Real-Time Imaging of Fluorescent Flagellar Filaments

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Bacteria swim by rotating flagellar filaments that are several micrometers long, but only about 20 nm in diameter. The filaments can exist in different polymorphic forms, having distinct values of curvature and twist. Rotation rates are on the order of 100 Hz. In the past, the motion of individual filaments has been visualized by dark-field or differential-interference-contrast microscopy, methods hampered by intense scattering from the cell body or shallow depth of field, respectively. We have found a simple procedure for fluorescently labeling cells and filaments that allows recording their motion in real time with an inexpensive video camera and an ordinary fluorescence microscope with mercury-arc or strobed laser illumination. We report our initial findings with cells of Escherichia coli. Tumbles (events that enable swimming cells to alter course) are remarkably varied. Not every filament on a cell needs to change its direction of rotation: different filaments can change directions at different times, and a tumble can result from the change in direction of only one. Polymorphic transformations tend to occur in the sequence normal, semicoiled, curly 1, with changes in the direction of movement of the cell body correlated with transformations to the semicoiled form.

The external part of the bacterial flagellum is composed of a short proximal hook and a long helical filament. In Escherichia coli and Salmonella enterica serovar Typhimurium, the filament is a polymer of a single protein called flagellin with a molecular weight of about 50,000. The filament is linked to the hook by two other proteins and capped at its distal end by a third. The hook is driven at its base by a reversible rotary motor. For reviews, see references 24 and 29. The shape of the filament depends upon the arrangement of the flagellin monomers, which depends in turn upon the amino acid sequence of the protein, temperature, pH, ionic strength, and torsional load. The monomers form 11 protofilaments that run along the surface of a cylinder about 20 nm in diameter, twisting slightly either to the left or to the right. The monomers bind in two different ways, forming short or long protofilaments (3). A filament made up of protofilaments of both types has curvature as well as twist and is helical, with the shorter protofilaments running along the inside of the helix and the longer ones running along the outside. The elastic strain energy is minimized if protofilaments of a given type are adjacent to one another. Given this rule, 12 different polymorphic forms are possible, two of which are straight (7, 8, 16). Four of these forms, the ones that we have observed in the present work, are shown in Fig. 1.

Cells either “run” (move steadily forward) or “tumble” (move erratically in place with little net displacement). Runs are relatively long (about 1 s, on average) while tumbles are relatively short (about 0.1 s, on average). These modes alternate, allowing cells to sample different directions in space. If a run happens to carry a cell in a favorable direction, e.g., up the gradient of a chemical attractant, the probability of tumbling is reduced. This biases the cell’s random walk, enabling chemotaxis (4).

The motion of flagella on live bacteria was first seen by Ehrenberg (9), who examined species with large flagellar bundles (groups of filaments rotating in synchrony), such as Chromatium okenii. Reichert (35), using dark-field condensers of high numerical aperture, observed the motion of flagellar bundles of even the smallest bacteria. However, observation of the motion of single filaments was limited to filaments that were sheathed, as in Vibrio metschnikovii, or to filaments of cells that were nearly immotile, as in old preparations of Proteus vulgaris. The first descriptions of changes in polymorphic form were presented by Pipper and Abraham (33), who noted that wavelengths of flagellar bundles could change by a factor of 2, with the overall length of the bundle remaining constant. Given our current understanding of polymorphic transformations, the “biplicity” observed by Pipper involved transformations between normal and curly 1 (Fig. 1). A normal filament is left-handed and has a pitch of about 2.5 μm and a diameter of about 0.5 μm, while curly filaments are right-handed and have a smaller pitch and diameter. The diameter of a curly filament is so small that, to Pipper, it often appeared straight (32, 34). From 1931 onwards, Pipper used the sun as a light source, equipping his microscope with a heliostat (30). In 1946, he decided that flagella were artifacts of locomotion (31). This led to a lively debate (36). The use of the dark-field microscope to observe moving flagella, including single filaments on fully motile cells, was perfected by Macnab, who used short-arc xenon or mercury lamps of high surface brightness, attenuating the light in the blue with a 530-nm long-pass filter (21, 23). Working primarily with S. enterica serovar Typhimurium, Macnab established our current understanding of transitions between runs and tumbles. Cells run when pushed from behind by a flagellar bundle of the normal left-handed waveform, with all of the filaments turning counterclockwise (CCW [when viewed from behind the cell]). Cells tumble when the filaments turn clockwise (CW) and the bundle comes apart. The filaments “operate as a coordinated bundle that actively disperses upon reversal of the rotation sense” (22). During this dispersal, filaments undergo transformations from normal to curly, with the change propagating rapidly from the cell body outwards. “The chaotic motion of the cell body, in reaction to a number of flagella which are rotating and in transition, constitutes the tumble” (26).
Polymorphic transformations also have been observed with isolated flagellar filaments attached rigidly at one end to glass and exposed to the flow of a viscous medium (13). The torque generated by the flow tends to unwind the filament, driving normal to semicoiled or curly transformations. This change in handedness redresses the torsional stress, so that the base of the filament returns to normal. Thus, the transformations are driven cyclically.

A serious difficulty with dark-field microscopy, observed with intact cells but not isolated filaments, is flare from the cell body, which obscures the view over distances of several micrometers. This difficulty was overcome by the use of video-enhanced differential-interference-contrast (DIC) microscopy, with a short-arc mercury lamp coupled to the microscope through a fiber-optic scrambler (6). This technique allows one to see filaments all the way to the cell body, except in the direction of shear of the Nomarski prism, where a shadow obscures the view over distances on the order of 1 μm. This method was used to demonstrate torsionally induced transformations from normal to straight or from curly to straight that occur in certain mutants defective in the protein to which the filament is attached at its base (10). However, a serious drawback with DIC microscopy is its shallow depth of field, which requires that filaments be observed close to a glass surface. Both dark-field microscopy and DIC microscopy are technically demanding, particularly if one wants to visualize individual filaments.

To our delight, we found that flagellar filaments can be readily stained with amino-specific Alexa Fluor dyes, available from Molecular Probes (Eugene, Oreg.). With readily stained with amino-specific Alexa Fluor dyes, available from Molecular Probes (Eugene, Oreg.), containing T-broth (1% Trisptone [Difco], 0.5% NaCl) and grown at 35°C. A single-colony isolate was used to inoculate 10 ml of T-broth in a 125-ml flask and the culture was grown to saturation on a rotary shaker (200 rpm) at 33°C. An aliquot was diluted 1:100 into another 10 ml of T-broth and grown 4.5 h to mid-exponential phase. The motility of the culture was checked by eye with a phase-contrast microscope. The bacteria were washed three times at room temperature by centrifugation (2,000 × g, 10 min) and gentle resuspension in 10 ml of buffer (0.01 M KPO4, 0.007 M NaCl, 10−3 M EDTA [pH 7.0]). The final suspension (0.3 ml) concentrated the bacteria 20-fold.

Alexa Fluor (488, 532, 546, or 594) carboxylic acid succinimidyl ester from Molecular Probes Protein Modification kit (one vial contained about 0.4 mg from kit A-10236, A-10235, A-10237, or A-10239) or 0.25 mg of Oregon Green 514 carboxylic acid succinimidyl ester (O-6139) was dissolved in 100 μl of buffer and added to the final suspension of bacteria. Sodium bicarbonate (25 μl [1 M]) was added to shift the pH to about 7.8. The suspension was stirred gently (by gyration at 100 rpm) for 1 h in the dark at room temperature. The bacteria were washed three times with buffer containing Brij 35 (10−2% Sigma, St. Louis, Mo.). This detergent was added to prevent the labeled cells from sticking to the walls of the Corex glass centrifuge tube. The final suspension (2 ml) diluted the bacteria fourfold. S. enterica serovar Typhimurium and a motile Streptococcus strain also were labeled by this procedure. Labeling of S. enterica serovar Typhimurium worked well, but with Streptococcus, motility was lost.

Visualization of slides. The suspension of labeled bacteria was diluted between 25- and 50-fold in buffer containing 10−9% Brij 35 and 0.1 M glucose (Sigma). Microscope slides and coverslips were used out of the box. About 50 μl was sealed between the coverslip and slide within a thin ring of Apeizom M grease (Fisher Scientific, Pittsburgh, Pa.). The coverslip was sealed carefully to eliminate air bubbles and then squeezed to form a layer about 50 μm thick. Samples were used immediately and for a period up to about 1 h. Through respiration, oxygen is used up. This reduces phototoxic effects, and glucose supports motility anaerobically (1).

Acquiring images. Cells were observed at room temperature with a Nikon Diaphot 200 epifluorescence microscope equipped with a shuttered black-and-white CCD camera (0.09-μlx sensitivity; V1070; Marshall Electronics, Culver City, Calif.) with Gamma 0.45 and gain maximum. Imagery was enhanced with a x60 objective (Nikon PlanApo 60/1.4 oil DM) and a x5 camera relay lens. Cells were illuminated either with a 100-W mercury arc with cube sets (Chroma Technology, Brattleboro, Vt.) 31007 fluorescein isothiocyanate (for Alexa Fluor 488 and B; for Alexa Fluor 546), 31003 phycoerythrin R and B (for Alexa Fluor 546), or 31004 Texas Red (for Alexa Fluor 594) or with a 514-nm-wavelength argon-ion laser (Stable 2017; Spectra-Physics, Mountain View, Calif.) with cube sets C5740 (for Alexa Fluor 5XX: excitation filter, D514/10; dichroic mirror, 527 DCLP; emission filter, E535LP). When the mercury arc was used, the camera was run with the shutter off, i.e., with an exposure time of 17 ms. The laser source was in the standard epifluorescence configuration (with parallel rays at the object plane), except that only the field of view of the video camera was illuminated (a circle about 60 μm in diameter). Laser power at the back focal plane of the objective was 100 to 300 mW. The laser was strobed at 60 Hz with an exposure time of 0.2 ms. This was done by inserting a slotted wheel in the laser beam and driving the wheel with a synchronous motor phase locked to the video vertical sync pulse. The shutter speed of the camera was set at 1,500 s, and the laser pulse was centered on this 2-ms window. For convenience, brightness and contrast were adjusted with an image processor (C-5510; Hamamatsu, Bridgewater, N.J.). Images of cells were captured at video rates (60 fields per s) in real time directly from the image processor by a G3 Power Macintosh (Apple Computer, Cupertino, Calif.) equipped with an LG-3 video capture board using Scion Image (v. 1.62c; Scion, Frederick, Md.).

Analyzing images. All measurements were made on a 17-in. Apple Studio Display monitor using Scion Image, with distances calibrated with an objective micrometer. Filaments were counted on cells stuck by their bodies to the coverslip and de-energized by exposure to light or by addition of the uncoupler FCCP [carbonyl cyanide p-(trifluoromethoxy)phenylhydrazone] (10−6 M; DuPont deNemours, Wilmington, Del.). Sticking was promoted by omitting the detergent from the final cell suspension. The filaments continued to display Brownian motion; their images were averaged for 1 s. Measurements of diam-
eter, pitch, and length of individual filaments were made on the same data set. Calculations of curvature and twist were from equation 1 of Calladine (7). Images and results were stored in a FileMaker Pro 4.1 database (FileMaker, Inc., Santa Clara, Calif.) and exported for graphing to Kaleidagraph 3.08 (Synergy Software, Reading, Pa.).

For freely motile bacteria, images were deinterlaced, and then selected areas were enlarged digitally ×2.5 to ×10. The numbers in the figures refer to fields, i.e., to images obtained at 60 Hz. In some figures, only alternate fields are shown. Some cells entered the field of view, tumbled, and then left the field of view, remaining in focus for the entire series of events. The trajectories of their cell bodies were traced, and the deflections generated by the tumbles were noted. Flagellar waveforms were identified by eye or, if difficult to discern, by measurements of diameter and pitch. Speeds of swimming cells were determined from the distance traveled per video field. The direction of wave propagation of filaments or bundles could not be determined, since the image acquisition rate was lower than flagellar rotation rates. Results were stored in a FileMaker Pro 4.1 database and exported for graphing to Kaleidagraph 3.08. Images were prepared for publication (but unenhanced) with Adobe Photoshop 5.5. Movie files are available at http://www.rowland.org.

RESULTS

Labeling. We tried four Alexa Fluor dyes, as well as Oregon Green. The name for each dye carries a number that corresponds to the peak of its excitation spectrum (in nanometers). With Alexa Fluor 488, both filaments and cell bodies were brightly labeled. Bacteria could be watched for a minute or more without substantial loss in brightness. However, excitation was in a spectral region that elicits a repellent response and readily de-energizes cells (21). As a result, the bacteria were in a spectral region that elicits a repellent response more without substantial loss in brightness. However, excitation was in a spectral region that elicits a repellent response and readily de-energizes cells (21). As a result, the bacteria tended to avoid the illuminated region and stopped swimming once trapped in the light spot. The number of filaments per cell (mean ± standard deviation, [SD], 2.78 ± 1.60) and the filament length (5.6 ± 2.9 μm) were smaller than those for the other dyes, presumably because the filaments were made brittle by the dye and/or the light.

Oregon Green 514 labeled filaments and cell bodies to a similar extent as Alexa Fluor 488, but it bleached more readily. Alexa Fluor 488, both filaments and cell bodies were brightly labeled. Bacteria could be watched for a minute or more without substantial loss in brightness. However, excitation was in a spectral region that elicits a repellent response and readily de-energizes cells (21). As a result, the bacteria tended to avoid the illuminated region and stopped swimming once trapped in the light spot. The number of filaments per cell (mean ± standard deviation, [SD], 2.78 ± 1.60) and the filament length (5.6 ± 2.9 μm) were smaller than those for the other dyes, presumably because the filaments were made brittle by the dye and/or the light.

Alexa Fluor 532 labeled filaments brightly and the cell bodies less brightly, so filaments could be seen as they passed over the cell body to form a bundle. The number of filaments per cell (3.37 ± 1.59) and the filament length (7.3 ± 2.4 μm) were larger than those for any other dye. Alexa Fluor 532 was developed for 532-nm lasers (Nd:YAG frequency doubled), which powers necessary for strobable illumination are quite expensive. Fortuitously, we already had a high-power argon-ion laser, which when turned to 514 nm worked well: the absorbance of Alexa Fluor 532 at 514 nm is about half that at 488 nm. This combination was used for most of the work reported here. There was no repellent response, so unless cells happened to tumble spontaneously while illuminated, they simply swam through the light spot without deflection or change in speed. The swimming speed of the cells studied with this dye was 30 ± 12 μm/s (mean ± SD).

Alexa Fluor 546 labeled filaments and cell bodies in a manner similar to that of Alexa Fluor 532. Alexa Fluor 546 is probably optimum for mercury-arc excitation: it was developed to match one of the mercury emission lines.

To the eye, Alexa Fluor 594 was the most impressive dye. Filipments were labeled brightly and appeared slightly thicker than with other dyes, presumably because of the longer emission wavelength. The cell bodies also were labeled, but they did not appear to be as bright as when labeled with dyes of shorter wavelength. The behavior of the bacteria was similar to that seen with Alexa Fluor 532. Also, our CCD camera was more sensitive at the longer wavelength. The number of filaments per cell was 3.26 ± 1.77, and the filament length was 5.9 ± 2.5 μm.

Flagellar waveforms. We measured waveforms of filaments on de-energized cells stuck to glass, using images that were averaged digitally over 30 video frames (60 fields). If the cells were de-energized with FCCP and then allowed to settle onto glass, all of their filaments were normal. However, if they were allowed to interact with the glass first and then were de-energized (or partially de-energized) by exposure to light, other waveforms were observed, as shown in Fig. 2. Of 512 filaments observed on 152 cells viewed as described for panels C and D, 465 were normal, 15 were semicoiled, 24 were curly 1, and 8 were curly 2. Presumably, these various waveforms were induced by torsion exerted by the flagellar motor on a filament pinned at one or more points to the underlying surface.

Changes of this kind were observed with filaments on energized cells stuck to glass, as shown in Fig. 3. A normal-to-curly 2 transformation appears in fields 1 to 8. The distal end of the filament is normal and extends to the left, held in contact with the glass by the CW-rotating motor. This transformation proceeds proximally to distally, and the length of the normal segment shortens. The filament then relaxes back distally to proximally from curly 2 to semicoiled, as seen in fields 10 to 14, presumably because the end of the filament is now free to turn and the torsion is reduced. Finally, the filament transforms from semicoiled to curly 1, as seen in fields 27 to 35.

Bundles on freely swimming cells also exhibited different waveforms, as shown in Fig. 4: in addition to normal (A), these included curly 1 (B and C) and semicoiled (D). One of the curly 1 bundles is loose (B), and another is tight (C). Measurements of diameter and pitch made from images of the kind shown in Fig. 2 to 4 are shown in Fig. 5A. The corresponding values of curvature and twist are plotted in Fig. 5B. The solid line is a half-sine wave (8).

Tumbles. Tumbles are brief events that enable cells to alter course (4). They are thought to begin when flagellar motors change their direction of rotation from CCW to CW and to end when they switch back again to CCW (19, 22, 26). We
analyzed 167 events in which a cell swam steadily into the field of view, moving in the plane of focus, tumbled, and then swam steadily out of the field of view, still moving in the plane of focus. Such events were relatively rare: of the cells that happened to swim into the field of view in the plane of focus and then tumbled, most left by moving out of focus. In other words, we selected events that could be analyzed in their entirety. The behavior of these cells is summarized in Table 1. The majority entered and left the field of view with normal bundles, but others displayed bundles that were semicoiled, curly 1, or of a hybrid waveform, i.e., with some filaments normal and others semicoiled or curly 1. A number of these events are shown in Fig. 6 to 11.

Figures 6 to 8 show tumbles generated by a single filament in cells with different numbers of filaments. The cell in Fig. 6 had only 1 long filament (and 1 short stub, not visible in this sequence). A transformation from normal to semicoiled is seen in fields 4 to 10, from semicoiled to curly 1 in fields 12 to 18, and from curly 1 back to normal in fields 24 to 30. This was a common sequence. This cell swam into the field of view moving toward 7 o’clock and left the field of view moving toward 5 o’clock. Notice that the curly 1 form wraps around the normal filament as it reverts back to the normal form and the tumble ends (fields 17 to 20). This cell swam into the field of view moving toward 5 o’clock and left the field of view moving toward 6 o’clock. Figure 8 shows a cell with a loose flagellar bundle (fields 2 to 6) from which a single filament emerges (fields 8 to 18), probably as curly 1. This filament appears to have rejoined the bundle by frames 20 to 24, where the bundle is tight. The change in the direction of motion generated by this maneuver was relatively small.

Figure 9 shows a tumble in which one filament (the one pointing towards 1 o’clock in fields 12 to 24) maintains a constant orientation and waveform, while all of the other filaments undergo polymorphic transformations. Evidently, this filament did not participate in the tumble. The shapes of the other filaments are difficult to discern: a semicoiled filament is prominent in field 18. This cell swam into the field of view moving toward 8 o’clock and left the field of view moving toward 5 o’clock.

Figure 10 shows a cell swimming with a curly 1 bundle with at least one filament of normal waveform (fields 1 to 4). In field 5, a curly filament appears that is wrapped around the bundle. It then unwraps (fields 6 to 8). More filaments leave the bundle (fields 11 to 15), with at least two remaining (field 15). Then all of the filaments rejoin the bundle (fields 16 to 20), which now appears normal. This cell swam into the field of view moving toward 10 o’clock and left the field moving toward 11 o’clock.

Figure 11 shows a cell with a normal bundle (field 2) that tumbles (fields 4 to 34) and then swims off in nearly the same direction, with its bundle displaying a mixed waveform (fields 36 to 40). In this case, all of the filaments appear to contribute to the tumble, although normal filaments are seen part of the time (fields 12 to 26).

The onset of a tumble was evident when the bundle loosened near the cell body. Soon thereafter, one or more filaments

![FIG. 3. Immobilized cell with a rotating filament undergoing polymorphic transformations. Successive fields are shown at 60 Hz (deinterlaced; total time span, 0.57 s). Fields 16 to 25 looked like field 26 and have been omitted.](http://jb.asm.org/)

![FIG. 4. Swimming cells with different kinds of flagellar bundles. Single fields are shown (deinterlaced). The waveforms of the flagellar bundles are normal (A), normal or curly 1 (both loose) (B), curly 1 (tight, but with one of the filaments on the cell at the right with a normal distal segment) (C), and semicoiled (with one filament with a normal distal segment) (D).](http://jb.asm.org/)
escaped from the bundle, and the cell body changed course.
Since these filaments exhibited right-handed waveforms (usu-
ally semicoiled first, and then curly 1), the motors driving them
must have switched from CCW to CW. The end of a tumble
can be defined in two different ways: when the cell body begins
to move in a new well-defined direction — this was the crite-
ron used in the tracking experiments (4) — or when the
filaments return to the bundle, i.e., when the bundle is consol-
dated. Cells tend to move in a new well-defined direction
before consolidation is complete. For 93 of the events listed in
Table 1, alterations in course were easily discerned, and the
period required was $0.14 \pm 0.08$ s. These alterations occurred
while one or more filaments were in the semicoiled conformation.
For these events, the corresponding time for bundle con-
solidation was $0.43 \pm 0.27$ s; for all of the events listed in Table
1, the time for bundle consolidation was $0.43 \pm 0.25$ s. In the
tracking experiments, the tumble length (the time before a new

TABLE 1. Tumble types of *E. coli* cells observed in this study

<table>
<thead>
<tr>
<th>Bundle characteristic:</th>
<th>No. of examples ($n = 167$)</th>
<th>Change in direction of cell (°)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Before tumble</td>
<td>After tumble</td>
<td></td>
</tr>
<tr>
<td>Normal</td>
<td>Normal</td>
<td>114 $57 \pm 42$</td>
</tr>
<tr>
<td>Normal</td>
<td>Semicooled</td>
<td>5 $57 \pm 37$</td>
</tr>
<tr>
<td>Normal</td>
<td>Curly 1</td>
<td>27 $55 \pm 31$</td>
</tr>
<tr>
<td>Semicooled</td>
<td>Normal</td>
<td>5 $57 \pm 37$</td>
</tr>
<tr>
<td>Curly 1</td>
<td>Normal</td>
<td>27 $55 \pm 31$</td>
</tr>
<tr>
<td>Curly 1</td>
<td>Curly 1</td>
<td>6 $49 \pm 33$</td>
</tr>
<tr>
<td>Normal</td>
<td>Normal and semicoiled</td>
<td>3 $104 \pm 75$</td>
</tr>
<tr>
<td>Normal</td>
<td>Normal and curly 1</td>
<td>2 $80 \pm 31$</td>
</tr>
<tr>
<td>Normal and semicoiled</td>
<td>Normal</td>
<td>1 $37$</td>
</tr>
</tbody>
</table>

*Mean ($\pm$ SD) change in direction of translation of the cell body from the end
of one run to the beginning of the next. With each tumble weighted equally, the
average change in direction was $58 \pm 40°$. 

FIG. 6. *E. coli* cell with one flagellar filament undergoing a polymorphic
transformation. Every other field is shown.

FIG. 7. *E. coli* with two flagellar filaments, one undergoing a polymorphic
transformation. Successive fields are shown.
direction was well defined) was $0.14 \pm 0.19$ s, in agreement with the value found here. It was noted in the tracking experiments (Fig. 2 of reference 4) that changes in course often were complete well before cells swam as fast as they did before the tumble. The time from the initial decrement in speed to 75% recovery for the events displayed in that figure was $0.26 \pm 0.10$ s, compared with $0.02 \pm 0.04$ s for the tumbles flagged by the tracker. (For this cell, all tumbles were relatively short.) Evidently, the initial run speed is not attained until the bundle is consolidated.

The change in direction from run to run for the events of Table 1 was $58 \pm 40^\circ$. These changes are plotted in Fig. 12 as a function of total number of filaments (two to five for panels A to D, respectively) and of the fraction of filaments out of the bundle. These data are summarized in polar plots (Fig. 13). Large changes in direction tended to occur when all of the filaments were involved in the tumble. When smaller numbers were involved, the distributions peaked more in the forward direction, and many of the events were below the threshold used to identify tumbles in the tracking experiments ($35^\circ$). When the number of filaments on a cell was large, e.g., five (Fig. 12D), tumbles generated by small numbers of filaments were rare.

**DISCUSSION**

**Labeling.** Succinimidyl esters of Alexa Fluor dyes appear to be ideal for labeling flagella of gram-negative bacteria, because they are (i) amino specific, (ii) relatively large and water soluble, (iii) highly fluorescent, and (iv) resistant to bleaching. Evidently the density of accessible amino groups on the outer surface of filaments is relatively large, while that on the outer surface of the cell body is relatively small. As a result, the filaments can be seen along their entire lengths. The N terminus of FliC (flagellin) is required for export, and both the N and C termini are required for filament assembly. The central region, comprising domains at the surface of the filament,
tolerates sizeable differences in length and composition (reviewed in reference 29). This region exhibits considerable diversity among flagellins of motile bacteria, varying in length from a few to 300 or 400 amino acid residues (11). This flexibility has led to the routine engineering of hybrid flagellins, including libraries of filament-bearing epitopes (20). Thus, it is not surprising that the surface of the flagellar filament can tolerate chemical modification. The outer subdomain of *E. coli* FliC contains at least three lysine residues (18), and that of *S. enterica* serovar Typhimurium contains at least eight (27, 37); however, seven of the latter are methylated. Of course, it is possible that the dye penetrates beyond the outer subdomain. In any event, the filaments of both species were relatively bright. The dyes do not appear to cross the outer cell membrane, presumably because of their size and solubility. Alexa Fluor 532 carboxylic acid succinimidyl ester, for example, is a pentacyclic rhodamine-like dye carrying one delocalized positive charge and two localized negative charges (sulfonic acid residues); its molecular weight is 724, somewhat above the permeability limit for known porins (28). As far as we could tell by practiced eye, the motility of labeled cells was normal. There was no apparent difference in behavior between cells treated with different dyes, which, a priori, might be expected to affect filament structure in different ways. The four Alexa Fluor dyes that we used have different net charges (−1 or −2) and different numbers and kinds of side groups (amino, carboxyl, chlorine, methyl, etc.).

**Flagellar waveforms.** In the models of Calladine (7, 8) and Kamiya et al. (16), the succession of waveforms expected for increasing twist is normal, coiled, semicoiled, and curly 1. Except for the absence of coiled, this is the order that we observed during tumbles. Curly 2 also was observed with filaments pinned to glass (Fig. 2C and D and Fig. 3). Semicoiled, curly 1, and curly 2 are right-handed and are expected to appear as the flagellar motor turns CW (26). Transformations to either left-handed or right-handed straight forms are generally prohibited, except in sag mutants, where the defect lies in HAP3, the protein to which the filament is attached at its base (10).

**Tumbles.** Cells of *E. coli* wild type for chemotaxis swim steadily along a smooth trajectory, suddenly change direction (or move erratically in place), and then swim steadily once again. These events were defined by three-dimensional tracking (4) as “runs,” “tumbles” (originally called “twiddles”), and “runs,” respectively. In the tracking experiments, the positions of the cell body were determined at intervals of 0.08 s, and the possibility of a tumble was checked whenever the cell appeared to change direction by 35° or more; see the addendum in the report by Berg and Brown (5). A tumble could be zero seconds long if a cell suddenly changed direction, or it could be several tenths of a second long if the cell continued to move erratically. Tumble lengths were distributed exponentially, with a mean of about 0.1 s; abrupt changes in direction were the most frequent. Changes in direction from run to run were nearly random, peaking slightly in the forward direction (mean ± SD,
68 ± 36° rather than 90 ± 39°, with the distribution for sudden changes in direction peaking even more sharply at 62 ± 26°). Later, it was recognized that runs occur when flagella spin CCW, and tumbles occur when they spin CW (19). However, the latter experiments were done with tethered cells, where one looks at only one flagellar motor at a time. Following the seminal work of Macnab and Ornston (26) described in the introduction, it was commonly thought that all of the motors turn CCW during a run and that most, if not all, turn CW during a tumble. A paradox arose when it was found that different motors in the same cell change direction independently, at random, provided that their filaments do not interact mechanically (15, 25). How, then, could all of the motors manage to turn CCW at the same time, as required if their filaments were to form a coherent bundle? This led to a “voting hypothesis,” in which it was supposed that a stable bundle forms when a certain fraction of motors rotate CCW (15). However, there was no direct evidence to support this hypothesis (see the appendix in the report by Ishihara et al. [15]). The present work resolves this paradox. Not all of the motors need to turn CCW for a cell to run, and only a few need to spin CW for a cell to tumble.

The deflection of the cell body tends to occur early in a tumble, while one or more filaments are in the semicoiled configuration, with the new run direction initiated before the bundle is consolidated, often while one or more filaments are still curly. This explains a phenomenon observed with wild-type cells in the tracking experiments: large changes in direction often occurred abruptly, within 0.08 s (classified as tumbles of length 0), while depressions in speed were more prolonged. The curly 1 waveform appears to be relatively flexible: curly 1 filaments often wrap around normal filaments or around a bundle (as in Fig. 7, fields 16 to 19, or Fig. 10, field 5). This compliance might make the curly filament bend (literally) to the will of the other filaments, and thus prevent it from interfering with the forward motion of the cell body that occurs after the cell has altered course, but before the bundle has consolidated.

We have not studied _S. enterica_ serovar Typhimurium as extensively as _E. coli_, but tumbles appear to involve the same transformations in either organism. In particular, we do see normal-to-semi-coiled transformations in _Salmonella_. These were not observed in dark field in the initial work of Macnab and Ornston (26) or in later studies in which video recordings were made with a silicon-intensified-target camera (17). We suspect that this discrepancy is due to the fact that the normal-to-semi-coiled transformation shorts the filament (compare fields 2 and 10 in Fig. 6) so that the semicoiled form is hidden by light scattered by the cell body.

_S. enterica_ serovar Typhimurium does appear to have more flagella than _E. coli_. Lino (14) shows distributions for cells of this species with a mean of between 6 and 7; for cells of _E. coli_ labeled with Alexa Fluor 532, we found a mean of 3.4. If the trend evident in Fig. 12D holds for _Salmonella_, then one might expect to see fewer tumbles that involve relatively small numbers of filaments. This might have contributed to the impression that tumbles involve all of the filaments in a bundle.

**Other vistas.** We have not looked at the behavior of cells with reduced numbers of filaments, with mutant flagellar filaments, or with normal filaments in highly viscous media. For example, more might be learned about correlations between changes in the direction of the cell body and filament polymorphic form if there were fewer filaments to complicate the issue. Nor have we looked beyond _E. coli_, _S. enterica_ serovar Typhimurium, or a motile _Streptococcus_ strain. Since labeling with the Alexa Fluor succinimidyl esters rendered the latter cells immotile, presumably because the gram-positive cell wall is permeable to these reagents, the use of this technique in such species might be limited to determinations of flagellar number and morphology. But even this is useful. The labeling technique is so simple, and the images are so vivid, even when seen with an ordinary fluorescence microscope, that the world of the flagellum is now more accessible.

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**References**