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# Domain-Specific Folding Kinetics of Staphylococcal Nuclease Observed through Single-Molecule FRET in a Microfluidic Mixer\*\*

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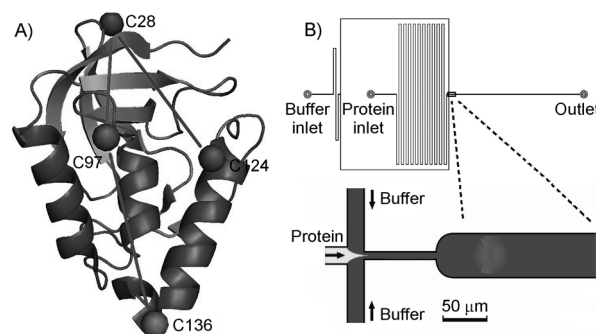
Characterization of the free-energy landscape in protein folding is key to understanding the “folding code” contained in the sequence.<sup>[1]</sup> Multidomain proteins represent a predominant fraction of the whole proteome in prokaryotic and—even more—in eukaryotic cells.<sup>[2]</sup> It is thus interesting to find out how these multidomain proteins fold. A number of multidomain proteins have been shown to fold with multistate kinetics before the native population appears,<sup>[3]</sup> and the information is usually extracted from elaborate modeling.<sup>[4]</sup> It would be nice if one could directly probe the heterogeneous populations of unfolded and folded ensembles along the folding pathway from the initial denatured state to the final native conformation.

Single-molecule fluorescence resonance energy transfer (smFRET) offers a powerful tool to probe the heterogeneous system of protein molecules.<sup>[5]</sup> Meanwhile, microfluidic laminar-flow mixers have facilitated kinetic measurements of biological systems by ultrafast mixing.<sup>[6]</sup> The combination of smFRET and a kinetic microfluidic mixer was first introduced by Lipman et al. in 2003 to rapidly trigger protein folding and to reveal the evolution of folded and unfolded species under non-equilibrium conditions.<sup>[7]</sup> Since then, this method has been developed to achieve a shorter mixing time,<sup>[8]</sup> and the shortest record of 0.2 ms was reported very recently.<sup>[9]</sup> Other types of microfluidic devices, such as a coaxial 3D mixer<sup>[10]</sup> and a mixer with enhanced photostability,<sup>[11]</sup> have also been proposed. The combined smFRET and microfluidic mixing technique has been applied to the protein folding inside the GroEL<sup>[12]</sup> and the folding of an intrinsically disordered protein.<sup>[9]</sup> Staphylococcal nuclease (SNase), consisting of an N-terminal  $\beta$ -sheet domain and a C-terminal  $\alpha$ -helical domain, has been studied as a model for multidomain protein folding and exhib-

ited a complex kinetic behavior.<sup>[4,13]</sup> Our previous equilibrium smFRET studies have shown that a domain-specific collapse occurs in the early stages of the refolding process.<sup>[13c]</sup>

Herein, we report our non-equilibrium smFRET studies in a microfluidic mixer to directly probe the transition rate from the unfolded state to the native folded state after the collapse. We labeled donor and acceptor dyes at selected sites to detect the kinetics of the conformational reorganization of the subdomains and the global molecule in the refolding landscape. By examining the unfolded and folded states, the kinetic measurements suggested that different domains adopt different searching pathways to reach the native conformation.

We constructed three mutants K28C/K97C, K97C/K136C, and K28C/H124C (Figure 1A) and labeled them with fluorescent dyes using the same procedure reported previously.<sup>[13c]</sup> The mutants were site-directed labeled with thiol-reactive fluores-



**Figure 1.** Labeling scheme on SNase and schematics of the microfluidic mixer. A) Mutants for the folding kinetic measurements: K28C/K97C spans the  $\beta$ -sheet domain, K97C/K136C spans the  $\alpha$ -helical domain, and K28C/H124C spans the two domains. B) Schematics of the microfluidic mixer. The denaturant was rapidly diluted in the neck region, where the protein also accomplishes the initial collapse. Then, the unfolded protein initiates its folding in the detection channel.

cence dyes of Alexa Fluor 546 and Alexa Fluor 647. A more detailed description of the mutation and labeling process is given in the Supporting Information. The mutant of K28C/K97C was designed to probe the  $\beta$ -sheet domain. K97C/K136C was for the  $\alpha$ -helical domain. Finally, the mutant of K28C/H124C was constructed to probe the conformational change of the whole molecule while maintaining a sensitive distance comparing to the Förster radius,  $R_0$ , between the donor–acceptor dye pair. The structure, the stability, and the enzymatic activity of the mutants K28C/K97C and K28C/H124C have been characterized already.<sup>[13c]</sup> We verified K97C/K136C using far-UV circular dichroism (CD) spectroscopy (Figure S1 of the Support-

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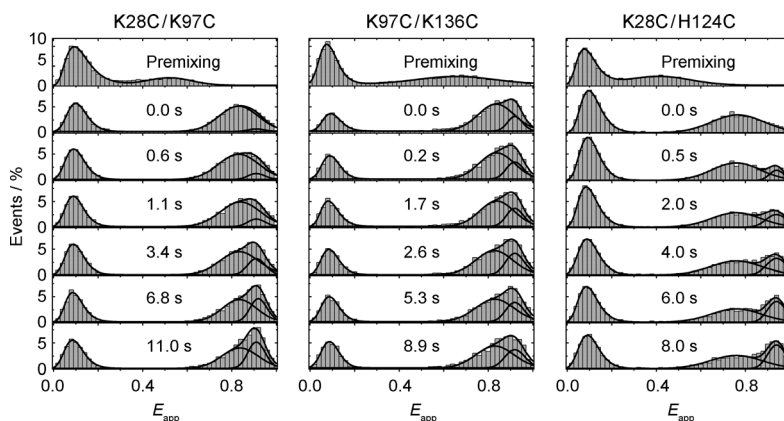
[\*\*] FRET: Fluorescence Resonance Energy Transfer

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ing Information) and the denaturation curves using the intrinsic fluorescence of Trp 140. The thermodynamic parameters were derived from the denaturation curves and are shown in Table S1 of the Supporting Information. Both results demonstrated that the K97C/K136C mutant kept the native structure well.

A microfluidic mixer was constructed (Figure 1B) following the essential ideas proposed by Pfeil et al.<sup>[8]</sup> and Lemke et al.,<sup>[11]</sup> with minor modifications to fit our need and capacity of micro-fabrication. The details and the characterization of the mixer have been described previously<sup>[14]</sup> (see also the Supporting Information). The major consideration of the design was to maintain a stable laminar flow for an optimal smFRET measurement with high photon bursts (maximum bursts of 80–100) and good signal-to-noise ratio (~100). The microfluidic mixer was aligned on a confocal microscope by an adaptor. The protein sample and the buffer were delivered into the inlets from two reservoirs, which were connected to compressed air. The mixing time of the microfluidic device was characterized to be 0.15 s<sup>[14]</sup> by the extremely fast process of protein collapse, suitable for current kinetic measurements. During the mixing time, the SNase molecule collapses with a very fast rate, while the transition from the unfolded to the folded state barely occurs.<sup>[14]</sup> The velocities at the detection points were measured by fluorescence correlation spectroscopy (FCS), simultaneously with the smFRET data acquisition, to calculate the definite time versus the position.<sup>[9,14]</sup> The labeled protein was in the unfolded state in 2 M GdmCl and was injected into the central inlet. The native buffer was delivered through the two side inlets which were originated from an entrance. The concentration of the labeled protein was less than 60  $\mu\text{M}$  in the detection channel by comparing the event-counting frequency with equilibrium experiments. The denaturant concentration was 0.49 M in the detection channel, which was determined by comparing the FRET efficiency of the unfolded state with that in the equilibrium experiments (see the Supporting Information). 1  $\mu\text{M}$  unlabeled wild-type SNase purified by desalting column was added into both lines to prevent protein adhesion.

The setup for the smFRET measurements was essentially identical to that reported previously (see also the Supporting Information).<sup>[13c]</sup> The residence time of the protein molecule in the laser focus was on the order of 1 ms. As individual labeled protein molecules flowed through the laser focus, which was 10  $\mu\text{m}$  above the coverglass in the detection channel, fluorescent photons from the donor and acceptor were counted separately by two avalanche photodiode (APD) detectors and the histograms of FRET efficiency were generated, with each histogram consisting of at least 3000 identified events at an appro-

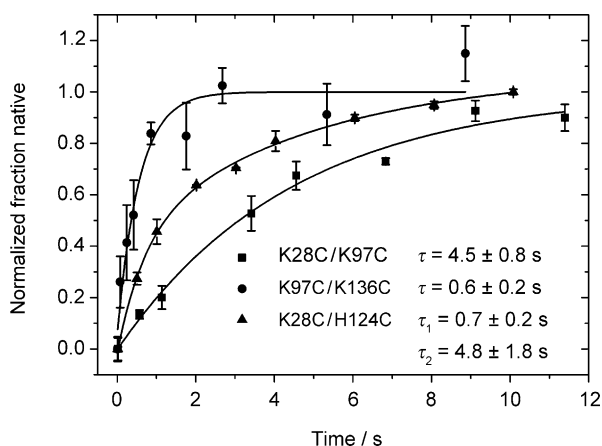


**Figure 2.** FRET efficiency histograms (at various times) for the mutants. The top panel shows the unfolded state in the central channel before mixing. The time zero was taken at the conjunction of the neck and the detection regions.

priate threshold (the sum of the photon counts from the donor and acceptor channels) to pick up the signal bursts. By scanning the laser focus along the central line of the detection channel, the kinetic evolution of the FRET efficiency distribution after the mixing was mapped (Figure 2).

The initial protein collapse was extremely fast. The time resolution of our microfluidic mixer was limited by the mixing process and was unable to probe the rate of the collapse.<sup>[15]</sup> During the mixing in the neck region, the shift of the peak of the unfolded state was the result of the protein collapse from a loose form to a more compact form.<sup>[14]</sup> In the detection channel, we observed the growth of the peak associated with the native state as well as the decrease of the population of the unfolded molecules. Finally, the FRET efficiency histogram approached its equilibrium under the given GdmCl concentration. The unfolded and folded states were fitted using the Gaussian function and lognormal function, respectively.<sup>[5a,b]</sup> We also fitted the folded states using a beta function<sup>[13c,16]</sup> and found that both functions delivered identical results within the experimental error. The peak near zero was the background generated from molecules without active acceptor or impurities in solution.<sup>[17]</sup> The peak of the subpopulations in the FRET efficiencies remained constant, indicating that the mean end-to-end distances of the donor and acceptor in both unfolded and folded states did not change during the folding reaction. This observation showed that single-molecule kinetic measurements can resolve the unfolded and folded species so that it excluded the complexity encountered in an ensemble experiment.<sup>[4]</sup>

SNase has been shown to fold through the pathway of multiple intermediate states using the ensemble stopped-flow method by tracking the Trp fluorescence.<sup>[4,13a]</sup> In our case, the data of a single domain could be well fitted using a single-exponential curve, while a double-exponential curve was needed to satisfactorily fit the data of the whole molecule (see Figure 3 and Figure S4 of the Supporting Information). For the  $\beta$ -sheet mutant of K28C/K97C, the fitting yielded a folding relaxation lifetime of  $\tau = 4.5 \pm 0.8$  s. For the  $\alpha$ -helical domain of



**Figure 3.** Single-molecule folding kinetics of the different subdomains and the whole molecule.

K97C/K136C, the fitted lifetime was  $\tau = 0.6 \pm 0.2$  s. Modeling the folding of the global molecule of K28C/H124C generated lifetimes of  $\tau_1 = 0.7 \pm 0.2$  s and  $\tau_2 = 4.8 \pm 1.8$  s, corresponding to the lifetimes of the single-domain mutants well. Our observed folding lifetimes were comparable to the folding kinetics measured by ensemble CD spectra.<sup>[18]</sup> Here, our results could be assigned in a straightforward way: the  $\alpha$ -helical subdomain folds about ten times faster than the  $\beta$ -sheet subdomain. This result provided an alternative view from the previous CD study which says that the sheet-like chain conformations precede the main-chain folding reaction.<sup>[18]</sup> The subtle difference comes from the fact that the CD study mainly detects the changes of the secondary structure during the processes of collapse and folding, while the smFRET study mainly detects the distance between the residues under investigation. Another conventional tool in the protein-folding study is the tryptophan fluorescence, where the change of the tryptophan fluorescence reflects the variation of the microenvironment around the tryptophan residues instead of the whole domain.<sup>[4]</sup> It has been reported that proline isomerization occurs on the order of tens of seconds,<sup>[19]</sup> but we were unable to find such a process due to our short time window. The data sets gathered by using different tools represent different aspects of the complicated protein folding, and they are complementary to each other. The trait of the smFRET technique resides on its capability of directly identifying the unfolded and folded species and directly assigning the rate to a selected process by site-directed mutagenesis and dye labeling.

It was pointed out by Pfeil et al. that diffusion would erode the relationship between the reaction time and the position by the time on the order of  $t = a^2/2D$ , where  $a$  is the half width of the detection channel and  $D$  is the diffusion constant of the protein.<sup>[8]</sup> With  $D = 73 \mu\text{m}^2\text{s}^{-1}$  for the SNase molecule and  $a = 25 \mu\text{m}$  for our apparatus,  $t = 4.3$  s was derived. According to this rough criterion, our measured reaction-time constants could have a deviation from the true values. It is easy to argue that diffusion would make the observed reaction-time constant smaller than it should be, and the slower the reaction, the

bigger the deviation. Therefore, if the effect of diffusion were not negligible, the difference of the refolding times between the  $\alpha$ -helical and the  $\beta$ -sheet subdomains would be even bigger than what we observed.

In summary, we coupled smFRET and microfluidic mixing to directly explore the substructural folding kinetics of a protein with two subdomains after the initial collapse. Our conclusion is that the  $\alpha$ -helical subdomain of SNase folds faster than the  $\beta$ -sheet subdomain, so that the folding kinetics of SNase is subdomain-specific.

## Experimental Section

Further information on Methods, as well as Figures S1–S4 and Table S1, is available in the Supporting Information.

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