Diffusion Dynamics, Moments, and Distribution of First Passage Time on the Protein-Folding Energy Landscape, with Applications to Single Molecules

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We study the dynamics of protein folding via statistical energy-landscape theory. In particular, we concentrate on the local-connectivity case with the folding progress described by the fraction of native conformations. We obtain information for the first passage-time (FPT) distribution and its moments. The results show a dynamic transition temperature below which the FPT distribution develops a power-law tail, a signature of the intermittency phenomena of the folding dynamics. We also discuss the possible application of the results to single-molecule dynamics experiments.

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The study of diffusion along a statistical energy landscape is a very important issue for many fields. In the field of protein folding, the crucial question is how the many possible configurations of a polypeptide chain dynamically converge to a particular folded state. Clearly, a statistical description is needed for a large number of configurational states. According to the energy-landscape theory of protein folding, there exists a global bias of the energy landscape towards the folded state due to natural evolution selection. Superimposed on this is the fluctuation or roughness of the energy landscape coming from competing interactions of the amino acid residues. The folding energy landscape is like a funnel, and there are in general multiple routes towards the folded state. Discrete paths emerge only when the underline energy landscape becomes rough, and local traps (minima) start to appear. In addition to the thermodynamics of the folding-energy landscape, the kinetics of folding along the order parameter that represents the progress of folding towards the native state can be discussed.

When the energy landscape is smooth, the average diffusion time is a good parameter for the characterization of the dynamical process. On the other hand, when the energy landscape is rough, there exist large fluctuations of the energies, and the diffusion time is expected to fluctuate very much around its mean. In this case we need to know the full distribution of the diffusion time in characterizing the folding process.

It is now possible to measure the reaction and folding dynamics of individual molecules in the laboratory. On complex energy landscapes such as those of biomolecules, reactions in general do not obey exponential laws, and activation processes often do not follow the simple Arrhenius relation. However, measurements on large population of molecules usually cannot distinguish whether such complex rate dynamics is from the inhomogeneous distribution among molecules or from intrinsic features of individual molecules. The study of the statistics of individual molecular reaction events can greatly clarify these more subtle reaction processes. The information on the diffusion-time distribution provides a way to help unravel the fundamental mechanism of single molecule reactions. Previously many works have focused on the average rate behavior, whereas very few physical studies and discussions addressed on the whole distribution of folding rates. In this paper, we concentrate on the statistics and distributions of the first passage time (FPT) to probe the folding energy landscape. A dynamic transition temperature is found above which the FPT distribution is Poisson-like, and below which the distribution develops a power-law tail, where non-self-averaging behavior in kinetics emerges. Moreover, we find that this dynamic transition temperature is close to the thermodynamic folding transition temperature.

The framework we adopt here was first introduced by Bryngelson and Wolynes. The problem of folding dynamics is modelled as random walks on a rough energy landscape. In this model, the energy landscape is generated by the random-energy model, which assumes that interactions among non-native states are random variables with given probability distributions. For this model there are $N$ residues in a polypeptide chain. For each residue there are $\nu + 1$ available conformational states, one being the native state. A simplified version of the polypeptide chain energy is expressed as

$$E = - \sum \epsilon_i(\alpha_i) - \sum J_{i,i+1}(\alpha_i, \alpha_{i+1}) - \sum K_{i,j}(\alpha_i, \alpha_j)$$

where the summation indices $i$ and $j$ are labels of amino acid residues, and $\alpha_i$ is the state of $i$th residue. The three terms represent the one-body potential, the two-body interactions for nearest-neighbor residues in sequence, and the interactions for residues close in space but not in sequence, respectively. In this random energy model the
energies for the non-native states and interactions are replaced by random variables with Gaussian distributions. By this random energy construction one can easily generate energy surfaces with roughness controlled by adjusting the widths of these probability distributions. Another feature of the model is the random-energy approximation, which assumes that energies for different conformational arrangements are uncorrelated. Using a microcanonical ensemble analysis, the average free energy and thermodynamic properties of the polypeptide chain can be obtained \[1\].

In this study, we use the fraction of native conformations \(\rho\) as an order parameter that represents the progress of the folding. When the kinetics is dominated by the activation folding process, the states are in general locally connected in \(\rho\). Therefore we assume the local connectivity condition here, assuring that the dynamics happens continuously with \(\rho\). The kinetic process is carried out with the use of Metropolis dynamics. Therefore the transition rate from one conformation state to a neighboring state is determined by the energy difference of these two states, and an overall constant \(\tau\) characterizes the time scale of residue interactions. The readers are referred to Ref. \[2,11\] for detailed derivations of the dynamic equations.

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The proteins with larger size tend to form domains which is beyond the scope of the current mean field approximation. For simplicity we assume \(\delta \epsilon = \delta L\) and \(\Delta L\). Therefore the ratio of the energy gap between the native state and the average of non-native states over the spread of non-native states, \(\delta \epsilon / \Delta \epsilon\), becomes an appropriate parameter, representing interactions. Note that the two-body energies \(\delta L\) and \(\Delta L\) already include the type of interactions due to the second and third term in Eq. (1): \(D(\rho, s)\) is the frequency-dependent diffusion coefficient \[3\]:

\[
D(\rho, s) = \left( \frac{\lambda(\rho)}{2N^2} \right) \left( \frac{R}{R + s} \right) (\rho) \left( \frac{1}{R + s} \right) (\rho),
\]

where \(\lambda(\rho) = 1/\nu + (1 - 1/\nu)\rho\). The average \(\langle \cdot \rangle_R\) is taken over \(P(R, \rho)\), the probability distribution function of transition rate \(R\) from one state with order parameter \(\rho\) to its neighboring states, which may have order parameters equal to \(\rho - \frac{1}{N}\), \(\rho\), or \(\rho + \frac{1}{N}\). The explicit expression of \(P(R, \rho)\) can be found in Ref. [2]. The boundary conditions for Eq. (2) are set as a reflecting one at \(\rho = 0\) and an absorbing one at \(\rho = \rho_f\). The choice of an absorbing boundary condition at \(\rho = \rho_f\) facilitates our calculation for the first passage-time distribution. One can also rewrite Eq. (2) in its integral-equation representation by integrating it twice over \(\rho\):

\[
G(\rho, s) = -\int_0^\rho dp' \int_0^{\rho''} dp'' [sG(\rho'', s) - n_i(\rho'')]
\]

\[
\times \exp \left( U(p', s) - U(p, s) \right) \frac{D(\rho', s)}{D(\rho, s)}
\]

The folding time is approximated by the first passage time (FPT) for the order parameter to reach \(\rho_f\). Thus one can easily write down the following relation for the FPT distribution function \(P_{FPT}(\tau)\):

\[
P_{FPT}(\tau) = \frac{d}{d\tau} (1 - \Sigma) = -\frac{d\Sigma}{d\tau}
\]

where \(\Sigma(\tau) = \int_0^\tau dp G(\rho, \tau)\). The \(n\)th moment of the FPT distribution function can be calculated via \(\langle \tau^n \rangle = n!(-1)^{n-1} \int_0^\tau dp G_{n-1}(p)\), where \(G(\rho, s) = G_0(\rho) + sG_1(\rho) + s^2G_2(\rho) + \cdots\). By Taylor-expanding Eq. (5) with respect to \(s\), we can solve for \(G_n(\rho)\) and therefore \(\langle \tau^n \rangle\) iteratively by matching the coefficients of \(s^n\). In the mean time, one can also solve for \(G(\rho, s)\) directly from Eq. (5) using the linear inversion technique. Observing that \(P_{FPT}(s) = 1 - s\Sigma(s)\), where \(P_{FPT}(s)\) and \(\Sigma(s)\) are Laplace transforms of \(P_{FPT}(\tau)\) and \(\Sigma(\tau)\), respectively, one can calculate \(P_{FPT}(s)\) and investigate \(P_{FPT}(\tau)\) via an inverse Laplace-transform technique.

We start the numerical calculations by setting \(R_0 = 10^8 s^{-1}\), \(N = 100\) and \(\nu = 10\), which are approximations to the physical values. We confine ourselves to the single domain proteins with size less than 100. The proteins with larger size tend to form domains which is beyond the scope of the current mean field approximation.
the importance of gap bias towards the folded state relative to the roughness of the landscape. One can show that only the relative ratios among $\delta \epsilon$, $\Delta \epsilon$ and $T$ are the controlling parameters in this problem. We set the initial distribution of the polypeptide chain molecules to be $n_i(\rho) = \delta(\rho - \rho_i)$, where $\rho_i$ is set to be 0.05. In our calculations we set $\rho_f = 0.9$. This means that 90 percent of the amino acid residues are in their native states.

The mean first passage time (MFPT) $\langle \tau \rangle$ for the folding process versus a scaled inverse temperature, $T_0/T$, is plotted in Fig. 1 for various settings of the parameter $\delta \epsilon/\Delta \epsilon$. We have an inverted bell-like curve for each fixed $\delta \epsilon/\Delta \epsilon$, and the MFPT reaches its minimum at a temperature $T_0$. At high temperatures, the MFPT is large although the diffusion process itself is fast (i.e., $D(\rho, s)$ is large). This long-time folding behavior is due to the instability of the folded state. The MFPT is also large at low temperature, which indicates that the polypeptide chain is trapped in low-energy non-native states. This is in agreement with simulation studies [8].

By comparing the MFPT minimum for various $\delta \epsilon/\Delta \epsilon$, we conclude that this minimum becomes smaller when the ratio of the energy gap versus roughness increases. This suggests that a possible criterion for selecting the subset of the whole sequence space leading to well-designed fast folding protein is to maximize $\delta \epsilon/\Delta \epsilon$. In other words, one has to choose the sequence subspace such that the global bias overwhelms the roughness of the energy landscape [14,3,9].

We also calculate the higher-order moments of the FPT distribution. In Fig. 2 we show the behavior of the reduced second moment, $\langle \tau^2 \rangle/\langle \tau \rangle^2$. We find that the reduced second moment starts diverging at a temperature around and below $T_0$, where the MFPT is at its minimum. This is an indication of a long tail in the FPT distribution. The divergence of the second moment also shows that the dynamics exhibits non-self-averaging behavior.

From the study of higher moments, we find the relationship $\langle \tau^n \rangle \approx n! \langle \tau \rangle^n$ when $T > T_0$. Therefore in the high-temperature regime the FPT distribution function is Poissonian and decays exponentially at large time. When $T < T_0$, it is hard to obtain more information from the moments because of their diverging behavior. However, we can study the problem by solving Eq. (5) directly. By investigating the behavior of the FPT distribution function in the Laplace-transformed space, we see that for $T < T_0$ the FPT distribution is very similar to a Lévy distribution in time space, which develops a power-law tail at large time: $P_{FPT}(\tau) \sim \tau^{-(1+\alpha)}$ for large $\tau$. In Fig. 3 we make a plot of the exponent $\alpha$ versus $T_0/T$ for the case $\delta \epsilon/\Delta \epsilon = 4.0$. We find that $\alpha$ is decreasing when the temperature is lowered, and $\alpha$ approaches 1 when $T$ goes to $T_0$, where the exponential kinetics is resumed.

![Fig. 1. MFPT versus reduced inverse temperature $T_0/T$ for various $\delta \epsilon/\Delta \epsilon$. As $\delta \epsilon/\Delta \epsilon$ increases, the minimum of MFPT decreases.](image1.png)

![Fig. 2. $\langle \tau^2 \rangle/\langle \tau \rangle^2$ versus reduced inverse temperature $T_0/T$ for various $\delta \epsilon/\Delta \epsilon$. At high temperature this value keeps finite and the folding process is self-averaging. As the temperature drops, the value starts to diverge and non-self-averaging behavior emerges.](image2.png)
FIG. 3. The exponent $\alpha$ versus $T_0/T$ for the case $\delta \epsilon/\Delta \epsilon = 4.0$ when $T_0/T > 1$. Below the transition temperature $T_0$, the FPT distribution is close to a Lévy distribution, which has a power-law tail $P_{FPT}(\tau) \sim \tau^{-(1+\alpha)}$ at large $\tau$.

From the results above, we find that for a fixed-energy landscape, there exists a dynamic transition temperature $T_0$. When the temperature is above $T_0$, the FPT distribution is Poissonian, indicating exponential kinetics, and in random-walk language we have normal diffusion on the energy landscape. Below $T_0$, the variance and higher moments diverge, and the FPT distribution shows a power-law decay behavior, exhibiting signs of anomalous diffusion. This indicates the process is non-self-averaging and distinct folding pathways emerge at various time scales. As a comparison, we have calculated the thermodynamic folding-transition temperature $T_f$ by identifying the maximum heat capacity. We find that $T_f$ is less than but close to $T_0$ for various settings of $\delta \epsilon/\Delta \epsilon$. This indicates that the thermodynamic and dynamic behavior in proteins are strongly correlated. However, recent simulation and experimental results [11,12] show that $T_f > T_0$ and folding is faster below $T_f$ but above $T_0$. This indicates a possible limitation of the present analytical study. Furthermore, the crossover behavior near $T_0$ has been shown to be smoother than what we observe here. This is probably due in part to the insufficient cooperative interactions in our study [11]. We will address these issues in detail in a future publication [13].

In single-molecule folding experiments, it is now possible to measure not only the mean but also the fluctuation and moments as well as the distribution of folding time [15]. Under different experimental and sequence conditions, one can see different behavior of the folding time and its distributions. A well-designed fast folding sequence with suitable experimental condition exhibits self-averaging and simple rate behavior. Multiple routes are parallel and lead to folding. A less well-designed sequence (with larger $\Delta \epsilon$) folds slowly and often exhibit non-self-averaging non-exponential rate behavior, indicating the existence of intermediate states or local traps. In this case, the folding process is sensitive to which kinetic path it takes, since a slight change in a folding pathway may cause large fluctuation in the folding time, which indicates intermittency. One can use single-molecule experiments to unravel the fundamental mechanisms and intrinsic features of the folding process. In typical experiments of bulk molecules, it is very hard to observe and analyze the intermittency, because the dynamics is averaged over an ensemble of molecules and furthermore, we cannot say if the bulk phenomena results are either from the intrinsic features of individual molecules or the inhomogeneous averages over the molecules.

It is worth mentioning that although we focus on the study of the protein-folding problem in this paper, the approach we use here is very general for treating problems with barrier crossings on a multi-dimensional complex energy landscape. The main ingredient for this model is Brownian motion on a rough multi-dimensional landscape, or equivalently, a random walk on a complex network with a frustration-inducing environment. Since this is quite general and universal, we expect our results may also be able to account for a large class of phenomena. In fact the experiments on glasses, spin glasses, viscous liquids and conformational dynamics already show the existence of non-exponential distributions at low temperature. In particular, a recent experiment on single-molecule enzymatic dynamics [10] shows explicitly the Lévy-like distribution of the relaxation time for the underlying complex protein energy landscape. An interesting study of anomalous diffusion and non-exponential dynamics has been made recently using a Fractional Fokker-Planck Equation (FFPE) [7] to describe dynamic processes characterized by Lévy distributions. Our results show that the approach we use here can serve as a microscopic basis for the use of such a FFPE.

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