Channel of viral DNA packaging motor for real time kinetic analysis of peptide oxidation states

Shaoying Wang a, b, Zhi Zhou a, Zhengyi Zhao a, b, Hui Zhang a, Farzin Haque a, Peixuan Guo a, *

a College of Pharmacy, Division of Pharmaceutics and Pharmaceutical Chemistry; College of Medicine, Department of Physiology & Cell Biology; and Dorothy M. Davis Heart and Lung Research Institute, The Ohio State University, Columbus, Ohio 43210, USA
b College of Pharmacy, Markey Cancer Center, University of Kentucky, Lexington, KY 40536, USA

ABSTRACT

Nanopore technology has become a powerful tool in single molecule sensing, and protein nanopores appear to be more advantageous than synthetic counterparts with regards to channel amenability, structure homogeneity, and production reproducibility. However, the diameter of most of the well-studied protein nanopores is too small to allow the passage of protein or peptides that are typically in multiple nanometers scale. The portal channel from bacteriophage SPP1 has a large channel size that allows the translocation of peptides with higher ordered structures. Utilizing single channel conductance assay and optical single molecule imaging, we observed translocation of peptides and quantitatively analyzed the dynamics of peptide oligomeric states in real-time at single molecule level. The oxidative and the reduced states of peptides were clearly differentiated based on their characteristic electronic signatures. A similar Gibbs free energy ($\Delta G_0$) was obtained when different concentrations of substrates were applied, suggesting that the use of SPP1 nanopore for real-time quantification of peptide oligomeric states is feasible. With the intrinsic nature of size and conjugation amenability, the SPP1 nanopore has the potential for development into a tool for the quantification of peptide and protein structures in real time.

© 2017 Elsevier Ltd. All rights reserved.

1. Introduction

Living systems contain wide varieties of nanomachines with diverse structures and functions. The ingenious design of viral DNA packaging motors and their intriguing mechanism of action has triggered a wide range of interests among scientists in many different areas [1–3]. The portal protein is one of the essential components of the viral packaging motors with a turbine-like shape [4,5]. SPP1 is a dsDNA phage that infects Bacillus subtilis. The DNA packaging motor of SPP1 consists of a terminase composed of small (gp1) and large (gp2) subunits, portal protein gp6, and a two head completion proteins gp15 and gp16 [6,7] that power the encapsulation of 45.9 kbp genomic DNA [8]. The central core of the SPP1 motor is a portal channel, termed connector, which mainly forms a 13-subunit assembly in vitro and 12-subunit assembly inside phage particles [7,9]. Explicit engineering of the SPP1 portal protein is possible due to its available crystal structure [9,10]. The connector has an overall diameter of 16 nm and a height of 10.5 nm. The narrowest constriction of the internal channel is ~3 nm (Fig. 1) [9,10]. Portal protein is a critical component in many dsDNA bacteriophages and herpes viruses, which plays a critical role in genome packaging and ejection. Structural studies have shown that similar cone-shaped dodecameric structure is shared in portals from herpes virus and different tailed bacteriophages, such as phi29, SPP1, T4, and T3, even though their primary sequences do not show homology with varied molecular weight from 36 kDa, 56 kDa, 60 kDa and 59 kDa, respectively.

Nanopore technology has recently emerged as a real-time and high-throughput single molecule detection method, holding great potential for sensing a wide range of analytes, molecular diagnostics and DNA sequencing applications [11–19]. Solid state nanopores generated by microfabrication generally have less reproducible pore sizes and lack chemical and location selectivity [20,21]. Protein nanopores harvested in bacteria are homogenous in size and can be functionalized with probes, but commonly used nanopores such as z-hemolysin, MspA, and aerolysin have an
internal channel diameter of only ~1.3 nm [22–24]. Larger protein pores such as ClyA [25] and FhuA [26] are being developed for single molecule analysis. In search for alternative larger sized channels, we previously developed membrane-embedded phi29 motor channels [27–29] for single molecule sensing of nucleic acids [27,28,30–33], chemicals [34], peptides [35] or binding assays of antibody based on channel conformational change [30,36]. Herein, we utilized our recently developed membrane-embedded SPP1 motor channel [37,38] for quantifying the translocation and dynamics of peptide oligomeric states at single molecule level.

Elucidating the oligomeric states of proteins and peptides is critical for understanding their biological functions. A wide range of biophysical methods, such as X-ray crystallography, NMR, Circular Dichroism, Dual Polarization Interferometry, and Mass Spectrometry have been used to investigate folding and dynamic structural changes of peptides and proteins. However, these methods require expensive instrumentation and specialized labor. Nanopores offer an attractive alternative as they are intrinsically single molecule in nature requiring ultra-low sample volumes, are label free, amplification free, and function using a simple detection process requiring no specialized expertise. While the translocation of DNA and RNA have been studied extensively in biological nanopores [12,20,22,39], studies on translocation of protein or peptides are beginning to emerge [35,40–44]. The proteome can be an accurate and direct indicator of current health status of patients [45]. For example, early diagnosis and monitoring the changes of amyloid-β peptide and α-synuclein are critical for the management of Alzheimer’s and Parkinson’s disease [46].

Several studies demonstrating peptide translocation using nanopores have been published [47–60], but quantitative analysis and translocation validation remain challenging, since there is no technique available for amplifying protein substrates as in DNA or RNA qualification using PCR. The availability of only trace amounts of peptide or protein for analysis after translocation is far beyond the sensitivity threshold of classical protein detection methods. In this report, we used single molecule fluorescence microscopy to validate data obtained from resistive pulse technique to quantitatively study peptide translocation through SPP1 connectors and elucidate the structural conformations of peptides at the single molecule level. Due to the availability of crystal structure, this new nanopore can be explicitly engineered via site directed mutagenesis, and has the potential to be applied for biomarker analysis and early disease diagnosis.

2. Materials and methods

2.1. Materials

The phospholipid 1,2-diphytanoyl-sn-glycerol-3-phosphocholine (DPhPC) was obtained from Avanti Polar Lipids, Inc. Organic solvents (n-decane and chloroform) were purchased from Fisher Scientific, Inc. and TEDIA, Inc., respectively. TAT peptide was custom-ordered from GenScript, Inc. All other reagents were purchased from Sigma or Fisher, if not specified.

2.2. Cloning and purification of the SPP1 connector

Gene gp6 encoding SPP1 portal was synthesized and cloned into PET3 vector between Nhel and BamHI by GenScript, Inc. His-tag was inserted into the C-terminal of the connector for purification. Then the plasmid was transformed into BL21 (DE3) for expression and purification. The purification procedure has been described previously [27,29]. Briefly, the transformed bacteria were cultured in 10 mL LB medium overnight at 37 °C. The bacteria were transferred to 1 L of fresh LB medium. When the OD600 reached ~0.5–0.6, 5 mM IPTG was added to the medium to induce protein expression. The bacteria were collected by centrifugation after 3 h continuous culture. Bacteria was lysed by passing through French press. The cell lysate was separated by centrifugation and supernatant containing expressed protein was collected. The supernatant was purified with Nickel affinity chromatography (Novagen) [61]. His Binding Buffer (15% glycerol, 0.5 M NaCl, 5 mM Imidazole, 10 mM ATP, 50 mM Tris–HCl, pH 8.0), and the cleared lysate was loaded onto a HisBind® Resin Column and washed with His Washing Buffer (15% glycerol, 0.5 M NaCl, 50 mM Imidazole, 50 mM Tris–HCl, pH 8.0). The His-tagged connector was eluted by His Elution Buffer (15% glycerol, 0.5 M NaCl, 0.5 M Imidazole, 50 mM Tris-HCl, pH 8.0). The purified connector from chromatography was further purified by a 15–35% glycerol gradient ultracentrifugation at 35 K for 2 h. Before incorporating into liposomes, the purified connector was dialyzed against buffer (0.5 M NaCl, 50 mM Tris-HCl, pH 8.0) to remove excess glycerol.

2.3. Insertion of the connector into planar lipid bilayer

The protocol for the incorporation of connectors into lipid bilayer has been reported [27–30,62]. Briefly, 150 μL of 10 mg/mL DPhPC lipids in chloroform was placed in a round bottomed flask and the chloroform was evaporated under vacuum using a rotary evaporator (Buchi). The dehydrated lipid film was then rehydrated with 300 μL buffer containing 250 mM sucrose and purified connectors (0.5–1 mg/mL). The suspension was thoroughly vortexed and maintained at 45 °C (above the phase transition temperature of DPhPC) followed by extrusion through 400 nm polycarbonate membranes (Avanti Polar Lipids) to generate uniform sized liposome/connector complexes.

Planar bilayer lipid membranes (BLMs) were generated in a BCH-1A horizontal BLM cell (Eastern Scientific). A Teflon partition...
with a 200 μm aperture was placed in the apparatus to separate the BLM cell into cis- (top) and trans- (bottom) compartments. The aperture was pre-painted with 0.5 μL of 3% (w/v) DPhPC in n-hexane. A conducting buffer (1 M KCl, 5 mM HEPES, pH 7.8) was added to both the top and bottom compartments of the BLM cell, and Ag/AgCl electrodes were placed in the buffer of each compartment. The electrode in the trans-compartment was connected to the headstage of an Axopatch 200B amplifier (Axon Instruments), and the electrode in the top compartment was grounded. A planar lipid bilayer was formed by painting the aperture with 0.5 μL of 3% (w/v) DPhPC in n-decane. 1 μL of the diluted liposome/connector complex was added to the cis-compartment directly to fuse with the planar lipid bilayer to generate membrane embedded nanoropes.

2.4. Electrophysiological measurements

The headstage and Axopatch 200B patch clamp amplifier were connected to a DigiData 1440 analog-digital converter (Axon Instruments, Inc.) to monitor and record electrochemical currents through BLMs [27–29]. The current recordings were low-pass filtered at a frequency of 5 kHz. The sampling frequency was 200 KHz in all experiments, unless otherwise specified. The data were recorded with pClamp 9.1 software (Axon Instruments, Inc.), and analyzed with the Clampfit module of pClamp 9.1 and OriginPro 8.1 (OriginLab Corporation).

2.5. Peptide translocation experiments

TAT peptide (Cys-Tyr-Gly-Arg-Lys-Lys-Arg-Gln-Arg-Arg-Arg) with a final concentration of 23 μg/mL was premixed with the conducting buffer before the insertion of connector. All experiments were conducted at least three times and similar results were obtained. For quantitative validation, Cy3 fluorophore was conjugated to TAT peptide by sulhydryl-maleimide chemistry and purified by HPLC. Multiple channels were inserted into bilayer lipid membrane and then Cy3-TAT peptide was added into cis-chamber. Samples were collected from the trans-chamber after 0, 20, 40, and 60 min. The errors represent mean ± standard deviation determined by a Gaussian fit of the data.

2.6. Single molecule fluorescence imaging

Samples collected from the patch clamp were incubated with the ozone pre-treated glass substrate for 10 min before imaging to ensure sufficient adsorption. A 532 nm green laser was used for the excitation of Cy3. A 60 × objective (N.A. = 1.4, oil immersion) was used for fluorescence imaging. The signals were recorded using Andor iXon 887 V electron multiplied CCD camera. Images were taken with an exposure time of 500 ms. The number of spots in the images were counted using iSM5 software [63]. These fluorescence spots in Fig. 4a at 0 point may come from the impurity of the buffers or minor imperefection of the glass surfaces, which give background noise. Comparing to the signals from the translocated peptides, the number of background fluorescence spots was not significant. In the plot of Fig. 4b, the background spot numbers from the glass surface itself was subtracted for each time points. For data analysis, the number of spots at 0 min time point was subtracted from the calculation when counting the translocation peptide signals. The errors represent mean ± standard deviation from three independent imaging from within one experiment. Three independent repeats were performed and similar trend was obtained.

3. Results

3.1. Characterization of SPP1 reengineered nanopore embedded in a lipid bilayer

Structural analysis revealed that the central region of the SPP1 connector shows slight hydrophobicity compared with the flanking regions at the N- and C-terminal ends which are more hydrophilic (Fig. 1a–b). We reengineered the SPP1 connector by inserting a 6 × His-tag at the C-terminal end along with a 6 × glycine linker for end-flexibility. The presence of His-tag enhanced the hydrophilicity of the C-terminal, thus making the hydrophilic–hydrophobic–hydrophilic layers of the connector more distinct, which is necessary to mimic the lipid bilayer architecture. After His-tag column purification, the purified protein was further purified by 15–35% glycerol gradient ultracentrifugation to further purify SPP1 portal complex from the single subunit (Suppl. Fig. 1). Bovine serum albumin (BSA) served as a marker to indicate the location of single subunit of SPP1 connector. BSA with molecular weight of 60 KDa, which is almost equal to a single subunit (58 KDa) of the SPP1 connector, centered at fraction 27, whereas the majority of SPP1 portal sample was at fraction 17, indicating that the single subunit and portal complex can be well separated.

To incorporate SPP1 portal channel into planar lipid membranes, a two-step procedure was employed [27]. DPhPC lipids in chloroform were first dehydrated to remove organic solvents and then rehydrated with buffer containing 250 mM sucrose and purified connectors. The multi-lamellar lipid-connector suspension was then extruded through 400 nm polycarbonate membrane filters to generate uniform unilamellar liposomes with the connector embedded in the membrane. The resulting lipidosome-connector complex was fused with a planar lipid membrane to generate planar membrane-embedded SPP1 nanopore. Since the connector is not a membrane protein, direct incubation of the connector with a planar lipid bilayer did not result in connector insertion into the lipid bilayer. Single channel conductance assay was performed to measure the electrophysiological properties of membrane-embedded SPP1 connectors. The connector insertion steps were observed as distinct stepwise increase in conductance as revealed in a continuous current trace (Fig. 2a). The insertion of single portal channel results in ~200 pA in current jump under an applied potential of ±50 mV in conducting buffer (1 M KCl, 5 mM HEPES, pH 8). Occasionally ~400 pA current jumps were observed, attributed to simultaneous insertion of two connectors. The average conductance of reengineered SPP1 connectors is 4.27 ± 0.27 nS (Fig. 2b). The conductance is uniform without displaying any voltage gating phenomena under the reported conditions of ±50 mV (Fig. 2c). At voltages greater than ±100 mV, SPP1 connector displayed discrete stepwise gating of the channel [37].

3.2. Characterization of peptide translocation through reengineered SPP1 connector

A positively charged 12 amino acid TAT peptide was used in the translocation studies with sequence Cys-Tyr-Gly-Arg-Lys-Lys-Arg-Gln-Arg-Arg-Arg. Due to the presence of cysteine at the N-terminus, the peptide forms a dimer by disulfide bond under physiological conditions, which is confirmed by mass spectroscopy (data not shown). In the absence of peptide, the current trace was quiescent. In contrast, when the peptide was premixed with the conducting buffer in both cis- and trans-chambers, a burst of transient blockage events was observed immediately after the insertion of connector in the lipid membrane (Fig. 3a). As the peptide concentration was increased from 1.25, 2.5, 3.75, to 5 μg/mL, the density of current blockage events increased with a
corresponding linear increase in peptide translocation rate from $2.33 \pm 1.54$, $7.5 \pm 1.66$, $10.17 \pm 3.58$, to $16.96 \pm 6.01$ events per second (Fig. 3b). One parameter used to characterize the translocation was the current blockage percentage, calculated as the ratio of current blockade resulting from peptide translocation to the open current of one portal channel, expressed as $\left[1 - \frac{\text{current peptide}}{\text{current open channel}}\right]$. The distribution of current blockage was relatively broad with a major peak centered at $55.1 \pm 3.0\%$, determined by Gaussian fit of the data (Fig. 3c). Another parameter was the dwell time ($t$), the time taken for the peptide to traverse from one end of the connector to the other end. The dwell time distribution followed an exponential decay with a rate constant of $0.84 \pm 0.09$ ms$^{-1}$ (Fig. 3d).

3.3. Quantitative validation of peptide translocation by single molecule fluorescence imaging

To validate the translocation of peptide through SPP1 connector, we conducted single molecule fluorescence imaging of samples collected from patch clamp experiments. HPLC purified Cy3 conjugated TAT peptide at a final concentration of $2.5$ ng/μL was added to the cis-chamber after stable insertion of SPP1 connectors in the lipid bilayer. Under an applied negative trans-membrane voltage, the positively charged peptide translocated through the portal channel to the trans-chamber. $50$ μL samples were collected from the trans-chamber at $0, 20, 40,$ and $60$ min after addition of Cy3-TAT peptide and loaded onto glass coverslips. The positively charged peptide can bind to the negatively charged glass surface through charge-charge interactions and appear as individual fluorescent spots (Fig. 4a). Fluorescence imaging revealed that the number of Cy3 spots in the field of view increased over the time course of $60$ min (Fig. 4b). In contrast, in a control experiment under the same conditions but in the absence of portal channel, very few Cy3 spots were observed, compared to the sample containing SPP1 connector (Fig. 4b). Since TAT is a membrane penetrating peptide, it is conceivable that a small fraction could potentially pass through the lipid bilayer, contributing to the small increase in the fluorescence background signal.

3.4. Kinetic analysis of TAT conformational status in real time

The focus of this study was to investigate whether the SPP1 connector facilitated the translocation of TAT peptides through the lipid bilayer.
motor channel can be used for kinetic analysis of peptides in real time. TAT peptide with and without a cysteine was used as a model system to study oxidation states. Structural analysis of TAT sequence using the computer program PEP-FOLD [64] revealed that without forming a disulfide bond, the TAT only adopts one α-helical conformation (Fig. 5). However, in the presence of a cysteine at the N-terminus, the peptide can be oxidized into a dimer by forming a disulfide bond between two N-terminal cysteines. The presence of two states, the oxidized and the reduced conformations, were confirmed by translocation studies with the SPP1 portal channel. Under an oxidized state, the current blockage distribution by TAT displayed a major peak, centered at 55.1 ± 3.0% (Fig. 5a), with a minor peak centered at 28.5 ± 1.9%. However, after adding the reducing agent TCEP ([tris(2-carboxyethyl)phosphine]) to break the disulfide bond, the blockage distribution significantly changed. The peak of 55.1 ± 3.0% decreased significantly over time, while the majority of the blockage distribution shifted to 28.5 ± 1.9%. This indicated that most of the peptide passing through the channel is in its single α-helical conformation (Fig. 5b).

Current blockage distribution of the reduced TAT peptide was similar to Cy3-TAT signature with TCEP treatment, since the conjugation of Cy3 prevented the de bond, the blockage distribution significantly changed. The presence of a de bond between two N-terminal cysteines. The presence of some 55% current blockade events are observed. This can be attributed to a few unreacted TAT dimers in solution or two Cy3-TAT monomers passing through the channel at the same time.

To quantify the kinetics parameter, we further examined the conformational changes of TAT peptide upon addition of TCEP and recorded the change of the current blockage profile in real time (Fig. 6). It was found that current blockade signature progressively shifted from predominantly 55.1 ± 3% (oxidized dimer) to 28.5 ± 1.9% (reduced monomer), representing the real-time kinetic process of disulfide bond reduction (500 μM TCEP, Fig. 6a). The TCEP reduction process can be described as a pseudo-first order reaction due to the nearly infinite TCEP amount and the reaction equation is described as follows:

\[ \text{TAT} - \text{TAT} + \text{TCEP} \xrightarrow{H_2O} 2 \text{TAT} + \text{O} - \text{TCEP} \]

\[ C_0 \quad C(t) \quad \varepsilon C_0^r \quad 0 \]

\[ C_i \quad C(t) - (C_0 - C_i) \quad 2(C_0 - C_i) + \varepsilon C_0 \quad C_0 - C_i \]

\[ \frac{dC_i}{dt} = -kC_i[(C(t) - C_0 - C_i)] \equiv -kC_iC(t) \]

thus,

\[ C_i \approx C_0^r \exp(-kC(t)t) \]

\[ C_0 \quad \text{and} \quad C_0^r \quad \text{are TAT dimer and monomer concentration prior to adding TCEP;} \quad C_i \quad \text{is TAT dimer concentration at any giving reaction time} \quad t; \quad k \quad \text{is the rate constant.} \]

\[ C(t) \quad \text{is the} \quad \text{reaction quotient} \quad \text{and} \quad \gamma \quad \text{is the reaction equilibrium constant.} \]

\[ \gamma = \frac{1}{2 \exp\left(\frac{Nk}{(4D_D)^3} \exp\left(-\frac{\pi^2}{Dt}\right)\right) - 1} \]

\[ Q_r = \frac{2(1 - \gamma)^3 C_0^2}{\gamma C_T^2(1 + \gamma)^2} \]

Here, \( C_T \) is the final TCEP concentration at diffusion equilibrium state. Experimentally, \( \gamma \) and \( Q_r \) can be derived from the concentration of TAT dimer and monomer, which is represented by the number of the dimer (with 40%–70% blockade, 5σ) and monomer (with 19%–38% blockade, 5σ) translocation events [66]. By calculating those two parameters at each reaction time, the transition of \( \gamma \) and \( Q_r \) as shown in Fig. 6b and d represents the temporal evolution of the reaction reduction. The red line in Fig. 6b is the fitting curve with equation (2) (500 μM TCEP as that in Fig. 6a). \( \gamma \) decreased over the course of reaction time, indicating a decrease in oxidized dimer conformation and concurrent increase in reduced monomeric state. Fitting the data revealed a rate constant \( k = 0.989 \pm 0.096 \text{ min}^{-1} \). The trend is also demonstrated in current blockade percentage vs. dwell time distribution of events over the course of the reaction time (Fig. 6c). \( Q_r \) (500 μM TCEP) in Fig. 6d, derived from \( \gamma \) with equation (3), reached a plateau when the reaction approached equilibrium. The plateau \( Q_r \) is equal to standard equilibrium constant \( K_r \), estimated to be 1.4 × 10^6 and corresponding to −23.3 kJ/mol Gibbs free energy (\( \Delta_C^0 - RT \ln K_r \)). All the reduction data from different concentration of TCEP show the same trend to approach equilibrium status (Fig. 6d). When the concentration of TCEP was increased from 100, 250, to 500 μM, the \( \Delta_C^0 \) were −24.5, −23.1, and −23.3 kJ/mol (Table 1), respectively. The \( \Delta_C^0 \) derived from our assay is very close to the \( \Delta_C^0 \) of −27 kJ/mol for protein disulfide bond reduction by TCEP obtained by conventional Raman and computational methods [67].
4. Discussion

With a conductance of 4.27 nS, SPP1 portal channel is one of the larger channels and only the second viral protein channel (apart from phi29 connector) [27] to be incorporated into a lipid membrane. In the presence of TAT peptide, a burst of current blockage events with characteristic current amplitude and dwell time were observed indicating the translocation of the peptides through the SPP1 connector. In this study, we further demonstrated that this new protein nanopore is capable of detecting kinetics of TCEP mediated reduction of disulfide bond and demonstrating peptide conformational changes in real time. From Fig. 6, the frequency of dimer forms (current blockage of ~55%) decreased gradually to monomer forms (current blockage of 30%), upon addition of TCEP, as the reaction time increased. The oligomerization states of peptides was determined in real-time at single molecule level, which is not possible by ensemble methods like NMR, Circular Dichroism, and optical second harmonic generation.

Fingerprints of various peptides translocation through phi29 nanopore was reported [35]. In this study, we further analyzed the kinetic of the peptide translocation. A similar Gibbs free energy ($\Delta G^0$) was obtained when different concentrations of substrates were applied, suggesting that the use of SPP1 nanopore for real-time quantification of peptide oligomeric states is feasible. Although there are many similarities among phi29 and SPP1 nanopore, such as similar shape and conductance, several different properties have been discovered. First, DNA translocation through SPP1 nanopore has not been observed (data not shown), whereas phi29 nanopore allow dsDNA or ssDNA translocation. Second, the orientation preference of these two nanopores in lipid bilayer [38].

Direct evidence of DNA translocation through $\alpha$-hemolysin was demonstrated twenty years ago by quantitative PCR [68]. Although several studies utilizing peptide and protein translocation through nanopores, have been published over the last several years [47–49,54–58], direct method for validating peptide translocations are lacking. A recent publication tried to solve the problem by conjugating a single-stranded DNA to unfolded protein and then amplify the DNA using PCR after translocation [69]. Another study demonstrated the translocation of hyaluronic acid oligosaccharides using high-resolution mass spectrometry [70]. Here, we developed a very simple method to provide evidence of peptide translocation using single molecule fluorescence imaging of samples obtained from the nanopore setup. This method can be easily adapted and employed by other laboratories to validate protein translocation and quantitatively study peptide dynamics through nanoparticles.

The current blockage signature based on the translocation profile can potentially be used to investigate the length, charge, hydrophobicity, secondary structures and ultimately the amino acid sequences of the peptides. The kinetics of protein folding and unfolding as well as entropic and energetic contributions can be further dissected in the future at the single molecule level.

Table 1
Parameters for the quantification of the oligomeric states of TAT peptide in real time.

<table>
<thead>
<tr>
<th>[TAT] (nM)</th>
<th>[TCEP] (mM)</th>
<th>$Q_r \times 10^3$</th>
<th>$K_0 \times 10^3$</th>
<th>$\Delta G^0$ (kJ/mol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>300</td>
<td>100</td>
<td>26.6</td>
<td>26.6</td>
<td>-24.5</td>
</tr>
<tr>
<td>300</td>
<td>250</td>
<td>11.8</td>
<td>11.8</td>
<td>-23.1</td>
</tr>
<tr>
<td>300</td>
<td>500</td>
<td>13.9</td>
<td>13.9</td>
<td>-23.3</td>
</tr>
</tbody>
</table>

Fig. 6. Real-time assessment of the conformational states of TAT peptide. (a) Continuous current trace showing transition of oxidized dimer states to reduce monomer states after addition of reducing agent TCEP. (b) Quantitative analysis showing the fraction ($\gamma$) of dimer and monomer states as a function of reaction time. (c) Current blockage vs. dwell time distribution over the course of reaction time. Applied voltage: 50 mV; Buffer: 1 M KCl, 5 mM HEPES, pH 8. (d) Quantitative analysis showing the reaction quotient $Q_r$ as a function of reaction time.
5. Conclusion

The reengineered membrane-embedded portal channel of bacteriophage SPP1 allows translocation of peptides with higher ordered structure to produce clear and reproducible electronic signatures. The translocation of peptides observed by single channel conduction assays were verified by optical single molecule fluorescence microscopy assays. The oligomer states of peptides were clearly differentiated in real-time at single molecule level. When different concentrations of substrates were applied, a similar ΔG° was obtained, suggesting that the use of SPP1 nanopore for real-time dynamic quantification of peptide folding is feasible.

Acknowledgements

The research was supported by NIH grant R01 EB012135 to P.G. P.G.’s Sylvia G. Frank Endowed Chair position in Pharmacutes and Drug Delivery is funded by the CM Chen Foundation. PG is the consultant of Oxford Nanopore Technologies and Nanobio Delivery Pharmaceutical Co. Ltd, as well as the cofounder of Shenzhen P&Z Bio-medical Co. Ltd and its subsidiary US P&Z Biological Technology LLC. The content is solely the responsibility of the authors and does not necessarily represent the official views of NIH.

Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.biomaterials.2017.01.031.

References


