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The bacterial flagellar protein export apparatus processively transports flagellar proteins even with extremely infrequent ATP hydrolysis

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For self-assembly of the bacterial flagellum, a specific protein export apparatus utilizes ATP and proton motive force (PMF) as the energy source to transport component proteins to the distal growing end. The export apparatus consists of a transmembrane PMF-driven export gate and a cytoplasmic ATPase complex composed of FliH, FliI and FliJ. The FliI₆FliJ complex is structurally similar to the $\alpha_3\beta_3\gamma$ complex of F_oF₁-ATPase. FliJ allows the gate to efficiently utilize PMF to drive flagellar protein export but it remains unknown how. Here, we report the role of ATP hydrolysis by the FliI₆FliJ complex. The export apparatus processively transported flagellar proteins to grow flagella even with extremely infrequent or no ATP hydrolysis by FliI mutation (E211D and E211Q, respectively). This indicates that the rate of ATP hydrolysis is not at all coupled with the export rate. Deletion of FliI residues 401 to 410 resulted in no flagellar formation although this FliI deletion mutant retained 40% of the ATPase activity, suggesting uncoupling between ATP hydrolysis and activation of the gate. We propose that infrequent ATP hydrolysis by the FliI₆FliJ ring is sufficient for gate activation, allowing processive translocation of export substrates for efficient flagellar assembly.

AAA⁺ family ATPases, which are involved in various cellular activities such as DNA replication, proteolysis and membrane fusion, usually form ring-shaped oligomers with a narrow central channel. AAA⁺ ATPases couple ATP binding and hydrolysis to the translocation of their substrates to the central channel. Coordination and cooperativity among subunits in the ring-shaped ATPases are critical for their biological activities^{1,2}.

The bacterial flagellum is a rotary nanomachine powered by PMF across the cytoplasmic membrane. It is composed of about 30 different proteins with their copy numbers ranging from a few to tens of thousands. The flagellum is divided into at least three parts: the basal body, the hook, and the filament. Flagellar assembly begins with the basal body, followed by the hook and finally the filament. The flagellar export apparatus of *Salmonella enterica* is a type III secretion system and consists of a membrane-embedded export gate made of FlhA, FlhB, FliO, FliP, FliQ and FliR and a cytoplasmic ATPase complex consisting of FliH, FliI and FliJ and transports flagellar proteins from the cytoplasm to the distal end of the growing flagellar structure for self-assembly. The flagellar export apparatus is evolutionally related to the following two nanomachines: the injectisome of pathogenic bacteria, which directly inject virulence factors into their host cells; and the F- and V-type ATPases^{3,4}.

The flagellar export apparatus utilizes both ATP and PMF as the energy sources for protein export^{5,6}. FliI is the ATPase of the export apparatus⁷ and forms a homo-hexamer with a narrow central pore^{8,9}. The FliI₆ ring has been structurally identified at the flagellar base by electron cryotomography¹⁰. FliJ binds to the center of the FliI₆ ring to form the FliI₆FliJ ring, which looks very similar to F₁-ATPase where the α/β and γ subunits correspond to FliI and FliJ, respectively^{11,12}. FliH binds to FliI¹³ and anchors the FliI₆FliJ ring complex to the export gate through interactions of FliH with a C ring protein FliN and FlhA^{14,15}. The export gate is intrinsically a proton-protein antiporter that uses the two components of PMF, $\Delta\psi$ and ΔpH , for different steps of the export process¹⁶. An interaction between FliJ and FlhA turns the export gate into a highly efficient $\Delta\psi$ -driven export apparatus¹⁶. Although FliH, FliI and FliJ are dispensable for protein export, they make the export gate highly more efficient



than their absence by which most of *Salmonella* cells cannot form flagella at all^{5,6}. However, the roles of PMF and the ATPase are still under strong debate because the actual mechanistic role of the ATPase has remained unclear.

In vivo fluorescent imaging of FliI-YFP by fluorescence microscopy with single molecule precision has shown that not only the FliI₆ ring but also several FliH₂FliI complexes are associated with the flagellar basal body (FBB) through interactions of FliH with FliN and FliA and that about 90% of the FliI-YFP spots show turnover between the FBB-localized and free-diffusing ones after photobleaching¹⁷. Neither the number of FliI-YFP associated with the FBB nor FliI-YFP turnover rate are affected by catalytic mutations in FliI, indicating that ATP hydrolysis by FliI does not drive the assembly-disassembly cycle of FliI during flagellar assembly¹⁷. In this study, to clarify the actual mechanistic role of ATP hydrolysis in flagellar protein export, we characterized the *Salmonella fliI(E211Q)* and *fliI(E211D)* catalytic mutants and an in-frame *fliI* deletion mutant with a reduced ATPase activity. We show that the export gate processively transports flagellar proteins during flagellar assembly even with extremely infrequent ATP hydrolysis. We also show that deletion of residues 401 to 410 of FliI considerably affects the energy coupling between ATP hydrolysis and activation of the gate by the FliI₆FliI ring.

Results

Effect of the E211Q and E211D mutations on flagellar protein export. The carboxyl group of Glu-211 in FliI, which corresponds to Glu-190 in the thermophilic *Bacillus* F₁-ATPase, polarizes a water molecule for the nucleophilic attack to the γ -phosphate of ATP (Fig. 1)^{11,18,19}. To investigate whether ATP hydrolysis by FliI limits the rate of flagellar protein export, we replaced Glu-211 with Asp and measured the ATPase activity. We used FliI(E211Q) as a negative control because the E211Q substitution results in the complete loss of its ATPase activity¹⁹. The ATPase activity of FliI(E211D) was measured quantitatively by subtracting the data from FliI(E211Q) as a background, and it was about 100 times lower than that of wild-

type FliI (Fig. 2a). The E211Q mutation in FliI considerably reduced both motility (Fig. 2b) and flagellar protein export (Fig. 2c, lane 7). In contrast, more than 80% of the *fliI(E211D)* mutant cells were motile, and their swimming speed was about 50% of the wild-type level (Fig. 2b and Fig. S1). Consistently, most flagellar proteins were detected in the culture supernatant of this *fliI(E211D)* mutant (Fig. 2c, lane 8). Interestingly, the levels of FlgD (1st row), FliK (3rd row) and FliD (6th row) secreted by the *fliI(E211D)* mutant were comparable to the wild-type levels while the secretion levels of FlgE (2nd row), FlgK (4th row) and FlgL (5th row) were four, two, and three-fold lower than wild-type levels, respectively. The E211Q and E211D mutations did not affect the cellular levels of FliI at all (Fig. 2c, 7th row). Therefore, these results indicate that the rate of ATP hydrolysis by FliI does not determine the rate of flagellar protein export.

We then quantitatively analyzed the number of the flagellar filaments produced by these two *fliI* mutants (Fig. 3a). The number of the filaments produced by wild-type cells ranged from 1 to 8 with an average of 4.4 ± 1.6 (Fig. 3b). About 83% of the *fliI(E211Q)* cells had no filaments while the remaining population had one or two filaments with length only about 25% of the wild-type (Fig. 3b and c). In contrast, more than 90% of the *fliI(E211D)* cells produced filaments of which number ranged from 1 to 7 with an average of 2.3 ± 1.5 (Fig. 3b). The average filament length was about half of the wild-type (Fig. 3c). These results suggest that the export gate can processively export flagellar proteins during flagellar assembly even with extremely infrequent ATP hydrolysis.

To test whether FliI(E211Q) and FliI(E211D) are incorporated into the export apparatus, we analyzed their dominant negative effect on motility of wild-type cells. Because FliI with a catalytic mutation requires FliH for its negative dominance¹³, we used a $\Delta fliH$ -*fliI fliB(P28T)* bypass mutant that can form some flagella even in the absence of FliH and FliI⁵ as a control. FliI(E211Q) and FliI(E211D) both inhibited the motility of wild-type cells (Fig. S2a) but not that of the $\Delta fliH$ -*fliI fliB(P28T)* bypass mutant (Fig. S2b), suggesting their assembly into the export apparatus in the presence of FliH.

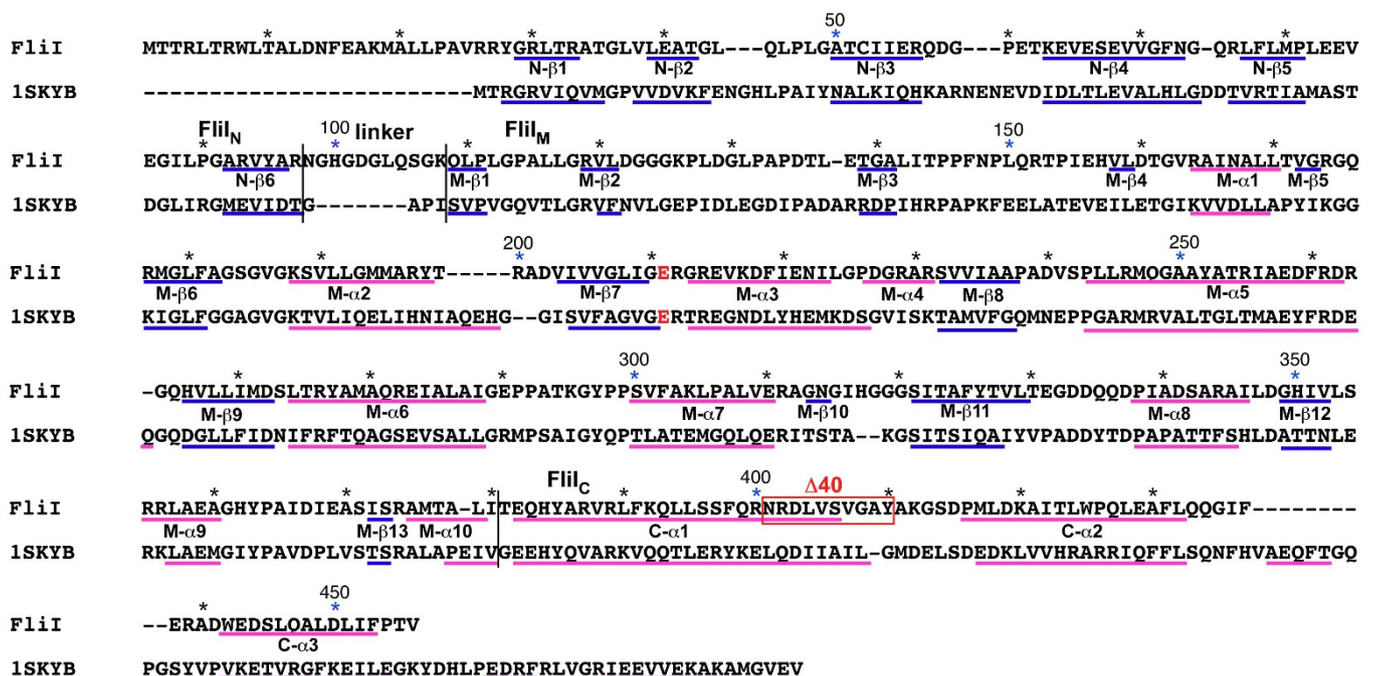


Figure 1 | Structure-base sequence alignment of FliI and F₁-β subunit from the thermophilic *Bacillus* PS3 (1SKYB). The regions of secondary structural elements are shown below each sequence: magenta line, α -helix; blue line, β -structure. The secondary structural elements are labeled with initials of three domains and numbers. A highly conserved Glu residue is highlighted by red. In-frame deletion of residues 401–410 in FliI (indicated as $\Delta 40$) is indicated by a red box.

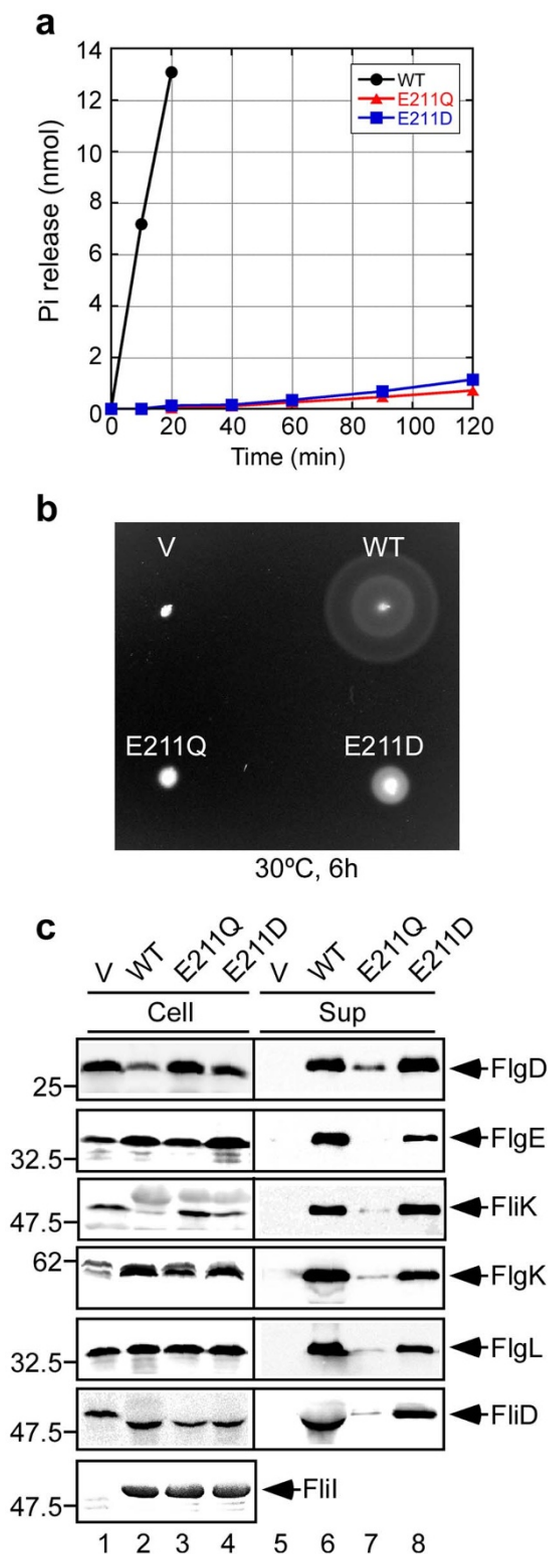


Figure 2 | Characterization of the *fliI*(E211Q) and *fliI*(E211D) mutants. (a) Effect of the E211Q and E211D mutations on the FliI ATPase activity. Measurements were carried out on purified His-FliI (indicated as WT, black filled circle), His-FliI(E211Q) (indicated as E211Q, red filled triangle) or His-FliI(E211D) (indicated as E211D, blue filled square) using malachite green assay. (b) Swarming motility of a $\Delta fliI$ mutant transformed with pTrc99A (V), pMM1702 (WT), pKK211 (E211Q) or pMM1702(E211D) (E211D) in soft agar plates. The plate was incubated at 30°C for 6 hours. (c) Effect of the E211Q and E211D mutations on flagellar protein export. Whole cell proteins (Cell) and culture supernatant fractions (Sup) were prepared from the above transformants, subjected to

SDS-PAGE, and analyzed by immunoblotting with polyclonal anti-FlgD (1st row), anti-FlgE (2nd row), anti-FliK (3rd row), anti-FlgK (4th row), anti-FlgL (5th row), anti-FliD (6th row) or anti-FliI (7th row) antibody. The positions of molecular mass markers are indicated on the left.

Effect of an in-frame deletion of FliI residues 401 to 410 on flagellar protein export. The export gate requires an interaction of FliJ with FlhA to drive protein export in a $\Delta\psi$ -dependent manner¹⁶, raising the possibility that ATP hydrolysis by FliI activates the export gate through the FliJ-FlhA interaction. Because FliJ binds to the C1 α -helix of the C-terminal domain of FliI¹², we constructed a series of FliI variants with sequential 10-amino-acid deletions within the C-terminal domain from residue 381 to 456 (FliI_C). All the deletion variants were too insoluble to measure the ATPase activity, except for FliI Δ 40 lacking residues 401 to 410 (Fig. 1). The ATPase activity of FliI Δ 40 was about 40% of wild-type FliI (Fig. 4a). To test whether this reduction in the ATPase activity resulted from a decrease in the probability of FliI ring formation, we analyzed the capability of FliI Δ 40 to form the ring in the presence of Mg^{2+} -ADP- AlF_4 , which stabilizes the FliI₆ ring structure¹⁹. FliI Δ 40 formed a homo-hexamers at the wild-type level (Fig. 4b). These results indicate that residues 401–410 of FliI are required for the full ATPase activity but not for FliI₆ ring formation.

We then measured the protein export activity of the *fliI* Δ 40 mutant. The *fliI* Δ 40 mutant displayed a very weak motility phenotype only after prolonged incubation (Fig. 4c). Consistently, very low levels of FlgD were detected in the culture supernatant (Fig. 4d, lane 6). The majority of *fliI* Δ 40 cells had no flagellar filament, and only a few % of cells had one short filament (Fig. 3). FliI Δ 40 exerted a dominant negative effect on motility of wild-type cells (Fig. S3a) but not on that of the *fliH-fliI flhB*(P28T) bypass mutant (Fig. S3b). Therefore, FliI Δ 40 is not functional in driving efficient protein export even though the (FliI Δ 40)₆ ring is assembled into the export apparatus.

Since FliI binds to the C-terminal cytoplasmic domains of FlhA (FlhA_C) and FlhB (FlhB_C), FliJ and chaperone substrate complexes, the FliI₆ ring is proposed to act as a static substrate loader to facilitate the initial entry of export substrate into the export gate¹⁷. To test whether the deletion of residues 401–410 in FliI affects these interactions, we carried out pull-down assays by GST affinity chromatography. FliI co-purified with GST-FlhA_C, GST-FlhB_C, GST-FliJ and GST-FliT94/FliD complex but not with GST alone, in agreement with previous reports^{14,20,21} (Fig. 5, left panels). The amounts of FliI Δ 40 co-purified with GST-FlhA_C, GST-FlhB_C, GST-FliJ and GST-FliT94/FliD complex were essentially the same as those of FliI (Fig. 5, right panels). These results indicate that the impaired export activity of the *fliI* Δ 40 mutant does not result from the impaired interactions of FliI with them. Therefore, we suggest that the (FliI Δ 40)₆ ring is simply wasting the energy of ATP hydrolysis even with its proper incorporation into the export apparatus through interactions with FliJ and FlhA.

Discussion

ATPases convert the chemical energy released by ATP hydrolysis into mechanical works required for their biological activities. In general the rate of ATP hydrolysis is coupled to their biological activity. SecA ATPase utilizes free energy derived from ATP hydrolysis to drive preprotein translocation through the SecYEG protein channel. Each catalytic cycle of ATP binding and hydrolysis is coupled to conformational changes in SecA that drive the stepwise translocation of preproteins across the membrane in a way similar to AAA⁺ ATPases such as ClpX and FtsH²². InvC, a FliI homolog of the *Salmonella* injectisome, forms a homo-hexamers, as has been visualized by electron cryomicroscopy at the base of the *Salmonella* injectisome²³, and is thought to induce the release of chaperone from the chaperone-substrate complex and to unfold the substrate for efficient

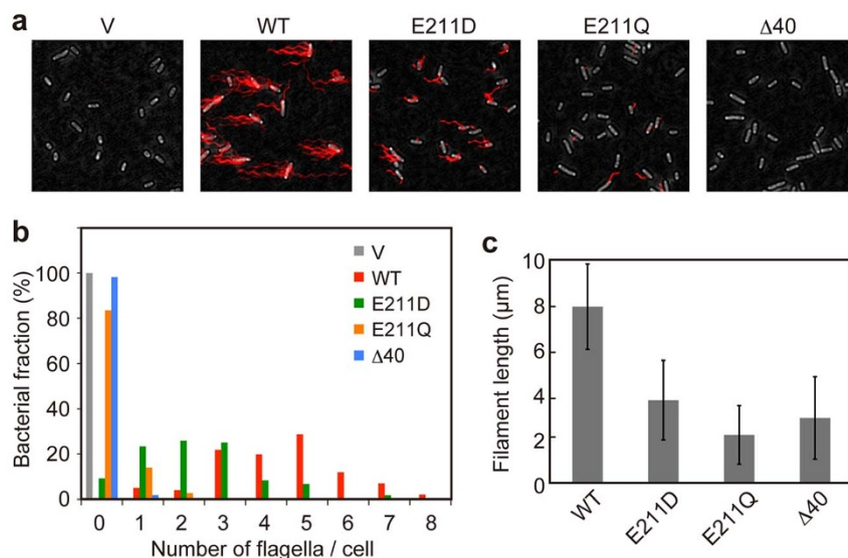


Figure 3 | Effect of FliI mutations on the number and length of the flagellar filaments. (a) Fluorescent images of the $\Delta fliI$ mutant harboring pTrc99A (V), pMM1702 (WT), pMM1702(E211D) (E211D), pKK211 (E211Q) or pMM140 ($\Delta 40$). Flagellar filaments were labeled with Alexa Fluor 594. The fluorescence images of the filaments labeled with Alexa Fluor 594 (red) were merged with the bright field images of the cell bodies. (b) Distribution of the number of the flagellar filaments in the $\Delta fliI$ mutant carrying pTrc99A (V, gray), pMM1702 (WT, red), pMM1702(E211D) (E211D, green), pKK211 (E211Q, orange) or pMM140 ($\Delta 40$, light blue). More than 100 cells for each transformants were counted. (c) Measurements of the length of the flagellar filaments produced by each transformants. Filament length is the average of more than 100 cells for the wild-type, *fliI*(E211D) and *fliI*(E211Q) cells and 20 cells for the *fliI $\Delta 40$ cell, and vertical lines are standard deviations.*

protein export in a way similar to the AAA⁺ ATPases²⁴. In contrast, flagellar assembly occurs even in the absence of FliI, indicating that FliI is not essential^{5,6}. Furthermore, FliI does not induce the release of flagellar chaperone from the chaperone-substrate complex in an ATP-dependent manner²¹. These observations suggest that the FliI₆ ring does not act as an unfoldase to facilitate the substrate entry into the export gate. In this study, we showed that more than 90% of the *fliI*(E211D) mutant cells have a couple of flagella with length nearly half of the wild-type (Fig. 3) despite the 100-fold lower ATPase activity than that of wild-type FliI (Fig. 2a). Interestingly, the levels of FlgD, FliK and FliD secreted by the *fliI*(E211D) mutant were at the wild-type levels (Fig. 2c, lane 8). Furthermore, the *fliI*(E211Q) mutant, in which the mutation does not affect ATP binding although resulting in the complete loss of the ATPase activity¹⁹, exported flagellar proteins to some degree (Fig. 2c, lane 7) and formed flagella albeit at very low probability (Fig. 3). Therefore, we suggest that the rate of ATP hydrolysis by FliI is not at all coupled with the protein export activity and that ATP consumption could be relatively small despite that 20,000–30,000 flagellin molecules have to be exported for the assembly of each flagellum.

Although deletion of FliH and FliI results in almost complete loss of flagellar formation, most of $\Delta fliH$ -*fliI* *flhB*(P28T) bypass mutant cells are capable of forming a couple of flagella even in the absence of FliH and FliI. This indicates that the *flhB*(P28T) bypass mutation increases the probability of entry of flagellar proteins into the export gate. Because an increased gate-opening probability for higher efficiency of protein entry could be deleterious to the cells due to leakage of small solutes, the interaction of export substrate with the gate would presumably induce the gate opening, with or without the FliI₆ ring⁵. Deletion of residues 401–410 in FliI (FliI $\Delta 40$) resulted in a considerably weak motility phenotype despite that the ATPase activity of FliI $\Delta 40$ was about 40% of the wild-type level (Fig. 4). Pull-down assays by GST affinity chromatography revealed that FliI $\Delta 40$ retained the ability to bind to FlhA_C, FlhB_C, FliJ and the FliT-FliD chaperone-substrate complex at the wild-type levels (Fig. 5). These results indicate that the $\Delta 40$ deletion causes uncoupling between ATP hydrolysis by the FliI₆ ring and flagellar protein export.

Therefore, we suggest that this deletion affects an activation step of the export gate by the FliI₆ ring rather than a gate-opening step. The export gate itself is an inefficient proton-protein antiporter, and since a specific interaction of FlhA with FliJ promoted by FliH and FliI switches the export gate to a highly efficient $\Delta\psi$ -driven export engine¹⁶, we propose that the FliI₆FliJ ring complex acts as an ATP-driven ignition key to turn on the export engine to be a $\Delta\psi$ -dependent, rapid and efficient protein transporter (Fig. 6).

How does the FliI₆FliJ ring complex act as the ignition key for the $\Delta\psi$ -driven export engine? The FliI₆FliJ ring complex is structurally very similar to the $\alpha_3\beta_3\gamma$ complex of F₁-ATPase^{11,12}. F₁-ATPase is an ATP-driven rotary motor, driving the γ subunit rotation within the $\alpha_3\beta_3$ ring. Three catalytic β -subunits in the $\alpha_3\beta_3$ ring undergo highly cooperative and sequential conformational changes in their C-terminal domains of the β -subunits coupled with ATP hydrolysis²⁵. It has been shown that ATP binds to the P-loop of FliI to induce a conformational change in FliI_C similarly to that observed in the β -subunits¹¹. FliJ binds to the C-terminal region of the C1 α -helix of FliI_C, which corresponds to the region of the β subunit responsible for the interaction with the γ subunit in F₁-ATPase¹². These similarities suggest that FliJ may also rotate within the FliI₆ ring. In fact FliJ actually shows a rotor like function within the central pore of the A₃B₃ complex of the *Thermus thermophilus* V-type ATPase²⁶. Because a highly conserved surface of FliJ is responsible for the interaction with FlhA²⁷, we propose that FliJ is a rotary ignition key driven by the FliI₆ ring to activate the export gate through interaction of the conserved surface of FliJ with FlhA. The ATP hydrolysis by FliI in the ring is likely to induce cooperative conformational changes in the FliI_C domains to rotate the FliJ ignition key.

This model can explain many aspects of our present results by analogy to F-ATPase. The E190D mutation in the β subunit considerably reduces the rotation rate of the γ subunit within the $\alpha_3\beta_3$ ring¹⁸. Therefore, markedly reduced efficiency of flagellar assembly by the *fliI*(E211D) mutant would be a consequence of a significant reduction in the rotation frequency of FliJ. The elementary step size of γ rotation is 120°, which consists of 80° and 40° substeps driven by ATP binding-ADP release and ATP hydrolysis-Pi release, respect-

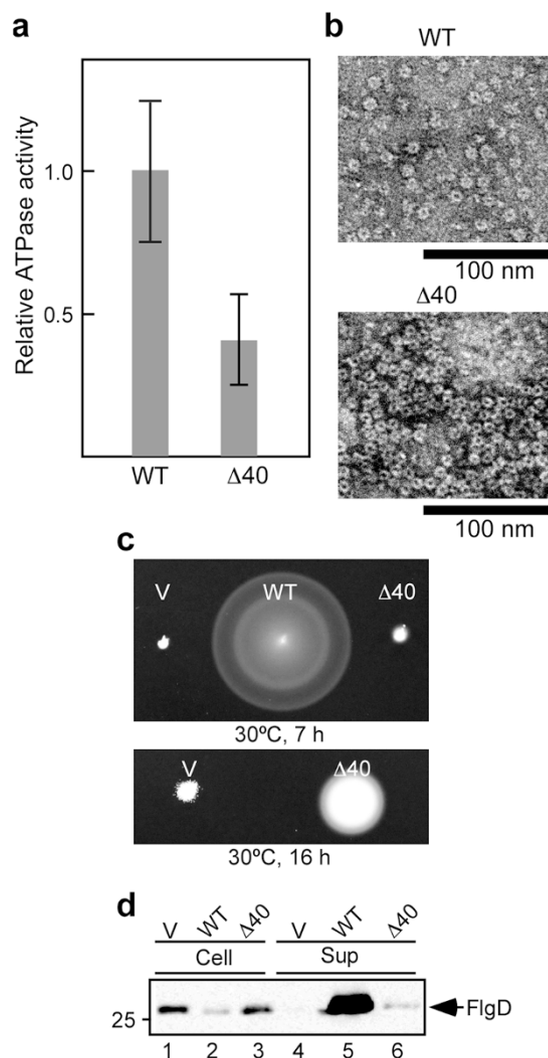


Figure 4 | Characterization of the *fliI*Δ40 mutant. (a) Relative ATPase activity of His-FliIΔ40. The ATPase activity of His-FliIΔ40 (indicated as Δ40) is normalized to that of His-FliI (indicated as WT). These data are the average of four independent experiments, and vertical lines are the standard deviations. (b) Electron micrographs of negatively stained samples of purified His-FliI and His-FliIΔ40 preincubated with 5 mM MgCl₂, 5 mM ATP (labeled Mg-ATP) or 5 mM MgCl₂, 5 mM ADP, 5 mM AlCl₃, 5 mM NaF and 100 μg/ml acidic phospholipids. (c) Swarming motility of the Δ*fliI* mutant transformed with pTrc99A (V), pMM1702 (WT), or pMMI40 (Δ40) in soft agar plates. (d) Immunoblotting, using polyclonal anti-FlgD antibody, of whole cell proteins and culture supernatant fractions prepared from the same transformants.

ively²⁵. So, just an ATP binding to the FliI(E211Q)₆ ring could rotate FliJ by 80°, and this may be sufficient to trigger the processive protein export to form a flagellum, although the frequency of this event would be very rare because no ATP hydrolysis occurs to make the FliI(E211Q)₆ ring ready for the next rotation event. Since the β-γ contact at the C-terminal domain of the β subunit is responsible for torque generation²⁵, it is possible that the Δ40 deletion impairs torque generation at the FliI-FliJ interface, resulting in a limited FliJ function as a rotary ignition key and therefore a very weak motility phenotype of the mutant cell. Thus, our data provides a clear insight into the mechanistic implication of the structural similarity between the FliI₆FliJ ring and F₁-ATPase and presents a very unique use of a rotary ATPase in a biological mechanism.

How does the export apparatus processively transport flagellar proteins even with extremely infrequent ATP hydrolysis? The cytoplasmic domain of FlhA (FlhA_C) forms a nonameric ring structure in the export apparatus^{28,29} and provides binding-sites for chaperone-substrate complexes^{30–32}. FliI also forms the FliH₂FliI complex together with a FliH₂ homo-dimer in the cytoplasm¹³. Not only the FliI₆ ring but also several FliH₂FliI complex associate with the FBB through interactions of FliH with FlhA and FliN¹⁷. The chaperone-substrate complexes bind to the FliH₂FliI complex through co-operative interactions among FliI, chaperone and export substrate^{20,21,30,33}. The FliH₂FliI complex shows rapid turnovers between the FBB and the cytoplasmic pool in an ATP-independent manner¹⁷, suggesting that the FliH₂FliI complex acts as a dynamic carrier to deliver the chaperone-substrate complexes from the cytoplasm to the FlhA_{C9} ring. Because FlhA_C is directly involved in the translocation of the export substrate³⁴, we propose that the FlhA_{C9} ring may play an important role in processive protein transport along with the FliI₆ ring that acts as a substrate loader upon the activation of the export gate by ATP hydrolysis by the FliI₆FliJ ring complex.

Methods

Bacterial strains, plasmids, and media. *Salmonella* strains and plasmids used in this study are listed in Table 1. L-broth (LB) and soft tryptone agar plates were prepared as described^{35,36}.

DNA manipulations. DNA manipulations were carried out as described before³⁷. Site-directed mutagenesis was performed as described previously³⁷. DNA sequencing was performed using BigDye ver 3.1 (Applied Biosystems), and the reaction mixtures were analyzed by a 3130 Genetic Analyzer (Applied Biosystems).

Protein expression and purification. His-FliI, His-FliI(E211Q), His-FliI(E211D) and His-FliIΔ40 were over-expressed in SJW1368 carrying pMM1702, pKK211, pMM1702(E211D) and pMMI40, respectively, and purified by Ni affinity chromatography using a Ni-NTA agarose column (QIAGEN) as described³⁶. GST-FlhA_C, GST-FlhB_C, FliJ and GST-FliT94 were overproduced in SJW1368 transformed with pMMA1001, pMMHB1001, pMMJ1001 and pMMT002, respectively, and purified by GST affinity chromatography using a glutathione Sepharose 4B column (GE Healthcare) as described before³⁷. 21. FliD was overproduced in BL21 (DE3) pLysS harboring pKOT134 and purified as described previously³⁸.

Measurements of FliI ATPase activity. The malachite green ATPase assay was carried out as describe previously⁷. At least four independent measurements were carried out.

Motility assays. For motility assay in soft agar, fresh colonies were inoculated on soft tryptone agar plates and incubated at 30°C. For measurements of free-swimming speeds of motile cells, overnight culture of *Salmonella* was inoculated into fresh LB and incubated at 30°C with shaking for 4 hours. The cells were washed twice with motility buffer (10 mM potassium phosphate pH 7.0, 0.1 mM EDTA, 10 mM L-sodium lactate) and resuspended in the motility buffer. *Salmonella* cells were observed under a phase contrast microscope (CH40, Olympus) at room temperature, and their motile behavior was recorded on hard disk drive. The swimming speed of individual cells was analyzed as described before³⁹.

Secretion assays. *Salmonella* cells were grown with shaking in 5 ml of LB at 30°C until the cell density had reached an OD₆₀₀ of ca. 1.4–1.6. Cultures were centrifuged to obtain cell pellets and culture supernatants. Cell pellets were resuspended in the SDS-loading buffer, normalized to a cell density to give a constant amount of cells. Proteins in the culture supernatants were precipitated by 10% trichloroacetic acid, suspended in the Tris/SDS loading buffer and heated at 95°C for 5 min. After SDS-PAGE, immunoblotting with polyclonal anti-FlgD, anti-FlgE, anti-FliK, anti-FliG, anti-FliL and anti-FliD antibodies were carried out as described before³⁵. Detection was performed with an ECL plus immunoblotting detection kit (GE Healthcare). At least three independent experiments were carried out.

Observation of flagellar filaments with a fluorescent dye. *Salmonella* cells were attached to a cover slip (Matsunami glass, Japan), and unattached cells were washed away with motility buffer. A 1 μl aliquot of polyclonal anti-FliC serum was mixed with 50 μl of motility buffer and then 50 μl of the mixture was applied to the cells attached to the cover slip. After washing with the motility buffer, 1 μl of anti-rabbit IgG conjugated with Alexa Fluor® 594 (Invitrogen) were mixed with 50 μl of motility medium and then the mixture was applied. After washing with the motility buffer, the cells were observed as described before⁴⁰. Fluorescence images were analyzed using ImageJ software version 1.48 (National Institutes of Health). To measure the filament length, each filament was traced by ImageJ freehand line tool and smoothed using the spline fit function.

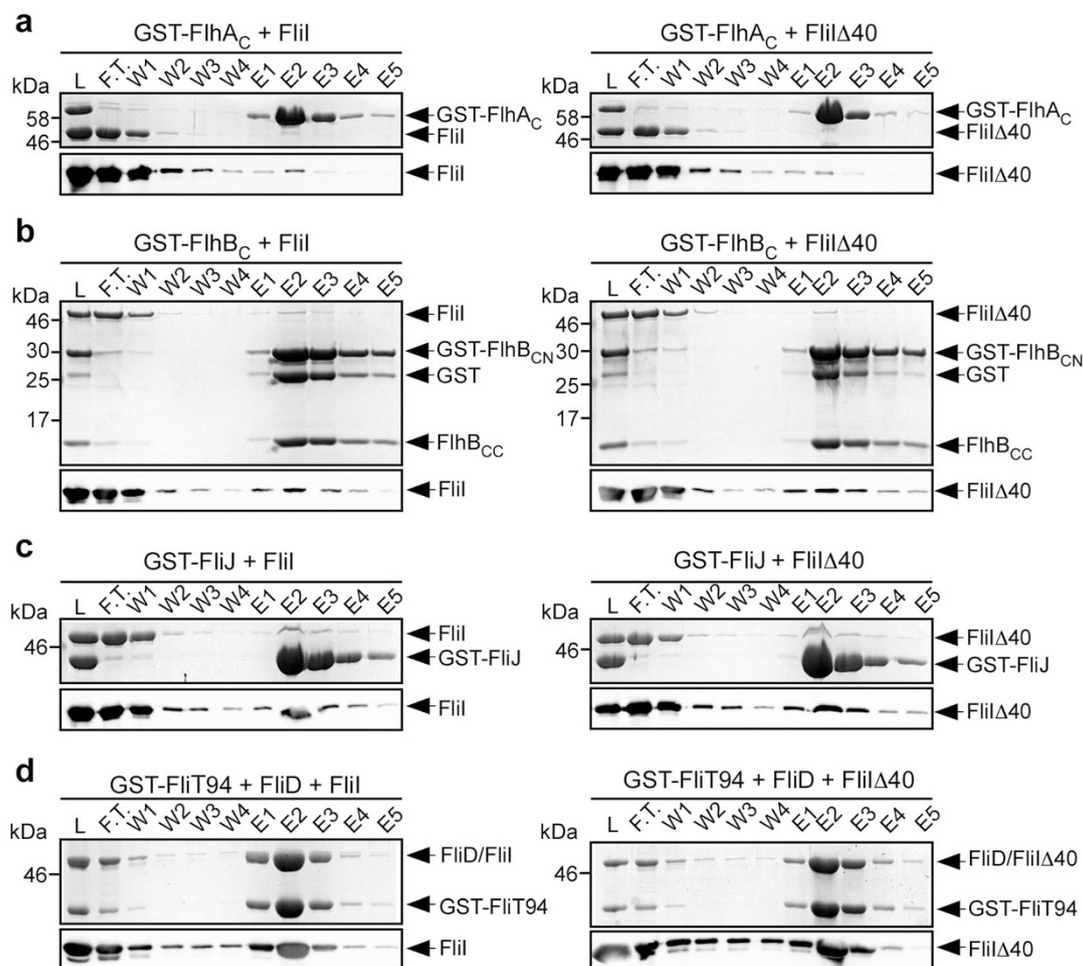


Figure 5 | Effect of in-frame deletion of residues 401–410 in FliI on the interactions of FliI with FlhA_C, FlhB_C, FliJ and the FliT94/FliD complex. (a) Interactions of GST-FlhA_C with FliI (left panel) and FliIΔ40 (right panel). Purified His-FliI or His-FliIΔ40 was mixed with GST-FlhA_C, and dialyzed overnight against PBS. These mixtures (L) were subjected to GST affinity chromatography. Flow through fraction (F.T.), wash fractions (W), and elution fractions (E) were analyzed by CBB staining (1st row) and immunoblotting with polyclonal anti-FliI antibody (2nd row). (b) Interactions of GST-FlhB_C with FliI (left panel) and FliIΔ40 (right panel). GST-FlhB_C is cleaved between Asp-269 and Pro-270 in an autocatalytic way⁴¹ but FlhB_{CC} remains to associate with GST-FlhB_{CN} even after cleavage. (c) Interactions of GST-FliJ with FliI (left panel) and FliIΔ40 (right panel). (d) Interactions of the GST-FliT94/FliD complex with FliI (left panel) and FliIΔ40 (right panel). The positions of molecular mass markers are shown on the left.

***in vitro* reconstitution of the FliI₆ ring and electron microscopy.** Purified His-FliI or His-FliIΔ40 were incubated in 35 mM Tris-HCl (pH 8.0), 113 mM NaCl, 1 mM DTT, 5 mM ADP, 5 mM AlCl₃, 15 mM NaF, 5 mM MgCl₂, and 100 μg/ml acidic phospholipids at 37°C for a few minutes. Samples were

applied to carbon-coated copper grids and negatively stained with 2% (w/v) uranyl acetate. Micrographs were recorded at a magnification of ×35,000 with a JEM-1011 transmission electron microscope (JEOL) operated at 100 kV.

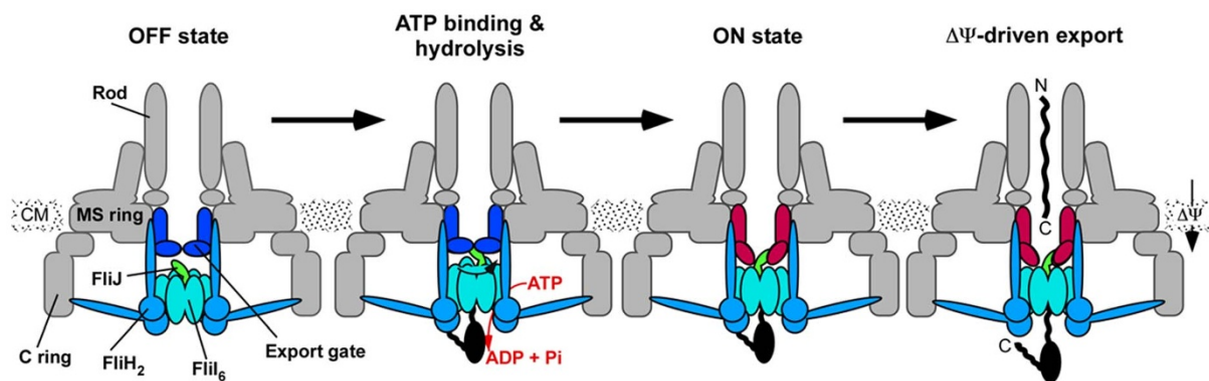


Figure 6 | An ignition key mechanism for flagellar protein export. ATP hydrolysis by the FliI₆ ring induces the FliJ rotation within the ring to cause a conformational change of the export gate to activate it, allowing rapid and efficient entry of the substrates into the gate. The export gate then efficiently utilizes the membrane potential component ($\Delta\Psi$) of proton motive force across the cytoplasmic membrane to unfold and transport the substrates into the central channel of the growing flagellar structure.



Table 1 | Strains and Plasmids used in this study

Strains and Plasmids	Relevant characteristics	Source or reference
<i>E. coli</i>		
BL21 (DE3)/pLysS	Overexpression of proteins	Novagen
<i>Salmonella</i>		
SJW1103	Wild type for motility and chemotaxis	(42)
SJW1368	$\Delta cheW$ - <i>flhD</i>	(43)
MKM30	$\Delta fljI$	(9)
MMH10117	$\Delta fljH$ - <i>fljI</i> <i>flhB</i> (P28T)	(5)
Plasmids		
pET19b	Expression vector	Novagen
pKK211	pTrc99A/His-FljI(E211Q)	(19)
pKOT134	pET3b/FljD	(44)
pMM1702	pTrc99A/His-FljI	(36)
pMM1702(E211D)	pTrc99A/His-FljI(E211D)	This study
pMM140	pTrc99A/His-FljI(Δ 401–410)	This study
pMMHA1001	pGEX-6p-1/GST-FlhA _C	(14)
pMMHB1001	pGEX-6p-1/GST-FlhB _C	(14)
pMMJ1001	pGEX-6p-1/GST-FljI	(34)
pMMT002	pGEX-6p-1/GST-FljT94	(20)

Pull-down assays using GST affinity chromatography. Pull-down assays using GST affinity chromatography were performed as described previously³². Flow through fraction, wash fractions, and eluted fractions were analyzed by SDS-PAGE with Coomassie Brilliant blue (CBB) staining and immunoblotting with polyclonal anti-FljI antibody. At least three independent experiments were carried out.

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Author contributions

T.M. and K.N. conceived and designed research; T.M., Y.V.M., M.K. and P.D.A. performed research; T.M., Y.V.M., M.K. and P.D.A. analysed the data; and T.M. and K.N. wrote the paper based on discussion with other authors.

Additional information

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