A new entropy model for RNA: part III. Is the folding free energy landscape of RNA funnel shaped?

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Abstract

The concept of a free energy (FE) landscape, in which the conformations of a polymer progressively take on the structure of the native state while spiraling down a FE surface that resembles the shape of a funnel, has long been viewed as the reason why a complex protein structure forms so rapidly compared to the number of conformations available to it. On the other hand, this landscape picture is less clear with RNA due to the multiplicity of conformations and the uncertainties in the current thermodynamics. It is therefore sometimes proposed that within the ensemble of suboptimal states of the RNA molecule, the vast majority of those states all closely resemble the native state and therefore simply overwhelm the few states that represent the global minimum FE. However, calculations of the free energy of observed structures often suggest that the most frequently observed cluster of structures are far from the minimum FE, particularly in the case of long sequences. If so, then such a FE surface is unlikely to be funnel shaped. We have been developing a version of vsfold that can evaluate the suboptimal structures of the FE surface (through a modified version called vs_subopt). Here we show that the ensemble of suboptimal structures for a number of known RNA structures can actually be both close to the minimum FE and also be the dominant observed structure when a proper Kuhn length is selected. Two state aptamers known as riboswitches can show neighboring FE states in the suboptimal structures that match the observed structures and their relative difference in FE is well within the range of the binding free energy of the metabolite. For the riboswitches and other short RNA sequences (less than 250 nt), the flow of the suboptimal structures (including pseudoknots) tended to resemble a rock rolling down a hill along the reaction coordinate axis. An important insight yielded by the cross-linking entropy (CLE) model is that the global entropy limits the size of domains. Hence, based on the CLE model, Levinthal’s paradox is overcome by the funnel shape in the FE, by a reduction in the number of degrees of freedom due to Kuhn length, and by limits on the size of the domains that can form. These concepts are also applicable to calculating transition rates between different suboptimal structures.

Introduction

In Part I of this series, we showed that the global aspects of the crosslinking entropy (CLE) model satisfy the fundamental requirements of a heat engine in reversible processes. In Part II, we showed the role of the Kuhn length of a polymer in influencing the local entropy in structure predictions, the local contribution of the CLE. Also, in a recent report,1 we showed that the CLE model is able to unify the lattice model, the Gaussian polymer chain model and the contact order model, under the same framework. Likewise, in previous work, we have also shown how the application program we are continuing to develop (vsfold5) is able to successfully predict RNA secondary structure16 and pseudoknots,17 at least when a good choice of Kuhn length is employed (Part II, Section 7). What has not been explored so far is the folding free energy (FE) landscape itself when using the CLE model. Is the landscape funnel shaped? In other words, is the optimal folded structure the final point of a process which is characterized by a series of suboptimal structures that are similar to the optimal structure and gradually descending through a funnel in the free energy landscape toward the global minimum. A lot of work has been devoted to studying the folding landscape of biopolymers, particularly for proteins,8-17 where some have attempted to construct a FE model that includes the Jacobson-Stockmayer (JS) model discussed in Part I.18 Some work has also been done for RNA,19-41 which often defers to the standard JS-model.20-23 However, the RNA problem has been less encouraging with the JS-model because known and observed RNA structures tend to be scattered within a mountain of suboptimal structures like a needle in a haystack, and there is no easily discernible pattern in the thermodynamic distribution of key suboptimal structures either.42With some RNA structures having passed through 3.5 billion years of natural selection to remove inefficiencies and instabilities in the RNA structures, relegating these known and observed RNA structures to positions deep within a list of suboptimal structures is rather incongruous. It may be possible that long RNA sequences require chaperones to fold correctly; however, the majority of shorter RNA sequences (like tRNA and riboswitches) should have solved this problem through natural selection. At least, it would be far more satisfying if a funnel shaped landscape could be shown for some representative
cases or the exceptions explained.

It has long been thought that the folding landscape for both RNA and protein folding is funnel shaped. Here we added the functionality to calculate suboptimal structures into the vsfold program we have been developing (through a modified version called vs_subopt). Using the CLE model, we show that the folding landscape of RNA is indeed funnel shaped, at least for the representative and important RNA molecules we have tested.

It is also well known that some RNA molecules are used by organisms as two state switches that bind metabolites in order to signal other transcription processes to stop or start, depending on the particular system and its purpose. Assuming the FE landscape is funnel shaped, it is our hypothesis that both states should be relatively stable, where trapping of the metabolite should require only modest changes in the total FE, since binding affinities are likely to be modest in magnitude given the CLE model, we show that the folding landscape of RNA is indeed funnel shaped, at least for the representative and important RNA molecules we have tested.

As a result, we show that domain sizes in RNA structures can be estimated based on the average base composition of the sequences. This is the only FE based model that predicts this based on a theoretical framework. The same framework also permits estimating the folding times between different suboptimal structures.

This is a theoretical model. This report is limited to a few hand-picked example structures not because these structures are the only cases that succeed; rather these examples represent what is generally the case and what is generally expected based on theoretical grounds.

Materials and Methods

Determination of suboptimal structures using vs_subopt

The general design and procedure used to calculate secondary structure and pseudoknots is explained in detail in Dawson et al. Therefore, we will focus on the method for evaluating suboptimal structures. Let $i$ and $j$ represent the identity of two bases that are numbered in a sequence such that $i < j$ and each successive base is counted sequentially from the 5' to the 3' end of the sequence. As vsfold calculates the secondary structure and pseudoknot structure, the information about the best (optimal) structure is retained at each position $i$ and $j$, which we denote by the ordered pair $(i, j)$. This residual structural information is used to help find the suboptimal structures.

The program searches in terms of junction points or levels in the hierarchy of secondary/pseudoknot structure. In Figure 1, a complex RNA structure including a pseudoknot is shown. Stems are defined in terms of a tail and a head of the structure, where the tail is the 5' and 3' most positions of the RNA sequence, Figure 1 (denoted in several places in Figure 1). Hence, if $(i, j)$ is the closing bp at the head of stem and $(i, j)$ at the tail, then $j - i < j - i$.

The zeroth level defines the arrangement of domains of secondary structure (or pseudoknots) and corresponds to the structures that extend off of level 0 in Figure 1. These are indicated by the labels domain 1, 2 (the pseudoknot) and 3 in Figure 1. In the Figure, the closing stem of domain 1 has the label [0] and we express the boundaries of this domain in this work by the notation $(i, j)[0]$, the tail of the closing stem for [0]. Level 1 of stem [0] begins at $(i, j)[0]$, in the region between $i < i, j < j$. For stem [0], the next level has stems labeled [0,0], [0,1] and [0,2], where the first index identifies the stem at level zero and the next index identifies the stem at level 1. Hence, [0,1,0,2] identifies a stem in domain 1 at level 3, connected to stem 0 of level 2, connected to stem 1 of level 1, connected to stem 0 of level 0. Likewise, the label [1] indicates the pseudoknot (PK), where its boundaries are $(i, j)[1]$. In the case of the PK, in the vsfold5 program, pointers are used to register which stem is assigned [1,0] and which [1,1] and from there, the hierarchy is the same. A level 1 search will have the label of the form $(i, j)[1]$, level 2 $(i, j)[2]$ and level 3 $(i, j)[3]$, and so on.

At each level, the region enclosed by the corresponding $i < i, j < j$ is scanned for alternative structures. The resulting suboptimal structures are then grafted onto the calculated stem. For intermediate levels, the best zeroth level is selected first, then the best first level and so on. If

![Figure 1. A schematic of the concept of levels in secondary structure and levels of the search. Level 0 represents the base domain of the structure. These are labeled domain 1, 2 and 3. The corresponding stems that close the domains are labeled (0), (2). All higher levels represent subdomains. These stems are expressed with progressively more indices, depending on the level. For example, stem (0,2,0) is at level 2 and (1,0) is at level 1. All searches except level zero begin from the head of the stem, level zero scans directly from the 5' to 3' ends of the sequence and begins with the optimal domain structure followed by successively suboptimal structures of the 5' to 3' sequence.](image-url)
the list is exhausted, then the program switches to next best structure from level 0, and so on. It is also possible to specify constraints in order to search a particular region of a specified structure as long as the search region is defined in terms of an actual stem.

In the examples of stems, some stems are shown with small interior loops (I-loops) that are ignored whereas other larger I-loops are treated as new levels. Short I-loops are often considered part of a stem in vsfold5. The precise rules for how vsfold defines these composite stems are explained in the distributed manual that comes with the vsfold5 distribution. In general, if the I-loop is short like a 1x1 or 2x2 I-loop, it is skipped in the evaluation of suboptimal structure because vsfold treats it as a bona fide stem.

The default setting of vs_subopt assumes the zeroth level as the desired suboptimal structures. This provides composite structures that are optimal for the sequence with a domain located between base i and base j. A fixed number of suboptimal structures are searched for until either no more structures are found, or the total number has been found within the default energy range (10.0 kcal/mol). The user can override the default settings by a variety of command line flags; -so_level n selects the level of interest (n=0,1,2,...), -so_max N adjusts the number of suboptimal structures to be scanned (default, N=20) and -FEspan E sets the energy range (from the optimal structure) to be scanned (default, 10 kcal/mol).

**Estimation of activation barriers and transition times**

In general, a chemical reaction involving bond formation or Van der Waals interactions happen on a time scan of ps or shorter. On the other hand, folding of RNA structures involves time scales greater than μs, typically ms and sometimes minutes. This is because different parts of a RNA sequence must diffuse together against the force of the chain entropy to form a specific base pairing interaction (explained in the model in Part 1, Section 5 and also in Dawson et al. and Cheng et al.)

The formation of contiguous bps into a stem, which is known as the stacking process, involves local diffusion processes and chemical interactions. From molecular dynamics simulations, one can observe a single bp that frays from the stem has a lifetime of approximately 5 to 10 ps. The process involves both the mechanical motion of the bp due to the local chain motions in solvent combined with chemical reaction when the stack is actually formed, a process of Van der Waals interactions happen on a time scan of ps or shorter. On the other time scale in the rate will usually be the diffusion of different parts of the RNA chain together against the force of the chain entropy. When the stack is actually formed, a process of Van der Waals interactions happen on a time scan of ps or shorter.

Hence the RNA chain folding interactions often happen on a much longer time scale than the stacking interactions but both are largely diffusion limited.

It follows that in the binding of base pair (bp) i and j, the dominant time scale in the rate will usually be the diffusion of different parts of the RNA chain together and local diffusion of the bases into stacks. For a diffusion rate constant and chemical reaction rate constant, the total rate (k) of a diffusion limited formation of chemical species (i.e., an RNA bp) is approximated by the following expression

\[ k = \frac{1}{k_{diff}} + \frac{1}{k_{chem}} \approx \frac{1}{k_{diff}} \]  

where \( k_{chem} \) should be seen as a combined local diffusion of water during the stacking process, rearrangement of the RNA chain, and chemical reaction rate.

In the concept of the CLE model, the statistical mechanics entity is the stem. Stems in the structures should independently disassemble and reassemble as blocks of stems with a global entropy weight. Evidence suggesting this can be found in the force extension experiments where a stem unzips or re-zips as a unit and in differential melting experiments where one can assign the melting transitions to particular stems. The global entropy places a strong weight on the folding time of a particular block. Because these stems separate independently, in this scenario, the maximum activation barrier (and the rate determining step), is dependent on the stem that exhibits the maximum entropy loss in the process of joining the two different parts of the single-strand RNA chain together. Therefore, in a model where the unit is a stem, estimates of the transition energies should be a function of the Kuhn length.

The rate can be estimated by viewing the process as the sum of the mutually dependent folding times of the individual stems \( r_i \)

\[ r_i = r_i + r_i + ... r_i \text{, with } r_i = \frac{1}{k_{diff} + \frac{1}{k_{chem}}} \]  

where \( n \) is the total number of stems, the dominant process is the global CLE, the reference is the time it takes for the structure to fold from a denatured state to the native stem state and the longest folding time will be a function of the maximum length of the subsequence separating bp \( (i,j) \); \( \max(N_j, j = i + 1) \).

The folding rates are sometimes handled by considering all the RNA folding pathways using some approach based on the Morgan-Higgs algorithm. Here, we are more interested in the overall time scale of the stem formation (formation of contiguous bps) in transforming between two different structures. For example, two general structures can be generated from the following toy example:

**Structures A and B are mutually exclusive stems-loops that share no common base pairs (bps). One stem has to unfold for the other stem to form. The activation barrier is then found by computing the change in free energy required to unpair stem A (deletion) and to form stem B (insertion); i.e., the minimum number of editing steps that permits transition between two different structural states. It should contain the total path because the transitions between structure A and B are in thermodynamic equilibrium and the one stem must unfold before the other can form. Therefore, to model the above simple process, we imagine the folding time to be the sum of the individual folding times for structures A and B; i.e., the sum of the two processes (\( r_{A\rightarrow B} = r_{A\rightarrow \text{unf}} + r_{B}\)). However, in general, we should expect some stems to be common to both structure A and structure B. Following Eq (2), let \( A^*B \) express those stems that are common to both structure A and structure B. Then the folding time for \( A^*B \) is expressed as the sum of the independently folded structures A and B minus the time for folding \( A^*B \).

\[ r_{A^*B} = \frac{1}{2}(r_{A\rightarrow \text{unf}} + r_{B}) - \frac{1}{2}(r_{A} + r_{B}) = \frac{1}{2}\left(\sum_{i=1}^{n} r_{i} + \sum_{j=1}^{m} r_{j}\right) - \frac{m+n}{2}\sum_{i=1}^{n} r_{i} (A \cap B) \]

**Therefore, to obtain the folding time, we first need to find the minimum number of stems that must change; i.e., the minimum number of stem-editing steps. This is done by comparing the similarity of the two suboptimal structures. We then assume a sequential set of steps in which the structure becomes partially unfolded in order to transition to the alternative structure. In this case, the folding rate is dependent on the total time for each of the stems (or parts of a stem) to come apart.**

\[ k_{chem} = \frac{\Delta G_{chem}}{k_{B}} \exp \left( \frac{\Delta G_{chem}}{k_{B}} \right) \]  

where \( \Delta G_{chem} \) is the free energy change for the chemical reaction.
where \( k_B \) is the Boltzmann constant, \( h \) is the Planck constant, \( T \) is the temperature, \( \omega_{\text{trans}} \) is the transmission coefficient (related to activities) and \( \Delta G_{\text{trans}} \) is the FE (including the CLE) to remove a stem. The transmission coefficient here is likely to be a function of various unknown activity coefficients and associated concentration.\(^{34,47}\) For a polymer in solution, \( \omega_{\text{trans}} \) is probably at least a few orders of magnitude smaller than 1. For RNA, \( \omega_{\text{trans}} \) has been lumped together with \( h k_B T \) and called the pre-exponential factor \( \tau_{\text{chem}} = \omega_{\text{trans}} h k_B T \).\(^{34,48}\) Present at present, there are no actual values for \( \omega_{\text{trans}} \). However, given the maximum folding rate of proteins (and perhaps RNA)\(^{35}\) is on the order of microseconds,\(^{32}\) this can be tuned for a single stem. For stems, because \( \Delta G_{\text{trans}} \) is much larger than the bp FE, the correction is on the order of \( \omega_{\text{trans}} = 10^{-2} \) to \( 10^{-3} \) compared to \( 10^{-4} \) for a single bp.

Since the transition time is found in the inverse rate, we write

\[
\frac{1}{k_{\text{chem}}} = \frac{1}{k} + \frac{1}{k_B} + \frac{1}{k_B} \sum_{n=1}^{\infty} \frac{1}{n!} \frac{\Delta G_{\text{trans}}}{k_B T} \exp \left( \frac{\Delta G_{\text{trans}}}{k_B T} \right)
\]

(5)

In general, this would require a detailed evaluation of all the stems and determination of the changes in FE for each case. However, if the Kuhn length is essentially constant for a region of the sequence, we can make a simpler estimate. In part II, we showed that the Kuhn length and the stem length should be essentially the same value. In such a case, all the stems are assumed to be of similar length and the total number of stems that change can be estimated from the total number of base pairs that must be removed divided by the Kuhn length,

\[
\Delta N_{\text{chem}} = \frac{\Delta N_{\text{bp}}}{\xi}
\]

(6)

where \( \Delta N_{\text{bp}} \) is the minimum number of editing operations that are required on the base-pairs (bp) to achieve the transition and \( \xi \) is the Kuhn length. Likewise, the average energy of each stem is just the sum of the FE from all the bps that are different, or the FE of stems that change, divided by \( \Delta N_{\text{chem}} \).

\[
\langle \Delta G_{\text{trans}} \rangle = \frac{\sum_{A \not= B} \Delta G_{\text{trans}} \xi}{\Delta N_{\text{chem}}} = \frac{\xi}{ \Delta N_{\text{chem}} } \sum_{A \not= B} \Delta G_{\text{trans}}
\]

(7)

where \( A \not= B \) specifies those stems in structures A and B that are not shared in common (the complement of \( A \cap B \)). \( \Delta (bp) \) refers to the associated bps in \( A \cap B \) (\( \Delta (bp) = A \cap B \)), and \( \Delta G_{\text{trans}} \) refers to the FE of \( A \cap B \), where both stacking and the global CLE are included.

Hence, the transition requires a larger energy than a single base pair, which is certainly too small and would suggest far too rapid a rate than a stem requires. Likewise, the transition energy is typically much smaller than the total change in energy of all the editing steps, which is often far too large and would require unreasonably long time intervals. It follows that the total time for a transition from structure A to structure B will simply to

\[
\langle \tau \rangle_{A \rightarrow B} = \frac{\Delta N_{\text{chem}}}{\Delta N_{\text{chem}}} \frac{h}{k_B T} \exp \left( \frac{\Delta G_{\text{trans}}}{k_B T} \right) + \Delta N_{\text{chem}} \tau_{\text{chem}}
\]

\[
- \Delta N_{\text{chem}} \frac{h}{k_B T} \exp \left( \frac{\Delta G_{\text{trans}}}{k_B T} \right) \tau_{\text{chem}}
\]

(8)

where \( \tau_{\text{chem}} \) is \( 1/k_{\text{chem}} \) (Eq 1) and \( \omega_{\text{trans}} = 1/\tau_{\text{chem}} \) (Eq 4). Currently, we make the crude estimate that \( \tau_{\text{chem}} \) is about 10 ns (based on estimates in Pocskhe et al. and might be as large as \( 1\mu s \) for a whole stem),\(^{25,26}\) and \( \omega_{\text{trans}} \) is 100, based very roughly on activities.

The rate of the reaction will be the inverse

\[
\langle \tau \rangle_{A \rightarrow B} = \frac{1}{\omega_{\text{trans}}} \frac{h}{k_B T} \exp \left( \frac{\Delta G_{\text{trans}}}{k_B T} \right) + \Delta N_{\text{chem}} \tau_{\text{chem}}
\]

Hence, the transition energy of a single stem changing is

\[
\langle \tau \rangle_{A \rightarrow B} = \frac{h}{\omega_{\text{trans}}} \exp \left( \frac{\Delta G_{\text{trans}}}{k_B T} \right) + \Delta N_{\text{chem}} \tau_{\text{chem}}
\]

(9)

Since base pair energies range from \( -0.5 \) to \( -2.5 \) kcal/mol, for a \( 5 \) nt, \( \langle \Delta G_{\text{trans}} \rangle \) is approximately of the order of \( -2.5 \) to \( -12.5 \) kcal/mol. Using Eq (8), this computes to a transition time ranging between 10 ps with \( \tau_{\text{chem}} = 10 \mu s \) at 310 K. For \( \tau = 10 \) nt, \( \langle \Delta G_{\text{trans}} \rangle \) (including the CLE contribution) is approximately of the order of \( -5 \) to \( -25 \) kcal/mol, which corresponds to a transition time of ranging between 500 ps to 14 h (at 310 K). It seems that 14 h is rate limiting for most biological processes.

The model used here is consistent with the view that the transitions tend to be cooperative in RNA. Co-operativity is well known in the melting of proteins.\(^{32}\) For RNA structures, this is seen in the peaks in differential melting curves where particular stems can be assigned to specific melting temperatures in small RNA molecules like tRNA\(^{38,39}\) and pseudoknots.\(^{40,41}\) Likewise, it can be seen in the force-extension experiments using optical-tweezers, where the stems suddenly unzip or re-zip (refold).\(^{30,34}\) These experiments all suggest that RNA unzips and re-folds by stems, not by bps. This can be understood as a direct consequence of the Kuhn length, which specifies that the bases act collectively as a group.

To estimate transition time between different suboptimal structures, we will use this method of estimation. The main economy of this strategy is i) that we don’t have to calculate all the folding pathways or even the specific stems of a given pair of suboptimal structures, and ii) that we can establish a baseline for the folding times that avoids any systematic issues of any particular algorithm that models the folding pathways. The main deficiencies in this estimation strategy are i) that the CLE model currently only uses one Kuhn length for a whole sequence, where many of the active RNA structures of interest clearly have variable stem lengths and therefore different Kuhn lengths, and ii) that the calculated value is the average stem FE, not the specific FE of any particular stem. The method proposed here, therefore, should be seen as a concept in which an average Kuhn length is applied over the entire sequence. This is only proposed as an estimation technique. Future work on \( \text{vsfold} \) and \( \text{vs_subopt} \) will attempt to address variable Kuhn lengths in the computation of RNA structure and are intended to adhere closer to the precise concept of Eq (3) through (5). Nevertheless, this estimation approach should provide a ball-park approximation of these transition times.

\[\text{Results and Discussion}\]

Figure 2A shows the result of a calculation of suboptimal structures within 10 kcal/mol of the minimum free energy using \( \bar{v} = 5 \) nt and level 0 search for yeast tRNA(Phe) based on the unmodified sequence of tRNA(Phe).\(^{35,52}\) The results are sorted by the predicted free energy (FE) with the minimum free energy shown at the top. Figure 2B groups the folding landscape according to similar folding structural intermediates. The optimal structure (minimum FE) is the familiar cloverleaf pattern of tRNA (Figure 2B, right most structure). The D-stem (Figure 2A, purple) and anticodon stem (Figure 2A, green) are already present in the folding landscape according to similar folding structural intermediates. The optimal structure (minimum FE) is the familiar cloverleaf pattern of tRNA (Figure 2B, right most structure). The D-stem (Figure 2A, purple) and anticodon stem (Figure 2A, green) are already present in the folding landscape according to similar folding structural intermediates. The optimal structure (minimum FE) is the familiar cloverleaf pattern of tRNA (Figure 2B, right most structure). The D-stem (Figure 2A, purple) and anticodon stem (Figure 2A, green) are already present in the folding landscape according to similar folding structural intermediates.
Figure 2. Calculations of the suboptimal structures including pseudoknots for tRNA(phe). A) The list of the predicted suboptimal structures (including the free energies) that are within 10 kcal/mol of the optimal structure listed in the order of the predicted energies. The RNA structures are listed in the Fontana-Schuster tree-notation plus a bracket notation [[ ]] for the pseudoknot notation. The bold colors represent known parts of the RNA structure and are labeled on the optimal tRNA structure: (purple) the D-stem (closing the D-loop), (green) the anticodon stem, (blue) the T-stem (closing the T-loop), and (black) the Acceptor stem. The red stem is known to occur in the 3D structure of tRNA. B) A rough schematic depiction of different secondary structures in the order that they appear in the calculated suboptimal structure list in A), where the right most labeled tRNA structure represents the optimal predicted structure. The structures on the top (red arrows) represent the corresponding structure's position and free energy on the approximate reaction coordinate. The purple and magenta arrows represent structures not directly on the reaction coordinate. The arrows propose some possible points where the improper stem unfolds and the structures refold and join the structures along the reaction coordinate.
numbers (10>7>6>5>2>1>0) corresponding to the order in the list of suboptimal structures in Figure 2A. The bottom side of Figure 2B shows the progression of two alternative pathways: 9>4 and 8>3. The majority of structures on the top side follow a natural progression along a gradient down to the native state and minimum free energy, corresponding to a natural energy-to-structure progression. The exception is structure 10, which has a higher FE because the acceptor stem is considerably shorter than the Kuhn length (≤5 nt) leading to local instability (Part II of this series). The alternative pathways (bottom of Figure 2B) are far fewer and those that appear have several features in common with the native state. Considering the folding times, these alternative pathways most likely connect somewhere with the general progression or become suppressed somewhere within the last 5 kcal/mol of the minimum FE. Whereas vsfold5 works in a 5' to 3' folding, vs_subopt folding looks more akin to a denature/refolding experiment where all parts of the sequence fold and compete with each other. The predicted transition rates from the misfolded structures to a neighboring structure along the path of the funnel (Figure 2B) are all on the order of μs for this RNA; e.g., structures 3 and 4 ending up at structure 2 and structures 8 and 9 ending up at structure 7. This may be a little fast, but even 1000 fold slower rate would still render these transitions on the order of ms. The results of experiments on tRNA(phe) are consistent with the assumption that any intermediates that might form during the folding process do not contribute significantly to slowing down the folding rate. In Figure 2B, four structures are shown below the curve depicting the FE along the rough schematic depiction of the 1D reaction coordinate: two with purple arrows (structures 4 and 9) and two with magenta arrows. The arrows (pointing from positions along the suggested reaction coordinate) are meant to propose possible points where the improper stem unfolds and the structures refold into one of the neighboring structures along the suggested 1D depiction of the reaction coordinate. These structures, which do not closely resemble the native state, may persist for some time as "blind alleys." Hence, the predictions suggest some potential trapped intermediates that may form and slow down the folding process. However, in the last three structures where the acceptor stem forms (upper side, red arrows), the structures all have the same major features as the native state. Moreover, the key structural features of the tRNA are largely conserved throughout the set of structures within 10 kcal/mol of the minimum FE. Melting studies of the tRNA do not show sig-

Figure 3. Calculations of the suboptimal structures including pseudoknots for tRNA(Ala) displayed in Fontana-Schuster notation. The optimal structure is shown in at the top of the list. The bold face markings indicate structures associated with the observed tRNA structure. The pseudoknot is not predicted because there are too many non-Watson-Crick pairs in the structure to make a clear distinction with current thermodynamic parameters. The coloring for the stems is the same as used in Figure 2.

| Structure | Free Energy (kcal/mol) |
|-----------|------------------------|
| 1 | -27.49 |
| 2 | -27.29 |
| 3 | -26.52 |
| 4 | -26.35 |
| 5 | -26.52 |
| 6 | -26.35 |
| 7 | -26.35 |
| 8 | -24.32 |
| 9 | -23.98 |
| 10 | -22.87 |
| 11 | -22.87 |
| 12 | -22.53 |
| 13 | -22.15 |
| 14 | -21.88 |
| 15 | -21.84 |
| 16 | -21.78 |
| 17 | -21.48 |
| 18 | -21.41 |
| 19 | -20.13 |
| 20 | -19.68 |
| 21 | -17.94 |
| 22 | -17.23 |

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significant trapping in intermediate states for these simple RNA molecules, though perhaps there is a very small fraction that was missed. In Figure 3, the tRNA sequence shows two major structural variations, including an incorrect anticodon stem, T-stem and acceptor stem. However, after the cloverleaf structure begins to form, the frequency of these alternative structures is greatly suppressed in favor of the correct anticodon- and T-stem. Therefore, with a good model for the entropy and suitable thermodynamic parameters such as Kuhn length, the structures largely fall into place. This shows that RNA is every bit as capable at finding the native state as proteins.

Moreover, this tendency is not unique to these two specific examples. We previously used *vsfold* to analyze all the tRNA sequences found in the genome of *Thermus thermophilus* HB8, and found that 80% of these structures were predicted with the expected cloverleaf structure. Failure was typically the result of neglecting non-Watson-Crick (non-WC) pairing, for which there are no established thermodynamic parameters at present. Using vs_subopt, given there is nothing particularly unusual about the sequence, we can often obtain the cloverleaf and a strongly reinforced homology resembling the native state all the way through the folding landscape and dominating most of the thermodynamically stable structures.

It is notable that there are only a handful of structures that lie with the 10 kcal/mol range and of the monomers and the entire search space must be searched, it is easy to show using Levinthal's paradox that even a fundamental RNA structure like ribosomal RNA 16S (approximately 1500 nt) would take longer than the lifetime of the universe for a search of all the folding conformations. Yet even given the large reduction (order N/ξ) in the number of degrees of freedom, scaling by N/ξ only changes the length scale. For example, we can simply propose a sequence with a structure that is 2 times longer in length and we will return to the same dilemma. Let p represent the ratio of the number of bps (Nωp) divided by the sequence length (N), p=Nωp/N, where 0≤p≤1/2. Based on Dawson et al., the global entropy can be approximated as:

\[ ΔG_{\text{global}} = pNk_B T \left( γ \ln(\Psi(N))-1 \right) \]

where γ(N) is the dimensionless function in the brackets and \( \Psi = γ/N^2 \) with λ=2 the chain-chain separation distance between the bps. Let ΔHω = −pNk_BT where H the absolute value of the average enthalpy of the base pairs in the sequence and let \( \bar{h}(T) = \bar{h}(k_B T) \) represent the dimensionless form of this average enthalpy. The local entropy correction to the FE for a free strand structure with a fixed Kuhn length (Part II) is

\[ ΔS_ω = \left( γ + 1/2 \right) Nk_B T \left( 1/\xi \right)(D/\xi^3) \]

where \( \xi = 3 \) nt approximates the Kuhn length in the free strand regions. The correction for stems in the structure (from Part II, Section 8, Eq 43) is

\[ ΔG_{\text{local}} = \left( γ + 1/2 \right) Nk_B T \left( f(ξ/ξ_0) - f(ξ/ξ_5) \right)(D/ξ^3) \]

where ξ is nt is the Kuhn length in the free strand regions. Hence, substituting \( ξ_0 = 3 \) and ξ = ξ_5 the local correction is

\[ ΔS_ω = \frac{\left( γ + 1/2 \right) Nk_B T \left( f(ξ/ξ_0) - f(ξ/ξ_5) \right)(D/ξ^3)}{D} \]

where, by inspection, if p=1/2, then Eq (11) becomes \( \bar{g} = f(ξ/ξ_0) \), and if p=0, then \( \bar{g} = f(ξ/ξ_5) \). The approximate FE of a domain of length N is therefore

\[ ΔG = ΔHω + ΔG_ω(N) + ΔG_μ \]

where \( \bar{h} = k_BT \) the conformation space decreases factorially with increasing \( ξ \). This also applies to simple base-pairing combinatorics (i.e., paired or unpaired), where 2^N combinations of bps reduced to 2^N/ξ combinations of stems.

Furthermore, the CLE model also limits the size of the domains that can form. If the number of degrees of freedom were equal to the number of monomers and the entire search space must be searched, it is easy to show using Levinthal's paradox that even a fundamental RNA structure like ribosomal RNA 16S (approximately 1500 nt) would take longer than the lifetime of the universe for a search of all the folding conformations. Yet even given the large reduction (order N/ξ) in the number of degrees of freedom, scaling by N/ξ only changes the length scale. For example, we can simply propose a sequence with a structure

| Base pairing | Average bp weight (C_b) [kcal/mol] | 3.0 | 5.0 | 7.0 | 9.0 |
|-------------|-----------------------------------|-----|-----|-----|-----|
| AU rich     | 1.5                               | 94  | 669 | 6530| 7.33E+04|
| AU rich     | 2.0                               | 374 | 6700| 1.64E+05| 4.64E+06|
| AU rich     | 2.5                               | 1491| 6.71E+04| 4.14E+06| 2.94E+08|
| GC rich     | 3.0                               | 5948| 6.72E+05| 1.04E+05| 1.83E+10|

Table 1. A list of estimates for the maximum domain size of RNA (in units of nucleotides [nt]) given an average base-pair binding free-energy (C_b; estimated from averaging the Turner rules for the stems in a particular sequence) and the Kuhn length (ξ) of the stems in the domain.
average Kuhn length is around 5 bps. Hence, Table I indicates that most RNA has a maximum domain size that can range from about 300 nt to 6000 nt, where a reasonable value is likely around 500 to 1000 nt. When $p \to 0$, Eq (12) vanishes. Around $p=0.25$, the predicted domain size does not increase ($p>0.25$) or decrease ($p<0.25$) dramatically, so there is little gain or loss by changing this quantity around $p=0.25$. This is due to the global FE costs of bp formation. It is also clear from Table I that it might be possible to make very large domains using GC rich sequences. Perhaps a relatively equal amount of ACGU is favored because it maximizes the randomness and therefore the amount of information that can be stored or perhaps a synergy in coupling with different biological processes requires time scales that favor smaller domains than the maximum conceivable.

The Kuhn length tends to drastically reduce the number of degrees of freedom but the global entropy (a function of the Kuhn length) sets limits on the size of the relevant search space. In combination, it becomes possible to estimate folding times that are consistent with the observed biologically relevant folding time-scales.\textsuperscript{17} Levinthal’s paradox is overcome by i) the funnel shaped FE landscape, ii) reduction of the number of degrees of freedom, and iii) the limits on the domain size due to faster growth of the global entropy compared to the base-pair free energy (Turner rules).\textsuperscript{16} The last point is not predicted by any other method.

Riboswitches are an example of a type of RNA that does not fold into a single type of molecule, but must exist in two different states. To have a reasonable likelihood of capturing a metabolite, the molecule likely spends some time in a configuration close to its cognate structure. Further, the riboswitch should be able to release that metabolite when specific conditions change. Hence, both states should be present in the observed two-state system, where the molecule hops back and forth through a FE barrier. In such a case, the folding landscape is likely to have both structures coexisting near the minimum FE with an activation barrier that prevents the structure from spending too much time in the alternative state, but with enough time that a metabolite can find the structure in a desirable configuration (perhaps like many biological systems, a rate of about ms$^{-1}$ to ms$^{-1}$). Therefore, a good test of the CLE model is to see if both states of the riboswitch are close together in the list of suboptimal structures and that the transition times are within a few ms at most.

Figure 4 shows two states of the \textit{Vibrio vulnificus add} Adenine riboswitch.\textsuperscript{66,67} Figure 4A is the optimal structure (#0) and is similar to the structure proposed for the bound metabolite (Adenine). However, the P1 stem is only partial. Figure 4B is the first suboptimal structure (#1) and represents the unbound state where stem P1 is lost,\textsuperscript{68} with $\xi=6$ nt. The two structures have rather similar FEs (\(\Delta G=0.1\) kcal/mol). Figure 4C,D compare the optimal structure (#0) and the second suboptimal structure (#2) for $\xi=7$ nt. For the $\xi=7$ nt case, the full P1 stem is found in the minimum FE structure and lost in the #2 suboptimal structure. Both Figure 4B and D show the same structure and contain the transition repression structure (right hand side of the structure). In Figure 4C,D, the FE difference between the structures #0 and #2 is \(\Delta G=6.0\) kcal/mol, which is a little high, but part of this is because the Kuhn length is too long for the structure on the left hand side of Figure 4A. The CLE model is the only approach that even considers the stiffness of the RNA in the FE calculation, and thus it provides more information on the nature of the RNA.

From Eq (10), the predicted transition time is about 1 ms, with $\tau_{\text{unopt}}=100$ and $\tau_{\text{opt}}=10$ ns for the two different structures. The alternative structures within 2 kcal/mol all tend to have a slower formation time by a factor of 4. With such a rate of fluctuation, diffusion of the metabolite (about 1 ms to ms within the volume of a cell)\textsuperscript{69,71} and binding energy difference (more than 2 kcal/mol) render this a system sufficiently rapid in fluctuation to uptake stray metabolites and slow enough to stall unwanted translation. In the determined tertiary structure, the pseudoknot in Figure 4A was found. At present, it is not known if the pseudoknot in Figure 4C exists.

Figure 5 shows the suboptimal structures of the \textit{Bacillus subtilis xpt} Guanine riboswitch.\textsuperscript{62,72} Figure 5A shows the optimal structure (#0) with $\xi=5$ nt, which resembles the observed bound state. Figure 5B is suboptimal structure (#3) with $\xi=5$ nt, which resembles a major part of the unbound form. With $\xi=5$ nt, the anti-terminator stem and shift between terminator and anti-terminator is not observed. The energy difference between the unbound structure (minimum free energy) and bound metabolite structure is on the order of 3 kcal/mol; within a necessary range of energy that a binding metabolite can successfully utilize. Further, the riboswitch should be able to release that metabolite when a rate of about $10^{-1}$ to $10^{-2}$ s, with the stems that are present. Nevertheless, as also shown in Part II, even a poor choice of Kuhn length can still sometimes yield success in a robust model.

The estimated transition time between the two structures in Figure 6B,C is about 30 ms, which may be rather fast based on current available information.\textsuperscript{18} However, the folding rate is very sensitive to Kuhn length, which is a function of the stem length (as shown in Part II). A 20%
Figure 4. Suboptimal structure results for the Vibrio vulnificus add Adenine riboswitch. A) The optimal structure (#0) for $\xi=6$ nt, where part of P1 is present and the structure resembles most of the features of the unbound structure. B) The first suboptimal structure (#1) for $\xi=6$ nt, where the P1 is removed and the structure resembles the bound form. The purple circled region represents the Shine-Dalgarno (SD) GAA sequence and the orange circled region represents the initiation codon. C) The optimal structure (#0) for $\xi=7$ nt, where P1 is complete and the secondary structure matches the unbound riboswitch. Here, a pseudoknot closes off the SD region but leaves the initiation codon free. D) The same suboptimal structure as in B) with $\xi=7$ nt. E) A list of the suboptimal structures and free energies within 2 kcal/mol of the minimum FE for this riboswitch (with $\xi=6$ nt), where the top most structure is the minimum FE and the FE is sorted in increasing order. The two states of this riboswitch are represented by the first and second suboptimal structure (energy difference 0.10 kcal/mol).
Figure 5. Suboptimal structure results for the Bacillus subtilis xpt Guanine riboswitch. A) Optimal structure (#0) at $\xi=5$ nt where the switch is in the bound form, where the label T is for the terminator stem. B) The second suboptimal structure (#2) at $\xi=5$ nt where the switch approximates the unbound form without the P1 stem. Here, the terminator stem is not removed. C) Optimal structure (#0) at $\xi=8$ nt, which is similar to A) except for the pseudoknot and represents the bound form. D) Suboptimal structure (#11) at $\xi=8$ nt, which resembles most of the features of the unbound form including the anti-terminator stem (AT). E) The suboptimal structures and free energies that are within 10 kcal/mol of the minimum free energy for this riboswitch for $\xi=5$ nt, where the top most FE is the minimum and the FE (right most) increases down the list. P1=paired stem 1, etc., T=terminator and AT=anti-terminator. The difference to remove the P1 stem is represented by the #0 and #2 structures (energy difference 3.24 kcal/mol with $\xi=5$ nt). To completely break up the terminator stem and remove the P1 stem, a transition energy of 11 kcal/mol is needed.
increase in the Kuhn length of the terminator stem would produce a transition time on the order of seconds. Furthermore, the transmission weight is currently unknown and may be much larger than we have estimated.

A total of 10 SAM-ribowitches were tested and similar observations were obtained. The results will be reported elsewhere.

SAM riboswitch structures similar to those shown in Figure 6A can also be found by simply recalculating using a different Kuhn length. Since binding a metabolite is likely to change the stiffness of the structure (the Kuhn length), this means changes in the Kuhn length also should be expected. Figure 7A shows the structure of the yusC riboswitch with the terminator stem (structure #0) and closely resembling the bound structure, and Figure 7(b) represents a neighboring suboptimal structure (#4) that approximately corresponds to the proposed unbound metabolite structure and has the antiterminator stem. In these calculations, the Kuhn length of $z=10$ nt, minimum stem length 3 and the `-Mg` option was necessary to make the precise yusC structure with all stems correctly matched. A further adjustment was the option `-cc_dist 5` that extends the internal-loops of contiguous stems to a maximum of 5×5 (where the default is 4×4 for $z=10$ nt). Figure 7C is the prediction (#0) when the Kuhn length is changed from $z=10$ to $z=8$ nt. The key features in the structure strongly resemble the unbound structure #5 found in Figure 6.

The precise extent that the Kuhn length changes in the presence of a metabolite is not known and similar suboptimal structures can also be found using $z=9$ nt. Changing the Kuhn length has some of the same effects as calculating the suboptimal structures. The merit in checking suboptimal structures with the same Kuhn length is that all other

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**Figure 6.** The results of the yusC SAM riboswitch. A) The first 8 structures predicted by vs_subopt for the yusCBA operon, where structure #1 represents the bound form and structures #5 through #8 represent the unbound form of the riboswitch. B) The bound form of the SAM riboswitch. C) The energetically nearby structure proposed for the unbound form of the SAM riboswitch. P1=paired stem 1, etc., T=terminator and AT=anti-terminator.
parameters are left unchanged. Therefore, it minimizes the systematic issues of estimating the differences in energy between structures using different Kuhn lengths. Nevertheless, comparing Figure 7A and C, it turns out that the free energy favors Figure 7C by roughly 10 kcal/mol. In the absence of the metabolite, the unbound structure should probably be favored.

Figure 7 also demonstrates another aspect of the CLE model; namely, the robustness of the predictions. The structures in Figure 7 were fitted with additional 5' sequence from the yusCBA operon, yet very similar cloverleaf structures are found for the bound state. This shows yet again that vsfold is valuable for doing RNA-biology research because the researcher does not have to manicure sequences to help the model find the domains. Rather, the CLE model is robust enough to find those domains on its own power. For the small expense of recalculating the same sequence using a different Kuhn length, the researcher learns far more information on the domain character and stiffness of a particular RNA sequence under study. A structure prediction model should aid researchers in understanding the physics of structure.

We have shown for representative structures of RNA that a funnel-shaped energy landscape with the optimal structure being the observed structure and the suboptimal structures largely pointing in the direction of the optimal structure was largely achievable using the CLE model. We also observed in several important riboswitches that the free-energy difference between the bound metabolite state and the unbound state of

![Diagram of RNA structures](image)

Figure 7. Results of the yusC SAM riboswitch for different Kuhn lengths. A) The switch is in the bound form (using $\xi=10$ nt and including Magnesium ion interactions), structure #0. B) The switch is in the unbound form of the structure #4. C) Calculation of the yusC SAM riboswitch using $\xi=8$ nt and including Mg$^{2+}$ interactions showing the unbound form of the structure (#0). The difference in energy between A) and C) is approximately 10 kcal/mol in favor of C), the unbound structure. All calculations use the -Mg option and -cc_dist 5 to expand the contiguous stem length.
the structure were well within the range expected for binding a metabolite and that these structures were partitioned between the two states and in close proximity, consistent with a two state mechanism where the neighboring state should be in close proximity in energy. The two states of the riboswitch are predicted to be close enough to one another in free energy that it is relatively easy for the metabolite to bind and stabilize the alternative structure. The CLE method is not only predicting structures, it is predicting reasonable energy differences.

A significant improvement here is that the structures are often found in a clear and straightforward order. Key native state structural elements form early and these structures often persist well before the expected native state structure is reached. Different Kuhn lengths generate different results. However, this gives the user some sense of the particular stiffness of the RNA under study. Such concepts are not even thought about by other approaches currently. In the case studies presented here, the optimal structures are certainly the observed structures in NMR and X-ray analysis and the structures ordered in terms of energy with the native state structures forming early and persisting largely throughout the energy landscape. This contrasts with the arrangement of suboptimal structure using conventional techniques, where the correct structure is often just one of many suboptimal structures bearing no particularly close free energy relationship to the optimal structure. This is often justified on the basis that the dominant structural features occur at a much higher frequency. However, it is generally thought that crystals of these structures are in their optimal structure and therefore at the minimum FE. A crystal composed of suboptimal structures would contain a high degree of disorder, especially if the difference between the observed structure and the structure that optimal structures would contain a high degree of disorder, especially where the correct structure is often just one of many suboptimal structures forming early and persisting largely throughout the energy landscape. This contrasts with the arrangement of suboptimal structure using conventional techniques, where the correct structure is often just one of many suboptimal structures bearing no particularly close free energy relationship to the optimal structure. This is often justified on the basis that the dominant structural features occur at a much higher frequency. However, it is generally thought that crystals of these structures are in their optimal structure and therefore at the minimum FE. A crystal composed of suboptimal structures would contain a high degree of disorder, especially if the difference between the observed structure and the structure that corresponds to the minimum FE exceeds 20 kcal/mol at typical temperatures.8,78 Granted, there might be a lot of states that are close together for the observed structure, but it would make more sense if 3.5 billion years of natural selection had already tuned the FE to have the maximum number of states all clustered around the minimum FE. Moreover, even if the commonly observed structure were so far from equilibrium, we should still observe the true optimal structures under at least some experimental conditions.

The distribution and type of species predicted by the CLE model appears to largely overcome these issues, even with the current severe limitations of using one single Kuhn length per calculation attempt. With the current model, the Kuhn length must be decided by the user. Based on the concepts outlined in Part II, stem lengths and Kuhn lengths are strongly correlated. Hence, the Kuhn length can often be discerned from the nature of the RNA itself, e.g., tRNA has short stems and should have a small Kuhn length (≅5 nt) whereas the SAM riboswitch (for example) tends to have long stems and therefore a longer Kuhn length (≅9-10 nt). We therefore think that further development of the model, particularly in the area of a variable Kuhn length (Part II and Dawson et al.), is likely to yield even more accurate and instructive insights, especially since the Kuhn length could also change during folding. In short, what we have been able to show is an approximation, and, given more flexibility in the parameterization (particularly z), better insights are likely to follow. At this point, the experimental data is far too unclear to address these matters further.

**Conclusions**

In this work, we have shown that the CLE model easily generates a folding landscape that is essentially funnel shaped for small RNA structures. For a good choice of Kuhn length, the observed structure turns out to be that predicted by the CLE model to lie at the bottom of the well. Substructures of the native state seem to be grouped consecutively in free energy in an understandable fashion and components of the native state often persist throughout the majority of the suboptimal structures and, therefore, a large part of the folding process.

We also observed that the free energy of the two different structures in a riboswitch can (at least) be relatively close in energy, where one state is the bound state and the other unbound. In general, riboswitches also typically tend to have structural distributions consistent with a two state system. The activation barrier appears to be accessible on a time scale of ms or less, enough time for a metabolite to diffuse to the location and bind. In the case of these two state systems, the folding would fall into either state and then oscillate between them. For such systems, it would have to take two wells on the funnel. Nevertheless, the structures fall into one or the other well and then hop through an activation barrier to the other well.

The observed tendency of the folding landscape to be funnel shaped is consistent with natural selection where, given sufficient time, the thermodynamics of the most essential biomolecules will surely be tuned to optimize the ensemble of suboptimal states (local minima) for folding efficiency, or use the predictable folding process to regulate this rate by mechanisms like trapping, or tune it to differentiate between a finite set of specific states for switching or recognition.

The CLE model is able to add unique insights into how we overcome Levinthal’s paradox. By limiting domain size, the maximum theoretically possible search space for RNA structure is finite for a given base composition. Natural selection may go even further in that the average base composition can, to some extent, bias the size of this cutoff. This also means that, computationally, there will be some cutoff where the computation can be done in linear time. Though this cutoff may still be somewhat prohibitive at this time, a cutoff would allow the application of parallel calculations at least over maximum-domain-size length scales.

The CLE model is robust, providing the researcher with more information than other available approaches about the stiffness of the RNA and providing a far deeper understanding about the stability of functional domains of RNA structure.

**Software**

A binary version of vs_subopt is available upon request to the corresponding author and upon written consent to the license agreement. Available formats are 64 bit Linux (x86_64), or 32 bit Linux, Window XP, and Mac OSX 10.5 and above.

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