Identification of Arginine Finger as the Starter of the Biomimetic Motor in Driving Double-Stranded DNA

Chenxi Liang and Peixuan Guo*

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ABSTRACT: Nanomotors in nanotechnology may be as important as cars in daily life. Biomotors are nanoscale machines ubiquitous in living systems to carry out ATP-driven activities such as walking, breathing, blinking, mitosis, replication, transcription, and trafficking. The sequential action in an asymmetrical hexamer by a revolving mechanism has been confirmed in dsDNA packaging motors of phi29, herpesviruses, bacterial dsDNA translocase FtsK, and Streptomyces TraB for conjugative dsDNA transfer. These elaborate, delicate, and exquisite ring structures have inspired scientists to design biomimetics in nanotechnology. Many multisubunit ATPase rings generate force via sequential action of multiple modules, such as the Walker A, Walker B, P-loop, arginine finger, sensors, and lid. The chemical to mechanical energy conversion usually takes place in sequential order. It is commonly believed that ATP binding triggers such conversion, but how the multimodule motor starts the sequential process has not been explicitly investigated. Identification of the starter is of great significance for biomimetic motor fabrication. Here, we report that the arginine finger is the starter of the motor. Only one amino acid residue change in the arginine finger led to the impediment and elimination of all following steps. Without the arginine finger, the motor failed to assemble, bind ATP, recruit DNA, or hydrolyze ATP and was eventually unable to package DNA. However, the loss of ATPase activity due to an inactive arginine finger can be rescued by an arginine finger from the adjacent subunit of Walker A mutant through trans-complementation. Taken together, we demonstrate that the formation of dimers triggered by the arginine finger initiates the motor action rather than the general belief of initiation by ATP binding.

KEYWORDS: DNA-packaging motor, viral assembly, phi29 DNA packaging, asymmetrical hexameric ATPase, revolving motor, sequential action, inchworm

INTRODUCTION

Biological motors, or biomotors in short, are nanoscale machines ubiquitous in many biological processes in both prokaryotes and eukaryotes, such as cell mitosis, DNA replication, RNA transcription, macromolecule trafficking, and viral genome packaging. Understanding the mechanism of biomotor function is essential for the studies on biological systems and for the development of new targeting drugs against bacterial or viral infections. One important component of motors is the ATPase, which belongs to one class of enzymes that catalyze the decomposition of adenosine triphosphate (ATP) into adenosine diphosphate (ADP) and a free phosphate ion. They serve as the main energy source for the mechanical work of biomotors, just like the engine for a car. For ring-shaped ATPases found in biomotors, conversion from chemical to mechanical energy usually takes place in a sequential manner among the subunits, coupled with conformational changes of the oligomer. While it is commonly believed that ATP binding triggers such conversion, how ATPases start the sequential process has not been explicitly investigated. The arginine finger (R-finger), Walker A (WA), and Walker B (WB) residues are three commonly conserved motifs in ATPases responsible for ATP binding and hydrolysis. For ring-shaped ATPases such as the AAA+ family

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family, R-fingers are found at the intersubunit interface regulating ATPase activities.19,20 For instance, various mutants were assayed for ATPase activity and the trans-acting Arg139 was identified to be the R-finger of TerL ATPase.21

The dsDNA packaging motor of bacteriophage phi29 has long been used as a model to study the viral DNA packaging pathway for a long time. In this motor, dsDNA is pushed through a one-way valve; an asymmetrical hexameric ring and a dodecameric connector channel follow a revolving mechanism (Figure 1).22−25 This revolving mechanism is common among different species of dsDNA translocases. The use of an arginine finger to control the sequential action26,27 through the asymmetrical hexamer of the revolving motor has been confirmed in different kinds of dsDNA translocating or packaging motors, such as the DNA packaging motor of phi29,28 herpesviruses,29,30 the DNA translocase FtsK of Gram-negative bacteria,31 and the TraB in Streptomyces for conjugative plasmid DNA transfer.32−34

The gene product 16 (gp16) of bacteriophage phi29 has long been employed to study the sequential process in ring-shaped ATPases (Figure 1).26,35,36 Based on the sequence and structural analysis, the R-finger of gp16 was determined to be Arg146, after the α4 helix. WA and WB residues are also located near the R-finger with a sequence of GXXGXGKS/T8 and hhhhDE, respectively. To translocate the viral genome double-stranded DNA (dsDNA) into the preformed capsid, the multimeric gp16 ATPases have to undergo concerted steps: ATP binding, substrate binding, ATP hydrolysis, and finally conformational changes leading to movement of the substrate. Even without knowing the exact sequence of these steps, we can reasonably assume that the halt of the very initial step will impair all the subsequent functional steps.

Through biochemical assays, we have shown that the R-finger of gp16 is indispensable for subunit dimerization, ATP binding, DNA binding, and viral assembly. Together with molecular modeling and simulations, we illustrated how the structural motifs of gp16 mediate the subunit–subunit interaction in a trans-acting manner. It has also been found that gp16 without an active R-finger can rescue ATP hydrolysis activity of an inactive Walker A mutant by trans-complementation. Therefore, we suggest that the R-finger of gp16 facilitates the dimerization of two adjacent subunits and initiates the sequential energy conversion process, finally leading to DNA translocation. We also anticipate such a structural initiation mechanism to be conserved among the ring-shaped ATPase family. Identification of this motor initiation mechanism will also benefit the assembly of artificial biomotors in vitro and our understanding of the complex mechanism behind biopolymer translocation.

RESULTS AND DISCUSSION

Indispensability of Arginine Finger for gp16 Dimerization. The formation of gp16 dimers in the absence of ATP has been reported extensively as tracking back to 1986.37 When the phi29 gp16 gene was engineered and expressed in E. coli, gp16 was found to form oligomers. The wild-type gp16 conjugated with enhanced Green Fluorescent Protein (eGFP) can form dimers, trimers, and other oligomers through Native-
At least six different bands indicate that different states of oligomers exist in the solution. However, the oligomer formation was impaired by the reengineering of R146 of the gp16, which is shown to be the R-finger of gp16.20,39 The ultracentrifugation of the engineered R-finger mutant showed only one peak representing monomer gp16 mutants in the 15%−35% glycerol gradient compared to other reengineered gp16 with weak representation of monomers and dimers. Interestingly, the reengineered R-finger shows intersubunit interaction between wild-type gp16 and other mutants, i.e., the reengineered Walker A mutant and Walker B mutant. These results demonstrate that the R-finger is indispensable for dimer formation of gp16. Given that dimer can be formed without ATP, it is possible for dimerization to precede ATP binding and hydrolysis.

However, the renatured gp16 monomer lost the biological activity within 40 min in solution due to protein/protein interaction.37 Although dimer formation is the key to gp16 function, adding gp16 monomer is required for gp16 to work in the motor since each motor contains only one dimer and four monomers at any transient movement.41 In other words, gp16 dimer is the initiation step via the contact of the arginine finger with the P loop of Walker A to form an ATP binding pocket (see below). The dimer is subsequently stabilized by the binding of ATP (or γ-S-ATP). Thus, the purified gp16 dimer alone could not have DNA packaging activity.20

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**Figure 2. Indispensability of arginine finger for ATP binding.** Phosphate imaging of 32P-ATP and different gp16-eGFP mutants. The arginine mutant R146A loses the ATP binding ability completely. The WT, Walker B motif (118/9DE), and arginine finger mutant (R146A) were incubated with 40bp DNA, pRNA, and 10 μM 32P-ATP with (+) and w/o (−) γ-S-ATP. "0" is the negative control without gp16. (A) Phosphate imaging overnight showing signal of 32P-ATP. (B) eGFP channel showing the signal of gp16-eGFP.

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**Figure 3. Indispensability of arginine finger for DNA binding.** The interaction of DNA labeled with Alexa647 with either the WT gp16 or the Walker A−arginine finger double mutant gp16 (WA-RF gp16) were tested by electrophoretic mobility shift assay (EMSA). When increasing the amount of WT gp16, more DNA bound to the WT gp16 protein (lane 2−7), while the Walker A−arginine finger mutant did not show binding to DNA (lanes 8−13). (A) eGFP channel showing the signal of gp16-eGFP. (B) Alexa647 channel showing the signal of red fluorescent-DNA. (C) Overlap image (merged) of A and B.
acetone and PEG were added to the gp16 solution to interfere with the dimer formation, the gp16 showed a significant increase of activity and displayed a much longer shelf life.\(^\text{42}\)

**Indispensability of Arginine Finger for ATP Binding.**
A re-engineered R-finger mutant R146A was investigated. In this mutant the 146th arginine was replaced by alanine.\(^\text{20,39}\)

The ATP binding affinity of this R-finger mutant, R146A, was evaluated through native PAGE together with the re-engineered Walker B mutant 118/9 DE, in which the 118th aspartic acid and 119th glutamic acid were reversed. Radioactively labeled \(^{32}\)P-ATP was used to determine if the mutant gp16 binds ATP (Figure 2). Surprisingly, the R146A showed no affinity for \(^{32}\)P-ATP in either the presence or absence of \(\gamma\)-S-ATP (Figure 2A), indicating that the re-engineered R146A mutation compromised its ATP binding ability.

**Indispensability of Arginine Finger for DNA Binding.**
The ATPase gp16 undergoes a conformational change after binding to ATP, leading to a higher affinity for DNA molecules. In a full function cycle of the motor, the bound ATP is subsequently hydrolyzed, a reaction that is required for the translocation of dsDNA. The nonhydrolyzed \(\gamma\)-S-ATP can be used to block the ATP binding/hydrolysis cycle and “freeze” the gp16 in a conformation that favors DNA binding. A mutation in R-finger also impaired the DNA binding ability of gp16.

EMSA was used to show that R146A lost its affinity for DNA completely (Figure 3). After the DNA was incubated with \(\gamma\)-S-ATP and wild-type gp16-eGFP, an upshift of the fluorescent DNA band was observed. Adding more wild-type gp16-eGFP resulted in a complete upshift of the fluorescent DNA band, while adding mutant without arginine finger did not lead to an upshift of the fluorescent DNA band, regardless of the amount added. It was rationalized that the lack of DNA binding in WAF mutant gp16 is a consequence of its inability to bind ATP due to arginine mutation.

**Indispensability of Arginine Finger for ATP Hydrolysis.**
It was shown that R146A impaired the ATP hydrolysis function of gp16 compared to wildtype (Figure 4). From a structural perspective, the R146A mutation could not alter either the WB motif, which is essential for ATP hydrolysis, or the WA motif, which is essential for ATP binding. Therefore, it is concluded that besides the Walker motifs, the R-finger is also essential for ATP binding and hydrolysis.

**Indispensability of Arginine Finger for DNA Packaging.**
The DNA packaging ability of the R146A was tested (Figure 5). The band of the packaged DNA can be seen using the wild-type gp16. However, the band of packaged DNA is almost invisible when the R146A mutant was added to the solution equivalent to 75% of the wild type gp16. Even 25% of R146A mutant had a negative effect on the packaging process, compared to the addition of the buffer (Figure 5). Apparently, R146A does not have the assembly ability since it loses its function in ATP binding, ATP hydrolysis, and DNA binding. However, this result indicates that R146A can still interact with the wild-type gp16, thus blocking the function of the wild-type gp16. Given that the R146A mutant also affects the dimerization of gp16, it is reasonable to assume that the wild-type gp16 can use its own R-finger to bind R146A, but the R146A in the dimer cannot bind to the next gp16, so that the packaging process is blocked.

**Arginine Finger Mutant to Rescue the ATP Hydrolysis Activity of Walker A Mutant by Trans-Complementation.**
On the basis of previous results, R146A loses all ATPase functions. However, the R-finger was found not directly involved in ATP binding and hydrolysis but instead to facilitate the dimerization of gp16. Hence, based on the structure of gp16 (Figure 6), the dimerization of gp16 should also be essential for ATPase function, and the lost function could be recovered through mixing R146A with a gp16 mutant with the intact R-finger. This hypothesis was tested by assessing the ATPase activity of a combination of different gp16 mutants. For example, the Walker A mutant G27D, in which the 27th glycine was replaced by aspartic acid, and the arginine finger mutant R146A (Figure 7A) were used for the assay. When R146A was mixed with G27D in different ratios, an ATP activity curve was observed representing a complementary effect (Figure 7AB). When R146A was mixed with G27D in a 1:1 ratio, the highest ATPase activity was observed (Figure 7AB). This indicates a complementary effect between R146A and G27D. We further showed that this complementary effect was not affected by the introduction of DNA or \(\gamma\)-S-ATP (Figure 7B). On the other hand, mixing E119A, in which the Walker B motif was mutated, with G27D in different ratios did not display the complementation effect, since in both Walker A and Walker B mutants the ATPase function was inactivated. Then a transaction will not rescue the ATPase activity. The result suggests that the arginine finger of gp16 can interact with another gp16 subunit, functioning as an ATPase dimer in a transacting manner (Figure 7A).

To further prove the complementary idea, a two-point mutation of the Walker A motif was introduced to generate the K30/S31A mutant (Figure 7C). This mutant replaced the 30th lysine and 31st serine with alanine. As expected, the K30/S31A also has a complementary effect with R146A. In addition, when mixing these two mutants together, the ATPase activity of gp16 mutants was significantly recovered to about 40% of the wild-type gp16. The results indicate that the gp16 functions as an ATPase in a dimeric form, and the ATP hydrolysis function was carried out by two gp16 subunits to work together.
CONCLUSION

We demonstrated that the arginine finger of gp16 is necessary for its ATPase function and DNA binding. Single-point mutation of the arginine finger in gp16 lost the ability of subunit dimerization, ATP binding, DNA binding, ATP hydrolysis, and DNA packaging. However, the loss of ATPase function can be rescued by the complementary effect of Walker A mutant G27D and K30/S31A. Since the arginine finger directly affects subunit dimerization, the fact that arginine finger mutation impairs the ATPase function and DNA binding indicates that the dimerization of gp16 is the prerequisite of all subsequent steps for a functioning gp16 ATPase ring. If, under natural conditions, the ATP binding and hydrolysis steps happen before the dimerization, these processes would not need an intact arginine finger to proceed. As a result, we conclude that the dimerization of gp16 through the R-finger is the initial step of the sequential process leading to DNA packaging in the bacteriophage phi29 motor.

EXPERIMENTAL METHODS

Engineering, Fabrication, And Purification of Motor Components. The engineering and purification of the eGFP-gp16 protein have been reported previously. The design of gp16 mutants is shown in Figure 6. The eGFP-gp16 arginine finger mutant (R146A), eGFP-gp16 Walker A mutant (G27D, K30/S31A), gp16 Walker B mutant (118/9 ED, E119A), and eGFP-gp16 mutant R146A were fabricated by engineering the gp16 gene (Keyclone Technologies). Briefly, the protein was overexpressed in E. coli BL21(DE3) with induction of 0.4 mM IPTG. The bacterial cells were harvested and resuspended in His-binding buffer (20 mM Tris–HCl, pH 7.9, 500 mM NaCl, 15% glycerol, 0.5 mM TCEP and 0.1% TEV protease). The recombinant eGFP-gp16 was purified by Ni-NTA chromatography after the above buffer exchange procedure. Background protein was removed by PAGE and further obtained by dialysis against the His-binding buffer. The resulting protein was concentrated to a final concentration of 1 mg/mL and stored at −80°C. The purified protein was used for crystallization. The crystals were soaked in 20% glycerol before cryoprotection and flash-frozen in liquid nitrogen and used for X-ray crystallography.

Figure 5. Indispensability of arginine finger for DNA packaging. The arginine finger mutant (R146A) was biologically inactive in DNA packaging as demonstrated by its inhibition to DNA packaging activity of WT gp16. An in vitro DNA packaging assay was carried out using a series of titrations of WT gp16 with buffer, Walker B mutant E119A, and arginine finger mutant R146A. (A) EtBr staining of the agarose gel showing the packaged DNA (black arrow). R146A gp16 by itself did not have DNA packaging activity. (B) Bar chart of the grayscale measurement of the packaged DNA. Data were normalized to the activity of only WT gp16.

Figure 6. Crystal structure of the N terminus of gp16 (top figure, PDB file: 5HD9) and the location of the mutations (bottom table). The red amino acids in the table indicate the mutation sites.

Figure 7. Fluorometry to study the complementary effect of ATP hydrolysis ability among different gp16 mutants. The fluorescent signals were measured every 30 min after mixing, and the data were normalized to the activity of WT gp16 after 120 min. (A) Test of the complementary effect of gp16 mutants: Walker A mutant (G27D) complement with Walker B mutant (E119A), and Walker A mutant (G27D) complement with Arginine mutant R146A. (B) Complementary effect of G27D and R146A w/ or w/o DNA and γ-S-ATP (n = 2). (C) Complementary effect of another Walker A mutant K30/S31A and R146A (n = 2).
The cells were then lysed by passing through a French press, and the lysate was clarified by centrifugation. 0.1% PEI was added to the clarified lysate to remove nucleotides and other proteins. Homogeneous eGFP-gp16 was purified by one-step Ni-resin chromatography.

**Native PAGE and Phosphate Imaging.** Wild-type and mutant gp16-eGFP were mixed with 0.05 μM DNA, 0.5 ng/μL rRNA, and 10 μM γ-S-ATP. A final concentration of 1 mM γ-S-ATP was introduced if needed. After incubation at room temperature for 30 min, all of the samples were loaded to a 4—15% acrylamide gel (Tris, glycine) and run for 90 min under 90 V. After the gel was exposed for eGFP signal, the gel was dried and then exposed for phosphate imaging overnight.

**Electrophoretic Mobility Shift Assay (EMSA).** A fluorescently tagged protein that facilitates detection and purification was shown to possess similar assembly and packaging activity as compared to wild type. 27,43 The EMSA method has been described previously. 36,44 Wild-type and mutant gp16 proteins were mixed with a 50bp DNA conjugated with Alexa647 in the presence of γ-S-ATP with a final concentration of 0.5 μM. Samples were incubated at room temperature for 20 min and then loaded onto a 1% agarose gel (44.5 mM Tris, 44.5 mM boric acid) and electrophoresed at 4 °C for around 1 h at 8 V/cm. The fluorescent signals of gp16 and DNA-Alexa647 were analyzed using a Typhoon FLA 7000 laser scanner.

**Measurement of ATPase Activity by Fluorometry.** A malachite Green Phosphate Assay Kit (Cayman Chemical) was used to examine the ATPase activity of gp16 in the presence or absence of the 33bp dsDNA and γ-S-ATP that was described previously. Purified gp16 was mixed with a final concentration of 1 ng/μL rRNA, MDCC-PBP containing 0.2 U/mL of purine nucleoside phosphorylase (PNPase), 0.04 mM 7-methyl guanosine, and 6% glycerol. The DNA and γ-S-ATP were added based on the experimental design with a final concentration of 0.1 and 0.2 μM, separately. Then a final concentration of 1 μM ATP was added. After addition of ATP to start the ATP hydrolysis, ATPase gp16 generated P, that was scavenged by MDCC. The P-bound MDCC-PBP released a fluorescent signal in 464 nm when excited at 425 nm. The fluorescent signals were measured by BioTek Synergy 4 plate reader.

**DNA Packaging Assay.** DNA packaging assay has been previously described. 45 In short, the viral prohead-connector complex, rRNA, gp16, and ph29 DNA-gp3 complex were mixed together. DNase was used to remove the unpackaged DNA, and after that proteinase K was digested with proteinase K to release the packaged DNA. The packaged DNA was observed through a 1% agarose gel in Tris borate EDTA (TBE) buffer. The wild-type gp16 was diluted with either a titration of R146A gp16 mutant or gp16 solution buffer as the control.

**AUTHOR INFORMATION**

**Corresponding Author**
Peixuan Guo — Center for RNA Nanobiotechnology and Nanomedicine, College of Pharmacy and College of Medicine, James Comprehensive Cancer Center, Dorothy M. Davis Heart and Lung Research Institute, and Biomedical Science Graduate Program, College of Medicine, The Ohio State University, Columbus, Ohio 43210, United States; orcid.org/0000-0001-5706-2833; Email: Guo.1091@osu.edu

**Author**
Chenxi Liang — Center for RNA Nanobiotechnology and Nanomedicine, College of Pharmacy and College of Medicine, James Comprehensive Cancer Center, Dorothy M. Davis Heart and Lung Research Institute, and Biomedical Science Graduate Program, College of Medicine, The Ohio State University, Columbus, Ohio 43210, United States

Complete contact information is available at: https://pubs.acs.org/10.1021/acsnano.1c02973

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**Notes**

The authors declare the following competing financial interest(s): P.G. is the licensor, consultant and grantee of Oxford Nanopore Technologies; the co-founder of Shenzhen P&Z Bio-medical Co. Ltd; and the co-founder of ExonanoR-NA, LLC and its subsidiary Weina (Guangdong) biomedical, Inc.

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