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Visualization of functional rotor proteins of the bacterial flagellar motor in the cell membrane

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Summary

The bacterial flagellar motor is a rotary motor driven by the electrochemical potentials of specific ions across the cell membrane. Direct interactions between the rotor protein FliG and the stator protein MotA are thought to generate the rotational torque. In this study, we used total internal reflection fluorescent microscopy to observe the localization of GFP-fused FliG in *Escherichia coli* cells. We identified three types of fluorescent punctate signals: immobile dots, mobile dots that exhibited simple diffusion, and mobile dots that exhibited restricted diffusion. When GFP-FliG was expressed in a $\Delta fliG$ background, most of the cells were not mobile. When the cells were tethered to a glass side, however, rotating cells were commonly observed and a single fluorescent dot was always observed at the rotational center of the tethered cell. These fluorescent dots were likely positions at which functional GFP-FliG had been incorporated into a flagellar motor. Our results suggest that flagellar basal bodies diffuse in the cytoplasmic membrane until the axial structure and/or other structures assemble.

Keywords: bacterial flagella; FliG; stator; GFP

Introduction

Many bacteria sense external signals from the environment and can swim toward favorable conditions by rotating locomotive organelles called flagella. The bacterial flagellum consists of a helical filament that acts as a propeller, a basal body that acts as a rotary motor embedded in the cytoplasmic membrane, and a hook that acts as a universal joint connecting the filament to the basal body.¹ The rotary motor of a bacterial flagellum is driven by the flux of specific ions across the cell membrane; *Escherichia coli* and *Salmonella enterica* serovar Typhimurium use H⁺, whereas marine *Vibrio* species and alkalophilic *Bacillus* species use Na^{+,2-4} In *Bacillus subtilis*, distinct stators for coupling either Na⁺ or H⁺ flux to force generation are thought to function in the same bacterial cell.⁵

The rotor in the flagellar basal body consists of an axial rod and two rings: the MS ring and the C ring. The MS ring is embedded in the cytoplasmic membrane, whereas the C ring is attached to the cytoplasmic surface of the MS ring. In Gram-negative bacteria, two additional rings, the L and P rings, surround the axial rod and are attached to the outer membrane and the peptidoglycan layer, respectively.⁶ These rings are thought to act as bearings for the axial rod. During flagellar assembly, FliF first multimerizes to form the MS ring in the cytoplasmic membrane.⁷ Next, FliG, FliM, and FliN assemble to form the C ring on the cytoplasmic surface of the MS ring.⁶ The axial proteins are then exported to the inside of the C ring to form the rod, hook, and flagellar filament.⁸

The stator of the flagellar motor consists of PomA and PomB in the Na⁺-driven motors of *Vibrio alginolyticus*, or MotA and MotB in the H⁺-driven motors of *E. coli*.⁹⁻¹¹ These stator complexes are thought to function as specific ion channels.^{12,13} The A and B stator subunits are incorporated into the motor as

a stator complex after completion of the hook-basal body; deletions in the Cterminal domain of the B subunit have been shown to block this incorporation process.^{14,15} FliG, a cytoplasmic protein that is thought to be directly involved in force generation by the flagellar motor, forms the switch complex with FliM and FliN that changes the direction of motor rotation between clockwise (CW) and counterclockwise (CCW) in response to chemotactic signals.¹⁶⁻¹⁹ Mutational analyses of the H⁺-driven flagellar motor have identified charged residues in the C-terminal region of FliG and the cytoplasmic domain of MotA that are important for force generation.^{20,21} In the crystal structure of the middle and Cterminal domains of *Thermotoga maritima* FliG, the corresponding charged residues are aligned in a C-terminal ridge that is exposed on the surface of the FliG molecule.²² It has been predicted that the electrostatic interactions between the charged residues of FliG and the cytoplasmic regions of MotA cause motor rotation.²³ The precise mechanism underlying motor rotation, however, is not fully understood.

FliG molecules of the flagellar rotor have been visualized at the base of the polar flagellum in *V. alginolyticus* using fusion proteins carrying green fluorescent protein (GFP).²⁴ Moreover, FliG molecules tagged with yellow fluorescent protein were detected as several fluorescent foci on the surface of the cell body of *S. typhimurium*, a bacterium that has peritrichous flagella.²⁵ In the present work, we constructed functional fusion proteins of GFP and the *E. coli* rotor protein FliG and observed the rotating motor using total internal reflection (TIR) fluorescent microscopy.

Results

Motility of cells producing GFP-FliG

The gfp gene was genetically fused to the 5' end of the fliG gene to form the *gfp-fliG* construct, the product of which was GFP-FliG. Wild-type *fliG* and *gfp-fliG* were cloned and placed under the control of arabinose-inducible promoters in the pYF2 and pYF1 plasmids, respectively. We investigated the motility of the cells producing GFP-FliG or wild-type FliG as a rotor component in semi-solid agar containing arabinose. In a medium containing 0.0005% arabinose, $\Delta fliG$ cells (DFB225) producing wild-type FliG from pYF2 showed the same degree of swarming as RP437 cells, a strain with wild-type motility and chemotaxis (Fig. 1a). $\Delta fliG$ cells producing wild-type FliG needed at least 0.0002% arabinose in the medium to achieve the same degree of motility as RP437 cells (data not shown), and the cells overproduced the FliG protein when they were placed in the medium containing 0.0002% arabinose (Fig. 1a). On the other hand, the $\Delta fliG$ cells producing GFP-FliG from pYF1 did not swarm in the semi-solid agar containing 0.0005% arabinose (Fig. 1a). Under a light microscope, however, $\Delta fliG$ cells producing GFP-FliG were observed to swim, although their swimming ability was much lower than that of wild-type cells (Table 1). Similar amounts of plasmid-derived wild-type FliG and GFP-FliG proteins were detected by immunoblot analysis of whole-cell extracts from the $\Delta fliG$ cells (Fig. 1b). On the other hand, the cells producing GFP-FliG showed a reduced level of flagellin expression (Fig. 2); the intensity of the band of flagellin derived from these cells was approximately ten times less than the intensity of the band derived from the cells producing wild-type FliG. In addition, the number and the length of the flagella observed with electron microscopy were greatly reduced in the cells producing GFP-FliG (data not shown). Thus, the fusion of the GFP molecule to FliG probably affected flagellar assembly.

Next, to investigate the function of individual motors, the cells were tethered to the surface of a coverslip using the FliC sticky filament, which has been shown to readily attach to glass surfaces.^{26,27} FliC sticky proteins were

produced from a second plasmid, pYS11.²⁷ The tethered cells robustly rotated on the coverslip in spite of their weak motility in liquid media. Immunoblot analysis of whole-cell extracts detected GFP-FliG at a position corresponding the expected molecular mass of 64 kDa (Fig. 1). Very weak bands were detected just under the GFP-FliG bands (Fig. 1b, open arrowhead); these bands may have resulted from a slight degradation or mistranslation of GFP-FliG. Degradation products with a molecular mass corresponding to that of wild-type FliG, however, were not detected, indicating that GFP-FliG was functional in individual flagellar motors. The rotational velocities of the tethered cells producing wild-type FliG and GFP-FliG were ca 6 and ca 3 revolutions/s, respectively (Table 2). For the wild-type FliG motor, the switching frequency and the percentage of time spent rotating CCW were 0.9 reversals/s and 65%, respectively. On the other hand, the switching frequency and the percentage of time the motor rotated CCW for the GFP-FliG motor were 0.4 reversals/s and more than 92%, respectively. Flagellar motors that incorporated GFP-FliG preferentially rotated CCW. Although the activity of the GFP-FliG motors was lower than that of the wild-type motors, the GFP-FliG motors were sufficiently functional for further analysis.

Subcellular localization of GFP-FliG

We observed the localization of GFP-FliG in the $\Delta fliG$ and $\Delta flhDC$ strains. Because the *flhDC* genes are master genes that control the expression of all of the flagellar genes, the $\Delta flhDC$ strain produces no flagellar components. Similar amounts of GFP-FliG protein were detected by immunoblot analysis of whole-cell extracts from the two cell lines (Fig. 1b, lanes 4 and 5). When cells at mid-log phase were immobilized on the surface of a polylysine-coated coverslip and observed using TIR fluorescent microscopy, we detected 5 ± 1 (mean \pm SD) fluorescent dots near the cell surface in each of the $\Delta fliG$ cells producing GFP-

FliG (Fig. 3; $\Delta fliG$, TIR). On the other hand, in the $\Delta flhDC$ cells producing GFP-FliG, no fluorescent dots were detected and a diffuse fluorescent signal was observed throughout the cell body (Fig. 3; $\Delta flhDC$, TIR). Using axial (brightfield) epi-illumination (epi-illumination), GFP-FliG was also detected throughout the cell bodies of the $\Delta flhDC$ cells (Fig. 3; $\Delta flhDC$, Epi). Therefore, the clustering of the GFP-FliG molecules was dependent on flagellar structures or one or more of the flagellar proteins. Under epi-illumination, we detected 11 \pm 3 (mean \pm SD) fluorescent dots in each $\Delta fliG$ cell though the background fluorescent signal was larger than that observed using TIR illumination (Fig. 3; $\Delta fliG$, Epi and TIR). The number of fluorescent dots in each cell was more than the reported number of flagella for an *E. coli* cell.^{28,30} This was probably due to flagellar basal bodies that lacked flagellar filaments.

Movement of GFP-FliG on the cytoplasmic membrane

In the $\Delta fliG$ cells producing GFP-FliG, we tracked the movement of the fluorescent dots in the cell membrane using TIR fluorescent microscopy. After the cells were immobilized on a polylysine-coated coverslip, we detected immobile and mobile fluorescent dots on the cell membrane (Fig. 4). In each cell, approximately two immobile and four mobile fluorescent dots were detected. The positional distributions of the centers of the immobile dots on the cell membrane were restricted to a very narrow area (Fig. 4d). On the other hand, as shown in Figures 4a and 4b, one fluorescent dot (dot 1) moved in the direction of the cell pole, whereas the movement of another fluorescent dot (dot 2) was restricted within a limited region for more than 12 s. We analyzed the movement of these mobile dots by plotting the mean squared displacement versus the time interval (MSD- Δt) and examining the relative deviation. The MSD- Δt plot for dot 1 appeared to fit a linear function, which is indicative particle undergoing simple diffusion (Fig. 4c). On the other hand, the MSD- Δt

plot of dot 2 is characteristic of a particle undergoing restricted diffusion (Fig. 4c). The relative deviation, RD (300, 90), of dot 1 and dot 2 were calculated to be 1.1 and 0.2, respectively. From a comparison with the theoretical RD (300, 90) reported by Kusumi et al.,²⁹ the trajectory of dot 1 was determined to be a result of simple diffusion, whereas the trajectory of dot 2 was determined to be a result of restricted diffusion. We analyzed the trajectories of 15 other fluorescent dots using RD (300, 90) or RD (200, 90). Three of these trajectories were determined to result from simple diffusion and the average RD value for these dots was 0.9. Eight of the trajectories were determined to result from restricted diffusion and the average RD value for these dots was 0.2. The diffusion modes of the four remaining dots were not determined, because there were not enough data points for analysis. Estimating the diffusion coefficients (D) of dot 1 and dot 2 using the slope of each of the MSD- Δt plots revealed that the diffusion coefficients of dot 1 and dot 2 were 5.6 x 10^{-11} cm²/s and 2.5 x 10^{-11} cm²/s, respectively. The average diffusion coefficient for all 17 of the mobile dots was $4.9 \ge 10^{-11} \pm 3.1 \ge 10^{-11} \text{ cm}^2/\text{s}.$

Observation of GFP-FliG in tethered cells

If the GFP-FliG protein retains its function and attaches to basal bodies as a component of the rotor, the rotational center of each tethered cell should be fluorescently labeled (Fig. 5a, central dot). To determine the position of the rotational center, we chose slowly rotating cells (approximately 1 revolution/s). For each of the GFP-FliG-producing cells, a single immobile fluorescent dot was observed at the midpoint of the cell body, which was the rotational center of the tethered cell (Fig. 5b, arrowhead C). In order to precisely determine the position of the central dot and the rotational center of the tethered cell, we determined the centers of the fluorescent signals from a central dot (arrowhead C) and a satellite dot (arrowhead S) (Fig. 5a and 5b) from more than 100 frames

(images were captured at 30 frames/s). As shown in the inset in Figure 5c, the center of the satellite dot followed a circular path with a radius of approximately 1.2 µm (shaded squares). The rotational center of the tethered cell was defined as the center of the ellipse calculated by least-squares fitting of the trajectory of the center of the satellite dot (Fig. 5c, large open square). In this cell, the standard deviations for the center of the fitted ellipse were \pm 12.8 nm along the X axis and \pm 19.2 nm along the Y axis. The positions of the center of the central dot were distributed within a range of approximately 80 nm along both the X and Y axes (Fig. 5c, shaded squares). The rotational center calculated from the path of the satellite dot was located within the area defined by the distribution of the positions of the central dot (Fig. 5c). Figure 5d shows the distributions of the positions of the central dots from 11 tethered cells. For each of these cells, the rotational center calculated from the movement of a satellite dot was located within the range defined by the distribution of the positions of the central dot. These results indicate that the GFP-FliG molecules localized at the rotational centers of the tethered cells.

Discussion

The bacterial flagellar motor is a rotary motor driven by the electrochemical potentials of ions. The interaction between the rotor and the stators is thought to generate torque, thereby converting the energy of ion flux into mechanical power. In the flagellar motor of *E. coli*, FliG is a rotor component that is thought to interact directly with the stator complex formed by MotA and MotB.²³ In the present study, we used the functional fusion protein GFP-FliG and TIR fluorescent microscopy to observe rotating motors and to characterize the motion of basal bodies in the cell membrane of *E. coli*.

To construct a flagellum, FliF first assembles to form the MS ring in

the cytoplasmic membrane. FliG then attaches to the cytoplasmic surface of the MS ring to form the C ring with FliM and FliN.^{6,7} In the $\Delta flhDC$ deletion strain, the MS ring is not formed because none of the flagellar proteins, including FliF, are produced.¹ Accordingly, no fluorescent dots were observed in GFP-FliG-producing $\Delta flhDC$ cells; a diffuse fluorescent signal was instead detected throughout the $\Delta flhDC$ cells (Fig. 3). In contrast, fluorescent dots were detected in GFP-FliG-producing $\Delta fliG$ cells, indicating that the formation of the fluorescent dots was dependent on the flagellar proteins. It can be concluded that the fluorescent dots corresponded to locations at which GFP-FliG bound to MS rings, and in some cases, assembled to form the rotor in the basal body of a flagellar motor.

Although we may not have detected all of the fluorescent dots due to the high background fluorescence, at least 11 fluorescent dots were observed in each cell. Interestingly, this number of dots is more than the reported number of flagella for an individual cell.^{28,30} The larger than expected number of fluorescent puncta was probably not due to the overproduction of GFP-FliG. In $\Delta fliG$ cells, the number of MS rings is probably the same as in RP437 cells because FliF is produced from the chromosome. This means that the number of fluorescent dots was limited by the number of MS rings. Thus, the maximum number of fluorescent dots would not exceed the number of MS rings even if GFP-FliG was overproduced. Therefore, *E. coli* cells must have complete flagellar structures and partial flagella that lack filaments, such as basal bodies that are not completely assembled and/or basal bodies that have lost the flagellar filaments. It has been reported that there are an average of 3.4 flagella in each *E. coli* cell.³⁰ This suggests that there are twice as many unfinished basal body structures as there are fully assembled basal bodies with intact flagella.

We observed mobile and immobile fluorescent dots in cells producing GFP-FliG. After the construction of the MS ring and the C ring, the export apparatus attached to the inside of the C ring transports the axial proteins

outside of the cytoplasmic membrane, where they assemble to form the rod, hook, and filament.⁸ Structures consisting of an MS ring and a C ring without the axial components can probably diffuse in the cell membrane, whereas basal bodies with axial structures that penetrate the peptidoglycan layer and/or the outer membrane are likely fixed in the cell membrane. The diffusion coefficient of the mobile fluorescent dots roughly estimated from the MSD- Δt plots was 4.9 x $10^{-11} \pm 3.1$ x 10^{-11} cm²/s. If each mobile fluorescent dot corresponded to a single flagellar basal body, the molecular size of the fluorescent dots was at least 3,200 kDa (26 molecules each of FliF, FliG, and GFP). In the cell membrane of *E. coli*, the diffusion coefficient (*D*) of a TatA-GFP fusion protein (9.6 kDa) was estimated to be $1.3 \times 10^{-9} \text{ cm}^2/\text{s}$ using fluorescence recovery after photobleaching.³¹ On the other hand, the *D* value of the MotA₄:MotB₂ complex, which is the stator of the bacterial flagellar motor, was estimated to be 8.8×10^{-1} ¹¹ cm²/s using single particle tracking.³² Because the D value is dependent on the Stokes radius (r) of the transmembrane domain in the lipid bilayer, it is not surprising that the D values of the complexes composed of GFP-FliG and MotA₄:MotB₂ are of the same order of magnitude (x 10^{-11} cm²/s). On the other hand, the observation that the D value of the GFP-FliG complex is smaller than that reported for the TatA fusion protein is reasonable, because TatA was predicted to have one transmembrane segment by the SOSUI program.

Some of the mobile fluorescent dots diffused within a restricted area (Fig. 4b and 4c, dot 2). Structures in the peptidoglycan layer and/or the outer membrane, such as L rings and P rings, might capture flagellar basal bodies that are diffusing in the cytoplasmic membrane. On the other hand, in eukaryotic cells, several elements have been proposed to restrict the movement of membrane proteins, including a membrane skeleton/cytoskeleton fence or transient confinement zones (TCZs).³³⁻³⁵ In bacteria, MreB forms actin-like filaments that run along the inner surface of the cytoplasmic membrane,³⁶ whereas FtsZ, a tubulin homolog that is required for cell division, forms a ring

structure at the site of division.³⁷ Membrane domains enclosed by cytoskeletallike structures or TCZ-like membrane domains might restrict the diffusion of flagellar basal bodies. This restriction might be involved in determining positions at which flagella form.

Only 3% of cells producing GFP-FliG were observed to swim, and the swimming speed of these cells was half of the speed observed for cells producing wild-type FliG from a plasmid. This phenotype may be attributable to a reduction in the number of flagella. The percentage of GFP-FliG-producing cells with flagella was low and the amount of flagellin in these cells was approximately ten times less than the amount of flagellin in the cells producing wild-type FliG. The *fliC* gene, which encodes flagellin, is a class III gene in the expression hierarchy of the flagellar genes. The expression of class III genes is repressed by the anti-sigma factor FlgM, and this repression is removed following the excretion of FlgM through the flagellar export apparatus.¹ The low expression level of flagellin suggests that anti-sigma factor FlgM was not efficiently pumped out of the GFP-FliG-producing cells; this would prevent the cells from downregulating the expression of the class II flagellar genes and increasing the expression of the class III genes. Our observations suggest that the activity of the export apparatus was reduced or the assembly of the export apparatus was inhibited by the GFP molecule fused to FliG.

When the GFP-FliG-producing cells were tethered to glass coverslips, they were able to rotate although the rotation speed was about half of that observed for cells producing wild-type FliG. A puncta of GFP-FliG was localized at the rotational center of each tethered cell (Fig. 5). This indicates that i) GFP-FliG was located at the site of the motor, ii) GFP-FliG assembled on an MS ring or in a flagellar basal body, and iii) the fusion protein functioned as a component of the rotor in the flagellar motor. Compared to cells with wild-type motors, the rotation of the motors was biased to the CCW direction and the switching frequency was low in the GFP-FliG-expressing cells. GFP is a large

protein (27 kDa) with a β -barrel structure.³⁸ Fusion of this protein to the N terminus of FliG may have affected the structure of FliG, the interaction between FliG and MotA, and/or the interaction between FliM and phosphorylated CheY, which switches the rotation of the motor from CCW to CW. Alternatively, the low level of expression of the class III genes or the late genes (the *che* and *mot* genes) might have affected the rotation of the motor.¹

In this study, we have successfully observed fluorescently labeled, rotating flagellar motors and the diffusion of flagellar basal bodies in the cytoplasmic membrane. We do not know the precise number of FliG molecules that are in an individual rotating motor. According to a rough estimate, the fluorescent intensity of one of the fluorescent dots appeared to be about ten times stronger than that of a single molecule of GFP (data not shown), suggesting that at least 10 molecules of GFP-FliG were present in each fluorescent dot. As shown in a recent report, ³² we may be able to determine the precise number of FliG molecules that are incorporated into a functional motor by measuring single-step photobleaching of GFP-FliG in a rotating cell. Additionally, development of this technique may allow us to examine the interaction between the flagellar motor components, such as the rotor and the stator, using fluorescence resonance energy transfer.

Materials and Methods

Bacterial strains, growth conditions, and media

The bacterial strains used in this study are listed in Table 3. *E. coli* cells were cultured in LB broth [1% (w/v) tryptone peptone, 0.5% (w/v) yeast extract, and 0.5% (w/v) NaCl] or TG broth [1% (w/v) tryptone peptone, 0.5% (w/v) NaCl, and 0.5% (w/v) glycerol] at 30°C. Chloramphenicol was added to a final

concentration of 25 μ g/ml and ampicillin was added to a final concentration of 50 μ g/ml.

Plasmids

The plasmids used in this study are listed in Table 3. The plasmids carrying the *fliG* gene under the control of the *araBAD* promoter were pBAD33-based plasmids. The SacI/HindIII fragment from pSL27 was inserted into the SacI/HindIII sites of pTY102, resulting in the pYF2 plasmid. A fragment encoding the *fliG* gene, which was amplified by a PCR using pYF2 as a template, the sense primer BsrGI-*fliG*, and an antisense primer that annealed to a region downstream of the multiple cloning site was inserted into the BsrGI/HindIII sites of pTY200; this plasmid was named pYF1.

Preparation of cells for observation with fluorescence microscopy

Overnight cultures of cells harboring the plasmids were inoculated into TG medium (1/100) in the presence of arabinose and cultured at 30°C for 4 h. The cells were harvested by centrifugation and suspended in motility medium: 10 mM potassium phosphate (pH 7.0), 0.1 mM EDTA, and 85 mM NaCl. Sixty μ l of the cell suspension was loaded into the space between a slide glass and a coverslip with a spacer. The slide was then inverted and incubated for 2 min. Another 60 μ l of motility medium was loaded into the space between the slide and the coverslip to remove the remaining unattached cells.

Microscopic system

We used an Olympus IX-71 microscope with a modified mirror turret to allow side entry of the excitation light into the microscope. A laser beam ($\lambda =$

488 nm) for excitation was focused at the back focal plane of the objective lens (Olympus PLAPON 60XOTIRFM, NA = 1.45). For objective TIR illumination,³⁹ the angle of the incident laser beam was adjusted with a mirror set in front of the focusing lens (f = 180) so that the laser beam would pass near the edge of the objective lens. In our experiments, the incidence angle was 71.4°, and thus the evanescent light reached a depth of approximately 70 nm (the value when the sample on the glass surface was water). The image was projected to an EM-CCD camera (DV860-BV, Andor Technology, South Windsor, CT) through a projection lens (5x); as a result, 1 pixel corresponded to 80 nm in the specimen plane. All images were captured at the normal video rate (30 frames/s). We confirmed the positional resolution of this microscope system was 5 nm in an experiment using a 100-nm fluorescent bead.

Determination of the rotational center of a tethered cell

used free software We the ImageJ (NIH Image. http://rsb.info.nih.gov/ij/) for analysis of all of the images. The position of the rotational center was defined as the center of a fitted ellipse calculated by the least squares method from the center of a fluorescent satellite dot. For the definition of the rotational center of a tethered cell, the positions of the center of the fluorescent satellite dot obtained from more than 100 video frames were used. The distribution of positions of the central dots was calculated from the centers of their fluorescent signals. To calculate the center of each dot, we analyzed video frames that contained more than 16 pixels that were brighter than the threshold intensity. We selected tethered cells rotating CCW for analysis.

Tracking the movement of fluorescent dots

To track the movement of the mobile fluorescent dots, $\Delta fliG$ cells

producing GFP-FliG were attached to polylysine coated-coverslips and observed with TIR fluorescent microscopy. Images were captured at 30 frames/s and the center of the fluorescent mobile dot was calculated in each video frame. For the trajectory of each fluorescent dot, the mean square displacement (MSD) was calculated according to the following formula:²⁹

$$MSD(n\delta t) = \frac{1}{N-1-n} \sum_{j=1}^{N-1-n} \{ [x(j\delta t+n\delta t) - x(j\delta t)]^2 + [y(j\delta t+n\delta t) - y(j\delta t)]^2 \},$$

time video frame (33 where the δt is ms) and $(x(j\delta t + n\delta t) - x(j\delta t), y(j\delta t + n\delta t) - y(j\delta t))$ describes the position of the dot following a time interval $\Delta t_n = n \delta t$ after starting at position $(x(j\delta t), y(j\delta t)); N$ is the total number of frames in the recording sequence, n and j are positive integers, and n determines the time increment. The diffusion coefficients (D_{2-4}) of the mobile dots were estimated by fitting the MSDs at $2\Delta t$, $3\Delta t$, and $4\Delta t$ using a straight line.²⁹ Mobile dots that were tracked for more than 200 frames were analyzed for the calculation of the diffusion coefficient. The relative deviation RD (N, n)was used for the determination of the diffusion mode of a mobile fluorescent dot.²⁹ The relative deviation is defined as

$$RD(N,n) = \frac{MSD(N,n)}{4D_{2-4}n\delta t}$$

where MSD(*N*, *n*) represents the MSD at the time interval $n\delta t$ from a sequence of *N* video frames. $4D_{2-4}$ is the expected average MSD value for fluorescent dots undergoing simple diffusion with a diffusion coefficient of D_{2-4} . The value of RD (300, 90) or RD (200, 90) was used for the determination of the diffusion mode of each mobile fluorescent dot.²⁹

Detection of GFP-FliG and flagellin in E. coli cells

E. coli cells harboring a plasmid encoding wild-type FliG or GFP-FliG were incubated at 30°C for 4 h in TG medium containing arabinose. The cells

were harvested by centrifugation and suspended in motility medium to an O.D.₆₆₀ of 10. The same volume of SDS loading buffer was added to the cell suspensions and the samples were boiled at 100°C for 5 min. The proteins were separated by SDS-PAGE and immunoblotting was performed using anti-FliG (a gift from D. Blair) and anti-GFP antibodies (GFP monoclonal antibodies, Clontech).

To detect flagellin, a 1/10 volume of trichloroacetic acid was added to the cell culture (O.D.₆₆₀ = 1) and the cells were harvested by centrifugation. The pellet was washed with acetone and dried. The pellet was suspended in a 1/10volume of SDS loading buffer to the original culture volume. The proteins were separated by SDS-PAGE and immunoblotting was performed using antiflagellin antibodies (laboratory stock).

Measurement of the swimming fraction and swimming speed

Cells were grown at 30°C for 4 h in TG medium containing arabinose. The cell suspensions were diluted 50-fold in fresh TG medium containing 20 mM serine, which was used to suppress CW rotation. Cell motility was observed under a dark-field microscope and recorded on videotape. Swimming speed was determined as described previously.⁴⁰ The average swimming speed was obtained by measuring at least 40 swimming tracks. The swimming fraction was measured from the same videotape.

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Figure legends

Figure 1. (a) Swarming behavior of cells producing GFP-FliG. pYF1 (encoding) *gfp-fliG*)-carrying DFB225 cells, pYF2 (encoding wild-type *fliG*)-carrying DFB225, pTY200 (encoding gfp)-carrying DFB225 cells, and pBAD33 (vector)carrying RP437 cells were grown in TG-0.27% (w/v) soft agar medium containing 0.0005% (w/v) arabinose at 30°C for 9 h. RP437, wild-type for chemotaxis; DFB225, $\Delta fliG$ strain. (b) Immunoblots for GFP-FliG or wild-type FliG in whole-cell extracts. Proteins were detected with anti-FliG antibodies or anti-GFP antibodies. Molecular mass values (kDa) are shown on the left side of the panels. The band between 47.5 kDa and 62 kDa is probably a non-specific band detected by the anti-GFP antibodies because the band was detected in all of the lanes. Lane 1, pBAD33-carrying RP437 cells grown in TG medium containing no arabinose; Lane 2, pYF2-carrying DFB225 cells grown in TG-0.0002% arabinose medium; Lane 3 and 4, pTY200-carrying and pYF1-carrying DFB225 cells grown in TG-0.0005% arabinose medium, respectively; Lane 5, pYF1-carrying RP3098 cells ($\Delta flhDC$) grown in TG-0.002% arabinose medium. Cells were grown at 30°C for 4 h.

Figure 2. Detection of flagellin. Proteins were separated and immunoblotting was performed using anti-flagellin antibodies. Lane 1, pBAD33-carrying RP437 cells grown in TG medium containing no arabinose; Lane 2, pYF2-carrying DFB225 cells grown in TG-0.0002% arabinose medium; Lane 3, pYF1-carrying DFB225 cells grown in TG-0.0005% arabinose medium; Lane 4 and Lane 5, pBAD33-carrying DFB225 cells and pTY200-carrying RP3098 cells grown in TG medium containing no arabinose, respectively.

Figure 3. Subcellular localization of GFP-FliG. Cells producing GFP-FliG were grown at 30°C for 4 h in medium containing 0.0005% arabinose ($\Delta fliG$) or

0.002% arabinose ($\Delta flhDC$), and were observed with a fluorescence microscope. TIR indicates the images obtained under TIR illumination, whereas Epi indicates images obtained under axial (brightfield) epi-illumination. BF indicates brightfield images. The adjacent fluorescent and brightfield images are images of the same cells. Bar, 1.6 µm.

Figure 4. Movement of the fluorescent dots. (a) A sequence of images (TIR illumination) of fluorescent dots formed by GFP-FliG. The figure shows a 2 s image sequence, and every 10th frame is shown. Images were captured at a normal video rate (30 frames/s). Bar, 1.6 µm. (b) Movement trajectories of fluorescent dot 1 and fluorescent dot 2 (arrowheads 1 and 2 in Fig. 4a, respectively). Trajectories of the centers of the fluorescent dots over 12 s are shown. Each trajectory color represents the movement of the dot during a period of 1 s, and the movements of the dots in first second through the sixth second are denoted in pink, blue, green, orange, turquoise, and red, respectively; this order of the colors is then repeated for the movements of the dots in the seventh second through the twelfth second. Start and finish indicate the position of the dot in the first video frame and the last video frame, respectively. (c) MSD- Δt plots for the trajectories of the fluorescent dots. Blue and red circles indicate the MSD- Δt plots for dot 1 and dot 2, respectively. The blue solid line and the red dashed line are the linear-fitted lines at $2\Delta t$, $3\Delta t$, and $4\Delta t$ for the MSDs (Δt is the video frame time: 33 ms). RD (N, n) was calculated as described in the Materials and Methods. (d) Trajectory of an immobile fluorescent dot in another cell.

Figure 5. Subcellular localization of GFP-FliG in a tethered cell and determination of the rotational center. (a) Schematic diagrams to illustrate the concept. A flagellar filament (sticky filament) extending from the cell body is immobilized on the coverslip and the cell body rotates around this point. Central and satellite fluorescent dots are localized at the rotational center and the cell

pole, respectively. These two dots were used to define the rotational center of the tethered cell. (b) A sequence of fluorescent images of a tethered cell. Images were captured at the normal video rate (30 frames/s) and every frame in the sequence is shown. C denotes the central dot and S marks the satellite dot. Bar, 1.6 µm. (c) The distributions of the positions of a satellite dot and a central dot in a tethered cell. Shaded squares indicate the distribution of the positions of the central dot (150 points). The small open squares indicate the centers of the ellipses fitted from each rotation of the satellite dot (we analyzed five rotations). The positions of the centers of the fitted ellipses were calculated using the least squared method from the center of the satellite dot. In this figure, the position of the rotational center (large open square) was defined as the average position of the five small open squares. The origin of this graph represents the center of the ellipse fitted based on the rotation of the satellite dot. Histograms indicate the distributions of the central dot along the X axis and Y axis. (d) Histograms of positional data obtained for central dots from 11 tethered cells. For each tethered cell, the positions of the central dot were obtained from 100 frames, and the range of the distribution of the central dots was normalized to the center of the fitted ellipse.

Strains	FliG variants	Swimming fraction $(\%)^{*1}$	Swimming speed (µm/s)* ²
DFB225	GFP-FliG	3 ± 3	12±3
DFB225	Wild-type	85 ± 6	26 ± 4
RP437	Wild-type* ³	96 ± 3	28 ± 6
		X	

Table 1. Motility parameters of the cells

*1. The average value calculated from more than eight visual fields.

*2. The average value calculated from more than 40 cells.

*3. Wild-type FliG encoded on the chromosome.

The cells were grown in media containing no arabinose (RP437), 0.0002% arabinose (DFB225 cells harboring the plasmid encoding *fliG*), or 0.0005% arabinose (DFB225 cells harboring the plasmid encoding *gfp-fliG*).



Strains	FliG variants	Rotational velocity* ¹	Rotational bias* ²	Switching frequency* ³
DFB225	GFP-FliG	3 ± 2	92 ± 15	0.4 ± 0.4
DFB225	Wild type	6 ± 2	65 ± 19	0.9 ± 0.8

Table 2. Rotational parameters of the cells

*1. Rotation velocity is shown in revolutions/s.

*2. Rotational bias is shown as the percentage of time spent rotating CCW.

*3. Switching frequency is shown as the number of reversals/s.

More than 40 tethered cells were investigated. Images were captured at 60 frames/s and 600 frames (10 s) were analyzed. The cells were grown in media containing 0.0002% arabinose (DFB225 cells harboring the plasmid encoding *fliG*), or 0.0005% arabinose (DFB225 cells harboring the plasmid encoding *gfp-fliG*).

	Description	Source or reference
<u>Strains</u>		
RP437	F ⁻ thi thr leu his met eda rpsL	41
	(wild type for motility)	
DFB225	RP437 $\Delta fliG$	42
RP3098	$\Delta(flhD - flhA)4$	43
<u>Plasmids</u>		
pBAD33	Cm ^r , PBAD	44
pBR322	Ap ^r , Tc ^r	45
pTY200	his6-gfp in pBAD33	24
pTY102	fliG of Vibrio alginolyticus in pBAD33	24
pSL27	<i>fliG</i> of <i>E. coli</i> in pAlter-1	41
pYF2	fliG of E. coli in pBAD33	This work
pYF1	his6-gfp-fliG of E. coli in pBAD33	This work
pYS11	fliC sticky in pBR322	27

Table 3. Bacterial strains and plasmids

Cm^r, chloramphenicol resistant; Ap^r, ampicillin resistant; Tc^r, tetracycline resistant; P*BAD*, *araBAD* promoter; his6, hexa-histidine tag



Fukuoka et al. Fig. 1



Fukuoka et al. Fig. 2



Fukuoka et al. Fig. 3



Fukuoka et al. Fig. 4



Fukuoka et al. Fig. 5

80^l

nm

nm