

Periodic reversal of direction allows Myxobacteria to swarm

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Many bacteria can rapidly traverse surfaces from which they are extracting nutrient for growth. They generate flat, spreading colonies, called swarms because they resemble swarms of insects. We seek to understand how members of any dense swarm spread efficiently while being able to perceive and interfere minimally with the motion of others. To this end, we investigate swarms of the myxobacterium, *Myxococcus xanthus*. Individual *M. xanthus* cells are elongated; they always move in the direction of their long axis; and they are in constant motion, repeatedly touching each other. Remarkably, they regularly reverse their gliding directions. We have constructed a detailed cell- and behavior-based computational model of *M. xanthus* swarming that allows the organization of cells to be computed. By using the model, we are able to show that reversals of gliding direction are essential for swarming and that reversals increase the outflow of cells across the edge of the swarm. Cells at the swarm edge gain maximum exposure to nutrient and oxygen. We also find that the reversal period predicted to maximize the outflow of cells is the same (within the errors of measurement) as the period observed in experiments with normal *M. xanthus* cells. This coincidence suggests that the circuit regulating reversals evolved to its current sensitivity under selection for growth achieved by swarming. Finally, we observe that, with time, reversals increase the cell alignment, and generate clusters of parallel cells.

gliding motility | stochastic model | pattern formation | cell alignment | oscillate

When fish school, birds flock, insects and bacteria swarm, a population of similar individuals cruises along similar, sometimes cyclic, paths. How do the members of a swarm anticipate the movement of others to coordinate movement with them? And how do members, when each is moving rapidly near to others, avoid impeding their neighbor's motion? These general issues, which have yet to be resolved for any schooling, flocking, or swarming organism, can be investigated in swarming bacteria (1). Such investigation might have practical value because swarming enables pathogens like *Proteus* (2), for example, to invade the urinary tract and to spread along the surface of catheters placed in the urethra (3). The behavior of swarming individuals might also suggest ways to speed high density automobile or pedestrian traffic and ways to avoid traffic jams (4). Bacterial sensory systems and behaviors are more amenable to analysis than those of animals. Because they are relatively small; several thousand bacteria can be followed in a time lapse movie of a swarm. Among bacteria that swarm are the hyperflagellated bacteria (5), all of the Myxobacteria (6), which lack flagella, and the Bacterioidetes that include *Cytophaga*, *Flavobacteria*, and *Bacteriodes* (7). Millions of *Myxococcus xanthus* cells are found to swarm outward for more than a week at a steady rate of ≈ 0.1 mm/hr, a rate close to half the speed of individual cell movement (8, 9). Swarming gives them a significant growth advantage: Whereas cells in the swarm center are competing with each other for nutrient and oxygen, cells at the swarm edge have practically unfettered access to both (10). In all of the movies made for Myxobacteria (11, 12, 13) and some of the Bacterioidetes (7), we

see the cells reverse their gliding direction, but none of these studies considered why. Individual *M. xanthus* cells are seen to reverse their gliding direction every few minutes (14, 15). This observation prompts the question: what benefit does *M. xanthus* gain when it expends an estimated 5% of the cell's ATP production (16, 17) only to move back and forth? We suggest that the oscillatory motion is actually necessary for the net migration of cells from the center of the swarm, where most are born, out to the swarm edge. By escaping from the central region of high cell density to the low density region at the swarm edge, *M. xanthus* moves to an unoccupied area where nutrient and oxygen are abundant and there is little contest for either (10)—a much greener pasture, to use Purcell and Berg's term (18).

Cell movement in *M. xanthus* is propelled by polar S engines (pulling type IV pili) at the leading end of the cell, and polar A gliding engines (for secretion of slime) at the trailing end of the cell. Evidence for sites of lateral cohesion of cells to the substrate has been presented in a context of challenging a strictly polar location of the A engines (19). However, we believe that the evidence presented fails to support the injection of a propulsive force at the sites of lateral cohesion as claimed, but instead support the view that lateral cohesions are necessary for propulsion by polar slime secretion. Arguments for these beliefs are summarized in the supporting information (SI) because they have been detailed elsewhere (20). Many of the proteins that constitute the polar A and the polar S engines have been characterized (20), and many single gene motility mutants have been examined (21, 22, 23). Social interactions, such as attachment by the S engines (pili) to fibrils on other cells and the following of slime trails left by the A engines, facilitate swarming (20). As long as fresh surface is available to them, the disk-shaped swarms of *M. xanthus* expand at a uniform rate, for weeks (8, 9). Uniform expansion implies that the flow of cells from the center of the swarm to the edge is steady. By using A⁻S⁻ mutants, it has also been shown that 90% of the swarm expansion rate results from cell movement; only 10% of the rate results from cell growth (9). Nevertheless, *M. xanthus* must be growing to swarm: When growth stops [even though a low level of nutrient may still be available, as in clone-fruiting medium (24)] the swarm stops expanding (10). Although the steady flow of cells to the edge is driven by growth, the spatial distribution of cells at the edge is the consequence of cell movement, which depends on the mechanics of the A and S gliding engines and on their social interactions. Because the cell flow to the swarm edge is constant over long periods of time, relative to the generation time of ≈ 3.5 h, time-lapse movies of several hours offer large

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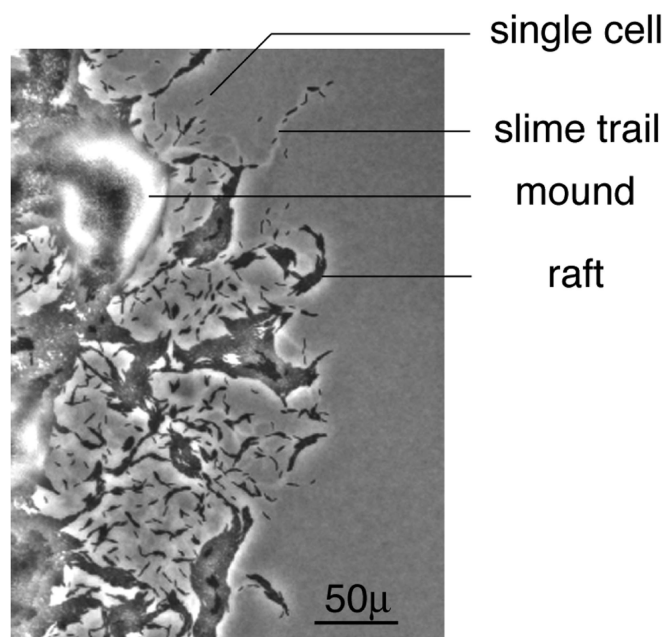


Fig. 1. Distribution of cells and multicellular structures at the swarm edge. In addition to some individual cells and slime trails, multicellular rafts and multicellular mounds are labeled. The swarm is expanding in the radial direction, which is to the right in this image. (Scale bar, 50μ .) This is the first frame of a 3 h time lapse movie (see Movie S1 in the SI).

samples of the patterns of cell movement to be found in the steady state.

Here we describe the rich dynamics of cell movement visible at the swarm edge observed in time-lapse movies. To find characteristic movement patterns in the images, we used a cell-based, stochastic computational model that was recently described (25). Because the model incorporates the mechanics of A and S motility as well as their social interactions, it accurately translates the macroscopic and readily measurable swarm expansion rates into microscopic cell fluxes that expose the average behavior of many individual cells.

Results

Dynamic Patterns at the Swarm Edge. A typical pattern of cells at the swarm edge is shown in Fig. 1. In every frame of the movie (see Movie S1 in the SI), there are many single, roughly $5\mu \times 0.5\mu$ cells. The single cells are always found on slime trails, because they are secreting propulsive slime from their poles (20). Two kinds of multicellular structures are also evident in Fig. 1 (see labels): Elongated rafts and mounds. Elongated rafts resemble two dimensional nematic liquid crystals and are com-

posed of several files of cells, with the long axes of the files adjacent and parallel to each other but the individual cell ends in adjacent files out of register with each other. Lack of registration is most evident at the ends of a raft where cells protrude. Mounds have several layers of cells stacked on top of one another. Each layer of a mound is roughly circular and concentric with the others, contrasting with the rafts that are rectangular and have fewer cells. The layering, evident in all mounds and on some rafts (indicated in Fig. 1) shows that cells readily glide over the surface of other cells. Single cells, small clusters of 2–3 cells, rafts and mounds are evident in every frame of the movie (see Movie S1 in the SI). Both multicellular structures exchange various numbers of cells with neighboring structures by branching or converging flows. In mounds and rafts, cells frequently move between adjacent layers. Because the movie was photographed at 2 frames/min for 3 h, the apparent smoothness of flow indicates that cell birth and death are negligible for 3 h. Finding smooth flow and the constant swarm expansion rate data (9) give firm support to the steady state assumption of the model. Individual cells are seen to glide in the direction of their long axis, consistent with a polar location for both the A and the S engines. Although cells can flex (26) and U-turns are very infrequent (27), cells never seem to move sideways, also consistent with a strictly polar engine location. Cells appear to be oriented in many directions; consequently the swarm disk would be expected to expand in radial directions; this justifies the assumption of a random initial cell orientation around the radius (25).

Solitary cells are observed to reverse their gliding direction in the movie with remarkable regularity: they reverse at 8.8 ± 2.1 min intervals (reversal data collection and analysis are in the SI). As noted elsewhere (23), reversing cells simply stop, pause for about a minute, then lead off with their formerly lagging pole. It has been proposed that reversal is effected by an MglA G protein switch (15). The proposal explains in detail the reversal behavior of RomR mutants (20), as described by ref. 28. Because *mglA* mutants lack the switch, they move abnormally, never reverse, and fail to swarm (23). Moreover, *mglA* mutant colonies are heaped up and sharp-edged, like colonies of a nonswarming *E. coli* strain (29). Switching the MglA G protein from its GDP to its GTP-state (30) is proposed to follow a pulse of FrzE~P from the frizzilator, a two-component system with negative feedback that causes it to oscillate with a definite period (31). As shown in Table 1 and the SI, none of the Frz deletion mutants have a sharp-edged, Mgl-like phenotype; all Frz mutants retain some capacity to reverse and to swarm. The data of Table 1 show that the rate of Frz mutant swarm expansion is correlated with a nonoptimal reversal period. Despite intensive genetic screens, no mutant that is still motile but has lost the ability to reverse, has ever been isolated. But, such a mutant can be simulated by deleting reversals from the computational model, and any value of the reversal period can be prescribed.

Table 1. Swarm expansion rates of wild-type and frizzy mutants

Strain	Rate of swarm expansion, mm/hr*	Gene deleted	Published reversal period [†]
DZ 2	0.155, $r = 0.99$	None	7.21 min
DZ 4480	0.028, $r = 0.99$	$\Delta 6-393$ Frz CD	34.1 min
DZ 4481	0.03, $r = 0.95$	Frz E	34.1 min
DZ 4482	0.123, $r = 0.99$	Frz G	4.15 min
DZ 4483	0.033, $r = 0.97$	Frz F	34.1 min
DZ 4486	0.055, $r = 0.98$	$\Delta 6-152$ Frz CD	1.41 min

* r values are the correlation of experimental points to the best-fitting straight line. The experimental protocol can be found in the SI.

[†]See ref. 32.

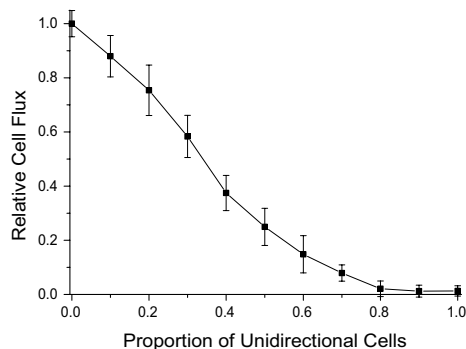


Fig. 2. Dependence of the flux on the proportion of unidirectional, wild-type cells. The average flux of 10 independent runs is plotted against the proportion of unidirectional cells in the mixture. Unidirectional cells were randomly distributed within the initial domain of cells, as described in *Methods*. A reversal period of 8 min was used for the reversing cells.

Reversals Facilitate Swarming. In the steady state, the number of cells that flow across the edge of the swarm per unit time (the cell flux), should be proportional to the rate at which the swarm expands on the macroscopic scale. The swarm expansion rate can be measured readily, and its value is used to set the model parameters. As shown in Fig. 2, the cell flux decreases gradually as the proportion of unidirectional (nonreversing) cells in a mixed population is increased in simulations. Such decreases are expected for wild-type cells (A^+S^+ in Fig. 2) and for an A^+S^- mutant (see SI). A^-S^+ mutant swarms have yet to be modeled, but they are expected to depend on reversals as well because their swarms expand at almost the same rate as the A^+S^- (9) and do have the same reversal period as A^+S^+ (23). The net fluxes diminish to zero in Fig. 2 when 100% of cells are unidirectional. Even though cells are moving, the swarms do not expand at all because unidirectional cells are easily jammed.

Seeking to understand why moving cells should need to reverse to swarm, we used the model to test how the flux would be expected to change as the length of the reversal period was changed. As shown in Fig. 3, both the A^+S^+ and the A^+S^- mutant reach their maximal fluxes at reversal periods between 5 and 12 min. The flux is significantly less on both sides of the 5 to 12 min range, and the wild-type maximum is narrower than that of the mutant. Because the flux of wild-type is twice that of the A^+S^- mutant in Fig. 3, the two engines are making equal contributions to the flux, just as they make roughly equal

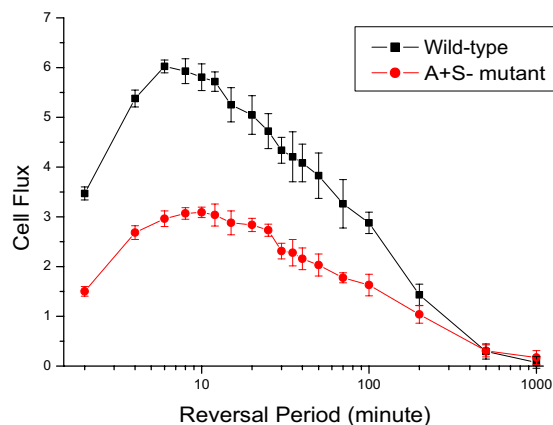


Fig. 3. The average cell flux of 10 independent runs is shown with error bars that represent the standard deviation. Flux is plotted against the reversal periods for A^+S^+ bacteria and for an A^+S^- mutant. The x-axis is scaled by \log_{10} .

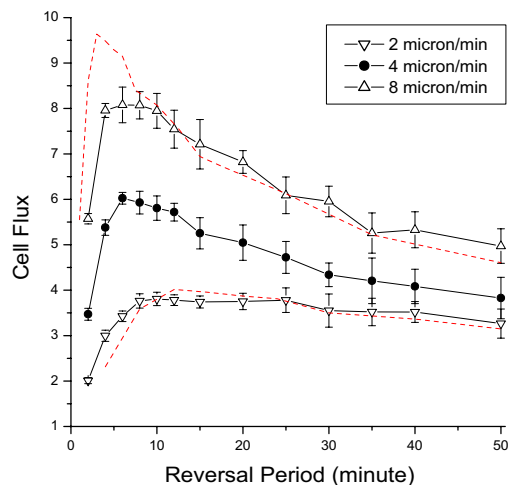


Fig. 4. Dependence of the optimal reversal period on the cell speed. Simulation results for different speeds are presented for A^+S^+ cells. The red curves are reconstructed from the data of normal speed ($4 \mu\text{m}/\text{min}$), by using the functions $Y = C1 * f(2x)$ for $8 \mu\text{m}/\text{min}$ and $Y = C2 * f(x/2)$ for $2 \mu\text{m}/\text{min}$ respectively to show the effect of scale change. Here x is the reversal period, $f(x)$ is the fitted function for the original curve of normal speed, and $C1$ and $C2$ are constant factors dependent on the average speed.

contributions to the maximum swarm expansion rates (9). Moreover, the wild-type (A^+S^+) and the A^+S^- mutant appear to have the same optimal reversal period within measurement error. For future reference, we note that the optimal value obtained by simulation corresponds to the observed reversal period of strains that have wild-type Frz systems (14, 23, 32).

Next, we investigated whether and how the optimum reversal period depended on the average speed at which the cells move. The speed of any cell is observed to vary from moment to moment as well as from cell to cell (9), possibly because cells differ in the number of pilus fibers, in the numbers of polar nozzles that happen to be secreting slime, and in whether they have stalled after colliding with another cell. The optimal reversal period was not found to be affected by noise in the distribution of reversal periods (data in the SI). Despite fluctuations around their mean, the average speed is found to be reproducible from one experiment to another (9, 33, 34). For these reasons and for simplicity, we assumed that all cells moved at the observed average speed of $4 \mu\text{m}/\text{min}$. To the first approximation, the speed would have been expected to fix the time scale for the simulation. The scaling expectations are shown by the dashed red lines in Fig. 4 for an $8 \mu\text{m}/\text{min}$ speed and a $2 \mu\text{m}/\text{min}$ speed. These speeds were chosen to lie on either side of the experimentally observed average cell speed of $4 \mu\text{m}/\text{min}$ (Fig. 4). Simulations are shown for both A^+S^+ (Fig. 4) and A^+S^- (SI). For both strains the $8 \mu\text{m}/\text{min}$ optimum (upright triangles) is shifted to a shorter reversal period, whereas the $2 \mu\text{m}/\text{min}$ optimum (inverted triangles) is shifted toward a longer reversal period. Whether deviations of both sets of points from the red line are significant (in which case they could reflect the model's nonlinear social interactions) or not, they clearly show that the optimal reversal period changes with the cell speed. Consistent with these data, we suggest that *Myxococcus* evolved an approximately 8 min reversal period because the cells had adopted an average speed of $4 \mu\text{m}/\text{min}$.

Reversal Facilitates Alignment of Cells. Fig. 1 shows the complex pattern of cells, rafts, and mounds found at the swarm edge when the 3 h movie starts. Over the movie's course, the field of the microscope starts with roughly 5,000 cells and expands outward by $\approx 20\%$. During expansion, the overall arrangement of cells at

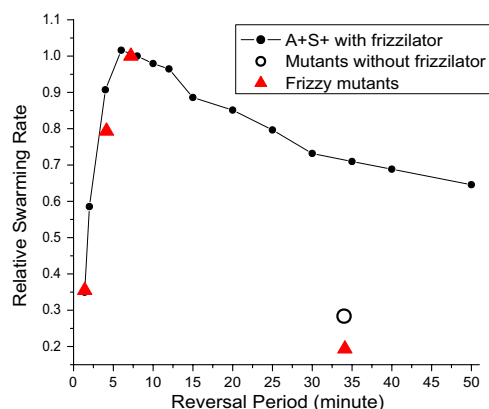


Fig. 6. Dependence of the swarm expansion rate on the reversal period. Observed expansion rates for WT and 3 deletion frizzy mutants (red triangles) are compared with those computed from the model (all are relative values). Measurement of the swarm expansion rate is described in the [SI](#).

$4\mu/\text{min}$. It was pointed out above that mutations in the frizzy genes are observed to change the average reversal period, significantly. Accordingly, the swarming model could be challenged with predicting the quantitative effect of changing the average reversal period on the flux, inasmuch as the reversal period of the frizzilator is an adjustable parameter of the model. For this purpose, we investigated two deletion mutants with short periods, one in Frz G and another in FrzCD, and three with long periods, comparing them all with wild-type cells. The measured changes in the reversal period (32) are presented in Table 1 along with our measurements of the corresponding swarm expansion rates, to which the flux is in direct proportion. As shown in Fig. 6, good quantitative agreement between observed and predicted rates was obtained for the whole-gene deletion mutation in FrzG and the N-terminal deletion of residues 6–152 of FrzCD that shorten the reversal period. However, the 3 whole-gene deletion mutants of FrzCD, of FrzF, and of FrzE, all with 34 min periods, fall below the value predicted by the frizzilator that is shown by the black line in Fig. 6. These three mutants are unique among the 5 tested in greatly and identically lengthening the reversal period to 34.1 min (32), see Table 1. Moreover, any one of those three mutations would have destroyed the capacity of the frizzilator to oscillate or to produce pulses of FrzE~P. With no pulses of FrzE~P to drive the MglAB switch, we suggest that MglAB would oscillate spontaneously between the MglA-GDP and MglA-GTP states with a period (observed) of 34 min. That period would arise from the kinetic constants of interacting MglB and MglA proteins. When the simulation is modified to approximate such an MglAB oscillator (that has no frizzilator, as detailed in the [SI](#)), a relative swarming rate of 0.28 is predicted (open circle in Fig. 6). That prediction is much closer to the Table 1 values of 0.21 observed for delFrzF, of 0.19 observed for delFrzE, and of 0.18 observed for delFrzCD.

Maximization of the cell flux also suggests that the reversal period evolved under selection for the most rapid swarming. Before 1941, *M. xanthus* had evolved in the soil of Iowa, whence it was isolated (38). Since 1941, wild-type strains DZ2 and DK1622 have been separately maintained in different laboratories. Yet, these strains exhibit similar swarm morphologies and have the same reversal period within measurement error, as indicated in Table 1. The similarity suggests that laboratory cultivation has not improved the swarm expansion rate beyond Beebe's natural isolate. The following events may have occurred after both engines had achieved their current strength and the current cell speed of $\approx 4\mu/\text{minute}$. Should the average reversal

period have ever been higher or lower than the current 8.8 min optimum, natural selection would have driven the period toward 8.8 min. The curves in Fig. 3 show that any change in the average reversal period (within the 2–50 min range) that brought it closer to an average of 8.8 min would exhibit a larger cell flux. Each such change, however small, would increase the cell flux and the swarm expansion rate, thereby promoting growth by limiting competition between cells for nutrients and would thus have a selective advantage. Targets for mutational refinement of the reversal period could include *frz CD*, *frz E*, *frz B*, *frz F*, *mglB*, and *mglA*—the protein components of the reversal generator (15). Among these targets, members of the protein complex that enables negative feedback are the most promising because their kinetics directly determine the reversal period. Mutations in those components strongly affect the swarm expansion rate (Table 1). The negative feedback helps to explain why a two-component system with adaptation by protein modification (methylation) serves as the reversal clock; such a system can readily be made to oscillate with the most appropriate period for cells that use the A and S engines to glide at $4\mu/\text{min}$. Because all these reversal period genes are conserved in the related myxobacterium, *Stigmatella aurantiaca* (39), they should have been present in the immediate precursor to *M. xanthus* and *S. aurantiaca*: this particular evolutionary scenario is certainly plausible. Traces of movement along this evolutionary path are evident in the fact that yeast or mammalian Ras, whose amino acid sequences differ from MglA protein, provide partial rescue to *ΔmglA* mutants (40).

Finally, we would like to emphasize that the flexibility and 10:1 aspect ratio of *M. xanthus* cells enhance their movement and that periodic reversals take advantage of those properties to organize cells. All swarming bacteria, including those that differentiate and move by many rotating flagella (5) and those that, like the Myxobacteria (11, 12, 13), use other engines, have long, thin cells for swarming. *M. xanthus* cells have a large aspect ratio and are unusually flexible for cells with peptidoglycan walls (25); they resemble *Bdellovibrio* (41) in that respect. As shown in the [SI](#), the swarm expansion rate should be higher for long, thin cells than for other shapes because collisions, which are inevitable among cells at high density that are moving in many directions, limit the swarming efficiency. Lacking the sophisticated senses that animals use to track their neighbors, swarming bacteria try to avoid collisions by reducing their collisional cross-section. They are also able to resolve the unavoidable collisions. It is evident watching the swarm movies that *M. xanthus* cells stall when one collides with another. The stall ends when one cell moves away or reverses. In end to side collisions, one cell, being pushed from behind by slime secretion, bends and comes along side the second cell (23). Once side by side, both cells can start to move again (see movie for Fig. 1 in the [SI](#)).

Like the type IV pili which pull cells into parallel alignments, reversals tend to orient cells along parallel tracks, thereby decreasing the probability of collision. With a reversal period of 8 min, we find (Fig. 5) that cell orientations are correlated over 10 cell lengths, or ≈ 50 microns. Without reversal, the order does not extend beyond one cell length. And, when the reversal period is too long, 50 min say, the order extends for only 3 cell lengths. The same reversal period that optimizes the flux of cells across the swarm edge also maximizes the cell order that is created. As shown in Fig. 3, optimal reversal plus the social interactions inherent in both the S engine and the A engines produce multicellular order. That order is reflected in the rafts and mounds that are seen at the swarm edge in Fig. 1. We suggest that the capacity of Myxobacteria to build species-specific fruiting bodies arises from their ability to swarm.

The *frz* genes are parts of a reversal clock that, according to our proposal, operates the MglAB reversal switch to change the gliding direction. Because the *frz* chemosensory genes are dif-

ferentiated orthologs of the chemotaxis genes of *E. coli* (42) that control the frequency of reversals of the flagellar motor, and because Frz proteins are wired up differently for the frizzilator than are the Che proteins for chemotaxis in *E. coli*, it appears that swarming and chemotaxis represent 2 different ways for motile bacteria to move to greener pastures (18).

Methods

The model for swarming of *M. xanthus* (27) has two parts describing the behavior of individual cells and the set-up for swarming (see the SI for more details).

A cell is represented by two segments connected by 3 nodes, having an observed aspect ratio of *M. xanthus* cells, $\approx 10:1$. The relative positions of the nodes can change according to an energy constraint to keep a stable cell shape. The cell is led forward by the motion of its head node. The direction is determined by local rules used in the model to reflect the properties of both engines and their social interactions (20, 27). Each cell in our model has a reversal clock, whose value increases until reaching T, the reversal period. Then the cell remains motionless for one minute, resets the clock to zero, and reverses.

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