

# Predataxis behavior in *Myxococcus xanthus*

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**Spatial organization of cells is important for both multicellular development and tactic responses to a changing environment. We find that the social bacterium, *Myxococcus xanthus* utilizes a chemotaxis (Che)-like pathway to regulate multicellular rippling during predation of other microbial species. Tracking of GFP-labeled cells indicates directed movement of *M. xanthus* cells during the formation of rippling wave structures. Quantitative analysis of rippling indicates that ripple wavelength is adaptable and dependent on prey cell availability. Methylation of the receptor, FrzCD is required for this adaptation: a *frzF* methyltransferase mutant is unable to construct ripples, whereas a *frzG* methyltransferase mutant forms numerous, tightly packed ripples. Both the *frzF* and *frzG* mutant strains are defective in directing cell movement through prey colonies. These data indicate that the transition to an organized multicellular state during predation in *M. xanthus* relies on the tactic behavior of individual cells, mediated by a Che-like signal transduction pathway.**

chemotaxis | morphogenesis | predation | rippling | chemosensory

Multicellularity has emerged several times in separate lineages of life, giving rise to plants, animals, and many other multicellular species (1). In most cases the origins of multicellularity have been obscured by time, but in general it is thought to arise through a gradual process of increasing cooperative traits between sister cells of the same organism. Many bacterial species also show multicellular organization in colony architecture or biofilm ultrastructure (2, 3). In bacteria, multicellularity is often induced by specific environmental conditions rather than as a function of programmed growth. In this report, we examine how the predatory bacterium, *Myxococcus xanthus* reorganizes its multicellular structure after invading a colony of suitable *Escherichia coli* prey.

*M. xanthus* is capable of forming several distinct multicellular structures (4). *M. xanthus* cells are motile on solid surfaces through gliding as a combination of individuals and groups ranging from 2 to 3 cells up to 10<sup>3</sup> cells in a cohesive pack. Gliding movement is driven by a combination of polar type IV pili and distributed focal adhesion sites (5–7). Under conditions of high cell density, *M. xanthus* cells form fruiting body aggregates of ≈10<sup>5</sup>–10<sup>6</sup> cells. Within fruiting bodies, low-nutrient availability induces vegetative rods to differentiate into nonmotile spores (8). During predation, *M. xanthus* cells organize into rippling wave structures. Rippling behavior coordinates cell movement across areas of several square millimeters and can involve >10<sup>8</sup> cells working as a unit.

Multicellular rippling in *M. xanthus* is a phenomenon that is superficially similar to many patterns displayed by other biological and chemical systems (9). Yet, it is unique in that counter migrating cells reflect off each other producing a convective rather than diffusion-based wave pattern (10). One model is that rippling constitutes an intermediate phase of cell organization before a more permanent fruiting body structure (11). However, rippling is not observed under all conditions that promote fruiting body formation (12). Additionally, rippling is induced by *M. xanthus* during predation on a variety of microbial species or degradation of macromolecular growth substrates such as peptidoglycan, protein, and chromosomal DNA (13, 14). An alternative hypothesis is that rippling occurs as a result of directed cell

movement to enhance contact with prey macromolecules to facilitate predation.

In swimming *E. coli* cells, individuals find optimal positions within spatial gradients of chemoeffectors by altering their behavior (smooth swimming and tumbling), a process controlled by the chemotaxis (Che) signal transduction pathway (15). Nonchemotactic mutants are less competent than chemotactic strains in both capillary assays and infection models (16, 17). Each individual cell makes decisions on the basis of its own perception of the local chemical environment, yet complex multicellular patterns can emerge, including expanding swarm rings and focal aggregates (18, 19). Chemical signals are sensed through an array of chemoreceptors and the signal is transduced to the flagellar motors to control the swimming behavior of cells (20). The sensitivity of the receptors in Che-like signal transduction is subject to methylation via the opposing biochemical activities of the methyltransferase (CheR) and methyltransferase (CheB). Altering the sensitivity of the receptors allows cells to adapt their behavior to subtle changes across a wide range of chemical concentrations. *E. coli* *cheR* and *cheB* mutants are ineffective at directing cell movement through chemical gradients as cells are locked in either a constitutively smooth swimming or a constitutively tumbling behavioral state (15).

In *M. xanthus*, the Che-like Frz pathway has previously been shown to regulate the frequency at which cells change the leading pole during surface gliding, similar to regulation of flagellar rotation in *E. coli* (21). Therefore, if predatory rippling in *M. xanthus* depends on the behavior of individuals similar to that seen during *E. coli* chemotaxis, then we predict that the Frz pathway will be required for controlling this behavior. If this hypothesis is correct, *M. xanthus* cells should display the two hallmarks of bacterial chemotactic behavior during predatory rippling: directed movement and adaptation to a stimulus.

## Results

**Directed Movement During Cellular Reorganization to Rippling.** To analyze predatory behavior we make use of *M. xanthus* strain DZ2, which has a low intrinsic level of autolysis and *E. coli* strain  $\beta$ 2155, which is a diaminopimelic acid (DAP) auxotroph whose growth is restricted in the absence of DAP, allowing us to analyze *M. xanthus* behavior in relation to an unchanging source of prey. During predation assays, *M. xanthus* cells organize into dramatically different structures depending on whether they are inside or outside of the prey colony (Fig. 1A). In the absence of prey, *M. xanthus* cells glide in amorphous groups such that no stable pattern of organization can be observed after 1 h of incubation (Fig. 1B). In contrast, cells within the prey colony organize into dynamic, parallel rippling waves and maintain this organization over time (Fig. 1C). In both the presence and absence of prey, cells are actively motile and glide at similar velocities: 2.7 +/-

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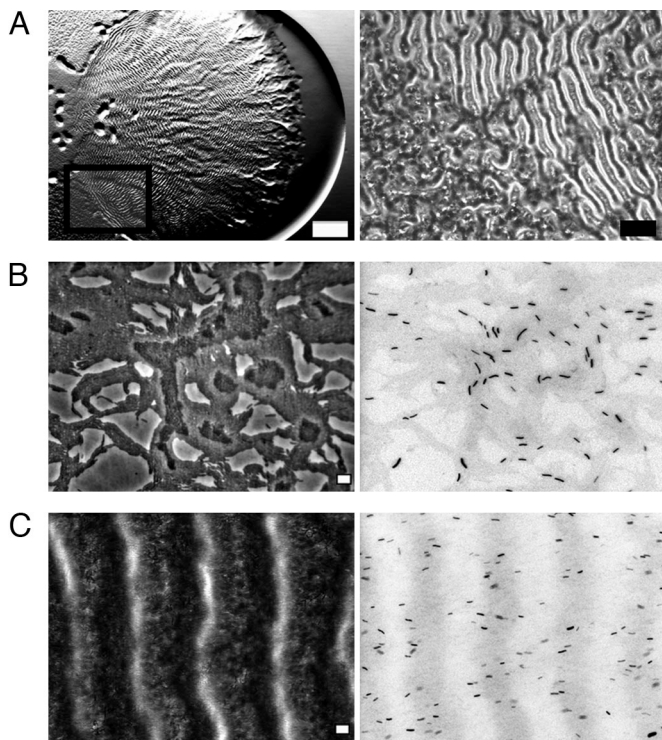
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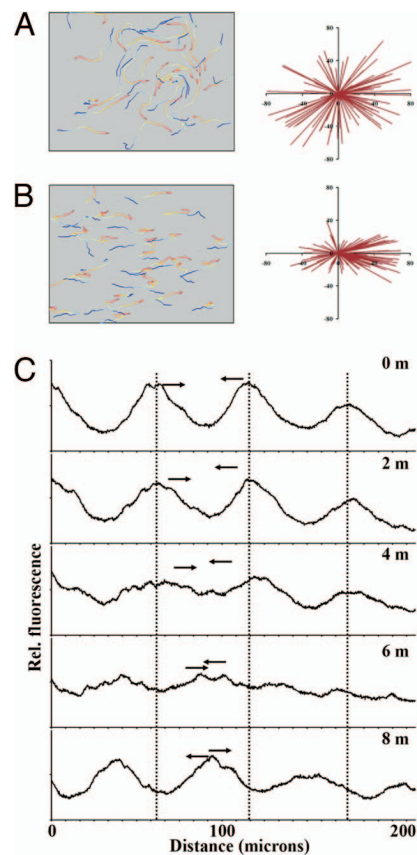


**Fig. 1.** Multicellular reorganization during predation. (A) Stereo microscopy (Left) of *M. xanthus* strain DZ2 penetrating an *E. coli* prey colony and inducing rippling behavior. The image is centered on the prey colony, where a majority of the prey cells have been lysed and are no longer visible, but a fraction of the remaining *E. coli* colony is visible as a dark crescent on the Right edge. The open box denotes the area of the phase contrast image (Right) which is centered on the near edge of the prey colony where the transition from nonrippling to rippling behavior can be observed. (White scale bar, 1 mm; Black scale bar, 100  $\mu\text{m}$ .) (B) Phase contrast (Left) and fluorescence microscopy (Right) of DZ2 mixed in a 50:1 ratio with GFP-labeled DZ10547, showing the characteristic mesh pattern of *M. xanthus* gliding behavior in the absence of prey. (C) Within the prey colony, the rippling pattern is stably maintained, although individual cells change position. (White scale bars in *b* and *c*, 10  $\mu\text{m}$ .)

0.9  $\mu\text{m}/\text{min}$  in the absence of prey and  $2.8 \pm 0.7 \mu\text{m}/\text{min}$  in the presence of prey.

To determine whether *M. xanthus* cells display directed movement during predatory rippling, *M. xanthus* strain DZ2 was mixed with a strain carrying GFP fused to the highly expressed *pilA* promoter and fluorescence microscopy was performed in the presence and the absence of prey [Fig. 1 *B* and *C* and supporting information (SI) Movies 1 and 2]. Tracking of cells was performed by mixing DZ2 cells with GFP-labeled cells on an agar-covered microscope slide and indicates that in the absence of prey, *M. xanthus* cell movement occurs in curved trajectories, with changes of direction occurring gradually through cell bending and collisions with other cells, rather than through oscillation of the leading cell pole (Fig. 2A). Net movement of cells was analyzed by plotting the total displacement of 100 individual cell displacement vectors after 30 min of movement (Fig. 2A). Net displacement occurs in all directions, with various magnitudes, and the multicellular makeup of the group consists of an amorphous pattern with no relationship to the initial pattern in the absence of prey stimuli (Fig. 1).

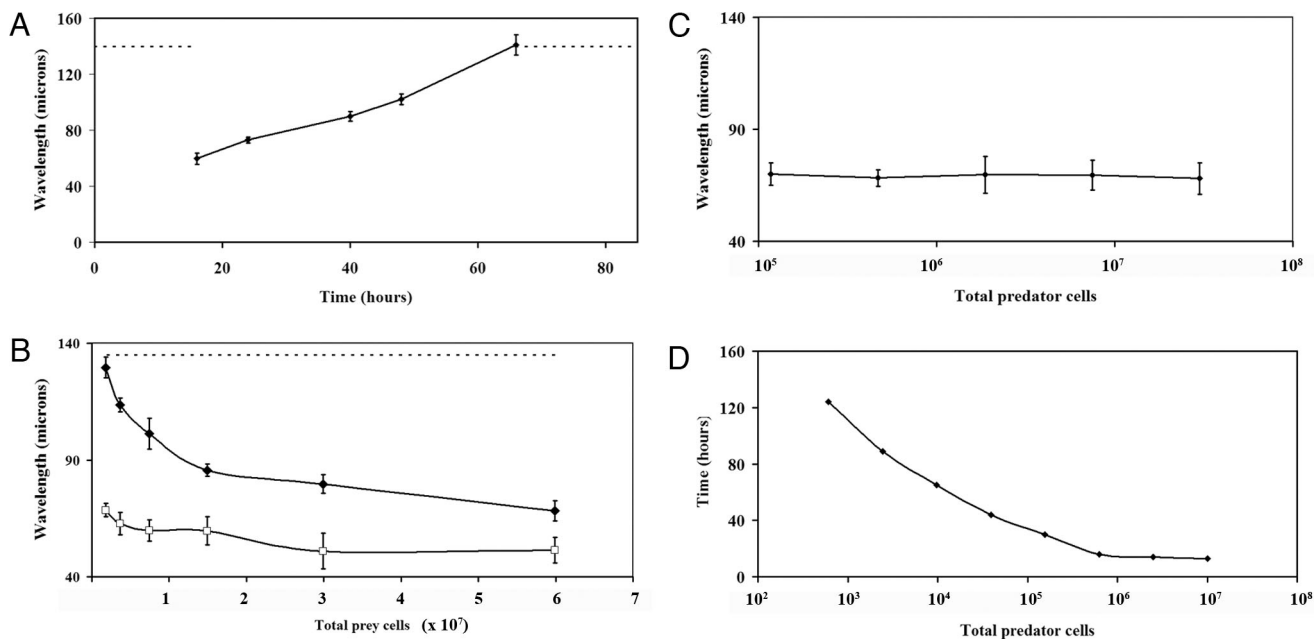
Previous studies on cell behavior during rippling led to the conclusion that there is no net cell displacement, i.e., no directed movement (22). However, those experiments were performed in closed systems where the ripple-inducing substrate was distributed throughout the sample. To examine rippling behavior in an open system, *M. xanthus* strain DZ2 cells were mixed with



**Fig. 2.** Tracking movement of *M. xanthus* cells. Tracking of GFP-labeled cells in (A) the absence of prey and (B) the presence of *E. coli* prey. Experimental setup is the same as in Fig. 1. Thirty cell tracks are shown for each condition. Cell movement over time is represented by the change in color of each track, with the beginning of the track labeled in dark blue and the end in dark red. Vector displacements of 100 cells for each condition are shown on the Right. The direction of migration through the *E. coli* prey colony is from Left to Right. (C) Distribution of cells was monitored at 2-min intervals by measuring fluorescence intensity across the entire field of view. The dashed line indicates the position of the wave crests at 0 min, and the arrows show the general direction of cell movement in the given area. The wave structure dissipates as *M. xanthus* cells leave the initial wave aggregates, but a similar pattern of aggregation emerges, shifted by  $\frac{1}{2}$  wavelength, at 8 min.

GFP-labeled cells and pipetted adjacent to an *E. coli* strain  $\beta 2155$  prey colony on an agar-covered microscope slide. *M. xanthus* ripples can be observed as dense bands of autofluorescence that aggregate and dissipate in a synchronized manner with a period of  $\approx 8$  min (Figs. 1C and 2C). New ripple aggregates that form are shifted by  $\approx \frac{1}{2}$  wavelength relative to the previous ripple crests (Fig. 2C). Analysis of individual cell behavior indicates that movement occurs along primarily linear paths that are perpendicular to the orientation of the multicellular ripple structure (compare Figs. 1C and 2B). Thus, there is little movement in the *y*-axis and cells are constrained to moving along a nearly one-dimensional path. Both the direction and magnitude of the displacement vectors are biased to the positive *x*-axis, the direction of swarm migration through the *E. coli* prey colony. Thus, although rippling behavior can last for days, the individual structures survive only for a few minutes and seem to be indicative of a shifting equilibrium of *M. xanthus* cell density rather than the intermediate construction of a larger multicellular structure.

**Adaptation to Prey Stimuli.** Although rippling is a stable multicellular behavior, it takes time for the cells to coalesce into waves



**Fig. 3.** Quantitative analysis of rippling pattern dynamics. (A) Rippling wavelength as a function of incubation time during predation. A 1- $\mu$ l suspension of  $10^6$  total *M. xanthus* strain DZ2 cells pipetted onto CFL media adjacent to a 3- $\mu$ l colony containing  $5 \times 10^7$  cells of *E. coli*  $\beta$ 2155 prey. The dashed line indicates times when quantifiable rippling was not observed. (B) Rippling wavelength at 40 h changes as a function of *E. coli*  $\beta$ 2155 prey cell density ( $2 \times 10^6$  to  $6 \times 10^7$  total cells) in *M. xanthus* strain DZ2 (diamonds), the wavelength is constitutively short in  $\Delta frzG$  (open squares), and no rippling is observed in  $\Delta frzF$  (-). (C) Rippling wavelength shows no change as a function of *M. xanthus* strain DZ2 cell density at 40 h ( $2 \times 10^5$  to  $4 \times 10^7$  total cells) incubated with  $5 \times 10^7$  cells of *E. coli*  $\beta$ 2155 prey. (D) Rippling induction time changes as a function of *M. xanthus* strain DZ2 cell density ( $6 \times 10^2$  to  $1 \times 10^7$  total cells) incubated with  $5 \times 10^7$  cells of *E. coli*  $\beta$ 2155 prey.

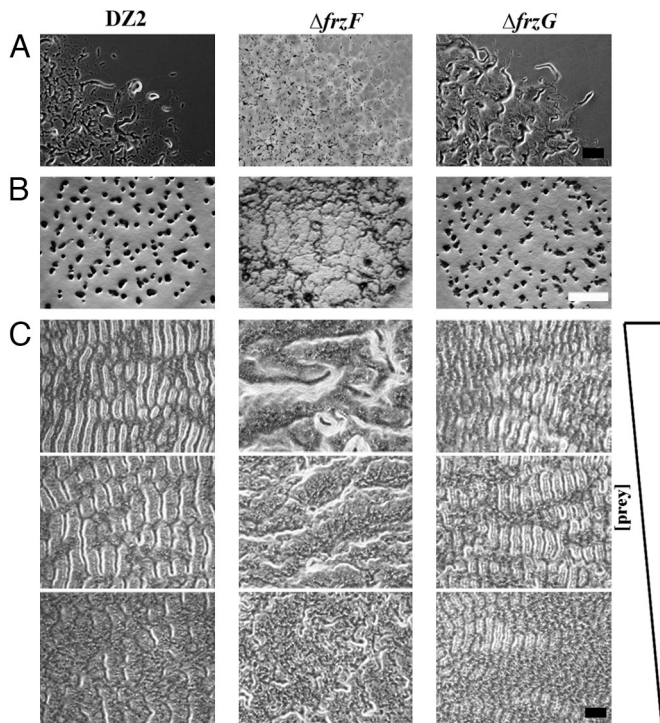
and rippling terminates soon after the available prey macromolecules have been consumed, such that rippling behavior lasts for a finite interval. To determine whether changes occur throughout the duration of rippling behavior, *M. xanthus* cells were pipetted adjacent to *E. coli* cells on a cloned fruiting light (CFL) low-nutrient agar plate and the distance between wave crests was measured from the beginning to the end of predation (Fig. 3A). There is entry of *M. xanthus* cells into the prey colony, but no measurable rippling during the first 18 h of the assay. After initiating, the ripples display a short wavelength of  $\approx 60 \mu\text{m}$ . There is a gradual increase from 18 to 66 h in the wavelength of the ripples with a maximum wavelength of  $140 \mu\text{m}$ . After 66 h, rippling behavior was still occasionally observed in scattered areas of the colony, but was not quantifiable.

To determine whether changes in rippling behavior occur over time as a result of adaptation by *M. xanthus* cells to the gradual loss of prey, predation assays were performed in which various levels of prey were provided, while the initial *M. xanthus* cell density was held constant (For quantification see Fig. 3B, for images see Fig. 4C). High densities of prey stimulate formation of a large quantity of ripples with a short wavelength between crests. Lower cell densities of prey result in longer wavelengths until a threshold is reached at which rippling is no longer observed. The output wavelength observed during rippling ranges from 70 to  $135 \mu\text{m}$ , consistent with the changes observed over time (Fig. 3A). It is expected that the presence of prey will also cause a gradual increase in the *M. xanthus* cell density as the prey are consumed and the *M. xanthus* cells grow and divide. Thus the changes in wavelength observed could be due, in part, to increasing predator cell density. To determine whether the *M. xanthus* cell density also impacts the rippling wavelength, we performed similar assays in which the predator cell density was varied, while the prey cell density was held constant (Fig. 3C). Changes in the initial *M. xanthus* cell density have little effect on

the output wavelength of rippling. At the cell densities tested, no significant change in ripple wavelength was observed.

Although the predator cell densities tested had no effect on ripple spacing, analysis of a wider range of initial cell densities indicates that the cell density of *M. xanthus* is critical for the timing of rippling induction. Fig. 3D shows the average time required for 10 parallel waves to appear. The resulting curve is biphasic, with a steep slope below  $5 \times 10^5$  cells and a shallow curve at higher cell densities. This indicates that at predator densities at and above  $5 \times 10^5$  cells, rippling is able to initiate rapidly, with the only delay being the result of factors such as biosynthesis of motility organelles, sensing of prey, and the time required to reorganize into ripples. At lower predator densities there is a longer delay, which is likely the result of insufficient *M. xanthus* cell density to facilitate rippling. Taken together these results indicate that rippling wavelength is adaptable and in proportion to prey cell availability.

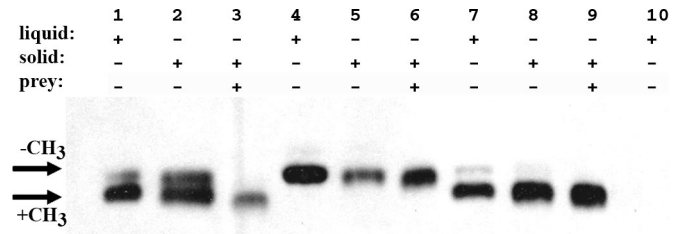
**Molecular Controls of Adaptation.** In *E. coli*, adaptation of cell motility to chemical stimuli occurs through the reversible methylation of the methyl-accepting chemotaxis proteins (MCPs) catalyzed by the methyltransferase, CheR, and the methyl-erase, CheB (23). The homologs in the *M. xanthus* Frz pathway that modify the FrzCD receptor are FrzF and FrzG, respectively (20). While cells lacking FrzF reverse direction of movement very rarely in comparison to wild type, a deletion of *frzG* does not lead to any obvious defect in the reversal frequency (24). To determine the impact of the Frz pathway on multicellular structures, we examined the wild-type,  $\Delta frzF$ , and  $\Delta frzG$  strains during gliding, fruiting body formation, and rippling behaviors (Fig. 4). During colony expansion, wild-type and  $\Delta frzG$  cells glide in a meshed pattern of individuals and groups (Fig. 4A). In contrast,  $\Delta frzF$  displays a dispersed pattern of individuals and very small groups. Under low-nutrient, high-cell-density conditions, distinct fruiting body aggregates can be observed forming in the



**Fig. 4.** Cooperation analysis in adaptation mutants. A 10- $\mu$ l aliquot containing  $10^7$  *M. xanthus* strain DZ2 cells was pipetted on CFL agar in the absence of prey. Stereo and phase microscopy images were captured at 72 h at (A) the colony edge at which group gliding behavior can be observed and (B) the colony center where fruiting body aggregation is observed. A  $\Delta$ *frzF* mutant is defective in forming gliding groups and also forms unorganized “frizzy” aggregates.  $\Delta$ *frzG* displays a phenotype with only subtle differences from the wild type in the absence of prey. (C) After 40 h in coculture with *E. coli* prey, wild-type cells modulate their rippling wavelength in response to the availability of prey. The  $\Delta$ *frzF* mutant does not form ripple structures. The  $\Delta$ *frzG* mutant forms numerous ripples at all prey cell densities in which rippling is induced. The direction of migration through *E. coli* prey is from Left to Right. (Black bar, 100  $\mu$ m in a and c; White bar, 1 mm (b).)

wild-type and  $\Delta$ *frzG* strains, but the  $\Delta$ *frzF* mutant displays disorganized, frizzy aggregates of cells (Fig. 4B) as described previously (20). In the presence of prey, wild-type *M. xanthus* ripples in accordance with the abundance of prey (Fig. 4C). The  $\Delta$ *frzF* mutant does not show rippling behavior at any of the prey cell densities tested. The  $\Delta$ *frzG* mutant, however, shows a distinct hyperrippling phenotype, in which ripple structures are tightly packed resulting in a constitutively short output wavelength regardless of the cell density of prey provided (for quantification, see Fig. 3B). This hyperrippling phenotype is consistent with an inability to adapt to decreasing prey availability. Rippling was not observed in either the wild-type strain or the  $\Delta$ *frzG* mutant below  $10^6$  total prey cells.

To determine whether the rippling defects observed in the  $\Delta$ *frzF* and  $\Delta$ *frzG* mutants are the result of a biochemical role in which the FrzCD MCP homolog is incorrectly methylated during predatory rippling, we isolated cells from rippling and nonrippling conditions and probed for the presence of the methylated and demethylated forms of the FrzCD receptor using  $\alpha$ -FrzCD antibody. Previous work has demonstrated that altering methylation of MCPs affects mobility under standard SDS/PAGE conditions and can be detected using immunoblot analysis (25). The apparent lower molecular weight form of FrzCD corresponds to a more highly methylated version of the receptor while demethylation of FrzCD produces an apparent increase in molecular weight (26).



**Fig. 5.** Immunoblot analysis of the methylation state of the FrzCD receptor. Cells of DZ2 (wild type, lanes 1–3),  $\Delta$ *frzF* (*cheR*, lanes 4–6), and  $\Delta$ *frzG* (*cheB*, lanes 7–9) were harvested after 24-h growth in liquid CYE (lanes 1, 4, and 7) and plated on CFL media ( $10^8$  total cells) either in the absence (lanes 2, 5, and 8) or presence of  $10^9$  *E. coli* prey cells (lanes 3, 6, and 9). Cells were harvested after 24 h at 32  $^{\circ}$ C and all samples were analyzed using SDS/PAGE to separate the methylated and demethylated forms of FrzCD, as described previously (26). *frzCD<sup>c</sup>*, a strain that expresses a truncated form of FrzCD, was used as a negative control (lane 10).

Immunoblot analysis of strain DZ2 after GYE in nutrient-rich liquid medium shows that both methylated and demethylated forms of FrzCD are detectable (Fig. 5). Both forms of FrzCD were also detected when cells were incubated on a solid surface for 24 h in the absence of prey. However, in the presence of prey, FrzCD can only be detected in the methylated state. In a  $\Delta$ *frzF* methyltransferase mutant, FrzCD was detected constitutively in the demethylated state under the three conditions tested (Fig. 5, lanes 4–6). In a  $\Delta$ *frzG* methyltransferase mutant, the FrzCD protein was detected primarily in the methylated state (Fig. 5, lanes 7–9). Together, these data indicate that the presence of prey can alter the methylation state of the FrzCD receptor and that both the FrzF and FrzG proteins are required to modify the FrzCD methylation state to bring about adaptation in response to prey.

**Adaptation Components Are Essential in a Predatory Taxis Assay.** *E. coli cheB* and *cheR* mutants have opposite behavioral phenotypes; *cheB* is tumbling and *cheR* is smooth swimming. Yet, both mutants are defective in their ability to direct movement in various chemotaxis assays (15, 27, 28). Similarly, the  $\Delta$ *frzF* and  $\Delta$ *frzG* mutants show opposite phenotypes with respect to rippling behavior;  $\Delta$ *frzF* does not ripple, whereas  $\Delta$ *frzG* ripples at a constitutively short wavelength. To determine the effect of these mutations on general predation ability, we devised a predatory taxis assay. In this assay, *M. xanthus* cells are pipetted into the center of a long strip of *E. coli* prey cells and the swarm expansion is measured in the dimension containing prey (*x*-axis) and the dimension lacking prey (*y*-axis) (Fig. 6A). In the absence of prey, *M. xanthus* strain DZ2 expands uniformly at a rate of  $\approx 100$   $\mu$ m/h (Fig. 6B). This rate is constant across a wide range of basal nutrient levels (0.01–10 g/liter casitone). In the presence of prey, the rate of swarm expansion in wild type is unchanged during the first 24 h when rippling is not observed. Soon after rippling induction, swarm expansion increases to a rate of  $\approx 190$   $\mu$ m/h. In a  $\Delta$ *frzG* mutant the rate of swarm expansion is  $\approx 100$   $\mu$ m/h both in the presence and absence of prey, even after the induction of rippling behavior (Fig. 6C). In the  $\Delta$ *frzF* mutant, swarm expansion is maintained at  $\approx 100$   $\mu$ m/h in the presence of prey. In the absence of prey, the rate of swarm expansion of  $\Delta$ *frzF* begins at 100  $\mu$ m/h during the first 24 h but decreases over time to a rate of 50  $\mu$ m/h during the last 24 h of the assay (Fig. 6D). Together, these data indicate that adaptation via methylation of FrzCD is required for directed movement during predation.

## Discussion

During bacterial chemotaxis, the methylation state of the receptors is regulated by the adaptation proteins CheB and CheR



washed twice and  $10^8$  *M. xanthus* cells were spread on CFL plates either with or without  $10^9$  *E. coli* strain  $\beta$ 2155 cells. Plates were incubated for 24 h at 32 °C. Rippling behavior was confirmed and cells were harvested from plates. Cells were resuspended in TE buffer and sonicated. The soluble protein fraction was normalized to 5  $\mu$ g/ml using Bio-Rad protein assay, boiled in SDS and separated using SDS/PAGE. Protein was transferred to polyvinylidene fluoride (PVDF) membranes and detected using anti-FrzCD antibody (primary) used at 1/10,000 and anti-IgG (rabbit)-HRP conjugate (secondary) at 1/15,000, similar to previously described method (21).

**Predataxis Assay.** Forty microliters containing  $4 \times 10^8$  *E. coli*  $\beta$ 2155 cells were pipetted in 2  $\mu$ l adjacent aliquots to generate a thin strip of prey with

dimensions of 20 mm  $\times$  2.5 mm on a CFL agar plate. A 1- $\mu$ l aliquot containing  $\approx 10^6$  *M. xanthus* cells was pipetted in the center of the prey strip. Swarm expansion was measured at the time indicated in two axes; the long axis of the *E. coli* strip is treated as the x-axis and the short axis as the y-axis. Measurements were collected from three independent samples for each strain.

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