Computing the partition function and sampling for saturated secondary structures of RNA, with respect to the Turner energy model

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Abstract

An RNA secondary structure is saturated if no base pairs can be added without violating the definition of secondary structure. Here we describe a new algorithm, RNAsat, which for a given RNA sequence $a$, an integral temperature $0 \leq T \leq 100$ in degrees Celsius, and for all integers $k$, computes the Boltzmann partition function $Z_T^k(a) = \sum_{S \in \mathcal{S}_{AT}(a)} \exp(-E(S)/RT)$, where the sum is over all saturated secondary structures of $a$ which have exactly $k$ base pairs, $R$ is the universal gas constant and $E(S)$ denotes the free energy with respect to the Turner nearest neighbor energy model. By dynamic programming, we compute $Z_T^k$ simultaneously for all values of $k$ in time $O(n^5)$ and space $O(n^3)$. Additionally, RNAsat computes the partition function $Q_T^k(a) = \sum_{S \in \mathcal{S}_T(a)} \exp(-E(S)/RT)$, where the sum is over all secondary structures of $a$ which have $k$ base pairs; the latter computation is performed simultaneously for all values of $k$ in $O(n^4)$ time and $O(n^3)$ space. Lastly, using the partition function $Z_T^k$ [resp. $Q_T^k$] with stochastic backtracking, RNAsat rigorously samples the collection of saturated secondary structures [resp. secondary structures] having $k$ base pairs; for $Q_T^k$ this provides a parametrized form of Sfold sampling [14]. Using RNAsat, (i) we compute the ensemble free energy for saturated secondary structures having $k$ base pairs, (ii) show cooperativity of the Turner model, (iii) demonstrate a temperature-dependent phase transition, (iv) illustrate the predictive advantage of RNAsat for precursor microRNA cel-mir-72 of C. elegans and for the pseudoknot PKB 00152 of Pseudobase [40], (v) illustrate the RNA shapes [18] of sampled secondary structures [resp. saturated structures] having exactly $k$ base pairs. A web server for RNAsat is under construction at bioinformatics.bc.edu/clotelab/RNAsat/
1 Introduction

In recent years, it has emerged that RNA plays a surprising and previously unsuspected role in many biological processes, including retranslation of the genetic code (selenocysteine insertion [5, 21], ribosomal frameshift [34]), post-transcriptional regulation via small interfering RNA and microRNA [39, 26], conformational switches [42], metabolite-sensing riboswitches which interact with small ligands and up- or down-regulate certain genes [3], small nucleolar RNAs which guide the methylation of specific ribosomal nucleotides [37], etc.

The function of structural RNA certainly depends on RNA tertiary structure, which in [2] has been shown to be largely determined by RNA secondary structure. The Turner nearest neighbor model [32, 46] consists of experimentally determined enthalpy and entropy values for stacked base pairs (both Watson-Crick and GU wobble pairs), as well as for hairpin loops, bulges and interior loops (arbitrary multi-loop free energies are approximated by an affine measure). Zuker’s algorithm [51] is a dynamic programming algorithm, which computes the minimum free energy (mfe) secondary structure of an input RNA nucleotide sequence, with respect to the nearest neighbor model. Though there are small discrepancies in certain energy parameters (e.g. treatment of coaxial stacking and dangles

1Dangles, either 5’ or 3’, are unpaired bases immediately adjacent to a paired base; dangles stack below or on top of existent stacked base pairs, hence contribute a stabilizing energy. At 37 degrees Celsius, a 3’ dangle can have an energy of up to approximately −2 kcal/mol. When computing minimum free energy structures and partition functions, dangles constitute an important factor.

Most current work on RNA secondary structure concerns the thermodynamic equilibrium minimum free energy structure (mfold, RNAfold, RNAstructure), or the low energy ensemble of structures (Sfold [13], RNAsubopt [45]), or multiple sequence/structure alignment (Foldalign [20], Dynalign [30]), or the general shape of RNA secondary structures (RNAshapes) [18, 38, 41], or applications and extensions of such software. One important application area concerns the development of noncoding RNA gene finders, such as RNAz [44, 43].

Considering molecular complexes, Dimitrov and Zuker [12] described how to efficiently compute a multi-species partition function for interactions between two species of nucleic acid sequences (e.g. between DNA and RNA). This led to the hybridization web server DINAMelt and software UNAFOLD [27]; see also [4, 35]. Computation of the partition function for molecular complexes allows one to compute concentrations of various hybridization products, clearly relevant in probe design for gene expression chips.

In contrast to much of the previously mentioned work, the current paper considers secondary structure energy landscape from a different point of view. We divide up the exponentially large collection of saturated [resp. all] secondary structures for a given RNA sequence into disjoint families $F_k$, each indexed by $k$. Choosing $F_k$ appropriately, we compute partition functions at temperature $T$ for the following: (i) $Z_k^T$, for saturated secondary
structures having \( k \) base pairs, (ii) \( Q_k^T \), for all secondary structures having \( k \) base pairs, (iii) \( \tilde{Z}_k^T \), for saturated secondary structures having \( k \) base pairs fewer than the maximum possible (i.e. Nussinov-Jacobson optimal), (iv) \( \tilde{Q}_k^T \), for all secondary structures having \( k \) base pairs fewer than the maximum possible (i.e. Nussinov-Jacobson optimal). For each of these cases, the partition function is computed with respect to the Turner energy model, and rigorous sampling is provided.

Our guiding motivation is to use the partition function and derived thermodynamic parameters (ensemble free energy, heat capacity, etc.) to better understand the nature and distribution of kinetic traps in the folding landscape of RNA. To this end, given a secondary structure \( S \) for an RNA sequence \( a_1, \ldots, a_n \), we say that \( S \) is locally optimal with respect to the Turner energy model, if \( E(S) \leq E(S') \), for all secondary structures \( S' \), obtained from \( S \) by either the removal or addition of one base pair.\(^2\) Unable currently to compute the partition function for secondary structures which are locally optimal with respect to the Turner energy model, we instead describe our software RNAsat, which computes the partition function for saturated secondary structures with respect to the Turner energy model. Computational experiments reported here with precursor microRNA cel-mir-72 of \( C. \) elegans and with pseudoknot PKB 00152 from Pseudobase \(^{40} \), suggest that saturated structures and locally optimal structures are quite distinct concepts, and that there are far fewer locally optimal structures than saturated structures. Nevertheless, the algorithms described are non-trivial, and provide a tool to better understand the folding landscape of RNA. The goal of the remaining paper is to explain the algorithms behind RNAsat; in future work, we will investigate additional applications. We begin by some definitions.

Given an RNA sequence \( a = a_1, \ldots, a_n \), a secondary structure is a well-balanced parenthesis expression with dots, where the nucleotides \( a_i, a_j \) form either Watson-Crick or GU wobble pairs in positions \( i, j \) corresponding to the well-balanced left and right parentheses. Formally, a secondary structure \( S \) on RNA sequence \( a_1, \ldots, a_n \) is defined to be a set of ordered pairs corresponding to base pair positions, which satisfies the following requirements.

1. **Watson-Crick or GU wobble pairs:** If \( (i, j) \) belongs to \( S \), then pair \( (a_i, a_j) \) must be one of the following canonical base pairs: (A,U), (U,A), (G,C), (C,G), (G,U), (U,G).

2. **Threshold requirement:** If \( (i, j) \) belongs to \( S \), then \( j - i > \theta \).

3. **Noneexistence of pseudoknots:** If \( (i, j) \) and \( (k, \ell) \) belong to \( S \), then it is not the case that \( i < k < j < \ell \).

4. **No base triples:** If \( (i, j) \) and \( (i, k) \) belong to \( S \), then \( j = k \); if \( (i, j) \) and \( (k, j) \) belong to \( S \), then \( i = k \).

In this paper, following convention, the threshold \( \theta \), or minimum number of unpaired bases in a hairpin loop, is taken to be 3. For any additional background on RNA and dynamic

\(^2\)Here \( E(S) \) is the energy of \( S \) using the Turner energy model \(^{32} \), \(^{46} \). In the Nussinov-Jacobson energy model, the notion of locally optimal and saturated coincide, and the corresponding problem was solved in \(^{6} \), \(^{7} \).
programming computation of secondary structures, the reader can consult the text [9] and the recent review [15].

In [49], M. Zuker defined a saturated structure to be a secondary structure in which no base pairs can be added without violating the definition of secondary structure (e.g. without introducing a pseudoknot). More recently, Evers and Giegerich [16] defined a saturated structure to be a secondary structure, such that no additional base pairs can be stacked and for which there are no isolated base pairs. For this notion, Evers and Giegerich described an algorithm which computes the total number of saturated structures – saturated in the sense of [16]. It is not difficult to construct examples of structures which are saturated in the sense of Zuker, but not in the sense of Evers-Giegerich, and vice-versa – i.e. these notions are distinct; see [8] for explicit examples.

A related notion is that of locally optimal secondary structure. To define this notion, we first define base pair distance, a metric between two secondary structures for the same RNA sequence. If \( S, T \) are secondary structures for an RNA sequence \( a_1, \ldots, a_n \), then \( d(S, T) \) is the number of elements in the symmetric difference of \( S, T \); i.e. the number of base pairs in \( S \), but not \( T \), or vice-versa. Structure \( T \) is a neighbor of \( S \) if the base pair distance between \( S, T \) is 1.

Given a fixed energy model, such as the base pair model of Nussinov-Jacobson [36] or the nearest neighbor model of Turner [32, 46], a locally optimal secondary structure is a structure \( S \), such that no neighbor \( T \) of \( S \) has lower energy. In [6, 7] we developed an algorithm and web server to compute the number of locally optimal secondary structures with respect to the Nussinov-Jacobson energy model; for this energy model, local optimality coincides with Zuker’s notion of saturated structure.

Locally optimal structures correspond to potential kinetic traps in the folding process. Indeed, if \( S \) is a locally optimal secondary structure, which differs from the minimum free energy (mfe) structure \( S_0 \), then an energetically unfavorable neighbor \( S' \) of \( S \) must appear in the folding path from \( S \) to mfe structure \( S_0 \). The barrier tree of Flamm et al. [17] illustrates this situation.

In this paper, we lift the algorithm of [6] to the Turner energy model; specifically we proceed as follows. Given an RNA nucleotide sequence \( a = a_1, \ldots, a_n \), for each value of \( k \), we compute the Boltzmann partition function \( Z^T_k(a) = \sum_{S \in \mathcal{SAT}_k(a)} e^{E(S)/RT} \), where \( \mathcal{SAT}_k(a) \) designates the collection of all saturated secondary structures \( S \) of \( a \) having exactly \( k \) base pairs, \( R \) is the universal gas constant with value 1.98717 cal/mol per degree Kelvin, \( 0 \leq T \leq 100 \) is temperature in degrees Celsius, and where \( E(S) \) denotes the free energy, using the Turner nearest neighbor energy model [32, 46]. By dynamic programming, we compute \( Z^T_k \) simultaneously for fixed \( T \), for all values of \( k \) in time \( O(n^5) \) and space \( O(n^3) \). Although the notion of saturated structure (in the sense of Zuker) is distinct from that of locally optimal structure within the Turner nearest neighbor energy model, the work of this paper is a step towards the goal of computing the partition function for all locally optimal secondary structures.
Our software, \textsc{RNAsat}, additionally computes the full partition function

$$Q^T(a) = \sum_{S \in \mathcal{S}(a)} \exp(-E(S)/RT)$$

and its parametrized form $Q^T_k(a) = \sum_{S \in \mathcal{S}_k(a)} \exp(-E(S)/RT)$, where \(\mathcal{S}(a)\) denotes the collection of all secondary structures of RNA sequence \(a\), and \(\mathcal{S}_k(a)\) designates the collection of all secondary structures \(S\) of \(a\) which have exactly \(k\) base pairs. The \(O(n^3)\) time and \(O(n^2)\) space algorithm to compute the partition function $Q^T(a)$ was first developed by McCaskill [33]. McCaskill’s algorithm has additionally been implemented by Hofacker et al. in the Vienna RNA Package [24], by D. Mathews [29], and by N. Markham and M. Zuker [28]. However, unlike these authors, we additionally compute in time \(O(n^4)\) and space \(O(n^3)\) the partition function $Q^T_k(a)$ for all secondary structures having exactly \(k\) base pairs, for all values of \(1 \leq k \leq n\). We omit details, since the method is similar, though simpler, to the method we present to compute the partition function of saturated structures having \(k\) base pairs.

\textsc{RNAsat} can sample from the collection of saturated secondary structures having \(k\) base pairs, using stochastic backtracking with $Z^T_k(a)$. Additionally, \textsc{RNAsat} can sample from the collection of all secondary structures having \(k\) base pairs, using stochastic backtracking with $Q^T_k(a)$. Our sampling procedure is mathematically rigorous, and is analogous to the method of Ding and Lawrence [14] with web server \textsc{Sfold} [13], except that we sample with respect to $Z^T_k(a)$ and $Q^T_k(a)$ for any temperature $0 \leq T \leq 100$, whereas Ding and Lawrence use McCaskill’s partition function $Q(a)$ to sample all secondary structures at 37 degrees Celsius. The algorithms of \textsc{RNAsat} to compute $Z^T_k(a)$ and sample saturated secondary structures having \(k\) base pairs are quite distinct from any existing algorithms. As well, despite the close relation of $Q^T_k(a)$ to $Q^T(a)$, to the best of our knowledge, no other software supports the computation of $Q^T_k(a)$ or samples, for given \(k\), from the collection of secondary structures having \(k\) base pairs.

Using partition functions, we develop the following applications. (i) We compute ensemble free energies for saturated secondary structures having \(k\) base pairs, thus settling the question of thermodynamic stability of saturated structures. (ii) We demonstrate that the Turner nearest neighbor energy model [46] leads to cooperative folding, in the sense of [11]. (iii) We compute and graph the relative density of states for saturated secondary structures having \(k\) base pairs as a function of temperature; these curves resemble temperature-dependent phase transitions. (iv) We illustrate the predictive advantage of \textsc{RNAsat} for precursor microRNA cel-mir-72 of \textit{C. elegans} and for pseudoknot PKB 00152 of Pseudobase [40]. (v) We compute frequencies of RNA shapes [18] of sampled secondary structures [resp. saturated structures] having exactly \(k\) base pairs.

The plan of the paper is as follows. Section 2 presents key definitions used in our dynamic programming algorithm for saturated secondary structures. Additionally, this section describes several applications of \textsc{RNAsat}. Section 3 presents the pseudocode and an explanation of how to compute the Boltzmann partition function $Z^T_k(a)$ for all saturated secondary structures having exactly \(k\) base pairs. Section 4 gives pseudocode for the algorithm to
sample, for each \( k \), from the ensemble of saturated secondary structures having \( k \) base pairs. The sampling is rigorously performed according to the Boltzmann probability; i.e. lower energy structures in the ensemble \( SAT_k(a) \) of saturated secondary structures of \( a \) having \( k \) base pairs are more heavily weighted than higher energy structures. Since sampling from the ensemble of saturated secondary structures is different from the procedure of [13], we include a proof of correctness of our algorithm. Although our software RNAsat as well as the pseudocode for sampling given in Figures 12, 13 and 14 is for the Turner energy model [46], the proof of correctness is given for the simpler Nussinov-Jacobson energy model, in the interests of readability.

Finally, we note that in this paper, when the temperature \( T \) is clear from context, we may write \( Z_k, Z \) [resp. \( Q_k, Z \)] in place of \( Z_k^T, Z^T \) [resp. \( Q_k^T, Q^T \)], etc.

### 2 Overview and applications

In [36], Nussinov and Jacobson introduced a simple energy model, where each base pair in a secondary structure \( S \) contributes an energy of \(-1.4\). In this model, known as the Nussinov-Jacobson energy model, the energy \( E(S) \) of secondary structure \( S \) equals \(-|S|\), where \(|S|\) denotes the number of base pairs in \( S \). In [36] a dynamic programming algorithm was given to compute the optimal secondary structure for a given RNA sequence; i.e. the optimal secondary structure with respect to the Nussinov-Jacobson energy model is that structure having the maximum number of base pairs (see [9] for implementation details).

Much earlier, a realistic nearest neighbor energy model was developed by Tinoco [23], with experimentally measured stacking free energies; i.e. energy contributions come from stacked base pairs rather than base pairs. The well-known Zuker algorithm [51] computes the minimum free energy (mfe) over all secondary structures by adding contributions of negative (stabilizing) energy terms for stacked base pairs and positive (destabilizing) energy terms for hairpin loops, bulges, internal loops and multiloops. Initially implemented with energy parameters from [23], successive refinements of Zuker’s algorithm have incorporated more accurate energy parameters from [32] with refinements from [46] for stacked base pairs and various loops (hairpin, bulge, internal loop, multi-branch loop) and 5’ and 3’ dangles, i.e. unpaired nucleotides which stack onto a base pair. The energy term contributed by a base pair depends on the base pair (if any) upon which it is stacks; for instance, Turner’s current rules [46] at 37 degrees Celsius assign stacking free energy of \(-2.24 \text{ kcal/mol}\) to \(5'\text{-AC}-3'\) of \(-3.26 \text{ kcal/mol}\) to \(5'\text{-CC}-3'\) and of \(-2.08 \text{ kcal/mol}\) to \(5'\text{-AG}-3'\) of \(3'\text{-GG}-5'\) and of \(3'\text{-UC}-5'\). Our software RNAsat uses

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3 The reader familiar with the Turner energy model should have no difficulty in constructing a similar argument for Proposition 1.

4 Of course, this model has trivial variants, such as assigning \(-3\) for GC, \(-2\) for AU and \(-1\) for GU pairs. In particular, Proposition 1 uses the notation \( bp(i, j) \) for the energy of base pair \((i, j)\) in the RNA sequence \(a_1, \ldots, a_n\).

5 This algorithm, known as the Nussinov-Jacobson algorithm, is also called the maximum circular matching algorithm [22].
free-energy values for stacked base pairs, dangles, hairpins, bulges, internal loops and multi-
loops from \textit{mfold} 3.0 when $T = 37^\circ$ Celsius and energy values from \textit{mfold} 2.3 otherwise. In particular, following the approach of Zuker’s \textit{mfold}, when loop size $N$ exceeds 30, the free energy, due to loss of entropy, is $c_T \cdot \ln(N/30)$, where $c_T$ is a temperature-dependent constant; e.g. $c_{37} = 1.079$.

Following [6], given an RNA sequence $a = a_1, \ldots, a_n$, a secondary structure $S$ for $a$ is defined to be $k$-saturated, if $S$ is locally optimal, and $S$ contains $k$ base pairs fewer than the maximum for $a$; i.e. $k$ fewer base pairs than that of the Nussinov-Jacobson optimal structure. If $m$ denotes the number of base pairs in the Nussinov-Jacobson optimal structure for RNA sequence $a$, then clearly any $k$-saturated secondary structure for $a$ has $m-k$ many base pairs, and any saturated secondary structure $S$ on $a$ having $k$ base pairs must be $(m-k)$-saturated.

In [6, 7], we developed a dynamic programming algorithm called \texttt{RNALOSS}, an acronym for RNA locally optimal secondary structures, computed with respect to the Nussinov-Jacobson energy model. This software computes, for a given RNA nucleotide sequence and each integer $k$, the number of $k$-saturated secondary structures. Since the Boltzmann partition function is essentially a weighted count, the algorithm of [6] can be used to compute, for given RNA sequence $a$ and all integers $k$, the Boltzmann partition function with respect to the Nussinov-Jacobson energy function for all saturated secondary structures on $a$ having $k$ base pairs.

In this paper, we lift the algorithm \texttt{RNALOSS} [6, 7] to an $O(n^5)$ time and $O(n^3)$ space algorithm \texttt{RNAsat} to compute, for a given RNA sequence $a$ and all integers $k$, the Boltzmann partition function $Z^T_k(a)$ with respect to the Turner energy model, for any integral temperature $T$ in degrees Celsius, $0 \leq T \leq 100$.

### 2.1 Visible nucleotides and visible positions

To fix ideas, consider the toy sequence \texttt{GGGGCCCCC}. The maximum number of base pairs is 3, and there is one 0-saturated structure (with 3 base pairs) given by $(((...)))$, twelve 1-saturated structures given by

\[
\begin{align*}
\text{GGGGCCCCC} & \quad \text{GGGGCCCCC} & \quad \text{GGGGCCCCC} & \quad \text{GGGGCCCCC} \\
\ldots(\ldots) & \quad \ldots(\ldots) & \quad \ldots(\ldots) & \quad \ldots(\ldots) \\
(\ldots) & \quad (\ldots) & \quad (\ldots) & \quad (\ldots) \\
((\ldots)) & \quad ((\ldots)) & \quad ((\ldots)) & \quad ((\ldots))
\end{align*}
\]

and three 2-saturated secondary structures, given by

\[
\begin{align*}
\text{GGGGCCCCC} \\
(\ldots) & \quad (\ldots) & \quad (\ldots) & \quad (\ldots) & \quad (\ldots)
\end{align*}
\]

Note that the Nussinov-Jacobson algorithm predicts an optimal secondary structure of $(((\ldots)))$ with three base pairs, whereas both \textit{mfold} and \textit{RNAfold} predict a minimum
free energy structure \((\ldots)\) of \(-1.40\) kcal/mol with one stacked base pair, one exterior base pair, and dangles.

With over 12 trillion saturated structures for the 54 nucleotide hammerhead ribozyme with EMBL accession number AF170517 (Rfam family RF00008 [19]), it is impossible in general to enumerate all saturated structures. Instead, we apply dynamic programming using the notions of visible nucleotides and visible positions defined in [6]. Given RNA sequence \(a = a_1, \ldots, a_n\), a position \(1 \leq i \leq n\) is visible in secondary structure \(S\) if for all base pairs \((x, y) \in S\), it is not the case that \(x \leq i \leq y\). A nucleotide \(X \in \{A, C, G, U\}\) is visible if \(X = a_i\), for some visible position \(i\). Define \(\text{VisNuc}(S) \subseteq \{A, C, G, U\}\) to be the set of visible nucleotides occurring in some position \(i < n - \theta\), i.e.

\[
\text{VisNuc}(S) = \{a_i \mid \text{ for all } (x, y) \in S \text{ where } (z < x \text{ or } z > y \text{ and } z < n - \theta)\}.
\]

To account for visible positions between \(n - \theta, \ldots, n\), we define \(\text{VisPos}(S) = b\), where \(0 \leq b \leq \theta + 1\), and \(b\) is the greatest value in \(0, \ldots, \theta + 1\), such that for all \(0 \leq x < b\), \(n - x\) is external to every base pair of \(S\). We say that structure \(S\) is \(s, b\)-visible if \(\text{VisNuc}(S) = s\) and \(\text{VisPos}(S, 1, n) = b\). For example, if \(a\) is the sequence AUUCCGGCA and if the minimum number of unpaired bases in a hairpin loop is \(\theta = 1\), then the four secondary structures

\[
\begin{align*}
\text{AUUCCGGCA} & \quad \text{AUUCCGGCA} \\
(\ldots)(\ldots) & \quad \ldots(\ldots) \\
\text{AUUCCGGCA} & \quad \text{AUUCCGGCA} \\
(\ldots)(\ldots) & \quad \ldots(\ldots)
\end{align*}
\]

are respectively \(\{G\}\), 2-visible, \(\{A, G, U\}\), 2-visible, \(\{C\}\), 1-visible and \(\{A, U\}\), 0-visible. For given RNA sequence \(a = a_1, \ldots, a_n\), for fixed temperature \(T\), for all integers \(k\), for all \(2^4 = 16\) possible sets \(s\) of visible nucleotides and for all possible values \(0 \leq b \leq \theta + 1 = 4\) of visible positions, our main algorithm, given in Figures 10 and 11, computes the partition function \(Z^T_k(a, i, j; s, b)\) where \(\text{SAT}_k(a, s, b)\) denotes the set of saturated secondary structures on \(a\) which are \(s, b\)-visible and have exactly \(k\) base pairs. The definitions of \(\text{VisNuc}\) and \(\text{VisPos}\) are precisely what is needed to compute \(Z^T_k(a, i, j; s, b)\), assuming that all values of \(Z^T_k(a, i', j', s', b')\) have been computed and stored in a table for \(|j' - i'| < |j - i|\). Using dynamic programming, for all integers \(k\), we compute as well the partition function

\[
Q^T_k(a, i, j) = \sum_{S \in \text{Sat}_k(a)} \exp(-E(S)/RT)
\]

where \(\text{Sat}_k(a)\) denotes the set of all secondary structures \(S\) having exactly \(k\) base pairs (i.e. in \(Q^T_k\), secondary structure \(S\) need not be saturated).

In [14], Ding and Lawrence describe how to sample secondary structures in a mathematically rigorous fashion, by using stochastic backtracking with the partition function as

\[
Q^T_k(a, i, j) = \sum_{S \in \text{Sat}_k(a)} \exp(-E(S)/RT)
\]

where \(\text{Sat}_k(a)\) denotes the set of all secondary structures \(S\) having exactly \(k\) base pairs (i.e. in \(Q^T_k\), secondary structure \(S\) need not be saturated).

\[\text{In this paper, as well as in mfold, RNAfold and RNAstructure, } \theta \text{ is taken to be 3. We have temporarily taken } \theta = 1 \text{ for perspicacity in this toy example.}\]
computed by McCaskill’s algorithm [33]. Though details are substantially more complicated, we have implemented a sampling algorithm for the set of saturated secondary structures having exactly $k$ base pairs; see Figures 12, 13 and 14. Though not described here, using our computation of $Q^T_k(a)$, our software RNAsat includes a rigorous sampling as well for the collection of all secondary structures having $k$ base pairs; i.e., we provide a parametrized version of Sfold [13]. In this fashion, RNAsat can additionally sample all [resp. saturated] secondary structures irrespective of number of base pairs.

2.2 Applications

For an input RNA sequence, fixed temperature $T$ and fixed value of $k$, we can generate (say) 1000 samples of saturated secondary structures, each having exactly $k$ base pairs and sampled using the Boltzmann partition function $Z^T_k$. Subsequent frequency analysis of the samples, when $T$ and $k$ are varied, allow us to investigate aspects of the ensemble of low energy saturated structures. In this section, we present (i) ensemble free energies for saturated secondary structures having $k$ base pairs, (ii) a demonstration of cooperative folding in the Turner energy model, (iii) a temperature-dependent phase transition for saturated structures, (iv) additional experiments including precursor microRNA cel-mir-72 from C. elegans and pseudoknot PKB 00152 from Pseudobase [40], (v) a frequency analysis of RNA shapes at various temperatures for all [resp. saturated] secondary structures of a 149 nt. SAM riboswitch having $k$ base pairs fewer than that of the Nussinov-Jacobson optimal structure (see [18] for the definition of RNA shape).

Ensemble free energy

Figure 1 depicts the ensemble free energy $G_k = -RT \ln Z^T_k$, where the universal gas constant $R$ is Avogadro’s number times the Boltzmann constant and $T$ is temperature in degrees Kelvin. Note the thermodynamic stability of saturated secondary structures, leading support to the suggestion that saturated structures form a potential kinetic trap in the folding process.

Cooperative folding

In [48] Zimm and Bragg introduce a nearest neighbor model for helix-coil transition of a protein, i.e. where the probability that a residue is in the helix (resp. coil) state is conditional on the state of its nearest neighbor. As indicated by the sigmoidal curves of the number of residues in coil state as a function of temperature, the Zimm-Bragg model illustrates cooperativity in the helix to coil transition of a polypeptide. (See Chapter 26 of [11] for a good treatment of cooperativity and the Zimm-Bragg and Ising models.) Figure 2 shows that the Turner model gives rise to a sigmoidal curve, typical of cooperative folding, while the Nussinov-Jacobson model does not. Assuming that the average number of base pairs in
the ensemble of secondary structures is inversely proportional to the UV absorbance in a spectrophotometer, notice the similarity between the Turner model curve in Figure 2 with the experimentally determined RNA melting curve of [25].

Apparent phase transition

Define the relative density of states $\rho_k^T$ for saturated secondary structures of a given RNA sequence having exactly $k$ base pairs at temperature $T$ to be the relative Boltzmann probability, i.e. $\rho_k^T = Z^T$, for each integer $k$. Since secondary structure $S$ for a $k$ is $k$-saturated exactly when $|S| = m - k$, where $m$ denotes the maximum number of base pairs in a secondary structure on $a$, this immediately allows us to compute the relative density of states $\tilde{\rho}_k^T = \tilde{Z}_k^T / Z^T$ for $k$-saturated secondary structures. Figure 3 displays overlaid graphs, for each fixed $k$, of the relative density of states for $k$-saturated structures as a function of temperature. The 0-saturated density is close to 1 at 0 degrees Celsius, gradually declines with rising temperature. With rising temperature, the 1-saturated density gradually increases to a peak, then diminishes, etc.

It should be mentioned that Figures 2 and 3 of the current paper are similar in form, although distinct, to Figure 2 of [12] and Figure 1 of [28]. Figure 2 of the current paper depicts the expected number of base pairs of a single RNA molecule as a function of temperature, while one of the curves in Figure 2 of [12] depicts the fraction of random coil as a function of temperature. Figure 3 of the current paper depicts the relative density of states of $k$-saturated secondary structures, for small values of $k$, as a function of temperature. Though unrelated, Figure 2 of [12] depicts the fraction of various species (duplex, hairpin, random coil) as a function of temperature – both graphs depict rising and falling concentrations as a function of temperature.

Experiments with microRNA cel-mir-72 and pseudoknot PKB 00152

In this section, we describe several additional experiments. Figures 4 and 5 illustrate the difference between the minimum free energy (mfe) secondary structure for 96 nt. precursor microRNA cel-mir-72, from C. elegans, with accession number MI0000043 from miRBase linked to Rfam [19]. The left panel of Figure 4 depicts the mfe structure, as computed with RNAfold from Vienna RNA Package 1.5. The right panel of this figure displays the consensus structure, as determined by 1000 saturated structures sampled using RNAsat. The consensus structure, which by definition consist of the base pairs occurring in strictly

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7Absorbance of UV light at 260 nm is due to especially pyrimidines when light impinges vertically to the plane of the base. Base pairing reduces likelihood of vertical impingement, hence the more base pairing that occurs in a secondary structure, the less UV absorbance will be detected. See [47] for more on UV absorbance.

8This expected number could alternatively be obtained by summing over all base pairs the Boltzmann probability of base-pairing, as computed by McCaskill’s algorithm – remark due to M. Zuker (personal communication).

9The consensus structure consists of those base pairs which appear in strictly greater than half the sampled structures.
more than half the sampled structures, clearly resembles the familiar approximate stem-
loop structure which precursor microRNAs are believed to adopt. Figure 5 displays dotplot
presentations of Boltzmann pair frequencies; in the upper triangular portion, square are
depicted in the position \((i,j)\) with area proportional to frequency that base pair \((i,j)\) occurs
in the sampled ensemble. In the lower triangular portion, the minimum free energy structure
is represented, in that a black square appears in position \((j,i)\) if base pair \((i,j)\) occurs in the
mfe structure, as computed using version 1.5 of RNAfold. The left panel of Figure 5 depicts
the base pair frequencies from 1000 sampled secondary structures, using the implementation
of McCaskill’s algorithm [33] in RNAsat; since RNAsat computes \(Q_T^k\) for each \(k\), by sampling,
we first compute the Boltzmann probability that a secondary structure has \(k\) base pairs, then
we sample accordingly. The middle panel depicts the dotplot of base pair frequencies from
1000 samples of approximately locally optimal secondary structures, produced by a greedy
algorithm suggested by one of the referees. The greedy algorithm first samples a low energy
structure using McCaskill’s algorithm, and subsequently adds that base pair which can be
added which would lead to the greatest decrease in energy. Clearly the dotplot in the left
and middle panels are similar. In contrast, the right panel depicts base pair frequencies from
1000 sampled saturated secondary structures. It follows that approximately locally optimal
structures, as determined by the greedy algorithm, resemble the structures sampled by the
method of Ding and Lawrence [14]. These appear to be distinct from sampled saturated
structures.

In the set of experiments displayed in Figures 6, 7, 8 and Table 3, we considered the
26 nt. pseudoknot PK2 of the upstream pseudoknot domain (UPD) of the 3’ UTR of RNA
beta, with EMBL accession number X03854 and start position 3088. This data is taken from
the Pseudobase database [40] with accession number PKB 00152. The RNA sequence and
secondary structure, with annotated pseudoknot, is as follows:

\[
UGGUGC\text{CAUCA}A\text{C}A\text{U}A\text{U}A\text{GUG}GGA
\]

Figure 6 displays the minimum free energy (mfe) secondary structure and corresponding dot
plot of Boltzmann pair probabilities, as computed using version 1.5 of Vienna RNA Package
RNAfold. RNAfold correctly detects one of the helices, computed to have energy of \(-10.5\)
kcal/mol, as illustrated as follows:

\[
UGGUGC\text{CAUCA}A\text{C}A\text{U}A\text{U}A\text{GUG}GGA
\]

The left panel of Figure 7 displays the dot plot of pair frequencies, computed by sampling
1000 structures using RNAsat. For each \(k\), RNAsat computes the Boltzmann probability
\(Q_T^k / \sum_k Q_T^k\) of all secondary structures having \(k\) base pairs, and then samples
appropriately. The same dot plot is produced by Sfold [14] (data not shown). Note that
this dot plot is identical to the right panel of Figure 5. The right panel of Figure 7 displays
the dot plot of pair frequencies computed by sampling 1000 locally optimal structures.
All locally optimal structures were computed by brute force, by first enumerating all secondary structures by using Vienna RNA Package RNAsubopt, and subsequently checking whether the structure is locally optimal with respect to the Turner energy model; i.e. no neighbor has lower energy. This produces a set LocOpt of all 780 locally optimal structures for PKB 00152. Note by contrast that there are 111,014 saturated structures of PKB 00152. We then sample 1000 locally optimal structures from LocOpt, each with probability \( Pr[S] = \exp(-E(S)/RT)/\sum_{S} \exp(-E(S)/RT) \), where the sum is taken over LocOpt. Among these 1000 sampled structures, with few exceptions, the mfe structure occurs. In contrast, Table 3 presents data on the frequency of occurrence of saturated structures, when 1000 saturated structures are sampled from the collection of 111,014 saturated structures of PKB 00152.

The left panel of Figure 7 displays the dot plot of 1000 saturated structures sampled by RNAsat. Notice how alternative base pairs appear, corresponding to the pseudoknot, a feature not detected using mfe calculation, McCaskill’s algorithm, or sampling over all secondary structures. The right panel of Figure 7 presents the histogram of energies of all sampled saturated structures.

**Shape frequencies of saturated and locally optimal structures**

In Tables 1 and 2, we present shape frequencies for sampled structures of the SAM riboswitch with EMBL accession number AL939119.1/177986-178133. As defined by Giegerich and co-workers [18, 38, 41], the shape of a given secondary structure is a homomorphic mapping from the secondary structure into a more compact structure, which succinctly describes basic features of the secondary structure; see [18] for the formal definition. For instance, the \( \pi \)-shape of the cloverleaf secondary structure of tRNA is \([\Box \Box \Box \Box]\), while the less succinct \( \pi' \)-shape is \([\Box \Box \Box \Box \Box \Box \Box \Box \Box \Box] \). By a simple linear time algorithm not described here, we can compute the \( \pi \)- and \( \pi' \)-shape of a given secondary structure; for reasons of space, we only present \( \pi \)-shapes in the data shown. Notice how the frequency of certain shapes for the SAM riboswitch depends on temperature (25, 37, 50 degrees Celsius), and on the number \( k \) of base pairs fewer than that of the Nussinov-Jacobson optimal structure for this riboswitch, which has 55 base pairs. Note the quite different shape frequencies of saturated structures versus general structures sampled by RNAsat, as displayed in Tables 1 and 2. Sample data is presented for different temperatures (25, 37 and 50 degrees Celsius) and for different values of \( k \), where \( k \) designates the number of base pairs fewer than that of the Nussinov-Jacobson optimal structure (in the case at hand, this number is 55). By ‘gen’, we mean a general sampling, i.e. without any restriction on the value of \( k \).

Collectively, the data for these examples suggest that there may in general be far fewer locally optimal structures than saturated structures, and that the ensemble of locally optimal structures resembles the Boltzmann ensemble of low energy structures, both quite different from the ensemble of saturated structures.
3 Computing the Boltzmann partition function $Z^T_k$

3.1 Recurrence relations for McCaskill’s algorithm

In [33], McCaskill described a cubic time algorithm to compute the partition function $Q = \sum S e^{-\beta E(S)}$, where the sum is over all secondary structures of a given RNA sequence $a_1, \ldots, a_n$. (Recall that we ‘Z’ to denote the partition function for saturated structures, while ‘Q’ denotes the partition function without the restriction to saturated structures.) In particular, McCaskill defined

$$Q_{i;j} = \sum S e^{-\beta E(S)}$$

where the sum in equation (1) is taken over all secondary structures $S$ of $a[i, j] = a_i, \ldots, a_j$.

Following [24], let $Q^B_{i;j}$ denote the partition function restricted to those secondary structures on $a[i, j]$ in which $(i, j)$ is a base pair. Similarly, $Q^M_{i;j}$ [resp. $Q^{M1}_{i;j}$] denotes the partition function for those secondary structures on $a[i, j]$ in which $a[i, j]$ is part of a multiloop having one or more [resp. exactly one] helix. With this notation, the recurrence relations for McCaskill’s algorithm are given as follows.

$$Q_{i;j} = Q_{i;j-1} + \sum_{i<x<j} Q^B_{i,x} Q^B_{x+1,j}$$
$$Q^B_{i;j} = e^{-\beta H(i,j)} + \sum_{i<x<y<j} Q^B_{x,y} e^{-\beta I(i,j,x,y)} + \sum_{i<x<j} Q^M_{i+1,x} Q^{M1}_{x+1,j-1} e^{-\beta a}$$
$$Q^M_{i;j} = \sum_{i<x<j} e^{-\beta(x-i-1)c} Q^M_{x+1,j} + \sum_{i<x<j} Q^M_{i,x} Q^B_{x+1,j} e^{-\beta b} + Q^M_{i,j-1} e^{-\beta c}$$
$$Q^{M1}_{i;j} = Q^{M1}_{i,j-1} e^{\beta c} + Q^B_{i,j} e^{\beta b}$$
$$Q_{i,i} = 1, Q^B_{i,i} = 0, Q^M_{i,i} = 0, Q^{M1}_{i,i} = 0$$

Here, $H(i, j)$ is the free energy of a hairpin loop closed by base pair $(i, j)$, and $I(i, j, x, y)$ is the free energy of an interior loop between base pair $(i, j)$ and base pair $(x, y)$. The free energy of a multiloop having $B$ base pairs and $U$ unpaired bases is approximated by the affine function $a + bB + cU$.

It is now straightforward to modify these recursion equations, in order to compute $Q_{i;j}(k)$, the partition function for all secondary structures on $a[i, j]$ having exactly $k$ base pairs. We
have the following.

\[ Q_{i,j}(k) = Q_{i,j-1}(k) + \sum_{1 \leq \ell < k} Q_{i,x}^Z(\ell)Q_{x+1,j}(k-\ell) \]
\[ Q_{i,j}^B(1) = e^{-\beta H(i,j)} \]
\[ Q_{i,j}^B(k + 1) = \sum_{i < x < y < j} Q_{x,y}^B(k)e^{-\beta H(i,j,x,y)} + \]
\[ \sum_{1 \leq \ell < k} \sum_{i < x < j} Q_{i+1,x}^M(\ell)Q_{i,j-1}^M(k + 1 - \ell)e^{-\beta a} \]
\[ Q_{i,j}^M(k) = \sum_{i < x < j} e^{-\beta(x-i-1)}Q_{i,x}^M(\ell)Q_{x+1,j}^M(k - \ell)e^{-\beta b} + \]
\[ Q_{i,j-1}^M(k)e^{-\beta c} \]
\[ Q_{i,j}^{M1}(k) = Q_{i,j-1}^{M1}(k)e^{\beta c} + Q_{i,j}^B(k)e^{\beta b} \]
\[ Q_{i,i}(k) = 1, Q_{i,i}^B(k) = 0, Q_{i,i}^M(k) = 0, Q_{i,i}^{M1}(k) = 0 \]

The difficulty of adding visibility predicates \( s, b \), necessary to compute the partition function \( Z_{i,j}(k) \) for saturated structures on \( a[i,j] \) having exactly \( k \) base pairs can be illustrated in an attempt to define the multiloop partition function \( Z_{i,j}^M(k, s, b) \) for \( k \) base pairs and visibility predicates \( s, b \). The term

\[ \sum_{i < x < j} e^{-\beta(x-i-1)}e^{Z_{i,j+1}^M(k, s, b)} \]

contributes to the partition function \( Z_{i,j}^M(k, s \cup \{a_{i+1}, \ldots, a_x\}, b) \), but there are other terms making it difficult to express how \( Z_{i,j}^M(k, s, b) \) depends on previously computed terms \( Z_{i,x}^M(\ell, s_0, b_0) \), \( Z_{x+1,j}^B(k - \ell, s_1, b_1) \), etc. For this reason, after giving an overview of \texttt{RNASat} we present the pseudocode in Figures 10, 11, 12, 13, 14.

### 3.2 Notation

Given an RNA sequence \( a_1, \ldots, a_n \) and fixed absolute temperature \( T \), Boltzmann partition function values are computed using dynamic programming, where (essentially) two-dimensional arrays \( Z_s, Z'_s, Z'_m, Z_m, Z_e \) are progressively filled along off-diagonals at increasing distance \( d = 1, 2, \ldots \) from the principal diagonal. In this manner, when defining the value \( Z[i,j] \), for \( 1 \leq i < j \leq n \), where \( Z \) collectively denotes any of \( Z_s, Z'_s, Z'_m, Z_m, Z_e \), the values of \( Z[i',j'] \) are already stored for all \( 1 \leq i' < j' \leq n \) such that \( (j' - i') < (j - i) \).

As described in Section 2.1, additional visibility parameters \( s, b \) are required to inductively account for all saturated structures. Each of \( s, b \) ranges over finitely many values; indeed \( s \subseteq \{A,C,G,U\} \) and \( 0 \leq b \leq \theta + 1 = 4 \). It follows that each of the arrays \( Z_s, Z'_s, Z'_m, Z_m, Z_e \) is not two-dimensional as initially described, but rather of the form \( Z[k, i, j, s, b] \).

Note that in the particular case \( s = \emptyset \) and \( b = 0 \), the secondary structures \( S \) considered in \( Z_s(k, i, j, \emptyset, 0) \) are such that \((i, j) \in S\); i.e. \( i \) and \( j \) base-pair together in \( S \).

The pseudocode description of the partition function algorithm in Figures 10 and 11 requires some explanation of notation used. The function \( \text{maxBP}(i,j) \) returns the maximum number of base pairs possible in a secondary structure on \( a_i, \ldots, a_j \), i.e. equal to the number
of base pairs in the Nussinov-Jacobson optimal structure on \( a_i, \ldots, a_j \). The function \( \text{nucl}(i, j) \) returns the set \( \{a_i, \ldots, a_j\} \); note that \( \text{nucl}(i, j) \subseteq \{A, C, G, U\} \) is a set, not a multi-set or list. The function, \( \text{basepairWithSet}(s, i) \) returns \( \text{true} \) if the nucleotide \( a_i \) at position \( i \) can base-pair with a nucleotide in the set \( s \), and \( \text{false} \) otherwise. This function is invoked when \( s \) is a set of visible nucleotides (i.e. external to any base pair).

Energy contributions for different types of loops (helix, bulge, interior loop and multi-loop) are defined as follows. For \( 1 \leq i < j \leq n \), \( \text{hairpin}(i, j) \) denotes the energy contribution of a hairpin loop on RNA subsequence \( a_i, \ldots, a_j \); i.e. \( a_i \) and \( a_j \) form a closing base pair of the loop region \( a_{i+1}, \ldots, a_{j-1} \), where the latter are not base-paired. The energy contribution for a stacked base pair, bulge or interior loop is denoted by \( \text{stack}(i_1, i_2, j_1, j_2) \), where \( (i_1, j_2) \) is the exterior closing base pair, and \( (i_2, j_1) \) is the interior closing base pair. Finally, the energy associated with a multi-loop is computed by the affine function \( \alpha_1 + i \cdot \alpha_2 + n \cdot \alpha_3 \), where \( i \) is the number of base pairs in the multi-loop and \( n \) the number of unpaired nucleotides. Both 5’ and 3’ dangles are taken into account by our algorithm, and we denote the energy contribution by \( \text{dangle}(i, j) \). Specifically, if \( i < j \), then the function \( \text{dangle}(i, j) \) returns the energy of the 3’ dangle of \( i + 1 \) on \( i \) plus the energy of the 5’ dangle of \( j - 1 \) on \( j \). Additionally, if \( i < j \), then by convention, the function \( \text{dangle}(j, i) \) returns the energy of the 5’ dangle of \( i - 1 \) on \( i \) plus the energy of the 3’ dangle of \( j + 1 \) on \( j \). Note that energy values of 3’ dangles are higher than those of 5’ dangles. Our manner of handling dangles is similar to that of [24] in RNAfold.

Although a single array \( Z \) suffices to compute the partition function with respect to the Nussinov-Jacobson energy model, this is no longer the case for the Turner model. In the latter case, for each \( k, i, j, s, b \), we must distinguish the partition function according to the last secondary structure element encountered (i.e. hairpin, stem, bulge, internal loop, multi-loop or exterior loop). Each of these 5 conditions requires a distinct array \( Z \).

- \( Z_s \): Secondary structures such that the first and the last paired nucleotides base-pair together. An example is \( \ldots(\text{***})\ldots \), where \( \cdot \) denotes an unpaired position, \( \text{***} \) denotes any valid substructure. The subscript \( s \) in \( Z_s \) suggests “stem”.

- \( Z'_s \): Secondary structures such that the leftmost and rightmost nucleotides of the subsequence base pair together. Fields \( s \) and \( b \) are irrelevant in this case and so omitted. The well-balanced parenthesis expression with dots (*** is an example.

- \( Z_e \): At least 2 hairpin loops occur in an exterior loop (unpaired nucleotides do not receive a penalty since they are external to any base pair). An example of this type of structure is given by \( \ldots(\text{***})\ldots(\text{***})\ldots \).

- \( Z^m_e \): Same as \( Z_e \), except that a penalty for unpaired bases occurring in a multi-loop is added for each nucleotide occurring outside the stem. An example of this type of structure is given by \( \ldots(\ldots(\text{***})\ldots(\text{***})\ldots) \).

- \( Z_m \): At least 2 hairpin loops appear in a multi-loop. In this case, a penalty is added for unpaired bases outside each stem.
It may seem that the case $Z'_s$ is included in the case $Z_s$; however, values stored in $Z_s(i, j, k, \emptyset, 0)$ differ from those of $Z'_s(i, j, k)$ because of dangle energy contributions (see algorithm description).

In order to reduce space complexity, we use the function $\text{segmentBasePair}(i_1, j_1, i_2, j_2)$ which returns $\text{true}$ if one can form a valid base pair for the segments $[i_1, j_1]$ and $[i_2, j_2]$ and $\text{false}$ otherwise. More formally, $\text{segmentBasePair}(i_1, j_1, i_2, j_2)$ holds exactly when $(\exists x, y \in \{i_1, \ldots, j_1\} \cup \{i_2, \ldots, j_2\})$ such that $x < y$ and $x, y$ can form a valid Watson-Crick or wobble pair.

The results of the $\text{segmentBasePair}$ function can be computed in $O(1)$ time using an 2-dimensional array $\text{nbSingleBp}$ which stores in a cell $\text{nbSingleBp}[i, j]$ the number of single base pairs (i.e. the number of secondary structures with one and only one base pair) over the subsequence $a_i, \cdots, a_j$. Indeed, one can note that the number of basepairs than can be formed over the two subsequence $[i_1, j_1]$ and $[i_2, j_2]$ is equal to $\text{nbSingleBp}[i_1, j_2] - \text{nbSingleBp}[i_1, i_2] - \text{nbSingleBp}[j_1, j_2] + \text{nbSingleBp}[j_1, i_2]$. Using this remark, we can compute this value and return $\text{true}$ if and only if the result is positive (false otherwise). Note that when only two index are given, the function $\text{segmentBasePair}$ operates for a single interval $[i_1, j_1]$. The same array $\text{nbSingleBp}$ can be used for this task.

### 3.3 Algorithm description

When considering substring $a_i, \ldots, a_j$ of the input RNA sequence, the partition function algorithm, given in Figures [10] and [11] treats 3 distinct cases, according to whether $i, j$ are base-paired, or $r, j$ are base-paired for some intermediate $i < r \leq j - \theta - 1$, or $j$ is not base-paired. Case 1, treated in lines 7 to 20, considers all saturated secondary structures containing base pair $(i, j)$, while Case 2, treated in lines 21 to 36, considered those containing base pair $(r, j)$, for some intermediate $r$, $i < r \leq j - \theta - 1$. Case 3, treated in lines 38 to 52, considered those saturated secondary structures in which $j$ does not base-pair with any position in $\{i, \ldots, j - \theta - 1\}$  – note that it can happen (later) that $j$ base-pairs with a nucleotide in $\{1, \ldots, i - 1\}$ or $\{j + \theta + 1, \ldots, n\}$. This latter case is the most difficult, and indeed is the reason for having introduced the visibility parameters $s, b$ in the first place.

In each block of of pseudocode, several possibilities are discussed. In Case 1, we consider the occurrence of a hairpin (lines 9-11), the closing of a multi-loop (line 12-14) and any stacked base pairs, bulges and internal loops (lines 15-19). The break statements at lines 16 and 18 prevent useless loops. Indeed, if the predicate $\text{segmentBasePair}(i, x, y, j)$ is $\text{true}$, then $\text{segmentBasePair}(i, x', y', j)$ is also $\text{true}$ for all $x \geq x'$ and $y' \leq y$. In Case 2, extension with unpaired bases on the left is examined in lines 23 to 26, while concatenation of helices appearing in a multi-loop or an exterior loop is handled in lines 27 to 36. Note that in this case lines 32-33 correspond to an initialization of the helical sequence, while lines 34-35 add a new stem to an already existent helical sequence. In Case 3, before any extension with an unpaired nucleotide on the right, we distinguish the “general case” (lines 41-45), such that $0 \leq b \leq \theta$, from the special case (lines 46 to 51), where $b = \theta + 1$. Any extension requires us to check that the rightmost nucleotide cannot already base-pair, before forming a base pair with the rightmost position; i.e. that $(j - \theta - 1, j)$ cannot base-pair.
3.4 Complexity

As earlier mentioned, $s$ and $b$ are of constant size ($s$ can take at most 16 values and $b$ can take at most 5 values). The partition function algorithm given in Figures 10 and 11 runs in time $O(n^4)$ with space requirement of $O(n^3)$. In practice, our implementation requires roughly 1 minute and 250 MByte to run for an input RNA sequence of 100 nucleotides.

4 Boltzmann sampling of saturated secondary structures

The Boltzmann partition function can be used to weight each secondary structure with respect to its energy contribution in the ensemble of all possible secondary structures. In analogy to Ding and Lawrence [14], who gave the first correct and mathematically rigorous sampling algorithm for RNA structures, we proceed as follows.

4.1 Notation

Here we define additional functions required for the sampling algorithm, depicted in Figures 12 and 13. The function $\text{remove}(i,s)$, where $i \in \mathbb{N}$ and $s \subseteq \{A,U,G,C\}$, returns the subset $s \setminus \{a_i\}$ of $s$ which does not contain the nucleotide at position $i$. The function $\text{compatible}(i,j,s)$ returns true if the set of all nucleotides present in $a_i, \ldots, a_j$ equals $s$, and returns false otherwise. The power set function $\mathcal{P}(s)$ returns the collection of all subsets of $s$, and function $\text{addBasePair}(i,j,\text{sample})$ adds the base pair $(i,j)$ to a secondary structure (called $\text{sample}$ below). All other functions have been previously defined. The $Z$-tables, i.e. $Z_e$, $Z_m$, etc., used in our sampling algorithm are the same as those used in our partition function algorithm. Additionally, in the case of exterior loop, we define a new array $Z'_e$ such that $Z'_e(i,j,k,s,b) = Z_e(i,j,k,s,b) + Z_e(i,j,k,s,b)$, for all $i,j,k,s$ and $b$. This array stores the values of the partition function for helical sequences belonging to an exterior loop, regardless of the number of helical sequences (one or more instead of at least two).

4.2 Algorithm description

The linear time sampling algorithm works by calling itself recursively. We begin with an empty secondary structure, denoted $\text{sample}$, and we stipulate the number $k$ of base pairs that the sample should ultimately contain. Since the Boltzmann probability at temperature $T$ for the collection of saturated secondary structures having $k$ base pairs equals the relative density of states $\rho_k^T = Z_k^T / \sum_k Z_k^T$, the value of $k$ is obtained by sampling. Moreover, $Z_k^T = \sum_{s,b} Z_k^T[k,1,n,s,b]$, and hence initial values of $s,b$ are obtained by sampling – say $s_0,b_0$. Given the visibility constraint $s_0,b_0$, in the sample to be produced, either $(1,n) \in \text{sample}$, or $(r,n) \in \text{sample}$ for some intermediate $1 < r \leq n - \theta - 1$, or $n$ is not base-paired in $\text{sample}$. Each of these alternatives determines visibility parameters, which must then agree with the initially determined values $s_0,b_0$. The structure $\text{sample}$ is then progressively filled.
by recursive calls of the sampling algorithm. At the end of the sampling algorithm the completed structure \textit{sample} having \(k\) base pairs is returned.

Subfunctions are designed to sample a specific (or a set of specific) secondary structure elements and to decide which ones will be called recursively. Three such subfunctions are needed. They are respectively associated with the sampling of base pairs in helices (Figure 12), decomposition of helix sequences in multi-loop (Figure 13) and decomposition of helix sequences in exterior loop (Figure 14). All functions follow the same approach. First, a random number in the real interval \((0, 1)\) is generated according to the uniform distribution (line 1 in Figures 12 and 13) and then this random number is used together with (previously computed, stored) partition function values to choose the next case according to the Boltzmann probability.

The sampling function \texttt{samplingHelix} (Figure 12) is the only function to write a base pair in the sample structure (line 3). The recursive call in the case of a stem structure is developed in lines 5 to 10, and the occurrence of a multi-loop is discussed in lines 12 to 17.

The function \texttt{samplingMultiLoop} (Figure 13) discusses the value of \(b\) and its possible ancestors. In lines 3 to 13, we assume that the upper bound for \(b\) is reached. The distance \(d\) from the rightmost nucleotide to the rightmost base-paired nucleotide is thus \textit{at least} \(\theta + 1\). We consider two possible values for its ancestor, namely \(\theta + 1\) (\(d > \theta + 1\)) and \(\theta\) (\(d = \theta + 1\)). When \(1 \leq b \leq \theta\) (lines 14 to 19) this discussion does not pertain since there is only one possibility (i.e. \(b - 1\)).

When \(b = 0\) (lines 20 to 36), the rightmost base is paired. This means that we need to extract the rightmost helix from the helical sequence of the multi-loop. Thus, we search for a valid cut point \(r\) (lines 21 and 22) and recursively call the helix sampling algorithm on the rightmost helix and recursively call the multi-loop sampling algorithm on the left part. Once the cut point \(r\) has been chosen, we take care of the left string status (line 23). If this segment can base-pair (lines 24 to 37), then the left part contains \textit{at least} one helix and hence the function calls itself recursively on this segment while \texttt{samplingHelix} is called on the right part. Line 28 to 32 treat the case of a left part which contains at least two helices. This implies that the multi-loop sampling is not terminated and then field \texttt{lastHelix} keeps the \texttt{false} value for the recursive sampling (line 30). In lines 33 to 37, we treat the case of left and right parts containing one and only one helix. Otherwise (lines 38 to 42), the left string cannot base-pair and the helix is the last one in the multi-loop. Before calling \texttt{samplingHelix} on the right side and terminating the sampling of the multi-loop we must check the correctness of this configuration (line 38). Hence we check first that more than one stem will appear in the multi-loop and next that the argument \(s\) is compatible with the visibility set of the left string.

When sampling an exterior loop (Figure 14), the principles are identical to those already described for function \texttt{samplingMultiLoop}. We need only to omit the energy contributions from dangle and to use array \(Z_s, Z_e\) and \(Z'_e\) instead of \(Z^m_s\) and \(Z^m_e\). This implies a simplification when sampling an exterior loop which contains at least two helices (lines 28 to 32).
4.3 Correctness of the sampling algorithm

To prove the correctness of the sampling algorithm, we need to demonstrate that a saturated structure can be decomposed into smaller saturated structures. This is done in the following Lemma 1. In this section, we consider a fixed RNA sequence $a_1, \ldots, a_n$, and use the notation $[i, j]$ to denote the interval $\{i, i+1, \ldots, j\}$ of integral values between $i$ and $j$.

**Lemma 1**

Assume that $S_{i,j}$ is a saturated secondary structure on RNA subsequence $a_i, \ldots, a_j$, and that $S_{i,j}$ has $k$ base pairs. Then exactly one of the following cases holds.

1. Nucleotides $i$ and $j$ base-pair and enclose a saturated secondary structure with $k-1$ base pairs.

2. There exists an index $i < r \leq j - \theta - 1$ such that $(r, j) \in S_{i,j}$, and there exist two saturated secondary structures $S_{i,r-1}$ and $S_{r,j}$ having respectively $k_1$ and $k_2$ base pairs on $a_i, \ldots, a_{r-1}$ and $a_r, \ldots, a_j$ respectively, where $k = k_1 + k_2 + 1$ and $S_{i,j} = S_{i,r-1} \cup S_{r,j} \cup [r, j]$.

3. Position $j$ is not base-paired and the secondary structure $S_{i,j-1}$ restricted to $a_i, \ldots, a_{j-1}$ is saturated.

**Proof.** Assume that nucleotides $i$ and $j$ base-pair with each other, and consider the secondary structure $S_{i+1,j-1}$ restricted to $a_{i+1}, \ldots, a_{j-1}$. If $S_{i+1,j-1}$ is not saturated, then for some $i < x < y < j$, the base pair $(x, y)$ can be added to $S_{i+1,j-1}$. But then $S_{i,j} \cup \{(x, y)\}$ would be a valid secondary structure, contradicting local optimality of $S_{i,j}$.

Assume now that nucleotides $i$ and $j$ do not base-pair together. Then either the nucleotide at position $j$ base-pairs with a nucleotide at position $i < r \leq j - \theta - 1$, or position $j$ is unpaired with any position in the interval $[i, j]$. In the former case, $S_{i,j}$ can be decomposed into two distinct secondary structures $S_{i,r-1}$ and $S_{r,j-1}$ along with the base pair $(r, j)$. If either of the substructures is not saturated, then for some $x, y$ satisfying $i \leq x < y \leq r - 1$ resp. $r \leq x < y \leq j - 1$, the base pair $(x, y)$ could be added to $S_{i,r-1}$ resp. $S_{r,j-1}$. But then $S_{i,j} \cup \{(x, y)\}$ would be a valid secondary structure, contradicting local optimality of $S_{i,j}$. In the latter case, if for $i \leq x < y \leq j - 1$ the base pair $(x, y)$ could be added to $S_{i,j-1}$, without violating the definition of secondary structure, then since $S_{i,j} = S_{i,j-1}$, the same is true for $S_{i,j}$, which contradicts local optimality of $S_{i,j}$. This concludes the proof. □

Lemma 1 states that every saturated secondary structure can be decomposed into smaller saturated secondary structures, themselves saturated. This is the justification for a recursive sampling algorithm for saturated structures. While RNAsat executes correct sampling with respect to the Turner energy model, for readability, we present the proof of correctness for the simpler Nussinov-Jacobson model. The following proposition gives the Boltzmann probabilities for each configuration described in Lemma 1.

---

10This is similar to the presentation style of [22], where details are given for the Nussinov-Jacobson model, although the authors implemented their algorithm for the Turner energy model.
Proposition 1
Let \(a_i, \ldots, a_j\) be a subword from the RNA sequence \(a_1, \ldots, a_n\). Let \(bp(i, j)\) denote the Nussinov-Jacobson energy associated with base pair \((i, j)\); in particular, if \(a_i, a_j\) do not form a Watson-Crick or wobble pair, then \(bp(i, j) = +\infty\). For all possible positive numbers of base pairs \(0 < k \leq \max\BP(1, n)\) and all possible visibility parameters \(s\) and \(b\), we have the following.

\[
Z(k, i, j, s, b) = e^{-\frac{bp(i, j)}{RT}} \sum_{s', b'} Z(k - 1, i + 1, j - 1, s', b') \\
+ \sum_{\text{comp. } k', r, s', b'} Z(k', i, r - 1, s', b') \cdot e^{-\frac{bp(r, j)}{RT}} \sum_{s'', b''} Z(k - k' - 1, r + 1, j - 1, s'', b'') \\
+ \sum_{\text{comp. } s', b'} Z(k, i, j - 1, s', b').
\]

In the terms (3) and (4), the sum is taken over all values \(s', b'\) which are compatible with values \(s, b\).

Proof. The proof is by simultaneous induction on \(k, s, b\). By definition, we have

\[
Z(k, i, j, s, b) = \sum_{S_{i,j} \text{ is loc. opt.}} e^{-\frac{E(S_{i,j})}{RT}}
\]

where \(S\) is constrained by parameters \(k, s\) and \(b\). Depending on the status of position \(j\) (base-paired or not), there are three cases. (i) \((i, j)\) is a base pair, (ii) \((r, j)\) is a base pair, for some \(i < r \leq j - \theta - 1\), (iii) \(j\) is not base-paired with any position in the interval \([i, j]\). This yields the following equation, where ‘s.t.’ abbreviates “such that”, and ‘bp’ abbreviates “base-pair”.

\[
Z(k, i, j, s, b) = \sum_{S_{i,j} \text{ s.t. } (i, j) \text{ bp}} e^{-\frac{E(S_{i,j})}{RT}} + \sum_{S_{i,j} \text{ s.t. } (r, j) \text{ bp}} e^{-\frac{E(S_{i,j})}{RT}} + \sum_{S_{i,j} \text{ s.t. } j \text{ not bp}} e^{-\frac{E(S_{i,j})}{RT}}
\]

Case 1. \((i, j)\) is a base pair of \(S_{i,j}\). Let \(S_{i+1, j-1}\) denote the restriction of \(S_{i,j}\) to \(a_{i+1}, \ldots, a_{j-1}\), obtained after removal of the extremal base pair \((i, j)\). The energy \(E(S_{i,j})\) of a secondary structure \(S_{i,j}\) on \(a_1, \ldots, a_j\) such that \((i, j) \in S_{i,j}\) can be decomposed as \(E(S_{i,j}) = bp(i, j) + E(S_{i+1, j-1})\). Hence the first sum can be written as

\[
\sum_{S_{i,j} \text{ s.t. } (i, j) \text{ bp}} e^{-\frac{E(S_{i,j})}{RT}} = \sum_{S_{i+1, j-1}} e^{-\frac{(bp(i, j) + E(S_{i+1, j-1}))}{RT}}
\]

\[
= e^{-\frac{bp(i, j)}{RT}} \sum_{S_{i+1, j-1}} e^{-\frac{E(S_{i+1, j-1})}{RT}}
\]

By Case 1 of Lemma [Lemma 1] we know that \(S_{i+1, j-1}\) is saturated. We then cluster such secondary structures \(S_{i+1, j-1}\), each having \(k - 1\) base pairs, according to their visibility parameters \(s'\) and \(b'\).
\[
\sum_{S_{i,j} \text{ s.t. (i,j) bp}} e^{-\frac{E(S_{i,j})}{RT}} = e^{-\frac{bp(i,j)}{RT}} \sum_{S'_{i,j-1}} \sum_{S'_{i+1,j-1}} e^{-\frac{E(S'_{i+1,j-1})}{RT}} \\
= e^{-\frac{bp(i,j)}{RT}} \sum_{S'_{i,j-1}} Z(k-1, i+1, j-1, s', b')
\]

This completes the proof for Case 2 of proposition 1.

Case 2. For some intermediate position \( r \), with \( i < r \leq j - \theta - 1 \) \((r, j) \in S_{i,j}\). By Case 2 of Lemma 1 the substructures \( S_{i,r-1} \) and \( S_{r+1,j-1} \) are saturated. The energy \( E(S_{i,j}) \) of a secondary structure on \( a_1, \ldots, a_j \), such that \((r, j) \in S_{i,j}\), can be decomposed as \( E(S_{i,j}) = bp(r, j) + E(S_{i,r-1}) + E(S_{r+1,j-1}) \).

\[
\sum_{S_{i,j} \text{ s.t. (r,j) bp}} e^{-\frac{E(S_{i,j})}{RT}} = e^{-\frac{bp(r,j)}{RT}} \sum_{S_{i,r-1}, S_{r+1,j-1}} e^{-\frac{(E(S_{i,r-1}) + E(S_{r+1,j-1}))}{RT}} \\
= e^{-\frac{bp(r,j)}{RT}} \sum_{S_{i,r-1}} \sum_{S_{r+1,j-1}} e^{-\frac{(E(S_{i,r-1}) + E(S_{r+1,j-1}))}{RT}}
\]

Let \( k' \) denote the number of base pairs of \( S_{i,r-1} \), and \( s' \) and \( b' \) denote the visibility parameters of \( S_{i,r-1} \). Note that \( s', b' \) must be compatible with \( s, b \); i.e. since \((r, j) \) is a base pair of \( S_{i,j}\), it must be that \( b = 0 \) and \( s = s' \cup \{a_{r-1-m} : 0 \leq m < b'\} \). Cluster the left substructures \( S_{i,r-1} \) by (compatible) visibility parameters \( s', b' \). By Lemma 1 the left substructure \( S_{i,r-1} \) is saturated, hence the sum can be rewritten as follows.

\[
\sum_{S_{i,j} \text{ s.t. (r,j) bp}} e^{-\frac{E(S_{i,j})}{RT}} = e^{-\frac{bp(r,j)}{RT}} \sum_{\text{comp. } k', r, s', b'} \sum_{S_{i,r-1}(k', r, s', b')} \sum_{S_{r+1,j-1}} e^{-\frac{(E(S_{i,r-1}(k', r, s', b')) + E(S_{r+1,j-1}))}{RT}} \\
= \sum_{\text{comp. } k', r, s', b'} Z(k', r-1, s', b') \sum_{S_{r+1,j-1}} e^{-\frac{(E(S_{r+1,j-1}))}{RT}}
\]

Since \((r, j) \) is a base pair, by induction we can apply Case 2 of this proposition to the right substructure \( S_{r,j-1} \). This concludes the proof of Case 3.

\[
\sum_{S_{i,j} \text{ s.t. (r,j) bp}} e^{-\frac{E(S_{i,j})}{RT}} = \sum_{\text{comp. } k', r, s', b'} Z(k', r-1, s', b') \cdot e^{-\frac{bp(i,j)}{RT}} \sum_{s'', b''} Z(k_2-1, i+1, j-1, s'', b'').
\]

Case 3. In this case, the rightmost position \( j \) is not base-paired with any position in the interval \([i, j]\). Hence, \( E(S_{i,j}) = E(S_{i,j-1}) \), where \( S_{i,j-1} \) denotes the secondary structure restricted \( a_1, \ldots, a_{j-1} \).

\[
\sum_{S_{i,j} \text{ s.t. } j \text{ not bp}} e^{-\frac{E(S_{i,j})}{RT}} = \sum_{S_{i,j-1}} e^{-\frac{E(S_{i,j-1})}{RT}}
\]
Let $s'$ and $b'$ denote the visibility parameters of $S_{i,j-1}$. Note that these parameters must be compatible with $s$ and $b$; i.e. since $j$ is unpaired in $S_{i,j}$, it must be that $b = \max(b' + 1, \theta + 1)$, and $s = s' \cup \{a_{j-\theta-1}\}$ if $b' = \theta + 1$ while $s = s'$ if $0 \leq b' \leq \theta$. (This is the reason for line 10 in Figures 13 and 14.) Cluster the secondary structures $S_{i,j-1}$ by these compatible parameters $s', b'$. Lemma 1 ensures that these secondary structures are saturated, hence by induction we have the following.

$$
\sum_{S_{i,j} \text{ s.t. } j \text{ not bp}} e^{-\frac{E(S_{i,j})}{RT}} = \sum_{\text{comp. } s',b'} \sum_{S_{i,j-1}(s',b')} e^{-\frac{E(S_{i,j-1}(s',b'))}{RT}} = \sum_{\text{comp. } s',b'} Z(k, i, j - 1, s', b')
$$

This concludes the proof of proposition 1. □

Proposition 1 implies the correctness of the sampling algorithm for locally optimal secondary structures with respect to the Nussinov-Jacobson energy model. This explanation should suffice to explain the underlying ideas in the pseudocode for sampling with respect to the Turner energy model, given in Figures 12, 13 and 14.

Acknowledgements

We would like to thank Michael Zuker for remarks and for the suggestion to use mfold 2.3 energy parameters for our temperature-dependent computations. We are indebted to two anonymous referees for their helpful remarks and criticisms, and to Yann Ponty for an algorithmic suggestion. The work of both authors was partially supported by the grant NSF DBI-0543506 to P.C.

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conserved RNA secondary structures predicts thousands of functional noncoding RNAs


Table 1: Shape frequencies for 100 sampled structures (with no restriction to being saturated) of the 149 nt. SAM riboswitch with EMBL accession number AL939119.1/177986-178133. Shapes with low frequency are not displayed for reasons of space economy. Notice how the frequency of certain shapes depends on temperature (25, 37, 50 degrees Celsius), and on the number $k$ of base pairs fewer than that of the Nussinov-Jacobson optimal structure for this riboswitch, which has 55 base pairs. By ‘sat’, we mean that there is no restriction on $k$; i.e. the sampling is general over all secondary structures, where the probability of sampling a $k$-saturated structure is $\hat{Q}_k^T / \sum_k \hat{Q}_k^T$. The shape frequencies for these structures should be compared with those for saturated structures, presented in Table 2.
Table 2: Shape frequencies for 100 sampled saturated structures of the 149 nt. SAM riboswitch with EMBL accession number AL939119.1/177986-178133. Shapes with low frequency are not displayed for reasons of space economy. Notice how the frequency of certain shapes depends on temperature (25, 37, 50 degrees Celsius), and on the number $k$ of base pairs fewer than that of the Nussinov-Jacobson optimal structure for this riboswitch, which has 55 base pairs. By ‘sat’, we mean that there is no restriction on $k$; i.e. the sampling is general over all saturated structures, where the probability of sampling a $k$-saturated structure is $\hat{Z}_k^T / \sum_k \hat{Z}_k^T$. The shape frequencies for these saturated structures should be compared with those for general structures, presented in Table 1.

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Table 3: Saturated structures for pseudoknot PKB 00152. One thousand saturated structures were sampled from the collection of all 111,014 saturated structures for the RNA sequence UGGUGCCAUCAACCAUAUGAUGGGA with secondary structure and annotated pseudoknot .(((.[[[[[[])[.]]]]]])]. For each structure, the frequency and energy in kcal/mol are additionally indicated.
Figure 1: Graph of the ensemble free energy $G_k = -RT \ln \tilde{Z}_k^T$ for $k$-saturated secondary structures for hammerhead type III ribozyme with Rfam [19] accession number AF170517 for both the Turner energy model (left panel) and the Nussinov-Jacobson energy model (right panel). As explained in the text, $\tilde{Z}_k^T$ is the sum over all $k$-saturated structures $S$ of $\exp(-E(S)/RT)$. The $x$-axis of each graph indicates the value of $k$, the saturation parameter – recall that a secondary structure $S$ is $k$-saturated if $S$ is saturated and has $k$ fewer base pairs than that of the Nussinov-Jacobson optimal structure. Ensemble free energy $G_k$ is given on the $y$-axis. It follows that energy values for $k = 0$ correspond to the ensemble free energy of 0-saturated structures, those having the maximum number of base pairs. Each graph displays six curves, corresponding respectively to temperature in degrees Celsius of 0, 20, 40, 60, 80, 100. The curve for $T = 0^\circ$ lies on the bottom, with curves for higher temperatures lying above those for lower temperatures. Note that the lowest ensemble free energy for the Turner model (left panel) occurs when $k = 1$. For the Turner energy model, ensemble free energy is in units of kcal/mol. Energy units are (roughly) in kcal/mol for the Nussinov-Jacobson (NJ) energy model, for the following reason. For each of the 57 seed hammerhead type III ribozymes in Rfam, we computed the minimum free energy using Vienna RNA Package RNAfold. Additionally, we computed the maximum number of base pairs using our implementation of the Nussinov-Jacobson algorithm. From this, we obtain an average minimum free energy per base pair of $-0.923385$ for Rfam family RF00008 of type III hammerhead ribozymes. Since this value is close to the value $-1$ assigned per base pair in the Nussinov Jacobson model, it follows that the energy units in the left and right graphs of this figure are comparable.
Figure 2: Graph of the expected number of base pairs as a function of temperature for signal recognition particle with Rfam [19] accession number X12643. Temperature in degrees Celsius is given on the x-axis, while the expected number of base pairs (base pairs/n), normalized by sequence length n, is given on the y-axis. Expected number of base pairs is computed by $\langle \text{base pairs} \rangle = \sum_k k \cdot Q^T_k / Q^T$, where $Q^T_k$ is the partition function at temperature $T$ for all secondary structures having exactly $k$ base pairs, and $Q^T = \sum_k Q^T_k$ is the partition function at temperature $T$ for all secondary structures. Although $Q^T$ is the value obtained by McCaskill’s algorithm [33], and can be obtained using RNAfold -p, the values $Q^T_k$ can only be obtained with our software.
Figure 3: (i) Left panel: Graph of relative density of states for $k$-saturated secondary structures of precursor microRNA \textbf{ath-MIR156c} of \textit{Arabidopsis thaliana}, for values of $0 \leq k \leq 10$, as a function of temperature in degrees Celsius. Notice that for $T = 0$, over 90% of the saturated structures are 0-saturated (i.e. have the maximum number of base pairs). As temperature rises, fewer saturated structures are 0-saturated, as indicated by the gradual decline of the first curve. At the same time, there is a gradual increase, as temperature increases, of the proportion of 1-saturated structures (i.e. have one fewer base pair than the maximum), etc. (ii) Right panel: Curve of specific heat for \textbf{ath-MIR156c}, produced by \texttt{RNAheat} from Vienna RNA Package. Temperature is given on the $x$-axis and specific heat on the $y$-axis.
Figure 4: Precursor microRNA cel-mir-72 of length 96 from *Caenorhabditis elegans* with sequence GUAGUCAUGACGACGUUCCCUAUAUCACGUGCAAGGUCGCUGAGACGCUAGGCAAGUGUUGCCAUAGCUGGAUGAUCGCUAU (data from Sanger Center miRBase linked by Rfam [19]). (Left) Minimum free energy structure, as computed with RNAfold from Vienna RNA Package 1.5. (Right) Consensus structure from 1000 sampled saturated structures, using RNAsat. Here, the consensus structure consists of those base pairs occurring in strictly greater than half the sampled structures.
Figure 5: Precursor microRNA cel-mir-72 of length 96 from Caenorhabditis elegans. (Left) Dot plot of base pair frequencies from 1000 samples over all secondary structures. This implementation of McCaskill’s algorithm [33] within RNAsat. (Middle) Dot plot of base pair frequencies from 1000 samples of approximately locally optimal secondary structures, produced by a greedy algorithm described in the text. (Right) Dot plot of base pair frequencies from 1000 samples of saturated secondary structures produced by RNAsat. It is of interest that the Boltzmann centroid output by the Sfold [13] server is empty; i.e. the centroid has no base pairs.
Figure 6: Pseudoknot PK2 of the upstream pseudoknot domain (UPD) of the 3′ UTR of RNA beta, with Pseudobase [40] number PKB 00152, EMBL accession number X03854 and start position 3088. The RNA sequence is UGGUGCCCAUCAACCAUAUGAUGGGA and the secondary structure with annotated pseudoknot is .(((.[[[[[[.[[[[.]))].]))])))).. The left panel displays the minimum free energy secondary structure predicted by version 1.5 of Vienna RNA Package RNAfold. The right panel displays the Boltzmann pair probabilities, as calculated using McCaskill’s algorithm [33] using RNAfold –p.
Figure 7: Dotplots of pseudoknot PKB 00152 as described in the caption of Figure 6, where the lower triangular portion depicts the minimum free energy structure, while the upper triangular portion describes pair frequencies according to the experiment performed. (Left panel) Pair frequencies, computed by sampling 1000 locally optimal structures, where all locally optimal structures were computed brute force by exhaustive search (see text for details); subsequently structure $S$ was sampled with probability $\exp(-E(S)/RT)/\sum S \exp(-E(S)/RT)$, where the sum is taken over all locally optimal structures. Note that this dot plot is identical to that of Figure 6 and is identical to the dot plot produced by Sfold [13]. (Right panel) Pair frequencies, computed by sampling 1000 saturated structures using RNAsat. Notice how alternative base pairs appear, a feature not detected using mfe calculation, McCaskill’s algorithm, or sampling over all secondary structures. This figure again illustrates the point that saturated structures are different in nature than locally optimal structures, which latter seem to be similar in nature to the Boltzmann low energy ensemble. One might speculate that this data suggests the existence of a folding funnel for RNA secondary structure.
Figure 8: (Left panel) Energy histogram for pseudoknot PKB 00152, the RNA sequence described in the caption of Figure 6. Histogram of energies of all saturated secondary structures. All saturated structures were determined brute force by exhaustive search. Note the energy variation for saturated structures. The histogram of energies of all locally optimal structures, as determined brute force by exhaustive search, is not shown because all locally optimal structures have the same energy as that of the mfe structure, i.e. $-10.5$ kcal/mol.

(Right panel) Graph of the relative density of states $\rho(k) = Z_k/Z$ as a function of $k$, the number of base pairs, for precursor microRNA cbr-mir-42 from C. briggsae. The lower curve with wider spread is that of cbr-mir-42, while the higher curve with more narrow spread is that of random RNA of the same length and dinucleotide frequency, as produced by our implementation of the Altschul-Erikson algorithm – see [1] [10].
Figure 9: Diagram for inductive extension of visibility predicates and overview of the three cases which arise in computing the partition function $Z^T_k(i, j, s, b)$ for saturated $(s, b)$-visible structures on the subsequence $a_i, \ldots, a_j$ having $k$ base pairs. The Feynman diagrams explaining the recursions are not different than those for McCaskill’s algorithm [33], with the exception that the number of base pairs and visibility predicates $VisNuc, VisPos$ depend in a precise manner on previously computed values which are stored in the dynamic programming table. For that reason, in this figure, we describe that dependence. The solid left rectangle in each of the four panels represents the set $s = VisNuc(S, i, j - 1)$ of visible nucleotides in positions $i, \ldots, (j - 1) - \theta - 1$. The dotted left rectangle in the first panel represents the $s' = VisNuc(S, i, j)$ of visible nucleotides in positions $i, \ldots, j - \theta - 1$. The solid right rectangle in each of the four panels represents the value $b = VisPos(S, i, j - 1)$ equal to the number of visible positions $(j - 1) - \theta, \ldots, j - 1$ which occur to the right of the rightmost position basepaired in interval $[i, j - 1]$. The dotted right rectangle in the first panel represents the value $b' = VisPos(S, i, j)$ equal to the number of visible positions $j - \theta, \ldots, j$ which occur to the right of the rightmost position basepaired in interval $[i, j]$. The black dot to the right of the rectangular regions represents the position $j$. By induction, we assume that $Z^T_\ell(i, j - 1, u, v)$ has been computed for all values $\ell, u \subseteq \{A, C, G, U\}$ and $b \leq \theta$. We now describe how the contributions from $(s, b)$-visible structures on $a_i, \ldots, a_{j-1}$ can be extended to contributions from $(s', b')$-visible structures on $a_i, \ldots, a_j$. (First panel) Schematic view of how to update the visibility parameters inductively. (Second panel) In case 1, $j$ basepairs with $i$, and the resulting structure is $(s', b')$-visible, where $s' = \emptyset, b' = 0$. Thus $Z^T_k(i, j, s', b')$ receives contributions from hairpin loops, bulges, internal loops and multi-loops closed by base pair $(i, j)$ having $k$ base pairs and visibility parameters $(s, b)$. (Third panel) In case 2, $j$ basepairs with intermediate $r$, for some $i < r \leq j - \theta - 1$, and the resulting structure is $(s', b')$-visible, where $s' = Vis(S, i, r - 1), b' = 0$. Thus $Z^T_k(i, j, s', b')$ receives contributions from hairpin loops, bulges, internal loops and multi-loops closed by base pair $(i, j)$ having $k$ base pairs and visibility parameters $(s, b)$. (Right panel) In case 3, $j$ does not basepair. In this case, the resulting structure is $(s', b')$-visible, where $s' = s \cup \{a_{j-\theta-1}\}$ if $VisPos(S) = \emptyset$, and otherwise $s' = s$. The new visible position $b' = \max(b, \theta)$. Thus $Z^T_k(i, j, s', b')$ receives contributions from hairpin loops, bulges, internal loops and multi-loops on $(i, j - 1)$ having $k$ base pairs and visibility parameters $(s, b)$. 

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Figure 10: Algorithm to compute the Boltzmann partition function for saturated secondary structure having $k$ base pairs, with respect to the Turner energy model.
for all \( 1 < nBP \leq maxBP(i,j) \)

for all \( s \in \mathcal{P}\{\{A,C,G,U\}\} \)

if \( \text{basepairWithSet}(s,j) = \text{false} \) {

for all \( 0 \leq b < \theta \) // general case

\[
\begin{align*}
Z_e(i,j,nBP, s, b + 1) & = Z_e(i,j - 1,nBP, s, b) \\
Z_m(i,j,nBP, s, b + 1) & = Z_m(i,j - 1,nBP, s, b) \cdot e^{-\frac{\alpha_3}{RT}} \\
Z_s(i,j,nBP, s, b + 1) & = Z_s(i,j - 1,nBP, s, b) \cdot e^{-\frac{\alpha_3}{RT}}
\end{align*}
\]

if \( \text{canBasePair}(j - \theta - 1, j) = \text{false} \) // special case

\[
\begin{align*}
s_0 = s \cup \text{nucl}(j - \theta - 1,j - \theta - 1) \\
Z_e(i,j,nBP, s0, \theta + 1) & = Z_e(i,j - 1,nBP, s, \theta + 1) \\
Z_m(i,j,nBP, s0, \theta + 1) & = Z_m(i,j - 1,nBP, s, \theta + 1) \cdot e^{-\frac{\alpha_3}{RT}} \\
Z_s(i,j,nBP, s0, \theta + 1) & = Z_s(i,j - 1,nBP, s, \theta + 1) \cdot e^{-\frac{\alpha_3}{RT}}
\end{align*}
\]

\[
\begin{align*}
Z_e(i,j,nBP, s0, \theta + 1) & = Z_e(i,j - 1,nBP, s, \theta + 1) \\
Z_m(i,j,nBP, s0, \theta + 1) & = Z_m(i,j - 1,nBP, s, \theta + 1) \cdot e^{-\frac{\alpha_3}{RT}} \\
Z_s(i,j,nBP, s0, \theta + 1) & = Z_s(i,j - 1,nBP, s, \theta + 1) \cdot e^{-\frac{\alpha_3}{RT}}
\end{align*}
\]

}
0. `samplingHelix(k,i,j,sample)`
1. `va = random()`
2. `current = 0`
3. `addBasePair(i,j,sample)`
4. if `k = 0` return // no more base pairs to add; end of sampling
5. for `x = i + 1` to `j - \theta - 1`
6. for `y = x + \theta + 1` to `j`
7. if `canBasePair(x,y) = true AND segmentBasepair(i,x,y,j) = false`
8. `current += \frac{stack(i,x,y,j)Z'(x,y,k-1)}{Z'(i,j,k)}`
9. if `va < current`
10. `samplingHelix(k-1,x,y,sample) // (i,j) belongs to a helix`
11. return
12. for `s \in P\{A,C,G,U\}`
13. for `b = 0` to `\theta + 1`
14. `current += \frac{e^{-\alpha_1(i,i)}e^{-\alpha_2(i,i)}Z_m(i+1,j-1,k-1,s,b)}{Z'(i,j,k)}`
15. if `va < current`
16. `samplingMultiLoop(k-1,i+1,j-1,s,b,sample,0) // (i,j) closes a multi-loop`
17. return

Figure 12: Sampling algorithm for helix. This function is called only after having determined that `i,j` can base-pair.
0. \text{samplingMultiLoop}(k, i, j, s, b, \text{sample}, \text{lastHelix})
1. \text{va} = \text{random}()
2. current = 0
3. if \( b = \theta + 1 \) // previous \( b \) can take two values
   4. for \( b' = \theta \) to \( \theta + 1 \)
      5. current + = \frac{-\alpha_3}{8\pi} \cdot Z_m(i, j-1, k, s, b') \frac{1}{Z_m(i,j,k,s,b)}
     6. if \( \text{va} < \text{current} \)
        7. \text{samplingMultiLoop}(k, i, j-1, s, b', \text{sample}, \text{lastHelix})
        8. return
   9. \( s' = \text{remove}(j - \theta - 1, s) \)
10. if \( b' = \theta + 1 \) AND \( s' \neq s \)
    11. current + = \frac{-\alpha_3}{8\pi} \cdot Z_m(i, j-1, k, s, b') \frac{1}{Z_m(i,j,k,s,b)}
    12. \text{samplingSingleLoop}(k, i, j-1, s', b', \text{sample}, \text{lastHelix})
    13. return
14. else if \( b > 0 \)
    15. current + = \frac{-\alpha_3}{8\pi} \cdot Z_m(i, j-1, k, s, b') \frac{1}{Z_m(i,j,k,s,b)}
    16. if \( \text{va} < \text{current} \)
        17. \text{samplingMultiLoop}(k, i, j-1, s, b-1, \text{sample}, \text{lastHelix})
    18. return
19. \text{else} // \( b = 0 \)
20. for \( r = i \) to \( j - \theta - 1 \)
21. if \( \text{canBasePair}(r, n) = \text{true} \)
    22. if \( \text{lastHelix} = \text{false} \) AND \( \text{segmentBasePair}(i, r-1) = \text{true} \)
    23. for \( s' \in P(s) \)
        24. for \( b' = 0 \) to \( \theta + 1 \)
            25. for \( k_1 = 0 \) to \( k-1 \)
               26. \( k_2 = k - k_1 \)
               27. current + = \frac{-\alpha_3}{8\pi} \cdot Z_m(i, r-1, k_1, s', b') \frac{\text{dangle}(r,j)}{8\pi} \cdot Z'_m(r, j, k_2)
            28. if \( \text{va} < \text{current} \) // multi-loop sampling is NOT terminated
               29. \text{samplingMultiLoop}(k_1, i, r-1, s', b', \text{sample}, \text{false})
               30. \text{samplingHelix}(k_2, r, j, \text{sample})
               31. return
               32. current + = \frac{-\alpha_3}{8\pi} \cdot Z_m(i, r-1, k_1, s', b') \frac{\text{dangle}(r,j)}{8\pi} \cdot Z'_m(r, j, k_2)
            33. if \( \text{va} < \text{current} \) // sampling last two helices in multi-loop
               34. \text{samplingMultiLoop}(k_1, i, r-1, s', b', \text{sample}, \text{true})
               35. \text{samplingHelix}(k_2, r, j, \text{sample})
               36. return
            37. \text{else if lastHelix=true AND compatible}(i, r-1, s) = \text{true}
               38. current + = \frac{-\alpha_3}{8\pi} \cdot Z_m(i, r-1, k_1, s', b') \frac{\text{dangle}(r,j)}{8\pi} \cdot Z'_m(r, j, k)
            39. if \( \text{va} < \text{current} \)
               40. \text{samplingHelix}(k, r, j, \text{sample})
               41. return

\text{Figure 13: Sampling algorithm for multi-loop.}
0. `samplingExteriorLoop(k,i,j,s,b,sample)`
1. `va = random()`
2. `current = 0`
3. if `b = θ + 1` // previous b can take two values
4. for `b' = θ to θ + 1`
5. `current += \frac{Z_e(i,j-1,k,s,b')}{Z_e(i,j,k,s,b)}`
6. if `va < current`
7. `samplingExteriorLoop(k,i,j-1,s,b',sample)`
8. return
9. `s' = remove(j - θ - 1, s)`
10. if `b' = θ + 1 AND s' ≠ s`
11. `current += \frac{Z_e(i,j-1,k,s',b')}{Z_e(i,j,k,s,b)}`
12. `samplingSingleLoop(k,i,j-1,s',b',sample)`
13. return
14. else if `b > 0`
15. `current += \frac{Z_e(i,j-1,k,s,b')}{Z_e(i,j,k,s,b)}`
16. if `va < current`
17. `samplingExteriorLoop(k,i,j-1,s-1,b,sample)`
18. return
19. else // b = 0
20. for `r = i` to `j - θ - 1`
21. if `canBasePair(r,j) = true`
22. if `segmentBasePair(i,r-1) = true`
23. for `s' ∈ P(s)`
24. for `b' = 0` to `θ + 1`
25. for `k_1 = 0` to `k - 1`
26. `k_2 = k - k_1`
27. `current += \frac{Z'_e(i,r-1,k_1,s',b')e^{-\text{dangle}(r,j)}}{Z_e(i,j,k,s,b)} · Z'_e(r,j,k_2)`
28. if `va < current`
29. `samplingExteriorLoop(k_1,i,r-1,s',b',sample,false)`
30. `samplingHelix(k_2,r,j,sample)`
31. return
32. else if `compatible(i,r-1,s) = true`
33. `current += \frac{e^{-\text{dangle}(r,j)}}{Z_e(i,j,k,s,b)} · \frac{Z'_e(r,j,k)}{Z'_e(i,j,k,s,b)}`
34. if `va < current`
35. `samplingHelix(k,r,j,sample)`
36. return

Figure 14: Sampling algorithm for exterior loop.