Using the Fast Fourier Transform to Accelerate the Computational Search for RNA Conformational Switches

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Abstract

Using complex roots of unity and the Fast Fourier Transform, we design a new thermodynamics-based algorithm, FFTbor, that computes the Boltzmann probability that secondary structures differ by \( k \) base pairs from an arbitrary initial structure of a given RNA sequence. The algorithm, which runs in quartic time \( O(n^4) \) and quadratic space \( O(n^2) \), is used to determine the correlation between kinetic folding speed and the ruggedness of the energy landscape, and to predict the location of riboswitch expression platform candidates. A web server is available at http://bioinformatics.bc.edu/clotelab/FFTbor/.

Introduction

In [1], we developed a dynamic programming algorithm, RNAbor, pronounced RNA neighbor, which simultaneously computes for each integer \( k \), the Boltzmann probability \( p_k = \frac{Z_k}{Z} \) of the subensemble of structures whose base pair distance to a given initial, or reference, structure \( S' \) is \( k \). (Here, \( Z \) denotes the partition function, defined as the sum of all Boltzmann factors \( \exp(-E(S)/RT) \), over all secondary structures \( S \) of a given RNA sequence, and \( R \) denotes the universal gas constant and \( T \) absolute temperature. Similarly \( Z_k \) denotes the sum of all Boltzmann factors of all structures \( S \), whose base pair distance to the initial structure \( S' \) is exactly \( k \).) RNAbor stores the value of the (partial) partition functions \( Z_k(i,j) \) for all \( 1 \leq i \leq j \leq n \) and \( 0 \leq k \leq n \), each of which requires quadratic time to compute. Thus it follows that RNAbor runs in time \( O(n^5) \) and space \( O(n^4) \), which severely limits its applicability to genomic annotation. This restriction is somewhat mitigated by the fact that in [2], we showed how to use sampling [3] to efficiently approximate RNAbor in cubic time \( O(n^3) \) and quadratic space \( O(n^2) \), provided that the starting structure \( S' \) is the minimum free energy (MFE) structure. We expect that a more efficient version of RNAbor could be used in applications in genomics and synthetic biology, to detect potential conformational switches – RNA sequences containing two or more (distinct) metastable structures.

In this paper, we describe a radically different algorithm, FFTbor, pronounced FFT neighbor, that uses polynomial interpolation to compute the coefficients \( p_0, \ldots, p_{n-1} \) of the polynomial

\[
p(x) = p_0 + p_1 x + p_2 x^2 + \cdots + p_{n-1} x^{n-1},
\]

where \( p_k \) is defined by \( p_k = \frac{Z_k}{Z} \). Due to severe numerical instability issues in both the Lagrange interpolation formula and in Gaussian elimination, we employ the Fast Fourier Transform (FFT) to compute the inverse Discrete Fourier Transform (DFT) on values \( y_0, \ldots, y_{n-1} \), where \( y_k = p(x^k) \) and \( \omega = e^{2\pi i/n} \) is the principal \( n \)th complex root of unity and \( p(x) \) is defined in (1). This gives rise to an improved version of RNAbor, denoted FFTbor, which runs in time \( O(n^4) \) and space \( O(n^2) \). Once two metastable structures \( S_1, S_2 \) are identified, we can subsequently evaluate the feasibility of transition between structures \( S_1 \) and \( S_2 \), by computing the barrier energy using algorithms, such as that described in Dotu et al. [4] or Flamm et al. [5].

Background

Let \( s = s_1, \ldots, s_n \) denote an RNA sequence, i.e. a sequence of letters in the alphabet of nucleotides \( \{A,C,G,U\} \). A secondary structure \( S \) is a set of base pairs \( (i,j) \), where \( 1 \leq i \leq j \leq n \) and \( i \neq j \) represents the minimum number of unpaired nucleotides in a hairpin loop (due to steric constraints, \( \theta \) is usually taken to be \( 3 \)°), such that if \( (i,j) \) and \( (x,y) \) both belong to \( S \), then \( i \neq x = y \) (a nucleotide is involved in at most one base pair) and \( i < x < y < j \) (no pseudoknots are allowed).

The secondary structure \( S \) is compatible with \( s \) if for every base pair \( (i,j) \) in \( S \), the pair \( (s_i, s_j) \) is contained in the set \( \beta = \{ (A,U),(U,A),(G,C),(C,G),(G,U),(U,G) \} \) of six Watson-Crick and wobble base pairs. Often we write that \( S \) is a secondary structure on \( s \), or equivalently, a secondary structure of \( s \), in place of stating that \( S \) is compatible with \( s \). Throughout this paper, by structure, we always mean a secondary structure which is compatible with an arbitrary, but fixed RNA sequence \( s = s_1, \ldots, s_n \).

Given two secondary structures \( S, T \) on \( s \), we define the base pair distance \( d_{bp} \) between \( S \) and \( T \) to be the number of base pairs that they have that are not in common, i.e.
\[ d_{BP}(S,T) = |S \cup T| - |S \cap T| \]  

Structures \( S, T \) are said to be \( k \)-neighbors if \( d_{BP}(S,T) = k \).

For \( 1 \leq i \leq j \leq n \), let \( S_{i,j} \) denote the restriction of \( S \) to interval \([i,j]\) of \( S \), i.e. the set of base pairs \( S_{i,j} = \{(x,y) : i \leq x < y \leq j, (x,y) \in S\} \). The notion of \( k \)-neighbor can also be applied to restrictions of secondary structures; i.e. a secondary structure \( T_{i,j} \) is a \( k \)-neighbor of \( S_{i,j} \) if

\[ d_{BP}(S_{i,j}, T_{i,j}) = \left| \{(x,y) : i \leq x < y \leq j, (x,y) \in S - T \} \right| = k. \]

In the following, we often omit the sequence \( s \) and initial secondary structure \( S^* \) in our notation, since these are arbitrary, but fixed. In particular, we write \( Z_k^i = Z_k^i(s) \) – see following paragraph for definitions.

Given an RNA sequence \( s = s_1, \ldots, s_n \) and any other structure \( S \) on RNA sequence \( s = s_1, \ldots, s_n \) satisfy

\[ d_{BP}(S,S^*) \leq |S^*| + \left\lfloor \frac{n-\theta-1}{2} \right\rfloor \leq n \]  

it follows that the full partition function

\[ Z = Z_{1,n} = \sum_{k=0}^{n} Z_{k,n}^i. \]  

Moreover, since \( \theta = 3 \), we need to compute at most the values \( Z_{1,3}^i, \ldots, Z_{1,n}^i \) – this observation will later prove useful. The Boltzmann probability \( P[d_{BP}(s,S^*) = k] \) that a secondary structure \( S \) on RNA sequence \( s = s_1, \ldots, s_n \) satisfy and base pair distance \( k \) from the initial structure \( S^* \) can be defined from the partition function by

\[ p(k) = \frac{Z_k^i}{Z_{1,n}^i}. \]

By graphing the probabilities \( p_k \) as a function of \( k \), we expect to see one or more peaks at base pair distance \( k \) when there is a meta-stable (low energy) structure \( S \) at base pair distance \( k \) from \( S^* \). See Figure 1 for an illustration.

**Recursions for structural neighbors**

For the rest of the paper, we consider both \( s \) as well as the secondary structure \( S^* \) on \( s \) to be fixed. We now recall the recursions from Freyhurst et al. [9] to determine the partition function \( Z_k^i \) with respect to the Nussinov-Jacobson energy \( E_0 \) model [9], defined by \(-1\) times the number of base pairs; i.e. \( E_0(S) = -1|S| \). Although we describe here the recursions for the

Nussinov-Jacobson model, for the sake of simplicity of exposition, both RNABidr [8] as well as our current software FFTbor, concern the Turner energy model, consisting of free energy parameters for stacked bases, hairpins, bulges, internal loops and multiloops. The full recursions for FFTbor are described for the the Turner energy model in the appendix.

The base case for \( Z_k^i \) is given by

\[ Z_k^i = 1, \text{ for } i \leq j, \]

since the only 0-neighbor to a structure \( S^* \) is the structure \( S^* \) itself, and

\[ Z_k^i = 0, \text{ for } k > 0, i \leq j \leq i + \theta, \]

since the empty structure is the only possible structure for a sequence shorter than \( \theta + 2 \) nucleotides, and so there are no \( k \)-neighbors for \( k > 0 \). The recursion used to compute \( Z_k^i \) for \( k > 0 \) and \( j > i + \theta \) is

\[
Z_k^i = Z_{k-i-\theta}^i + \sum_{\{r,j\}, w+w'-k-b(r)} \exp(-E_0(r,j)/RT) Z_{k-w-w'}^i \left( Z_{\{i,j\}_{w+w'}^i, T_{i,j}} \right) \]

where \( E_0(r,j) = -1 \) if positions \( r,j \) pair in sequence \( s \), and otherwise \( E_0(r,j) = +\infty \). Additionally, \( b_0 = 1 \) if \( j \) is base-paired in \( S_{i,j}^* \) and \( 0 \) otherwise, and \( b(r)=d_{BP}(S_{i,j}^*, S_{i-r-1}^* \cup S_{r+1-j}^* \cup \{(r,j)\}) \).

This holds since in a secondary structure \( T_{i,j} \) on \( s_i, \ldots, s_j \) that is a \( k \)-neighbor of \( S_{i,j}^* \), either nucleotide \( j \) is unpaired in \( [i,j] \) or it is paired to a nucleotide \( r \) such that \( i \leq r < j \). In this latter case it is enough to study the smaller sequence segments \([i,r-1] \) and \([r+1,j-1] \) noting that, except for \((r,j)\), base pairs outside of these regions are not allowed, since there are no pseudoknots. In addition, for \( d_{BP}(S_{i,j}^*, T_{i,j}) = k \) to hold, it is necessary for \( w+w'=k-b(r) \) to hold, where \( w = d_{BP}(S_{i,r-1}^*, T_{i,j-1}) \) and \( w' = d_{BP}(S_{r+1-j}^*, T_{r+1-j}) \), since \( b(r) \) is the number of base pairs that differ between \( S_{i,j}^* \) and a structure \( T_{i,j} \) due to the introduction of the base pair \((r,j)\).

**Methods**

Given RNA sequence \( s \) and compatible initial structure \( S^* \), we define the polynomial

\[ Z(x) = \sum_{k=0}^{n} c_k x^k \]  

where coefficients \( c_k = Z_k^i \). Moreover, because of (3) and the fact that the minimum number of unpaired bases in a hairpin loop \( \theta \) is 3, we know that \( c_n = 0 \), so that \( Z(x) \) is a polynomial of degree strictly less than \( n \). If we evaluate the polynomial \( Z(x) \) for \( n \) distinct values

\[ Z(a_i) = y_1, \ldots, Z(a_n) = y_n, \]

then the Lagrange polynomial interpolation formula guarantees that \( Z(x) = \sum_{k=1}^{n} y_k P_k(x) \), where the polynomials \( P_k(x) \) have degree at most \( n-1 \) and are given by the Lagrange formula...
pair distance located adjacent to the phenylalanyl-tRNA synthetase operon leader, with GenBank accession code CP000948.1/1887748-1887820 (complement). The base pair distance to MFE structure is at most 18877820.

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Figure 1. FFTbor output for the RNA attenuator for the phenylalanyl-tRNA synthetase (pheST) operon in E. coli K-12 substr. DH10B, located adjacent to the phenylalanyl-tRNA synthetase operon leader, with GenBank accession code CP000948.1/1887748-1887820 (complement). The y-axis represents base pair distance to the minimum free energy structure \( S' \); x-axis represents Boltzmann probability \( p(k) \) that a structure has base pair distance \( k \) to \( S' \). (Left) Probability \( P(d_{BP}(S,S')=k) \) that base pair distance to MFE structure is \( k \). (Center) Cumulative probability \( P(d_{BP}(S,S') \leq k) \) that base pair distance to MFE structure is at most \( k \). (Right) Finite difference (Derivative) \( P(k \leq d_{BP}(S,S')=k+1) \) of probability that base pair distance to MFE structure is \( k \).

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\[ P_k(x) = \frac{\prod_{i \neq k} (x-x_i)}{\prod (x_k-x_i)}. \] (10)

Since the polynomials \( P_k(x) \) can be explicitly computed, it follows that we can compute the coefficients \( c_k \) of polynomial \( Z(x) \). As we describe below, the evaluation of \( Z(x) \) for a fixed value of \( x \) can be done in time \( O(n^2) \) and space \( O(n^2) \). It follows that the coefficients \( c_k = Z_{1,n} \) can be computed after \( n \) evaluations of \( Z(x) \), where the space for each evaluation of \( Z(x) \) is re-used; hence these evaluations can be performed in time \( O(n^2) \) and space \( O(n^2) \).

Finally, Lagrange interpolation is clearly computable in time \( O(n^2) \). Although this approach is theoretically sound, there are severe numerical stability issues related to the interpolation method [10], the choice of values \( a_1, \ldots, a_n \) in the interpolation, and floating point arithmetic (round-off error) related to the astronomically large values of the partition functions \( Z_{1,n} \) for \( 0 \leq k < n \). After many unsuccessful approaches including scaling (see File S1), we obtained excellent results by interpolating the polynomial \( p(x) \), defined in equation (1), rather than the polynomial \( Z(x) \), defined in equation (9), and performing interpolation with the Fast Fourier Transform (FFT) [11] where \( x_0, \ldots, x_{n-1} \) are chosen to be \( mh \) complex roots of unity, \( z_k = e^{\frac{2\pi i k}{m}} \).

One advantage of the FFT is that interpolation can be performed in \( O(n \log n) \) time, rather than the cubic time required by using the Lagrange formula (10) or by Gaussian elimination. Fewer numerical operations implies increased numerical stability in our application. Details now follow.

Recursions to compute the polynomial \( Z_{i,j}(x) \)

Given an initial secondary structure \( S^o \) of a given RNA sequence \( s \), our goal is to compute

\[ Z_{1,n} = \sum_{S \text{ such that } d_{BP}(S,S^o) = k} e^{-\frac{E_0(S)}{RT}}. \] (11)

where \( S \) can be any structure compatible with \( s \). As previously mentioned, the recurrence relation for RNAbor with respect to the Nussinov energy model \( E_0 \) is

\[ Z_{i,j}^k = Z_{i,j-1}^{k-h_0} + \sum_{x \leq (r<s)} \left( e^{-\frac{E_0(t,r)}{RT}} \sum_{n' \leq k < b(r)} Z_{i,j-1}^{n'} Z_{i+1,j-1}^{h_0} \right) \] (12)

where \( E_0(t,r) = -1 \) if \( r \) and \( j \) can base-pair and otherwise \(+\infty\), and \( h_0 = 1 \) if \( j \) is base paired in \( S^o_{[i,j]} \) and 0 otherwise, and \( b(r) = d_{BP}(S^o_{[i,j]}, S^o_{[i+1,j-1]} \cup \{ (r,j) \}) \).

The following theorem shows that an analogous recursion can be used to compute the polynomial \( Z_{i,j}(x) \) defined by

\[ Z_{i,j}(x) = \sum_{k=0}^n c_k(i,j)x^k \] (13)

where

\[ c_k(i,j) = Z_{i,j}^k \sum_{S \text{ such that } d_{BP}(S,S^o_{[i,j]}), E_0(S) = -1} e^{-\frac{E_0(S)}{RT}}. \]

Here, in the summation, \( S \) runs over structures on \( s_i, \ldots, s_j \), which are \( k \)-neighbors of the restriction \( S^o_{[i,j]} \) of initial structure \( S^o \) to interval \([i,j]\), and \( E_0(S) = -1 \mid S \) denotes the Nussinov-Jacobson energy of \( S \).

**Theorem 1**: Let \( s_i, \ldots, s_n \) be a given RNA sequence. For any integers \( 1 \leq i \leq j \leq n \), let

\[ Z_{i,j}(x) = \sum_{k=0}^n c_k(i,j)x^k \]

where

\[ c_k(i,j) = Z_{i,j}^k. \]

Then for \( i \leq j \leq i+\theta \), \( Z_{i,j}(x) = 1 \) and for \( j > i + \theta \) we have the recurrence relation
\[ Z_{i,j}(x) = Z_{i,j-1}(x) \cdot x^0 + \sum_{i \leq r < j} \left( \frac{-F_0(r)}{R} \cdot Z_{i,j-1}(x) \cdot Z_{r+1,j-1}(x) \cdot x^0 \right). \] (14)

where \( b_0 = 1 \) if \( j \) is base-paired in \( S'_{i,j} \) and 0 otherwise, and

\[ b(r) = d_{B,F}(S'_{i,j} \cup S'_{i,j-1} \cup \{ (i,j) \}). \]

Proof: First, some notation is necessary. Recall that if \( F \) is an arbitrary polynomial [resp. analytic] function, then \( [x^k]F(x) \) denotes the coefficient of \( x^k \) [resp. the \( k \)th Taylor coefficient in the Taylor expansion of \( F \)] for instance, in equation (1), \( [x^4]p(x) = p_4 \), and in equation (9), \( [x^4]Z(x) = c_k(i,j) \).

By definition, it is clear that \( Z_{i,j}(x) = 1 \) if \( i \leq j \leq i + \theta \), where we recall that \( \theta = 3 \) is the minimum number of unpaired bases in a hairpin loop. For \( j > i + \theta \), we have

\[ [x^k]Z_{i,j}(x) = c_k(i,j) = Z_{i,j}^k \]

By induction, the proof of the theorem now follows. \( \square \)

Notice that if one were to compute all terms of the polynomial \( Z_{1,n}(x) \) by explicitly performing polynomial multiplications, then the computation would require \( O(n^2) \) time and \( O(n^2) \) space. Instead of explicitly performing polynomial expansion in variable \( x \), we instantiate \( x \) to a fixed complex number \( z \in \mathbb{C} \), and apply the following recursion for this instantiation:

\[ Z_{i,j}(z) = Z_{i,j-1}(z) \cdot z^0 + \sum_{i \leq r < j} \left( \frac{-F_0(r)}{R} \cdot Z_{i,j-1}(z) \cdot Z_{r+1,j-1}(z) \cdot z^0 \right). \] (15)

In this fashion, we can compute \( Z(z) = Z_{1,n}(z) \) in \( O(n^2) \) time and \( O(n^2) \) space. For \( n \) distinct complex values \( z_0, \ldots, z_{n-1} \), we can compute and save only the values \( Z(z_0), \ldots, Z(z_{n-1}) \), each time re-using the \( O(n^2) \) space for the next computation of \( Z(z_k) \). It follows that the computation resources used to determine the \( \langle \text{column} \rangle \) vector

\[ Y = (y_0, \ldots, y_{n-1})^T = \left( \begin{array}{c} y_0 \\ y_1 \\ \vdots \\ y_{n-1} \end{array} \right) \] (16)

where \( y_0 = Z(z_0), \ldots, y_{n-1} = Z(z_{n-1}) \) is thus quartic time \( O(n^4) \) and quadratic space \( O(n^3) \).

Polynomial interpolation using the FFT

Let \( \omega = e^{2\pi i/n} \) be the principal \( n \)th complex root of unity. Recall that the Vandermonde matrix \( V_n \) is defined to be the \( n \times n \) matrix, whose \( i,j \) entry is \( \omega^{ij} \); i.e.

\[ V_n = \begin{pmatrix} 1 & 1 & 1 & \cdots & 1 \\ 1 & \omega & \omega^2 & \cdots & \omega^{n-1} \\ 1 & \omega^2 & \omega^4 & \cdots & \omega^{2(n-1)} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 1 & \omega^{n-1} & \omega^{2(n-1)} & \cdots & \omega^{(n-1)(n-1)} \end{pmatrix} \]

The Fast Fourier Transform (FFT) is defined to be the \( O(n \log n) \) algorithm to compute the Discrete Fourier Transform (DFT), defined as the matrix product \( Y = V_nA \):

\[ \begin{pmatrix} y_0 \\ y_1 \\ \vdots \\ y_{n-1} \end{pmatrix} = V_n \begin{pmatrix} a_0 \\ a_1 \\ \vdots \\ a_{n-1} \end{pmatrix} \]

On page 837 of [11], it is shown that the \( (i,j) \) entry of \( V_n^{-1} \) is \( \frac{\omega^{-ij}}{n} \) and that

\[ a_j = \frac{1}{n} \sum_{k=0}^{n-1} y_k \omega^{-kj} \] (17)

for \( j = 0, \ldots, n-1 \).

Since we defined \( Y \) in (16) by \( Y = (y_0, \ldots, y_{n-1})^T \), where \( y_0 = Z(z_0), \ldots, y_{n-1} = Z(z_{n-1}) \) and \( z_k = \omega^k \exp\left( \frac{k \pi i}{n} \right) \), it follows that the coefficients \( c_k = Z_{k,n}^2 \) in the polynomial

\( Z(x) = c_0 + c_1 x + \cdots + c_{n-1} x^{n-1} \) defined in (8) can be computed, at least in principle, by using the FFT. It turns out, however, that the values of \( Z_{k,n}^2 \) are so astronomically large that the ensuing numerical instability makes even this approach infeasible for values of \( n \) that exceed 56 (data not shown). Nevertheless, our approach can be modified as follows. Define \( Y \) by
\( Y = (y_1, \ldots, y_n)^T \), where \( y_1 = \frac{Z(s_1)}{Z}, \ldots, y_n = \frac{Z(s_n)}{Z} \) and \( Z \) is the partition function defined in (4). Using the FFT to compute the inverse DFT, it follows from (17) that we can compute the probabilities \( p_0, \ldots, p_{n-1} \) that are coefficients of the polynomial \( p(x) = p_0 + p_1 x + \cdots + p_{n-1} x^{n-1} \) defined in equation (1). For genomics applications, we are only interested in the \( m \) most significant digits of each \( p_k \), as described in the pseudocode below.

**Algorithm for FFTbor**

This pseudocode computes the \( m \) most significant digits of probabilities \( p_k = \frac{Z_k}{Z} \).

**INPUT**: RNA sequence \( s = s_1, \ldots, s_n \), and initial secondary structure \( S' \) of \( s \) and integer \( m \).

**OUTPUT**: Probabilities \( p_k = Z_k / Z \) to \( m \) significant digits for \( k = 0, \ldots, n-1 \).

1. generate roots of unity \( \omega^k \) for \( k = 0, \ldots, n-1 \), where \( \omega = \exp \left( \frac{2 \pi i}{n} \right) \) and \( i = \sqrt{-1} \).
2. note that the partition function \( Z = y_0 = Z(\omega^0) \).
3. for \( k = 0 \) to \( n-1 \)
   4. compute \( y_k = Z(\omega^k) \) using recursion (15)
   5. \( y_k = 10^m \frac{y_k}{Z} \) // normalize \( y_k \)
   6. compute \( P = (p_0, \ldots, p_{n-1})^T \) where \( p_j = \frac{\sum_{k=0}^{n-1} a_k \omega^{-kj}}{n} \) by using FFT in (17)
   7. for \( k = 0 \) to \( n-1 \)
   8. \( p_k = \frac{10^m p_k}{10^m} \) // truncate to \( m \) most significant digits

**Speed-up in our implementation of FFTbor.** In this subsection, we show that we need only evaluate the polynomial \( Z(x) \), as defined in equation (8), for \( n/2 \) of the complex \( m \) roots of unity. It is first necessary to recall the definition of complex conjugate. Recall that the complex conjugate of \( z \) is denoted by \( \overline{z} \); i.e. if \( z = a + bi \) where \( a, b \in \mathbb{R} \) are real numbers and \( i = \sqrt{-1} \), then \( \overline{z} = a - bi \).

**Lemma 1**: If \( Z(x) \) is the complex polynomial defined in equation (8), then for any complex \( m \) root of unity \( \omega \), it is the case that \( Z(\omega) = \overline{Z(\omega)} \). In other words, if \( x \) is a complex \( m \) root of unity of the form \( a + bi \), where \( a, b \in \mathbb{R} \) and \( b > 0 \), and if \( Z(a + bi) = A + Bi \) where \( A, B \in \mathbb{R} \), then it is the case that \( Z(a - bi) = A - Bi \).

**Proof**: Letting \( i = \sqrt{-1} \) if \( \theta = \frac{2\pi}{n} \) then \( \omega = e^{i\theta} = \cos(\theta) + i \sin(\theta) \) is the principal \( m \) complex root of unity, and \( 1 = \omega^0, \ldots, \omega^{m-1} = \omega^{m-1} \) together constitute the complete collection of all \( m \) complex roots of unity – i.e. the \( n \) unique solutions of the equation \( x^n - 1 = 0 \) over the field \( \mathbb{C} \) of complex numbers. Clearly, for any \( 1 \leq r < n \), \( e^{i\theta r} = 1^r e^{i0} = e^{(2\pi/r)0} = e^{(0\pi/r)0} = e^{0(\theta r)0} = e^{0(0\pi)(0\theta)} = e^{0(0\theta)} \). Moreover, if \( x = e^{i\theta} = a + bi \) where \( b > 0 \), then we have \( e^{-i\theta} = a - bi \). It follows that for any \( m \) root of unity of the form \( a + bi \), where \( b > 0 \), the number \( a - bi \) is also an \( m \) root of unity.

Recall that \( Z(x) = \sum_{k=0}^{n-1} c_k x^k \), where \( c_k \in \mathbb{R} \) are real numbers representing the partition function \( Z_k \) over all secondary structures of a given RNA sequence \( s_1, \ldots, s_n \), whose base pair distance from initial structure \( S' \) is \( k \). Thus, in order to prove the lemma, it suffices to show that for all values \( k = 0, \ldots, n-1 \), if \( a + bi \) is a complex \( n \)th root of unity, where \( a, b \in \mathbb{R} \) and \( b > 0 \), and if \( (a + bi)^k = C + Di \) where \( C, D \in \mathbb{R} \), then \( (a - bi)^k = C - Di \). Indeed, we have the following.

\[
(a + bi)^k = \sum_{k=0}^{m} \binom{m}{k} a^{m-k} (-bi)^k
\]

It follows that each term of the form \( a^{m-k}(-bi)^k \), for \( k = 0, \ldots, m \), is the complex conjugate of \( a^{m-k}(-bi)^k \), and thus \( (a + bi)^m \) is the complex conjugate of \( (a - bi)^m \). Since \( Z(a + bi) \) is a sum of terms of the form \( c_k (a + bi)^k \), it follows that \( Z(a - bi) \) is the complex conjugate of \( Z(a + bi) \). This completes the proof of the lemma.

Lemma 1 immediately entails that we need only evaluate \( Z(x) \) on \( n/2 \) many of the complex \( m \) roots of unity – namely, those of the form \( a + bi \), where \( b \geq 0 \). The remaining values of \( Z(x) \) are obtained by taking complex conjugates of the first \( n/2 \) values. This, along with a precomputation of powers of the complex \( m \) roots of unity, leads to an enormous performance speed-up in our implementation of FFTbor.

**Results**

**Applications of FFTbor**

In this section, we consider two applications of FFTbor: (i) correlation between kinetic folding speed and the ruggedness of the energy landscape near the minimum free energy structure, (ii) computational detection of riboswitch expression platform candidates.

**Kinetic folding speed and energy landscape ruggedness.** The output of FFTbor, as shown in Figure 2, is a probability distribution, where the \( x \)-axis represents the base pair distance from an arbitrary, but fixed secondary structure \( S' \), and the \( y \)-axis represents the Boltzmann probability \( p(k) = \frac{Z_k}{Z} \) that a secondary structure has base pair distance \( k \) from \( S' \). Arguably, this probability distribution is an accurate one-dimensional projection of the rugged, high dimensional energy landscape near structure \( S' \), of the sort artistically rendered in the well-known energy landscape depicted in Figure 1 of [12]. In the sequel, we may call the FFTbor probability distribution a structural neighbor profile, or simply structural profile \( S' \). A hypothesis behind theoretical work in biomolecular folding theory in [13] is that kinetic folding
sloths down as the energy landscape becomes more rugged. This is borne out in our computational experiments for RNA using FFTbor, as reported in Figure 2.

We randomly chose two TPP riboswitch aptamers from the seed alignment for Rfam family RF00059. The first sequence has EMBL accession code BX842649.1/277414-277318 and is comprised of the 97 nt sequence ACCUGAGCGU AGGGGU-GUGUG GUGAAGUCAG CGCAGUAA AUAACCUUUUG AAGCGUGUAG AGAUAAGGCU CCGCAGGGA AGCAAGAUA GAAGAU, while the second sequence has EMBL accession code AACY022101973.1/389-487 and is comprised of the 99 nt sequence UUAAGGCUAGG-GUGGACCA AAUUGGCAUG AGGGUUAUA CGAUCGAA UGUAAUAGGG CAAUACCGCU AAUAGGGCUGGUA CUU UGAACCUGAU CCGGUUAAUA CCGGCCGAGGAAUGAUUUU CUCUCAGGC. Rfam consensus and minimum free energy structures for both sequences are depicted in Figure 3. Despite the fact that there is no sequence homology according to pairwise BLAST [14], this figure clearly demonstrates that consensus and minimum free energy structures closely resemble each other, and that the structures of both TPP riboswitch aptamers are quite similar, with the exception of the leftmost hairpin loop [resp. multiloop]. The MFE structures differ from the consensus structures principally by the addition of base pairs not determined by covariation in the Rfam alignment. Indeed, if we let S0,S1 denote the Rfam consensus structure resp. MFE structure for the 97 nt sequence with EMBL accession code BX842649.1/277414-277318, then S0/S1 has 4 base pairs, and S1/S0 has 7 base pairs. If we let T0,T1 denote the Rfam consensus structure resp. MFE structure for the 99 nt sequence with EMBL accession code AACY022101973.1/389-487, then T0/T1 has 1 base pair, and T1/T0 has 5 base pairs.

We ran FFTbor on each of the TPP riboswitch aptamer sequences, with the MFE structure of each sequence taken as the initial structure S* for that sequence. For the first sequence, BX842649.1/277414-277318, the FFTbor output suggests that there are low energy structures at a distance from the MFE structure, which might compete with the MFE structure and hence slow the kinetics of folding. In contrast, for the second sequence, AACY022101973.1/389-487, the FFTbor output suggests that there are no such competing low energy structures, hence the second sequence should fold more quickly than the first.

In order to more systematically determine the relation between kinetic folding speed and the ruggedness of an energy landscape near the MFE structure, we need to numerically quantify ruggedness. To this end, in the following we define the notion of expected base pair distance to a designated structure. Let S* be an arbitrary secondary structure of the RNA sequence s = a1, ..., an. The expected base pair distance to S* is defined by

\[ E[(d_{bp}(S, S^*)) : S \in \mathcal{S}(a_1, \ldots, a_n)] = \sum_{\mathcal{S}} P(S) d_{bp}(S, S^*) \]  

where \( \mathcal{S}(a_1, \ldots, a_n) \) denotes the set of secondary structures for s = a1, ..., an, \( P(S) = \frac{e^{-E(S)/RT}}{Z} \) is the Boltzmann probability of S, and \( d_{bp}(S, S^*) \) denotes base pair distance between S and S*. If we run FFTbor on an input sequence s and secondary structure S*, then clearly \( E[(d_{bp}(S, S^*)) : S \in \mathcal{S}(a_1, \ldots, a_n)] = \sum_k k \cdot p(k) \), where \( p(k) = \frac{Z_k}{Z} \), obtained from the program output. If S* is the empty structure, then FFTbor output is simply the probability distribution of the number of base pairs per secondary structure, taken over the Boltzmann ensemble of all structures.

For the benchmarking assay, we took all 61 selenocysteine insertion sequence (SECS) sequences from the seed alignment of Rfam family RF00031 [23]. Average length was 64.32 ± 2.83 nt. For each sequence, we ran both FFTbor and a Monte Carlo folding algorithm, developed by E. Freyhult and P. Clore (unpublished). Using the Monte Carlo algorithm, we determined the mean first passage time (MFPT), defined as the average taken over 50 runs, of the number of Monte Carlo steps taken to fold the empty structure into the MFE structure, where an absolute upper bound of 5 million steps was allowed in the simulation. After unsuccessful attempts due to ruggedness of the energy landscape near the MFE structure, by using the Hartigan-Hartigan dip test of unimodality [24], expected base pair distance from MFE structure, total variation distance between FFTbor output and the exponential distribution estimated by the method of moments [25], etc., we ran FFTbor when starting from the empty structure (rather than the MFE structure) as initial structure. As mentioned above, in this case, FFTbor output is simply the probability distribution for the number of base pairs per structure, taken over the ensemble of all secondary structure for the input RNA sequence. Surprisingly, we found that there is a significant correlation of 0.4836192 with one-tailed p-value of 0.00018249 between the standard deviation of the FFTbor output (when starting from the empty structure) and logarithm base 10 of the mean first passage time. Table 1 and Figure 4 explain this phenomenon in detail.

In the right panel of Figure 4, we applied FFTbor to each of the two randomly chosen TPP riboswitch aptamers BX842649.1/277414-277318 and AACY022101973.1/389-487, starting from the empty reference structure S' = \emptyset. The mean for the FFTbor
structural profile near the empty structure is $m_1 \sim 23.0203$ [resp. $\mu_1 = 27.5821$], the standard deviation $\sigma$ for the FFTbor structural profile is $\sigma_1 = 2.22528791$ [resp. $\sigma_1 = 1.98565959$], and the Kinfold MFPT is 311,075 [resp. 61,575]. This anecdotal evidence supports the hypothesis that small standard deviation in FFTbor distribution is correlated with fast folding.

Additionally, in following a suggestion of one of the anonymous referees, we randomized the TPP riboswitches BX842649.1/277414-277318 and AACY022101973.1/389-487 by using our implementation of the Altschul-Erikson dinucleotide shuffle algorithm [26], and then applied FFTbor to these sequences, starting from the empty structure. The mean $m_1$ and standard deviation $\sigma_1$ for the FFTbor distribution for randomized BX842649 are respectively $m_1 = 19.93$ and $\sigma_1 = 2.88$, while those for randomized AACY022101973 are $m_2 = 24.39$ and $\sigma_2 = 24.00$. Running Kinfold, with a maximum of 500,000 steps with 30 replicates (as explained in the text), we found that for randomized BX842649, all 30 runs converged yielding a mean first passage time (MFPT) of 13022.58 with standard deviation of 15221.78. In contrast for randomized AACY022101973, only 15 out of 30 runs converged within 500,000 steps, and discounting these nonconvergent data, we obtain an average mean first passage time (MFPT) of 94446.93 with standard deviation of 157107.43. This additional test provides more anecdotal evidence supporting our hypothesis that small standard deviation $\sigma$ in FFTbor probability density is correlated with fast folding, as measured by MFPT.

Riboswitch expression platform prediction. A bacterial riboswitch is a portion of the 5’ untranslated region (UTR) of messenger RNA, that performs gene regulation by undergoing a conformational change upon binding with a ligand, such as thiamine pyrophosphate (TPP).
guanine, thiamine pyrophosphate, lysine, etc. [27]. This conformational change may either turn on or off the corresponding gene by either transcriptional or translational regulation of the messenger RNA [28], depending on the particular riboswitch. The common feature shared by all riboswitches is that a gene is regulated by conformational change upon ligand binding. Bacterial riboswitches are often found upstream of operons, regulating groups of genes, as in purine de novo synthesis and salvage [29].

A riboswitch consists of two equally important parts: an upstream aptamer, capable of highly discriminative binding to a particular ligand, and a downstream expression platform, capable of undergoing a radical conformational change upon binding of a ligand with the discriminating aptamer. Since aptamers have been

**Table 1. Pearson correlation between various aspects of selenocysteine insertion sequences from the seed alignment of Rfam family RF00031 [23].**

|        | μ   | σ   | σ/μ | len | MFE | logMFPT |
|--------|-----|-----|-----|-----|-----|---------|
| μ      | 1   |     |     |     |     |         |
| σ      |     | −0.43722448 |     | 1   |     |         |
| σ/μ   |     |     | −0.69141183 | 0.943650913 |     |         |
| len    |     |     |     | 0.707683898 | −0.158951202 | −0.364591789 | 1 |
| MFE    |     |     |     | −0.569474125 | 0.739515083 | 0.759622716 | −0.368485646 | 1 |
| logMFPT|     |     |     | −0.036291124 | 0.48436192 | 0.376230235 | 0.40565529 | 0.399015556 | 1 |

For each of the 61 RNA sequences, we ran FFTbor, starting from empty initial structure \(S'\), and we ran a Monte Carlo folding algorithm, developed by E. Freyhult and P. Clote (unpublished). Using the Monte Carlo algorithm, we determined the mean first passage time (MFPT), defined as the average taken over 50 Monte Carlo runs per sequence (log base 10 of the standard deviation of number of Monte Carlo steps per run was approximately 9% of log10MFPT on average). The table shows the correlation between each of these aspects. Some correlations are obvious – for example, (i) the standard deviation \(σ\) is highly correlated with the coefficient of variation \(\frac{σ}{μ}\) (ii) the mean \(μ\) is negatively correlated with the coefficient of variation \(\frac{σ}{μ}\) (iii) the mean \(μ\) is negatively correlated with the minimum free energy (MFE) – if most low energy structures in the ensemble have many base pairs, then it is likely that the minimum free energy is very low (i.e. since MFE is negative, the absolute value of MFE increases); (iv) sequence length is negatively correlated with MFE – as sequence length increases, the minimum free energy (MFE) decreases. However, it may appear surprising that (v) the mean \(μ\) number of base pairs per structure is independent of MFPT (correlation −0.036291124), although (vi) MFE is correlated with MFPT (correlation 0.399015556) – i.e. from (iii), lower MFE is correlated with a larger average \(μ\) number of base pairs per structure, from (iv) higher MFE is correlated with longer folding time, but from (v) the average \(μ\) number of base pairs per structure is independent of folding time. The most important insight from this table is that (vii) standard deviation \(σ\) is correlated with mean first passage time – the correlation is statistically significant, with one-tailed \(p\)-value of 0.00018249.
under strong evolutionary pressure to bind with high affinity (e.g. $K_D \approx 5 \text{nM}$ for guanine [30]), there is strong sequence conservation found in the aptameric region of orthologous riboswitches. In contrast, while secondary structure is conserved in the terminator loop of the expression platform in purine riboswitches, there is relatively low sequence conservation (data not shown). While a number of methods exist to computationally predict riboswitch aptamers [31,32,33,34,35] (and especially INFERNAL [36], which latter is used to predict riboswitch aptamers in Rfam), it is an important biological problem to determine the expression platform, since the structure of the expression platform can suggest whether there is transcriptional regulation via a terminator loop or translational regulation via the sequestration of the Shine-Dalgarno sequence [28]. Determination of the precise location and structure of the expression platform is difficult due to low conserved sequence identity (in-house computations, data not shown). Although this problem remains open, we report here how FFTbor may provide help to biologists in the selection and prioritization of riboswitch candidates.

Figure 5 depicts the gene OFF secondary structure of the xpt G-box purine riboswitch in *B. subtilis*; structure taken from that in Figure 1A of [27].

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Figure 6. Graph of the expected distance from target secondary structure, as a function of window offset position in the 5' untranslated region (UTR) of the xpt gene of *B. subtilis*, i.e. GenBank accession code CP002906.1/c2165402-2165042 *B. subtilis* subsp. *subtilis* RO-NN-1. In a moving window application, FFTbor computed the Boltzmann probability $p(k)$ that secondary structures of the current window contents have base pair distance $k$ from the target (or initial) structure $S'$. In each case, the size of the window was set to equal the length of $S'$. (Left) Target structure $S'$ comprises the entire secondary of the xpt riboswitch, as depicted in Figure 5, with the exception that the leading and trailing unpaired positions were removed, as explained in the text – see displayed dot bracket structure in (19). (Center) Target structure $S'$ comprises only the aptamer secondary structure, as displayed in dot bracket structure in (20). (Right) Target structure $S'$ comprises only the expression platform secondary structure, as displayed in dot bracket structure in (21). The number of points displayed on the $x$-axis differs in each case, since the window size differs, as explained above. The very well-defined minimum in each panel corresponds to the exact location of the entire riboswitch (left panel), aptamer (center panel) and expression platform (right panel). Note that the base line value for the expected base pair distance in the left panel (entire riboswitch) is approximately 70, while that for both the center panel (aptamer) and right panel (expression platform) is approximately 35.

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is only partial, since there are regions with no base pairs depicted, despite the fact that additional base pairs could be added. By using blastn, it is found that this 161 nt purine riboswitch can be found on the complement strand of GenBank accession number CP002906.1/c2165302-2165142 in complete genome of B. subtilis subsp. subtilis RO-NN-1. Figure 6 depicts the result of three computational experiments with FFTbor. The left panel displays the expected base pair distance to the following secondary structure

(19)

as a function of window offset, where window size equals the size of this target structure. This structure was obtained by removing all leading and trailing unpaired positions from the structure depicted in Figure 5, except for the leftmost [resp. rightmost] unpaired position adjacent to the leftmost [resp. rightmost] base-paired position. The reason for removal of the leading and trailing unpaired positions was that the structure of [27], depicted in Figure 5, is clearly only partial, as earlier mentioned. The center panel displays the expected base pair distance to the following secondary structure

(20)

as a function of window offset, where window size equals the size of this target aptamer structure. Similarly, the right panel displays the expected base pair distance to the following secondary structure

(21)

as a function of window offset, where window size equals the size of this target expression platform structure. Figure 6 determines the precise location of the xpt riboswitch, both aptamer and expression platform.

If the biologically functional target structure is unknown, one can instead attempt a similar moving window computation, where the target structure is taken to be the minimum free energy structure of the current window contents. In this case, one may hope to determine a bimodal distribution, as displayed in Figure 7. Given an input RNA sequence, or genomic region, the web server http://bioinformatics.bc.edu/clotelab/FFTbor creates a movie as follows, described here for the xpt riboswitch previously discussed. We extended the 161 nt xpt G-box purine riboswitch described in Figure 5, with GenBank accession number CP002906.1/c2165302-2165142, to a sequence of length 200 nt, by appending flanking downstream genomic nucleotides. Running FFTbor on all prefixes of the resulting sequence of lengths 70,72,74, …, 200, we produced a movie, displayed on the webserver http://bioinformatics.bc.edu/clotelab/FFTbor. Figure 5 displays the output of FFTbor on the 166 nt prefix, clearly showing a bimodal distribution. Attempting to automate the identification of non-unimodal FFTbor output, we have applied the Hartigan-Hartigan dip-test [24], implemented in R; however, the dip-test appears to be too sensitive, in that a probability distribution is reported to be non-unimodal, even when visual inspection indicates that it appears overwhelmingly to be unimodal (data not shown). It is for this reason that the web server http://bioinformatics.bc.edu/clotelab/FFTbor produces a movie of prefixes, where the user can start/stop the movie, move forward/backward, or download all raw data output by FFTbor.

Figure 7. The 161 nt xpt G-box purine riboswitch described in Figure 1A of [27], found on the complement strand of GenBank accession number CP002906.1/c2165302-2165142 in complete genome of B. subtilis subsp. subtilis RO-NN-1. We extended this 161 nt sequence to a sequence of length 200 nt, by appending flanking downstream genomic nucleotides. The web site http://bioinformatics.bc.edu/clotelab/FFTbor displays a movie of all prefixes of the resulting 200 nt sequence, where prefix lengths range from 70,72,74, …, 200. doi:10.1371/journal.pone.0050506.g007
Benchmarking results

Total variation distance for density of states. Recall that the total variation distance between two probability distributions $P = \{ p_x : x \in \Omega \}$ and $Q = \{ q_x : x \in \Omega \}$, defined on the same sample space $\Omega$, is defined by

$$\delta(P, Q) = \sum_{x \in \Omega} |p_x - q_x|.$$ 

The density of states for an RNA sequence $s$ with respect to an initial structure $S^*$ of $s$ is defined to be the probability distribution $P = (p_0, \ldots, p_{n-1})$ where $p_k = Z_{k+1}^* Z$. In all our tests, for RNA of length up to 400 nt, we found the total variation distance between

| Len  | 1              | 2              |
|------|----------------|----------------|
| 200  | 123.2 ± 16.2   | 61.8 ± 8.0     |
| 250  | 331.1 ± 27.2   | 166.1 ± 13.7   |
| 300  | 723.4 ± 59.9   | 365.2 ± 30.1   |
| 350  | 1,380.8 ± 95.2 | 698.4 ± 46.9   |
| 400  | 2,239.1 ± 210.9| 1,129.5 ± 104.3|
| 450  | 3,635.0 ± 857.4| 1,980.9 ± 126.5|
| 500  | 5,076.7 ± 1,292.1| 3,389.8 ± 788.4|

Figure 8. Run times in seconds for RNAbor and FFTbor, on random RNA of length 20, 40, 60, ..., 300 in step size of 20 nt. Each algorithm was run with the empty initial structure $S^*$, see rows RNAbor (empty), FFTbor (empty), and with the minimum free energy structure as the initial structure $S^*$, see rows RNAbor (MFE) and FFTbor (MFE). Note that for both RNAbor and FFTbor, the run time increases when $S^*$ is the MFE structure, rather than the empty structure. Notice the radical improvement in the run time of FFTbor over that of RNAbor.

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Figure 9. (Left) Table showing parallel run times in seconds for FFTbor, using OpenMP http://openmp.org/. Column headers 1, 2 indicate the number of cores used in the computational experiment. For each sequence length 200, ..., 500, five random RNAs were generated using equal probability for each nucleotide A,C,G,U. Run time in seconds, plus or minus one standard deviation, are given for a 24-core AMD Opteron 6172 with 2.10 GHz and 64 GB RAM, with only 1 (resp. 2) cores used. (Right) Graph showing parallel run time of FFTbor on an AMD Opteron 6172 with 2.10 GHz and 64 GB RAM, using respectively 1,2,3,4,6,9,12,15,20 cores.

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than 300 nt were not tested, due to sequences of length
were generated, while for values of
from the defining recursions, the algorithmic time complexity of
were both run, each starting with empty initial structure
structure. Each data point in the table comprises the average run
environment, that supports parallelizations of C/C
and flexible multi-platform shared-memory parallel programming
in Figure 9 describes average run time in seconds (± one standard
deviation) for running FFTbor on random RNA of length
sequences of length up to
random sequences of length
values, for each fixed sequence length in Table 2,
with different coefficients of variable x but with approximately
the same exponent of x (data not shown, but easily computable
from data in Table 2).

**Conclusion and Discussion**

In this paper, we have used a dynamic programming
computation to evaluate the polynomial

\[ Z(x) = \sum_{k=0}^{n} c_k x^k \]  

(22)

on the complex \( m \) roots of unity \( 1, e^{2\pi i/n}, \ldots, e^{2\pi i(n-1)/n} \), where the coefficients \( c_k = Z_{1,n}^k \) are equal to the sum of Boltzmann factors over all secondary structures of a given RNA sequence, whose base pair distance to a given initial structure \( S^* \) is \( k \). Recall the definition of polynomial

\[ p(x) = \frac{Z(x)}{Z_{1,n}} = \sum_{k=0}^{n} p_k x^k \]  

(23)

obtained from \( Z(x) \), whose coefficients are Boltzmann probabilities \( p_k = Z_{1,n}^k \) that a secondary structure has base pair distance \( k \) to \( S^* \). By using the fast Fourier transform to compute the inverse
discrete transform, we can approximate to \( m \) decimal places the
coefficients \( p_k = Z_{1,n}^k \) of \( p(x) \), and thus the \( m \) most
significant positions of \( c_k = Z_{1,n}^k = p_k Z_{1,n} \). Interpolation is
performed for \( p(x) \), rather than \( Z(x) \), due to issues concerning
numerical instability. The computational advantage of FFTbor
over its predecessor RNAbor [1] is that the new algorithm runs in
quartic time \( O(n^4) \) and quadratic space \( O(n^2) \), in contrast to the
\( O(n^3) \) run time and \( O(n^2) \) space required by RNAbor. We have
additionaly provided a parallelization of FFTbor using OpenMP
primitives. Additionally, we have described applications of FFTbor
to determine the correlation between kinetic folding speed and the

**Table 2.** Table showing parallel run times of FFTbor, using OpenMP http://openmp.org/. Column headers 2, 3, etc. indicate the number of cores used in the computational experiment.

| Len | 2  | 3  | 4  | 6  | 9  | 12 | 15 | 20 |
|-----|----|----|----|----|----|----|----|----|
| 200 | 61.8 ± 8.0 | 41.6 ± 6.0 | 31.6 ± 4.2 | 21.1 ± 2.9 | 15.0 ± 2.3 | 11.3 ± 1.1 | 9.6 ± 1.3 | 7.6 ± 1.5 |
| 250 | 166.1 ± 13.7 | 111.5 ± 8.7 | 84.1 ± 7.1 | 56.9 ± 4.0 | 38.8 ± 3.8 | 30.3 ± 2.6 | 24.6 ± 1.6 | 18.9 ± 2.7 |
| 300 | 365.2 ± 30.1 | 246.4 ± 20.5 | 184.5 ± 14.9 | 125.1 ± 9.6 | 85.5 ± 6.6 | 64.9 ± 6.5 | 53.6 ± 5.9 | 42.1 ± 5.2 |
| 350 | 698.4 ± 46.9 | 470.1 ± 32.6 | 352.0 ± 23.0 | 242.6 ± 15.8 | 163.3 ± 12.4 | 125.1 ± 6.7 | 104.2 ± 8.9 | 76.2 ± 3.9 |
| 400 | 1,129.3 ± 104.3 | 757.5 ± 68.9 | 571.6 ± 53.9 | 391.2 ± 36.4 | 265.1 ± 24.2 | 207.2 ± 18.4 | 165.6 ± 14.5 | 125.9 ± 14.9 |
| 450 | 1,980.9 ± 126.5 | 1,326.3 ± 85.1 | 1,000.0 ± 59.0 | 688.9 ± 44.8 | 469.2 ± 29.1 | 355.1 ± 25.4 | 289.8 ± 21.2 | 223.1 ± 18.2 |
| 500 | 3,389.8 ± 788.4 | 2,067.9 ± 99.2 | 1,555.0 ± 72.2 | 1,074.3 ± 53.7 | 728.1 ± 40.9 | 548.3 ± 24.0 | 451.5 ± 25.7 | 338.1 ± 22.7 |

For each sequence length 200, . . . , 500, five random RNAs were generated using equal probability for each nucleotide A,C,G,U. Run time in seconds, plus or minus one
standard deviation, are given for a 24-core AMD Opteron 6172 with 2.10 GHz and 64 GB RAM. Least-squares fit of the data indicates a quadratic dependency of run time
on sequence length (despite the obvious \( O(n^4) \) theoretical run time), and a power law dependency of approximately \( x^{-0.49} \) on the number of cores \( x \).

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P, as computed to 6 decimal places by RNAbor and by FFTbor, to
be 0. It follows that FFTbor can reliably be used in place of
RNAbor to determine Boltzmann probabilities \( p(k) = P(d_{BP}(S,S^*) = k) \).

**Run time comparison of RNAbor and FFTbor.** As visible
from the defining recursions, the algorithmic time complexity of
RNAbor is \( O(n^5) \) and space complexity is \( O(n^4) \), where \( n \) is the
length of input RNA sequence. In contrast, the time complexity of
FFTBor is \( O(n^4) \) and space complexity is \( O(n^2) \). Figure 8 displays
run time curves for both RNAbor and FFTbor, when the initial
structure \( S^* \) is taken to be either the empty structure or the
minimum free energy (MFE) structure.

Here, we compare the run time of RNAbor [1] and the
(unparallelized version of) FFTbor, using a Dell Power Edge 1950,
2 x Intel Xeon E5430 Quad core with 2.80 GHz and 16 GB
RAM. For \( n = 20, 40, 60, \ldots, 300 \), in step size of 20 nt, we
generated \( n \) random RNA sequences of length \( n \) with equal
probability for each nucleotide A,C,G,U (i.e. a 0th order Markov
chain). For values of \( n \leq 200 \), random sequences of length \( n \) were
generated, while for values of \( 220 \leq n \leq 300 \), only 10
sequences of length \( n \) were generated. RNA sequences larger
than 300 nt were not tested, due to \( O(n^5) \) memory constraints
required by RNAbor. For each RNA sequence, RNAbor and
FFTBor were both run, each starting with empty initial structure
\( S^* \), and also with initial structure \( S^* \) taken to be the MFE
structure. Each data point in the table comprises the average run
time for three independent evaluations.

**OpenMP parallelization of FFTbor.** OpenMP is a simple
and flexible multi-platform shared-memory parallel programming
environment, that supports parallelizations of C/C++ code – see
http://openmp.org/. Using OpenMP primitives, we created
multiple threads to evaluate the polynomial \( Z(x) \) on different
complex \( m \) roots of unity. The table in the left panel of Figure 9
and Table 2 together present benchmarks, executed on a 24-core
AMD Opteron 6172 with 2.10 GHz and 64 GB RAM, for the
speedup of FFTbor as a function of the number of cores. The table
in Figure 9 describes average run time in seconds (± one standard
deviation) for running FFTbor on random RNA of length
200,250,300,400,450,500 with either 1 or 2 cores. Table 2
presents similar data for running FFTbor on 2,3,6,4,12,15,20
cores. Although FFTbor clearly has quartic \( O(n^4) \) run time as a
function of RNA sequence length, least-squares fit of run times
from Table 2 instead shows a quadratic run time for RNA
sequences of length up to 500 nt. There appears to be a power law
dependence of FFTbor speedup, as a function of number of cores.
ruggedness of the energy landscape, and to predict the location of riboswitch expression platform candidates.

It is important to point out that the algorithm and software RNAbor is more precise than that of FFTbor – in particular, RNAbor not only computes the partition function values \( Z_{1,\ldots,n}^k \) for all \( 0 \leq k \leq n \), but as well as computes the structures \( S_k \), defined to be the minimum free energy structure over all \( k \)-neighbors of initial structure \( S_k \). In contrast, FFTbor only computes the \( m \) most significant digits of the probabilities \( p_k = Z_{1,\ldots,n}^k / Z_n \), for \( 0 \leq k \leq n \), where by multiplication of \( p_k \) by the partition function \( Z = Z_{1,\ldots,n} \), one obtains an approximation of the partition function values \( Z_{1,\ldots,n}^k \).

There is no possibility that FFTbor can compute the structures \( S_k \), nor can at present we see how to use FFTbor to sample structures from the Boltzmann ensemble of structures having base pair distance \( k \) from \( S_k \).

In [37,38], we introduced the a related parametric RNA structure algorithm, RNAmutants, which computes the partition function \( Z_{1,\ldots,n}^k \) and minimum free energy structure \( MFE(k) \) over all secondary structures of all \( k \)-point mutants of a given RNA sequence \( s = s_1, \ldots, s_n \). In [39], RNAmutants was extended to sample low energy structures over \( k \)-point mutants within a certain range of GC-content. Some of the ideas in [39] foreshadowed the results of this paper, and in the future, we intent to apply interpolation and the FFT to similarly provide a more efficient version of RNAmutants. Nevertheless, this future, more efficient version will be incapable of efficiently sampling low energy structures over \( k \)-point mutants, analogous to the current differences between RNAbor and FFTbor.

### Supporting Information

**File S1 Supplementary information.**

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FFTbor depends heavily on the use of the Fast Fourier Transform implementation FFTW of Frigo and Johnson [40] at http://www.fftw.org/.

### Author Contributions

Conceived and designed the experiments: PC YP. Performed the experiments: PC ES SS. Analyzed the data: PC ES. Contributed reagents/materials/analysis tools: PC ID ES SS. Wrote the paper: PC. Interpolated ideation: YP. Idea for handling numerical instability (FFT, scaling): PC. Designed the software used in analysis: PC ES SS.

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