Force-induced denaturation of RNA

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We describe quantitatively a RNA molecule under the influence of an external force exerted at its two ends as in a typical single-molecule experiment. Our calculation incorporates the interactions between nucleotides by using the experimentally-determined free energy rules for RNA secondary structure and models the polymeric properties of the exterior single-stranded regions explicitly as elastic freely-jointed chains. We find that in spite of complicated secondary structures, force-extension curves are typically smooth in quasi-equilibrium. We identify and characterize two sequence/structure-dependent mechanisms that, in addition to the sequence-independent entropic elasticity of the exterior single-stranded regions, are responsible for the smoothness. These involve compensation between different structural elements on which the external force acts simultaneously, and contribution of suboptimal structures, respectively. We estimate how many features a force-extension curve recorded in non-equilibrium, where the pulling proceeds faster than rearrangements in the secondary structure of the molecule, could show in principle. Our software is available to the public through a ‘RNA-pulling server’.

I. INTRODUCTION

In recent years, single-molecule experiments employing optical tweezers, atomic force microscopy, and other techniques have successfully probed basic physical properties of biomolecules through the application of forces in the pN range (see, e.g., Bockelmann et al. (1997); Essevaz-Roulet et al., (1997); Mehta et al. (1999); Reid et al. (1997, 1998); Smith et al. (1996); Yang (2000)). Both, simple elastic properties of the polymers (such as persistence length and longitudinal elasticity) and structural transitions (e.g. unfolding of protein domains) were characterized by recording and analyzing force-extension curves (FEC’s). For nucleic acids, a prominent experiment of the latter type is the ‘unzipping’ of double-stranded DNA (Bockelmann et al., 1997; Essevaz-Roulet et al., 1997). The resulting FEC’s display clear sequence-specific features (e.g. local maxima), which may be attributed to small regions of the sequence that are more strongly bound than their neighbors (Essevaz-Roulet et al., 1997; Lubensky and Nelson, 2000; Thompson and Siggia, 1995). In contrast, long single-stranded DNA, which, like RNA, may fold into complicated branched structures by forming intra-strand basepairs, showed extremely smooth FEC’s in a very recent experiment by Maier et al. (2000). Thus, depending on its structure, DNA may show a broad range of FEC’s from very rugged to completely featureless. However, it is unclear how quantitatively the structure determines the outcome of the FEC measurement.

Here, we address this question theoretically, focusing on the case of RNA and restricting ourselves to secondary structure (i.e. basepairing patterns only instead of full, tertiary structure). In this context, RNA seems to be a more interesting object than DNA, since RNA naturally occurs in many different and functionally important structures, while DNA is primarily found as a double strand. One may hope that pulling experiments generate new insights into the RNA folding problem (Tinoco and Bustamante, 1999), including the folding pathways (Chen and Dill, 2000; Isambert and Siggia, 2000; Thirumalai and Woodson, 2000). Also, force-induced denaturation of RNA is currently studied experimentally (C. Bustamante and I. Tinoco, private communication). The limitation to secondary structure allows us to draw upon the experimentally determined ‘free energy rules’ for RNA secondary structure (Freier et al., 1986; Mathews et al., 1999; Walter et al., 1994), which yield minimum free energy structures that agree reasonably well with experimentally and phylogenetically determined ones (Mathews et al., 1999). Furthermore, it permits us to employ and extend the efficient dynamic programming algorithms (Hofacker et al., 1994; McCaskill, 1990; Zuker and Stiegler, 1981) which can compute the exact partition function (including all possible secondary structures) and reconstruct the minimal free energy structures in polynomial time. Experimentally, the secondary structures may be probed in specific ionic conditions (e.g., those with only monovalent ions) such that the tertiary contacts are strongly disfavored (due to electrostatic repulsion of the sugar-phosphate backbone) (Tinoco and Bustamante, 1999).

The type of experiment that we consider is sketched in Fig. 1. The distance \( R \) between the two ends of an RNA molecule is held fixed, e.g., by attaching them to two beads whose positions are controlled by optical tweezers, and the force \( f \) acting on the beads is recorded as a function of \( R \). As long as the external change in force/extension is applied at a much slower time scale than that of structural transitions of the molecule, the equilibrium FEC is measured. In the main part of the present article, we assume that this is always the case. Experimentally, this condition is usually checked by retracing the FEC (e.g., a hysteresis effect is a clear sign of a non-equilibrium situation).

Besides the above-mentioned free energy parameters for RNA secondary structure, we need a polymer model for single-stranded RNA as input in order to make quan-
The measurement of fluctuations about the equilibrium is therefore not very useful. More promising options include the measurement of the fluctuations about the equilibrium and non-equilibrium FEC’s, where the pulling proceeds faster than (some of) the rearrangements in the structure. While the present approach is extended readily to include equilibrium fluctuations (Gerland, U., R. Bundschuh, and T. Hwa, in preparation), a quantitative treatment of the dynamics of force-induced denaturation of RNA presents a challenge to theoreticians.

The organization of the paper is as follows. In the next section, we explain the details of our model and the way we calculate the FEC’s. Readers interested in the results only should directly proceed to section III. The discussion in section IV explores the possibility of using experimental FEC’s of appropriately designed sequences as an alternative way to determine the RNA free energy parameters. In addition, we estimate to what extent features may be expected in non-equilibrium FEC’s.

II. MODEL AND METHODS

We assume that the force \( f(R) \) acting on the beads (see Fig. 1) is measured as a function of the fixed distance \( R = |\mathbf{R}| \), where \( \mathbf{R} \) denotes the end-to-end vector of the RNA molecule, and that \( R \) is varied very slowly so that thermal equilibrium is always maintained. In practice, the force measurement requires a device acting as a spring, hence the distance cannot be kept exactly constant. However, we consider the situation where the stiffness of this spring is much higher than that of the single-stranded RNA, which has already been pulled out. This condition could only be violated in the very early part of the pulling experiment, which is not the focus of the present investigation. We may therefore neglect the presence of the spring altogether, which amounts to working in the ‘fixed-distance ensemble’\(^1\). Another difference between our model and actual experiments is that we neglect the presence of additional spacer sequences, which are used to connect the RNA molecule to the force-measuring device (e.g. the beads). Again, we assume that they are stiffer than the liberated single-stranded RNA, since we are interested in the size of the features in the FEC, which are observable in an ideal measurement.

The partition function at fixed extension, \( Z_N(R) \), for a given RNA sequence consisting of \( N \) nucleotides, may be written as a sum over the number \( m \) of exterior open bases (as represented by open circles in Fig. 1). For each \( m \), the secondary structure contributes a factor \( Q_N(m) \).

\(^1\) In the ‘fixed-distance ensemble’, only the average force is well-defined, whereas the fluctuations about the average diverge. This reflects the fact that it takes increasingly higher forces to compensate thermal fluctuations on shorter and shorter timescales, in order to keep the extension exactly fixed. Therefore, if one is interested in the fluctuations (of either the force or the extension), the external spring should not be neglected, which would amount to working in a mixed ensemble between ‘fixed-distance’ and ‘fixed-force’.
to the partition function, according to the free energy rules for RNA/DNA secondary structure to be detailed shortly below. This contribution needs to be weighted by the probability \( W(R; m) \) that the chain of \( m \) exterior open bases has end-to-end vector \( R \), given by an appropriate polymer model for the single strand. Together, they yield

\[
Z_N(R) = \sum_{m} Q_N(m) W(R; m)
\]

(1)

The normalization \( \int d^3 R W(R; m) = 1 \) assures that the integral of \( Z_N(R) \) over space yields the usual partition function \( Z_N \) for \( N \) nucleotides without any external constraints. Eq. (1) clearly separates the contribution of the secondary structure, which is entirely contained in \( Q_N(m) \), from the contribution of the exterior single strand contained in \( W(R; m) \). Note that the polymer properties of the interior single strands (i.e. the single strands not subject to the external force) are contained in \( Q_N(m) \) through the loop-entropy parameters, which are part of the free energy rules derived from experiments (see Walter et al. (1994) and references therein).

**Secondary structure.** The number of possible secondary structures for a given sequence of length \( N \) grows exponentially with \( N \). To each structure \( S \), a Boltzmann weight \( \zeta(S) \) may be assigned with the help of the free energy rules (Walter et al., 1994) which contain a large number of experimentally determined energy and enthalpy parameters, e.g., those for the stacking of basepairs, formation of internal, hairpin, bulge or multi-loops, and dangling ends. Due to the large number of possible structures, the full partition function \( Z_N = \sum_S \zeta(S) \) is impossible to evaluate by enumeration, except for very small \( N \). However, one can make use of recursion relations that express the partition function for a subsequence with the help of the partition functions for even shorter subsequences (McCaskill, 1990; Zuker and Stiegler, 1981), and proceed to compute the full partition function exactly in \( O(N^3) \) time. These recursion relations owe their existence to the fact that the class of secondary structures was defined to include only nested structures, e.g. two basepairs \( (i, j) \) and \( (k, l) \) with \( i < k < j < l \) are not admitted (the occurrence of such pairings is called a pseudoknot and contributes relatively little to the free energy of natural RNAs (Tinoco and Bustamante, 1999)). One implementation of this algorithm with very detailed free energy rules is the ‘Vienna package’ (Hofacker et al., 1994), publically available at [http://www.tbi.univie.ac.at/]. In the following, we describe the modifications that we made to this package in order to obtain \( Q_N(m) \) and the corresponding minimum free energy structures.

The Vienna package calculates the auxiliary partition function \( \Pi(i, j) \) for the substrand (i.e., a contiguous segment of the sequence) from base \( i \) to base \( j \), under the condition that base \( i \) and base \( j \) are paired. These quantities can be used to calculate the partition function \( Q(j; n) \) of the substrand from base 1 to base \( j \), under the condition that the exterior part of the configurations is \( 0 \leq n \leq j \) bases long. The recursion formula for \( Q \) is

\[
Q(j+1; n) = Q(j; n-1) + \sum_{i=n-\Delta+1}^{j} Q(i-1; n-\Delta) \Pi(i, j+1),
\]

obtained by splitting the partition function \( Q(j+1; n) \) up according to all possible binding partners of base \( j+1 \). This formula, together with the appropriate boundary conditions for \( j = 0 \) and \( n = 0 \), can be solved recursively by calculating \( Q(j; n) \) first for all \( n \) at a given \( j \) and then for increasing \( j \). In the end, we have \( Q_N(m) = Q(N; m) \) for the \( m \) exterior bases in \( O(N^3) \) time.

To produce the minimum free energy structures at fixed \( m \), we use an equivalent recursive scheme, but replacing the summations by maxima to obtain first the minimum free energy (Zuker and Stiegler, 1981). Then, we determine the corresponding structure by going through the scheme in reverse and reconstructing at each step which of the terms was maximal.

**Polymer model.** The simplest polymer model for the exterior single strand (the open circles in Fig. 1) is the Gaussian chain (de Gennes, 1979). However, as shown below, the force-induced denaturation of RNA occurs at forces of order 10 pN, where the exterior single strand is strongly stretched and the Gaussian model breaks down. In this regime, an elastic freely jointed chain (EFJC) model\(^2\) yields a good fit to experimental FEC’s (Montanari and Mézard, 2000; Smith et al., 1996).

The distance along the backbone between two adjacent nucleotides is the segment length of the chain. We denote it by \( l \) and assign an elastic energy \( V(r) = \frac{C}{2} (r - l)^2 \) per segment, where \( r \) represents the end-to-end vector of the segment. Instead of attempting (the very cumbersome) exact computation of the end-to-end vector distribution \( W(R; m) \) of the chain, we employ an asymptotic expression that becomes exact in the limit of large \( m \) and is sufficiently accurate for our purposes even for small \( m \). It can be derived along the line of a similar calculation for the case of the regular (i.e. non-elastic) freely jointed chain given in (Flory, 1967).

The result is conveniently expressed in terms of the quantity \( q(h) = \int d^3 r e^{-h r} r^{-1} / \int d^3 r e^{-V(r)/k_B T} \), where \( k_B \) denotes the Boltzmann constant and \( h \) is a vector of length \( h \) with fixed (but arbitrary) orientation in space. The asymptotic expression is then

\[
W(R; m) \approx C \frac{h}{2 \pi R} \left[ q(h) \right]^m e^{-h R},
\]

(2)

where \( C \) is a normalization constant and \( h \) is determined from \( R = m \frac{h}{2m} \log q(h) \). We incorporate the effect of

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\(^2\) Here, the constant \( \Delta = 3 \) accounts for the fact that each stem branching from the exterior single strand contributes an additional segment, whose length is approximately equal to the length of three single stranded bases.

\(^3\) Self avoidance in the exterior single strand may be neglected, again because of its highly stretched state.
FIG. 2 (a) Force-extension curve (FEC) for a group I intron (solid line, see text for details) and a homopolymeric RNA of the same length, N = 251 (dashed). The depicted secondary structure is the minimum free energy structure at R = 10 nm. (b) FEC for a hairpin composed of randomly chosen basepairs (solid) and a homogenous hairpin of AU-basepairs (dashed). In both cases the total sequence length is the same length, N = 251 (dashed). The depicted secondary structure is the minimum free energy structure at temperature, as supplied with the Vienna package. The salt concentrations at which these free energy parameters were measured are [Na\(^+\)] = 1M and [Mg\(^{++}\)] = 0M.

III. RESULTS

Fig. 2a and b show the FEC’s (solid lines) for two RNA sequences with practically the same total length and composition, both computed as described in the last section using the same set of parameters. Strikingly, the first curve is almost completely smooth with no significant features, while the second is extremely jagged with large ‘jumps’ in the force. This dissemblance is entirely due to the difference between the secondary structures into which the two sequences fold. The sequence in Fig. 2a originates from the group I intron of the methionine tRNA of Scytomena hofmannii with a sequence length of N = 251 (GenBank# U10481). Its dominant secondary structure (according to our algorithm\(^1\)) at an extension of R = 10 nm is also depicted in Fig. 2a. The sequence in Fig. 2b was artificially generated by concatenating a randomly chosen sequence with its reverse complement, so that it folds into a single hairpin composed of random basepairs. Its FEC is very similar to the experimental force curve obtained upon unzipping double stranded DNA by Essevaz-Roulet et al. (1997); the sawtooth-like oscillations correspond to a ‘molecular stick-slip process’ (Bockelmann et al., 1997). Why does the group I intron not display an abundance of features in the FEC like the hairpin does? Its secondary structure consists of many structural elements (e.g. stem-loop structures), the opening of which one might expect to produce clear signatures in the FEC.

\(^{1}\) The known native secondary structure of this sequence contains two helical regions forming a pseudoknot. Since pseudoknots are excluded from our approach (as explained above), we removed it from the structure computationally by replacing 6 basepairs in the less stable of the two helical regions (positions 79–84 and 157–162) by artificial bases which are excluded from base pairing. With this modification, the minimum free energy structure at zero force (as determined by our algorithm\(^1\)) is almost identical with the secondary structure known from comparative sequence analysis (Patell et al., 2000), available at [http://www.rna.icmb.utexas.edu](http://www.rna.icmb.utexas.edu) outside of the pseudoknot region. Beyond the distance at which the pseudoknot is pulled apart, our modification of the sequence should not effect the FEC significantly. This expectation is supported by our numerical observation that the FEC’s for the unmodified sequence (ignoring the pseudoknot) and for our modified sequence become close to identical beyond a distance of R ≈ 70 nm.
Indeed, in their theoretical study of force-induced denaturation of DNA/RNA, Thompson and Siggia (1995) concluded that the opening of individual basepairs in double stranded DNA cannot readily be observed, but the opening of stem-loop structures in RNA should be.

One fairly obvious effect that could cause the smooth FEC is thermal superposition of alternative secondary structures. Since one may expect that typical RNA structures (such as the one depicted in Fig. 2a) are less well-designed than a perfect hairpin, force-induced denaturation should make more alternative structures accessible in the former case than in the latter. In our analysis below, we find that this effect is indeed non-negligible, but the largest loss of features originates from another, more subtle mechanism, which we call the ‘compensation effect’, and which persists even when no alternative secondary structures are allowed. The compensation effect depends on the fact that when several structural elements are pulled at in parallel, the optimization process that determines the minimum free energy structure with a given number $m$ of external open bases may reclose stretches of basepairs which had already been opened at a lower value $m' < m$.

In our approach (see ‘Model and Methods’ above), the information on the secondary structure energetics for a given sequence is entirely contained in the function $Q(m)$. With the help of the polymer model (contained in $W(R, m)$) this information is translated into a FEC via Eq. (4) (a). Our investigation therefore comprises two steps. First, we seek to understand what property of $Q(m)$ determines the size of the fluctuations in the FEC, and second, how this property depends on the secondary structure.

The first question is addressed most readily for the special case of the random hairpin of Fig. 2b. It is known that in the fixed-force ensemble, unzipping of a random hairpin may be mapped onto the problem of a particle in a tilted one-dimensional random potential (de Gennes, 1979; Lubensky and Nelson, 2000). The random potential is correlated and has the statistical properties of a one-dimensional random walk. In the fixed-distance ensemble, we may perform a very similar mapping (see Fig. 3). Here, the bias for the direction of movement of the particle is not caused by a tilt of the potential, but instead by a spring that is attached to the particle. The position of the other end of the spring is externally controlled, i.e. it is determined by $R$, the given end-to-end distance of the RNA molecule.

In the following, we review the relation between the parameters of the particle-in-a-random-potential problem, i.e. the spring constant $\gamma$ and the variance of the random potential, and the parameters of the unzipping problem. This will also serve us to introduce our notation for the subsequent discussion. We may write the free energy $G(m) = -k_B T \log Q(m)$ of the random hairpin as $G(m) = -\sum_{i=1}^{N-m} \eta(i)$, where the $\eta(i)$ are random with mean $\langle \eta \rangle = \varepsilon$ and variance $\langle \eta(i)\eta(j) \rangle = \langle \eta \rangle^2 = \delta_{ij}(\Delta \varepsilon)^2$. Here, $\varepsilon$ represents the mean binding energy per base, which depends on the GC-content of the hairpin, the temperature, and the salt concentrations, and $\Delta \varepsilon$ measures the fluctuations of $\varepsilon$, both along a given hairpin and between different realizations of the random sequence. The difference between two free energies that are apart, $\Delta G(\ell) = G(m) - G(m-\ell)$, then has the variance

$$\text{var}(\Delta G(\ell)) = \ell (\Delta \varepsilon)^2 .$$

In the particle picture (see Fig. 3), $m - \ell$ corresponds to the position of the particle, and $m$ to the position of the other end of the spring. For fixed $m$, the particle therefore sees the effective potential

$$\Delta G(\ell) + \frac{\gamma}{2} \varepsilon^2 ,$$

i.e. Eq. (4) determines the variance of the random potential. The spring constant $\gamma$ is determined by $\varepsilon$ as follows. If $\Delta \varepsilon$ were zero, the unzipping force would take a constant value $f_0$ (cf. the dashed line in Fig. 2b, which shows the FEC of a homogeneous AU-hairpin). The dependence of $f_0$ on $\varepsilon$ can be calculated analytically by evaluating the sum in Eq. (4) by the saddle point method (see also (D.K. Lubensky and D.R. Nelson, in preparation)). The result is shown in Fig. 4 (solid line). Now $\gamma = f^2 \Gamma$, where $\Gamma$ is the local spring constant of a non-binding RNA of $m$ bases at force $f_0$. Since the spring constant of a homopolymer scales with the inverse of the number of segments, we write $\Gamma = \Gamma_0/m$, where $\Gamma_0$ depends only on $f_0$, but not on $m$. Graphically, $\Gamma_0(f_0)$ is the slope at $f = f_0$ of the dashed line in Fig. 2b (FEC of a homopolymeric RNA), multiplied by 251 (the number of bases in that example). In this way $\Gamma_0(f_0)$ may also be determined from an experimental FEC.

5 For these mappings, alternative structures of the hairpin sequence are neglected, which is a good approximation due to the perfect design of the hairpin. Also, the nearest-neighbour correlations in the random potential caused by the stacking energies are not taken into account, since they would not change the qualitative predictions of the model.
FIG. 4 Threshold force $f_0$ for unzipping of a homogeneous hairpin as a function of the binding energy per base $\varepsilon$ (solid line). The dashed line indicates the Gaussian approximation $f_0 = (6k_BT\varepsilon/l^2b)^{1/2}$, which is obtained by using the end-to-end distance distribution $W(R;m)$ of a Gaussian chain. Note that the Gaussian approximation breaks down already at low forces, and the more detailed treatment according to Eq. (2) is necessary. The stacking energy for AU-pairs at $T = 20^\circ$C is $2\varepsilon \approx 1.21$ kcal/mol corresponding to a threshold force $f_0 \approx 11$ pN, which agrees with the value observed in Fig. 2b (dashed line).

When the fluctuations in the random potential are not too weak, the particle follows the other end of the spring in discrete jumps. The typical size of a jump, $\Delta \ell_{\text{jump}}$, is given by the value of $\ell$ for which the two terms in Eq. (5) are of equal size, $\Delta \ell_{\text{jump}} \simeq (2m \Delta \varepsilon/\ell^2 \Gamma_0)^{2/3}$. A typical jump then leads to a drop in the force by $\delta f \simeq \Gamma \ell \Delta \ell_{\text{jump}}$, i.e.

$$\delta f \simeq (4 \Gamma_0 \Delta \varepsilon^2/R)^{1/3}.$$

This is valid as long as the thermal broadening of the particle position, $\Delta \ell_T \simeq (2m/\ell^2 \Gamma_0)^{1/2}$, is less than the typical jump size $\Delta \ell_{\text{jump}}$. In the opposite case, the particle is sliding more or less smoothly, and $\delta f \propto \Delta \varepsilon$.

Eq. (3) furnishes an estimate for the size of the fluctuations in the FEC for the case of a random hairpin. However, since we used an arbitrary function $G(m)$ as input, the above argument may be made in general for any structure, as long as Eq. (4) holds sufficiently well. Alternatively, if for a particular structure the dependence of $\text{var}(\Delta G(\ell))$ on $\ell$ is determined numerically, this could be used to replace Eq. (4) and Eq. (5) would have to be modified accordingly.

We now address the question of how the fluctuations in $G(m)$ depend on the secondary structure. An essential difference between unzipping of a hairpin and force-induced denaturation of a typical RNA structure is that in the latter case, several stems are being pulled on simultaneously for most of the extension interval (see Fig. 2b), which shows the number of stems as a function of the extension for the group I intron studied above). To analyze the effect of multiple stems, we constructed artificial sequences that form a given number $n$ of random hairpins in a row (i.e. the sequences are a concatenation of $n$ random hairpin sequences, each of which is constructed as explained above). For each $n$ in the range $1 \leq n < 10$, we computed $G(m)$ and the FEC’s for 1000 different sequence realizations, all with an approximate total length of $N = 1000$. As an example, Fig. 5 shows the FEC’s for three sequences, which fold into $n = 1$, 3, and 8 hairpins, respectively. Clearly, the fluctuations in the force curve decrease with increasing $n$. We obtained $\text{var}(\Delta G(\ell))$ as an average over the 1000 realizations and a small interval of $m$. Some of the resulting curves are shown in Fig. 6.

6 In principle, a situation where several stems are pulled on in parallel can also arise in the process of unzipping a single long hairpin, due to accidental palindromic regions in the single strand which has already been pulled out. However, these non-native interactions have to overcome the energetic advantage of the native single-hairpin interactions, in order for the effect to become relevant. Hence the palindrome needs to be extremely GC-rich. For a single hairpin consisting of random basepairs, we estimated that a non-negligible palindrome would typically occur only in sequences of at least several thousand bases in length, which is beyond the length of the sequences studied here.
Although the dependence of \( \text{var}(\Delta G(\ell)) \) on \( \ell \) is not completely linear, the deviation from linearity over the small range of \( \ell \)-values relevant here (typically, \( 0 \leq \ell \leq 12 \)) is not very large. For the sake of simplicity, we chose to interpret the data with the theory for a linear \( \text{var}(\Delta G(\ell)) \) developed above. To this end, we define an effective \( \Delta \varepsilon \) for each \( n \) from the slope of \( \text{var}(\Delta G(\ell)) \) at \( \ell = 4 \).

Fig. 7 shows that \( \Delta \varepsilon^2 \) decreases monotonically with the number of stems that are being pulled on simultaneously. This decrease is almost entirely due to the compensation effect, which we may intuitively understand as follows. When a single hairpin is being unzipped, the stick-slip process described in (Essevaz-Roulet et al., 1997) is topologically inevitable, since the basepairs have to be opened in the order they occur. A strongly bound region that is followed by a weakly bound one, then always leads to a rise and subsequent drop of the FEC. However, with several hairpins, only the total number of exterior open bases is externally constrained, while the individual hairpins may freely open and reclose basepairs (for equilibrium FEC’s there is no kinetic constraint). Therefore, if in a particular hairpin a strongly bound region is followed by a weakly bound one, both regions can open together and another hairpin can reclose a few basepairs to compensate for the released single-strand. Obviously, with a growing number of hairpins this mechanism will be increasingly effective. Clearly, in the fixed-force ensemble, the compensation effect is equivalent to an average over the FEC’s of the individual hairpins. Moreover, with a large number of hairpins, the fixed-force and the fixed-distance ensembles become equivalent (D.K. Lubensky and D.R. Nelson, in preparation).

To analyze the force fluctuations quantitatively, we calculated the FEC’s for all of the 1000 sequence realizations of the \( n \) parallel hairpins, and defined \( \Delta f(R) \) as the standard deviation of the force at extension \( R \) (the so-defined \( \Delta f \) is smaller than the typical size of a force jump, \( \delta f \), but should have the same scaling behavior). Fig. 8 shows a plot of the force fluctuations against the free energy fluctuations, where the horizontal axis, \( \Delta \varepsilon (2\beta^2 R/\Gamma_0)^{1/4} = (\Delta \ell_{\text{jump}}/\Delta \ell_{\text{T}})^{3/2} \), is scaled such that it separates the jumping regime from the sliding regime at a crossover value of one. The vertical axis is scaled such that the data should collapse onto a straight line in the jumping regime according to Eq. (6). In order to guide the eye, Fig. 8 also displays artificial data (crosses) for which \( G(m) \) was generated by drawing random numbers \( \eta(i) \) and taking \( G(m) = -\sum_{i=1}^{N-n} \eta(i) \) (the different points are for different values for the mean and variance of \( \eta(i) \)). The circles mark the data points for
the parallel hairpins and the rectangular symbol in the lower left indicates in what region the group I intron is situated. 

For the artificial data (crosses) the above scaling arguments should apply rigorously. Indeed, the artificial data falls onto a straight line in the jumping regime (the solid line represents a linear fit to the points with abscissae larger than two) and in the sliding regime $\Delta f$ is proportional to $\Delta \epsilon$ (not shown). For the real data, Fig. 8 shows that passing from a single hairpin through structures with several parallel perfect hairpins to a typical natural RNA may be viewed as passing from the jumping regime to the sliding regime for a particle in a (correlated) random potential. At the same time, the FEC's change from jagged to smooth.

As mentioned above, thermal superposition of alternative secondary structures also contributes to the smoothening of the FEC's: since the structural elements in each suboptimal structure open at different values of $m$, the thermal average over all these structures smooths $G(m)$. In order to assess the importance of this effect, we suppressed it by taking only the minimum free energy secondary structures into account instead of calculating the full partition function $Q(m)$. For the group I intron, the FEC without the contribution of suboptimal structures is shown in Fig. 9a. Compared to the full thermodynamic curve (shown in Fig. 9a), some structure is gained, but not nearly as much as in the FEC for the random hairpin of the same length, Fig. 9b. This indicates that the compensation effect is the dominant source for the smoothing of the FEC.

IV. DISCUSSION

In the last section, we found that the equilibrium FEC's for typical RNA molecules (like the group I intron that served us as an example) are quite smooth and do not reveal any features that can be associated with the opening of structural elements. The compensation effect is the primary cause for this result, and we expect it to be responsible, in part, also for the experimental observation of extremely smooth FEC's for single-stranded DNA by Maier et al. (2000). Nevertheless, the measurement of equilibrium FEC's for RNA or single-stranded DNA might still be useful, e.g. for an experimental determination of the RNA/DNA free energy parameters. Usually, these are extracted from melting curves of oligomers Freier et al., 1989, which requires variation of the temperature away from the melting point of the oligomers, where the free energy and its temperature derivative are determined. The free energy parameters at the temperature of interest are then obtained by extrapolation, which introduces an error inherent to the method. For pulling experiments, the temperature can be kept constant at the value of interest, which is an obvious advantage. Here, the limiting factor is only the precision of the force measurement. The quantitative relationship between stacking energy and threshold force expressed by Fig. 6 furnishes the necessary link between force and energy. Measuring FEC's for periodic hairpins composed of different building blocks, would lead to curves like the dashed line in Fig. 3 with different values for the threshold force. From these values, the stacking energies could then be determined, which might lead to more accurate parameters at the desired temperature and salt concentrations.

There are (at least) two options to obtain FEC's with more features, which in turn might allow one to obtain information on RNA secondary structure from pulling experiments. One could either record non-equilibrium FEC's or analyze the fluctuations around the equilibrium curve. For our theoretical investigation, the latter option is not available as long as we work in the fixed-distance ensemble, since the force fluctuations around the thermodynamic average diverge in that ensemble. We will pursue this option in a separate publication by working in a mixed ensemble (Gerland, U., R. Bundschuh, and T. Hwa, in preparation). Here, we briefly consider non-equilibrium FEC's, where the rate of external increase in the force/extension is higher than (some of) the rates

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7 The rectangular area marks the range of points that we obtained by determining $\delta f$, $\Delta \epsilon$, and $\Gamma_0$ by averaging over different extension intervals, all within the range 50–110 nm, which is a region where the mean force is relatively constant (this is required in order to separate fluctuations in the force from a gradual change in the mean value).
FIG. 10 Sketch of the assumed pathway for the formation of a stem-loop structure in the presence of a stretching force $f$. A generalized reaction coordinate $x$ is plotted along the horizontal axis and the free energy $G$ along the vertical axis. The work that has to be exerted against the force in order to pull in the single strand needed for the formation of the stem-loop structure is denoted by $\Delta W$. In principle, the entropy difference between the random coil state on the left and the transition state also contributes to the barrier height, however, we assume that at typical stretching forces it is negligible compared to $\Delta W$.

associated with internal rearrangements in the secondary structure. In the case of long proteins, either naturally occurring as an array of globular domains (Rief et al., 1998) or synthesized protein arrays (Yang, 2000), mechanical stretching experiments resolved the unfolding of up to 20 individual domains. These experiments were performed under non-equilibrium conditions (Rief et al., 1998) with typical pulling speeds of $1 \mu$m/s.

In order to estimate whether non-equilibrium conditions are attainable for RNA with reasonable pulling speeds, we need a rough idea of the timescales involved in secondary structure rearrangements of RNA. For this, we again assume that RNA and single-stranded DNA behave similarly, so that we may draw on an experiment by Bonnet, Krichevsky, and Libchaber (Bonnet et al., 1998), measuring the opening and closing rates of DNA stem-loops using fluorscence correlation spectroscopy. From their results, we extract 10 piconewton forces as an estimate for the closing time (at $T = 20^\circ$C) of a stem-loop structure with three basepairs and a loop of four nucleotides, which may be considered as a minimal secondary structure element. We expect that the formation of the stem-loop takes place in a single step whose reaction pathway goes through a transition state where the basepairs of the stem have not yet formed, but the corresponding bases are already closely together (see Fig. 10). In the presence of an external force, the closing time must then be multiplied with an Arrhenius factor $e^{\Delta W/k_bT}$, where $\Delta W$ is the work that has to be exerted against the force to pull in the amount of single strand needed for the formation of the stem-loop (Rief et al., 1998). With a typical force of 6 pN we obtain $\Delta W \approx 4$ kcal/mol, which results in a closing time on the order of 10 ms. This timescale has to be compared to the time it takes to stretch out the stem-loop. At a pulling speed on the order of $1 \mu$m/s, the two timescales are comparable and hence, both the formation of new secondary structure elements and the restoration of already opened ones are likely to be suppressed\(^8\). Although it is beyond the scope of this paper, we want to note that in the presence of pseudoknots and/or tertiary interactions, the formation or re-formation of structural elements is expected to be slowed down even further, due to long search times for the interaction partners.

To obtain an impression of how many features a non-equilibrium FEC might show for the group I intron, we change our equilibrium algorithm, such that the re-binding of bases is disabled once they have been unbound, and include only the contribution of the minimum free energy structures instead of all possible secondary structures. This is clearly a very crude approximation. In a proper treatment, only those kinetic processes whose energy barrier is higher than a certain threshold as determined by the pulling speed should be suppressed. Also, we did not account for the fact that the opening of basepairs occurs at higher forces in non-equilibrium as a consequence of Kramers theory (Evans and Ritchie, 1997). Nevertheless, the FEC shown in Fig. 10 gives an idea of the large number of structural transitions that take place during force-induced denaturation (for comparison, the equilibrium FEC is shown again in Fig. 9a). We therefore believe that non-equilibrium stretching experiments of RNA could lead to interesting and useful results.

We made most of the software tools developed for the present work available to the public by creating a ‘RNA pulling server’ at http://bioinfo.ucsd.edu/RNA

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\(^8\) This estimate does not apply for the rezipping of partially opened, perfectly complementary long hairpins, which is faster than closing of a stem-loop. However in real RNA structures, long stems are usually interrupted by internal or bulge loops, which we expect to reclose on similar timescales as the stem-loops.
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