Growth of Flagellar Filaments of Escherichia coli is Independent of Filament Length

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Growth of flagellar filaments of *Escherichia coli* is independent of filament length

Running title: Flagellar growth independent of length

Linda Turner*, Alan S. Stern, and Howard C. Berg†

*Rowland Institute at Harvard, Cambridge, Massachusetts 02142*

Corresponding authors.

* Mailing address: Rowland Institute at Harvard, 100 Edwin Land Blvd., Cambridge, MA 02142. Phone: (617) 497-4658. Fax: (617) 497-4627. E-mail: turner@rowland.harvard.edu
† Mailing address: Department of Molecular and Cellular Biology, 16 Divinity Ave., Cambridge, MA 02138. Phone: (617) 495-0924. Fax: (617) 496-1114. E-mail: hberg@mcn.harvard.edu.
ABSTRACT

Bacterial flagellar filaments grow at their distal ends, from flagellin that travels through a central channel ~2 nm in diameter. The flagellin is extruded from the cytoplasm by a pmf-dependent pump. We measured filament growth in cells near mid-exponential phase with flagellin bearing a specific cysteine-for-serine substitution, allowing filaments to be labeled with sulfhydryl-specific fluorescent dyes. First, we labeled filaments with a green maleimide dye and then, following an additional period of growth, with a red maleimide dye. The contour lengths of the green and red segments were measured. The average lengths of red segments (~2.3 µm) were the same regardless of the lengths of the green segments from which they grew (ranging from less than 1 to more than 9 µm in length). Thus, flagellar filaments do not grow at a rate that decreases exponentially with length, as formerly supposed. If flagellar filaments were broken by viscous shear, the broken filaments continued to grow. Identical results were obtained whether flagellin was expressed from fliC on the chromosome under control of its native promoter, or on a plasmid under control of the arabinose promoter.
INTRODUCTION

A bacterial flagellum has three parts, a basal body embedded in the cell wall (a rotary motor) and two external structures, a short proximal hook (a universal joint), and a long helical filament (a propeller, polymerized from the protein flagellin). A flagellum is assembled from the inside out, beginning with the MS ring in the cytoplasmic membrane and the C ring in the cytoplasm. A rod connects the MS ring to the hook, and hook-associated proteins connect the hook to the filament (12). The axial components are exported by a type III secretion system, each new component added at the end of the growing structure (3, 13). This export is powered by a protonmotive force (14, 15). Two of the hook-associated proteins couple the hook and the filament, allowing the hook to flex and the filament to rotate rigidly (4), while a third protein caps the distal end of the filament, preventing loss of flagellin and orchestrating its polymerization (7, 20). The hook associated proteins are exported continuously, appearing in the culture medium, even of wild-type strains, suggesting that caps lost from broken filaments can be replaced (6).

In seminal work summarized in 1974, Iino (8) used electron microscopy to determine the numbers and lengths of flagellar filaments on cells in a culture of Salmonella growing in exponential phase. He concluded that the rate of elongation decreases exponentially with length. Support for this idea was obtained in vivo by dark-field microscopy by Aizawa and Kubori (1), but the data were limited and only long filaments could be seen. Having shown that flagellin moves through the filament rather than via the external medium, Iino assumed that transport efficiency decreases with
filament length, like the flow of liquid through a pipe driven by a constant pressure head. This idea has been reinforced by recent theoretical work (17).

Several years ago, we found that flagellar filaments of Gram-negative bacteria could be labeled with fluorescent dyes in a manner that did not interfere with their motility (18). The original work was done with amino-specific dyes (succinimidyl esters), which also labeled outer cell membranes. A milder and more specific procedure was developed by adding cysteine to flagellin and using sulfhydryl-specific (maleimide) dyes (19). These methods enabled us to visualize flagellar filaments all the way to the cell surface. By using dyes of different color and measuring the lengths of segments of different color on individual filaments, we now have monitored filament growth. We have found, to our surprise, that E. coli filaments grow in a manner that is independent of their length, not at a rate that is an exponentially decreasing function of length, as commonly supposed (1, 8, 17). We also have confirmed that broken filaments continue to grow, a proposition that has remained in doubt (16).

MATERIALS AND METHODS

Strains, growth and labeling.

E. coli strain AW405 (2) is wild-type for chemotaxis and swims vigorously. Strain HCB1737 is isogenic with AW405 except for a single cysteine substitution in fliC (S219C). The wild-type fliC allele of strain AW405 was replaced by the mutant allele using the λ-Red system, described previously (22). This strain also swims vigorously. Strain HCB1668 is a Tn5 fliC null derivative of AW405 in which fliC (S353C) is
expressed on the plasmid pBAD33 under control of the arabinose promoter, maintained by addition to the growth media of kanamycin (50 µg/ml) and chloramphenicol (34 µg/ml). This strain swims and also swarms (19) vigorously. Details of the serine to cysteine replacements are given under **Strain Construction** in **Supplemental Material.**

**Strains HCB1737 and HCB1668 were cultured from single-colony isolates from frozen stocks (-75°C), placed in 10 ml LB broth (10 g Bacto-Tryptone, 5 g Yeast Extract and 5 g NaCl per liter) in 125-ml Erlenmeyer flasks, and grown to saturation at 30°C with gyration at 150 rpm. 1/100 dilutions of the LB cultures were made in 10 ml of T broth (10g Bacto-Tryptone and 5g NaCl per liter) and grown in 125ml Erlenmeyer flasks at 30°C with aeration by gyration at 150 rpm for 4 h to a cell density of ~4.1 x 10^8 cells/ml. For HCB1668, arabinose at 0.5% final concentration was added to the T broth. Cultures were checked for motility using a phase-contrast microscope (Nikon Optiphot), and washed with 10 ml aliquots of motility buffer (0.01M potassium phosphate, pH 7.0, 10^{-4} M EDTA, 0.067M NaCl) (MB) containing 0.0001% Tween-20 (MBT) by 12 min centrifugation at 1,400 x g followed by gentle resuspension, 4 times, manipulations spanning a period of 68 min. Then the cells were labeled with Alexa Fluor 488 maleimide dye, as described in (19), for ~90 min rather than 60 min with double the amount of dye, so that even very short filaments could easily be seen by fluorescence. Once labeled, 2 ml of MBT was added to the cells, and the suspension was divided into two 1-ml aliquots in separate centrifuge tubes. One aliquot was sheared by passing it 5 times in and out of a 3 ml syringe equipped with a 22 gauge needle. The extent of shearing was checked by microscopy. A small sample (diluted 10-fold) was imaged in fluorescence – loose filaments were seen stuck to the coverslips – and in phase-
contrast – cell bodies appeared dark – and in combined illumination – filament stubs could be seen on cell bodies. After shearing, the two aliquots (unsheared and sheared) were treated in parallel. Cell aliquots were brought to 10 ml volumes with MBT and washed by centrifugation twice (as above), then centrifuged a final time and resuspended in 5 ml of T broth, manipulations spanning a period of 51 min. The entire 5 ml of T broth was used as an inoculant for continued culturing in 250ml Erlenmeyer flasks containing 20 ml of T broth (with antibiotics and arabinose, when required). The culture flasks were stored overnight (~14 h) at 7°C, then incubated at 30°C with aeration by gyration at 150 rpm for 3 h to a cell density of ~3.1 x 10^8 cells/ml. After growth, 10 ml of each culture was washed into MBT by 12 min centrifugation at 1,400 x g followed by gentle resuspension, 4 times (as above), and then labeled with Alexa Fluor 546 maleimide dye, using the procedure described for the 488-nm dye (as above). Excess dye was removed by repeated centrifugation and resuspension in MBT (as above). The final suspension of the pellets was in 2.5 ml of motility buffer, MB, without Tween-20.

**Growth curves**

The growth and labeling procedure described above was repeated in precisely the same way with more cells and flasks, and 1.5 ml samples were withdrawn from the different flasks at different times for measurements of OD_{610} and construction of growth curves. The conversion from OD_{610} to cells/ml was based upon a calibration of the spectrometer (Perkin Elmer Lambda EZ142) by serial dilution and plating.

**Microscopy.**

Microscope slides and coverslips were used out-of-the box. Tunnel slides were constructed using Scotch-brand double-sided sticky tape, by placing two layers of tape
about 7 mm apart across the width of the microscope slide and placing a 22 x 22 mm
No. 1 coverslip on top of the tape. Poly-L-lysine 0.01% (Sigma P4707) was added to
the tunnels, which were allowed to sit for ~5 min in a humidity chamber and then rinsed
with ~0.5 ml MB. The cell suspensions, described in the previous section, were diluted
with an equal volume of MB and then added to fill the tunnels. The tunnel slides were
inverted and placed in the humidity chamber for ~5 min and then rinsed with MB. The
phase-contrast microscope was used to monitor the density of cells stuck to the
coverslips, to determine if the final cell density was suitable for fluorescence imaging of
the flagellar filaments. If too many cells were stuck, a new preparation was made using
a more dilute cell suspension. Glutaraldehyde (0.25% in MB) was added to the tunnel
slides for 2-3 min to stop flagellar rotation and to cross-link the filaments to the surface
of the coverslip. Tunnel-slide preparations were rinsed with MB, then stored in the
humidity chamber prior to microscopy. Fluorescence imaging was with a Nikon inverted
microscope (Diaphot 200) equipped with a 100 W mercury arc, with a dual filter cube
(FITC/Texas red, Chroma Technologies #51006), a 100x oil-immersion objective, a 2.5x
relay lens, and a Nikon D-70 camera. Images were captured using Nikon Image
Capture software running on a PC laptop. Exposure times were sample dependent and
usually between 2 and 10 s.

Image Analysis.

RGB colored images (Fig. 1A) were transferred to a Macintosh computer and analyzed
using ImageJ software. Each image file was 3872 X 2592 pixels at 300 pixels/inch.
The image magnification was 40 pixels/micron. For analysis, each RGB image was split
via the standard ImageJ command “Split Channels” to recover the green images in
black and white, with green appearing white and other colors black (Fig. 1B). Green filament lengths, indicating old growth, were identified on these black and white images. Red filament lengths, indicating new growth, were identified on the original RGB images (Fig. 1C). The ImageJ free-form line tool was used to trace filament contours. Contour lengths were smoothed using the ImageJ cubic spline fit function, drawn on the image, recorded as a length measurement on a spreadsheet, and then analyzed and graphed using KaleidaGraph software. Rough estimates of errors for measurements of contour lengths were ~5% when filaments were short and ~2.5% when filaments were long. If a green filament had no red filament section, then a zero red segment length was recorded. Every in-focus image was analyzed and every green segment in each image was measured until a total of 1250 filaments had been logged.

RESULTS

Filament growth, protocol A.

To learn how growth of flagellar filaments depends upon initial length, we used two derivatives of the wild-type strain AW405 with flagellin bearing a serine to cysteine substitution. Expression was either from the chromosome under control of the native promoter or from a plasmid under control of the arabinose promoter. The results for the two strains were essentially identical. The experimental protocol is shown in Fig. 2A. Cells were cultured to late-exponential phase, checked for maximal motility, then labeled to saturation with a sulfhydryl-reactive green fluorescent dye. The flagellar filaments of these cells were all green, creating a population of filaments that could be traced through subsequent generations. A second growth period was initiated. After 3 h of growth, the cells were labeled a second time with a red fluorescent dye, so new
growth was red. 99.85% of the cells of strain HCB1737 showed new (red) flagellar growth, indicating that the green labeling procedure did not kill cells. Three types of colored filaments were seen by fluorescence (Fig. 1A), as indicated in the cartoon (Fig. 2): a, filaments that were both green and red; b, filaments that were entirely green; and c, filaments that were entirely red. About 14% of filaments bearing green dye were entirely green, showing no evidence of growth during the second culture period; although, red filaments grew elsewhere on the same cells. The other 86% of the filaments bearing green dye were green at their proximal ends and red at their distal ends, indicating new filament growth during the second culture period. Many filaments were entirely red, indicating that their growth was initiated after cells had been labeled green. These filaments were not considered in the analysis, because the time for their initiation could not be determined. However, lengths of red-only filaments were measured on a subset of 200 unsheared cells of strain HCB1737, yielding a mean of 2.52 μm with an SEM of 0.09, values not significantly different from the red growth on green filaments.

The contour lengths of 1250 filaments were measured for each strain, with two measurements for each filament: the length of the green segment and the length of the corresponding red segment. The complete data set for strain HCB1737 is shown in Fig. 3A. This is a scatter plot in which the lengths of red segments are plotted versus the lengths of green segments from which they grew. The cloud of points is broad and roughly rectangular, centered around 4.5 μm in the green dimension and around 2.5 μm in the red dimension, indicating a varied range of growth for each filament length. The
cloud does not skew downward to the right as expected were filament growth to
decrease with length. There are several red points of value zero spread along the
green axis (the 14% of green filaments that did not grow further).

More could be learned about these data by examining distributions binned according to
segment lengths, as shown in Fig. 4 (top row). The distribution of green segments (Fig.
4A) was roughly normal with a mean of ~4.3 µm, but included a few filaments as long as
13 µm. Following the second period of growth, the distribution of red segments (Fig.
4B) had a mean of ~2.3 µm. The distribution of total lengths (Fig. 4C) was nearly
normal with a mean of 6.6 µm. These results are summarized in the first column of
Table 1.

As expected from Fig. 3A, distributions of red segments binned in successive 1 µm
intervals and sorted according to the lengths of the green segments from which they
grew (data not shown) were all fairly broad. Remarkably, they all had similar means, as
plotted in Fig. 5. Means ranged between 1.8 µm and 2.5 µm, almost all within 3
standard errors of 2.16 µm, the mean of all points, and most within one standard error.
This implies that new growth is much the same, regardless of the length of old growth,
i.e., that the growth of flagellar filaments of *E. coli* does not depend upon filament
length. This was true whether flagellin was expressed from the chromosome under
control of its native promoter (strain HCB1737) or from a plasmid under control of the
arabinose promoter (strain HCB1668).
Growth curves

As noted in MATERIALS AND METHODS, the two culture periods were separated by washing and labeling steps that required cells to remain at room temperature in dilute phosphate buffers for about 4 h, followed by storage in growth medium at 7°C for about 14 h. The motility was checked along the way, and found to be maximal. But it was important to know whether the two growth periods were equivalent. So we repeated the procedure with strain HCB1737 following all the same steps and measured the optical densities of the cultures during both growth periods. The results are shown in Fig. 6. During both periods the cells grew exponentially with a doubling time of ~56 min up to the time point of 180 min. At later times the rate of growth was smaller, as the cells entered late exponential phase. Although these results imply that labeling did not kill cells, we checked this directly by growing cells, dividing the culture into two parts, labeling one, and measuring growth curves for both without delay, using similar inocula. The two growth curves were identical (data not shown), indicating that labeling did not affect viability.

Filament growth, protocol B.

To learn whether broken filaments continue to grow, half of the population of cells that had been labeled green were subjected to viscous shear and then were treated in the same way as the cells that were not sheared, as indicated in the cartoon (Fig. 2B). A scatter plot of the lengths of red segments versus the lengths of the green segments from which they grew is shown in Fig. 3B. The cloud of points is no longer rectangular.
It has shifted to the left: there are many more points near the left boundary, and the right boundary no longer extends beyond 7 µm. So it is evident that filaments were broken by shear. The upper edge of the cloud now slants downward to the right.

Again, more could be learned about such data by examining distributions binned according to segment lengths, as shown in Fig. 4 (bottom row). The distribution of green segments (Fig. 4D) was exponential, with the shortest lengths the most probable. The mean for this distribution is 1.6 µm, and the distribution only extends out to ~7 µm.

So, as asserted above, long filaments were broken. Following the second period of growth, the distribution of red segments (Fig. 4E) had a mean of 2.6 µm, including segments as long as 8 µm. So the new growth was roughly the same whether the filaments had been sheared or not. The distribution of total lengths (Fig. 4F) was more nearly normal with a mean of 4.2 µm. These results are summarized in the second column of Table 1.

Evidence that filaments broken by shear continue to grow is contained in the length distributions of Fig. 4 panels A, D and F. The aliquot of cells that were sheared, with the green-segment length distribution shown in panel D, was taken from the suspension of cells that were not sheared, with the green-segment length distribution shown in panel A. A comparison of these two distributions enables us to distinguish most of the filaments that were broken by shear from those that were too short to be broken by shear. There are only 46 filaments of length < 1 µm in the initial distribution, Fig. 4A, but 483 in the final distribution, Fig. 4D. Therefore, approximately 437 of these
filaments must originally have been longer than 1 µm, i.e., must have been broken.

Following the second period of growth, only 82 filaments showed a total length < 1 µm, Fig. 4F. Hence the majority of the 437 broken filaments must have grown. A formal extension of this analysis that includes data for filaments of length > 1 µm is given in Supplemental Material.

As expected from Fig. 3B, distributions of red segments binned in successive 1 µm intervals and sorted according to the lengths of the green segments from which they grew (data not shown) were all fairly broad. But now their means decreased somewhat with green segment length, as shown in Fig. 7. The average growth decreased approximately linearly from slightly less than 3 µm to slightly less than 2 µm over the entire range of initial lengths of less than 1 µm to 7 µm. This was true for both strains HCB1737 and HCB1668.

DISCUSSION

We studied growth of flagellar filaments by a method that allowed us to distinguish new growth (segments labeled red) from old growth (segments labeled green) in individual filaments. We concluded that filament growth for samples that had not been sheared did not depend upon filament length, Fig. 5. For samples that had been sheared, shorter filaments grew somewhat longer extensions, Fig. 7. A comparison of length distributions between a sample of a cell culture that had not been sheared (Fig. 4A) with one that had been sheared (Fig. 4D) and then grown further (Fig. 4F) showed that broken filaments continue to grow.
An earlier study of this kind was made by Iino (9), who grew cells at 37° on minimal medium with glucose or on minimal medium with glucose followed by saline, and then on minimal medium with glucose plus a complete mixture of L-amino acids, in which phenylalanine was replaced by p-fluorophenylalanine (FPA). This analogue had been shown by Kerridge (11) to promote growth of curly filaments. The number of flagella per cell decreased from ~4.4 to ~2.5, but newly grown curly waves were shorter in filaments with more normal waves and reached zero at about 5 normal waves (~12 µm).

However, it was not possible to distinguish normal waves newly grown on FPA from those present before treatment. In this work, Iino also concluded that broken filaments continue to grow. In a later study conducted without FPA, Iino (8) concluded that flagella grew at a rate that decreased exponentially with length. His analysis was based upon the assumption that the order of relative lengths of filaments remained fixed at successive time points during exponential growth; for example, that the longest filaments at one time point were derived from the longest filaments at an earlier time point. This is not true for our data. Pick a green length from the abscissa of Fig. 3 and note the wide distribution of red lengths denoting subsequent growth. These distributions look much the same regardless of green length. New growth is not correlated with old growth. We would expect results obtained with Salmonella to be similar to those obtained with E. coli. But our results are different.

If Iino’s theory (8) is used to compute the times required for filaments to grow to the lengths shown on the abscissa of Fig. 5 (using the data Iino gives for a generation time
of 56 min) and then the filaments are allowed to grow for an additional 3 h, we find the additional lengths to be much shorter at the 7.5 µm point than at the 0.5 µm point, only 39% as long. The plot for this (steep) decline is nearly linear (slightly concave upwards), as in Iino's Fig. 8. Our data for unsheared filaments, Fig. 5, do not show such decline.

There is a decline, albeit slight, for sheared filaments, Fig. 7. It is linear but not steep, and appears with both strains. We do not understand the reason for this decline. Presumably, it has to do with replacement of the cap protein, FliD. However, if FliD moves slowly through sheared filaments, why does it not also move slowly through unsheared ones? Might the mode of operation of the pump change when the filament lacks a cap?

In vitro, filaments grow at a constant rate, with pauses thought to occur when defective flagellin monomers are transiently incorporated (10). In between pauses, the average rate of extension is proportional to the concentration of free flagellin, with a distribution of rates from filament to filament that is fairly broad. In our experiments, filaments continue to grow in length and number, suggesting that the supply of flagellin is adequate and sustained, but the distributions of rates from filament to filament also are broad. The results are the same whether the flagellin is expressed from the chromosome under control of its native promoter (strain HCB1737) or from a plasmid under control of the arabinose promoter (strain HCB1668). Note that we did not measure rates, per se, as one might do with a time-lapse movie of growing filaments,
but rather the extension produced over a specified time span. Thus, we do not know
how variable rates of growth might have been or whether there were pauses.

In vivo, flagellin monomers move along the axis of a flagellar filament through a pore
only 2 nm in diameter (21), crystallizing out under the FliD cap at the distal end. To
pass through this pore, the flagellin must be unfolded. The N- and C-terminal ends of
the molecule are \(\alpha\)-helical, so if we assume that the molecule goes through the pore as
an extended \(\alpha\)-helical chain, then it is \(~74\) nm long! This is 14 times longer than the
distance between subunits comprising flagellar protofilaments (5). Since there are 11
protofilaments, it takes \(11 \times 14 = 156\) subunits to lengthen the filament by enough for
the pore to contain one more flagellin \(\alpha\)-helix. So if one pumps in 157 subunits, 156 will
crystallize and one will remain in the pore. Because growth does not vary with length,
on average, our results imply that this pmf-driven export pump (14, 15) is a constant-
rate device, not one that runs at constant pressure. Under the conditions of our
experiment, it extrudes one flagellin molecule every 2 s. Variations from flagellum to
flagellum could arise from variations in pmf, in pump structure, or in rates at which
flagellin monomers are delivered to the pump.

If pumps keep pumping, what limits the lengths of flagellar filaments? Long filaments
probably break.

ACKNOWLEDGEMENTS

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REFERENCES


TABLE 1.

Lengths of colored segments in strains HCB1737 and HCB1668 (Mean ± SEM, µm)

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<th>HCB1737*</th>
<th>HCB1668**</th>
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<tr>
<td></td>
<td>Not sheared</td>
<td>Sheared</td>
</tr>
<tr>
<td></td>
<td>(n=1250)</td>
<td>(n=1250)</td>
</tr>
<tr>
<td>Green</td>
<td>4.29 ± 0.05</td>
<td>1.60 ± 0.03</td>
</tr>
<tr>
<td>Red</td>
<td>2.33 ± 0.05</td>
<td>2.58 ± 0.06</td>
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<tr>
<td>Green+Red</td>
<td>6.63 ± 0.07</td>
<td>4.18 ± 0.06</td>
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*After the 2nd culture period, the not-sheared cell population had on average 4.76 ± 0.17 filaments/cell (n=100) and the sheared cell population had on average 4.23 ± 0.18 filaments/cell (n=100).

** After the 2nd culture period, the not-sheared cell population had on average 4.87 ± 0.20 filaments/cell (n=100) and the sheared cell population had on average 5.54 ± 0.24 filaments/cell (n=100).
FIGURE LEGENDS

Figure 1. Examples of flagellar filaments of strain HCB1737 imaged by fluorescence microscopy. Cells were labeled with Alexa Fluor 488 dye (green), cultured for an additional period of time, and then labeled with Alexa Fluor 546 dye (red). A a full-color image. B an image of the same scene in which the green pixels appear white and other colors appear black, a technique used to more clearly distinguish green from red. The green segments were traced and are shown in white over white. C an image of the same scene in which the red segments of green/red filaments were traced and are shown in white over red.

Figure 2. Experimental protocol. Time flows from left to right. Protocol A (top row): Cells were grown to late exponential phase, labeled to saturation with a green fluorescent dye, grown again to late exponential phase, and labeled with a red fluorescent dye. Three kinds of filaments were observed: a, filaments with green proximal ends and red distal ends (common); b, filaments that were entirely green (rare); c, filaments that were entirely red (common). Initiation of growth of filaments of the latter type occurred after cells had been labeled green. The filaments in the experiments were normal and helical; for simplicity, they are shown here as straight. Protocol B (bottom row): The same as A except an aliquot of cells that had been labeled green was subjected to viscous shear; two of the filaments shown were shortened and one was not. The parent population for the cells that were sheared was the population studied in protocol A, so a record was made of cells both before and after shearing.

Figure 3. Plots of the lengths of red segments versus the lengths of the green segments from which they grew, for a single culture of strain HCB1737. A Filaments of unsheared cells. B Filaments of sheared cells. The corresponding plots for strain HCB1668 are shown in Supplemental Material, Fig. S1.
Figure 4. Distributions of segment lengths of filaments on cells of strain HCB1737, binned at 1-µm intervals. Green segments are shown in green, red segments are shown in red, and total lengths (green + red) are shown in grey. The distribution of the subset of green segments that failed to produce red extensions is shown by the white bars in panels A and D, while the total numbers of such green segments, which equals the numbers of red segments of zero length, are shown by the white bars in panels B and E. The top row (A-C) is for cells that were not sheared, and the bottom row (D-F) is for cells that were sheared. The corresponding plots for strain HCB1668 are shown in Supplemental Material, Fig. S2.

Figure 5. Plots showing the mean values and standard errors of the lengths of red segments as a function of the lengths of the green segments from which they grew (plotted at bin mid-values). Closed symbols (●) and solid line, filaments of unsheared cells of strain HCB1737, from left to right n = 49, 103, 158, 240, 266, 207, 138, 56, 33. The weighted line fit has a reduced χ² value of 1.77. Open symbols (○) and dashed line, filaments of unsheared cells of strain HCB1668, from left to right n = 72, 170, 178, 205, 155, 162, 142, 114, 52. The weighted line fit has a reduced χ² value of 0.63.

Figure 6. Growth curves for the two culture periods. The logarithm of cell density (cells/ml) is plotted as a function of culture time (min). Open symbols (○) and dashed line, first growth period; closed symbols (■) and solid line, second growth period Cells were harvested at log 8.6 for the first growth period (4.1 x 10⁸ cells/ml) and at log 8.5 for the second growth period (3.1 x 10⁸ cells/ml). During the exponential phase (straight-line fits) the doubling time was ~56 min.

Figure 7. Plots showing the mean values and standard errors of the lengths of red segments as a function of the lengths of the green segments from which they grew (plotted at bin mid-values). Closed symbols (●) and solid line, filaments of sheared cells of strain HCB1737, from left to right n = 488, 389, 198, 105, 70. The weighted line fit has a reduced χ² value of 0.52. Open symbols (○) and dashed line, filaments of
sheared cells of strain HCB1668, from left to right n = 407, 370, 243, 132, 98. The weighted line fit has a reduced $\chi^2$ value of 0.75.
Grow, label green

Grow again, label red

Grow, label green, shear

Grow again, label red
Filaments of unsheared cells

Red segment length (μm)

Green segment length (μm)
Filaments of sheared cells

Red segment length (µm)

Green segment length (µm)