Predicting RNA pseudoknot folding thermodynamics

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Received March 24, 2006; Revised April 10, 2006; Accepted April 18, 2006

ABSTRACT

Based on the experimentally determined atomic coordinates for RNA helices and the self-avoiding walks of the P (phosphate) and C₄ (carbon) atoms in the diamond lattice for the polynucleotide loop conformations, we derive a set of conformational entropy parameters for RNA pseudoknots. Based on the entropy parameters, we develop a folding thermodynamics model that enables us to compute the sequence-specific RNA pseudoknot folding free energy landscape and thermodynamics. The model is validated through extensive experimental tests both for the native structures and for the folding thermodynamics. The model predicts strong sequence-dependent helix-loop competitions in the pseudoknot stability and the resultant conformational switches between different hairpin and pseudoknot structures. For instance, for the pseudoknot domain of human telomerase RNA, a native-like and a misfolded hairpin intermediates are found to coexist on the (equilibrium) folding pathways, and the interplay between the stabilities of these intermediates causes the conformational switch that may underlie a human telomerase disease.

INTRODUCTION

RNA pseudoknot structure is defined as a structure with base pairing between a loop and other regions of the RNA. The simplest RNA pseudoknot is the H-type pseudoknot, where a hairpin loop base pair with a single stranded region outside the hairpin. A simple pseudoknot is composed of two helix stems and two loops that span across the helix stems. RNA pseudoknots play an indispensable role in the structures and functions of many RNAs (1–13). For example, in the translation of many viruses, the downstream pseudoknots play a crucial role to promote the frameshifting (14–20). The biological functions of a pseudoknot are directly related to the folding stability and the conformational changes. In an RNA pseudoknot, the stability and folding thermodynamics are largely determined by the interplay between the loops and the helical stems. For example, pseudoknot folding can be cooperative or noncooperative, which involves several intermediates in the folding process. Some of the folding intermediates may be native-like, or misfolded, which cannot be formed from the native helix stems (21-23). To predict how RNA pseudoknot folds, including the formation of all the possible the native-like and misfolded intermediates and the folding stability and cooperativity, it is essential to have an accurate model that can predict not only the native state, but also all the possible native-like and misfolded states. The availability of such a model would be useful to extract the information about the conformational changes and the folding energetics from the experimentally measured thermal melting curves.

To predict the folding thermodynamics requires (i) the energy parameters for the evaluation of the energy for a given conformation and (ii) the chain entropy. While the energy of a pseudoknot can be obtained from the nearest neighbor interaction model, which gives the energy (entropy) as the sum of the energy (entropy) parameter for each base stack in the helices, the chain entropy calculation requires a model. The evaluation of the chain entropy has become one of the bottlenecks for modeling RNA pseudoknot folding.

For a pseudoknot, the two loops span across the deep and the shallow grooves of the helix stems, respectively (See Figure 1a). The (narrow) deep and the (wide) shallow grooves correspond to the major and the minor grooves of the helix stems, respectively. Therefore, the loop conformations are strongly dependent on the helix that the loop spans across. The stem–loop correlation, such as the volume exclusion makes the loop entropy calculation very complex, because the loop and helix conformations must be treated together when evaluating the loop entropy. As a result of the stem– loop correlation, the conformational entropy of pseudoknots are nonadditive, i.e. the pseudoknot entropy cannot be calculated as the additive sum of the entropies of the helix and the (helix-free) loops.

Despite the extensive experimental studies on RNA pseudoknot folding thermodynamics (21–29), our ability to quantitatively predict RNA pseudoknot structure and folding

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Figure 1. (a) A schematic diagram for a simple H-type pseudoknot. Loops L_1 and L_2 span the deep narrow (major) and the shallow wide (minor) grooves, respectively. (b) The secondary structure for the gene 32 mRNA pseudoknot of bacteriophage T2 and (c) the corresponding atomic structure for P and C_4 atoms in the virtual bond representation. The atomic coordinates data are from the NMR structure (PDB ID: 2TPK). (d) The virtual bond representation for the conformation of a nucleotide strand.

stability is very limited (30–34). This is mainly due to lacking of the thermodynamic parameters, especially the chain entropy parameters. The dynamic programming described in Ref. (30) provides an efficient computational algorithm for the structure prediction for pseudoknotted RNAs. Based on ad hoc assumptions, Gultyaev et al. (35) compiled a table for the loop entropies for different loop length and helix length. Later, using the Gaussian chain approximation, Aalberts et al. (36) developed a model to estimate the loop entropies. The model is based on the polymer physics and neglects the atomic details of the loop conformations and the excluded volume interactions. Based on the model, Aalberts et al. performed an analysis for the asymmetry in the pseudoknot structure. Recently, based on the lattice models for chain conformations, Lucas et al. (37) and Kopeikin et al. (38,39) developed theories for pseudoknots and for simple RNA tertiary folds, respectively. These lattice-based physical models can rigorously treat the excluded volume effects but cannot treat atomic details and are thus unable to treat realistic RNA conformations.

In the present study, we present a virtual bond model for RNA pseudoknot conformations and conformational entropy (40–42). The model can account for the atomic details of the conformations. Specifically, we model the helix stems using the NMR measured coordinates (43) and model the loop conformations across the deep and shallow grooves as self-avoiding walks of the nucleotide virtual bonds in a diamond lattice. The atomic coordinates of the helices and the lattice representation for the loops are matched at the loophelix junction, where steric viability is accounted for. When computing the conformational entropy, we explicitly account for the volume exclusion between different nucleotides. From the statistical mechanical model, we compute the free energy landscapes and the base pairing probability at different temperatures, from which the native and intermediate structures, the equilibrium folding pathways, the folding cooperativity and the stabilities can be obtained. RNA pseudoknots can be classified into several different types. In this study, we treat not only the simple H-type pseudoknots, but also the more complex pseudoknots, e.g. the TYMV and TMV pseudoknot structures.

THEORY AND MODEL

Structural model for pseudoknot

A simple (H-type) pseudoknot consists of two stems (S_1 and S_2 in Figure 1a) and two loops (L_1 and L_2 in Figure 1a). In Figures 1b and c, we show the secondary structure and the corresponding positions of the P and the C_4 atoms for the gene 32 mRNA pseudoknot of bacteriophage T2 (43). As shown in the Figure, stems S_1 and S_2 are linked by loops L_1 and L_2 , which span across the deep and shallow grooves, respectively.

The loop conformations are dependent on the helix stems through the chain connectivity and the stem–loop volume exclusion. As a result, the pseudoknot loop entropy is dependent on multiple parameters: the loop entropy is determined not only by the loop length, but also by the helix stem length and structure. Due to the enormously large parameter space (different possibilities for the helix stem length and the loop length), it is practically impossible to rely only on experiments to obtain exhaustively all the possible pseudoknot loop entropies. We need a computational model to calculate the pseudoknot loop entropy. Moreover, to extract the loop entropy parameter from the experiments also requires a model. In this work, we develop a statistical mechanical model to compute the pseudoknot entropy, from which we can obtain the pseudoknot folding free energy landscapes, the folding stability and the conformational changes.

We use the virtual bond model to describe the Hpseudoknot conformations. Since the C-O torsions in the nucleotide backbone have high propensity to be in the *trans* (*t*) rotational isomeric state, the P-O₅-C₅-C₄ bonds and the C₄-C₃-O₃-P bonds in a nucleotide backbone can be treated as planar (40). This makes it possible to describe the nucleotide backbone conformations through two effective bonds: the P-C₄ and the C₄-P bonds. The $P^{(i)}-C_4^{(i)}$ (i = 1, 2and 3) and the $C_4^{(i)}-P^{(i+1)}$ (i = 1 and 2) in the 5' \rightarrow 3' direction are called virtual bonds (see Figure 1d). In our model, we use the experimentally determined atomic coordinates to model the virtual bonds of the helix stems. For the loop region, of which the virtual bonds are more flexible, we use selfavoiding random walks in a diamond lattice to model the conformations.

Helix. For the helix stems, we use the P and C₄ atomic coordinates in the NMR structure of gene 32 mRNA pseudoknot of bacteriophage T2; see Figure 1c. In gene 32 mRNA pseudoknot, the lengths of stems S_1 and S_2 are respectively 5 and 7 bp. For helix stems of other lengths, we use the gene 32 mRNA pseudoknot helix as a template to generate the P and C₄ atomic coordinates through the virtual bond torsional angles (η , θ) and the bond angles (β_P , β_C) (42); see Figure 1d. The virtual bond torsional angles (η , θ) in the helix are (170°, 210°) (41) and the bond angles (β_P , β_C) are (105 ± 5°, 95 ± 5°) for a rigid A-RNA helix (44).

Loop. As shown in Figures 1b and c, loops $L_1(A_8)$ and $L_2(U_{21} \rightarrow G_{27})$ span across the deep narrow and shallow

wide grooves, respectively. The two loops have quite different spatial configurations. For example, the loop end-end distance, defined as the distance between the C_4 atom at the helix-loop junction and the C_4 atom on the other end of the stem, shows different stem length-dependence for the two loops. Shown in Figure 2a are the results from our virtual bond model for the end-end distances, denoted as D_{L_1} and D_{L_2} for loops L_1 and L_2 , respectively. The two loops show very different behaviors. As the helix stem length is increased, D_{L_2} increases monotonically while D_{L_1} decreases until stem length $S_2 = 6$ then increases. Our findings here are consistent with the experimental observation (45).

The above intriguing differences in the stem lengthdependence of D_{L_1} and D_{L_2} are due to the different geometries of the narrow and wide grooves. For the deep narrow groove, the minimum distance between the two paired helical strands is between two non-pairing nucleotides separated by a few base pairs apart instead of between two base paired nucleotides. So the minimum end-end distance for L_1 occurs when the helix stem S_2 has a length of a few base pairs. Moreover, for narrow deep grooves, the minimum end-end distance is small, so L_1 can be as short as a single nucleotide, as shown in Figures 1b and c. On the other hand, for the shallow wide groove, the minimum inter-strand distance occurs between two pairing nucleotides and hence the end-end distance of L_2 increases monotonically with the helix stem S_1 . In addition, the minimum end-end distance is large, so a single nucleotide loop cannot form a viable structure for L_2 .

In our model, we use diamond lattice to configure the loop conformations. This is because the three torsional angles of the diamond lattice bonds (g^+, t, g^-) represent the typical rotational isomeric states t and g^{\pm} of a polymer and thus the diamond lattice conformations may well represent the conformational ensemble of a realistic loop. We model loop conformations as self-avoiding walks of the virtual bonds (i.e. the P and the C₄ atoms) in the diamond lattice. The bond length of the diamond lattice is equal to the length of a virtual bond 3.9 Å. Furthermore, in order to connect the loop to the helix, we map the P and C₄ coordinates of the helices onto the



Figure 2. (a) The C_4 - C_4 end–end distance for the loops for different helix stem lengths. Loops L_1 and L_2 span across the deep and the shallow grooves of the helix stems, respectively. We obtain the C_4 coordinates for short stems using the NMR determined values for the 32 mRNA pseudoknot of bacteriophage T2. For longer stems, we obtain the end–end distance from the helix coordinates generated from the virtual bond model. (b) The entropies for loops across the deep and the shallow grooves. The entropy L_2 , the loop across the shallow groove, shows a non-monotonic loop length-dependence.

closest diamond lattice sites. Such coarse-grained approach causes a root-mean-square deviation (RMSD) of about 2.2 Å for the helix.

Pseudoknot entropy

At the center of the statistical thermodynamics is the partition function Q, defined as the sum over all the possible structures:

$$Q = \sum_{s} \Omega_{s} e^{-E_{s}/k_{\rm B}T}$$
 1

where *s* denotes a structure. Here a structure is defined by the helices. So a given structure would have fixed loop lengths but can have different loop conformations, which cause different pseudoknot conformations. Ω_s denotes the number of pseudoknot conformations accessible to the structure s, and E_s is the energy (enthalpy) of s. The sum \sum_s is for all the possible structures that contain RNA secondary structures and pseudoknotted structures. For a given pseudoknot structure s, E_s can be evaluated from the nearest neighbor model using the experimentally measured enthalpy parameters for the base stacks, while Ω_s can be obtained from the computational model developed here. Given the fixed coordinates of the helix stems, the entropy of the pseudoknot is determined by the loops. So the main focus for Ω_s calculation is to compute the loop entropies in the presence of the fixed helix stems in the structure.

We decompose a pseudoknot into two subunits, each consisting of a stem and the corresponding loop: $S_1 + L_2$ and $S_2 + L_1$. Due to the loop-helix volume exclusion, the loop entropies are dependent on the length of the corresponding helix stems (namely, stem S_1 for loop L_2 , and stem S_2 for loop L_1). The strategy in our theory is to first compute the loop entropies for L_1 and L_2 separately, then combine the subunits to obtain the entropy of the whole pseudoknot.

To ensure the steric compatibility for the connections between the subunits, we configure the P and the C_4 atoms in the loop-helix junction regions onto the closest diamond lattice sites. For example, for the pseudoknot structure shown in Figure 1b, we fit the P and the C_4 atomic coordinates for C7, G9, A20 and U28 onto the diamond lattice. Furthermore, in order to consider the excluded volume between the loops and helices, we also fit stems S_1 and S_2 onto the diamond lattice.

We compute the number of conformations for each subunit $(S_1 + L_2 \text{ and } S_2 + L_1)$ separately through exact computer enumeration for the self-avoiding walks in the diamond lattice. We denote the conformational counts as ω_{L_1} and ω_{L_2} for the two loops, respectively. If we neglect the excluded volume interaction between the two loops, the pseudoknot conformational count can be computed as

$$\Omega_s = \omega_{L_1} \cdot \omega_{L_2}$$

where ω_{L_i} denotes the number of the conformations of loop L_i . Figure 3 shows that the conformational entropy computed from Equation 2 gives accurate results as tested against the exact computer enumeration. From Equation 2, the key for the pseudoknot conformational entropy calculation is to compute the conformational count ω_{L_i} for the loops.



Figure 3. Comparison for the calculated conformation entropy from the exact computer enumeration (Times) and the result from Equation 2 (Triangle), where L_1 and L_2 are the lengths (nt) for the loops across the the deep groove of S_2 and the shallow groove of S_1 , respectively, and Ω_s is the number of the pseudoknot loop conformations. The helix lengths for S_1 and S_2 are fixed at 5 and 7 bp, respectively (See Figure 1b).

In Table 1, we show our results from exact computer enumeration for the two loops. The results are given as the loop entropy parameters defined as the following.

$$\Delta S_{L_i} = -k_{\rm B} \ln \omega_{\rm coil \ (L_i)} / \omega_{L_i}$$

Here ΔS_{L_i} is the loop entropy for loop L_i , $\omega_{\text{coil}(L_i)}$ is the number of the corresponding coil conformations for L_i . To obtain $\omega_{\text{coil}(L_i)}$ and ω_{L_i} , we enumerate the number of self-avoiding loop conformations on the diamond lattice. The volume exclusion between different monomers in the loop and between the loop and the helix (S_1 for L_2 and S_2 for L_1) are rigorously accounted for. We note that due to the discreteness of the virtual bond configurations in the diamond lattice, certain conformations with long helix stems and very short loops cannot be realized in the diamond lattice. However, these loops, which are denoted by * in Table 1, may be viable in realistic structures. For these special loops, the conformations would be very restricted, so we assume their loop entropies are equal to the minimal loop entropies.

We denote the loop length by l (nt). For large loops (l > 12 nt), the enumeration of the loop conformations is not possible because of the exponentially increasing computational time. According to the results from the exact computer enumeration for $l \leq 12$ nt loops, we fit the following formula for the loop entropies:

$$\ln \omega_{L_i} = a \ln (l - l_{\min} + 1) + b(l - l_{\min} + 1) + c;$$

$$\ln \omega_{\text{coil} (L_i)} = 2.14 \ l + 0.10$$
4

where l_{\min} is the minimal loop length to form a H-pseudoknot for the given helix stem (S_1 for L_2 and S_2 for L_1). The l_{\min} values for different helix stems are obtained from the experimental data (35,45). The above equations combined with Equation 3 give the loop entropies for a pseudoknot (46–48).

In Table 2, we present our results for (l_{\min}, a, b, c) for L_1 (across the narrow groove of S_2) and L_2 (across the wide groove of S_1). To test the accuracy of the above fitted loop

Stem size (bp)	Loop size (nt)											
	1	2	3	4	5	6	7	8	9	10	11	12
(S_2)	L_1 (across the deep groove of S_2)											
2			_	6.2	6.4	6.4	6.6	6.8	6.9	7.1	7.2	
3		6.4*	6.4*	6.4	6.6	6.6	6.8	6.9	7.1	7.3	7.5	
4	4.4*	4.4*	4.5	5.4	5.6	6.0	6.3	6.6	6.9	7.1	7.3	
5	2.3	4.4	4.6	5.7	6.0	6.5	6.9	7.2	7.5	7.8	8.0	
6	2.3	4.4	4.8	5.8	6.0	6.5	6.8	7.1	7.4	7.6	7.8	
7	2.3	4.4	5.0	5.9	6.2	6.8	7.0	7.3	7.6	7.8	8.0	
8		4.4	5.2	5.7	6.4	6.7	7.1	7.3	7.5	7.7	7.9	
9		5.5*	5.5	6.4	6.7	7.2	7.5	7.9	8.1	8.3	8.5	
10		6.9*	6.9*	6.9	7.5	7.7	8.1	8.3	8.6	8.8	8.9	
11				_	8.7	8.8	8.9	9.1	9.2	9.3	9.3	
12				_	9.8	9.2	9.5	9.6	9.7	9.8	9.8	
(S_1)	L_2 (acro	oss the shall	low groove	of S_1)								
2	_	_		7.6	7.0	7.0	7.1	7.2	7.3	7.4	7.5	7.7
3		6.5*	6.5*	6.5	6.6	6.7	6.9	7.1	7.2	7.4	7.6	7.7
4			9.2*	9.2*	9.2	8.9	8.9	8.9	9.0	9.0	9.1	9.2
5				9.8*	9.8*	9.8	9.1	8.9	8.8	8.8	8.8	8.8
6				11.9*	11.9*	11.9*	11.9	11.0	10.4	10.1	9.9	9.8
7		_			12.4*	12.4*	12.4*	12.4	11.4	11.0	10.7	10.5
8		_			12.1*	12.1*	12.1*	12.1	11.6	11.4	11.2	11.1
9						13.7*	13.7*	13.7*	13.7	12.6	12.0	11.5
10		_				13.7*	13.7*	13.7	12.7	12.2	11.8	11.5
11	_		_						15.9	14.1	13.0	12.4
12	_	_	_	_	_	_	_	_	18.7	15.8	14.2	13.2

Table 1. Pseudoknot loop entropies parameters defined in Equation 3.

The upper half of the Table gives $(-\Delta S_{L_1}/k_B)$ for loop L_1 as a function of the S_2 helix stem length and the L_1 loop length. The lower half of the Table gives $(-\Delta S_{L_1}/k_B)$ for loop L_2 as a function of the S_1 helix stem length and the L_2 loop length. See Figure 1 for the stem–loop construct of a pseudoknot. The * entries in the Table indicate the long stem and short loop structures that cannot be realized in the diamond lattice but may be viable for a realistic pseudoknot. For these restricted loops, we use the entropies of the minimal loop lengths for the same helix length.

Table 2. The parameters for the loop entropies in Equation 4.

S_2 stem le	ngth (bp)										
	2	3	4	5	6	7	8	9	10	11	12
l_{\min}	4	2	1	1	1	1	2	2	2	5	5
а	0.12	0.39	-2.14	-2.22	-2.40	-2.61	-1.17	-1.66	-1.43	-0.14	0.77
b	1.96	1.92	2.15	2.11	2.18	2.21	2.03	2.09	2.09	2.06	1.84
С	0.52	-3.89	-2.09	-2.25	-2.33	-2.32	-1.96	-1.98	-2.93	0.15	-0.65
S_1 stem le	ngth (bp)										
	2	3	4	5	6	7	8	9	10	11	12
l_{\min}	4	2	3	4	4	5	5	6	6	9	9
a	0.95	0.32	1.77	3.99	7.73	8.38	4.52	9.05	4.77	2.74	4.69
b	1.84	1.92	1.82	1.55	1.29	1.16	1.61	1.15	1.68	2.05	1.80
С	-0.67	-3.90	-5.76	-5.86	-12.67	-11.45	-7.58	-11.45	-6.78	1.38	-1.11

The upper half is for loop L_1 and the lower half is for loop L_2 .

entropy formula, in Figure 2b we show the comparison between Equation 4 and the results from the exact computer enumeration. We find that the above fitted formula can indeed provide good approximations for the loop entropies.

From Table 1 and Figure 2b, we find that the loop entropy $|\Delta S_{L_i}| = k_{\rm B} \ln \omega_{\rm coil} (L_i)/\omega_{L_i}$ shows an intriguing loop lengthdependence. As the loop size *l* is increased, we find that (i) $|\Delta S_{L_1}|$ and (ii) $|\Delta S_{L_2}|$ for short helix stem S_1 would monotonically increase. However, for loop L_2 with longer helix stem S_1 , we find a non-monotonic behavior: $|\Delta S_{L_2}|$ first decreases for small *l* and then increases for large *l*. Such non-monotonic behavior for L_2 , the loop spanning across the minor groove, is evident from the exact computer enumeration results (e.g. ΔS_{L_2} for stem length $S_1 = 4$ bp in Table 1 and in Figure 2b). In the following, we explain why the pseudoknot loop entropies show such distinctive behavior.

If the helix has zero or weak excluded volume, the loop entropy $|\Delta S_{L_i}|$ would increase monotonically with the loop size l, because $\omega_{\text{coil }(L_i)}$ increases faster with the loop length *l* than ω_{L_1} . With the helix stem S_2 , L_1 spans across the deep narrow groove of the helix S_2 and can thus have a small end-end distance. For a small end-end distance (compared to the loop length), the probability for the loop to bump into the helix is small, i.e. the excluded volume of the helix is rather weak. Therefore, $|\Delta S_{L_1}|$ increases monotonically with l, for L_2 , which span across the wide shallow groove of helix S_1 , the loop end-end distance for a long helix S_1 can be large. Consequently, for a short loop length l, the volume exclusion between the loop (L_2) and the helix (S_1) can be very strong. For such case, a small increase of the loop length l can greatly weaken the loop-helix volume exclusion, causing a notable increase in the number of loop conformations ω_{L_2} . This would result in a decrease in $|\Delta S_{L_2}| = k_{\rm B} \ln \omega_{\rm coil (L_2)} / \omega_{L_2}$. Therefore, $|\Delta S_{L_2}|$ decreases with l for a long S_1 stem. For large loops, the loop-helix volume exclusion becomes relatively weak and thus $|\Delta S_{L_2}|$ increases with l.

The above excluded volume-caused non-monotonic ΔS_{L_2} is supported by the results from the exhaustive computer enumeration. First, the results for $S_1 = 4$ bp in Table 1 shows a non-monotonic loop length-dependence; see also Figure 2b. Second, for longer S_1 stems, the loop-helix excluded volume interaction is strong and thus the non-monotonic behavior would be more pronounced. For example, $|\Delta S_{L_2}|$ for $S_1 = 6$ bp decreases from 11.9 to 9.8 k_B as the loop length l is increased from 7 to 12 nt. This implies a non-monotonic behavior, because for a very large l (\gg end–end distance of the loop), the loop-helix excluded volume interaction is weak, resulting in a monotonically increasing loop entropy.

Though we are lacking in exact computer enumeration data for very large loops due to the extremely demanding computational time for exhaustive enumeration, broad experimental tests presented in the following sections suggest that the loop entropy parameters given in Table 1 and Equation 4 may be reliable for the loop sizes tested. In addition, the entropy parameters derived here might be suffice for biologically significant sequences, which are unlikely to involve very large loops (length \gg 12 nt). With the increasing availability of the experimentally measured loop entropy data, further refinements for Equation 4, which is accurate only for short and mid-size loops, can be possible.

Our current model has two advantages. First, the model is based on the virtual bond representation of the atomic P and C_4 atoms of the nucleotides, so the model can treat atomic details. Second, the model explicitly accounts for the excluded volume interactions between the helices and the loops and between the loop nucleotides. In the following sections, based on the pseudoknot entropy parameters that we derived, we compute the partition function and the native structure as well as the folding free energy landscape for RNA pseudoknots. Through extensive experimental comparisons on the native structures and the melting curves, we validate the entropy model.

Partition function

In this section, we develop a recursive algorithm (42,49,50) to compute the partition function for all the possible structures that contain pseudoknots and secondary structures.

In order to account for the excluded volume interactions between different structural subunits, we classify different conformational types. A chain segment from nucleotides ato b is connected to the rest of the molecule at the terminal nucleotides a and b. To account for the conformational viability between the chain segments, we classify the conformational types according to the excluded volume near a and b. We assume that a lone base pair is not stable, so we classify structures according to base stacks.

(i) We classify conformations according to whether the terminal nucleotides a and b are involved in base pair or not: the conformation is 'closed' if both a and b are involved in base pairing (such as the structures shown in Figures 4b, c and f) and is 'open' otherwise (such as the structure shown in Figures 4a and d). It is important to note that for a closed conformation, a and b can either base pair with each other (such as the hairpin structure shown in Figure 4f) or with different nucleotides (such as the pseudoknot structure shown in Figure 4f).

(ii) For the open conformations, according to whether the nucleotides adjacent to the terminal nucleotides (*a* and *b* in Figure 4), i.e., nucleotides a_1 and b_n in Figure 4, are involved in base pairing, we classify four types of conformations (see Figure 4d):

type — LR	if a_1 is adjacent to a and b_n is adjacent to	b;
type — L	if only a_1 is adjacent to a ;	
type — R	if only b_n is adjacent to b ;	
type — M	if neither a_1 nor b_n is adjacent to a or b .	5

For convenience, we use the following rules in our notations: C = 'partition function for the closed conformations', O = 'partition function for the open conformations', subscript '2' = secondary structures, '3' = pseudoknots, '23' = secondary structures + pseudoknots. For example, $C_2(a, b)$ and $O_2^t(a, b)$ (t = L, R, M and LR) are the partition functions for the closed secondary structures and the open secondary structures for a chain segment from a to b, respectively.

Closed conformations. The partition function for the closed conformations from *a* to *b* can be calculated as the sum of the partition functions for all the possible closed secondary structures without pseudoknot $C_2(a, b)$ and for all the possible structures with pseudoknots $C_3(a, b)$:

$$C_{23}(a,b) = C_2(a,b) + C_3(a,b)$$
6

 $C_2(a, b)$ can be calculated from the dynamic program by growing the chain step by step as described in equations 12–16 in the Appendix. $C_3(a, b)$ can be computed from the sum over all the possibilities for the helix stem lengths S_1 and S_2 and the loop lengths L_1 , L_2 and L_3 as shown in Figure 4f:

$$C_3(a,b) = \sum_{L_1} \sum_{L_2} \sum_{L_1} \sum_{L_2} \sum_{L_3} e^{-\Delta G(S_1, S_2, L_1, L_2, L_3)/k_B T}$$

where $\Delta G(S_1, S_2, L_1, L_2, L_3)$ is the free energy of the pseudoknot, which can be obtained from the loop entropy parameters in Tables 1 and 2 and the enthalpy and entropy parameters for the helices. In the Results and Discussion section, we illustrate the detailed calculation of the pseudoknot free energy.

Open conformations. The partition function for the open conformations (with pseudoknots) $O_{23}^t(a, b)$ (t = conformational type L, M, R, LR; see Figure 4d) can be computed from the same recursive relations as the ones that we have derived for the secondary structures; see Equations 13–16 in the Appendix with C_2 and O_2 replaced by C_{23} and O_{23} , respectively. From the recursive relations and Equation 17, we can compute efficiently the partition function $O_{23}^t(a, b)$ for any chain segment from a to b.



Figure 4. An open conformation (**a**) and a closed conformation with the closing stack connected to a loop (**b**) or to a stack (**c**). The closed conformation in (b) is formed from the open conformation in (a) through the closure of the unstacked loop of length l in (b). (**d**) The four types of open conformation (L, M, R and LR). (**e**) The partition function for *M*-type conformations for a chain from *a* to *b* can be computed as the sum of the partition function for a shorter chain from *a* to *b* - 1. (**f**) A closed conformation for RNA secondary structure and pseudoknots. Here we use a polymer graph to represent $C_{23}(a,b)$. The straight dark lines in the polymer graph represent the nucleotide backbone. The curved links represent the base pairs between the nucleotides. The shaded region between two curved links in the polymer graph corresponds to a helix stem.

Total partition function. Through the recursive relation, we can treat not only the simple pseudoknots, secondary structures, but also more complex structures with both secondary structures and pseudoknots (see Figure 5a) and structures with multiple pseudoknots (see Figure 5b). As shown in Figure 4d, we can treat structures with one or more closed conformations connected in series, where each closed conformation can be in the form of a secondary structure or a pseudoknot as shown in Figure 4f, or a mixture of secondary and pseudoknotted structures.

The total partition function Q(a, b) for a chain from a to b is given by the sum of the partition functions for all the different types of conformations:

$$Q(a,b) = 1 + C_{23}(a,b) + \sum_{t=L,R,M,LR} O_{23}^{t}(a-1,b+1)$$
 7

The first term accounts for the contribution from the unfolded coil state. The computational time scales with the chain length N as $O(N^6)$ and the memory scales as $O(N^2)$.



Figure 5. The density plots for the base pairing probabilities and the predicted stable structures for (a) TYMV and (b) TMV pseudoknots at $T = 37^{\circ}$ C with the coaxial stacking. The upper diagrams are the corresponding polymer graphs for the predicted native structures. The straight dark lines in the polymer graph represent the nucleotide backbone and the curved links represent the base pairs.

RESULTS AND DISCUSSION

Pseudoknot free energy calculation

For a pseudoknot with helix stem lengths S_1 and S_2 and loop lengths L_1 , L_2 and L_3 as shown in Figure 4f, the pseudoknot free energy $\Delta G(S_1, S_2, L_1, L_2, L_3)$ can be computed as the sum of the free energies of the stems, loops and possibly the coaxial-stacking.

$$\Delta G(S_1, S_2, L_1, L_2, L_3) = \Delta G_{S_1} + \Delta G_{S_2} - T\Delta S_{L_1} - T\Delta S_{L_2} + \Delta G_{cs} + \Delta G_{assemble}$$
8

Here ΔS_{L_i} is the entropy of loop L_i (see Equation 3), ΔG_{S_i} (i = 1, 2) is the free energy of helix stem S_i , and ΔG_{cs} is the free energy of the coaxial stack between the two helices (51).

 $\Delta G_{\text{assemble}}$ in Equation 8 is introduced to account for the entropy change as the two subunits $(L_1 + S_2 \text{ and } L_2 + S_1)$ are assembled into the pseudoknot. In Equation 3 (and Table 1), ω_{L_1} and ω_{L_2} are computed with the presence of the helices, therefore, the number of conformations for the assembled pseudoknot can be calculated as $\Omega_s = \omega_{L_1} \omega_{L_2}$ as in Equation 2. On the other hand, $\omega_{\text{coil} (L_i)}$ for L_1 and L_2 are calculated as two separate chains. As the two (coil) chains are assembled to form a longer chain, due to the different ways for the connection between the two chains, the total number of coil chain conformations is given by $\omega_{\text{assemble}} \cdot \omega_{\text{coil} (L_1)} \cdot \omega_{\text{coil} (L_2)}$, where $\omega_{\text{assemble}} \sim 3 \times 3 = 9$ is the number of conformations for the junction nucleotide (= two virtual bonds, each with three possible orientations) between the two chains. The corresponding free energy for the ω_{assemble} effect is $\Delta G_{\text{assemble}} = k_{\text{B}}T \ln \omega_{\text{assemble}} \sim k_{\text{B}}T \ln 9 \simeq 1.3 \text{ kcal/mol at } T = 37^{\circ}\text{C}.$

In Equation 8, we have neglected the loop enthalpy and the stability of the single strand segment L_3 , which is assumed to be short and flexible. From the NMR structural studies (43,52), the two stems in the pseudoknot can be coaxially stacked. Especially for $L_3 = 0$ the two stems can be juxtaposed into coaxial positions and we need to consider the coaxial stacking in the stability calculation.

For a given pseudoknot structure, we obtain (i) the loop entropies ΔS_{L_i} (i = 1, 2) from Tables 1 and 2, (ii) the helix free energy $\Delta G_{S_i} = \Delta H_{S_i} - T\Delta S_{S_i}$ from the nearest neighbor interaction model with the enthalpy ΔH_{S_i} and the entropy ΔS_{S_i} parameters computed from the Turner rule (53), and (iii) the coaxial stacking free energy ΔG_{cs} from the sequencedependent parameters in Ref. (51).

Using Equation 8, we can compute the free energy for simple H-pseudoknots, such as the T4-35 pseudoknot shown in Figures 6c and 7a. For more complex structures, which involve multiple pseudoknots or a mixture of the pseudoknots and secondary structures, such as the TYMV and TMV structures shown in Figures 5a and 5b respectively, we can estimate the total free energy of the structure as the sum of the free energy of the component structures (pseudoknots, secondary structures). In this section, using T4-35, TYMV and TMV pseudoknots as three representative examples for the different levels of complexity, we show how to compute the pseudoknot free energy using our loop entropy parameters. For all the illustrative calculations in this section, we assume temperature $T = 37^{\circ}$ C.

T4-35 pseudoknot. For the T4-35 pseudoknot shown in Figures 6c and 7a, there are two stems S_1 (=5 bp) and S_2 (=7 bp) and two loops L_1 (=1 nt) and L_2 (=4 nt).

- (i) (*Stems S*₁ and *S*₂): The nearest neighbor interaction model with the base stack thermodynamic parameters (53) gives the stabilities (in kcal/mol) for stems *S*₁ and *S*₂: Δ*G*_{*S*₁} = -6.6; Δ*G*_{*S*₂} = -11.2.
 (ii) (*Loops L*₁ and *L*₂): (*L*₂, *S*₁) = (4 nt, 5 bp), (*L*₁, *S*₂) =
- (ii) (Loops L_1 and L_2): (L_2 , S_1) = (4 nt, 5 bp), (L_1 , S_2) = (1 nt, 7 bp). From the lower (upper) half of Table 1 for L_2 (L_1), we find $\Delta S_{L_2} = -9.8 k_B$ and $\Delta S_{L_1} = -2.3 k_B$.
- (iii) $(T4-28 \ pseudoknot)$: The stability of the T4–28 pseudoknot (without coaxial stacking) at $T = 37^{\circ}C = 310 \text{ K} (k_{\text{B}}T = 0.62 \text{ kcal/mol})$ is equal to:

$$\Delta G = \Delta G_{S_1} + \Delta G_{S_2} - T(\Delta S_{L_1} + \Delta S_{L_2}) + \Delta G_{\text{assemble}}$$

= (-6.6) + (-11.2) + (6.1) + (1.4) + (1.3)
= -9.0 (kcal/mol) 9

The above calculated free energy is close to the experimentally measured -8.1 kcal/mol in the mixed 3 mM Mg²⁺ and 50 mM Na⁺ ion condition (24). If we include the coaxial stacking (through the base stack 5'AG-CU3'), which has $(\Delta H, \Delta S) = (-12.5 \text{ kcal/mol}, -32.6 \text{ cal/mol/K})$ (51), the pseudoknot stability would be changed to $\Delta G =$ -2.5 kcal/mol, which disagrees with the experimental result. Therefore, coaxial stacking is unlikely to play a role in the T4–35 pseudoknot. The NMR studies for T2 pseudoknot (43) indicate that the C7–G16 base pair in the helix–helix junction is over-rotated by 18° compared with a continuous A-form helix. This may result in the disruption of the coaxial stacking in the junction. Therefore, our theoretical prediction is in accordance with the experiment.

TYMV pseudoknot. As shown in Figure 5a, the TYMV pseudoknot consists of two parts: a hairpin denoted as C_2 -1 from C1 to G15 and a pseudoknot denoted as C_3 -1 from C16 to A39.

- (i) (*Hairpin* C₂-1): The nearest neighbor interaction model gives the stability $\Delta G_{C_2-1} = -3.9$ kcal/mol.
- (ii) (Pseudoknot C_3 -1):
 - (a) (*Stems* S_1 and S_2): The nearest neighbor model gives the stabilities (in kcal/mol) for stems 1 and 2: $\Delta G_{S_1} = -6.0$ and $\Delta G_{S_2} = -10.5$
 - (b) (*Loops* L_1 and L_2): (L_2 , S_1) = (3 nt, 3bp), (L_1 , S_2) = 3 nt, 6 bp). From the lower (upper) half of Table 1 for L_2 (L_1), we find $\Delta S_{L_2} = -6.5k_B$ and $\Delta S_{L_1} = -4.8k_B$.
 - (c) (*Pseudoknot* C_3 -1): The stability for pseudoknot C_3 -1 is equal to

$$\Delta G_{C_3-1} = \Delta G_{S_1} + \Delta G_{S_2} - T(\Delta S_{L_2} + \Delta S_{L_1}) + \Delta G_{\text{assemble}}$$

= -8.2 (kcal/mol)

(iii) (*TYMV pseudoknot*): The total stability for the TYMV pseudoknot is given as below.

$$\Delta G_{\text{tot}} = \Delta G_{C_2-1} + \Delta G_{C_3-1} + \Delta G'_{cs}$$

= $\Delta G_{C_2-1} + \Delta G_{C_3-1} + \Delta G(5'GC - GC3')$
= $-3.9 + (-8.2) + (-4.3) = -16.4 \text{ (kcal/mol)}$

where $\Delta G'_{CS}$ is the coaxial stacking free energy between stem 0 in the hairpin C_2 -1 and stem 1 in the pseudoknot C_3 -1. The $\Delta G'_{CS}$ parameter is from Ref. (51).

If we include the (5'AG-CU3') coaxial stacking between stems 1 and 2 within the C_3 -1 pseudoknot (51), the total stability would become -18.9 kcal/mol.

TMV pseudoknot. As shown in Figure 5b, the TMV pseudoknotted structure consists of three H-pseudoknots (C_3 -1, C_3 -2 and C_3 -3).

- (i) (*Stems*): From the nearest neighbor model and the thermodynamic parameters for the base stacks (53), we obtain the stabilities (in kcal/mol) for the six helix stems (two in each pseudoknot): $\Delta G_{S_1} = -7.3$; $\Delta G_{S_2} = -5.9$; $\Delta G_{S'_1} = -4.1$; $\Delta G_{S'_2} = -10.4$; $\Delta G_{S''_1} = -5.6$; $\Delta G_{S''_1} = -8.3$.
- (ii) $(Loops L_1, L_1', L_1'')$: These loops span across the deep narrow grooves of stems 2, 2', and 2'', with [loop length (nt), stem length (bp)] = [1, 4], [1, 6], and [5, 5], respectively. From the upper half of Table 1, we have $\Delta S_{L_1} = -4.4 k_{\rm B}$; $\Delta S_{L'_1} = -2.3 k_{\rm B}$; $\Delta S_{L''_2} = -6.0 k_{\rm B}$.
- $\Delta S_{L_1} = -4.4 \ k_{\rm B}; \ \Delta S_{L'_1} = -2.3 \ k_{\rm B}; \ \Delta S_{L''_1} = -6.0 \ k_{\rm B}.$ (iii) (Loops L_2, L_2', L_2''): These loops span across the wide shallow grooves of stems 1, 1', and 1", with [loop length (nt), stem length (bp)] = [4, 4], [3, 3], and [7, 4], respectively. From the lower half of Table 1, we have $\Delta S_{L_2} = -9.2 \ k_{\rm B}; \ \Delta S_{L'_2} = -6.5 \ k_{\rm B}; \ \Delta S_{L''_2} = -8.9 \ k_{\rm B}.$



Figure 6. The comparison between the calculated melting curves and the experimentally measured results for eight pseudoknots sequences (**a**) T4–28, (**b**) T4–32, (**c**) T4–35 (24,26), (**d**) G80 (23), (**e**) mIAP (27), (**f**) the telomerase pseudoknot domain (PK_{WT}), (**g**) (28) and (**h**) (29) are two mutants of the telomerase pseudoknot domain. The Y-axis is the heat capacity. The calculated melting curves have been normalized such that the theoretical and the experimental curves have the same peak value. The ion conditions are 50 mM NaCl and 1.0 mM Mg²⁺ for (a), (b) and (c), 1 M KCl for (d), 50 mM KCl for (e), 200 mM KCl for (f), (g) and 200 mM NaCl for (h). Because our calculated enthalpic and entropic parameters for the base stacks are from Turner's rules for 1 M NaCl salt condition, our model generally overestimates the melting temperatures. Results with and without coaxial stacking (CS) are both presented.



Figure 7. The density plot for the base pairing probabilities, the predicted stable structures, and the density plot for the free energy landscapes for T4–35 pseudoknot at different temperatures. In the free energy landscape F(n, nn), darker color means lower free energy. *n* and *nn* denote the numbers of the native and the non-native base pairs, respectively. At $T = 70^{\circ}$ C, the partially unfolded pseudoknot structure (Z) coexists with the hairpin structure (X).

- (iv) (*Pseudoknots* C_3-1 , C_3-2 , C_3-3): From the above parameters for the helices and the loops, we use Equation 8 to obtain the stability (in kcal/mol) for each pseudoknot. We have $\Delta G_{C_3-1} = -3.5$, $\Delta G_{C_3-2} = -7.7$, $\Delta G_{C_3-3} = -3.4$.
- (v) (*TMV pseudoknot*): The stability for TMV pseudoknot is equal to

$$\Delta G = \Delta G_{C_3-1} + \Delta G_{C_3-2} + \Delta G_{C_3-3} + \Delta G_{cs21'} + \Delta G_{cs2'1''}$$

= (-3.5) + (-7.7) + (-3.4) + (-3.5) + (-1.1)
= -19.2 (kcal/mol)

where ΔG_{csij} is the free energy of coaxial stacking between stems *i* and *j*. The coaxial stacking between $C_3 - 1$ and $C_3 - 2$ is through the 5'AC-GU3' base stack between stem 2 and stem 1', and the coaxial stacking between $C_3 - 2$ and $C_3 - 3$ is through the 5'AU-UG3' base stack between stem 2' and stem 1". We note that the coaxial stacking parameter for 5'UA-UG3' is not listed in Ref. (51), so we use the stacking energy from the Turner rule (53). The two stems in each H-pseudoknot element may also be coaxially stacking. The coaxial base stack 5'GU-AC3' between stems 1 and 2 can stabilize C_3 -1 by -3.5 kcal/mol, and the base stack 5'UA-UA3' between stem 1' and stem 2' can stabilize C_3 -2 by -2.2 kcal/mol. For C_3 -3, however, the coaxial stacking, if formed, would involve a noncanonical base pair (C-A) and would be unstable. So we ignored the coaxial stacking for C_3 -3. The total extra stability provided by the coaxial stacking within the H-pseudoknot elements (C_3 -1 and C_3 -2) could reach (-3.5) + (-2.2) = -5.7 kcal/mol.

Pseudoknot structure prediction

The probability P_{ij} for the formation of a base pair (i, j) can be calculated from the conditional partition function Q_{ij} for the ensemble of conformations with base pair (i, j) formed: $P_{ij} = Q_{ij}/Q$. From the distribution of the base pairing probability, we can deduce the stable structures for a given temperature *T*. Specifically, the structural prediction involves the following two steps. We first compute P_{ij} for all the possible base pairs (i, j). From P_{ij} , we identified the helices as the consecutive (i, j) pairs with large P_{ij} values. For example, the density plot for P_{ij} in Figure 5a for the TYMV pseudoknot has three trains of dark dots (= large P_{ij} 's), indicating three stables helices (stems 1, 2 and 3).

We predict the stable structures at $T = 37^{\circ}C$ for pseudoknot-forming sequences which have known experimentally measured native structures. We quantify the accuracy of our predictions using the sensitivity (SE) and the specificity (SP) parameters (54,55). The SE is defined as the ratio between the number of the correctly predicted base pairs and the number of the base pairs in the experimental determined structure, and the SP is defined as the ratio between the number of the correctly predicted base pairs and the number of base pairs in the predicted structure. A perfect prediction would have (SP, SE) = (1.0, 1.0) and a completely failed prediction would have (SP, SE) =(0.0, 0.0). NMR experiments suggest that the two stems in an H-type pseudoknot can be coaxial (43,52,56) or bent (18,57), so we perform our computations for two cases (with and without coaxial stacking) separately.

Viral ribosomal frameshift signals and viral ribosomal readthrough signals pseudoknots. In Table 3, we summarize the accuracy of our theoretical predictions for the native structures with coaxial stacking for 15 viral ribosomal frameshift signals and 7 viral ribosomal readthrough signals pseudoknots (58). We truncate the sequences and retain only the pseudoknot portions; see Table 3 for the sequences. For the viral ribosomal frameshift signals, BEV, EAV, HCV-229E, LDV-C and RSV sequences are not included because the computational time is demanding for these long (>100 nt) sequences. From Table 3, we find that, except for the BLV and CABYV sequences, our model can give good predictions for all the other 20 sequences. If the second lowest free energy structure is also considered, the predictions for BLV and CABYV would be exact (see Table 3). For BLV, the predicted alternative structure is a hairpin formed through the disruption of the helix stem close to the 3' end in the native pseudoknot. The hairpin is predicted to be more stable than the native pseudoknot by 0.7 kcal/mol. For CABYV, the model predicts a misfolded hairpin with the helix stem formed by the base pairs from G1495-C1511 to G1501-C1505. The misfolded hairpin is predicted to be 1.6 kcal/ mol more stable than the native pseudoknot. The inaccuracy in the predictions for the BLV and CABYV pseudoknots may be caused by the neglected tertiary interactions, such as base triples. The A \cdot (C-G) and C \cdot (C-G) base triples have been found to significantly stabilize the ribosomal frameshifting pseudoknots (18-20), such as ScYLV and BWYV, and may also play important roles in the BLV and CABYV stabilities.

As a test, we also predict the pseudoknots without considering the coaxial stacking. We find that removing the coaxial stacking causes no change in the predicted native pseudoknot except for three sequences: PEMV, PLRV-S and FeLV, for which the predicted structures become significantly less accurate with (SE, SP) = (0.60, 0.86), (0.50, 0.40) and (0.57, 0.53), respectively. This clearly shows that the coaxial stacking is an important stabilizing force for these pseudoknots. More specifically, PEMV and PLRV-S are stabilized by the coaxial stacking 5'UC-GA3' and FeLV is stabilized by the coaxial stacking 5'CC-GG3'. The energy parameter for the 5'UC-GA3' coaxial stacking is not listed

Table 3. The accuracies of the structure predictions for the viral ribosomal frameshifting signal and the viral ribosomal readthrough signal pseudoknots (58).

Viral ribosomal frameshifting signals 1 BChV 26 G1595-C1620 1.00 2 BLV 27 G1604-U1630 0.67 3 BWYV 26 C1566-G1591 1.00	1.00
1 BChV 26 G1595-C1620 1.00 2 BLV 27 G1604-U1630 0.67 3 BWYV 26 C1566-G1591 1.00	1.00
2 BLV 27 G1604–U1630 0.67 1.00 3 BWYV 26 C1566–G1591 1.00	0.06 (1)
1.00 3 BWYV 26 C1566-G1591 1.00	0.86(1)
3 BWYV 26 C1566-G1591 1.00	1.00 (2)
2 20 01000 01001 1.00	1.00
4 BYDV–NY–RPV 27 G1706–C1732 1.00	1.00
5 CABYV 27 G1494–C1520 0.00	0.00(1)
1.00	1.00 (2)
6 EIAV 35 G1797–C1831 1.00	1.00
7 FIV 35 G1893–C1927 1.00	1.00
8 IBV 69 G81–U149 1.00	0.78
9 MMTVgag/pro 34 G2090–U2123 1.00	1.00
10 PEMV 28 U2042–C2069 0.90	1.00
11 PLRV–S 26 G1781–G1806 1.00	1.00
12 PLRV–W 26 G1676–G1701 1.00	0.88
13 PRRSV–LV 59 G7402–G7459 1.00	0.95
14 PRRSV-16244B 58 G7701-G7758 1.00	1.00
15 SRV1gag/pro 37 G2337–C2373 1.00	1.00
Viral ribosomal readthrough signals	
16 AKV-MuLV 50 G2261-C2310 1.00	0.79
17 BaEV 50 G2607-C2656 1.00	1.00
18 Cas–Br–E–MuLV 50 G2239–C2288 1.00	0.75
19 FeLV 50 G2196–G2245 1.00	0.82
20 GaLV 49 G2202–G2250 1.00	0.79
21 Mo-MuLV 50 G1982-C2031 1.00	0.79
22 SNV 50 G183-C232 1.00	0.83

The calculations are performed at the standard conditions (1 M NaCl, 37° C). These sequences are truncated to keep the pseudoknot part. In the calculation, we consider the coaxial stacking. The SE is defined as the ratio between the number of the correctly predicted base pairs and the number of the base pairs in the experimental determined structure, and the SP is defined as the ratio between the number of the correctly predicted base pairs and the number of base pairs in the predicted structure (54,55).

in Ref. (51), so we use the stacking energy parameter from the Turner rule (53). The above two coaxial stacks contribute about -2.3 kcal/mol and -4.1 kcal/mol to the stability of the pseudoknots at $T = 37^{\circ}$ C, respectively. In addition, the PEMV and PLRV-S may be stabilized by the tertiary interactions (12) such as (C.C-G for PEMV) and (C.G-C for PLRV-S).

Pseudoknots with high binding affinity to the human immunodeficiency virus type 1 reverse transcriptase (HIV-1-RT). To further test the model, as shown in Table 4, we predict the native structures for 18 pseudoknot sequences [from Figure 2 in Ref. (59)] that have high binding affinity to HIV-1-RT as selected by the SELEX experiments. We find that, except for the sequence 2.9, our model can correctly predict the native structures for all the rest 17 sequences (out of the 18 sequences). For sequence 2.9, our model predicts that an alternative hairpin is more stable than the pseudoknot suggested in the functional studies (59). The theory–experiment difference for sequence 2.9 may come from other stabilizing interactions, such as triple bases (18–20,29) that are neglected in the current model.

Because of the use of the more physical entropy parameters, the present model gives more reliable predictions for these sequences (30). If we delete the coaxial stacking in our calculations, we find that the model predicts the same structures as listed in Table 4 except for sequence 2.5a (entry Sequence number

1.1

1.3a

29

2.4a

2.7a

2.11

1.7

1.17

2.1b

2.10

1.8

1.6

1.9h

2.12

2.5a

2.6b

2.2b

2.3a

ID

1

2

3

4

5

6

7

8

9

10

11

12

13

14 15

16

17

18

 Table 4. The accuracies of the structure prediction for the pseudoknot sequences with high affinity to HIV-1-RT (59).

Length

37

37

34

37

36

37

37

37

39

37

39

37

37

37

41

42

42

45

SP

0.91

0.80

0.42

0.82

0.80

0.80

0.67

0.80

0.80

0.90

0.70

0.78

0.77

1.00

0.91

1.00

0.60

1.00

14 PK17

23

SE

0.91

0.89

0.63

1.00

1.00

1.00

1.00

0.89

1.00

1.00

1.00

1.00

0.90

1.00

1.00

1.00

1.00

1.00

PK	1 and PK5-17 J	pseudokn	ots (61	,62).		
ID	Abbreviation	Length	SE	SP	T_m (Experiment)	T_m (Prediction)
1	PK1	19	1.00	1.00	73°C	83°C
2	PK5	26	1.00	1.00	64°C	67°C
3	PK6	25	1.00	1.00	63°C	67°C
4	PK7	25	1.00	1.00	65°C	67°C
5	PK8	24	1.00	1.00	_	_
6	PK9	23	1.00	1.00	59°C	72°C
7	PK10	25	1.00	1.00	_	_
8	PK11	24	1.00	1.00	63.5°C	68°C
9	PK12	23	1.00	1.00	_	
10	PK13	22	1.00	1.00	_	_
11	PK14	22	1.00	1.00	69.5°C	71°C
12	PK15	22	1.00	1.00	67°C	71°C
13	PK16	25	1.00	1.00		

Table 5. The accuracies for the prediction for the melting temperatures for the

In the calculation, the coaxial stacking is not included. In the experiment, the ion concentration is the mix 5 mM Mg^{2+} and 50 mM Na^+ , which may be close to the 1 M NaCl condition used in the calculation. For each of the sequences listed in the Table, we find a single transition in the calculated melting curve.

1.00 1.00 66°C

68°C

experimental value (-5.4 kcal/mol) (61). Adding the coaxial stacking free energy of -3.0 kcal/mol (5'CG-CG3') would also over-estimate the stability. Therefore, the coaxial stacking is unlikely to be formed in PK1. Though the predicted results cannot exclude the possibility of partial coaxial stacking, to simplify our calculation, we would not include the coaxial stacking in our further folding thermodynamics calculations for these sequences.

TYMV and TMV pseudoknots. The TYMV and TMV sequences are parts of the tRNA-like structures in viral RNAs (63–65). Their predicted native pseudoknots are more complex than the simple H-pseudoknots discussed above. As shown in Figures 5a and b, TYMV pseudoknot consists of a closed secondary structure (C_2 -1) and a H-pseudoknot (C_3 -1), and the TMV pseudoknot includes three H-pseudoknot elements ($C_3 - i$, i = 1, 2 and 3).

Shown in Figures 5a and b are the predicted native structures for the TYMV and TMV pseudoknots at $T = 37^{\circ}$ C. The predicted structures agree exactly with the experimental results both with and without considering the coaxial stacking within each pseudoknot.

Other pseudoknots. In Figure 6, we show the predicted native structures for other pseudoknot-forming sequences. The predicted structures agree exactly with the experimental measured ones, except for sequences PK_{WT} and PK_{DC} , which, as shown in Figures 6f and g, each has two predicted structures. For both PK_{WT} and PK_{DC} , one of the two predicted structures (structure II) agrees exactly with experiments and the other predicted structure (structure I) is formed through a single nucleotide sliding in the bulge of stem 2 in structure II.

In conclusion, nearly for all the 63 RNA pseudoknots that we have tested, the model can give good predictions as compared with experiments.

Folding thermodynamics

From the temperature-dependence of the partition function Q(T), we can predict the heat capacity melting curve C(T) for a given sequence: $C(T) = \frac{\partial}{\partial T} [k_B T^2 \frac{\partial}{\partial T} \ln Q(T)]$. We have

In the calculation, we consider the coaxial stacking.

15 in Table 4), for which the predicted native structure (without coaxial-stacking) has a poor (SE, SP) value of (0.50, 0.50). This clearly demonstrated the importance of the coaxial stacking in stabilizing the native pseudoknot for sequence 2.5a.

In Tables 3 and 4, the SP parameters for some predicted structures are not as good as SE. This is because these sequences (such as IBV) have a long loop L_2 across the shallow groove, and our model predicts that some of the nucleotides in the loops can form intra-loop base pairs. These predicted intra-loop base pairs are not explicitly listed in the experimentally measured structures, so the SP values appear to be less optimal.

Pseudoknots PK1 and PK5–PK17. Pseudoknots are found to be functionally active in viral RNAs (45) and ribosomal RNAs (60). A systematic study for the thermodynamic and structural properties of a specific type of short pseudo-knots was reported in Refs. (6,52,61,62). These pseudoknots are denoted as PK*i* (i = 1, 5, 6, ..., 17), where PK5-PK17 have the same helix stems but different loops. We here focus on the pseudoknot-forming sequences PK1 and PK5–17 [PK2–PK4 are hairpin-forming (61,62)]. Table 5 shows the results for the accuracies for our predicted native structures (PK1 and PK5–17) are in exact agreements with the experiments.

Our stability calculation can provide useful evidence for the existence/absence of coaxial stacking. For example, without considering the coaxial stacking, our theory predicts a stability of -4.2 kcal/mol for PK5. The predicted result is quite close to the experimental results of -4.9 kcal/mol (optical) or -4.3 kcal/mol (NMR) (62). Adding the coaxial stacking energy -4.1 kcal/mol (5'CC-GG3') (51) would overestimate the stability. So the PK5 pseudoknot is unlikely to have the full coaxial stacking. On the other hand, the deficiency in the stability without coaxial stacking suggests that we cannot exclude the possibility of partial coaxial stacking (52). For the PK1 pseudoknot, our predicted free energy without coaxial stacking is -6.4 kcal/mol, which is close to the calculated the melting curves for PK1 and PK5–17 (totally 14) sequences (6,61,62), and other 8 pseudoknot-forming sequences in Figure 6: T4–28, T4–32 and T4–35 (24,26), G80 (23), mIAP (27), the pseudoknot domain of human telomerase RNA (hTR) (PK_{WT}) (28) and its two mutants (PK_{DC} and ΔU 177) (29).

Table 5 shows the comparison between the predicted and experimental melting temperatures for PK1 and PK5–17 sequences. The Table shows good theory–experiment agreements. Since our pseudoknot stability calculation and the theory–experiment comparison indicate that the coaxial-stacking is unlikely to exist in these pseudoknots, we do not include coaxial stacking in the melting curve calculation for these sequences.

In Figure 6, we show the theory–experiment comparisons for eight sequences. For these sequences, since the two helical stems can be coaxial or bent, we compute the melting curves with and without coaxial stacking separately. As shown in the Figure, for pseudoknots T4–28, T4–32 and T4–35, the calculated melting curves with and without coaxial stacking give similar results, and the curves without coaxial stacking show slightly lower melting temperatures.

For the G80 and mIAP pseudoknots, however, the predicted results without the coaxial stacking give much better agreement with the experiments, indicating that coaxial stacking may not exist in these pseudoknots. The predicted absence of coaxial stacking is in agreement with the experiments (23,27), which suggest that the two stems are bent in G80 and mIAP.

For PK_{WT} and its mutants PK_{DC} and $\Delta U177$, the predictions with coaxial stacking are better than those without coaxial stacking, suggesting that coaxial stacking may play a role in stabilizing PK_{WT} and its mutant. Moreover, in the experiment (28), the measured stability for the native structure of wildtype at $T = 37^{\circ}$ C is -17.8 kcal/mol. Our model predicts that, without coaxial stacking, the predicted PK_{WT} structure II in Figure 6f, which agrees exactly with the experimental structure, has stability of $\Delta G_{II} = -12.3$ kcal/mol, and structure I has stability $\Delta G_I = -13.2$ kcal/mol. Adding the stability -3.9 kcal/mol for the coaxial stacking (5'UC-GA3') would give better agreement with the experiment.

Our theory–experiment agreements shown in Table 5 for the melting temperatures and in Figure 6 for the melting curves are not exact. A major problem is the ion effect. Ions play important roles in RNA pseudoknot folding stability (66,67). In our calculations, the enthalpic and entropic parameters for the helix stems are for the 1 M NaCl condition. In the experiments, however, different concentrations of Mg^{2+}/Na^+ or K^+ mixture are used. For lower experimental ion concentrations, such as 50 mM Na⁺ and 1.0 mM Mg²⁺ in Figures 6a–c; 50 mM K⁺ for Figure 6e; 200 mM K⁺ for Figures 6f and g; 200 mM Na⁺ for Figure 6h, the calculated melting temperatures are higher than the experimental results (See Figure 6). Nevertheless, our theory can predict not only the native structures but also the structural transitions as shown in the melting curves.

Coaxial stacking in pseudoknots

Pseudoknots, such as T4–28, T4–32, T4–35, G80 and mIAP that are predicted not to form coaxial stacking all have a short

(mostly 1 nt) loop L_1 across the deep groove. Short loops tend to form rigid conformations due to the volume exclusion from the helix stem. Such loop-helix interactions can strongly restrict the configurations of the helix–helix connection, such as coaxial stacking. For example, in the gene 32 mRNA pseudoknot of bacteriophage T2 (43), which has a single-nucleotide loop L_1 , the helix–helix junction is overrotated compared to the continuous A-form helix, causing the coaxial stacking impossible.

Longer flexible loops can make the formation of coaxial stacking or partial coaxial stacking possible. For example, pseudoknots PK_{WT} , PK_{DC} and Δ U177, which are predicted to have coaxial stacking, all have larger loops. The PK5 pseudoknot, which has flexible loops L_1 and L_2 with lengths of 4 and 6-nt, respectively, can possibly form partial coaxial stacking.

Free energy landscape and conformational switch

We introduce the free energy landscape to make a direct structure-free energy connection. The free energy F(x) for a macrostate is defined through a structural parameter (or a parameter set) x. Such a macrostate is a collection of conformations described by x. We choose x = (n, nn) = the number of (native, nonnative) base pairs. Here a base pair is called 'native' if it exists in the native structure and 'nonnative' otherwise. The free energy landscape F(n, nn) can be computed from the following conditional partition function Q(n, nn):

$$F(n,nn) = -k_B T \ln Q(n,nn); Q(\mathbf{x}) = \sum_{\operatorname{conf}(\mathbf{x})} e^{-E/k_B T}$$
 11

where $\sum_{\text{conf}(x)}$ is the sum over all the possible conformations with *n* native and *nn* nonnative base pairs, respectively. The free energy landscape provides a full view for the stability of all the conformations.

T4-derived pseudoknots. For the T4-derived pseudoknots (T4–28, T4–32 and T4–35 in Figures 6a–c), the stability calculation in the previous section shows that the predicted stability without coaxial stacking gives a better agreement with the experiment. So we assume no coaxial stacking for the T4-derived pseudoknots in the free energy landscape calculation. The predicted structures, as shown in Figures 6a–c, agree exactly with the NMR structures (24,26).

Shown in Figure 7 is the temperature-dependent of the free energy landscape and the conformational transitions for T4– 35. The pseudoknotted native structure (N) of T4–35 is predicted to be the single predominant state at $T = 0^{\circ}$ C. As the temperature is increased, stem 1 is partially unfolded at T = 70° C and a resultant native-like hairpin (X) emerges and coexist with the partially unfolded state (Z). At $T = 70^{\circ}$ C, the intermediate states Z and X are equally stable. This suggests that the first transition from the pseudoknot structure to native-like hairpin X happens at about 70° C, which is close to the 67° C transition temperature observed in the experiment (26). As the temperature is further increased, stem 2 is disrupted and the pseudoknot is fully unfolded. Since stem 2 is only slightly more stable than stem 1, the melting temperatures for the breaking of stem 1 and of stem 2 are close to each other, resulting in an apparent single peak in the melting curve (Figure 6c).

Since the unfolding pathway of T4-35 RNA pseudoknot involves the intermediate states (Z and X), a simple twostate analysis for the experimental data would give inaccurate results for the enthalpic and entropic parameters (24). In fact, a simple two-state assumption may result in an underestimation for the folding enthalpy. For example, the experiment gives a total folding enthalpy of about -100.7 kcal/mol (26), but the simple two-state assumption gives a enthalpy change of about -78 kcal/mol (24). From the nearest neighbor interaction model (53), we find that the enthalpy change without considering the coaxial stacking is equal to -93.5 kcal/mol, which is close to the -100.7 kcal/mol experimental result (26). If we further consider the contribution of the unpaired terminal nucleotides (5'CA-G5'), which has $(\Delta H, \Delta S) = (-9.0 \text{ kcal/mol}, -23.4 \text{ cal/mol/K})$ (53), our predicted enthalpy would be -102.5 kcal/mol, which agrees with the experimental result (-100.7 kcal/mol) very well.

G80 and mIAP pseudoknots. For the G80 and mIAP sequences, two well separated peaks are found in the melting curves (see Figures 6d and e). Our free energy landscape calculation shows that the first peak in the melting curve of G80 (mIAP) corresponds to the pseudoknot \rightarrow hairpin transition through the disruption of pseudoknot stem 1 (stem 2), while

the second transition corresponds to the subsequent unwinding of stem 2 (stem 1) in the hairpin; see Figures 8a and b for the equilibrium unfolding pathways for the two sequences. Experiments show that stem 1 melts before stem 2 for the G80 sequence (23) and stem 2 is disrupted before stem 1 for the mIAP sequence (27). Our predicted pathways are in accordance with the experimental findings. By comparing the unfolding pathways for G80 and mIAP in Figure 8, we find that the pseudoknot can unfold either through unzipping from stem 1 (close to the 5' end) or from stem 2 (close to the 3' end).

Human telomerase RNA (hTR) PK_{WT} and the PK_{DC} and $\Delta U177$ mutants. For PK_{WT} (28), our predicted structure II shown in Figure 9 agrees with the experimentally proposed structure (28), and the predicted structure I slightly differs from II by a single-nucleotide shift for the A-U base pairs in stem 2. The melting curve of PK_{WT} (Figure 6f) shows a major peak and a minor shoulder. Our free energy landscape and structural calculations (see Figure 9a) show that the major peak (at lower temperature) corresponds to the formation of two co-existing intermediates: a native-like hairpin structure formed through the disruption of stem 2 and a misfolded hairpin through the complete rearrangement of the base pairs. At $T = 80^{\circ}$ C, the native-like and the misfolded hairpin intermediates co-exist. The minor shoulder at higher



Figure 8. The density plot for the base pairing probabilities and the predicted stable structures for (a) the G80 and (b) mIAP pseudoknots at different temperatures.



Figure 9. The density plot for the base pairing probabilities and the predicted stable structures for (a) modified telomerase pseudoknot domain (PK_{WT}) in human and (b) its mutant (PK_{DC}) at different temperatures.

temperature in the melting curve corresponds to the unfolding of the hairpins (28). Our predictions are consistent with the experiment (28), except that an additional minor transition for the disruption of a triple base pairs (28,29) is observed in experiment. Such additional transition is absent in our prediction because the theory does not treat triple base pairs.

As shown in Figure 9, the folding/unfolding of PK_{WT} involves the native-like hairpin intermediate. The pseudoknot \rightarrow native-like hairpin switch is found to be functionally important in human telomerase RNA (hTR) (28,68,69). The predicted native-like hairpin in Figure 9 is in accordance with the experimentally proposed structure, except that the experimentally proposed hairpin contains tandem base pairs (28,68). The tandem base pairs are not treated in the present form of our model. Experimental mutational studies show that these tandem mismatches have only minor contributions to the hTR function (70).

Our predicted misfolded hairpin in the folding/unfolding of PK_{WT} may also be important for function. Phylogenetic analysis shows that the nucleotides 5' 107G-111C 3' and 5' 115U-119C 3' in the helix stem of the predicted misfolded hairpin intermediate are highly conserved (>80%) in the published vertebrate telomerase RNA sequences (71). The conserved nucleotides suggest that the misfolded hairpin may serve as the other (hairpin) candidate for the functionally important

pseudoknot \rightarrow hairpin conformational switch. For the PK_{DC} mutant, the 107GC \rightarrow AG mutation would disrupt the most stable base stacking (5'GC-GC3') in stem 2 of the wild-type (PK_{WT}) and thus destabilize stem 2. As a result, the mutation would cause a decrease in the melting temperature for the disruption of stem 2 (= the first transition in the unfolding process); see Figure 6f for the wild-type PK_{WT} and Figure 6g for the mutant PK_{DC}. In addition, the mutation significantly destabilizes the misfolded hairpin intermediate (see Figure 9a at T = 80°C) that emerges in the unfolding process of PK_{WT}. As a result, the misfolded hairpin is absent in the unfolding process of PK_{DC}.

In contrast, for $\Delta U177$, the deletion of U177 eliminates the 1 nt bulge loop in the (wild-type) stem 2 and thus stabilizes stem 2, which would cause an increase in the transition temperature for the melting of stem 2; see also the melting curves in Figures 6f and h. The prediction agrees with the experiment (29). From the above results for the mutants, we find that the two mutants (107GC \rightarrow AG and $\Delta U177$) cause significant changes in the pseudoknot stability. Such changes have been suggested to cause human telomerase diseases (28,29,68,72,73).

Tetrahymena thermophila telomerase RNA pseudoknot. In Tetrahymena thermophila, the telomerase RNA pseudoknot



Figure 10. The density plot for the base pairing probabilities and the predicted stable structures for (a) the telomerase pseudoknot domain in *Tetrahymena thermophia* and its mutants (b), (c), (d), (e) and (f) at different temperatures. We consider the coaxial stacking in the predictions.

is found to be functionally important. Mutants that disrupt the pseudoknot structure are found to cause low telomerase activity (74). We use the sequences in Figure 1b of Ref. (74) for the predictions. Calculations with and without the coaxial stacking give the same native structures; see Figure 10. Our model predicts that the 71–72G, 84–85C and 84–87C mutations will completely disrupt the pseudoknot structure, and the two compensatory mutations, Comp 1 (71–72G and 84–85C) and Comp 2 (84–87C and 71–74G) can restore the wild-type pseudoknot structure. These predictions are consistent with the experimental results (74). Moreover, the experiments (74) show that the mutants (71–72G, 84–85C and 84–87C) can reduce the telomerase activity to a very low level, which shows that a stable pseudoknot structure is crucial for the functional activity of the telomerase RNA (74,75).

CONCLUSION

Predicting the folding thermodynamics for RNA pseudoknots has been greatly hampered by the lacking of a physical model that can give accurate thermodynamic parameters (especially loop entropy parameters). In the present study, we develop a polymer statistical mechanical model to compute the conformational entropy from first principle. From the rigorous polymer chain conformational count, we derive a set of pseudoknot loop entropy parameters, from which we can compute the pseudoknot partition function and the free energy landscapes and the folding thermodynamics for RNA pseudoknots. Experimental tests for the predicted native structures and the thermal melting curves for a wide range of different pseudoknots show that the model is reliable. Furthermore, from the free energy landscapes and the base pairing probabilities, we predict the equilibrium folding pathways, folding stabilities for RNA pseudoknots. We find that the T4-pseudoknots, the mIAP and the G80 pseudoknots unfold though sequential disruptions of the helix stems without the formation of nonnative intermediates, while the pseudoknot domain of the hTR unfolds through the formation of the native-like and misfolded hairpin intermediates. These folding intermediates are found to be functionally important and mutations that alter the stabilities of these intermediates can cause disease.

ACKNOWLEDGMENTS

The authors acknowledge grant support from NIH (GM063732 to S-J.C). Funding to pay the Open Access publication charges for this article was provided by NIH through grant GM063732.

Conflict of interest statement. None declared.

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APPENDIX

RNA secondary structure partition function

Closed conformations. For RNA secondary structures without pseudoknots, the hierarchical relationship of the secondary structure results in the following recursive relation for the

partition functions (42); see also Figures 4b and c:

$$C_{2}(a-1,b+1) = (e^{-\Delta G_{\text{stack}}/k_{B}T})\{C_{2}(a,b) + e^{\Delta S_{\text{unstacked}}(b-a-1)/k_{B}} + \sum_{t,l} e^{\Delta S_{\text{unstacked}}(l)/k_{B}}O_{2}^{t}(a,b,l)\}$$
12

where ΔG_{stack} is the free energy of the closing stack formed by the base pairs (a, b) and (a - 1, b + 1)in Figure 4b, and $\Delta S_{\text{unstacked}}(l)$ is the entropy for a type-*t* loop of length *l* (see the central large loop in Figure 4b). $C_2(x, y)$ denotes the partition function of the closed conformations from nucleotide *x* to nucleotide *y*, $O_2^t(a, b, l)$ denotes the partition function of the type-t open conformations (t = L, M, R, LR) from nucleotide *a* to nucleotide *b* shown in Figure 4a.

Open conformations. For RNA secondary structures, $O_2^t(a, b, l)$ can be conveniently calculated recursively from the partition functions for shorter chains (42).

$$O_{2}^{L}(a, b, \pm l) = O_{2}^{L}(a, b-1, l-1) + O_{2}^{LR}(a, b-1, l) + C_{2}(a +1, b-2)$$
13

$$O_2^M(a,b,l) = O_2^M(a,b-1,l-1) + O_2^R(a,b-1,l)$$
14

$$O_2^R(a,b,l) = O_2^R(a+1,b,l-1) + O_2^{LR}(a+1,b,l) + C_2(a+2,b-1)$$
15

$$O_2^{LR}(a,b,l) = \sum_{a < x < b} C_2(x,b-1)$$

$$\times \{O_2^L(a,x,l-2) + O_2^{LR}(a,x,l-1)$$

$$+ C_2(a+1,x-1)\}$$
16

To illustrate the meaning of the above equations, as an example, in Figure 4e we show a diagrammatic illustration for Equation 14.

From Equations 12 and 13–16, the recursive relations for the partition functions for the closed and the open conformations are inter-related. The computation based on the above recursive relations can efficiently give the partition function for any chain segment from a to b:

 $C_2(a,b) =$ the partition function for the closed conformations : $O_2^t(a,b) = \sum_l O_2^t(a,b,l)$

= the partition function for the open conformations