Three reversible and controllable discrete steps of channel gating of a viral DNA packaging motor

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1. Introduction

Linear double-stranded DNA viruses package their genomic DNA into a preformed protein shell called the procapsid [1,2]. This DNA encapsulation task is an intriguing step of the viral replication cycle, accomplished by nanomotors using ATP as energy [3–7]. The ingenious design of viral DNA packaging motors and the novel mechanism of action has provoked a broad range of interest among scientists in virology, molecular biology, structure biology, nanotechnology, biophysics, biomaterials, nanomedicine, RNA biochemistry, and therapeutics. In bacteriophage phi29, the nanomotor consists of a protein channel, DNA packaging ATPase gp16, and a ring composed of six pRNA (packaging RNA [8]) to gear the motor [8–10] using one ATP to package 2 [3] or 2.5 [11] base pairs of DNA. The protein hub of this motor is a truncated cone shaped connector [12–14], which contains a 3.6 nm wide central channel that allows the 19.3 kb dsDNA genome to enter during maturation and to exit during the infection process (Fig. 1A–C). The defined phi29 DNA packaging motor, constructed 24 years ago [15], is one of the strongest biomotors [16] assembled in vivo. Elucidation of the mechanism of motor action will impact areas of biology, engineering, medicine, and various other nanotechnological fields. The novelty and ingenious design of such machines have inspired the development of biomimetics. In vitro, the biomimetic motor could be integrated into synthetic nanodevices [17–21]. In vivo, the artificial nanomotors could be used to load drugs, deliver DNA/RNA, pump ions, transport cargos, or drive the motion of components in the heart, eye or other sensing organs in the body. Although the protein sequence of each subunit of the connector holds little sequence homology, and each subunit varies in size [14,22–24], the connector of many viruses displays significant morphological similarity [25]. The structure of the phi29 connector has been previously solved by X-ray crystallography [13,14,26], which shows that it is composed of twelve protein subunits which form a ring with a wider end of 13.8 nm outer diameter buried within the viral procapsid and an extruding narrow end of 6.6 nm. The central channel is 6.0 nm at the wider end and 3.6 nm at the narrow end (Fig. 1C).

Many phenomena concerning procapsid expansion during the life cycle related to DNA packaging of bacteriophage have been reported [27–34]. DNA packaging significantly involves the connector. It has also been reported that the connector is a vital component in the regulation of procapsid shape and size [35,36].
Based on logical analysis, it is reasonable to believe that procapsid expansion is linked to the connector conformational change. However, direct evidence of a conformational change of the connector has never been reported.

We have recently inserted the reengineered connector into a lipid bilayer [20,37] (Fig.1E). The translocations of ions and dsDNA through the channel demonstrate the potential to use the connector for DNA sensing and translocation through a membrane. Ion channels may be classified by the nature of their gating [40], such as voltage-gated [41], ligand-gated [42], stretch-gated [43], or other gating [40]. Voltage-gated ion channels are activated by changes in electrical potential difference near the channel, while the ligand-gated ion channels are open or closed in response to the binding of a chemical messenger (i.e., a ligand). There is also recent progress on synthetic ion channels, which are sensitive to the environmental stimuli, such as temperature [44], voltage [45], pH [46,47], or their combination [48]; but they have a gating mechanism different from protein channels. Here we report a real-time direct observation of the gating of the phi29 DNA packaging nanomotor connector protein channel. It is also interesting to find that this viral protein channel gating can be induced by both voltage and ligand binding, which is similar to the other ion channels.

2. Materials and methods

2.1. Materials

The phospholipid, 1,2-diphytanoyl-sn-glycerol-3-phosphocholine (DPhPC) (Avanti Polar Lipids, Alabaster, AL), Nickel-NTA nanogold (1.8 nm; Nanoprobes), n-decane (Fisher), chloroform (TEDIA) were used as instructed by the vendor. All other reagents were from Sigma, if not specified. The construction and purification of phi29 C-terminal tagged connectors have been reported previously [20,49].

2.2. Regineering, expression and purification of phi29 connector

The construction of the plasmid harboring the gene coding for gp10, the overexpression of gp10 and the purification of phi29 connector have been reported recently [50,51]. The deletion of the tunnel loop (N229–N246) of gp10 was performed by two-step PCR. First, primer pair F1–R1 and primer pair F2–R2 were used to amplify the DNA sequence coding for gp10 (1–228) region and gp10 (247–309) respectively. In the second round of PCR, F1 and R2 were used as primer pairs to link and amplify the PCR product in the first round. The second PCR product was digested with Ndel/Xhol and ligated into Ndel/Xhol sites of the vector pET-21 a(+) (Novagen).

The deletion of the N-terminal (1–14) of gp10 was performed by PCR. The sequence coding for gp10 (15–309) region was amplified by a pair of primers. The forward primer contained Ndel restriction site; the reverse primer contained XhoI restriction site and a 6-histidine affinity tag. The PCR product was digested with Ndel/Xhol and ligated into Ndel/Xhol sites of the vector pET-21 a(+) (Novagen).

The connector mutants constructed were expressed, and then were purified with Nickel affinity chromatography [52]. Cells were resuspended with His Binding Buffer (15% glycerol, 0.5 w NaCl, 5 mM Imidazole, 10 mM ATP, 50 mM Na2HPO4–NaH2PO4, pH 8.0), and the cleared lysate was loaded onto a His•15°N resin column (Novagen) and washed with His Washing Buffer (50 mM Na2HPO4–NaH2PO4, 15% glycerol, 0.5 w NaCl, 50 mM Imidazole, 10 mM ATP, pH 8.0). The His-tagged connector was eluted by His Elution Buffer (50 mM Na2HPO4–NaH2PO4, 15% glycerol, 0.5 w NaCl, 0.5 w Imidazole, 50 mM ATP, pH 8.0).

Fig. 1. Illustration of the phi29 connector channel structure. (A) Side view without showing the C- and N-terminal fragment [14]; (B) Side view and (C) bottom view with complete protein sequence [51]; (D) Model of phi29 DNA packaging motor within the procapsid; (E) Illustration of dsDNA translocation through a connector channel reconstituted in an artificial lipid bilayer for the measurement of conductance.
2.3. Electrophysiological measurements

A bilayer was formed on a thin Teflon film partition (aperture 200 μm in diameter) which separates a bilayer lipid membrane (BLM) chamber into a cis- and trans-chamber (compartment). The cis-chamber refers to the grounded compartment to which the connector-reconstituted liposome is added. Connector channel insertion into the bilayer has been described previously using vesicle fusion [20]. Briefly, connector-reconstituted liposomes were prepared using a deactivation–rehydration method and further extruded to form unilamellar liposomes. The reconstituted liposomes were further diluted by 10-fold using the conducting buffer before applying to BLM chambers. The final concentration of added protein was 5–50 μg/mL.

For electrophysiological measurements, both compartments in the BLM chamber were filled with a conducting buffer (1 M NaCl, 5 mM Tris, pH 7.8, if not specified). A pair of Ag/AgCl electrodes connected directly to both compartments was used to measure the current traces across the bilayer lipid membrane. The current trace was recorded using an Axopatch 200B patch clamp amplifier coupled with the Axon DigiData 1322A analog–digital converter (Axon Instruments) or the BLM workstation (Warner Instruments). All voltages reported were those of the trans-compartment. Data was low pass-band filtered at a frequency of 5 kHz or 1 kHz and acquired at a sampling frequency of 10 kHz. The Pclamp 9.1 software (Axon Instruments) was used to collect the data, and the software Origin Pro 8.0 was used for data analysis.

2.4. Probing connector conformational change using antibody

C-His connector and rabbit polyclonal antibodies (where) specific for His tag were added to the bottom chamber and measured under a voltage of −75 mV at the bottom chamber (a buffer containing 150 mM NaCl at pH 7.8).

2.5. Probing connector conformational change using Nickel-NTA nanogold

Conductance measurements were carried out under asymmetric ionic conditions [35]. The chamber without nanogold contained 1 mM NaCl, 20 mM Tris (pH 7.6). For the chamber that contained nanogold, the NaCl concentration was reduced to 150 mM to avoid the high salt effect that might interfere with the binding of nanogold to the His-tagged connector. DNA was premixed in both chambers with the conducting buffer. After forming the bilayer, a final concentration of 150 pM Ni-NTA nanogold (1.8 nm; Nanoprobes) was added to the chamber and incubated for 5–10 min to ensure an even particle distribution before the connector-reconstituted liposomes were added to the chamber.

3. Results

3.1. Discrete one-third step-wise conformational change during high-voltage gating

The connector protein embedded in the lipid bilayer was stable and displayed a uniform conductance under a wide range of experimental conditions including different salt concentrations and varying pH [37], and has a uniform conductance in each of the conditions. At a constant holding potential (−75 mV), the channel conductance is uniform (Fig. 2A). However, when a higher potential (−150 mV or −200 mV) was applied, a 3-step channel size reduction was often observed immediately thereafter (Fig. 2C and D), which is very similar to the gating behaviors of membrane proteins. In the given experiment conduction (1 mM NaCl with 5 mM Na2HPO4 buffer, pH 12), the step size of a single C-His connector is around 190 pA at +75 mV (Fig. 2A), and −190 pA at −75 mV (Fig. 2B). The channel was stable over minutes, and even hours without any blockade or vibration. When the voltage was switched to −150 mV, the current through the channel doubled in accordance with the voltage increase from −190 pA to −380 pA. Immediately after that, the same channel closed after a three-step reduction in 5 s (Fig. 2C). Each reduction step size was 120 pA, accounting for 31% of the entire channel. This 3-step channel gating can be triggered by the high voltage despite the buffer conditions. It is interesting to note that after the 3-step reduction, there was still a residual current of 20 pA opening, equal to 5.3% of one channel. Similar results were observed at higher potentials of 200 mV (Fig. 2D).

3.2. Reopening of the gated channel by voltage dropping

It was also observed that the gating of the C-His connector protein channel was reversible. When a transmembrane voltage of 0 pA was applied for 10–30 s, the gated channel was observed to reopen (Fig. 3). After the channel shutting down was triggered by higher voltage (−100 mV in Fig. 3A, and −150 mV in Fig. 3B), a lower voltage was applied to allow the connector protein to recover. It was found that the conductance of the channels restored to their original state within 10 s.

3.3. Discrete one-third step-wise conformational change of connector induced by protein binding to the C-terminal

A C-terminal His-tagged connector and rabbit polyclonal antibody (Ab) specific against His tag were used to measure the conductivity under a voltage of −75 mV with the negative pole placed at the bottom chamber (the trans-side). Since the channel exercises a one-way traffic mechanism, orientation of the connector was determined by their capability to translocate dsDNAs which was placed in only one, either top or bottom of the two chambers via the polarity of applied electrical current (see Ref. [38] for details). Addition of the Ab to the bottom chamber (Fig. 4B) resulted in six discrete 31% current reduction steps for two connectors, indicating binding events induced by interaction of one Ab molecule with one of the two connector channels. As a negative control, two connectors were inserted into BLM with the His-tagged C-terminal exposed to the bottom chamber. DNA was premixed with conducting buffer before the insertion of the connector channels. When the Ab was added to the top chamber, no binding events were observed (Fig. 4A). The results indicate a three-step conformational change of connector after the anti-His antibody bound to the His tag at the C-terminal.

3.4. Discrete one-third step-wise conformational change of the connector channel induced by Ni-NTA nanogold binding to the C-terminal

A 1.8-nm Ni-NTA nanogold particle was used to bind to the His-tag at the C-terminal of the connector. The binding was promoted via a Ni-NTA/His-tag interaction. The nanogold was premixed with the conducting buffer (asymmetric conditions; see Methods) at only one side of the chamber (the trans-side). In the presence of a single or multiple connector channels, the nanogold only bound to a connector with its His-tagged C-terminal end oriented toward the trans-chamber to the bottom, and did not bind to connectors with the N-terminal facing the trans-side. Clear discrete step-wise closing of the channels with a corresponding decrease in conductance was observed when a single connector with the appropriate orientation for nanogold binding was present (Fig. 4C). Binding of each nanogold particle resulted in ~31% reduction in channel current. Such observations enabled the direct counting of the number of nanogold particles bound to each connector. The step size of each current reduction, resulting from the binding of one nanogold particle, was nearly identical, even in the case of membranes with multiple channels (data not shown).

3.5. Gating behavior of mutant connector with the internal 18-amino acid flexible loop or the N-terminal 14-amino acid removal

Phi29 connector protein contains three flexible fragments, the N-terminal fragment amino acid (aa) 1–14, the internal fragment aa 229–246, and the C-terminal fragment aa 287–309 (Fig. 5A). The structures of these three fragments were not included in the crystal structure due to their flexible nature. The flexibility of
the connector channel is expected to play a critical role in motor motion and DNA translocation, instead of channel structure construction. To investigate whether these three loops were involved in channel gating, three connector mutations, gp10-C-His/N14, gp10-C-His/D229e246, and gp10-C-Strep/C25 were constructed, with the deletion of the loop aa 1e14, 229e246, and 285e309, respectively (Fig. 5B). The step-wise gating was observed under both the higher holding potentials and a ramping voltage (from −150 mV to +150 mV, or from −200 mV to +200 mV). Fig. 6A and B shows the events happened on a gp10-C-His/NΔ1−14 connector channel, which has a similar channel conductance with the C-His connector. Gating occurred immediately when the ramping began at −150 mV, as shown by the step-wise reductions in the slope of the current trace (Fig. 6B). The slope restored to its original state under −50 mV, indicating that the channel conductance reopened completely. Complete gating happens at a positive potential higher than +110 mV. Such a ramping voltage was applied periodically to test the possibility of the channel gating and reopening (Fig. 6C and D). It was found that both C-His (Fig. 6D) and gp10-C-His/NΔ1−14 can be closed completely after several ramping circles, while the gp10-C-His/D229—246 were much less likely to be gated completely (Fig. 6E). The current trace through gp10-C-His/Δ229—246 was still linear (Fig. 6F) under 120 ramping cycles after 3 h, indicating the amazing stability of the channel conductance and significant differences between the channels with/without intact internal loops.

3.6. The effect of the C-terminal region in regulating the conformational change

The aforementioned results suggest that phi29 connector implemented a conformational change with three discrete steps. It was proposed that these three steps of conformational change is regulated by the interaction of DNA with the connector, and the binding or contact of components to the C-terminal would result in the similar discrete steps of conformational change. This proposal was supported by a conductance assay using mutant connector conjugated with a His-tag to its C-terminal and incubated with anti-His tag antibody or nanogold coated with Ni-NTA (Fig. 4). It was hypothesized that the translocation of DNA into the procapsid or the internal pressure of the fully packaged DNA within the procapsid might lead to the contact of DNA with the C-terminal
flexible domain, inducing a conformational change with two subsequence functions: 1) to prevent the dsDNA from exit; and 2) to prepare a new channel configuration to facilitate the injection of DNA during the host cell infection process. To further confirm this hypothesis, a connector mutant with the removal of a 25-amino acid segment at each C-terminal of the 12 subunits was constructed. Similarly, a strep-tag was conjugated to the C-terminal fused to amino acid #287. Conductance assay of the mutant connector by electrical ramping revealed that three discrete steps of conformational changes were also observed (Fig. 6G). However, the discrete steps of 32% conductance change were not observed in the presence of streptavidin.

4. Discussion

Viral DNA packaging has been investigated extensively in many viral systems, but the actual mechanism remains elusive. During the last several decades, many models have been proposed to interpret the mechanism of motor action. These include 1) Gyrase-driven supercoiled and relaxation [6,54,55]; 2) Force of osmotic pressure [56]; 3) Ratchet mechanism [57]; 4) Brownian motion [58]; 5) Five-fold/six-fold mismatch connector rotating thread [59]; 6) Supercoiled DNA wrapping [60]; 7) Sequential action of motor components [11,61]; 8) Electro-dipole within central channel [14]; and, 9) Connector contraction hypothesis [62]. All these models are very intriguing, however, none have been fully supported by conclusive experimental data. In other cases, various models were validated in one viral system but disproved in another. The five-fold/six-fold mismatch connector rotating thread model [59] has been popular for more than 30 years, since this model could bring about a new mechanical motor prototype. Even within the last several years, numerous laboratories, including our own, have persevered to search, interpret, match, link, and even design a five-fold ring to adapt findings to this fascinating and extraordinary model [11,63–65]. Unfortunately, recent studies in many viral DNA packaging motors reveal that the stoichiometry of motor components is not an odd number but actually an even number [19,66–68]. In 1998, Guo and co-workers first proposed and revealed that the mechanism of DNA packaging is simply via a mechanism similar to the hexameric AAA + ATPase that translocate dsDNA during DNA replication and repair [9]. The finding of even-number structures is consistent with the mechanism of many other well-studied DNA-tracking motors [69–72] and the AAA + ATPase family. Most recent publications [52,65,73–75] all support our long-term findings [19,68,76] that pRNA dimer is indeed the building block for hexameric structure [76]. X-ray crystallography also confirmed that pRNA forms a dimer in solution and the dimer forms a tetramer in the absence of the procapsid [65], supporting the theory that the dimer is indeed the building block of the hexamer and the sequential action in hexamer assembly is 2 → 4 → 6. Furthermore, explicit results conclude that the connector does not rotate during the DNA translocation process [77,78]. Recently, Guo and co-workers proposed a “Pushing through a one-way valve model” [2,38]. In this model, the connector remains as non-rotating valve to allow dsDNA travel one way toward the procapsid; DNA translocation is induced by a DNA packaging enzyme or terminase, which pushes a certain length of DNA into the procapsid segment by segment. This model strongly agrees with many recent findings. Most recently, it

Fig. 3. Controllable and switchable closing and reopening of the connector channel. (A) The C-His channel with three steps of gating at high voltage (–100 mV or –150 mV) was reopened after the voltage was reduced to 0 mV. (B) The channel closed at –150 mV was reopened to its original size at a lower voltage of –75 mV.
was reported that under an external electrical force, the channel of the phi29 nanomotor favored DNA entry but blocked DNA departure [38]. The one-way traffic property implies that during the packaging of dsDNA via the active motor, dsDNA travels in only one direction from the narrow external end (N-terminal) toward the wider internal end (C-terminal) of the channel. Essentially, the channel functions as a one-way valve. Thus, these results suggest that dsDNA packaging occurs through the combination of two separate tasks. The first task is the active pushing mechanism [2,38,79,80] provided by the ATPase, gp16, bound to pRNA (the fulcrum) of phi29, actively pushing dsDNA coupled with hydrolysis of ATP. In other dsDNA phage systems, two proteins (terminases) act in combination where the large subunit grips onto the small subunit for use as a fulcrum as with pRNA of phi29. The second task is the control of DNA migration direction with a one-way valve mechanism of the channel [38]. The mechanism providing force through the one-way valve connector also agrees with the finding in T4 phage system by Black and co-workers [79,80]. They found that dsDNA was crunched and the compressed if the DNA entry was blocked at the front end [79,80]. Although the authors interpreted that the force for the crunching and compression is due to torsional force from the coiled DNA at the external end, it is not contradictory to support our finding that the ATPase gp16 functions similar to the DNA-tracking AAA+ family that twisting or rotating the dsDNA [81]. Our new model is also supported by the finding that in the T4 DNA packaging system, both ends of DNA are able to stay outside of the procapsid [79,80]. If the motor would have implemented a pulling instead of pushing function, it would be difficult to interpret how the initiation of DNA translocation begins when both ends are located outside the channel [79,80].

The new model raises the question of how dsDNA is ejected during infection if the channel only allows the dsDNA to travel in one direction. It has been proposed that the motor connector adopts a conformational change after DNA packaging is complete [29]. Such a conformational change renders the channel capable to eject the dsDNA from the viral procapsid. It has been reported that the conformation of the phi29 connector is substantially changed after DNA packaging [29,82,83]. Significant rearrangement of the connector after DNA packaging, a similar feature reported in other phage systems [84–86], may also change the channel configuration to favor the reverse exit of DNA during infection. In bacteriophage p22, a conformational switch of the portal protein primes genome injection [33]. From these results, we confirmed that the connector channel indeed exercises conformational changes. Such conformational changes were induced by molecule binding to the C-terminal wider end that was located within the capsid, and a high electrical voltage shift. The conformational change exhibited three discrete steps, with each step reducing the channel size by 31%.

Our results support the proposal that these three steps of conformational change are regulated by the interaction of DNA with the connector. We demonstrated that the binding or contact of components to the C-terminal would result in the similar discrete steps of conformational change. Conductance assay using mutant connector conjugated with a His-tag to its C-terminal and incubated with anti-His tag antibody or nanogold coated with Ni-NTA revealed three discrete steps of channel change with blockade of about 32% (Fig. 4). It is reasonable to believe that the translocation of DNA into the procapsid or the internal pressure of the fully packaged DNA within the procapsid might lead to the contact of DNA with the C-terminal flexible domain, inducing a conformational change with two subsequent functions: 1) to prevent the dsDNA from exit; and 2) to prepare a new channel configuration to facilitate the injection of DNA during the host cell infection process. The proposed second step of action has also been evidence in the
P22 system [33]. To further support this, a connector mutant with the removal of a 25-amino acid segment at each C-terminal of the 12 subunits were constructed with a strep-tag conjugated to the C-terminal. With this mutant, the discrete steps of 32% conductance change were not observed in the presence of streptavidin. However, conductance assay of the mutant connector by electrical ramping revealed that three discrete steps of conformational changes were also observed (Fig. 6G). The finding suggests that conformational change was a result of the transition of the entire connector structure, and the C-terminal only served as a trigger. This conclusion agrees with the finding in the connector structure of bacteriophage P22 [23]. A unique topology of the C-terminal domain was reported to be a ~200-Å-long α-helical barrel that inserts deeply into the virion and is highly conserved in the Podoviridae family. They proposed that the barrel domain would facilitate genome spooling into the interior of the procapsid during DNA packaging, and in analogy to a ri...
leads to the opening or closing of the channel, which help to control the packaging or release of the viral genome.

The controllable opening and closing of the connector protein has been achieved with different polarity, which also resembles the voltage-gated ion channels. However, they have different functions in the biological environment. The ion channels may have to open and close multiple times in their life cycle, whereas the phi29 DNA connector protein might require fewer. The significantly different gating behavior of the internal flexible loop-connected connector suggests that the flexible loops may play a key role in the voltage gating. Results described above show that after the removal of these loops, both the occurrence and extent of gating reduced tremendously. These loops can induce conformational changes to adjust the channel size in response to an applied potential. Further studies are necessary to investigate if other domains of the connector protein also contribute to its gating and conformational changes.

5. Conclusions

The real-time direct observation of the viral connector protein opens a new door to study the mechanism of DNA packaging motors. This “modularity” allows us to use these simple and inexpensive model systems to study the regulation of DNA trafficking, the role of the viral portal protein in infection, and pharmaceutical control of the infection. The gating is also reversible and controllable by voltage or ligand binding, making the modified connector protein an ideal nano-valve for constructing nanomachines with potential applications in nanotechnology and nanomedicine, such as for drug loading and controlled release, as well as for high-throughput single pore DNA sequencing.

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