Refined Mechanism of *Mycoplasma mobile* Gliding Based on Structure, ATPase Activity, and Sialic Acid Binding of Machinery

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**ABSTRACT** *Mycoplasma mobile*, a fish pathogen, glides on solid surfaces by repeated catch, pull, and release of sialylated oligosaccharides by a unique mechanism based on ATP energy. The gliding machinery is composed of huge surface proteins and an internal "jellyfish"-like structure. Here, we elucidated the detailed three-dimensional structures of the machinery by electron cryotomography. The internal "tentacle"-like structure hydrolyzed ATP, which was consistent with the fact that the paralogs of the \(\alpha^-\) and \(\beta^-\)-subunits of F\(_1\)-ATPase are at the tentacle structure. The electron microscopy suggested conformational changes of the tentacle structure depending on the presence of ATP analogs. The gliding machinery was isolated and showed that the binding activity to sialylated oligosaccharide was higher in the presence of ADP than in the presence of ATP. Based on these results, we proposed a model to explain the mechanism of *M. mobile* gliding.

**IMPORTANCE** The genus *Mycoplasma* is made up of the smallest parasitic and sometimes commensal bacteria; *Mycoplasma pneumoniae*, which causes human "walking pneumonia," is representative. More than ten *Mycoplasma* species glide on host tissues by novel mechanisms, always in the direction of the distal side of the machinery. *Mycoplasma mobile*, the fastest species in the genus, catches, pulls, and releases sialylated oligosaccharides (SOs), the carbohydrate molecules also targeted by influenza viruses, by means of a specific receptor and using ATP hydrolysis for energy. Here, the architecture of the gliding machinery was visualized three dimensionally by electron cryotomography (ECT), and changes in the structure and binding activity coupled to ATP hydrolysis were discovered. Based on the results, a refined mechanism was proposed for this unique motility.

**KEYWORDS** electron microscopy, electron cryotomography, F-type ATPase/synthase, image averaging, sialylated oligosaccharide

*Mycoplasmas* are parasitic and occasionally commensal bacteria that lack peptidoglycan layers and have small genomes (1, 2). *Mycoplasma mobile*, a fish pathogen, has a membrane protrusion and a gliding machinery at one pole and glides in the direction of the protrusion (Fig. 1A) (3–6). The average speed is 2.0 to 4.5 \(\mu\)m/s, or 3 to 7 times the cell length/s, with a propulsive force up to 113 pN (see Movie S1 in the supplemental material) (7–10). This motility, combined with the ability to adhere to the host cell surface, likely plays a role in infection, as has been suggested for another species, *Mycoplasma pneumoniae* (4, 11, 12). The motor proteins involved in this

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motility are unlike the motor proteins involved in any other form of bacterial or eukaryotic cell motility (3–6, 13, 14).

The cell surface can be divided into three parts beginning at the front end, i.e., the head, neck, and body (Fig. 1B) (3, 5, 6, 15–17). Three large proteins, Gli123, Gli349, and Gli521, with respective masses of 123, 349, and 521 kDa, are involved in this gliding mechanism and are localized exclusively at the cell neck (7, 15, 16, 18–21). Approximately fifty-nanometer legs corresponding to Gli349 can be seen protruding from the neck surface by electron microscopy (EM) (22, 23). The internal structure of the machinery is a “jellyfish”-like structure composed of 10 jellyfish-structure proteins (JSPs) (22, 24, 25). Interestingly, the amino acid sequences of MMOB1660 and MMOB1670, belonging to JSPs show high similarity to the respective α- and β-subunits of F1-ATPase, the catalytic subunit of proton pumps. However, these proteins are unlikely to function in an F-type proton pump, because F-type proton pumps require additional subunits (26) and the M. mobile genome has another locus containing a complete set of pump subunits. The energy for motility should be supplied through ATP, although gliding can be driven also by GTP and dATP (27–30), and the direct binding targets for gliding are the sialylated oligosaccharides (SOs) found on the surface of animal tissue (31–36). On the basis of the above information, we proposed a working model called the “centipede” or “power stroke” model, in which the cells are propelled by a “foot,” a part of Gli349 that, through repeated cycles driven by the hydrolysis of ATP, catches and releases SOs (3, 5, 7, 37). However, the force generation by ATP hydrolysis in the cell body and the subsequent transmission to the cell surface are still unclear. Elucidation of the detailed three-dimensional composition of the gliding machinery will be crucial in order to fully clarify the gliding mechanism.

In this study, we analyzed the structure of the gliding machinery by electron cryotomography (ECT) and studied the reactions involved in ATP hydrolysis; based on our findings, we proposed a mechanism for M. mobile gliding.

RESULTS

Detailed structures of the gliding machinery observed by ECT. To visualize the gliding machinery three dimensionally, intact cells (Fig. 2A; see also Movie S2 in the supplemental material), permeabilized cells (Fig. 2B and C; Movie S3), and the internal jellyfish-like structures (Fig. 2D; Movie S4) were analyzed by ECT. Cultured cells were...
collected, suspended in a growth medium, put on a grid, blotted with paper filters, frozen quickly, and observed by ECT. The contrast of the internal structures in an intact cell was low, because the thick and dense cell bodies inhibited transmission of the electron beams (Fig. 2A). To solve this problem, we prepared permeabilized cells using 0.7% Tween 20 (Fig. 2B and C) and then succeeded in observing filamentous structures on the cell surface, with a length ranging from 60 to 100 nm and featured with a thick end. (Fig. 2E, left, and Fig. S1A). Two types of filamentous structures were also observed as undercoats beneath the cell membranes of the head and neck regions (Fig. 2C, middle, and Fig. 2E, middle and right). Moreover, a novel structure was found to bridge the filament to the cell membrane for a distance of 12.0 ± 1.3 nm (n = 144 positions in 16 cells) (Fig. 2E, right, and Fig. S1B). Although the jellyfish-like structure was observed in the permeabilized cells, the contrast of the jellyfish-like structure was not sufficient for further analyses (Fig. 2C). Therefore, we removed the cell membrane by treatment with 0.08% Triton X-100, resulting in a clearer image of the jellyfish-like structure
On the basis of these observations, we concluded that each cell has 28 tentacle structures (Fig. S1Bii), each 110 nm long (Fig. S1Biii) and that the tentacle structures contain globular particles with a clear periodicity of 32 nm (Fig. S1Biv). A three-dimensionally rendered image of a permeabilized cell is presented in Fig. 2G and Movie S5. The undercoats and the tentacle structures were analyzed for periodicity by image averaging and its Fourier transform (Fig. 2H). Common periodicities were observed in the Fourier transforms of the neck undercoat and the tentacle structure but not in the front undercoat, suggesting that the first two are derived from a common structure. The tentacle image was obtained as an average of 199 images of a 23.6-nm-thick slice, featured with 31-nm repeating units, including a particle 13-nm wide and 21-nm long connected by axial stalks 5 nm in diameter and featured with lateral protrusions at two positions (Fig. 2H). The neck undercoat image was obtained as an average of 61 images featured with possible projections to the cell membrane. The structures and their dimensions are schematically summarized in Fig. 1B and C.

The jellyfish-like structure hydrolyzes ATP with conformational changes. As the jellyfish-like structure contains the paralogs of F1-ATPase catalytic subunits, we assayed the ATPase activity of the isolated jellyfish-like structure at room temperature (RT) and found that the structure hydrolyzes ATP with a $K_m$ of 66 μM (Fig. 3A). If this activity is derived from the β-subunit paralog, MMOB1670, the maximum turnover rate is calculated to be 0.09 molecules/s, based on the estimation of protein amount from SDS-PAGE analysis. Considering that the amino acid sequences suggested that the other protein components of the jellyfish-like structure do not exhibit ATPase activity (22), MMOB1670 should hydrolyze ATP. This ATPase activity was inhibited by the addition of

![FIG 3](image-url)  
Reactions of the jellyfish-like structure with ATP. (A) ATPase activity of the isolated jellyfish-like structure and speed of the gliding heads under various ATP and sodium azide concentrations. The ATPase activities under 0, 15.4, and 154 mM sodium azide are indicated by blue, orange, and red filled circles, respectively, and the gliding speeds without sodium azide are indicated by solid squares. These data were fitted as solid and broken lines for ATPase activities and gliding speeds by Michaelis-Menten kinetics with $K_m$ values of 66 μM and 60 μM and $V_{max}$ values of 7.6 nM/s and 1.4 μm/s, respectively. The concentration of ATPase β-subunit paralogs was estimated as 82 nM, and the maximum turnover rate was estimated as 0.09/s per β-subunit paralog. (B) Negatively stained EM image of the jellyfish-like structure in the presence of ATP. The bell structure is marked by a white cross. (C) Averaged images (left) and graph of interval lengths (right) of the tentacle structures. The interval lengths of tentacle structures, shown by a broken line, in the presence of various ATP analogs are presented in the graph. The measurement process is shown in Fig. 52.
sodium azide. The $K_m$ and the maximum turnover rate were calculated to be 84 and 76 $\mu$M and 0.063 and 0.033 molecules/s under 15.4 and 154 mM sodium azide, respectively.

Next, the possibility of conformational changes in the jellyfish-like structure was examined by averaging the EM images in the presence of various ATP analogs (Fig. 3B and C and S2). Two particles on a linear tentacle structure in the presence of each of AMPPNP, ADP-Vi, ADP-AlFx and ADP and in the absence of nucleotide (none) were selected and integrated into nine classes from 108, 101, 109, 102, and 109 images, respectively. The distances between the minimum peaks of particles in the density profiles were measured and averaged for the nine classes in each nucleotide state. The interval of the particles was approximately 2 nm longer under the conditions that included AMPPNP, ADP-Vi, and ADP-AlFx than the interval in the presence of ADP and the absence of nucleotides. The significance of these differences was revealed by Student's $t$ test ($P < 0.05$). These results show that the structural change occurs at the tentacle structure, depending on the ATP hydrolyzing forms.

**Preparation of the gliding head.** To examine coupling between SO binding and ATP hydrolysis, we intended to isolate the gliding machinery. In a previous study, *M. mobile* cells were elongated by treatment with detergents and their gliding was analyzed (8). In the present study, we found that in the elongated cells, the small portion of the back end of the cell (i.e., the end opposite the gliding direction) that begins to pull away from the main body of the cell rarely fully detaches and glides away (Fig. S3A), a result which has also been observed in *M. pneumoniae* and may arise from the same mechanism (38). Basically, an *M. mobile* cell usually has only one gliding organelle at a pole, but during the preparation for cell division, it sometimes has two gliding machineries at both poles (39, 40). The gliding force directing both sides occasionally generates tension in the elongated cell, resulting in the detachment of one machinery from the other part. Since the pole in the gliding direction is named the “head,” we call the detached machinery the “gliding head.” The above observation showed that the small part of the cell can glide without the other parts, suggesting that it might be possible to artificially isolate the gliding machinery. To investigate this possibility, we elongated the cultured cells by treatment with 0.1% Tween 60, sonicated and subjected them to sucrose-gradient centrifugation, and then isolated the fraction containing the gliding heads (Fig. 4A and Movie S6). The gliding heads were reactivated for gliding by placing them on an SO-coated glass slide and adding ATP, as described previously for a permeabilized cell model called the “gliding ghost” (27–29). Immunofluorescence microscopy, Western blotting, and peptide mass fingerprinting (PMF) demonstrated that the gliding heads retained Gill21, Gill39, Gill123, MMOB1650, -1670, -4530, -1660, and -1620, which should be involved in the gliding machinery (Fig. 4B, S3B, and S4A). The negatively stained EM image showed that most of the cell pieces in the fraction contained the gliding machinery, based on the observation of the internal jellyfish-like structure. The machinery was observed more easily in the gliding head than in the intact cells due to the removal of the cytosol, which improved the image contrast compared to that of the intact cells. However, the contrast was not better than that of the permeabilized cells, because the structures of the gliding heads were still densely packed.

**Speed and glass binding of the gliding head.** We next examined the movements of gliding heads under various concentrations of ATP (Fig. 3A). The gliding speed increased with the ATP concentration, adhering simply to the Michaelis-Menten kinetics with a 1.4 $\mu$m/s maximum speed and 60 $\mu$M $K_m$, the latter of which was similar to the value for ATPase activity of the jellyfish-like structure (66 $\mu$M). The gliding speed did not show an obvious sigmoidal format, suggesting that the leg does not catch SOs without nucleotides. Next, the effects of sodium azide on the gliding heads were examined (Fig. 4D). Both the number on glass and speed of gliding head decreased and reached to 4.5% and 49.6% of the original, respectively, at 50 s after the addition of 154 mM sodium azide.
sodium azide, suggesting that the activities of gliding heads were inhibited by sodium azide.

To clarify the coupling of ATP hydrolysis and displacements, we examined the effects of ATP analogs on the SO binding of the gliding head. To distinguish the SO binding involved in gliding from nonspecific binding, we examined the binding of the...
gliding head to solid surfaces coated by SOs under various conditions (Fig. S4B). These results showed that the glass binding of the gliding heads occurred in a manner similar to the binding of gliding cells, suggesting that the catch of SOs by feet in different ATP forms can be monitored through the binding of gliding heads to the glass. Then, the gliding heads were mixed with various concentrations of ATP, ADP, ADP-Vi, or ADP-AlFx, inserted into a tunnel of glass slides, and traced to determine the number of bound gliding heads (Fig. 4E). The numbers of bound gliding heads at 10 min after the insertion are presented as the ratio to that in 1 mM ATP. The presences of ADP-Vi and ADP-AlFx decreased the binding activity to 35% and 50%, respectively, but ADP increased it to 220% of that without ADP at 3 mM ADP. These results show that the leg catches SOs in the ADP form, and releases them in the ATP and ADP+P_i forms.

**DISCUSSION**

**Gliding machinery.** Although several structures involved in the gliding machinery have been visualized by conventional EM methods (22, 23, 41, 42), the positional relationship among the surface structure, the cell membrane, and the jellyfish-like structure have remained unclear. The present study using ECT was able to resolve this issue, because the ECT analysis allowed three-dimensional visualization of the cells under natural conditions (Fig. 2) (43). Many flexible filaments were observed on the surface of the gliding machinery, with a length ranging from 60 to 100 nm and featured with a thick end. These are consistent with the characteristics reported for isolated Gli349 molecules, which are 97 nm long and shaped like eighth-notes in music, each with a terminating globule that is designated a foot (42). It is likely that the legs observed in the present study were composed of Gli349 molecules. These molecules on the cell surface appeared flexible and did not show obvious alignment with the direction of the cell. The slice images from ECT and their average showed that the tentacle structures are positioned beneath the cell membrane of the neck part at a 12-nm distance and are bridged to the cell membrane through the two novel bridges (Fig. 2H, arrowheads). Probably, these novel bridges hold the internal jellyfish-like structure near the membrane and may play roles to transmit the force generated in the internal structure to the cell surface. The tentacle structures observed in negatively stained EM images tend to appear as bundles (Fig. 3B), suggesting that they are bundled laterally by an unknown structure. The lateral protrusions found in the averaged image of tentacles shown by triangles in Fig. 2H may be responsible for this bundling. A novel periodical undercoat was found at the head part, suggesting that this undercoat connects the “bell,” the front part of the jellyfish-like structure, to the front end of the cell membrane.

The three-dimensional image of a whole cell revealed the precise number of tentacle structures, the number of particles per tentacle structure, the lengths and diameters of the tentacle structures, the size of the particles, and the interval between the particles. In a previous study, the diameter and interval of the particles on the tentacle structures were reported to be 20 and 30 nm, respectively, in good agreement with the present study (22). The number of particles was estimated to be 113 to 225 per cell in the previous study (22), although we estimated it to be approximately 480 based on the results that the individual cells had 28 tentacle structures containing 17 particles in the present study. This difference may have been due to damage to the tentacle structures during the process of the negative staining in the previous study. The number of Gli349 molecules has been estimated as 450 per cell, based on titration using a monoclonal antibody in a previous study (19), in good agreement with the number of particles estimated in the present study. Therefore, a single leg may be driven by a single particle on the tentacle structure.

**Updating possible gliding mechanism.** Previously, we proposed a working model to explain the gliding mechanism, including a repetitive cycle consisting of catch, pull, and release of SOs by the foot composed of Gli349, based on previous data as listed below (3, 5, 37). (a) On the machinery surface, large proteins Gli349 and Gli521 work as leg and crank, respectively (41, 42). (b) The foot, the distal domain of Gli349, has
binding activity to SOs (31). (c) Gliding is driven by strokes 70 nm long, 1.5 pN in force, and the direction is slightly tilted from the cell axis (7, 27, 28, 32). (d) The propelling force is used also to detach the foot from SOs after a stroke, and the foot generates drag before detaching (15, 18, 34). (e) The foot can be detached to promote forward direction with 1.8 times less force than for a backward direction (44). (f) ATP is required to detach the foot from SOs, as shown by the gliding ghost experiments (29). (g) The foot catches SOs after thermal fluctuation with some cooperativity (32–34).

In the present study, the experimental data showed structural and functional changes of gliding machinery coupled to ATP hydrolysis. First, the intervals between the particles on the tentacles of jellyfish-like structure were approximately 5% longer in the presence of ATP-related “analogs,” including AMPPNP, ADP$+\text{AlF}_4^-$, and ADP$+\text{V}_4^-$, than those in ADP and no nucleotide (Fig. 3C). Second, the binding activity of gliding head, probably occurring on the foot of Gli349 to SOs, was accelerated by ADP rather than ATP-related analogs. This observation is consistent with the results of previous studies that gliding cells stayed on the glass when they were permeabilized by Triton (29). Probably, the gliding motors stayed in the ADP form under this condition. These changes in binding activity are likely linked to the structural changes found in the jellyfish-like structure. Actually, we found novel bridges linking the jellyfish-like structure to the cell membrane (Fig. 2E, right, and Fig. 2H, arrowheads).

Here, we update our model, based on the new information obtained in the present study (Fig. 5). (i) The unit in the ADP$+\text{P}_i$ form catches SOs at a front position after thermal fluctuation. (ii) The force applied to the leg from the front triggers the release of $\text{P}_i$, resulting in tight binding and force generation in the tentacle structure. The generated force transmits to the foot through several different proteins, stabilizes binding, and pulls the cell forward (stroke) to a direction slightly tilted from the cell axis.
(iii) Cell movements occurring as a result of other legs pull the units forward and set the parts other than the leg back to the initial conformation. The conformational change transmitted to the motor releases ADP. (iv) A new ATP molecule comes to the motor and allows the foot to release the SOs. The foot can be removed preferentially in the forward direction, and this directed binding property causes directed movement with a directed stroke. Then, the detached foot rebinds to SOs at the proper position after conversion from the ATP to ADP + Pi forms and some thermal fluctuation.

**Perspectives.** The model discussed here is consistent with all data obtained so far, but we need more information to discuss this in a concrete way. First, we need better resolution images including the jellyfish-like structure itself as well as the structures around the cell membrane. We do not even know the assembly of three large proteins on the surface, although the individual structural outlines have been visualized (41, 42, 45, 46). Second, the conformational changes occurring with ATP hydrolysis should be traced, including cooperativity in the machinery units.

Here, we found ATPase activity in the fraction of jellyfish-like structures. This activity should be linked to the gliding motility, based on the following observations. (a) The affinities to ATP estimated from Michaelis-Menten kinetics were in similar ranges for the ATPase activity of the jellyfish-like structure and the gliding speed of the “head” (Fig. 3A). (b) The ATPase activity was also inhibited by sodium azide, even with low affinity, ~100 mM dissociation constant ($K_d$), as observed in the gliding head, because azide is known to react specifically to the catalytic subunits of the F$_1$-ATPase $\beta$-subunit and its related proteins featured by a P-loop or a “Walker A” motif (47). These characteristics were comparable to the inhibitory effects on binding and the gliding speed of the gliding head (Fig. 4D). (c) The jellyfish-like structure changes the conformation depending on the bound ATP analogs (Fig. 3C). (d) Among the 21 proteins identified in the gliding head, only the F$_1$-ATPase paralogs of the jellyfish-like structure can be suggested for ATPase from the amino acid sequence (see Fig. S4Ai in the supplemental material).

If the ATPase activity is derived from the $\beta$-subunit paralog, its maximum activity can be calculated as 0.09 turnovers/s. This number can provide the gliding speed of 4.5 $\mu$m per s if we assume that mycoplasma cells can glide with a stroke of 70 nm coupled with the ATP hydrolysis, occurring in one of the 450 legs (19). Alternatively, the ATPase activity may be stimulated when the legs catch SOs in the gliding mechanism. If the catalytic subunits of the F$_1$-ATPase paralog play a central role in the gliding mechanism, this fact may suggest that the gliding mechanism evolutionally originated from the combination between F$_1$-ATPase and adhesin. It also suggests a novel evolutionary destination for F$_1$-ATPase (3, 5, 25).

**MATERIALS AND METHODS**

**Strains and culture conditions.** A mutant strain of *M. mobile*, P476R gli521, which binds to SOs more tightly than the wild-type strain, was used in this study (7, 14, 18, 29). Cells were cultured as previously described (21, 48). The nongliding mutant strains, m13 and m23 mutated for the gli349 gene, were reported previously (7, 15, 18, 21).

**Optical microscopy.** Mycoplasma cells and cell fractions were observed by phase-contrast microscopy using an IX71 microscope (Olympus, Tokyo, Japan). The images and movements were recorded with a Wat-120N charge-coupled-device (CCD) camera (Watex, Yamagata, Japan). All video data were analyzed by using ImageJ software, version 1.7 (http://rsb.info.nih.gov/ij/). Fluorescence microscopy was performed for the signals by enhanced yellow fluorescent protein (EYFP) fusion and immunofluorescence, as previously described (8, 24).

**EM.** For ECT of the permeabilized cells and the jellyfish-like structures, cultured cells were collected by centrifugation at 12,000 x g for 4 min at RT and suspended in a fresh growth medium at a 1,500-fold-higher concentration. A cell suspension of 1.5 $\mu$l was placed on an EM grid for 10 min, and 1.5 $\mu$l detergent solution (1% Tween 20 or 0.16% Triton X-100, 1 mg/ml DNase, and 5 mM MgCl$_2$ in phosphate-buffered saline (PBS) consisting of 75 mM sodium phosphate (pH 7.3) and 68 mM NaCl) was added. In the case of the gliding heads and intact cells, 2.6-$\mu$l suspensions derived from 150- and 15-ml cultures, respectively, were applied to the EM grids. The images were observed and captured at 0.570 nm per pixel by a Titan Krios FEG transmission electron microscope (FEI, Eindhoven, Netherlands) operated at 300 kV on an EM Falcon 4 k by 4 k direct electron detector (FEI). Single-axis tilt series were collected covering an angular range from $-70^\circ$ to $+70^\circ$ with a nonlinear Saxton tilt scheme at 5- to 10-$\mu$m underfocus using the Xplor 3D software package (FEI). The other experimental procedures were
performed as previously described (49, 50). Images were generally binned 2-fold, and three-dimensional (3D) reconstructions were calculated using the IMOD software package (51). Fourier transform was calculated by Imagej 1.37v (http://rsb.info.nih.gov/ij/). For negatively stained EM, the jellyfish-like structures were prepared on the grid as previously described (22). To examine the effect of ATP analogs, the Triton solution was removed after washing by buffer A (2 mM MgCl₂, 20 mM Tris-HCl at pH 7.5), and the jellyfish-like structure was treated with buffer A containing an ATP analog at 2 mM ADP, AMPPNP, ADP-V₅, or ADP-AIF₅. The grid was stained for 1 min by 2% ammonium molybdate (wt/vol) and air dried (22, 52). The specimens were observed by a transmission electron microscope, JEM-1010 (JEOL, Tokyo, Japan) at 80 kV, equipped with a FastScan-F214 (T) CCD camera (TVIPS, Gauting, Germany). Image averaging was done by EMAN, version 1.9 for both EM methods. Surface-rendering images were obtained using the three-dimensional modeling software Amira 5.2.2 (Visage Imaging, San Diego, CA).

To analyze the change in the periodicity of the chain structure, the negatively stained images were captured at 0.744 nm per pixel (see Fig. S2 in the supplemental material). More than 100 images 45 pixels wide and 216 pixels long including a tentacle were manually selected for each ATP analog. The obtained images were classified and averaged into nine classes. The averaged images were profiled for a rectangle 18 pixels wide and 83 pixels long. The distances between two negative peaks corresponding to the centers of particles were measured for all classes and averaged.

**ATP**ase assay. Mycoplasma cells were collected by centrifugation at 12,000 × g for 10 min at RT, washed twice with PBS-G (PBS containing 10 mM glucose), and suspended in PBS-G to be 50-fold concentrated from the culture. The cells were treated with Triton solution (0.3% Triton X-100, 1 mg/ml DNase, and 5 mM MgCl₂ in PBS) for 1 min at RT. The Triton-insoluble fraction was collected by centrifugation at 20,000 × g for 20 min at 4°C and suspended in buffer A. This process was repeated.

ATPase was assayed by a continuous spectrophotometric method using a 2-amino-6-mercapto-7-methylpurine ribonucleoside–purine nucleoside phosphorylase reaction to detect the released inorganic phosphate (EnzChek kit; Life Technologies, Carlsbad, CA, USA) (53). The reaction mixture was as follows: 20 μg/ml jellyfish-like structure, 2 mM MgCl₂, 20 mM Tris-HCl (pH 7.5) in a total volume of 0.2 ml at RT. Sodium azide was added at a concentration of 15.4 or 154 mM when stated. The protein amount of the F₁-ATPase β-subunit paralog was estimated from the densitometry of SDS-PAGE.

**Gliding head.** Mycoplasma cells were collected by centrifugation, washed with PBS-G, and suspended in PBS-G containing 0.1% Tween 60 to be 15-fold concentrated from the culture. After 30 min, the cells were applied to a sonicator (UR-20P; Tomy Seiko, Tokyo, Japan) for 1 s, 60 times on ice. The sonicated fractions were treated with 1 mg/ml DNase, 1 mg/ml RNase, and 10 mM MgCl₂ and subjected to a stepwise gradient centrifugation consisting of 0%, 20%, 30%, 40%, 50%, and 60% sucrose layers. After centrifugation at 18,000 × g for 15 min at 4°C, the fraction of gliding heads was recovered from the top of the 30% sucrose layer and washed with buffer A. The gliding head suspension was inserted into a tunnel at time zero, and the bound heads within a 32- by 43-μm² field were traced under the microscope. The tunnel was assembled using two pieces of glass, precoated with 10 mg/ml fetuin in buffer A for 10 s, and then washed with 50 μl buffer A (8). The numbers of heads in the three fields were averaged for each time point and presented as a ratio to the number of gliding heads in the presence of 1 mM ATP at 10 min, with standard deviations. The cell behaviors were analyzed as previously reported (8, 54). SDS-PAGE, PMF, and Western blotting were performed as previously described (22). The monoclonal antibodies used were also described previously (16, 20, 54).

**Data availability.** The cryoEM data have been deposited in the Electron Microscopy Data Bank under accession code EMD-0893.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.02846-19.

**MOVIE S1,** AVI file, 3.9 MB.
**MOVIE S2,** MPG file, 9.5 MB.
**MOVIE S3,** MPG file, 9.6 MB.
**MOVIE S4,** MPG file, 5.1 MB.
**MOVIE S5,** MPG file, 7.8 MB.
**MOVIE S6,** AVI file, 3.1 MB.
**FIG S1,** PDF file, 0.8 MB.
**FIG S2,** PDF file, 1.7 MB.
**FIG S3,** PDF file, 0.6 MB.
**FIG S4,** PDF file, 0.6 MB.

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