Dynamics of pattern formation in bacterial swarms

Edward B. Steager, Chang-Beom Kim, and Min Jun Kim

Department of Mechanical Engineering and Mechanics, Drexel University, 3141 Chestnut Street, Philadelphia, Pennsylvania 19104, USA

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To gain a more thorough understanding of the dynamics of swimming bacteria, a nonlabeled cell tracking algorithm was used to study the velocity field of flagellated bacteria, Serratia marcescens, swimming on a soft agar plate. The average velocities for local regions regularly arranged over the entire flow field were investigated. The velocity field of the bacteria typically featured the combination of curvilinear translation and vortex modes. They repeated these patterns for short periods of time, forming several groups and dissipating. To further investigate the flow patterns generated by the collective motion of the swimming bacteria, the velocity field on the swarm was spatially correlated. The highest velocities and correlation lengths have been found to occur in the region from 0.5 to 1 mm from the swarm edge, followed by a steady decline as distance from the edge increases, and a sudden decrease in motion typically occurs between 2 and 4 mm from the swarm edge. © 2008 American Institute of Physics. [DOI: 10.1063/1.2953245]

INTRODUCTION

Bacteria such as Serratia marcescens or Escherichia coli swim in liquid environments by rotating thin helical flagellar filaments. Other types of locomotion in the form of surface translocation can be found on semisolid surfaces and are referred to as swimming or gliding. It has been commonly believed that all their motions are powered by the reversible rotary motor embedded in the cell membrane. On soft agar surfaces, bacterial colonies have been observed to grow rapidly through a collective mechanism.

Swarming is a crucial behavior of organisms which span their territories or population in an environment. Among them, bacterial swarming is a form of flagella-dependent surface motility. The flagellar motions driven by the bacterial rotary motor eventually result in the collective motion of the swimming bacteria. Swarming bacteria are hyperflagellated, elongated, and migrate cooperatively. Characteristic dynamic patterns of whirls and jets have been observed both in swimming bacteria as well as in high-cell density swimming populations, and the interplay of the roles of sensory mechanisms and hydrodynamic interactions is widely researched. Although communication mechanisms between swarming S. marcescens are not fully understood, research suggests that pattern formation in the active swarm may be influenced largely by hydrodynamic interactions between densely packed cells. Large-scale dynamic coherence has been observed in swimming populations of Bacillus subtilis. This phenomenon of dynamic coherence has been confirmed by simulation by accounting for hydrodynamic interactions of suspensions of self-propelled particles at low Reynolds number. Results of bacteria swimming in thin fluid films also provide evidence for the pure hydrodynamic origin of collective swimming. Further experimental work with small copper rods reveals that inanimate objects exhibit swarmlike formations due to nonspecific interactions. In fact, these rods do not demonstrate swimming behavior unless the ends are rounded, resembling the elongated bacillus shape of swimming bacteria. Analytical models of bacterial swarming also reveal that swimming and swarming formations may be a result of physical interactions between individual bacteria. This research lends credence to an effort to understand the larger scale dynamics of the swarm. That is, the study of swarming formations may reveal dynamics consistent with a collective effort to rapidly populate environments which may otherwise remain unpopulated.

Recently, associated with the function of the flagella, bacteria have been used as actuators in microscale engineered devices. Bacterial monolayers, referred to as bacterial carpets, were created on microstructures by using a blotting method, and the swarming pattern is directly reproduced on the bacterial carpet. Controlled manipulation may be exerted by exploiting the existing bacterial chemical sensory mechanisms. However, the swarm plate dynamics have not been well understood. In this study, we employ a nonlabeled cell tracking method to trace the bacterial swarming flow field without fluorescence labeling or microspheres as tracers. This research is intended to lend fundamental insight into ongoing bacterial actuation studies with the specific goal of understanding the large-scale collective dynamics of the swarm through a study of pattern formation.

METHODS

The bacteria S. marcescens (ATCC 274, American Type Culture Collection, Manassas, VA) were cultured and grown on a swarm plate. 10 g of Difco Bacto tryptone, 5 g of yeast extract, 5 g of NaCl, and 6 g of Difco Bacto agar were dissolved into 1000 ml of de-ionized water to prepare 0.6% agar plates for swimming bacteria, followed by autoclaving the solution and dividing into 100 ml sterile bottles. This solution was stored at room temperature and solidified and was reliquefied using a microwave on the lowest power set-
are covered with numerous (10–100) petrichously or laterally situated flagella.20

Ideally, every individual bacterium would be traced and analyzed for velocity field analysis. If this were possible, a typical particle tracking velocimetry scheme could be used for velocity calculations. However, as shown in Fig. 2(b), individual bacteria swarming in close proximity have overlapping or otherwise unclear boundaries due to close contact and inherent resolution limits of light microscopy. Also, the distance of the bacterial monolayer measured from the leading edge of the swarm to the development of a multilayer varied from 15 to 100 μm along the swarm edge. To create a solution to these difficulties, small interrogation windows (20×20 pixel, 4.3×4.3 μm²) for locally averaged velocity analysis were defined. It should be noted that this window only contains a total of five to ten swarming bacteria on average. As shown in Fig. 2(a), the interrogation windows were repeated with 10 pixel shifts (50% overlap) both in the horizontal (x) and vertical (y) directions over the entire investigation region. For the current study, the total number of interrogation windows is 1296 (36×36 windows). Each interrogation window included several bacteria which were generally heading toward similar directions. Along the edges of these windows, only the fraction of the individual cell bodies present in the window was included for calculation. To investigate an averaged velocity to compensate for all bacterial motions involved in an interrogation window, two temporally consecutive (images at time \( t \) and \( t+Δt \), \( Δt = 1/15 \) s in this study) but spatially identical interrogation windows were compared by shifting the latter interrogation window (at time \( t+Δt \)) by 1 pixel in every major or diagonal direction with respect to the center of the former interrogation window (time \( t \)). The minimum change in pixel intensity value was found by subtracting one window from the other and consequently obtaining the new center of the shifted interrogation window in the image at time \( t \). This indicates that shifting with the bulk movement of a small group of bacteria generally caused the minimum difference. After one process of single-pixel shifting, the next shifting process started over with respect to the new center of the shifted interrogation window (still at time \( t \)) with the new interrogation window which is again temporally consecutive and spatially the same. This process of single-pixel shifting was performed in order of “top to bottom” and “left to right” and was repetitively performed ten times for every time interval for all interrogation windows over the entire investigation region and also for all 300 images. By shifting ten times for each window, a very precise measurement can be made for each velocity vector, but to ensure accuracy, the time interval \( Δt \) must be small enough so that temporally consecutive windows do not shift to a degree greater than the scale of the interrogation window itself.

To further investigate the patterns of the entire flow field generated by the collective motion of the swarming bacteria, the velocity field on the swarm was spatially correlated. The spatial correlation function offers an insight to the degree to which the swarming bacteria are coordinated in some direction for any given area of interest. The spatial correlation function is defined as...
\[ \Omega(s) = \frac{\langle \tilde{V}(r) \cdot \tilde{V}(r + s) \rangle}{\langle \tilde{V}(r) \cdot \tilde{V}(r) \rangle} \]

where the angular brackets represent an ensemble average, \( \tilde{V} \) is the averaged velocity for each interrogation window, \( \vec{r} \) is the position vector of the center of each window, and \( s \) is the vector between the centers of two windows being interrogated. Once two velocity vectors are highly correlated with each other, i.e., “aligned” with an acute angle between two vectors, the inner product for the velocities yields a positively large value, while orthogonally oriented velocities yield zero and obtuse-angled vectors yield negative values. The correlation length can be determined by integrating \( \Omega \) with respect to the in-between distance \( s \).

Although cell culturing is performed with the same batches of nutrients and chemicals, swarms demonstrate noticeable variations. This is likely due to minute changes in the characteristics of the immediate surroundings of the bacteria such as variations in surface moisture, surface topography, or local nutrient content. Due to these inherent variations between different swarms and along the edges of a single swarm, multiple images and data sets were captured for an averaged analysis of swarm motility.

**RESULTS AND DISCUSSION**

The computed flow fields qualitatively matched the motion of the original image sequences upon visual inspection with the computed vectors superimposed on the original video. The velocity field of the swarming bacteria typically featured the combination of curvilinear translation and vortex modes. The bacteria formed several groups over the entire investigation region for a short time period, mostly translating along rather curved paths. They joined temporarily and joined other streams, forming another translational motion, the near-edge regions have the largest average velocities. Also, typically between 1 and 4 mm from the edge, the bacteria move more freely due to the relative motility of the neighboring bacteria. An analogy can be made between the classic parabolic profile of viscous flow in a channel and the region between the edge of the swarm and 0.45 mm. It should also be considered that the bacteria secrete serratexin, a surface active exolipid which wets the agar surface and enhances motility.

As the region of interest passes 0.45 mm from the swarm edge, the motility decreases. This may be due to entanglement of flagella between adjacent cells as density increases. Also, typically between 1 and 4 mm from the edge, a multilayer wave of immotile bacteria consistently overcomes the swarm. There is no coordinated movement after this wave passes in the sense that the bacteria do not exhibit collective motion in the form of streams and vortices. However, there is still an extremely slow progression of cells toward the swarm edge. Keeping in mind that the cells in this region are as yet quite alive and continue to reproduce, this is most likely due to the forces of biomass production causing branched off from the main streams and joined other streams, but some formed vortices, which again merged into an adjacent stream or disappeared shortly.

There is a consistent increase in average velocity from the edge of the swarm to roughly 0.45 mm. At the edge of the swarm, individual bacteria are unable to move due to the semisolid, highly viscous nature of the agar. As the region of interest progresses further from the edge of the swarm, the bacteria move more freely due to the relative motility of the neighboring bacteria. An analogy can be made between the classic parabolic profile of viscous flow in a channel and the region between the edge of the swarm and 0.45 mm. It should also be considered that the bacteria secrete serratexin, a surface active exolipid which wets the agar surface and enhances motility.

![Figure 3](image1.png) Instantaneous bacterial swarming flow field (a) at the edge and (b) 4 mm from the swarm edge. The cells revealed complex flow patterns composed of curvilinear motions and vortices. The arrow (→) denotes a speed of 35 μm/s.

![Figure 4](image2.png) (Color online) Average velocities at different locations from the swarm edge using multiple data sets. Due to strong vortices and translational motion, the near-edge regions have the largest average velocities.

![Figure 5](image3.png) Average velocities at different locations from the swarm edge using multiple data sets. Due to strong vortices and translational motion, the near-edge regions have the largest average velocities.
a form of plug flow of entangled, immotile cells. Due to the fact that Fig. 4 represents the average of several swarms, this appears as a linearly declining velocity when, in fact, for each individual swarm, velocity slowly decreases after 0.45 mm, then collective motion suddenly stops as the immotile wave is reached.

To further quantify the complex combination of the bacterial behaviors on the swarm plate, the spatial correlation function for the velocity field was obtained as a function of distance between individual interrogation windows over 0.5 and 4 mm investigation domains.

The correlation length was determined by integrating the normalized correlation between the averaged velocities for each interrogation region. The figure indicates that immediately neighboring bacteria are headed in the same direction with less directional correlation as distance between bacteria increases. Where the values of the spatial correlation function fall below 0.1 (distances of 30–70 µm), the distance between flanks of both sides of the humps in the curves indicates the diameter of vortices.\(^{17}\) The average intervortex distance ranged from 30 to 35 µm.

The correlation length was determined by integrating the correlation function over distance (Fig. 6). The mean correlation length was 9.4 ± 1.2 µm for the region within 1 mm from the edge. Bacteria very near the swarm edge (10–100 µm) are slowed due to viscous interactions with the set of bacteria which are directly situated on the leading edge of the swarm (0–10 µm from the swarm edge). These bacteria of the edge of the swarm are themselves unable to move and may remain in place for tens of seconds before being pushed by the swarm. However, bacteria at 0.5 mm are less restricted and thus able to attain greater correlation length. The maximum correlation occurs at roughly 0.9 mm. The slight increase from the edge to 0.5 mm is again likely due to the lack of influence of immotile bacteria at the swarm edge. The correlation length decreases past 0.9 mm from the edge due to flagellar entanglement, and the sudden drop around 3–4 mm can again be attributed to the inactive waves of entangled bacteria that follow the freely moving near-edge region.

This complex pattern is driven by surging bacterial crowds whose direction is not predictable. Sudden formations of helically directed surges generate large velocity gradients at a local region and replenish the stream with the local cells which then return into transient vortex domains. The average velocities and spatial correlation lengths reveal that the swarm enables rapid population of semisolid surfaces. Bacteria with the greatest velocities and correlations are active along the edge. The consequence of this is that collectives of cells act as battering rams against the immotile bacteria along the swarm edge. This hammering motion pushes the otherwise immotile cells along the very edge of the swarm enabling wetting of the surface. The high momentum due to surging motion of several hundred or even thousand cells enables the swarm to rapidly proceed. The seemingly random nature of the surges averages across the edge of the swarm with several surges occurring within a minute near any given point along the edge. The cells decrease in motility and eventually stop moving completely in the region between 2 and 4 mm inside the swarm from the edge. It is hypothesized that the cells in the swarm interior are not responsible for rapid population and progression of the swarm and thus motility in the form of streams and vortices is not required.

**CONCLUSION**

In this study, we used a non labeled cell tracking algorithm for characterizing the flagellated bacterial swarming motion on a soft agar plate without fluorescence labeling or help of microspheres as tracers. This method calculates average velocities for local regions regularly arranged over the entire flow field composed mostly of curvilinear bacterial stream and vortices.

The tracking method was further applied to study the motility of swarming bacteria as a function of distance from the swarm edge. It was discovered that the average velocities were not highest at the very edge of the swarm but actually reached the maxima in the areas roughly 450 µm from the edge. The maximum for the correlation length was also found to occur roughly 900 µm from the swarm edge. This indicates that the bacterial flows align for the longest distances in this region. The data suggest that the high velocity, surging motions of the swarming cells cause a viscous push to the swarm edge and enable rapid population of the surface.
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