Effect of the MotB(D33N) mutation on stator assembly and rotation of the proton-driven bacterial flagellar motor

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Received April 1, 2014; accepted May 28, 2014

The bacterial flagellar motor generates torque by converting the energy of proton translocation through the transmembrane proton channel of the stator complex formed by MotA and MotB. The MotA/B complex is thought to be anchored to the peptidoglycan (PG) layer through the PG-binding domain of MotB to act as the stator. The stator units dynamically associate with and dissociate from the motor during flagellar motor rotation, and an electrostatic interaction between MotA and a rotor protein FliG is required for efficient stator assembly. However, the association and dissociation mechanism of the stator units still remains unclear. In this study, we analyzed the speed fluctuation of the flagellar motor of Salmonella enterica wild-type cells carrying a plasmid encoding a nonfunctional stator complex, MotA/B(D33N), which lost the proton conductivity. The wild-type motor rotated stably but the motor speed fluctuated considerably when the expression level of MotA/B(D33N) was relatively high compared to MotA/B. Rapid accelerations and decelerations were frequently observed. A quantitative analysis of the speed fluctuation and a model simulation suggested that the MotA/B(D33N) stator retains the ability to associate with the motor at a low affinity but dissociates more rapidly than the MotA/B stator. We propose that the stator dissociation process depends on proton translocation through the proton channel.

Key words: Salmonella, stator, torque generation, proton channel, speed fluctuation
nificantly reduce the probability of GFP-MotB localization to the motor as well. These suggest that electrostatic interactions between MotA Arg-90 and FliG Asp-289 and between MotA Gln-98 and FliG Arg-281 are required for efficient stator assembly around the rotor.

At least 11 copies of the MotA/MotB (hereafter MotA/B) complex are incorporated around the rotor to be the stators. Since MotB contains a highly conserved peptidoglycan-binding (PGB) motif in its C-terminal domain (MotBc), the MotA/B complex is thought to be anchored around the rotor through the PGB motif. High resolution single molecule imaging techniques have shown the turnover of GFP-fused stator unit between the motor and the membrane pool during rotation, providing evidence that the stator unit is not always kept anchored to each specific binding site but goes through a rather dynamic association/dissociation process. Recently, it has been shown that the flagellar motor responds to changes in external load to control the number of functional stators in the motor and that MotB acts as a load-sensitive structural switch to regulate the assembly and disassembly of the stators in response to the load changes. Interestingly, the MotA/B complex senses even a small change in the external load to regulate its proton conductivity during flagellar motor rotation.

MotB(D33N) and MotA still form the stator complex but the proton conductivity is lost and therefore the complex is non-functional as the stator. The MotA/B(D33N) complex exerts a strong dominant negative effect on the motility of wild-type cells in Salmonella. This is in agreement with the observation by an epillumination fluorescence microscopy that GFP-fused MotB(D33N) is localized to the flagellar motor. These suggest that the MotA/B(D33N) complex lacking the proton conductivity can be incorporated into the motor. Interestingly, over-expression of the MotA/B(D33N) complex reduces the number of functional stators to one or two, while the wild-type motor kept rotating even when the expression level of the stator complex is based on the result of model simulation. Computer simulation was performed by a simple kinetic model proposed before (see Computer simulation in Results). Kinetic parameters were manually optimized to reproduce experimental results. The sampling interval in the simulation was the same as that of the bead assays (1 ms), and the expression level of the stator complex is based on the result of immunoblotting. All calculations were performed using a macro developed in Microsoft Excel.

Materials and Methods

Bacterial strains, plasmids, media

A Salmonella strain, SJW46(fliCΔ204–292)), was used in this study. This strain has intact flagellar motors with sticky flagellar filaments and chemotaxis system. A plasmid, pNSK9-D33N, encodes wild-type MotA and MotB(D33N) on pTrc99A (Pharmacia). Luria broth (LB) and motility buffer were prepared as described before.

Bead assay

Overnight culture of SJW46 cells transformed with pTrc99A or pNSK9-D33N was diluted into fresh LB with or without IPTG and incubated for 4 h at 37°C. Bead assays were carried out as previously described. The position of 1-μm bead attached on the sticky filament of SJW46 was determined by a quadrant photodiode at 1-ms intervals. The average value and the standard deviation of rotation rate were calculated according to Muramoto et al. The rotational diffusions of the flagellar filament with bead were calculated as described before which was evaluated from the rotation rate and a half of the sampling interval.

Immunoblotting

SJW46 carrying pTrc99A or pNSK9-D33N was grown in LB under the same condition with the bead assay. Cultures were centrifuged to obtain cell pellets. The cell pellets were resuspended in SDS-loading buffer, normalized in cell density to give a constant amount of cells. Immunoblotting using polyclonal anti-MotB antibody was carried out as previously described.

Computer simulation

Computer simulation was performed by a simple kinetic model proposed before (see Computer simulation in Result). Kinetic parameters were manually optimized to reproduce experimental results. The sampling interval in the simulation was the same as that of the bead assays (1 ms), and the expression level of the stator complex is based on the result of immunoblotting. All calculations were performed using a macro developed in Microsoft Excel.

Results

Speed fluctuation of Salmonella flagellar motor

When MotA/B(D33N) was expressed in the wild-type cell, a much higher expression level than that of MotA/B was required to reduce the number of functional stators around the rotor to one or two. This indicated that the binding affinity of the MotA/B(D33N) complex for the rotor is much lower than that of the wild-type MotA/B complex. To test this, we changed the expression level of the MotA/B(D33N) complex in the Salmonella SJW46 cell,
The expression level of MotA/B was judged by immunoblotting with polyclonal anti-MotB antibody. Even in the absence of IPTG, the expression level of MotB(D33N) was 10-fold higher than MotB expressed from the chromosome, and it increased further by adding IPTG (Fig. 2). The rotation of the motor frequently slowed down and/or stopped, but interestingly, the motor speed was restored to the original level after those slowdown and stop events (Fig. 1B–D).

Figure 3 shows that the σω/<ω> value of the motor containing MotA/B(D33N) significantly increases when the rotation speed is reduced. The wild-type motor stably rotated at about 70 Hz, and the value of σω/<ω> ranged from 0.1 to 0.2. The rotation rate of the motor containing MotA/B(D33N) ranged from 10 Hz to 50 Hz in the absence of IPTG, and the values of σω/<ω> were between 0.1 and 0.4. When MotA/B(D33N) was over-expressed by adding IPTG, the motor speed was reduced down to a few hertz, and the value of σω/<ω> was increased up to 0.5 (Fig. 3). Even in the
absence of IPTG, the expression level of MotB(D33N) was 10-fold higher than MotB expressed from the chromosome (Fig. 2). As shown in Figure 1, the motor speed fluctuated but never completely stopped for a significant period of time. These observations suggest that the MotA/B(D33N) complex retains the ability to assemble into the motor but with much lower affinity than the MotA/B complex and rapidly dissociates from the motor to be replaced with the wild-type stator.

**Computer simulation**

To reproduce the speed fluctuation in the presence or absence of the MotA/B(D33N) stator, we used a simple kinetic model proposed by Muramoto *et al.* It was assumed that 11 stators independently function in the motor and that a single wild-type stator unit generates 150 pN nm but the D33N stator unit produces neither positive nor negative torque. The wild-type and D33N stators associate with or dissociate from the rotor with the following rate constants: \( k_{wt+} \) and \( k_{wt-} \), association-rate and dissociation-rate constants of the wild-type stator; \( k_{d33n+} \) and \( k_{d33n-} \), association-rate and dissociation-rate constants of the D33N stator. The event occurs at an interval of \( 10^{-4} \) sec (\( \Delta t \)), and the rotation speed is sampled at an interval of \( 10^{-3} \) sec. The association and dissociation probabilities of \( P_{wt+}, P_{wt-}, P_{d33n+}, \) and \( P_{d33n-} \) are given as \( P_{wt+} = k_{wt+}, \Delta t \), \( P_{wt-} = k_{wt-}, \Delta t \), \( P_{d33n+} = k_{d33n+}, C_{d33n} \Delta t \), and \( P_{d33n-} = k_{d33n-}, \Delta t \), where \( C_{wt} \) and \( C_{d33n} \) are the copy number of wild-type stators and D33N stators, respectively. A random number (RND) from 0.0 to 1.0 is generated for each stator unit. For example, if a stator binding position around the rotor is vacant and \( P_{wt+} < \text{RND} \), the wild-type stator associates with the rotor and generates torque. If the position is occupied by the wild-type stator and \( P_{wt+} < \text{RND} \), the stator dissociates from the rotor and the position becomes vacant.

If \( k_{wt+} = k_{d33n+} \) and \( k_{wt-} = k_{d33n-} \), the number of wild-type stator units rapidly reduced to zero when the copy number of the D33N stators was 10-fold higher than that of the wild-type stators (\( C_{d33n}/C_{wt} = 10 \)) (Fig. 4A upper panel). In contrast, if \( k_{wt+} = k_{d33n+} \) but \( k_{d33n-} \) was 10-fold larger than \( k_{wt-} \), the apparent number of stators temporarily decreased but was restored even when the copy number of D33N stators was 25-fold higher than that of wild-type stators (\( C_{d33n}/C_{wt} = 25 \)) (Fig. 4A lower panel). The relationship between \( <\omega> \) and \( \sigma_{\omega}/<\omega> \) obtained by this simulation is in a relatively good agreement with the present experimental result (Fig. 4B and 3). This suggests that the D33N mutation significantly affects the dissociation constant of the stator unit but not the association constant, to cause the large fluctuation of the motor speed in the presence of both the MotA/B and MotA/B(D33N) complexes. Note that the dissociation constant of the wild-type stator assumed here (= 0.4 s\(^{-1}\)) is 10-fold higher than the value predicted from fluorescent observation of GFP-MotB (= 0.04 s\(^{-1}\))\(^{33,36}\). This raises the possibility that the GFP tag significantly affects the dissociation rate of the MotA/B complex from the motor.

**Discussion**

The MotA/B complex is assembled into the flagellar motor to be the stator through the binding of the PGB domain of MotB to the PG layer\(^{23–25}\). At least 11 stator units can be assembled into the motor, and their association and dissociation are dynamic in response to changes in external load\(^{23–25}\). It has been shown that depletion of PMF induces the dissociation of GFP-MotB from the motor in *E. coli*\(^{36}\). Consistently, the D32A mutation in *E. coli* MotB interferes with the assembly of GFP-MotB to the flagellar motor\(^{36}\). In contrast, the *Salmonella* MotA/B(D33N) complex with totally impaired proton conductivity can still associate with the
rotor and significantly reduces the motor speed when over-expressed in wild-type cells\cite{17}. In agreement with this, depletion of PMF does not abolish the localization of GFP-MotB to the flagellar base\cite{14}. In this study, we showed that a large speed fluctuation of the flagellar motor occurs when both the wild-type and non-functional MotA/B complexes are expressed in *Salmonella* wild-type cells. A high-level expression of the MotA/B(D33N) complex in wild-type cells caused a large speed fluctuation and pausing of motor rotation (Fig. 1).

Several studies concerning unstable rotation of the flagellar motor have been reported to date, and possible sources to generate the speed fluctuation have been proposed as those including the fluctuations in input energy, the number of stator units and the rotational Brownian motion\cite{27,33,34,37,38}. The rate limiting process of proton translocation is thought to limit the maximal motor speed when the motor rotates under low-load conditions, but the effect of proton influx becomes smaller with an increase in the drag force against the motor\cite{39,41}. The motor speed is proportional to the number of stator units under high-load conditions, but just one stator unit can spin the motor at the maximum speed under low-load conditions\cite{18,42,43}. The MotB(D33E) mutation, which reduces the proton influx to about a half of the wild-type MotA/B complex, causes a marked reduction in the maximum motor speed by more than 10 fold and shows a large speed fluctuation only under low load conditions, implying that the stator is a load-sensitive proton channel that efficiently couples proton translocation with torque generation\cite{27}.

Since the characteristics of the flagellar motor rotating with a 1-μm bead belong to high-load regime in the torque-speed relationship of *Salmonella*, the unstable rotation speed observed in this study is unlikely to be caused by the fluctuation of the proton influx through the MotA/B proton channel. Our present results indicate that the MotA/B(D33N) complex competes with the wild-type MotA/B complex for the stator-binding sites of the motor, associates with the rotor, and reduces the number of functional stator units in a rather dynamic way.

The motors were never completely paralyzed even when the expression level of the MotA/B(D33N) complex reached above 10 fold of the wild-type MotA/B complex (Fig. 2), suggesting that the time period that a MotA/B(D33N) complex occupies a stator binding position is rather short. A computer simulation reproduced the observed speed fluctuations only when the dissociation rate of the MotA/B(D33N) stator was 10-fold higher than that of the wild-type stator, without changing the association rate (Fig. 4). Because the D33N mutation interferes with proton translocation through the proton channel\cite{6,28}, we propose that the arrest of proton transduction through the channel suppresses a conformational change or flexibility of the stator required for its stable association with the rotor to make its detachment from the rotor faster. Even in the wild-type motor, when a stator unit becomes malfunctioned during motor rotation, this characteristic may allow the motor to efficiently exchange the broken stator with the functional one.

In our simulation, the stator units were assumed to be independent of each other, and the cooperative association and dissociation of the stator to the motor was not adopted. The simulation reproduced the large fluctuation in the number of functional stators as shown in the experimental results (Fig. 3 and 4A), suggesting that the cooperativity is not really necessary in the mechanism of stator assembly. However, this does not completely eliminate the possibility that a stator dynamics influences its neighbors. To make this clear, direct observation of the stator dynamics around the motor at higher temporal and spatial resolution is required.

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**Figure 4** Computer simulation of motor rotation by a dynamic exchange of the wild-type and mutant stators. (A) Temporal change in the number of wild-type stator units at \( k_{d33n} / k_{wt} = 1 \) and \( C_{d33n} / C_{wt} = 10 \) (upper panel), and at \( k_{d33n} / k_{wt} = 10 \) and \( C_{d33n} / C_{wt} = 25 \) (lower panel). Initial numbers of wild-type stator units were assumed to be 5. (B) Values of \( \sigma / <\omega> \) plotted as a function of \( <\omega> \) when the value of \( C_{d33n} / C_{wt} \) was varied from 0 to 30. The calculation for (B) was done with parameters as follows: \( k_{d33n} = 1 \) copy/s\(^{-1}\), \( k_{wt} = 4 \) s\(^{-1}\), \( k_{d33n} = 0.4 \) s\(^{-1}\), and \( C_{wt} = 200 \) copy.
Acknowledgement

We thank Y. Sowa for helpful discussion and comments. This research has been supported in part by JSPS KAKENHI Grant 24770141 to S.N., 23115008 on Innovative Areas ‘Spying minority in biological phenomena’ to T.M. and 21227006 and 25000013 to K.N.

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