# Folding with Downhill Behavior and Low Cooperativity of Proteins

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ABSTRACT The downhill folding observed experimentally for a small protein BBL is studied using off-lattice Gō-like model. Our simulations show that the downhill folding has low cooperativity and is barrierless, which is consistent with the experimental findings. As an example of comparison in detail, the two-state folding behavior of proteins, for example, protein CI2, is also simulated. By observing the formation of contacts between the residues for these two proteins, it is found that the physical origin of the downhill folding is due to the deficiency of nonlocal contacts which determine the folding cooperatively. From a statistics on contacts of the native structures of 17 well-studied proteins and the calculation of their cooperativity factors κ<sub>2</sub> based on folding simulations, a strong correlation between the number of nonlocal contacts per residue  $N_N$  and the factors  $\kappa_2$  is obtained. Protein BBL with a value of  $N_N = 0.73$  has the lowest cooperativity factor  $\kappa_2 = 0.34$  among all 17 proteins. A crossover around  $N_N^c \sim 0.9$  could be defined to separate the two-state folders and the downhill folder roughly. A protein would behave downhill folding when its  $N_N = N_N^c$ . For proteins with their  $N_N$  values are about (or slightly larger than)  $N_N^c$ , the folding behaves with low cooperativity and the barriers are small, showing a weak two-state behavior or a downhill-like behavior. Furthermore, simulations on mutants of a two-state folder show that a mutant becomes a downhill folder when its  $N_N$  is reduced to a value smaller than  $N_N^c$ . These could enable us to identify the downhill folding or the cooperative two-state folding behavior solely from the native structures of proteins. Proteins 2006;63:165-173. © 2006 Wiley-Liss, Inc.

Key words: protein BBL; downhill folding; Gō-like model; cooperativity; the number of non-local contacts per residue

## INTRODUCTION

Natural proteins fold to their unique folded structures or native states relying on the interactions encoded in their amino acid sequence. <sup>1–5</sup> A widely accepted folding mechanics is based on the funnel-like landscape theory. <sup>5</sup> The folding of protein molecules resembles a diffusive process on rugged landscape of free energy, and there is an ensemble of pathways to the native state. <sup>6,7</sup> Different

proteins show different folding kinetics under different conditions. Many small or single-domain proteins behave with so-called two-state folding behavior, that is, hopping between the denatured states and the native state without any accumulation of intermediate states due to the existence of a main barrier between these two kinds of states.8 The corresponding folding transition resembles to a firstorder-like phase transition with high cooperativity.9-11 For large or multidomain proteins, the folding shows a behavior with multiple states or intermediate states. However, the funnel theory also suggested another kind of folding, which proceeds without crossing free energy barrier along the reaction coordinate. 5,7,12 Recently, Garcia-Mira and colleagues observed experimentally such a folding behavior, namely the downhill folding, for a small protein BBL (PDB code: 1BBL). 13 They showed that the cooperativity of folding is low and there is no free energy barrier from their thermodynamic measurements. In contrast to the binary character of two-state folder, the order parameter Q of equilibrium states for this protein decreases continuously as the temperature increases, implying a conformational rheostat which is a new concept of functional states of proteins. 13-15 Such a protein is called a downhill folder. Very recently, Muñoz and coworkers further checked the robustness of the downhill folding and showed experimentally that singly and doubly labeled BBL do not aggregate, and unfold reversibly. 16

Thus, some interesting issues have been raised. What features does the downhill folding show, and can we simulate the downhill folding using simplified or all-atomic models and characterize the folding mechanism? How do we interpret the difference between the experimental finding for protein BBL and other two-state proteins? More importantly, can we find out what are the physical reasons that make the downhill folder, such as protein BBL, show the downhill folding behavior? Can we identify a protein being a two-state folder, a downhill or downhill-like folder solely from the nature of its native structures? The answer to these questions will be helpful for under-

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standing the folding behaviors, and also the protein functions and protein design as well.

In this work, a study on the downhill folding from aspects of both molecular dynamics simulations and statistics on the contacts between residues for a number of proteins is reported. The Gō-like model 17-25 is used for our simulations. For protein BBL, a wide peak of specific heat and the absence of folding transition imply that the cooperativity is low. The free energy as a function of the similarity Q values to the native state clearly shows the absence of energetic barrier. As a comparison, the folding of two-state folders, for example, the chymotrypsin inhibitor II (CI2) and some other proteins, is also worked out. Our results show that the downhill folding is rather different from the two-state folding behavior, and the nonlocal contacts dominate essentially the cooperativity of folding. A statistics on two kinds of interactions, namely the local and nonlocal contact interactions, between the residues for the native structures of 17 proteins is made. In addition, the cooperativity factors  $\kappa_2^{26-28}$  of these 17 proteins are also calculated based on the folding simulations. A strong correlation between the numbers of nonlocal contacts per residue  $N_N$  and the cooperativity factor  $\kappa_2$ is found, and a crossover around  $N_N^c \sim 0.9$  could be defined to separate the two-state folder and the downhill folder roughly. This is further validated by folding simulations on mutants of a two-state folder which become a downhill folder when their  $N_N < N_N^c$ . Thus, a protein could be roughly classified into downhill folding or two-state folding only by checking its number of nonlocal contacts per residue, that is, the value of  $N_N$ .

# MODEL AND METHODS

#### Gō-Like Model

The Gō-like energy  $^{17}$  has been widely used to describe the interactions of residues both in lattice  $^{29,30}$  and off-lattice model proteins.  $^{18-25}$  Residues are represented by single beads centered in their  $C\alpha$  atoms, and all beads are connected into a polymer chain by virtual bonds. The interactions include terms related to the virtual bonds, angles, dihedral angles, and nonbonded pairs of the beads, that is, the residues. Such kind of Gō-like interactions has been successfully applied in the modeling of folding processes of small and single domain proteins.  $^{18-20,31-34}$  The terms and parameters of the Gō-like interactions are similar to those used by Clementi and colleagues.  $^{18}$  For a certain conformation of the protein, the total Gō-like potential energy is thus given by the expression:

$$\begin{split} V_{\rm total} &= V_{\rm bonded} + V_{\rm bond-angle} + V_{\rm dihedral} + V_{\rm nonbonded} \\ &= \sum_{\rm bonds}^{N-1} K_r (r-r_0)^2 + \sum_{\rm bond-angles}^{N-2} K_{\rm \theta} (\theta-\theta_0)^2 \\ &+ \sum_{\rm dihedrals}^{N-3} \{ K_{\rm \phi}^{(1)} [1-\cos(\varphi-\varphi_0)] + K_{\rm \phi}^{(3)} [1-\cos3(\varphi-\varphi_0)] \} \\ &+ \sum_{i < j-3}^{\rm native} \epsilon \left[ 5 \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{12} - 6 \left( \frac{\sigma_{ij}}{r_{ij}} \right)^{10} \right) \right] + \sum_{i < j-3}^{\rm non-native} \epsilon \left( \frac{\sigma_0}{r_{ij}} \right)^{12}. \end{split} \tag{1}$$

Here, N is the total number of residues; r,  $\theta$ , and  $\phi$  are the bond length between two subsequential residues; bondangle is formed by three values in the native structure of the protein; and  $r_{ij}$  is the spatial distance between two  $C\alpha$ atoms that have at least four residues between them along the chain. In the summation for all native contacts, a 10 to 12 Lennard–Jones (LJ) potential is used, and  $\sigma_{ij}$  is the  $C\alpha - C\alpha$  distance between the contacting residues *i* and *j* in the native structure. Here, the native contact is defined as the contact between two residues i and j in the native structure if any a heavy atom pair from each residue are less than 5 Å apart. In the summation for all non-native contacts, the parameter  $\sigma_0$  introduces the excluded volume repulsion between residue pairs that do not belong to the given native contact set. In our simulations,  $\sigma_0=4~{\rm \AA}$  is used. The interaction parameters are taken as  $K_\gamma=100\epsilon$ ,  $K_{\theta} = 20\epsilon$ ,  $K_{\phi}^{(1)} = \epsilon$ ,  $K_{\phi}^{(3)} = 0.5\epsilon$ , respectively. During the folding, a native contact between two residues i and j is formed if the distance between their  $C\alpha$  atoms is shorter than  $\kappa$  times their native distance  $\sigma_{ij}.$  Here  $\kappa=1.2$  is used. Based on such a simplified model, the cooperative folding behavior of proteins can be reproduced in a semiquantitative manner, which suggests that the Gō-like potential is workable.24,28

#### **Langevin Dynamics**

We use Langevin dynamics<sup>22,24,25,35</sup> to simulate the folding process starting from a linear coil. The Langevin equation of motion is:

$$m\dot{\mathbf{v}}(t) = \mathbf{F}_{conf}(t) - \gamma \mathbf{v}(t) + \Gamma(t)$$
 (2)

where  $\dot{\mathbf{v}}$ ,  $\mathbf{v}$ , and m are velocity, acceleration, and mass of a bead, respectively;  $\mathbf{F}_{conf} = \nabla E_p$  is the conformational force;  $\gamma$  is a friction (viscosity) constant. Here, we use  $\gamma = 0.5\tau^{-1}$  with  $\tau$  being the time scale.  $\Gamma$  is the random force, which is produced from the Gaussian distribution with a standard variance related to temperature by:

$$<\Gamma(t)\Gamma(t')> = 6\gamma k_B T\delta(t-t').$$
 (3)

Here,  $k_B$  is the Boltzmann constant, T is the temperature, t is the time, and  $\delta(t-t')$  is the Dirac delta function. Every bead is subject to a random force at each integration time step. The components of random force are independently generated by setting  $\Gamma_{\alpha} = \Phi_{\alpha} \sqrt{2m\gamma k_BT/\delta t}$  where  $\Phi_{\alpha}$  is a random value taken from a standard Gaussian distribution (zero mean and unit variance);  $\alpha$  denotes the uncorrelated components of random force in the x,y, or z directions; and  $\delta t$  is the integration time step. At the beginning of every simulation, the initial velocities are assigned as zero.

We use the leap-frog algorithm to integrate Eq. (2). The time scale of the model protein here is always controlled by a quantity  $\tau = \sqrt{m\lambda^2/\epsilon_0}$  with the length scale  $\lambda = 3.8$  Å, the energy scale  $\epsilon_0 = 1$ . The time step of simulations is set as  $\delta t = 0.005\tau$ . Simulation times in this study are presented in units of  $\delta t$ . The energy parameter  $\epsilon$  and temperature T are given, in units  $\epsilon_0$  and  $\epsilon_0/k_B$ , respectively. The length is measured in units of  $\lambda$ . To simplify the notation, other units are chosen such as m=1 and  $k_B=1$  in the present

simulation, as used by Thirumalai and coworkers.<sup>36</sup> An approximate correspondence between the model time and real protein kinetic time scales has been discussed by Thirumalai and coworkers.<sup>36</sup>

## Weighted Histogram Analysis Method

The weighted histogram analysis method (WHAM) is used to calculate relevant thermodynamic quantities based on the statistical physics.  $^{18,37,38}$  WHAM yields an optimal estimate of the density of state of the system. In the canonical ensemble at temperature T, the probability distribution,  $P_T$ , of potential energy E is given by

$$P_T(E) = (1/Z_c)w(E)e^{-E/k_BT}$$
 (4)

and the probability distribution of other thermodynamic quantities, such as Q, follows

$$P_T(Q) = (1/Z_c) \sum_E w(Q, E) e^{-E/k_B T}$$
 (5)

where w is the density of state of the system and is obtained by solving two self-consistent equations. <sup>38</sup>  $Z_c$  is the canonical partition function. The specific heat is defined by the energy fluctuation, <sup>25</sup> and the free energy at fix temperature  $F_T(Q)$  is estimated by the logarithm of probability  $P_T(Q)$ . <sup>18</sup>

#### **Quantification of the Cooperativity**

In this work, a factor  $\kappa_2$  is used to measure the cooperativity of the folding. Such a factor has been discussed by Chan and colleagues<sup>26–28</sup> and can be defined as

$$\kappa_2 = 2T_f \sqrt{k_B C_v(T_f)} / \Delta E_{\rm cal}$$
 (6)

where  $T_f$  is the folding transition temperature,  $C_v(T_f)$  is the specific heat at  $T_f$ , and  $\Delta E_{\rm cal}$  is the calorimetric energy defined by the difference between the energy of denatured state and native state. A detailed discussion can be found in the related papers by Chan and coworkers. <sup>26–28</sup> In this article, the factor  $\kappa_2$  of 17 proteins are calculated based on folding simulations.

#### **Computational Mutations**

Mutation is a useful method for protein science. For some specific mutations of a protein, the stable structures of the related mutants may not be available from the PDB database although they can be solved experimentally using X-ray or NMR techniques. For our theoretical studies, because only some sketchy structures of mutants are needed, a computational approach to the structures of mutants would be a good alternative of experiments. In our work, the structures with global minimized energy are defined as the stable structures of the mutants using simulated annealing, which is realized by all-atomic simulation software Amber. <sup>39</sup> A similar realization using Insight-II for finding stable structures has also been reported in literature. <sup>40</sup>

In detail, the procedure of computational mutations for a certain protein is as follows: For our interested proteins, for example, protein monomeric  $\lambda$ -repressor, we regularly

prefer to mutate the residues with large side-chains into residues with small ones. Thus, the number of the nonlocal contacts for residues could be reduced. Because only a few residues are mutated, the structure of the mutant may change slightly compared with the native structure of the wild type. For every mutated sequence, the unfoldingannealing and minimization start from the native structure of the wild type. [The mutated sequence is threaded onto the native structure of the wild type.] This initial structure begins its unfolding at 500 K until its value of cRMSD = 10 Å (here, cRMSD is the root mean square deviation of the coordinates of  $C\alpha$  atoms of protein<sup>41</sup>), and then a gradually annealing to room temperature within 3000 ps is made. A 4000-cycle minimization is further carried out after the annealing to get a structure with the minimal energy, and the resultant structure is taken as a candidate of the stable/native structure of the mutant. Here, the process of unfolding-annealing and minimization is repeated several times (4 times for mutants M2 to M6 and 50 times for mutant M1) for finding a structure with energy as low as possible. Among all the candidates, the one with the lowest energy is chosen as the final stable structure of the mutant. Note that the Amber 1999 force field,<sup>39</sup> periodic boundary condition, constant pressure, and all other default parameters of Amber are used in our simulations.

It is noted that taking the native structure of the wild type as the starting structure of the mutants for our computational mutations is quite reasonable. Threading of the mutated sequences onto different unfolded conformations of the native structure of the wild type shows that the energies of these conformations after minimization are larger than the lowest energy of the mutated sequence threaded onto the native structure of the wild type after minimization. Here, over 10 unfolded conformations with their related values of 10 Å < cRMSD < 20 Å are used. Therefore, a partial unfolding of the native structure of the wild type with cRMSD = 10Å for process of unfoldingannealing and minimization to find the stable structure of the mutant is workable. These actually indicate that all the related conformations are around the basin of the native structure of the wild type on the energy landscape. Thus, although the number of nonlocal contacts is reduced, the stable structure of the mutant does not change so significantly (see Fig. 5).

# RESULTS AND DISCUSSIONS Folding Behaviors

The folding kinetics of protein BBL is characterized by the fraction of native contacts Q versus the time at various temperatures. At temperature  $T=0.8T_{\rm f},\,1.0T_{\rm f},\,1.1T_{\rm f},\,$  and  $1.2T_{\rm f},\,$  after an extremely short relaxation the system reaches different equilibrium states with  $Q=0.9,\,0.7,\,0.45,\,$  and  $0.3,\,$  respectively (see Fig. 1). It is found that the Q values fluctuate around these equilibrium states, and there is a maximal fluctuation at  $T=T_f$  [Fig.1(b)]. {Here, the value of  $T_f$  could be defined at the peak of specific heat [see Fig. 2(a)]}. There is no cooperative hopping between native and denatured states, so no obvious folding transi-

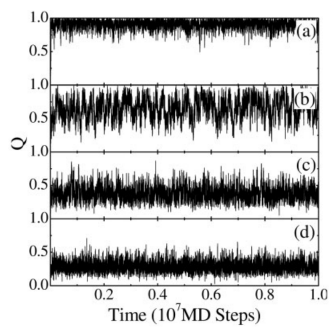


Fig. 1. Four folding trajectories for protein BBL at different temperatures: (a)  $T=0.8T_p$  (b)  $T=1.0T_p$  (c)  $T=1.1T_p$  and (d)  $T=1.2T_p$  Here Q is the fraction of native contacts.

tion. The curve of specific heat is wide and the peak is not sharp. These are consistent with the absence of cooperative transition, indicating that the cooperativity of folding for protein BBL is rather low. For each temperature, the free energy profile shows only one minimum around the equilibrium Q value which is dependent of the temperature [see Fig.2(b)]. The equilibrium Q value increases (or the minimal position of the free energy) monotonically from its value Q=0.3 at the denatured state to its value Q>0.9 at folded state, behaving as a conformational rheostat. There is no metastable state in the whole range of temperatures, and the free energy profiles show no energetic barrier clearly. These relate to the characteristics of downhill folding observed in experiment.  $^{13}$ 

As a comparison, the folding behavior of protein CI2 is simulated [see the insets in Fig. 2(a,b)]. The folding of CI2 shows a typical two-state behavior, that is, below  $T_f=1.05$  the Q value reaches to Q>0.9, and above  $T_f$  the Q value is about Q=0.2, while around  $T_f$  the Q value changes between the folded and unfolded states. The specific heat has a much sharper peak, which is about 10 times higher than that of protein BBL [see Fig.2(a)]. The free energy profile shows a large barrier around Q=0.5 [see Fig.2(b)]. These are also observed for SH3 domain and other proteins (data not shown). The folding cooperativity of the two-state folders is very high.

#### **Local and Nonlocal Contacts**

Cooperativity is a main signature of the folding mechanism. What results in the difference of the cooperativity between the downhill folders and the two-state folders? For the Gō-like model, the energetic effects are considerably optimized, while the topology of a protein is believed

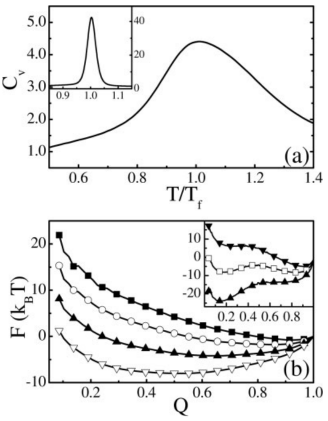


Fig. 2. The thermodynamics of folding. (a) The specific heat versus temperature T for protein BBL. The inset shows the specific heat for protein Cl2. (b) Free energy profile for protein BBL at four temperatures  $T=0.8T_r$  (solid squares),  $T=0.90T_r$  (circles),  $T=1.00T_r$  (solid triangles), and  $T=1.10T_r$  (down triangles). The inset shows the free energy profile for protein Cl2 at temperatures  $T=0.95T_r$  (solid down triangles),  $T=1.00T_r$  (squares), and  $T=1.05T_r$  (solid triangles). Here  $T_r$  for BBL and Cl2 are 0.85 and 1.05, respectively.

to be the controlling factor for the thermodynamics and kinetics of the folding. The native topology is characterized by the distribution of contacts which can be divided into local contacts and nonlocal ones. A contact is classified as local (or nonlocal) contact  $^{43,44}$  if the distance of the two contacted residues along the sequence, that is,  $L_c$ , is smaller (or larger) than a certain value, say five or six residues. Our criterion for this classification is defined as  $L_c^c=6$ . Note that the distance along the sequence is relevant to the contact order. Previously, it was argued that the nonlocal interactions dominate the cooperativity of the protein folding  $^{44,45}$  because the nonlocal interactions introduce some global ordering into the system.

For proteins BBL and CI2, it is found that the nonlocal and local contacts have different distributions during the folding, and the evolutions of the degrees of these contacts formed at the transition temperature  $T_f$  are also different (see Fig. 3). For protein BBL, the local contacts with  $L_c \leq 6$  and some nonlocal contacts with Lc > 8–12 are formed quickly after a short initial stage [Fig.3(a)]. The nonlocal contacts with  $L_c = 13$ –22 are half formed, and those with  $L_c > 23$  are basically not formed. The local and nonlocal contacts reach their equilibrium values at about  $4\times 10^4$ 

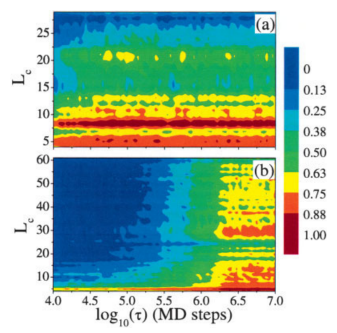


Fig. 3. The formation probabilities of contacts with different loop lengths  $L_c$  during the folding. The colors show the probabilities of contacts averaged over 100 trajectories for protein (a) BBL and (b) Cl2 at their  $T_f$ 

MD steps almost simultaneously. There is basically no further formation of contacts as the time evolves. At temperature below or above  $T_{\rm f}$ , the evolutions and distributions of the nonlocal contacts are similar. However, the situation is quite different for protein CI2. The local contacts are formed first, and all the nonlocal contacts begin to be formed cooperatively at about  $6.2\times10^6$  MD steps [Fig. 3(b)]. This formation time is apparently about 150 times longer than that of the local contacts. The nonlocal contacts show an all-or-none behavior when the temperature is below or above the transition temperature  $T_{\rm c}$ 

 $T_f$ . Clearly, these relate to the folding mechanisms of two kinds of proteins. It is noted that the formation of the local contacts for two proteins is similar, indicating that the formation of local contacts may be independent of the folding manners. Nevertheless, from our results, two proteins with well-defined folding mechanisms behave distinctly in their nonlocal contacts. Such differences in the features of the nonlocal contacts may act as an indicator for their folding behaviors. Furthermore, the topological structure of a protein may determine the formation of the nonlocal contacts which dominates the folding behavior.

The formation of nonlocal contacts can greatly limit the conformational searching of the protein molecule due to the geometrical restrictions. A few of nonlocal contacts may essentially outline the global topology of the chain. Therefore, the nonlocal contacts tend to make the convex shape of the landscape which induces a folding with a cooperative manner. These contacts generally relate to the so-called folding nucleus. The richness of nonlocal contacts could strengthen the cooperativity, and the deficiency of

the nonlocal contacts will decrease the barrier, even waive off the barrier totally. Microscopically, the larger the number of nonlocal contacts per residue, the more stable the local packing structure is, although the packing of structural units of the protein is initiated by the formation of local contacts. Thus, the folding is not in a gradual mode, but is cooperatively progressed. However, it is found that for the downhill folder there is no nucleus, so no barriers. Some nonlocal contacts are formed early and are not broken, and some never be formed. Nevertheless, for protein CI2, the formation of nonlocal contacts is flexible with an all-or-none feature, thus a two-state behavior.

#### **Statistics on the Native Contacts**

To verify that the lack of the cooperativity for BBL is due to the deficiency of nonlocal interactions, let us calculate the statistics on the contact distribution and the folding behavior over some well-studied proteins. 46 The statistics on the number of nonlocal contacts per residue  $N_N$ , the number of local contacts per residue  $N_L$ , the ratio of them  $R_{LN} = N_L/N_N$ , and the related results of various features based on our simulations are listed in Table I. One can see clearly the differences between the downhill folder and other proteins. The downhill folder, such as the barrierless folder BBL, has a small number of nonlocal contacts per residues as  $N_N = 0.730$ , which is rather smaller than those of proteins of two-state folding or three-state folding, for example,  $N_N = 1.875$  for CI2. As a result, the ratio  $R_{LN} =$ 1.111 of BBL is rather larger than those of the other proteins, for example,  $R_{LN}=0.250$  for CI2. It is obvious that protein CI2 has large propensity in nonlocal contacts, and protein BBL is deficient in nonlocal contacts. This statistics suggests that the number of nonlocal contact per residue is relevant to the folding mechanism of the downhill folder.

Note that the statistics listed in Table I is independent of the chain lengths. As an example, let us consider the β-hairpin, a short secondary structural element, which is rather short compared with BBL. The β-hairpin is argued to fold with two-state manner.47 In our simulation, a barrier in the free energy profile is also found (data not shown). The statistics for the  $\beta$ -hairpin gives  $N_N = 1$ , which is slightly larger than that of the downhill folder BBL and is obviously smaller than those of other two-state folders. At the same time, the ratio  $R_{LN}$  is quite small and indicates a two-state folder. Actually, the free-energy barrier of  $\beta$ -hairpin is located at the region with rather small values of Q = 0.1.47 Consequently, the folding trajectory is, in some sense, similar to the downhill folder after passing this barrier. In addition, some other proteins are also checked. It is found that some of these proteins would fold in a weak cooperative manner and have low barriers when their values of number of nonlocal contacts per residue  $N_N$  are around 1, such as the three-helix bundle,  $^{48,49,50}$  the protein En-HD,  $^{51-53}$  and so on.

Thus, whether a protein is a two-state folder or a downhill folder can easily be identified from the distribution of the contacts, namely the number of nonlocal contacts per residue,  $N_N$ , from the native conformation

TABLE I. The List of Various Statist	ical and Folding Feature	es for 16 Cooperative Proteins and BBL

		Structure			Chain				
Index	Folding Behavior	Class	Protein Name	PDB Code	Length	$R_{LN}$	$N_L$	$N_N$	$\kappa_2$
1	Two-state	α	Monomeric λ-repressor <sup>a</sup>	1LMB	80	0.832	1.050	0.263	0.613
2		α	ACBPa	2ABD	86	0.856	1.105	1.291	0.603
3		α	Cytochrome c <sup>a</sup>	1HRC	104	0.457	0.769	1.683	0.590
4		α	En-HD <sup>b</sup>	1ENH	54	0.869	0.981	1.130	0.362
5		β	Tenascin <sup>a</sup>	2AIT	74	0.157	0.419	2.662	0.854
6		β	$CspB^a$	1CSP	67	0.140	0.299	2.134	0.692
7		β	α-spectrin SH3 <sup>a</sup>	1SHG	57	0.123	0.298	2.421	0.829
8		β	β-haipin <sup>d</sup>	_	16	0.267	0.250	0.938	0.512
9		α/β	$CI2^{a}$	1COA	64	0.250	0.469	1.875	0.800
10		α/β	Ubiqution <sup>a</sup>	1UBQ	76	0.284	0.553	1.947	0.728
11		α/β	ADA2h <sup>a</sup>	1PBA	81	0.365	0.617	1.691	0.731
12	Three-state	α	Thre-helix Bundle <sup>c</sup>	1BDD	46	1.049	0.935	0.891	0.358
13		α	Barstar <sup>a</sup>	1BTA	89	0.526	0.899	1.708	0.762
14		β	$\mathrm{CD2^{a}}$	1HNG	98	0.135	0.306	2.265	0.743
15		α/β	Barnase <sup>a</sup>	1BNI	108	0.260	0.537	2.065	0.698
16		α/β	$\mathrm{CheY}^{\mathrm{a}}$	3CHY	128	0.405	0.797	1.969	0.790
17	Downhill	α	BBL	1BBL	37	1.111	0.811	0.730	0.337

<sup>&</sup>lt;sup>a</sup>Referred to Ref. 46

 $R_{LN}$  is the ratio of number of local contacts per residue;  $N_L$  to the number of nonlocal contacts per residue  $N_{N^5}$   $\kappa_2$  is a value to quantify cooperativity of protein folding. <sup>26–28</sup> Note that protein barnase was described as a two-state folder in equilibrium denaturation, <sup>57</sup> while was argued as a three-state from kinetics. <sup>58</sup> However in our folding simulations, the results are more similar to the observation in Ref. 58 Thus, we classify protein barnase as a three-state folder, which is also consistent with the classification in Ref. 46

without any detailed folding simulations. That is, the statistics on the contacts can provide some judgment for possible folding mechanisms of proteins. Note other statistics with different definitions, for example, with different cutoff distances, for the contacts is also made (data not shown). It is found that they do not change our conclusion, only affect the value of  $N_N^c$  slightly (see following discussion).

#### Quantification of the Cooperativity

In order to get a quantitative relationship between  $N_N$  and the folding cooperativity, a factor  $\kappa_2$  is used to measure the folding cooperativity. The results are listed in the last column of Table I. A strong correlation between  $N_N$  and  $\kappa_2$  with a correlation coefficient r=0.878 is found [see Fig.4(a)]. The values of  $\kappa_2$  for various proteins are close to the fitting line. Clearly, protein BBL has the smallest value of  $N_N$ , that is, the lowest cooperativity among all the proteins studied. It is noted that the cooperativity of protein CI2 is high and the value of  $N_N$  is large. A plot of  $R_{LN}$  versus  $\kappa_2$  is shown in Figure 4(b). It is found that the correlation coefficient is r=-0.807, and the points are dispersive relatively.

Thus,  $N_N$  can be used to characterize the cooperativity of the folding and is considered as an important topological parameter describing the proteins. From the cooperativity factors  $\kappa_2$  and the statistics on the values of  $N_N$  shown in Table I, a crossover around  $N_N=0.9$  could be defined to separate the two-state folder and the downhill folder.

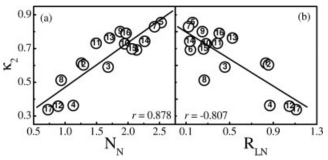


Fig. 4. The cooperativity factor  $\kappa_2$  versus the number of nonlocal contact per residue  $N_N$  (a), and the ratio between the number of local contact and the number of nonlocal contact  $R_{LN} = N_L/N_N$  (b) for 17 proteins listed in Table I. The number in the circle is the index of the related protein in Table I. r is the correlation coefficient.

#### **Exemplification by Mutants**

Finally, to further illustrate the relationship between the parameter  $N_N$  and the cooperativity and to validate the crossover of  $N_N$  for different folding behaviors, a study based on mutation is made. If the downhill folder is solely determined by the value of  $N_N$ , one can change the two-state folder into a downhill one by reducing the related value of  $N_N$  via mutations, and vice versa. Taking an example, protein monomeric  $\lambda$ -repressor (PDB code: 1LMB) has 80 residues and is an  $\alpha$ -class protein with its  $N_N=1.263$ . The mutation for this protein is easy since its value of  $N_N$  is close to that of  $N_N^c$  (see Table I).

Using the method of computational mutations, six mutants, that is, M1 to M6, have been obtained (see Fig. 5). The  $N_N$  values of M1 to M5 are distributed between those

<sup>&</sup>lt;sup>b</sup>Referred to Refs. 51–53

 $<sup>^{\</sup>rm c}$ Referred to Refs. 48–50

 $<sup>^{\</sup>rm d}\beta\text{-haipin}$  is the C-terminal fragment (41–56) of protein GB1.  $^{47}$ 

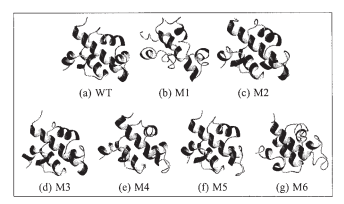


Fig. 5. The wild-type structure and the stable structures of protein monomeric  $\lambda$ -repressor and its six mutants. The mutations for mutant M1 are Leu50Gly, Phe51Ala, Leu65Gly, and Ile80Gly, and the mutations for mutant M2 to M6 include some of Gln33Ala, Leu50Ala, Phe51Ala, Leu57Ala, Leu65Ala, Phe76Ala, and Ile80Ala, respectively. Because only a few residues are mutated, the changes of the structures are small. The values of cRMSD between the stable structures of M1 to M6 and the structure of the wild type are 8.451 Å, 7.090 Å, 7.392 Å, 7.222 Å, 7.626 Å, and 6.858 Å, respectively.

of the native structure and M1 structure. Clearly, M1 has the smallest  $N_N$  value, that is,  $N_N=0.825$ , and M6 has a  $N_N$  value larger than that of wild type, that is,  $N_N = 1.375$ . For each mutant, the related folding simulations are made using the Gō-like model, and the heights of free energy barriers at their related  $T_f$ s and cooperativity factors  $\kappa_2$ are obtained (see Fig. 6). From Figure 6, it is seen that the heights of the free energy barriers and the cooperativity factors  $\kappa_2$  for these six mutants decrease as the related values of  $N_N$  decrease. Note that the free energy barrier of the wild type obtained from our simulations is about 2.5  $k_BT$ . For M1, the height of the barrier is only about 0.6  $k_BT$ and the cooperativity is quite low with  $\kappa_2 = 0.373$ , indicating clearly a downhill or downhill-like folding behavior at temperature  $T_f$ . This is really consistent with our argument mentioned above. The inset in Figure 6 shows the free energy profile with a very small barrier after the mutation. Thus, we believe that when the value of  $N_N$  of a protein is smaller than  $N_N^c$ , the folding is of downhill and the height of the free energy barrier will be smaller than 1  $k_BT$ , and when it is extremely smaller than  $N_N^c$ , such a barrier will disappear.<sup>54</sup> As a result, the protein can be a conformational rheostat.

#### Remarks on Recent Experiments

In order to make a further check, let us compare our results with recent related work. In a review article,  $^{55}$  Eaton and coworkers made a summary on the fast and ultrafast folding behavior of proteins. They argued that generic N-residue single-domain proteins fold at or near the speed limit, predicted to be approximately N/100  $\mu s$  by both experimental and theoretical approaches, and may have no free energy barrier. As listed in the review, indeed, the free energy barriers for some fast or ultrafast folding proteins are quite small. Thus, 12 fast folding proteins with  $\tau_{\rm folding} < 100~\mu s$  were considered to be the candidates of downhill folder.  $^{55}$  Among them, seven proteins fold very

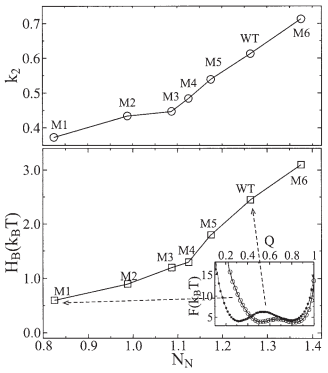


Fig. 6. The cooperativity factors  $\kappa_2$  (circle) and the height of free energy barrier  $H_B$  (square) at  $T_r$  for the wild type (labeled as WT) and six mutant structures of protein Monomeric  $\lambda$ -repressor shown in Figure 5. The inset shows the free energy profiles for the wild type (solid circle) and the mutant M1 (circle).

fast, with  $\tau_{\rm folding} < 10~\mu s$ . Using our definition, their related values of  $N_N$  are calculated and plotted against the folding times  $\tau_{\rm folding}$  in Figure 7.

Among the seven ultrafast folding proteins, three [tryptophan cage (1L2Y), BBA5 (1T8J), and Villin headpiece subdomain (1VII)] have their values of  $N_N = N_N^c$ , and other three [albumin binding domain (1PRB),  $\alpha_3$  D (2A3D), and B domain of protein A (1BDC)] have their values  $N_N = N_N^c$ . Therefore, the former three proteins could be potential candidates of downhill or downhill-like folders because they fold very fast with  $\tau_{\rm folding} < 10~\mu s$  and the barriers of folding are quite small as listed by Eaton and coworkers.<sup>55</sup> Our simulations for proteins BBA5, HP-36, and Trp-cage are also consistent with such a prediction. Differently, the later three proteins show weak two-state folding behavior and have low barriers because they are in the crossover between the two-state and the downhill folding (see Ref. 55 and the references therein). Note that as argued by Sosnick and coworkers, proteins or peptides with folding speed  $\tau_{folding} < 6~\mu s$  are not enough to be barrierless. To fold in a downhill manner, the proteins or peptides should fold extremely rapidly.<sup>56</sup> Here, in this work, our criterion based on the structural features of proteins may roughly outline the crossover or the boundary of the potential candidates for downhill folding if the proteins fold fast enough and their  $N_N = N_N^c$ . It is worthy to note that although protein Cytochrome  $b_{652}$  (PDB ID: 1QQ3) has  $N_N$  $\leq N_N^c$  and  $au_{
m folding} < 10 \mu 
m s$ , its folding rate was measured in

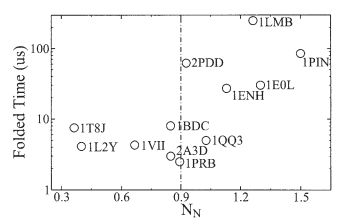


Fig. 7. The number of nonlocal contacts per residue  $N_N$  and the folding time  $\tau_{\rm folding}$  of 12 ultrafast folding proteins collected by Kubelka and colleagues. The labels are the PDB identities of the protein structures. They are tryptophan cage (1L2Y), BBA5 (1T8J), villin headpiece subdomain (1VII), WW domain Fin (1PIN), WW domain FBP28 (1E0L), peripheral subunit binding (2PDD), albumin binding domain (1PRB), engrailed homeodomain (1ENH),  $\alpha_3$ D (2A3D),  $\lambda$ -repressor (1LMB), B domain of protein A (1BDC), and cytochrome  $b_{652}$  (1QQ3). The dashed line indicates the rough crossover  $N_N^2$  = 0.9. Note that 1BDC is used here because it was discussed in Ref. 54 and its related  $N_N$  value is calculated based on the first structure among 10 structures in the file of 1BDC. Differently, in our simulations for Table I, the averaged NMR structure (corresponding to the code 1BDD of the same protein) is used.

2.2M GndHCl instead of the natural physiologic condition. This implies that the case may be different and complicated.

#### **CONCLUSION**

In this work, the downhill folding of protein BBL is simulated with the Gō-like model, and is compared with the two-state folding. There are obvious differences both in kinetics and in thermodynamics. The basic feature of the folding for protein BBL is of weak cooperativity, and there exists no obvious free energy barrier. The thermodynamic stable state of downhill folder, corresponded to the single minimum in the free energy landscape, changes continuously from high Q to small Q values as the temperature increases. That is to say, unlike some typical two-state proteins, downhill folder is more like a molecular rheostat. In addition, in our study the downhill folder candidates can be identified from the statistics on the features of contacts. If the number of nonlocal contacts per residue is smaller far than the crossover around 0.9, the protein can be identified as a candidate of downhill folder. On the contrast, for the two-state proteins, the nonlocal contacts are always rich, which leads to a high cooperativity of folding. Therefore, the content of the nonlocal contacts per residue could act as an important quantity to predict the folding behaviors of proteins.

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