

A molecular brake, not a clutch, stops the *Rhodobacter sphaeroides* flagellar motor

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Many bacterial species swim by employing ion-driven molecular motors that power the rotation of helical filaments. Signals are transmitted to the motor from the external environment via the chemotaxis pathway. In bidirectional motors, the binding of phosphorylated CheY (CheY-P) to the motor is presumed to instigate conformational changes that result in a different rotor-stator interface, resulting in rotation in the alternative direction. Controlling when this switch occurs enables bacteria to accumulate in areas favorable for their survival. Unlike most species that swim with bidirectional motors, *Rhodobacter sphaeroides* employs a single stop-start flagellar motor. Here, we asked, how does the binding of CheY-P stop the motor in *R. sphaeroides*—using a clutch or a brake? By applying external force with viscous flow or optical tweezers, we show that the *R. sphaeroides* motor is stopped using a brake. The motor stops at 27–28 discrete angles, locked in place by a relatively high torque, approximately 2–3 times its stall torque.

Controlling the output of molecular motors is critical for many cellular processes governing cell growth, division, and transport. In addition, regulating their force-generating capabilities is a prerequisite for the operation of useful bionanodevices (1). Here, by examining the mechanics required to control the *Rhodobacter sphaeroides* flagellar motor, we provide an example of how molecular machines are controlled at the nanoscale.

The bacterial flagellar motor [reviewed in ref. 2 and 3] is a molecular machine made from several hundred proteins embedded in the cell envelope. It drives the rotation of an extracellular helical filament, enabling bacteria such as *Escherichia coli* to swim. The motor is powered by the cell's ion-motive force, usually maintained by electron transport (respiration/ photosynthesis). The flow of ions through peptidoglycan-bound stator complexes, down an electrochemical gradient into the cytoplasm, is responsible for rotor rotation via electrostatic interactions at the rotor-stator interface (4). Stator units can independently engage/disengage from the rotor (5), resulting in stepwise changes in speed (6).

By controlling the output of the motor, *E. coli* can accumulate in areas favorable for their survival (7, 8). External conditions are sensed and transmitted via the chemotaxis pathway to the motor. This simple pathway controls the intracellular concentration of phosphorylated-CheY (CheY-P). In the absence of positive stimuli, CheY-P molecules bind to the rotor switch complex, which consists of approximately 26 copies of FliG, approximately 34 copies of FliM, and approximately 100 copies of FliN. Elevated CheY-P levels increase the probability that a change in direction of motor rotation occurs (9), causing a free-swimming bacterium to tumble (10). When motor rotation in its original direction recommences, the cell swims on a new trajectory. By controlling the frequency of motor switches, delaying tumbles when swimming up attractant gradients, cells achieve a biased random walk.

Many other bacterial species, such as the metabolically diverse α -proteobacterium *R. sphaeroides*, have more complicated chemosensory systems (reviewed in ref. 7). *R. sphaeroides* possesses

multiple chemotaxis protein homologues, whose interactions are limited by differences in subcellular localization, expression level, and selective phosphotransfer (reviewed in ref. 11). A minimum of 2 CheY homologues can support chemotaxis in *R. sphaeroides*, CheY₆ plus either CheY₃ or CheY₄. Although CheY₆-P can stop the motor, it cannot mediate chemotaxis alone (12). Unlike *E. coli*, which has a switching motor, these CheYs control the behavior of a single unidirectional proton-driven flagellum termed fla1 (13). The cell is reorientated by relaxation of the flagellar filament from helical to coiled form when the motor stops (14).

Here, the mechanism used by the chemotaxis pathway to stop motor rotation was investigated. Data were obtained by either monitoring motor rotation at a high temporal and spatial resolution using back-focal-plane (BFP) interferometry or by manipulating motors using viscous flow or an optical trap.

We found that the motor rotates in the counterclockwise (CCW) direction at similar torques and speeds to *E. coli*, and that without chemotactic stimulation (Δche strain), the motor rotates continuously. We also showed that the chemotaxis system does not control motor speed (termed chemokinesis) as previously reported (15, 16).

By applying external force to chemotactically stopped motors, we discovered that the *R. sphaeroides* motor stops using a brake. We estimated that the motor can resist approximately 2–3 times more torque when stopped than it generates when rotating at high load. Furthermore, we found that beads are held at 27–28 discrete angles around the motor, in agreement with the 26 steps per revolution previously observed using a Na⁺-driven chimeric flagellar motor in *E. coli* (17).

Results

Characterizing Motor Output. The output of the *R. sphaeroides* motor has previously been analyzed using free-swimming or tethered cells. The *R. sphaeroides* bead assay (Fig. 1A) was developed to improve the resolution of stop events and investigate motor speeds/torques.

Fig. 1B (black trace) shows a typical speed-time trace from a wild-type cell; rotation is punctuated by stop events. Beads (0.83- μ m diameter) were monitored by BFP interferometry, sampled at 10 kHz and 200-point median filtered (sliding window in 1-point increments); giving a resolution of 20 ms at worst.

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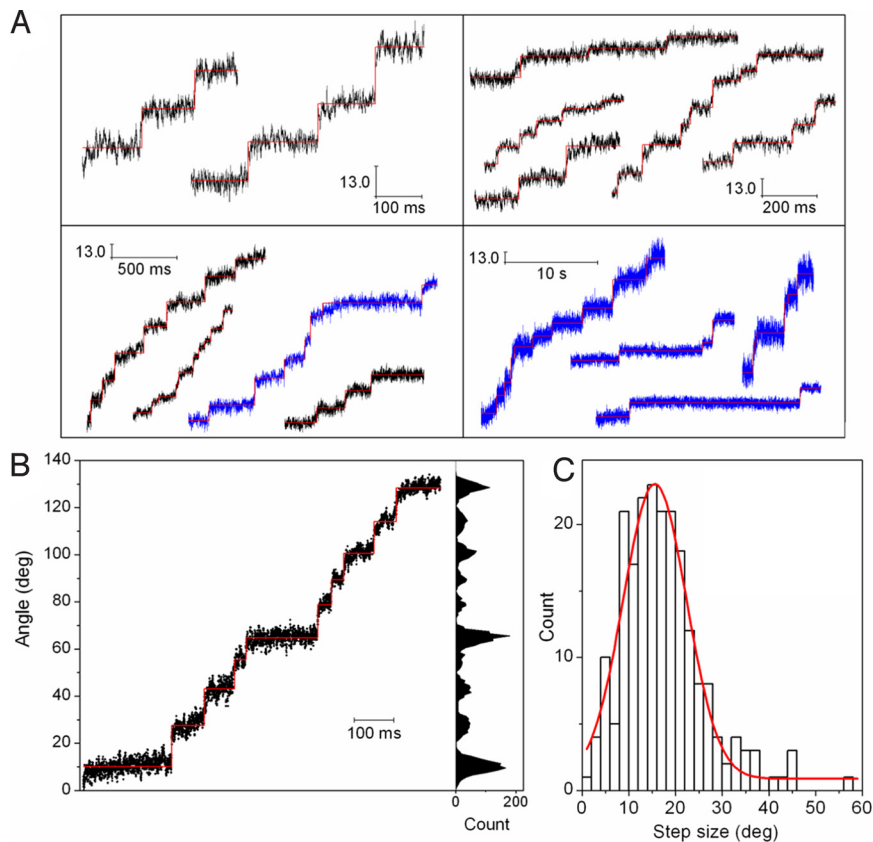


Fig. 4. Step analysis. (A) Examples of angle-time traces showing steps during stop events from wild-type (blue) and WT+CheY6 (black) cells. Wild-type data were obtained with either a 632-nm laser (10-kHz sampling rate) or a 1,064-nm laser (2-kHz sampling rate); WT+CheY6 motors were recorded with a 1,064-nm laser at 5-kHz sampling rate (in all cases 0.83- μ m beads were used). The output of a step-finding algorithm is superimposed (red). (B) A particularly clear stepy angle-time trace and its accompanying histogram of dwell angles (bin size = 1 $^\circ$). (C) The histogram of step sizes obtained by the step-finding algorithm ($n_{\text{steps}} = 5,361$; histogram bin size = 2 $^\circ$). The peak (Gaussian fit) at $13.0 \pm 4.5^\circ$ corresponds to approximately 27–28 steps per revolution.

the motor rotates continuously in the CCW direction unless signaled to stop via phosphorylated CheYs. Only the motor stop frequency, and not its speed, is controlled by the chemotaxis system. Stops began and ended within approximately 50 ms, close to the resolution time of our measurements using the bead assay (≈ 20 ms). If the motor were to lock instantaneously, the tether would relax exponentially from an initial twist angle $\theta = T/\kappa_\theta$, to $\theta = 0$, with a time constant $\tau = f/\kappa_\theta$; where T is the motor torque, f the drag coefficient and κ_θ the tether stiffness. The initial speed of this relaxation would be $\omega = \kappa_\theta \theta / f = T/f$, the same as the speed at which the motor was running. With our estimates of T , f , and κ_θ we obtain $\theta = 16 \pm 8^\circ$, with time constants $\tau = 0.4 \pm 0.2$ and 190 ± 80 ms for 0.83- μ m beads and tethered cells, respectively. Thus, slow forward rotation at the beginning of some stops in tethered cell traces could be explained by tether relaxation, but in most cases, the movement is larger than 16 $^\circ$ and slower than 190 ms. Tether relaxation cannot explain the observation of slow ($\ll 1$ Hz) stepy forward rotation lasting several seconds in 0.83- μ m bead traces (Fig. 3).

By applying external torque to the motor using 2 independent techniques, we found that during chemotactic-induced stops, motors resisted rotation indicating a braking mechanism. Using the cell body as a handle for an optical trap we estimated that the brake can resist a torque up to 2–3 times the stall torque of the motor. In contrast, recently Blair et al. showed that *Bacillus subtilis* uses a molecular clutch to disable its flagella during biofilm formation (19). Tethered cells overexpressing the clutch-initiating protein, EspE, behaved as although they were unpowered, analogous to a strain lacking torque-generating units

(Δ motAmotB). The difference may reflect requirements for long-term stopping during biofilm formation versus fast reversible control needed for chemotaxis.

Consistent with a brake mechanism, beads attached to flagellar stubs remain at a fixed angle for several seconds during stops, exhibiting only small fluctuations (significantly smaller than the step size of $\approx 13^\circ$). Some motors showed slow stepping rotation during stops, stopping at 27–28 discrete angles around the motor. This is in agreement with the approximately 26 steps per revolution observed previously using a Na $^+$ -driven chimeric flagellar motor in *E. coli*, measured at low sodium-motive force and with controlled expression of a small number of torque-generating units (17).

In bidirectional motors, the binding of CheY-P to the switch complex is presumed to instigate a conformational change in FliM, which in turn affects the orientation of the charged ridge in the C terminus of FliG (20, 21). The different electrostatic interactions resulting from this altered rotor-stator interface [between FliG and the cytoplasmic loop of the stator (MotA)], are thought to be responsible for rotation in the alternative direction (22).

Similarly, this altered rotor-stator interface may stop the *R. sphaeroides* motor: Conformational changes during a stop must lock the motor, either in 1 of the states through which it normally cycles during torque generation or in a state that is only present during a chemotactic stop. In either case, the observed steps in angle could be due either to rapid transitions between the stopped and cycling states or to a temporary reduction in the amplitude of the locking potential that permits a thermally

activated jump from 1 minimum to the next. The exaggerated stepping behavior of a strain overexpressing CheY₆, which can bind in a phosphorylated or unphosphorylated state (23) is compatible with both stepping mechanisms.

The similar periodicity of steps observed in chemotactically stopped *R. sphaeroides* motors with a full complement of stators and in chimeric motors with few stators present (17), indicates that the step size is governed solely by the periodicity of the rotor ring. The locked state is described by an interaction potential between the rotor and each stator, and these potentials will each share the approximately 27-fold periodicity of the rotor. Thus, the rotor will step in a combined potential that is the sum of the individual rotor-stator potentials. The amplitude of the combined potential depends upon both the number and relative phase of the individual potentials, which might in part account for the observed variability in the locking strength at $T_{max} = 0.5\text{--}3.2T_{stall}$. However, the periodicity of the combined potential stays the same irrespective of the number and relative phase of the individual potentials, which might explain why step size is independent of the number of stators, as observed. If the step size is found to be independent of the number of stators in a running motor and in the locked motors observed in this study, an alternative explanation could be that the stators are separated by angles close to an integer multiple of the rotor periodicity (24).

At this stage, the potentials discussed above are hypothetical. In practice they are likely to be because of interactions between the charged residues that have been shown to affect motor function in *E. coli* (K264, R281, D288, D289, and R297 in FliG, and R90 and E98 in MotA) (4), all of which are conserved in *R. sphaeroides*. Analysis of strains harboring point mutations at these sites may reveal details of the locking mechanism.

Our estimated tether stiffness ($\kappa_{\theta} = 3,800 \pm 2000$ pNm/rad²) is an order of magnitude higher than that of *E. coli*, which was attributed primarily to flexibility of the hook (25, 26). This is consistent with electron micrographs showing that the *R. sphaeroides* hook is straight, not curved like *E. coli* (27). Recent models of the flagellar motor have proposed elastic links between stators and the cell wall, each with a stiffness of approximately 200 pNm/rad² (24, 28). Approximately 11 such stators in parallel (6) would have a combined stiffness within the range of our estimates for the *R. sphaeroides* tether, but ≈ 10 times stiffer than the *E. coli* tether. Thus, it is possible that the *R. sphaeroides* hook is stiffer than or comparable to the stator springs, and that the latter dominate the tether stiffness. In this case, variation in the number of stators would contribute to variability in tether stiffness. However, until direct evidence is obtained for the existence of compliant stator springs these considerations remain speculative.

High-resolution measurements of rotating beads coupled with the application of external torque have revealed how the *R. sphaeroides* flagellar motor is stopped. This provides an insight into how nature uses fundamental mechanical principles to operate motors effectively at the nanoscale.

Methods

Strains. *R. sphaeroides* strains (Table S1) were grown from frozen stocks (made from single colonies) in succinate medium (29) at 30 °C to an OD of 0.45–0.65, either aerobically with shaking without illumination, or when specified, photoheterotrophically in an anaerobic cabinet with illumination at 50 $\mu\text{mol}/\text{m}^2/\text{s}$. Media were supplemented with 25 $\mu\text{g}/\text{mL}$ kanamycin and 1 mM IPTG (overnight induction) when using pIND4. Although the *R. sphaeroides* genome encodes 2 sets of *fla* genes (30), only the *fla1* set has ever been observed under these conditions.

Bead and Tethered Cell Assay. Carboxy-modified latex spheres (4% wt/vol, Interfacial Dynamics Corp.) were covalently coupled to goat anti-rabbit IgG (Abcam) in 50 mM Mes (pH 6) plus 50 mg/mL EDAC (Invitrogen). Cells with truncated filaments were immobilized to the glass surface via polylysine (17). The filament stubs were coated with anti-flagellin rabbit IgG for attachment to antibody labeled beads. In the tethered cell assay, cells were immobilized to the coverslip via anti-flagellin antibody as described in ref. 13. All experiments were performed in 10 mM Na-Hepes, pH 7.2, except where stated, when Hepes was supplemented with 1 mM sodium propionate (Sigma).

Speed, Stopping, and Running Torque Measurements. The position of rotating beads was detected by BFP interferometry (31) using a 632 nm Helium Neon or 1,064 nm Ytterbium fibre laser (18). All experiments were performed at 23 °C.

Bead angle was computed as described in ref. 17. Speeds were calculated from angle versus time traces ($d\theta/dt$) and filtered with a 200-point median filter (sliding window in 1-point increments), or where stated, by means of power spectra of the combined x and y signals (31).

For Fig. 1B, cells were selected as follows. (i) The mean speed (M_{CCW}) for each record was calculated by fitting a Gaussian to the peak in the speed histogram corresponding to intervals during which the bead was spinning. (ii) Stop events were measured from when speed decreased to 0.1 M_{CCW} to when speed increased $>0.4 M_{CCW}$. (iii) Stop events were screened by a 45° moving window and the largest number of stops registered within a 45° sector subjected to a binomial test. Cells with nonuniformly distributed stop angles were discarded.

Deceleration times were defined as the time it takes for the speed of the bead attached to the filament stub to decrease from 0.7 M_{CCW} to 0.1 M_{CCW} (or from 0.1 M_{CCW} to 0.7 M_{CCW} for acceleration times).

Torque ($T = f \cdot 2\pi \cdot M_{CCW}$) was estimated by calculating the viscous drag (f) on the bead, $f = 8 \pi \eta r^3 + 6 \pi \eta r l^2$ where η is viscosity, r is bead radius, and l is eccentricity. For the bead sizes used, contribution from the filament stub is negligible and was therefore neglected (32). The eccentricity of rotation was 126 ± 28 and 299 ± 91 nm for 0.83- μm and 1.90- μm beads, respectively, measured as described previously in refs. 6, 18.

Lock Torque Measurements. The trap and rotating tethered cells were aligned as previously described (18). T_{max} was calibrated against T_{stall} at the start of each run while keeping the trap at a fixed position, until the rotating cell body was just stalled. The laser was shuttered and a chemotactic stop induced by removal of chemoattractant. The shutter was opened and the trap used to move the cell body in a series of 45° angles (up to a maximum of 315°). The trap was once again shuttered and the motors allowed to resume rotation. The procedure was repeated with different values of T_{max} : (0.1, 0.2, 0.5, 1.0, 2.0, 3.2, 5.0, and 10.0) $\times T_{stall}$. The position of the cell body was recorded using a video camera (LCL-902K, Watec) at 50 Hz. Cells were rejected if they appeared to be sticking or took longer than 2 min (the maximum recovery time without mechanical manipulation) to recover. At $T_{max} = 10T_{stall}$, some motors appeared to break permanently (no recovery within 12 min), similar to when the *E. coli* motor is driven backwards (33).

For T_x calculation, we confirmed that the *R. sphaeroides* tether behaves like a linear spring, validating the use of the equipartition method ($\langle \kappa_{\theta} \Delta \theta^2 \rangle = k_B T$) to estimate κ_{θ} directly. In addition, κ_{θ} was estimated by fitting the relaxation angle of the cell body with an exponential of time constant (τ), where $\tau = f/\kappa_{\theta}$ (f being the drag coefficient of the tethered cell). f of each individual cell was calculated from the speed of the cell's rotation before and after external torque was applied. It was not possible to accurately estimate the rebound angle at $T_{max} = 10T_{stall}$ because at such large torques, the cell body moved from its initial centre of attachment, introducing a large error.

Steps. Step analysis was performed using a computer algorithm as described in ref. 17. In theory the peak value of 'quality factor' (as a function of the ratio of the length of the data trace and number of steps fitted) gives a fit that best represents real steps in the data (34). In practice a fit that slightly overfits the data (but still gives a high 'quality factor') was chosen (34, 17).

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