

Determining the DNA stability parameters for the breathing dynamics of heterogeneous DNA by stochastic optimization

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(Dated: September 30, 2011)

We suggest that the thermodynamic stability parameters (nearest neighbor stacking and hydrogen bonding free energies) of double-stranded DNA molecules can be inferred reliably from time series of the size fluctuations (breathing) of local denaturation zones (bubbles). On the basis of the reconstructed bubble size distribution, this is achieved through stochastic optimization of the free energies in terms of Simulated Annealing. In particular, it is shown that even noisy time series allow the identification of the stability parameters at remarkable accuracy. This method will be useful to obtain the DNA stacking and hydrogen bonding free energies from single bubble breathing assays rather than equilibrium data.

Keywords: DNA breathing, heteropolymer, stochastic optimization, Simulated Annealing, robustness

I. INTRODUCTION

The Watson-Crick double-helical form of DNA¹ is not a static structure: even at standard salt conditions and room temperature the base pairs may intermittently open up and expose the otherwise protected core of the nucleotides. Such local denaturation bubbles are usually quite short-lived, however, the propensity of double-stranded DNA towards formation of longer-lived bubbles can be increased by elevating temperature or lowering the salt concentration.²⁻⁶ In naturally underwound circular DNA denaturation bubbles are stabilized by partial twist release,^{7,8} while in modern single DNA molecule setups bubble formation may be facilitated by the exertion of longitudinal stretching forces.⁹⁻¹³ The preferred location of bubbles is connected with the stability landscape of the genome, as quantified by maps of stability parameters, which are functions of the specific, underlying sequence of GC and AT base pairs.¹³⁻¹⁸ In a biological context, bubbles correspond to so-called DNA Unwinding Elements (DUE), which are central in processes such as gene regulation, DNA replication, and transcription.¹⁹ Similarly, in higher organisms the thermodynamic stability landscape of DNA is related to the coding versus non-coding properties of the genome.^{20,21} The denaturation of a long DNA chain from double-strand to two separate single-strands is a physical phase transition, whose order is determined by the magnitude of the critical exponent c for the entropy loss of a flexible polymer loop, see the discussion below.^{2-5,12,22,23} The opening-closing dynamics of denaturation bubbles can be quantified by simple

nonequilibrium models based on the gradient of the DNA stability free energy landscape.²⁴⁻²⁸

Melting profiles of DNA can be obtained from a host of experimental techniques. These include UV spectroscopic methods,⁶ circular dichroism,⁶ fluorescence resonant energy transfer measurements,²⁹ calorimetry,³⁰ or nuclear magnetic resonance,³¹ among others. Single DNA manipulation techniques such as unzipping have recently been shown to provide high accuracy results for the stability parameters and their salt dependence.¹³ From the respective melting or unzipping curves the DNA stability parameters are deduced, which in bioinformatics serve to predict the melting profiles of arbitrary, given DNA sequences.³² Up until now the different sets of stability parameters differ considerably from each other.¹³⁻¹⁸ Alternative methods to measure these may help to pin down optimized parameters. One way could be to use dynamic information from bubble breathing. Indeed, by fluorescence correlation spectroscopy the breathing dynamics of single DNA bubbles has been monitored, producing the breathing-induced fluorescence-fluorescence correlation function, that is pronouncedly non-exponential.^{33,34} Given the recent progress in experimental methods, we expect that time series of single bubble dynamics will soon become available, in which opening or closing events of individual base pairs can be monitored. A high potential for such time records lies in nano-channel approaches as the one reported in Ref. 35, after new labeling techniques will become available shortly.

In what follows we pursue the question whether the bubble size distribution obtained from single breathing time series may, in principle, be used to obtain reliable information on the DNA stability parameters. We show that indeed by stochastic analysis methods such as Simulated Annealing (SA) accurate estimates for the stability

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parameters may be obtained for known DNA sequences.

The paper is structured as follows. We first introduce the general statistical model of DNA base pairing, before proceeding to present the methodology of SA. In the subsequent section we present our results, before drawing our conclusions.

II. STATISTICAL MODEL FOR DNA DENATURATION

A. Thermodynamics

The size of denaturation bubbles typically ranges from a few broken base pairs (bps) at physiological temperature in linear, unconstrained DNA, to some 200 broken bps closer to the melting temperature of the DNA.^{2,3,5,14,34} Bubbles of some hundred broken base pairs also occur in naturally underwound DNA.^{8,19} Following the notation of Ref. 14, the stability of DNA is characterized by the free energies $\epsilon_{hb}(\text{AT})$ and $\epsilon_{hb}(\text{GC})$ for the Watson-Crick hydrogen bonds between complementary nucleotides (A and T, G and C, respectively) as well as the independent stacking free energies ϵ_{st} for disrupting the stacking interactions between nearest neighbor bps. These stacking energies depend on the nature of the two vicinal bps, as well as on their orientation along the DNA molecule (3' to 5'). The free energies are functions of temperature and salt concentration. Depending on the used set of stability parameters more or less pronounced asymmetries in the stacking free energies are observed.¹³⁻¹⁸ In addition to the hydrogen bonding and stacking free energies, there is an additional energetic cost for initiating a bubble in the first place. Roughly speaking, this term originates from the fact that two stacking contacts need to be broken, while only one single broken bp yields an entropic gain. This is either taken into consideration by the cooperativity factor σ_0 , or the so-called ring factor ξ , see below.³⁶

The L33B9 sequence³⁷ we are analyzing in the present work is given as follows,

$$5' - \text{cCGCCAGCGGCCTTTACTAAAGGCCGCTGCGC} - 3', \quad (1)$$

where the double-strand is completed by adding the complementary single strand. The sequence (1) is linear, and the high content of more stable GC bps at the two ends ensures that these ends preferentially remain closed. A denaturation bubble forms in the center of the chain that is rich in weaker AT bonds. We therefore view the two extremities denoted by the lower case symbol c as completely clamped. Labeling the sequence of bps by the coordinate x , ranging from $x = 0$ to $x = M + 1$, we thus have $M = 31$ internal bps, which are allowed to open up, while the bps at $x = 0$ and $x = M + 1$ remain closed by definition. In a mathematical sense, the bps at the two extremities represent reflecting boundary conditions. Furthermore, we call x_L and x_R the momentary positions of the two closed bps embracing the denaturation bubble

to the left and right, such that the bubble size becomes $m = x_R - x_L - 1$. In terms of the Boltzmann factors for hydrogen bonding of the bp at position x ,

$$u_{hb}(x) = \exp\left(\frac{\epsilon_{hb}(x)}{k_B T}\right), \quad (2)$$

and the stacking interactions between the bps at positions $x - 1$ and x ,

$$u_{st}(x) = \exp\left(\frac{\epsilon_{st}(x)}{k_B T}\right), \quad (3)$$

the bubble partition function becomes ($m \geq 1$):

$$\mathcal{Z}(x_L, m) = \frac{\xi'}{(1+m)^c} \prod_{x=x_L+1}^{x_L+m} u_{hb}(x) \prod_{x=x_L+1}^{x_L+m+1} u_{st}(x). \quad (4)$$

At $m = 0$, we take $\mathcal{Z}(m = 0) = 1$. In Eq. (4), the factor $(1+m)^{-c}$ takes care of the entropy loss upon formation of a closed polymer loop. For a self-avoiding chain in three dimensions, the critical exponent becomes $c = 1.76$.²² Corrections of c may occur due to interactions with the rest of the chain,²³ however, for the short DNA construct used here, such effects are not expected to be relevant. The ring factor is $\xi \approx 10^{-3}$,¹⁴ and we define $\xi' = 2^c \xi$. The ring factor may be interpreted as the cooperativity parameter, divided by the Boltzmann factor for stacking, $\xi = \sigma_0 / \exp(\epsilon_{st}/k_B T)$.¹⁴ In principle, the ring factor depends on the position. However, a bubble will statistically always form at the weakest link. Considering this we have used a constant value of ring factor, ξ in the present work. With above notation, the equilibrium distribution for finding a bubble of size m and with the leftmost broken bp located at position $x + 1$, is given by

$$P_{\text{eq}}(x_L, m) = \frac{\mathcal{Z}(x_L, m)}{\mathcal{Z}(0) + \sum_{m=1}^M \sum_{x_L=0}^{M-m} \mathcal{Z}(x_L, m)}. \quad (5)$$

B. Nonequilibrium: bubble breathing

Powered by thermal fluctuations, the bubble size becomes a random process as a function of time. Varying stepwise by further unzipping of one bp at position x_L or x_R , or by zipping at $x_L + 1$ and $x_R - 1$, the bubble size m performs a random walk along the coordinate x , the bubble breathing dynamics.^{24-28,34} This process is described by the master equation³⁴

$$\frac{\partial P(x_L, m, t)}{\partial t} = \mathbb{W}P(x_L, m, t), \quad (6)$$

where $P(x_L, m, t)$ is the probability distribution for finding a bubble of size m with the leftmost open bp at position $x_L + 1$, at time t . The matrix \mathbb{W} contains the transfer rates for all possible transitions in the (x_L, m) space, for details see Ref. 34. In the long time limit, the solution P of the master equation (6) equilibrates to

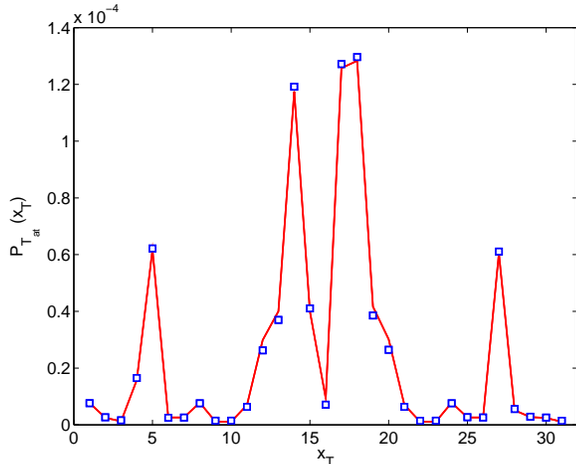


FIG. 1. (Color online) Theoretical probability distribution for finding a tagged bp at position x_T open (solid line), compared with the result from the converged SA scheme (blue open squares). The underlying DNA sequence is given in Eq. (1).

the distribution P_{eq} of Eq. (5). To generate individual bubble breathing time series for $m(t)$ and $x_L(t)$, as well as construct the distribution P_{eq} , one may employ the Gillespie algorithm.^{38,39}

Following the experimental setup in Ref. 33, one may study the dynamics of a *tagged* bp located at $x = x_T$. In the typical experimental scenario fluorescence occurs if the bps in a δ -neighborhood of the fluorophore position x_T are open. Measured fluorescence time series thus correspond to the stochastic variable $I(t)$, with the properties $I(t) = 1$ if at least all bps in $(x_T - \delta, x_T + \delta)$ are open, and $I(t) = 0$ otherwise.³⁴ In what follows we probe whether a single bp is open or closed, i.e., we choose $\delta = 0$.

III. STOCHASTIC OPTIMIZATION

Given the probability distribution $P_{\text{eq}}(m, x_L)$, constructed from an experimental or simulations time series $m(t)$, $x_L(t)$, for a bubble in the DNA construct under consideration: can we reliably extract the stability parameters? Here we show that stochastic optimization is the method of choice.

Finding system parameters in a complex landscape is a generic task across disciplines.^{40–46} Typically, a given problem is cast in such a manner that the sought-for optimum corresponds to an extremum of a functional in the complex search space. For instance, to obtain the global minimum in a rugged potential energy surface, one starts from any arbitrary point on this landscape and then moves on in the search space, following certain rules, such as accepting a move if the gradient norm for the new position decreases. This process converges to a point for which the gradient norm is zero. To ver-

ify whether this point is a minimum, one needs to check if the eigenvalues of the Hessian matrix at that point are all positive. A completely deterministic optimization procedure such as this minimization of the gradient norm, however, will generally fail to determine the global minimum if the search space features multiple minima. Once a local minimum is found, the deterministic search method will simply terminate. Such a misguidance is avoided by true global optimizers, whose search is not solely driven by a gradient. In particular, stochastic optimization techniques turn out to be very successful. Originally proposed by Kirkpatrick and coworkers to solve the traveling salesman problem,^{47,48} SA represents such a true global optimizer, and has been applied to a broad range of problems across disciplines, see, for instance, Refs. 49–57. In SA, the search space is initially sampled at a high temperature (T_{at}). The associated thermal fluctuations at a suitable value of T_{at} will lift the optimizer out of local minima such that the search may continue towards increasingly deeper minima. Once the temperature becomes sufficiently small and/or the search is carried out over a sufficient time span, the entire search space is probed. Due to this ergodic property the global minimum is indeed found unequivocally.⁴⁸

Typically, an SA analysis is started at a sufficiently high temperature. This makes nearly all moves acceptable, as the criterion for accepting or rejecting a move is determined by the Metropolis criterion. In our case, the associated cost function, which is being minimized, is the sum of the squares of the difference of the occupation probabilities at the various positions,

$$\text{cost}_i = \sum_{i=1}^M (P_{\text{eq}}(x_i) - P_{T_{\text{at}}}(x_i))^2, \quad (7)$$

where $P_{T_{\text{at}}}(x)$ denotes the distribution at position x found in the current SA step, when the simulation temperature is T_{at} . If, on going from one SA step (i) to the next ($i+1$) the magnitude of the cost function decreases, we at once accept that move. If it increases, we do not discard the move outright. Instead, we subject it to the Metropolis test⁵⁸: if the quantity $\Delta = \text{cost}_i - \text{cost}_{i-1}$ has a positive value, the probability for accepting the move is determined by the function

$$F = \exp\left(-\frac{\Delta}{T_{\text{at}}}\right). \quad (8)$$

For positive Δ , F is always between 0 and 1. For each evaluation of F , we invoke a random number rand between 0 and 1. If $F > \text{rand}$, we accept the move. If not, the move is rejected. Thus, at very high T_{at} , F will be close to 1 and most moves will be accepted, such that a greater region of the search space will be sampled. As the simulation proceeds, T_{at} is decreased by the *annealing schedule*. Once the correct path towards the global minimum is followed, we need not search the entire space and concentrate on a small region, which will guide us specifically to the global minimum. That is, as

T_{at} is lowered, a decreasing number of moves pass the Metropolis test. Ultimately, in our problem we recover the stability parameters from the SA analysis.

In SA, the crucial factor which determines the success of optimization is the annealing schedule, which is basically the rate at which the simulation temperature is decreased in successive annealing steps. In the present study we have kept the initial temperature at 1000. The rate of cooling was kept at 10% of the value of the present step. We have also ensured that after every 30 SA steps, the system is re-heated to the initial starting value, i.e., the simulation temperature is forcibly increased to a higher value. This is done to remove any possibility of being trapped in a local minimum (coming out of which will be difficult if the simulation temperature is low). In successive SA steps, along with the temperature, the individual stability parameters are changed by the following strategy. If u is a parameter chosen for change in SA, it is updated by the rule: $u' = u + u \times (-1)^n \times \delta \times r_n$, where n is a random integer, δ is the amplitude of allowed change (kept at 0.01), and r_n is a random number between 0 and 1. The new u' (changed stability parameter) is used to generate the updated distribution profile. The magnitudes of the different optimization parameters are collected in Table I.

TABLE I. Magnitude of optimization parameters used in SA.

Parameter	Magnitude
Annealing Schedule	10%
Initial Simulation Temperature	1000
Magnitude of Change δ	0.01

IV. RESULTS AND DISCUSSION

In a first step, the equilibrium distribution for a tagged bp at location x_T in the DNA sequence (1) was determined from the theoretical stability parameters from Ref. 14. SA was then employed for successive convergence of $P_{T_{\text{at}}}$ to this theoretical distribution through variation of the 12 independent free energy parameters (compare Table II), by minimizing the cost function. The SA analysis was terminated once the value of the cost function becomes smaller than 10^{-4} . Fig. 1 shows the quite accurate convergence of the SA scheme in terms of the equilibrium distribution.

To visualize the progress of the SA procedure, we display in Fig. 2 the progress of the approximation of the twelve DNA stability parameters of hydrogen bonding and base stacking (compare also Table II) for 8000 SA steps, for three separate SA runs starting with different initial simulation temperatures. For each simulation the initial free energy values are chosen via random perturbation of the experimental u values,¹⁴ following our SA strategy. In all cases the convergence is quite accurate. Two parameters do not change during the SA scheme, these correspond to the two pairs of bps, that do not occur in the employed sequence (1). To be sure that the search proceeds without being held up in local basins, the annealing temperature was raised after every 30 SA steps and then allowed to follow the usual annealing schedule. The sudden jumps in the profile are a result of this effort. At an abruptly elevated temperature, newer moves start to get accepted and hence the zigzag pattern.

In terms of the free energy values for hydrogen bonding and base stacking, the average results from 1000 SA runs are shown in Table II. We also indicate which combinations of nearest neighbor pairs actually occur in the underlying sequence (1). The convergence of the SA algorithm in all cases is quite remarkable. In addition to the free energy parameter we also optimized the loop exponent c and the ring factor ξ . The resultant simulation profiles (Fig. 3) show a good convergence towards theoretical values.

In typical experimental data the distribution of the bubble opening probability will be noisy, due to finite sampling and measurement errors. To check if our SA algorithm is robust against such noise we randomly perturbed the theoretically expected equilibrium distribution by a gaussian random processes with amplitude and width being the P_{eq} and 10% of P_{eq} , respectively. Fig. 4 shows how this noisy data was quickly smoothed out to

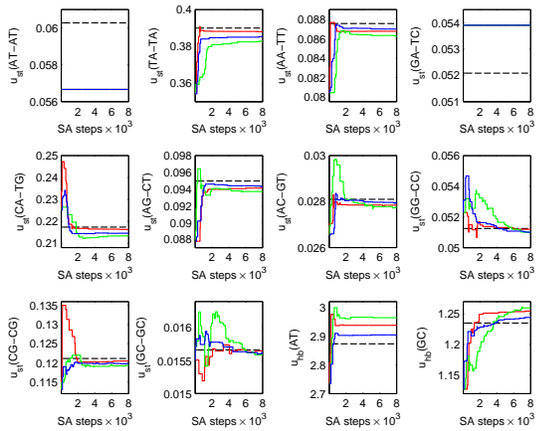


FIG. 2. (Color online) Evolution of the free energy parameters of hydrogen bonding and base stacking as function of SA steps (full lines) from three separate SA runs. The black dashed horizontal lines represent the expected experimental values taken from Ref. 14, towards which convergence is expected to occur. Note the different scales on the vertical axes. The values for two pairs of bps, AT-AT and GA-TC, do not change in the SA procedure; these two pairs do not occur in the underlying sequence (1) and are thus not subject to the SA optimization criteria, i.e., they do not converge.

TABLE II. Comparison of experimental¹⁴ and simulated free energy data. Each simulation data is a mean of 1000 different SA outputs. The rightmost column shows the presence (\checkmark) or absence (\times) of particular free energies in sequence (1). Units of free energies (ϵ_{st} and ϵ_{hb}) reported here are kcal/mol. The last two rows of the table gives a comparison of the ring factor ξ and critical exponent c .

	Experimental	SA results	
ϵ_{st} (AT-AT)	-1.729409	-1.767474	\times
ϵ_{st} (TA-TA)	-0.579800	-0.588968	\checkmark
ϵ_{st} (AA-TT)	-1.499484	-1.510239	\checkmark
ϵ_{st} (GA-TC)	-1.819371	-1.798201	\times
ϵ_{st} (CA-TG)	-0.939677	-0.922743	\checkmark
ϵ_{st} (AG-CT)	-1.455363	-1.462615	\checkmark
ϵ_{st} (AC-GT)	-2.199241	-2.175124	\checkmark
ϵ_{st} (GG-CC)	-1.829370	-1.801741	\checkmark
ϵ_{st} (CG-CG)	-1.299554	-1.318516	\checkmark
ϵ_{st} (GC-GC)	-2.559130	-2.549840	\checkmark
ϵ_{hb} (AT)	0.649775	0.651781	\checkmark
ϵ_{hb} (GC)	0.129955	0.113848	\checkmark
ξ	0.001	0.001034062	
c	1.76	1.758298	

reach the theoretical distribution profile. We show snapshots of the process for different SA steps. In each figure, the original noisy data, the equilibrium distribution profile and the evolving profile at the particular SA step are shown. At 1500 SA steps, the noisy data completely matches with the equilibrium distribution.

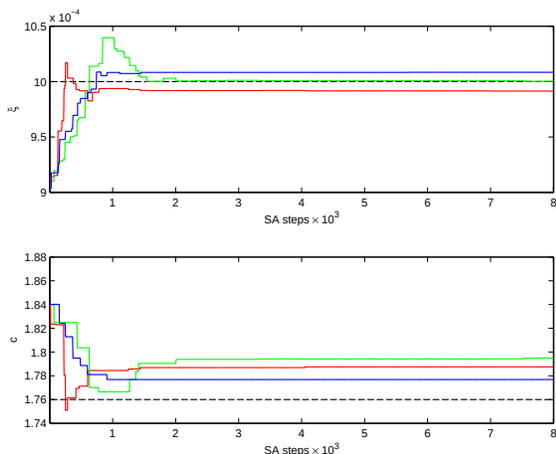


FIG. 3. (Color online) Evolution of ring factor ξ and critical exponent c from three different SA runs. The black dashed horizontal lines represent the expected literature value towards which convergence is expected to occur.

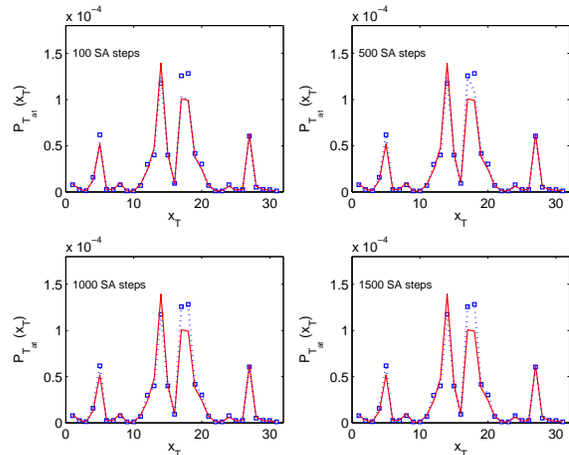


FIG. 4. (color online) Plot of $P_{T_{at}}(x_T)$ against x_T at various SA steps. In each panel the red solid lines represents the original noisy data, and the blue dashed line is the output of SA runs. The blue open squares stand for the theoretical distribution $P_{eq}(x_T)$. The plot for 1,500 SA steps already matches quite well the expected distribution $P_{eq}(x_T)$.

V. CONCLUSION

Generalising our previous approach,⁵⁹ we here demonstrate the outstanding ability of stochastic optimization to determine the stability parameters of double-stranded DNA from time series of the breathing dynamics of individual bps. Even for a short DNA sequence such as L33B9 [Eq. (1)] with only 31 internal bps, the convergence of the chosen SA scheme to all present base stacking and hydrogen bonding free energies is recovered with appreciable accuracy. Even when the input data are perturbed randomly, mimicking noisy experimental or simulations data, the stochastic optimization technique works successfully.

Optimization based on the bubble distribution $P_{eq}(x)$ is not the only way to extract the DNA stability parameters. For instance, one might use average values for the zipping and unzipping rates of individual bps and relate their ratio to the underlying free energy difference. Alternatively, once from high throughput fluorescence correlation experiments an accurate result for the fluorescence autocorrelation function becomes available, one might use this function as basis for the optimization. In principle, one might also modify our approach to analyse data from DNA unzipping. This, however, requires detailed knowledge on the change of the stacking and hydrogen free energies upon stretching of the DNA strands.

In general, it may be worthwhile to also explore the possibility to apply other techniques such as the genetic algorithm,⁶⁰ parallel tempering,⁶¹ or ant colony optimization,^{62,63} and to compare these methods.

ACKNOWLEDGMENTS

ST acknowledges the financial support from UGC, New Delhi, for granting a Junior Research Fellowship [UGC/800/Jr. Fellow (SC)]. PC wishes to thank The Centre for Research on Nano Science and Nano Technology, University of Calcutta for a research grant [Conv/002/Nano RAC (2008)]. RM acknowledges funding through the Academy of Finland's FiDiPro scheme. SKB acknowledges support from Bose Institute through an initial start up fund.

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