



A Mechanism of Modulating the Direction of Flagellar Rotation in Bacteria by Fumarate and Fumarate Reductase

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Abstract

Fumarate, an electron acceptor in anaerobic respiration of *Escherichia coli*, has an additional function of assisting the flagellar motor to shift from counterclockwise to clockwise rotation, with a consequent modulation of the bacterial swimming behavior. Fumarate transmits its effect to the motor *via* the fumarate reductase complex (FrdABCD), shown to bind to FliG—one of the motor's switch proteins. How binding of the FrdABCD respiratory enzyme to FliG enhances clockwise rotation and how fumarate is involved in this activity have remained puzzling. Here we show that the FrdA subunit in the presence of fumarate is sufficient for binding to FliG and for clockwise enhancement. We further demonstrate by *in vitro* binding assays and super-resolution microscopy *in vivo* that the mechanism by which fumarate-occupied FrdA enhances clockwise rotation involves its preferential binding to the clockwise state of FliG (FliG_{cw}). Continuum electrostatics combined with docking analysis and conformational sampling endorsed the experimental conclusions and suggested that the FrdA–FliG_{cw} interaction is driven by the positive electrostatic potential generated by FrdA and the negatively charged areas of FliG. They further demonstrated that fumarate changes FrdA's conformation to one that can bind to FliG_{cw}. These findings also show that the reason for the failure of the succinate dehydrogenase flavoprotein SdhA (an almost-identical analog of FrdA shown to bind to FliG equally well) to enhance clockwise rotation is that it has no binding preference for FliG_{cw}. We suggest that this mechanism is physiologically important as it can modulate the magnitude of ΔG^0 between the clockwise and counterclockwise states of the motor to tune the motor to the growth conditions of the bacteria.

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Introduction

Bacteria such as *Escherichia coli* swim by rotating their flagella. Each flagellum rotates by a motor (reviewed in Refs. [1–4]), embedded in the cell membrane. The motor's direction of rotation is the target of navigation control by the chemotaxis system because it dictates the swimming mode of the bacteria, with counterclockwise rotation pushing the bacterium forward and clockwise rotation causing cell reorientation [1–5]. The direction of rotation is determined by a “switch,” made of circular layers of three proteins, FliN, FliM and FliG.

An unexpected finding made over two decades ago was that fumarate, a dicarboxylate that serves as an electron acceptor in anaerobic respiration [6], acts as a switching and clockwise factor under aerobic conditions. Thus, in *E. coli* under aerobic conditions, fumarate enabled the flagella of cytoplasm-free cell envelopes to switch their direction of rotation [7,8] and enhanced switching and clockwise rotation in intact bacteria [9,10]. These effects were due, in part, to reduction of the standard free energy difference (ΔG^0) between the clockwise and counterclockwise states of the switch [9], but they were neither due to the function of fumarate as an electron acceptor [8] nor due to

electron-transport-related energization [11]. The mechanism by which fumarate enhances clockwise rotation and switching has remained obscure. Years later, the target of fumarate for these functions was identified. The membrane-bound enzyme fumarate reductase (also known as quinol-fumarate oxidoreductase and as anaerobic complex II) was found to interact with FliG at the switch [11]. This enzyme functions in the terminal step of the anaerobic electron transfer chain, at which electrons from the menaquinol cofactor are used to reduce fumarate [12]. Its expression level is 10- to 15-fold higher under anaerobic conditions than under aerobic conditions [13]. The presence of fumarate increases the expression level 1.5-fold further [13,14]. Fumarate reductase is a membrane-linked heterotetramer ([12] for a review), composed of FrdA, a

cytoplasmic flavoprotein subunit; FrdB, a cytoplasmic iron–sulfur protein; and a transmembrane anchor domain that consists of two subunits, FrdC and FrdD (Fig. 1a; the enzyme will be termed hereafter FrdABCD). FrdA is the subunit that binds fumarate [22]. Intriguingly, the almost identical aerobically expressed enzyme, succinate dehydrogenase (SdhCDAB), also binds to FliG, but this binding has no effect on flagellar rotation [11]. Succinate dehydrogenase is a component of the tricarboxylic acid cycle, where it converts succinate to fumarate, and also a component of the aerobic electron transport chain, where it reduces quinone to quinol [12]. The enzyme is a structural and functional homolog of fumarate reductase, and both enzymes are considered to be evolutionarily related and part of the same enzyme

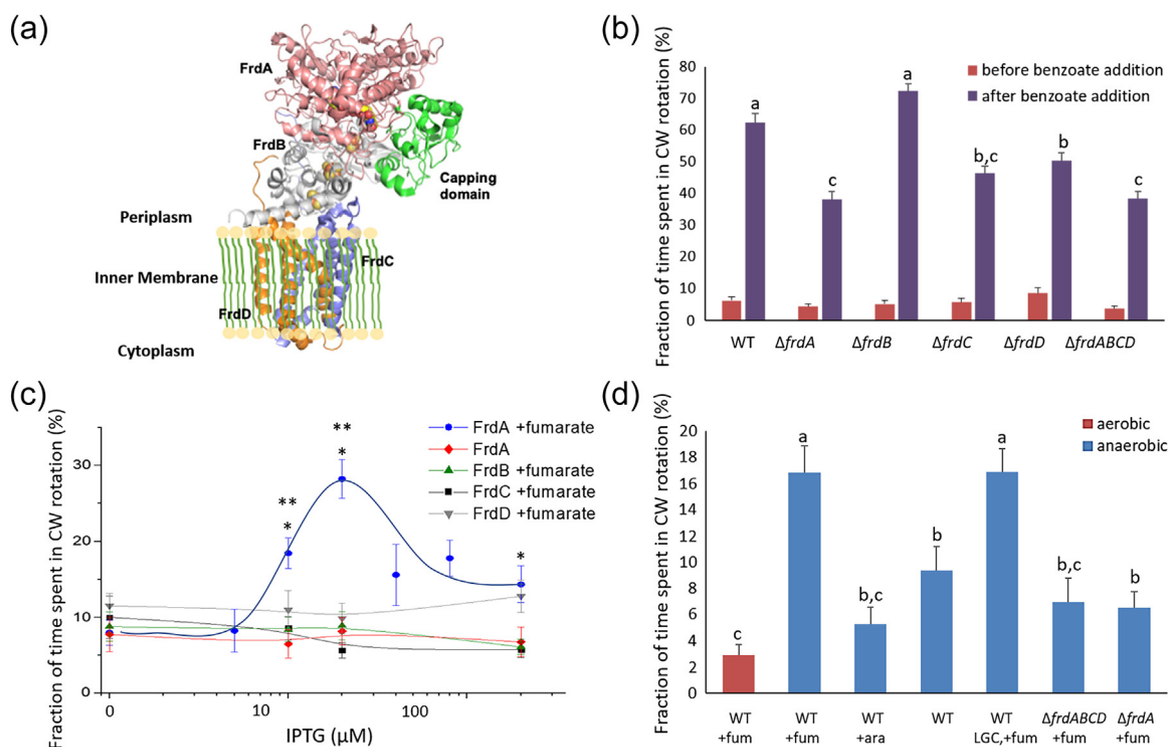


Fig. 1. FrdA alone can enhance clockwise rotation. (a) Structure of *E. coli* FrdABCD. Based on PDB 3P4P. (b) Clockwise rotation in mutants lacking individual FrdABCD subunits. See Table 2 for strains. The fraction of time spent in clockwise rotation was calculated in 1-min intervals immediately before and after the addition of benzoate (50 mM, pH 7.0). Columns marked with the same letter were statistically indifferent. Columns marked with dissimilar letters were significantly different ($P < 0.05$) according to Kruskal–Wallis statistical test ($n = 113$ –190 cells). (c) Effect of FrdA level on clockwise rotation. The expression of the subunits shown was done from the IPTG-induced plasmids listed in Table 2 in B275 $\Delta frdABCD$ background. Where indicated, fumarate (40 mM) was present in the medium during growth but not during the experiment. The curve was hand-drawn. * $P < 0.001$ (for 10 and 20 μ M IPTG) or $P < 0.01$ (for 200 μ M IPTG) relative to absence of fumarate at the same IPTG concentration, and ** $P < 0.05$ or $P < 0.001$ (for 10 or 20 μ M IPTG, respectively) relative to zero IPTG in the presence of fumarate according to Kruskal–Wallis statistical test ($n = 44$ –95 cells). (d) Anaerobic growth in the presence of fumarate increases clockwise probability. The wild-type strain used was B275. See Table 1 for the other strains. Prior to tethering, the cells were washed and resuspended in fumarate-free motility buffer. Low-growth control stands for wild-type cells that grew to $OD_{600} = 0.09$, similar to the maximal growth OD of $\Delta frdABCD$ cells under anaerobic conditions. Columns marked with the same letter were statistically indifferent. Columns marked with dissimilar letters were significantly different according to Kruskal–Wallis test ($P < 0.001$, except for the difference between aerobic WT + fum and $\Delta frdA$ + fum, for which $P < 0.05$; $n = 70$ –170 cells). Abbreviations: ara, arabinose; fum, fumarate; LGC, low-growth control; WT, wild type.

family [12]. In fact, FrdABCD and SdhCDAB can functionally replace each other when overexpressed because they can catalyze the same reactions [76]. Due to the high degree of structural and functional similarity of the two enzyme complexes, they bind fumarate equally well [22]. It is unclear how one of them, FrdABCD, enhances clockwise rotation, whereas the other, SdhCDAB, does not. This study looks for the cause of this difference and for the mechanism by which fumarate enhances switching and clockwise rotation.

Results

Intactness of the FrdABCD complex is not essential for clockwise enhancement

FrdABCD was shown to interact with FliG and be involved in multiple functions of the flagellar motor, including enhancement of clockwise rotation [11]. To determine whether the FrdABCD complex, as a whole, is required for this enhancement or whether one or more of its subunits are sufficient, we first examined how deletion of each subunit affects clockwise rotation. We focused on the repellent response of mutants lacking each of the four subunits of FrdABCD (one at a time) because the effect of FrdABCD deletion on clockwise rotation is best observed in response to a repellent [11]. To determine the direction of flagellar rotation, we tethered individual cells *via* their flagellum to glass and recorded their direction of rotation around their immobilized flagella [23]. Consistent with Förster resonance energy transfer studies suggesting that the binding site for FliG is on FrdA [24], we found that both *frdA* and *frdABCD* deletions similarly reduced the extent of clockwise rotation following stimulation with the repellent benzoate (Fig. 1b). In contrast, a *frdB* deletion did not reduce the rotation in this direction and, perhaps, even increased it to a level slightly higher than that of the wild-type parent (though the increase was statistically insignificant). Since FrdB links FrdA to the membrane part of the complex, that is, to the FrdCD units (Fig. 1a) [12], the absence of FrdB means that FrdA is free in the cytoplasm. The ability of the Δ *frdB* mutant to generate high clockwise levels therefore suggests that FrdA, which lacks enzymatic activity by itself [25], is nevertheless fully functional in generating clockwise rotation in its soluble, cytoplasmic form. This is consistent with the observation that the enhancement of clockwise rotation by fumarate, though done *via* FrdABCD, is independent of the electron-transfer activity of the latter [8,11]. The clockwise responses of the Δ *frdC* and Δ *frdD* mutants to benzoate were stronger than that of the Δ *frdA* mutant (though the difference was statistically significant for the Δ *frdD* mutant only). However, they were lower than the wild-type parent's response, probably

due to the futile interaction of FrdA with free FrdB in these mutants (FrdB cannot be linked to the membrane in the absence of either FrdC or FrdD [12]). Thus, it appears that an intact FrdABCD complex is not obligatory for proper clockwise rotation of the flagellar motor and that FrdA in its soluble form is sufficient for enhancing clockwise rotation.

Overproduced FrdA generates clockwise rotation

To validate the above conclusion, we examined the effect of overproduction of each Frd subunit (each tagged with 6xHis at the *N*-terminus; for simplicity, the tag will not be mentioned hereafter) in the absence of the other subunits, thus ensuring that the measurement is of the subunit itself and not of the subunit in a complex. Aerobic expression of FrdA in a Δ *frdABCD* mutant (strain B275 Δ *frdABCD*) to approximately the level measured in wild-type cells under anaerobic conditions (Supplementary Fig. S1) generated a strong clockwise bias of the motor, provided that the cells had been grown in presence of fumarate (Fig. 1c). This dependence on fumarate is consistent with the role of FrdABCD to transmit the effect of fumarate to the motor [11]. In contrast, expression of the other three subunits did not affect the level of clockwise rotation (Fig. 1c). These results not only endorse the conclusion that FrdA is sufficient for clockwise rotation but also further suggest that when the concentration of soluble, fumarate-bound FrdA is sufficiently high as in cells under anaerobic conditions (conditions under which the FrdABCD complex is maximally expressed [13]), it can promote clockwise rotation even in the absence of any other stimulation.

Notably, expression of FrdA to higher levels decreased the fraction of time spent in clockwise rotation (Fig. 1c). Two potential causes could account for this. One is reduced concentration of soluble FrdA due to FrdA aggregation at high concentrations. We put this possibility to the test by examining both the soluble and insoluble fractions of the lysed cells. Western blots of these fractions demonstrated that, although the level of FrdA in the insoluble fraction increased with the IPTG concentration, its level in the soluble fraction did not decrease (Fig. S1C), thus excluding reduced FrdA concentration due to aggregation as a cause of the observed decrease in clockwise rotation at high IPTG concentrations. The other potential cause is competition of fumarate-free FrdA with fumarate-bound FrdA for FliG. The fraction of fumarate-free FrdA molecules is expected to increase with the level of overexpression. Since fumarate is required for the effect of FrdA on clockwise rotation (Fig. 1c) but, apparently, not for its binding to FliG (see below), it is probable that the fumarate-free FrdA molecules would compete with the fumarate-bound FrdA molecules for FliG but would not produce clockwise rotation. The consequence of this futile binding to FliG would be reduced clockwise rotation.

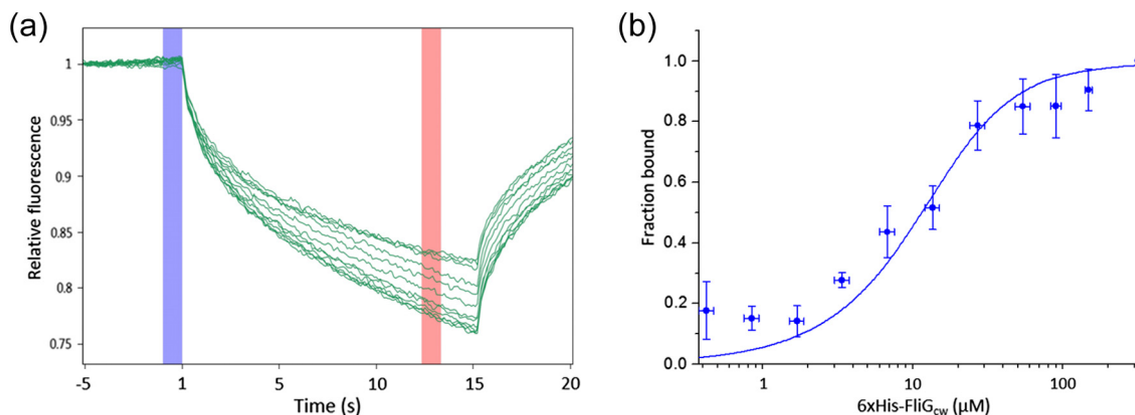


Fig. 2. Binding of FrdA to FliG_{cw} monitored with MST. The FrdA concentration was kept constant at 15 μM and FliG_{cw} was serially diluted 1:1 beginning with 138 μM . The natural fluorescence of the FAD cofactor was utilized for measuring the MST signal. (a) MST traces of a representative experiment. Averaged values were taken before and after heating (blue and red areas). (b) Binding curve obtained by averaging the values at each FliG_{cw} concentration in four separate experiments. The curve is a fit ($R^2 = 0.955$) according to the law of mass action equation [27], yielding $K_d = 4 \pm 1 \mu\text{M}$ (\pm SEM, $n = 4$).

Anaerobic growth increases clockwise probability due to elevated FrdA

To determine whether or not the dependence of clockwise rotation on the FrdA level is physiologically relevant, we compared the rotation of tethered cells grown aerobically with the rotation of cells grown anaerobically, where the FrdABCD complex is expressed to higher levels [13]. Consistent with the results of FrdA expression described above, the clockwise level of anaerobically grown cells was significantly higher than that of aerobically grown cells ($P < 0.001$; Fig. 1d, two left columns). This effect was fumarate dependent as absence of fumarate in the growth medium reduced clockwise rotation ($P < 0.001$; Fig. 1d, fourth column). As expected, the clockwise levels of the negative-control cells, anaerobically grown ΔfrdA and $\Delta\text{frdABCD}$ cells (Fig. 1d, two right columns), were significantly lower than that of the anaerobically grown wild-type cells ($P < 0.001$), although they were somewhat higher than aerobically grown cells, probably due to the anaerobic growth conditions. Since, under anaerobic conditions, $\Delta\text{frdABCD}$ cells maximally grew to $\text{OD}_{600} = 0.09$, we performed a control with wild-type cells grown to the same low OD. The clockwise level of these control cells was similar to that of wild-type cells grown anaerobically to $\text{OD}_{600} = 0.4$ (Fig. 1d, second and fifth columns). As another control, we studied wild-type cells that grew anaerobically in the presence of arabinose instead of fumarate. These cells grew similarly well in the presence of either one of these compounds ($\text{OD}_{600} = 0.6$ and 0.4 in the presence of arabinose and fumarate, respectively). However, the level of clockwise rotation was low when arabinose replaced fumarate (Fig. 1d, third column). The switching frequency of the rotation followed the same trend as clockwise rotation, that is,

higher switching frequency when clockwise rotation was higher (Fig. S2 versus Fig. 1d), consistent with the function of fumarate as a clockwise and switching factor [7–10]. It should be noted that the rotation of anaerobically grown cells was measured aerobically. We therefore performed a control to verify that FrdA, which can only be expressed to high levels under anaerobic conditions [13], had not degraded during the ~ 1 h aerobic tethering assay. We subjected lysates of cells exposed to air for 1.5 h after anaerobic growth to Western blots followed by mass spectrometry. Clearly, 1.5-h air exposure essentially did not affect the FrdA level (Fig. S3). Thus, anaerobic growth conditions elevate the FrdA level with a resultant clockwise rotation, an effect that is maintained for at least 1.5 h after air exposure.

FrdA preferentially interacts with the clockwise conformation of FliG

Our observations that FrdA binding to FliG (in the presence of fumarate) is sufficient for clockwise enhancement raised two potential causes of this enhancement. One is that FrdA binding to the counterclockwise state of FliG (FliG_{ccw}) results in a shift of the latter to its clockwise state (FliG_{cw}). The other is that FrdA preferentially binds to FliG_{cw} with a consequent stabilization of this state. To examine whether any of these possibilities holds, we compared the binding of FrdA to FliG_{cw} and FliG_{ccw} both *in vitro* and *in vivo*.

In vitro, we measured the binding of purified FrdA to purified wild-type FliG, which is essentially FliG_{ccw} [26], and to mutant FliG (containing a PAA deletion at positions 169–171) locked in the clockwise conformation [26], that is, FliG_{cw}. Employing microscale thermophoresis (MST) [27], we found that FrdA bound to FliG_{cw} with $K_d \approx 4 \mu\text{M}$ (Fig. 2). Remarkably,

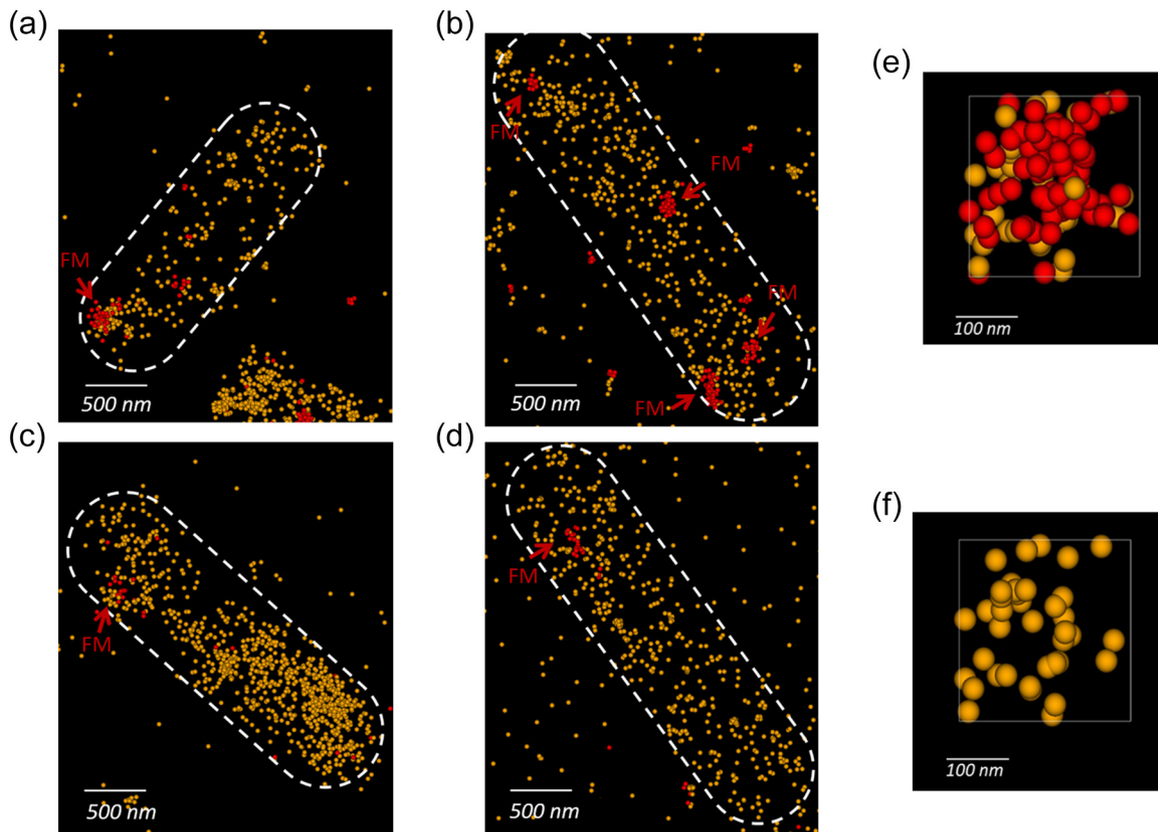


Fig. 3. Representative images of cells containing fluorescently labeled FrdA and FliM. (a and b) Combined PALM/STORM imaging of cells containing wild-type motors (strain DFB190 Δ frdABCD containing the plasmids pH3–Dendra2–FrdABCD and pFliM_{WT}–mPlum). Yellow dots represent individual Dendra2 (FrdA) molecules; red dots represent individual Alexa 647-labeled wild-type FliM molecules; FM indicates the location of the flagellar motors. (c and d) As in panels a and b, but for cells containing motors with FliM_{CW} molecules (strain DFB190 Δ frdABCD containing the plasmids pH3–Dendra2–FrdABCD and pFliM_{CW}–mPlum). (e) A ring structure of FrdA molecules formed around the motor. (f) As in panel e, but with the motor fluorescence omitted to reveal the ring-like structure.

Table 1. FrdA molecules are more crowded near clockwise motors

	Wild-type motors	Clockwise motors
FrdA–Dendra2 molecules/cell ^a	250 ± 86	230 ± 191
Density near motors (molecules/ μ m ²) ^b	140 ± 98	250 ^d ± 150
Density in other areas (molecules/ μ m ²) ^c	120 ± 57	160 ± 75

Counting of molecules was done automatically.

^a The values shown are the mean number of FrdA–Dendra2 molecules \pm SD in 28 wild-type and 32 clockwise cells (including cells in which Alexa 647-labeled FliM molecules were not detected).

^b The values shown are the mean number of FrdA–Dendra2 molecules \pm SD found within a 0.3×0.3 - μ m square around the Alexa 647 label for 24 and 15 wild-type and clockwise motors, respectively (in 19 and 10 cells, respectively).

^c The cells were those from the preceding row.

^d $P < 0.05$ relative to density in other areas and relative to density near wild-type motors according to Student's paired *t*-test and non-parametric Mann–Whitney test, respectively.

this K_d is very similar to the K_d of CheY binding to FliM [28,29], suggesting that the interactions of FrdA and CheY with the motor are similarly strong. Unlike FliG_{CW}, when we studied FliG_{WT} at the same concentration range, we could neither obtain a consistent binding curve nor reach binding saturation. This could potentially be due to too weak FrdA binding to FliG_{CW}, to the possibility that FliG_{CW}, but not FliG_{WT}, might oligomerize *in vitro* [30], or to the theoretical possibility that FliG_{WT}, but not FliG_{CW}, contains more than one conformation of FliG.

In vivo, we employed super-resolution microscopy, more specifically photoactivated localization microscopy (PALM), which provides up to ~ 25 nm resolution [31]. We fluorescently labeled FrdA with the photoswitchable protein, Dendra2 [32], and expressed the frdABCD operon under anaerobic conditions from a low-copy number plasmid in the presence of fumarate. In addition, we labeled FliM at the motor with Alexa 647 (detailed in Materials and

Methods). To obtain FliG_{cw} without a mutation in *fliG*, we expressed FliG in a motor that is almost constantly in the clockwise conformation due to mutation in *fliM* [*fliM*(R60C)] [33]. Representative examples are shown in Fig. 3a–d. The number of FrdA–Dendra2 molecules in the cell seemed to vary largely between cell to cell (Table 1). A portion of the molecules were clustered, usually 1–3 clusters per bacterium (e.g., Figs. 3a, c), as was reported for

some other oxidative phosphorylation complexes in *E. coli* [34]. Such clusters were hardly detected in a negative control strain, in which Dendra2 was encoded in the same vector under the control of the same *frdABCD* promoter but without being fused to FrdA (Fig. S4). Automated clustering analysis indicated an order of magnitude less clusters in the negative control (0.2 ± 0.4 clusters/cell, 269 ± 181 molecules/cell, mean \pm SD of 22 control cells).

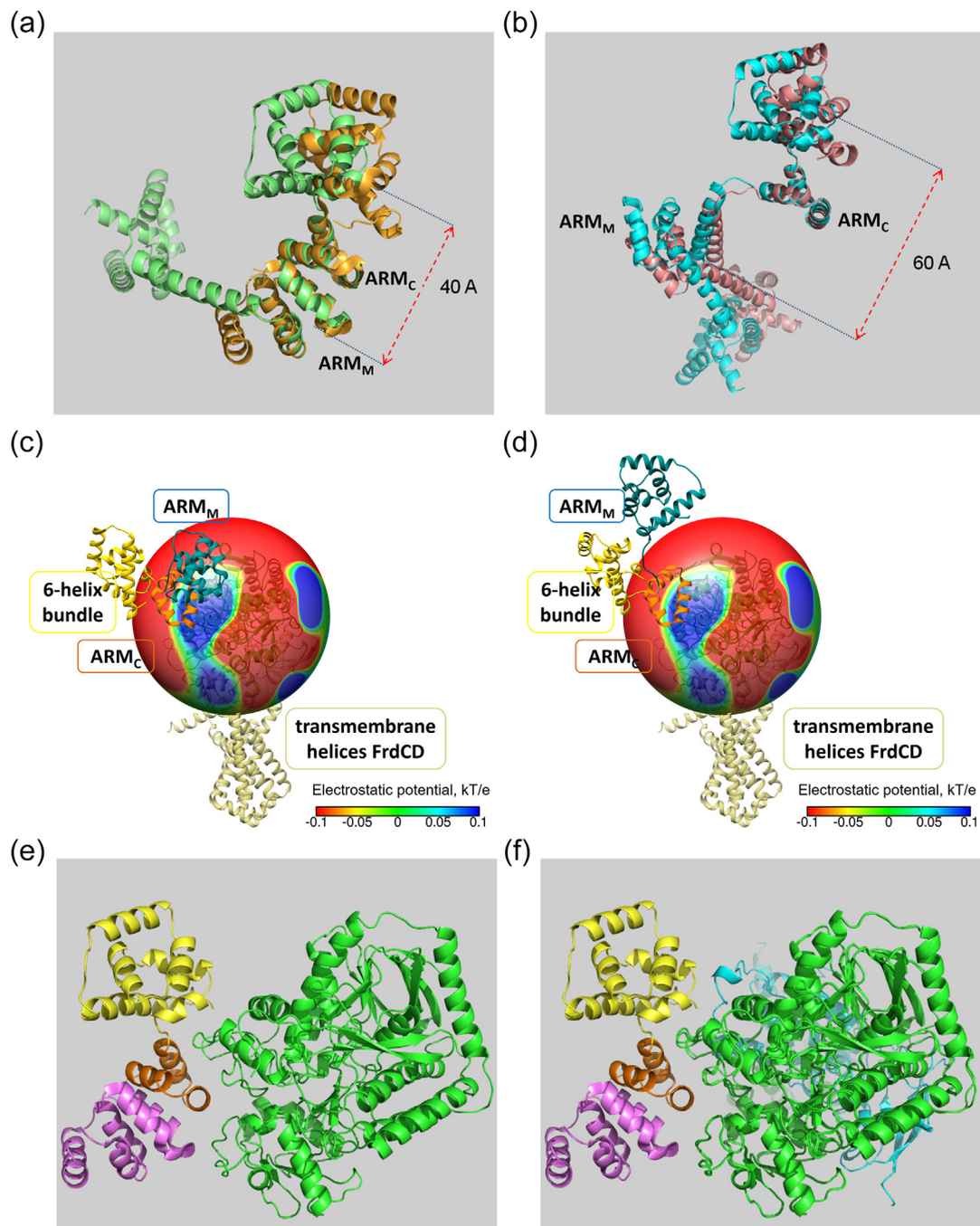


Fig. 4 (legend on next page)

Clearly, the density of FrdA–Dendra2 molecules near clockwise motors (i.e., within an area of $0.09 \mu\text{m}^2$ around the motor—see [Materials and Methods](#)), but not wild-type motors, was significantly higher than in other areas, although the total number of molecules was similar in both cell types ([Table 1](#)). In some of the cells, we observed FrdA molecules arranged around the motor in the form of a ring (e.g., [Fig. 3e, f](#)). What can be concluded is that FrdA is indeed associated with the flagellar motor *in vivo* under anaerobic conditions, and the association is more intense with clockwise motors.

Molecular basis for the preferred binding to FliG_{cw}

The results above suggested that FrdA is sufficient for clockwise enhancement and that it preferentially binds to the clockwise state of FliG. To obtain information for the structural causes of these results, we carried out molecular dynamics (MD) conformational sampling of the FliG protein combined with continuum electrostatics and docking analysis. Employing the structures of full-length FliG from *Aquifex aeolicus* (PDB code [3HJL](#)) and of the middle-to-C-terminal domain of FliG from *Thermotoga maritima* (PDB codes [4FHR](#), [3AJC](#)) as representatives of the counterclockwise ([3HJL](#)) and the clockwise ([3AJC](#)) states of *E. coli* FliG [[35](#)], we found multiple conformational transitions between the clockwise and counterclockwise states. These two structures are significantly different at the C-terminal domain, primarily due to the rotation of a six-helix bundle (helices C1–6) relative to the armadillo repeat motif (ARM_C) in this domain [[36](#)] ([Fig. 4a, b](#)). Conformational analysis of MD trajectories revealed that, of the many conformations obtained, only four conformation clusters appeared: FliG_{ccw} and FliG_{cw} each in the compact ([Fig. 4a](#)) and extended ([Fig. 4b](#)) states (Supplementary Information for detailed analysis). The calculations further showed that FrdA

preferentially interacts with compact FliG_{cw}, that the interaction involves both ARM_M and ARM_C motifs, and that the binding of FrdA to FliG_{cw} is driven by positive electrostatic potential generated by FrdA and the negatively charged areas of FliG ([Fig. 4c, d](#); Supplementary Information for details). These electrostatics calculations combined with docking analysis further suggested that the interaction between FliG_{cw} and isolated FrdA ([Fig. 4e](#)) is weaker than the interaction between FliG_{cw} and FrdABCD ([Fig. 4f](#)) due to the smaller average intermolecular contact surface area in the case of FrdA (estimated to be 990 versus 1200 \AA^2 for FrdA and FrdABCD, respectively). Our modeling indicates that the stronger binding of the whole complex to FliG_{cw} is because FrdB also contributes to the electrostatic and hydrophobic interaction ([Fig. 4f](#); Supplementary Information for details). This stronger binding is well consistent with the measured K_d values for the binding of FrdA ([Fig. 2](#)) and FrdABCD [[11](#)] to FliG (4 and $0.4 \mu\text{M}$, respectively).

A related question is what the function of fumarate is in FrdA-mediated clockwise enhancement. According to recent structural data [[37](#)], the presence of a dicarboxylate ligand of FrdA rotates the capping domain in relation to the flavin domain of FrdA. Our docking results suggest that residues 275–280 of the capping domain are exposed to the FliG binding area and interact with the ARM_M motif. These data suggest that the fumarate-triggered rotation or shift of the capping domain increases the total contact surface area between FliG_{cw} and FrdA with a consequent increased affinity between them.

Molecular basis for the effectiveness of FrdA versus SdhA in clockwise enhancement

One of the puzzles related to the interaction of FrdABCD with FliG is why the interaction of the structurally and functionally similar protein complex SdhCDAB, although binding with similar affinity to FliG, does not have any effect on clockwise rotation.

Fig. 4. Molecular basis for the preferred interactions as revealed by MD simulations combined with docking analysis. (a) “Compact” conformation of FliG_{cw} (green) and FliG_{ccw} (orange). (b) “Extended” conformation of FliG_{cw} (cyan) and FliG_{ccw} (salmon). Note the difference between compact (a) and the extended (b) conformations with respect to the distance between the center of the M domain and the center of the C-terminal domain. (c and d) The compact conformation of FliG_{cw} interacts with FrdA better than the extended conformation. The panels show the distribution of the electrostatic potential mapped on a sphere of a 40-Å radius around a FrdA subunit interacting with FliG in its compact (c) and extended (d) conformations. FrdA values of the electrostatic potential on the color maps are given in kT/e units. It is evident that the extended conformation is less preferable for binding to FrdA (d). Higher binding affinity can be achieved in the compact conformation when both the ARM_C and ARM_M motifs of FliG enter the “blue area” around FrdA (c) at the initial step of the binding process, followed by the final stage (e, f). In these initial “encounter complexes” (c and d), minimal FliG–FrdA surface-to-surface distance ($\sim 15 \text{ \AA}$) is found between Lys121 and Lys281 of FrdA and Glu212 from ARM_C helix_1 of FliG. Electrostatic interaction energies between FliG_{cw} and FrdA are estimated to be in the range of -8 to -30 kJ/mol ([Table S1](#)), suggesting attraction and efficient steering at long distances. (e) Proposed binding mode of a FrdA monomer to FliG_{cw}. In this final step of the binding, helix_1 of the ARM_C motif fits into the groove on the surface of FrdA surrounded by a “lysine cluster.” Green, FliG; yellow, C-terminal six-helix bundle; pink, the ARM_M motif; orange, the ARM_C motif. (f) Proposed binding mode of FrdABCD to FliG_{cw}, showing possible involvement of the loop Lys118-Pro137 (cyan) of FrdB in contacts with the C-terminal six-helix bundle of FliG_{cw}.

To address this enigma, we compared between FrdABCD and SdhCDAB with respect to their predicted interaction with FliG. Our modeling showed that, unlike FrdA within FrdABCD, which binds specifically to FliG_{cw}, SdhA binds equally well to both FliG_{cw} and FliG_{ccw} (Supplementary Information and Fig. S10). It thus seems that the mechanism by which FrdA enhances clockwise rotation is selective binding to FliG_{cw} with a consequent stabilization of the motor's clockwise state. In contrast, the similar binding of SdhA to both conformations of FliG stabilizes both directions of rotation, explaining the failure of SdhA binding to FliG to enhance clockwise rotation.

Discussion

Over two decades ago, fumarate was found to be a switching and clockwise factor in *E. coli* [7–10], which acts by decreasing ΔG^0 between the clockwise and counterclockwise states [9]. A decade later, the enzyme FrdABCD, but not its aerobically expressed homolog SdhCDAB, was identified as the protein that transmits the effect of fumarate to the motor [11]. These findings were intriguing because they neither offered clues to the mechanism by which fumarate and FrdABCD affect the direction of flagellar rotation nor explained how both “twin” enzymes similarly bind to the motor but, nevertheless, only one of them affects the rotation. The findings, made herein, appear to provide answers to these questions and are discussed below.

Function of FrdABCD at the motor

The observations, made in this study, of reduced repellent-stimulated clockwise enhancement in a $\Delta frdA$ mutant similar to a $\Delta frdABCD$ mutant and elevated clockwise rotation in response to FrdA overproduction, but not to overproduction of other subunits (Fig. 1), indicated that FrdA is the subunit responsible for the clockwise-enhancement effect of FrdABCD. The finding that clockwise enhancement was high in a $\Delta frdB$ mutant, where FrdA is free in the cytoplasm, further suggested that FrdA, in its soluble form, is sufficient to enhance clockwise rotation. The results, however, do not distinguish between whether, under normal physiological conditions (i.e., intact FrdABCD), FrdA acts as a soluble molecule or within the FrdABCD complex. Hitherto, there is no information about whether or not FrdA exists in the cell under physiological conditions in a soluble form. (Obtaining such information is not trivial because preparation of a cell extract might release FrdA from the complex, making it difficult to distinguish how much free FrdA exists *in vivo*.) Nevertheless, the findings that the intact FrdABCD complex binds to FliG [11] suggest that, under physiological conditions, FrdA binds to, and acts on, the motor from within the intact FrdABCD complex. This conclusion is well in line with the 10-fold stronger

binding of FrdABCD to FliG compared to that of FrdA, also revealed from our docking analysis combined with electrostatics calculations.

Mechanism of clockwise enhancement by FrdA and fumarate

An obvious question is how FrdA, which lacks any enzymatic activity [25], affects the rotation of the flagellar motor by merely binding to FliG and how fumarate affects FrdA. In principle, FrdA could affect the rotation by one of the following ways: (1) It could selectively bind to FliG_{cw} and stabilize this conformation; (2) it could bind to FliG_{ccw} and stimulate its transition to the clockwise conformation; and (3) it could bind to either conformation of FliG and affect somehow the entire ring organization with a resultant clockwise rotation. Our results, which demonstrated preferred FrdA binding to FliG_{cw} both *in vitro* (Fig. 2) and *in vivo* (Fig. 3, Table 1), supported the first option. Furthermore, our MD conformational analysis indeed revealed that FrdA preferentially interacts with FliG_{cw} in its compact conformation and that the interaction is electrostatic in nature (Fig. 4). Importantly, our *in silico* analysis revealed that, unlike the selective binding of FrdA to FliG_{cw}, SdhA binds equally well to both FliG_{cw} and FliG_{ccw} (Supplementary Information and Fig. S10), explaining why SdhCDAB binding to FliG does not generate clockwise rotation [11]. Thus, the mechanism of clockwise enhancement by FrdA is probably preferred binding to FliG_{cw} with a consequent stabilization of the motor's clockwise state. As our docking results suggest, the function of fumarate is likely to cause a conformational change [37–40] that increases the total contact surface area between FrdA and FliG_{cw} with a consequent increased affinity between them.

The proposed mechanism requires that, for fumarate being effective, at least one of the FliG subunits would be in its clockwise conformation. This is likely to be indeed the case at physiological temperatures, as evident from the occurrence of some clockwise rotation even in the absence of fumarate (Fig. 1d). This clockwise rotation results from binding of the signaling molecule CheY to its target at the motor—the switch protein FliM ([2] for a review), followed by FliM interaction with FliG at a specific binding surface of the latter [41]. However, CheY *per se* is apparently neither involved in, nor necessary for, the functions of fumarate and FrdA. This is because, at temperatures of $\sim 2^\circ\text{C}$, where some switching to clockwise rotation occurs even in the complete absence of CheY and of any other cytoplasmic chemotaxis protein, fumarate can still effectively enhance switching and clockwise rotation [9]. Consistently, at room temperature where clockwise rotation does not occur unless CheY is present, fumarate is effective only in the presence of CheY [9,10].

In vivo, FliM binds to the middle domain of FliG (FliG_M) in different bacterial species [42]. Analysis of

available structural data [26,36,42,43] shows that this binding is not affected by the arrangement of FliG_M and FliG_C, meaning that both the extended and compact states of FliG can bind to FliM similarly well. This binding has bearings on FrdA interaction with FliG because the movement of the latter is more restricted due to the FliG-FliM_M packing and protein crowding within the switch complex, resulting in stronger interaction between FliG and FrdA. The lower amount of high-dielectric water and hydrophobic environment in the highly viscous switch complex should also enhance electrostatic interactions [44] in the FliG–FrdA binding.

Each flagellar motor contains ~25 FliG molecules arranged in a ring [45]. It is reasonable to assume that even in a counterclockwise motor, one or more of the FliG subunits would temporarily be in the clockwise conformation, depending on the level of active CheY in the cell, and that fumarate-occupied FrdA would bind to these FliG_{cw} subunits and stabilize them. An expected outcome is that these stable FliG_{cw} units in a counterclockwise ring would make the latter thermodynamically unfavorable and increase the probability of a ring shift to the clockwise state, as was shown earlier for genetically locked FliG_{cw} [46]. As a result, ΔG^0 between the clockwise and counterclockwise states would decrease in the presence of fumarate, as was indeed found experimentally [9].

Physiological significance

The results of this study demonstrated a remarkable case in which a subunit of a respiratory enzyme, which is mainly expressed under anaerobic conditions, directly interacts with a subunit of the flagellar motor when it is in a specific conformation, and affects its function. A legitimate question is what advantage this interaction confers on the bacterial cell. Since this interaction modulates the magnitude of ΔG^0 between the clockwise and counterclockwise states of the motor [9] and depends on the level of FrdA (Fig. 1c), it is reasonable that one of the functions of this interaction is to tune the motor according to the growth conditions of the bacteria. Anaerobic conditions elevate the expression level of FrdABCD [13] with a consequent elevation of clockwise rotation to moderate levels (Fig. 1d). The outcome of such clockwise levels would be moderate tumbling frequencies of the cells, which are expected to better propagate in viscous media, such as agar, than cells with very low or very high tumbling frequencies [47]. This can be advantageous to *E. coli* in the mammalian digestive tract, where *E. coli* inhabits the polysaccharide-rich mucus layer lining the epithelium and where the environment is generally anaerobic [48,49] and respiration on fumarate via FrdABCD is essential [50,51]. Furthermore, in such energy-poor environments, swimming toward oxygen and energy sources is essential. This navigation, which is mediated by the chemotaxis receptor Aer [52,53], is expected to be

more efficient with moderate tumbling frequencies, and the cells are expected to respond to the Aer-mediated signals with higher sensitivity. Consistent with this notion, an additional function of Aer is upregulation of the expression of anaerobic respiratory enzymes, including fumarate reductase [54].

Materials and Methods

Materials

Amino acids were purchased from Calbiochem (Darmstadt, Germany). Ampicillin, kanamycin, chloramphenicol, tetracycline, fumarate (sodium salt), IPTG, L-arabinose, imidazole, benzoate (sodium salt), poly L-lysine, resazurin (sodium salt), ethylenediaminetetraacetic (EDTA), and EDTA-free cOmplete™ Protease Inhibitor cocktail were purchased from Sigma-Aldrich (St. Louis, MO, USA). Phosphate-buffered saline (PBS) was purchased from Biological Industries USA, Inc. (Cromwell, CT, USA). Paraformaldehyde was purchased from Thermo Scientific Inc. (Rockford, IL, USA). Vitamin B₁ hydrochloride was purchased from Fluka (Gillingham, England). BugBuster™ Protein Extraction Reagent (used to lyse cells for Western blots) was purchased from Novagen, Inc. (A Brand of EMD Biosciences, Inc., an Affiliate of Merck KGaA, Darmstadt, Germany). Bacto Casamino Acids and Bacto Agar were purchased from Difco (Franklin Lakes, NJ, USA). Bovine trypsin was purchased from Promega (Madison, WI, USA). LC–MS/MS solvents were purchased from Bio-Lab Chemicals (Jerusalem, Israel). All the compounds were of the highest purity grade available. Antibodies: anti-flagellin antibody (serotype H48) was a gift from the National Center for Enterobacteriaceae (Central Laboratories, Ministry of Health, Jerusalem, Israel). Anti-FrdABCD antibody was prepared in earlier studies in our laboratory [[11]; anti-mCherry monoclonal antibody was purchased from Abcam (Cambridge, MA, USA); and Alexa Fluor 647 anti-mouse antibody was purchased from Jackson ImmunoResearch Laboratories, Inc. (West Grove, PA, USA).

Bacterial strains

The strains and plasmids used in this study are listed in Table 2. For construction of B275 Δ frdABCD and DFB190 Δ frdABCD, a P1 phage lysate [55] prepared on RP437 Δ frdABCD was used to infect B275 and DFB190, respectively. Positive colonies were selected for tetracycline resistance and for their inability to grow on a fumarate and glycerol minimal medium under anaerobic conditions. For construction of B275 Δ frdA, B275 Δ frdB, B275 Δ frdC, and B275 Δ frdD, the P1 phage lysates, prepared on

Table 2. Strains and plasmids used in this study

Strain/plasmid	Relevant genotype/description	References
Strains		
BL21(λ DE3)pLysS	<i>hsdS gal ompT r_B⁻ m_B⁻</i> [Δ C1857(Ts)indl Sam7 min5 lacUV5-T7 gene 1]	[15], Novagen
JW4115	<i>rmB3 lacZ4787 hsdR514 (araBAD)567 (rhaBAD)568 ΔfrdA</i>	[16]
JW4114	<i>rmB3 lacZ4787 hsdR514 (araBAD)567 (rhaBAD)568 ΔfrdB</i>	[16]
JW4113	<i>rmB3 lacZ4787 hsdR514 (araBAD)567 (rhaBAD)568 ΔfrdC</i>	[16]
JW4112	<i>rmB3 lacZ4787 hsdR514 (araBAD)567 (rhaBAD)568 ΔfrdD</i>	[16]
RP437	<i>his thr leu metE thi eda rpsL</i> ; wild type for chemotaxis	[17]
RP437 Δ sdh Δ frd	Δ sdh Δ frd, RP437 parent	[11]
B275	<i>thr leu metE thi lac rpsL</i> ; wild type for chemotaxis	[18]
B275 Δ frdABCD	Δ frdABCD, B275 parent	This study
B275 Δ frdA	Δ frdA, B275 parent	This study
B275 Δ frdB	Δ frdB, B275 parent	This study
B275 Δ frdC	Δ frdC, B275 parent	This study
B275 Δ frdD	Δ frdD, B275 parent	This study
DFB190	Δ fliM, RP437 parent	[19]
DFB190 Δ frdABCD	Δ fliM Δ frdABCD, RP437 parent	This study
Plasmids		
pCa24N	empty vector, Cap ^R	[20]
pCa24N–FrdA	FrdA expression vector, N-terminal 6 \times His, pCa24N based, Cap ^R	[20]
pCa24N–FrdB	FrdB expression vector, N-terminal 6 \times His, pCa24N based, Cap ^R	[20]
pCa24N–FrdC	FrdC expression vector, N-terminal 6 \times His, pCa24N based, Cap ^R	[20]
pCa24N–FrdD	FrdD expression vector, N-terminal 6 \times His, pCa24N based, Cap ^R	[20]
pEWG1	FliG expression vector, N-terminal 10 \times His, Amp ^R	[11]
pCa24N–FliG _{CW}	FliG(Δ 169–171) expression vector, N-terminal 6 \times His, pEWG1 based, Amp ^R	This study
pH 3	FrdABCD, <i>frdABCD</i> operon with natural <i>frd</i> promoter, pBR322 derivative, low copy number plasmid Amp ^R	[21]
pH 3–Dendra2–FrdABCD	Dendra2–FrdABCD expression vector (FrdA is linked to Dendra2) with natural <i>frdABCD</i> promoter, pH 3 based, Amp ^R	This study
pH 3–Dendra2	Dendra2 expression vector with natural <i>frdABCD</i> promoter, pH 3 based, Amp ^R	This study
pFliM _{WT} –mPlum	FliM–mPlum expression vector with arabinose P _{BAD} promoter	This study
pFliM _{CW} –mPlum	FliM(R60C)–mPlum expression vector with arabinose P _{BAD} promoter	This study

corresponding KEIO deletion strains (Table 2), were used to infect B275. All the insertions were verified by PCR amplification and sequencing.

Plasmids

The plasmid pH 3–Dendra2–FrdABCD was constructed in two ligation steps. First, the plasmid pH 3 (Table 2) was amplified by PCR using primers 5'-ACAGCGGTACCGTCGTGCAAACCTTTCAAGCCGATC-3' and 5'-ATTCCTCCAGATTGTTTTTATCCCACAGCCACGTAC-3'. This added a KpnI site 5' to *frdA* gene of the resulted linear DNA, which ends immediately downstream to the *frdABCD* promoter. Next, the sequence encoding for Dendra2 protein was amplified from pDendra2 Vector [Clontech Laboratories (Takara Bio Company), Mountain View, CA, USA], with the primer sequences 5'-GTGAA CACCCCGGAATTAACCTGATCAAG-3' and 5'-ACAGCGGTACCTGCAGGCGGCCCTCTTGTA-3', while the start codon was replaced by GTG start codon as in *frdA* gene and its stop codon was removed. The amplified PCR fragment was digested with KpnI, leaving one end blunt, and ligated with KpnI predigested plasmid from the previous step, inserting *dendra2* gene downstream the *frd* promoter and in-frame with the *frdA* gene. The functional status of Dendra2–FrdA fusion protein was verified by the

ability of the DFB190 Δ frdABCD mutant, transformed with this plasmid, to grow under anaerobic conditions. The plasmid pH 3–Dendra2 was constructed similarly but the stop codon was added at the 5' of the *dendra2* reverse primer.

The plasmid pCa24N–FliG_{CW} was prepared by deleting residues 169–171 [26,56]. This was done by PCR amplification of the plasmid pCa24N–FliG [20] with the primers 5'-CTGGCGGAGCTGACCGAAGTACTGAATGG-3' and 5'-TGCACGCCGCCAAAGGTGGCGATA-3', which, upon ligation of the amplified sequence, resulted in a plasmid containing FliG without the sequence encoding for aforementioned residues.

To construct the plasmids pFliM_{WT}–mPlum and pFliM_{CW}–mPlum, the sequence encoding for mPlum fluorescent protein was amplified by PCR from pRSET–mPlum (Invitrogen™, Thermo Fisher Scientific, Waltham, MA, USA) with the primers 5'-ACAGCAGATCTGGCATGGTGAGCAAGGGCGAGGAGGTC-3' and 5'-ACGCAAGCTTAGGCGCCGGTGGAGTGGC-3', which add a BglII site upstream and stop codon and HindIII site downstream of the mPlum coding sequence. Next, the amplified PCR fragment was digested with BglII and HindIII and ligated with similarly digested plasmids pMWT (encoding *Salmonella* wild-type FliM) and pMCW1 [encoding *Salmonella* clockwise-locked FliM(R60C)] [57]. The functional

status of FliG-mPlum fusion protein was verified by the ability of the DFB190 strain (Table 2), transformed with this plasmid, to form expanding rings on Tryptone broth semi-solid agar plates. Expression level of FliM-mPlum fusion protein in DFB190 Δ frdABCD was optimized by modulating the D-arabinose concentration in order to provide maximal restoration of motility during the anaerobic growth (2.5–3.5 μ M arabinose). Transforming DFB190 Δ frdABCD with FliM_{CW}-mPlum resulted in cells rotating clockwise 90% of the time.

All genetic constructs were confirmed by DNA sequencing. Fluorescence of the fusion proteins was verified by microscopic visualization. All FliM-mPlum fusions were found non-fluorescent due to the anaerobic conditions (the maturation process that makes mPlum fluorescent requires oxygen [58,59]).

Growth and media

All bacterial cultures for protein isolation and genetic manipulations were grown in Luria-Bertani broth [60] at 37 °C. For determining the direction of flagellar rotation of aerobically grown cells, overnight cultures were diluted 1:100 in fresh Tryptone broth and grown at 30 °C to OD₆₀₀ of ~0.5. For determining the direction of flagellar rotation of anaerobic cells and their control aerobic cells, overnight cultures were diluted 1:50 in fresh M9 salt medium supplemented with glycerol (20 mM), the required amino acids (1 mM each), and, where indicated, with sodium fumarate (40 mM) or L-arabinose (40 mM). For anaerobic conditions, the tube was filled to the end and tightly closed, then shaken at 30 °C and 100 rpm. Conversion of colored resazurin (1 μ g/ml) to colorless resorufin in a parallelly grown wild-type culture was used as an indication for reaching anaerobic conditions. For super-resolution experiments, the anaerobic bacterial cultures were similarly grown at 30 °C, but the M9 salt medium was supplemented with Casamino acids (0.5%), vitamin B₁ (0.5 mg) and sodium fumarate (40 mM). For overexpression of Frd proteins, the overnight cultures were diluted 1:100 in fresh Tryptone broth supplemented with fumarate (40 mM) and grown aerobically at 30 °C to OD₆₀₀ of ~0.1, followed by further 3-h incubation with 0–1000 μ M IPTG.

Protein isolation

For lysis, the bacterial cultures were suspended in buffer containing 50 mM Tris (pH 8), 0.5 M NaCl, 5% glycerol, DNase, lysozyme, 1 mM PMSF and EDTA-free cComplete™ Protease Inhibitor cocktail and lysed by sonication. All the proteins were concentrated by ultrafiltration (Amicon Ultra centrifugal filter units, Sigma-Aldrich, St. Louis, MO, USA).

His-tagged-FrdA

The pCA24N-FrdA vector was expressed in RP437 Δ sdh Δ frd (Table 2) by induction with 0.2 mM IPTG overnight at 15 °C. His-FrdA was captured on a HisTrap_FF-crude_5ml column (GE Healthcare Life Sciences) equilibrated with 50 mM Tris (pH 8.0), 0.5 M NaCl and 20 mM imidazole and eluted with the same buffer containing 0.5 M imidazole. The pooled FrdA-containing fractions were loaded onto a HiLoad_16/60 Superdex column (GE Healthcare Life Sciences) equilibrated with PBS. The purification was carried out by the Dana and Yossie Hollander Center for Structural Proteomics (ISPC).

His-tagged-FliG

10xHis-FliG_{WT} and 6xHis-FliG_{CW} were expressed from the plasmids pEWG1 and pCa24N-FliG_{CW}, respectively, after being transformed into strain BL21(IDE3) pLysS (Table 2). The expression was induced with IPTG (0.2–0.4 mM) at 15 °C overnight. The proteins were purified successively by HisTrap-HP affinity chromatography (GE Healthcare Life Sciences), HiLoad 16/60 Superdex 75 prepgrade chromatography (GE Healthcare Life Sciences) both in PBS, and Tricorn Q 10/100GL chromatography (GE Healthcare Life Sciences) using 20 mM sodium phosphate (pH 7.2) as a binding buffer. The proteins were eluted with a linear gradient to the same buffer containing 1 M NaCl. Some batches of 6xHis-FliG_{CW} were purified by HisTrap-HP affinity chromatography only.

Determination of the direction of flagellar rotation by tethering assay

The direction of flagellar rotation was determined by the tethering assay [23]. Bacterial cultures were transferred several times through a syringe needle to shear the flagella, washed twice in motility buffer [10 mM KPi (pH 7.0), 0.1 mM EDTA and 0.1 mM L-methionine] and immobilized on a microscope coverslip using an anti-flagellin antibody. The coverslip was placed in a flow chamber, and the rotation of the tethered cells was monitored with a phase-contrast microscope (Zeiss, Oberkochen, Germany) and recorded using uEye UI-2410-M-GL camera (IDS Imaging Development Systems GmbH; Obersulm, Germany) at 75 frames/s. The recordings were analyzed with a custom-made software [61].

MST

MST measurements were performed on a Monolith NT 115 instrument (NanoTemper Technologies GmbH, München, Germany). All measurements were performed at 90%–60% LED power and 60%

MST power with 30-s laser-on time and 5-s laser-off time. His-tagged-FrdA (~95% purity) was used at the concentration of 15 μM . The natural fluorescence of the flavin cofactor was utilized to follow the migration of the protein. His-tagged-FliG_{wt} or His-tagged-FliG_{cw} (~95% purity) was titrated in 1:1 dilutions beginning with 138 or 319 μM . The experiments were performed in PBS buffer with 0.05% Tween and measured in standard capillaries (NanoTemper Technologies). All binding reactions were incubated for 10 min at room temperature before they were loaded onto the capillaries. Curve fitting was carried out by MicrocalTM Origin version 6 software (Northampton, MA, USA).

In-gel digestion and protein identification by LC-ESI-MS/MS

Protein bands were excised from a Coomassie-stained SDS gel. They were subsequently reduced, alkylated and digested with bovine trypsin at a concentration of 12.5 ng/ μl in 50 mM ammonium bicarbonate at 37 °C, as described [62,63]. The peptide mixtures were extracted with 80% CH₃CN and 1% CF₃COOH, followed by evaporation of the organic solvent in a vacuum centrifuge. The resulting peptide mixtures were reconstituted in 80% formic acid and immediately diluted 1:10 with Milli-Q water before analysis. Liquid chromatography–tandem mass spectrometry (LC–MS/MS) was performed using a 15-cm reversed-phase fused-silica capillary column (inner diameter, 75 μm) made in-house and packed with 3 μm ReProSil-Pur C18AQ media (Dr. Maisch GmbH, Ammerbuch-Entringen, Germany). The LC system, an UltiMate 3000 (Dionex, Sunnyvale, CA, USA), was used in conjunction with a LTQ Orbitrap XL (Thermo Fisher Scientific, Bremen, Germany) operated in the positive ion mode and equipped with a nano-electrospray ion source. Peptides were separated with a 2-h gradient from 5% to 65% acetonitrile (buffer A, 5% acetonitrile, 0.1% formic acid and 0.005% TFA; buffer B, 90% acetonitrile, 0.2% formic acid and 0.005% TFA). The voltage applied to the union to produce an electrospray was 1.2 kV. The mass spectrometer was operated in the data-dependent mode. Survey mass spectrometry scans were acquired in the Orbitrap (Thermo Fisher Scientific, Bremen, Germany) with the resolution set to a value of 60,000. The seven most intense ions per scan were fragmented and analyzed in the linear ion trap. Raw data files were searched with MASCOT (Matrix Science, London, UK) against the Uniprot *E. coli* database. Search parameters included a fixed modification of 57.02146 Da (carboxyamidomethylation) on Cys, and variable modifications 15.99491 Da (oxidation) on Met and 0.984016 Da (deamidation) on Asn and Gln. The search parameters also included: maximum two missed cleavages; initial precursor ion mass tolerance, 10 ppm; fragment ion mass toler-

ance, 0.6 Da. Samples were further analyzed in Scaffold (Proteome software, Portland, OR, USA).

Super-resolution microscopy

The cells were first optimized with respect to growth conditions and fixation procedure by comparing the photoconversion efficiency of Dendra2 at each set of conditions before and after fixation, as recommended [64]. At the chosen conditions (see above for growth and below for fixation), the fluorescence intensity and the photoconversion efficiencies were similar before and after fixation. Immunostaining of FliM–mPlum fusion protein using anti-mCherry monoclonal antibody and Alexa Fluor 647-conjugated anti-mouse antibody was performed according to the manufacturers' instructions as follows. Immediately after air exposure, anaerobically grown bacterial cultures were precipitated and resuspended in 4% (v/v) paraformaldehyde in PBS, followed by 5-min incubation with shaking for fixation and permeabilization. The cells were spun down (470 g, 10 min, room temperature) and then washed with PBS (two centrifugations at 470 g for 3 min followed by centrifugation at 1300 g for 3 min). The cells were then blocked with 3% BSA in PBS for 30 min at room temperature with gentle rocking, followed by aspiration of the blocking buffer and 1-h incubation with the primary antibody anti-mCherry (5 $\mu\text{g}/\text{ml}$) in 1% BSA in PBS with gentle rocking. Following three washes in PBS (twice at 470 g and once at 1300 g, as above), the cells were incubated in the dark with gentle rocking for 45 min with the secondary antibody Alexa Fluor 647-conjugated anti-mouse antibody (1.4 $\mu\text{g}/\text{ml}$) in PBS, and washed with PBS three times as above. The cells were then immobilized for 20 min on pre-washed polylysine (0.2% w/v)-coated coverslips, gently washed three times with 0.5 ml PBS, and covered with freshly prepared imaging buffer [65], which contains 10 mM NaCl, 10% (w/v) glucose, 5 mM cysteamine, and oxygen scavengers (7 μM glucose oxidase and 56 nM catalase) in 50 mM Tris (pH 8.0). Super-resolution images were recorded with Vutara SR 200 microscope (Bruker, Billerica, MA, USA). Images were recorded using a 60 \times , NA 1.2 water immersion objective (Olympus, Tokyo, Japan) and Evolve 512 EMCCD camera (Photometrics, Teledyne Technologies, Tucson, AZ, USA) with the gain set at 50. Combined PALM [31] and STORM [66] imaging was performed in two steps. First, STORM was recorded for 5000 frames, while the photoblinking of Alexa Fluor 647 was induced with a 640-nm laser, set to 1.2 kW/cm². Next, PALM was recorded. Photoswitching of Dendra2 was activated by a 405-nm laser, whose intensity was stepwise and slowly increased. The intensity of the exciting 561-nm laser was kept constant at 20 kW/cm². All the recordings were carried out at 10 frames/s. Acquisition was carried out until all Dendra2 molecules were

converted and detected (15,000–20,000 frames). Data were analyzed by Vutara SRX software (Bruker). To avoid over-counting in PALM records, single-emission events detected in sequential frames and localized within the range of the resolution of a single molecule were grouped as one detection event. In addition, to avoid over-counting because of photoblinking of single Dendra2 molecules, clusters of molecules whose time of appearance was too close to be individually detectable by the software were eliminated manually. Over-counting due to altered blinking behavior of Dendra2 in the imaging buffer was ruled out by preparing control samples for imaging in PBS [32] and finding that the number of molecules per bacterium in PBS (270 ± 20 , mean \pm SEM; $n = 21$ cells) was not different from the number of molecules detected in imaging buffer (cf. Table 1). Quantitative analysis of spatial distribution of Dendra2 molecules was performed using Image-Based Algorithm for two dimensions of Vutara SRX Software. Minimal cluster count was set to be seven molecules. Minimal density threshold was separately set for each bacterium as three times the average density of Dendra2 in this cell. Clusters identified in this way were in good agreement with those detected visually. Motors were identified by groups of at least 12 single localizations forming two-dimensional round or ring-like shapes. FrdA molecules were considered interacting with the motor if found within a 0.3×0.3 - μm square surrounding the center of the motor. This area was calculated on the basis of the diameter of FliM ring (~ 47 nm), the dimensions of the proteins involved (FrdA, 70 Å; Dendra2, 29.6 Å, according to the crystal structures available at the Protein Data Bank), and the resolution (35 nm).

Computational methods

MD (50–100 ns) [67–69] and detailed conformational analysis were performed for unbound *E. coli* FliG models. Homology models of *E. coli* FliG were built from the indicated structural data by using Modeller v8 [70] as published in ModBase [71]. In our computational modeling of possible interactions between FrdA(BCD) and FliG, we applied a rigid-body docking protocol by implementing Potential Smoothing and Search (PSS) algorithm, as described [72,73], to all conformational states of FliG identified in the MD study. For the FliG–FrdA complexes, we performed multiple short-time MD runs (10–20 ns) for refinement only. Docking was used in combination with Poisson–Boltzmann electrostatics methods ([74,75] and references cited therein). Multi-step binding process in the FliG–FrdA system was modeled as described [75].

Statistical analysis

All statistical analyses were carried out by Instat software (GraphPad Software, San Diego, CA).

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jmb.2019.08.001>.

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Abbreviations used:

ARM, armadillo repeat motif; FliG_{ccw}, FliG in its counter-clockwise state; FrdABCD, fumarate reductase; MD, molecular dynamics; MST, microscale thermophoresis; PALM, photoactivated localization microscopy; SdhCDAB, succinate dehydrogenase.

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