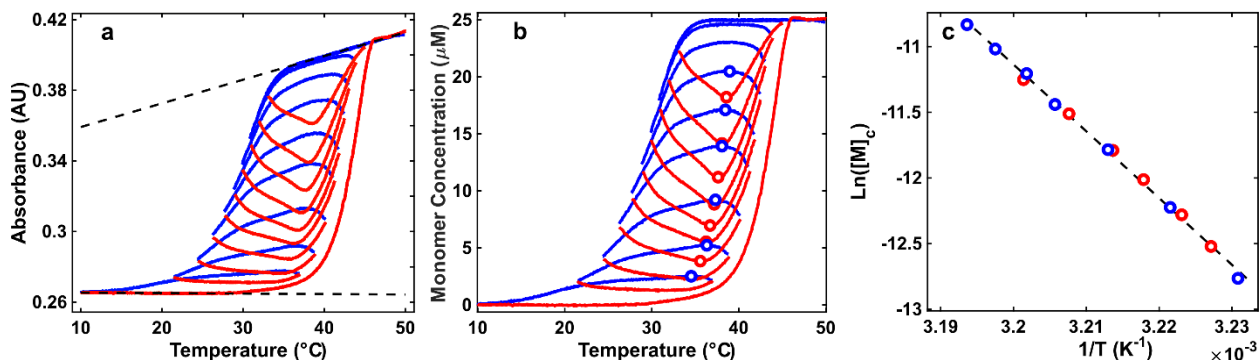


Figure 4: **a)** Raw absorbance data for a 15mer polyA-CA coassembly with 25 μM polyA₁₅ and 15 mM CA at pH 4.5, blue lines represent cooling traces and red lines represent heating traces. Unfolded (top black line) and folded (bottom black line) are also shown. **b)** Processed TE data with extrema of each cycle shown as points. **c)** Van 't Hoff analysis of experimental TE points, plotted along with confidence interval of one standard deviation.

Analysis of experimental TE data

We performed a TE experiment on a mixture of CA and polyA chains by repeatedly raising and lowering the temperature over a window of roughly 15°C that shifted from (29-44) to (14-33) °C in 9 cycles while we monitored the spectroscopic absorbance at 260 nm (*Figure 4a*, see *Supplementary Information* for a guide to selecting sliding T-windows). The lower and upper absorbance regions were fitted to linear baselines and assigned 100% and 0% folded, i.e. $[M_1]=0$ and 25 μM , respectively. The converted data are shown in *Figure 4b*, with blue and red indicating cooling and heating, respectively, and open circles placed at the extrema. These experimental arcs have a remarkable similarity to the calculations shown in *Figure 3a*. The y- and x-values of the extrema correspond directly to critical monomer concentration, $[M]_c$, and temperature pairs. As discussed above, $[M]_c$ values are equivalent to the equilibrium dissociation

constant, K_e , for adding a polyA to the end of an elongating fibre, for this system. A van 't Hoff plot of $\ln([M]_c) = \ln(K_e)$ vs $1/T$ is linear with a slope of $-\Delta H_e/R$ and y-intercept of $\Delta S_e/R$ (Figure 4c), giving $\Delta H_e = 100 \pm 2 \text{ kcal mol}^{-1}$ and $\Delta S_e = 301 \pm 7 \text{ cal mol}^{-1} \text{ K}^{-1}$. Notably, although the values of ΔH_e and ΔS_e determined by TE mapping differ from those obtained by kinetic fits to TH data by factors of 1.6 (Supplementary Table 2), repeating the TH analysis with ΔH_e and ΔS_e fixed to the TE-derived values gives good agreement with experimental data (Figure 2b), illustrating the insensitivity of the kinetic fits to these thermodynamic parameters. In general, we would strongly recommend that, even if assembly kinetics are the main interest, the combination of TE and thermal hysteresis experiments provide more robust solutions than thermal hysteresis alone, as TE resolves ambiguity in the fitted rate constants and ratios thereof.

Furthermore, the thermodynamic parameters provide a basis for comparing polyA-CA fibres to other nucleic acid structures. For example, polyA/polyT (dA₁₅dT₁₅) duplex dissociation is predicted to have approximately $\Delta H = 108 \text{ kcal mol}^{-1}$ and $\Delta S = 335 \text{ cal mol}^{-1} \text{ K}^{-1}$ under similar solution conditions to those used here.²² It is intriguingly similar to the values we measured for polyA-CA assembly ($100 \text{ kcal mol}^{-1}$ and $301 \text{ cal mol}^{-1} \text{ K}^{-1}$). At first glance, we would have expected polyA-CA fibres to show much higher enthalpies and entropies than dAdT duplexes, since there are three strands rather than two, each dA forms twice as many hydrogen bonds and immobilizes a CA molecule in the putative polyA-CA structure (Figure 1). However partial vacancy of CA binding sites may help to reconcile these observations, as elaborated below.

Stoichiometry of polyA-CA fibres.

One of the great advantages of quantitative thermodynamic data is that much can be learned about the system of interest through careful analyses of how energetic parameters vary with changing conditions. For instance, the presumptive structure of polyA-CA fibres shows that one molecule of CA is present for every deoxyadenosine residue in each polyA chain. In other words, when one of the dA₁₅ polyA chains binds the end of an elongating fibre, it should be accompanied by 15 CA molecules. While equilibrium dialysis experiments are consistent with this structure,¹⁷ they have relatively low precision and the stoichiometry is very difficult to measure with accuracy. This property is of great interest since a CA:polyA stoichiometry of less than 15 would reveal the existence of defects, which could potentially be targeted with other small molecules. Thermodynamic data can help to resolve this issue, since the apparent dissociation constant, K_e , for a polyA chain binding to the end of the fibre should vary with CA concentration in a predictable way. For instance, if a polyA chain always brings with it c molecules of CA, i.e.



(following the nomenclature of Equation 1), then the full equilibrium dissociation constant for the process is given by

$$K_{(T)}^\circ = \frac{[M_n][M_1][CA]^c}{[M_{n+1}]} \quad (3)$$

This is something of an over-simplification, as elaborated below, but for now it serves to illustrate the dependence of K_e on [CA]. For polyA-CA fibres, CA is always in great excess so that its concentration is effectively constant for any set of assembly conditions. The apparent polyA dissociation constant K_e is related to the full equilibrium constant according to

$$K_e = \frac{[M_n][M_1]}{[M_{n+1}]} \Big|_{[CA]} = K_{(T)}^{\circ} [CA]^{-c} \quad (4)$$

with the temperature dependence of the standard equilibrium constant ($K_{(T)}^{\circ}$) described by

$$K_{(T)}^{\circ} = \exp \left(- \frac{(\Delta H_{(T)} - T \Delta S_{(T)})}{RT} \right) \quad (5)$$

Therefore, measuring K_e at a series of different CA concentrations should produce offset van 't Hoff plots where the vertical distance between each line follows the stoichiometry of CA. To proceed, we noted that stabilization of polyA-CA fibres at high [CA] is largely entropic in nature, since it is primarily driven by differences in the entropy of dilution when dissociation of a polyA chain concomitantly releases c molecules of CA into solution.

We repeated the TE experiment at four CA concentrations between 7.5 and 15 mM (*Supplementary Figure 1*). Van 't Hoff plots of the resulting K_e values are shown in *Figure 5*. Fitting *Equation 4* to this data set allows us to directly obtain the stoichiometry of CA. To account for the possibility of a temperature dependent enthalpy value we extracted global values of ΔH_e and ΔC_p . The heat capacity change of binding, ΔC_p , accounts for any temperature-dependent differences in the slopes of the different experiments according to:

$$\Delta H_e(T) = \Delta H_e(T_0) + \Delta C_p(T - T_0) \quad (6)$$

$$\Delta S_e(T) = \Delta S_e(T_0) + \Delta C_p \ln \left(\frac{T}{T_0} \right) \quad (7)$$

The extracted $\Delta C_p = -0.6 \pm 0.3 \text{ kcal mol}^{-1} \text{ K}^{-1}$ indicates that the enthalpy of adding a polyA chain to a growing fibre has only a slight temperature dependence. This is perhaps unsurprising, since ΔC_p values associated with nucleic acid folding are largely sequence dependent and have been observed to vary from slightly negative to positive values.²³ The global fit was in good agreement with experimental data points (*Figure 5*, *Supplementary Table 3*). Surprisingly, the extracted stoichiometry coefficient, $c = 10.4 \pm 0.6$, implies that 30% of possible CA binding sites are unoccupied in polyA-CA fibres under these conditions.

Master equations for high-valence assembly

The thermodynamics of multivalent supramolecular assembly can be summarized in terms of two main trends: the “principle of maximum occupancy” which refers to the tendency of systems to evolve toward the most stable state with full occupancy of binding sites, and the “entropy factor” which favours the state of the system with the largest number of product species.²⁴ For most of the supramolecular systems studied to date, the valency (number of binding sites per monomer) is relatively small (<6), the principle of maximum occupancy dominates, and the all sites are generally filled in the assembled materials.^{25 26} However, for high-valence monomers, such as the polyA chains studied here, the entropy factor strongly opposes the principle of maximum occupancy and more complex behaviour emerges. For example, each dA₁₅ chain creates an additional 15 potential CA binding sites, on average, as it adds to the end of growing fibre; one site must be created for each additional dA residue to achieve the theoretical 1:1 dA:CA stoichiometry. The number of ways to fill c of the 15 binding sites with c molecules of CA is given by the binomial coefficient²⁷

$$N_c = \frac{15!}{c!(15-c)!} \quad (8)$$

While there is only $N=1$ way completely fill all 15 binding sites ($c=15$), there exists a total of $N=32,766$ distinct ways fill the sites with $1 \leq c \leq 14$ molecules of CA. A simplified model of this energy diagram is seen in *Figure 6b*, where partially filled states are higher in energy but are more numerous. Therefore, even though a polyA chain with 15 bound CA molecules may represent the single lowest energy configuration, there exists such an enormous number of partly-filled configurations that these dominate, with a broad distribution of CA uptake and just 10 of the 15 sites being filled on average as seen in *Figure 6c*.

This explanation implies that polyA chains can bring a variable number of CA molecules with them when they attach to the end of a growing fibre, which is inconsistent with *Equation 4*, where the stoichiometry is fixed. To resolve this inconsistency, we developed a simple combinatorial model to describe polyA-CA fibre elongation. There is a free energy penalty for bringing an unbound polyA chain in close proximity to the end of a fibre, $\Delta G_{\text{polyA}} = \Delta H_{\text{polyA}} - T\Delta S_{\text{polyA}}$. This is compensated by energetically favourable binding of CA molecules to the newly-created 15 binding sites. All CA molecules are assumed to bind with equal free energy $\Delta G_{\text{CA}} = \Delta H_{\text{CA}} - T\Delta S_{\text{CA}}$. The total free energy change for a polyA chain binding along with a specific configuration of c CA molecules is $\Delta G_{\text{polyA}} + c\Delta G_{\text{CA}}$. Overall, the apparent equilibrium dissociation constant for polyA chain binding is given by²⁸

$$(K_e)^{-1} = K_{\text{polyA}}(1 + K_{\text{CA}}[\text{CA}])^{15} \quad (9)$$

where $K_{\text{polyA}} = \exp(-\Delta G_{\text{polyA}}/RT)$ and $K_{\text{CA}} = \exp(-\Delta G_{\text{CA}}/RT)$. The average number of CA molecules can be calculated using the following equation

$$\langle c \rangle = 15 \frac{K_{\text{CA}}[\text{CA}]}{1 + K_{\text{CA}}[\text{CA}]} \quad (10)$$

and the fraction of bound states with a given number of CA molecules can be calculated by

$$\theta_c = \left(\frac{15!}{c!(15-c)!} \right) \frac{K_{\text{CA}}[\text{CA}]^c}{(1 + K_{\text{CA}}[\text{CA}])^{15}} \quad (11)$$

We fit *Equation 9* to the TE data, obtaining excellent agreement, and extracting ΔH_{polyA} , ΔS_{polyA} , ΔH_{CA} , and ΔS_{CA} (*Figure 5*, *Supplementary Table 3*). These parameters allowed us to calculate the fractions of polyA chains with different numbers of CA molecules bound at different temperatures and $[\text{CA}]$, providing a highly detailed description of assembly (*Figure 6c*). Under highly stabilizing conditions of high $[\text{CA}]$ and

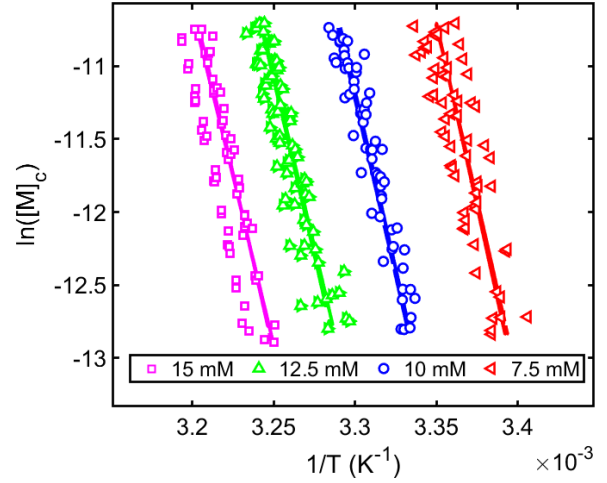


Figure 5: Van 't Hoff plot of TE data at different cyanuric acid concentrations. Coloured symbols represent experimental data from TE traces, solid-coloured lines represent a global fit of Equation 4 and dashed coloured lines represent a global fit of Equation 9. Solid and dashed lines are virtually superimposed on each other. Experimental errors are smaller than the size of the symbols.

low temperature, the Equations predict that almost all binding sites are filled, in agreement with previous dialysis experiments.¹⁷ Importantly, *Equation 9* and *Equation 10* explain why we observe 10 bound CA, and not more or less, even though experiments were performed at different [CA]. All experiments used 25 μM polyA, which means that we only detected K_e values between about 3 μM and 22 μM in all cases. This implies that the $K_{\text{CA}}[\text{CA}]$ values are nearly identical in all experiments (since K_{polyA} does not change much with temperature) From *Equation 10*, this implies that $\langle c \rangle$ is very similar in all experiments, ranging from 10 to 11, and in excellent agreement with the simple fit described in the previous section.

High valence supramolecular systems have many useful properties that are only just beginning to be explored, such as the ability to self-heal, responsiveness to stimuli, and simple, inexpensive chemical derivatization. Examples include small molecule-directed nucleic acid assembly (CA + polyadenosine or polydeoxyadeonsine^{14, 17}; melamine + polythymine²⁹) and non-covalent polymer crosslinking via multiple metal chelation^{25, 30} or host/guest interactions^{31, 32}. *Equation 9* and *Equation 10* can serve as starting points for quantitatively describing assembly in such systems, where simple probabilistic considerations ensure that some of the binding sites will remain vacant under many conditions. Furthermore, we find that TE-derived data are sufficient to extract the relevant thermodynamic parameters robustly, providing a new avenue for gaining insight into these complex materials.

Generality of the Method

Our aim for the TE mapping method is that it can be used as a general tool to determine the thermodynamic parameters of supramolecular assembly when standard thermal melting and annealing experiments are unsuitable for thermodynamic analysis. Towards this end, we have also tested the method on a tetrameric intermolecular guanine quadruplex (G4) in aqueous buffer, and zinc-porphyrin self-assembly in mixture of methylcyclohexane and chloroform. In both cases, we obtained series of concave-up and concave-down arcs, similar to those of the polyA-CA fibres (*Supplementary Figure 4*). In parallel, we used computer simulations to model the TE experiment for different types of self assembling systems and observed two patterns of behaviour: either all the extrema aligned with the equilibrium curve or the maxima for the cooling curves and minima for the heating curves were offset from one another (*Supplementary Figure 4*). This provides a useful guide for interpreting TE data on new systems of interest: when the extrema align, they can be used to trace out the equilibrium curve (as for polyA-CA fibres and the intermolecular G4). When they are offset, they cannot be directly equated to equilibrium temperature/concentration pairs (as for the zinc porphyrin system), although the data are still information-rich, as detailed in the *Supplementary Information*. Fortunately, many slowly assembling supramolecular structures are amenable to the TE approach and in these cases it provides thermodynamic information that is not readily available from other sources. For example, a polyA-CA $[\text{M}]_c$ dataset similar to the one reported here would require a scan rate of $<0.001 \text{ K min}^{-1}$ in traditional melting measurements, leading to experiments on the impractically long timescale of a month. Our study demonstrates how the ready availability of high-quality thermodynamic dynamic data can lead to new insights, such as the prevalence of unfilled CA binding sites in polyA-CA fibres, and provides an opportunity to test theoretical developments, such as our master equation for high-valence assembly. These advances would not have been realistically possible for polyA-CA structures using previously-existing methods. It is worthwhile noting that quite a few other types of slowly assembling systems have been reported including viral capsids, amyloid fibrils and other supramolecular polymers,^{10, 33, 34} and that other slowly-equilibrating systems such as spin-crossover^{35, 36} and nanoparticle agglomeration³⁷ could benefit from the TE mapping approach.

Furthermore, we may be substantially underestimating the number of potentially interesting slowly assembling systems. Anecdotally, molecules that initially show hysteresis in thermal melts are often disfavoured for further characterization, since adequate tools for thermodynamic analysis have been lacking, leading to underreporting of these types of structures. We believe that the TE mapping method presented here is a big step towards filling this gap in the experimentalist's toolbox. It can be applied to a wide variety of systems using common benchtop laboratory equipment and measurement times are on the order of 10 hours. The experiments are straightforward to set up and a typical analysis (eg van 't Hoff plot), can be performed entirely using standard spreadsheet software (*see Supplementary Information*). We believe that the TE mapping method will prove generally useful to the supramolecular chemistry community.

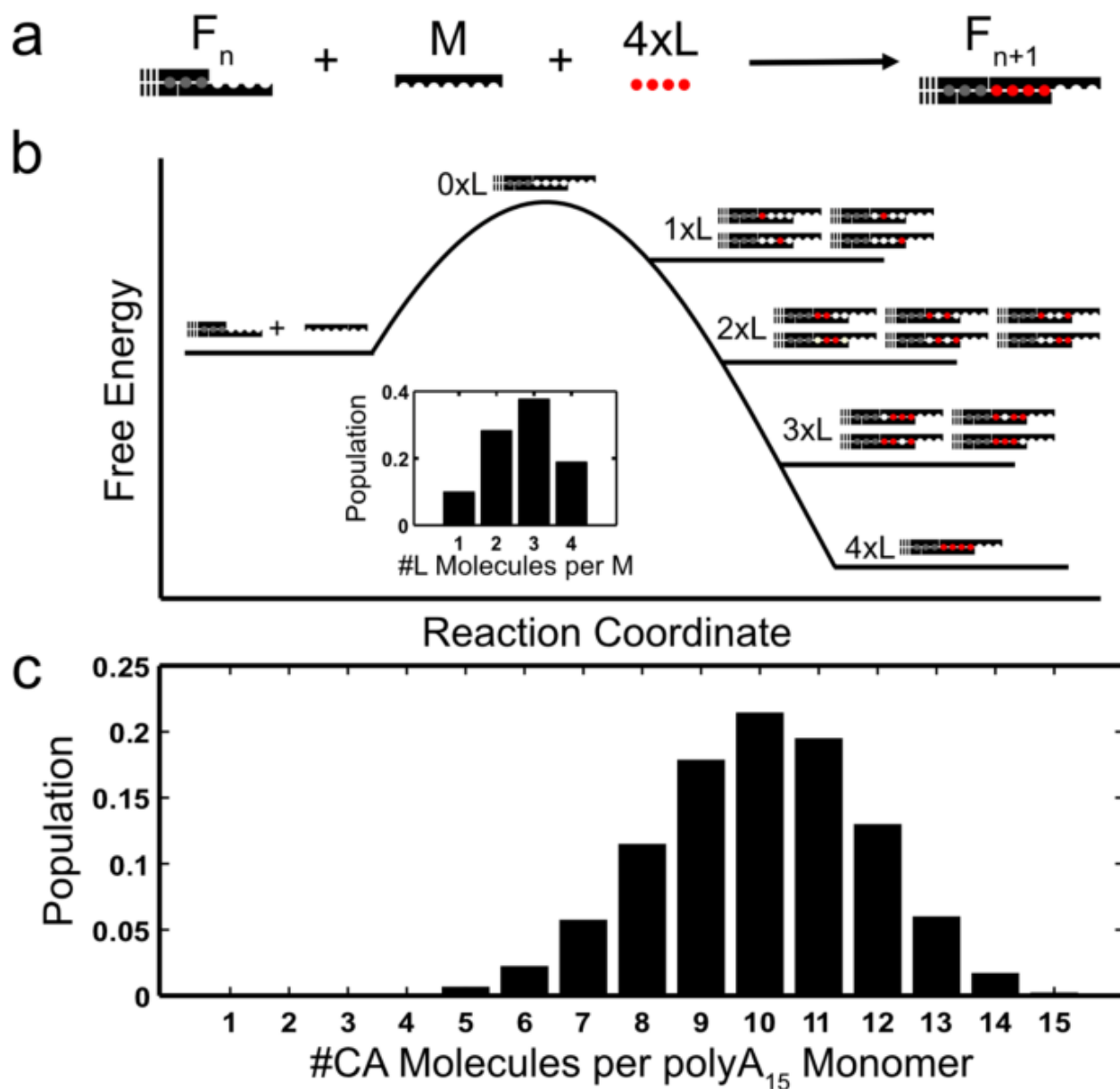


Figure 6: **a)** A simple constant stoichiometry assembly model where the end of a growing fiber (F_n) assembles with one monomer M and 4 ligand molecules L to create a fiber of length $n+1$ (F_{n+1}). **b)** A free energy diagram of a variable stoichiometry assembly model where the end of a growing fiber (F_n) can assemble with a monomer M and any number of ligand molecules L up to a maximum of 4, the insert represents the populations of each stoichiometry. **c)** The populations of each stoichiometry for the self assembly of polyA-CA fibers at 25°C with a concentration of 12.5mM CA.

Online Methods

Materials

Cyanuric acid (CA), tris(hydroxymethyl)aminomethane (Tris), magnesium chloride hexahydrate ($\text{MgCl}_2 \cdot 6 \text{H}_2\text{O}$), sodium chloride (NaCl), glacial acetic acid and urea were used as purchased from Sigma-Aldrich. Boric acid was obtained from Fisher Scientific and used as supplied. Acrylamide/bis-acrylamide (40% 19:1) solution, ammonium persulfate and tetramethylethylenediamine (TEMED) were used as purchased from BioShop Canada Inc.

1xTBE (Tris-boric acid-EDTA) buffer was composed of 45 mM Tris, 45 mM boric acid and 2 mM EDTA at pH 8.3. 1xAcMg buffer was composed of 40 mM acetic acid, 7.6 mM $\text{MgCl}_2 \cdot 6 \text{H}_2\text{O}$, with pH adjusted to 4.5. Buffers and samples were prepared with Milli-Q water.

d(A₁₅) oligonucleotides were synthesized on a Mermade-12 synthesizer, purified by denaturing polyacrylamide gel electrophoresis (PAGE 20%, 1xTBE running buffer, 8 M urea) and desalted with Gel-Pak desalting columns from Glen Research. Purity of the strand was confirmed by HRMS (Calculated mass: 4635.18; Observed mass: 4634.28).

Stock solutions of 20 mM CA were prepared by dissolution in 100 mL of Milli-Q water in a volumetric flask and adjusted with acetic acid to pH 4.5. To properly dissolve and degas the solutions, they were heated at 65 °C and sonicated, then cooled down to room temperature before being used.

Samples of 100 µL of polyA₁₅ (25 µM) and CA (7.5, 10.0, 12.5 and 15.0 mM) in pH 4.5 $\text{Mg}(\text{OAc})_2$ buffer (7.6 mM) were made in quadruplicates. A thin layer (~30 µL) of silicon oil was applied on top to prevent evaporation during experiments.

Instrumentation

UV-Vis absorbance-based quantification of d(A₁₅) was performed on a Nanodrop Lite spectrophotometer from Thermo Scientific. DNA purification by PAGE was carried out on a 20 x 20 cm vertical acrylamide Hoefer 600 electrophoresis unit.

UV-Vis absorbance studies were performed using a 1.0 mm quartz cuvette and monitored at 260 nm on an Agilent Cary 300 Series UV-Vis Spectrophotometer equipped with a Peltier temperature controller and water recirculator. A variable temperature range which started from 50-40 °C and went down to 10-4 °C was scanned at a rate of 0.5 °C/min and with a max equilibration time of 30 min. Argon gas and drierite were used to dry the chamber at temperatures below 10 °C.

Data availability

The MATLAB code for performing the analyses herein is available from the corresponding author upon request.

Code availability

The data for performing the analyses herein are available from the corresponding author upon request.

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Author Contributions

H.S. and A.M. conceived the project. C.L.B. acquired all experimental data. C.D.H. performed simulations, data analysis and interpreted the results with A.M. A.M. and H.S. supervised the projects. C.D.H. and A.M. wrote the paper.

Competing Interests

The Authors declare no competing interests.