

Barriers Impeding Active Mixing of Swimming Microbes in a Hyperbolic Flow

Helena Yoest ^{1†}, John Buggeln ^{1†}, Minh Doan ^{1†}, Payton Johnson ^{1†}, Simon A. Berman², Kevin A. Mitchell ² and Thomas H. Solomon ^{1*}

¹Department of Physics and Astronomy, Bucknell University, Lewisburg, PA, United States, ²Department of Physics, University of California, Merced, Merced, CA, United States

We present experiments on the motion of swimming microbes in a laminar, hyperbolic flow. We test a theory that predicts the existence of *swimming invariant manifolds* (SwIMs) that act as invisible, one-way barriers that block the motion of the microbes. The flow is generated in a cross-channel in a PDMS cell, driven by syringe pumps. The swimming microbes are euglena and tetraselmis, both single-celled, eukaryotic algae. The algae are not ideal smooth-swimmers: there is significant rocking in their motion with occasional tumbles and a swimming speed that can vary. The experiments show that the swimming algae are bound very effectively by the predicted SwIMs. The different shapes and swimming behavior of the euglena and tetraselmis affect the distribution of swimming angles, with the elongated euglena having a larger probability of swimming in a direction parallel to the outflow directions. The differences in swimming orientation affect the ability of the microbes to penetrate the manifolds that act as barriers to passive tracers. The differing shapes of the euglena and tetraselmis also affect probabilities for the microbes to escape in one direction or the other along the outflow.

Keywords: manifolds, active matter, dynamical systems, mixing, fluid dynamics, microfluidics

1 INTRODUCTION

In the 1980s and 1990s, the tools of dynamical systems and chaos theory were applied to developing an understanding of fluid mixing in laminar flows [1, 2]. In particular, manifolds were identified [3–5] that were found to act as barriers that block the motion and mixing of impurities in a wide range of laminar flows. Furthermore, lobes (or "turnstiles") formed from these manifolds were found to explain long-range transport in flows with simple periodic time dependence [6–8]. These dynamical tools have been extended to identify local barriers and "Lagrangian coherent structures" that guide and organize transport and mixing in spatially- and temporallycomplicated fluid flows [9–14]. These results have led to a range of applications in diverse physical systems such as the spread of pollution in the oceans and atmosphere [12, 15], transport barriers in nuclear fusion plasmas [16], blood transport in cardiovascular flows [17] and morphogenesis in developing embryos [18].

For most of these studies, mixing is passive, characterized by impurities that simply follow a flow without any deviations and without any influence on the flow itself. But mixing in real systems is often active, involving impurities that deviate from the flow and/or impurities that feed back and alter the flow. The behavior of propagating reaction fronts in a fluid flow is an example of active mixing, since the reaction fronts are able to move across a fluid even in the absence of flows. The theory of

OPEN ACCESS

Edited by:

Arlette R. C. Baljon, San Diego State University, United States

Reviewed by:

Eric Climent, UMR5502 Institut de Mecanique des Fluides de Toulouse (IMFT), France Sanjeeva Balasuriya, University of Adelaide, Australia

*Correspondence:

Thomas H. Solomon tsolomon@bucknell.edu

[†]Present Address:

Helena Yoest, Applied Physics Laboratory, Johns Hopkins University, Laurel, MD, United States John Buggeln, Department of Physical Therapy, University of Delaware, Newark, Delaware Minh Doan, FPT University, Hanoi, Vietnam Payton Johnson, Velico Medical, Beverly, MA, United States

Specialty section:

This article was submitted to Soft Matter Physics, a section of the journal Frontiers in Physics

Received: 24 January 2022 Accepted: 22 March 2022 Published: 25 April 2022

Citation:

Yoest H, Buggeln J, Doan M, Johnson P, Berman SA, Mitchell KA and Solomon TH (2022) Barriers Impeding Active Mixing of Swimming Microbes in a Hyperbolic Flow. Front. Phys. 10:861616. doi: 10.3389/fphy.2022.861616

1

manifolds developed for passive mixing has been extended to the problem of front propagation in fluid flows [19, 20]. In this case, burning invariant manifolds (BIMs) permeate the fluid system and act as one-way barriers that both hinder and guide the behavior of fronts in a wide range of fluid flows. We have done numerous experiments that have demonstrated the role of BIMs as one-way barriers in a wide range of two-dimensional [21–23] and three-dimensional laminar fluid flows [24].

Another class of active impurities are those that are selfpropelled, such as biological organisms ranging from bacteria [25–28] to birds and people [29], and artificial swimmers ranging from microscopic Janus particles [30–32] to ships. A modified and extended version of the burning invariant manifold theory [33, 34] predicts similar manifolds—referred to as swimming invariant manifolds (SwIMs)—that act as barriers that inhibit and organize the mixing of self-propelled tracers in a flow. Edges of SwIMs act as one-way barriers when projected into (x, y) position space, similar to the one-way blocking of BIMs of reaction fronts in (x, y). In fact, the burning invariant manifold theory is a special case of the more general swimming invariant manifold approach. We have previously tested parts of this theory experimentally in a hyperbolic flow with rod-shaped, prokaryotic swimming bacteria, some of which were genetically mutated to inhibit tumbling [33].

In this paper, we present the results from additional experiments that test the applicability of SwIM theory to larger, less idealized swimmers with different shapes (both rod-shaped and circular) and different swimming behavior. We choose one of the simplest, non-trivial flows: a laminar, hyperbolic fluid flow in a cross channel. The hyperbolic flow is particularly special since the vorticity is zero everywhere. The self-propelled particles are two different types of eukaryotic swimming algae—freshwater algae (euglena) and marine algae (tetraselmis) that are both an order of magnitude larger in size than the prokaryotic bacteria studied earlier. We examine trajectories of these swimming microbes in the flow to see if they are bounded by the one-way barriers that are predicted by the SwIM theory. We also analyze orientation and escape probabilities in the context of the SwIM theory.

In **Section 2**, we present background about dynamical systems theory of manifolds as applied to the mixing of passive and self-propelled tracers. We discuss the experimental methods used to generate the hyperbolic fluid flow in **Section 3**, along with the techniques used for handling the microbes. Trajectories of the microbes in the flow are presented in **Section 4**, highlighting the SwIMs that act as barriers that impede their motion. We also present a statistical analysis of trajectories through (x, y, θ) phase space and orientation distributions in **Section 4**, interpreted using the SwIM framework. In **Section 5** we discuss these results, as well as continuing work that is needed to further explore the range of applicability of SwIM theory for active mixing.

2 BACKGROUND

2.1 Passive Mixing

The motion of passive tracers in a fluid flow is governed by the differential equations $\dot{x} = u_x$ and $\dot{y} = u_y$, where u_x and u_y are the

x- and *y*-components of the velocity field. For a hyperbolic flow (**Figure 1**) with fluid going inward in the *y*-direction and outward in the *x* direction, the velocity field is described by the equations

$$u_x = Ax, \tag{1a}$$

$$u_y = -Ay. \tag{1b}$$

There is a *passive* fixed point at (x, y) = (0, 0) where u_x and u_y are both zero (Figure 1B). Attached to this fixed point are two passive invariant manifolds. The unstable manifold of the fixed point is defined by starting with an infinite number of passive tracers initially located infinitesimally close to the fixed point and allowed to evolve in time. In the long-time limit, the locations to where those tracers have moved define the unstable manifold which, for a timeindependent, hyperbolic flow is the line y = 0 (horizontal, red dashed line in Figure 1B). Similarly, the stable manifold of the fixed point is defined using the same approach, but integrating the trajectories backward in time. Alternately, the stable manifold is the locus of points that ultimately will end up infinitesimally close to the fixed point. For the hyperbolic flow of Eq. 1a, Eq. 1b, the stable manifold of the fixed point is the line x = 0 (vertical, red dashed line). The passive stable and unstable manifolds act as transport barriers for passive mixing. For the time-independent hyperbolic flow of Figure 1, the stable and unstable manifolds divide the flow into four quadrants, and passive tracers initially in one quadrant remain in that quadrant. For a time-independent flow, the identification of manifolds as transport barriers for passive mixing is a somewhat trivial result since passive tracers do not deviate from the streamlines in the flow. But the nature of manifolds as barriers persists to the less-trivial case of flows with periodic time-dependence [8, 35]. Studies during the past two decades have extended the ideas of manifolds as barriers to aperiodic and even turbulent flows, typical of flows in the oceans and atmospheres [9-14].

2.2 Self-Propelled Tracers

An example of a tracer that is *not* passive is one that is selfpropelled such as a biological organism (e.g., swimming microbes, birds, people), synthetic microscale swimmers such as Janus particles [30–32], and larger-scale devices (e.g., cars, boats, drones, planes). In many cases, the self-propelled tracer is itself subject to flows in the surrounding medium; e.g., fish in a river, bacteria in the lungs or bloodstream, ships on the ocean.

Figure 2 shows an ellipsoidal swimmer. The aspect ratio $\gamma = l_{\parallel}/l_{\perp}$ is the ratio of the long and short dimensions of the ellipsoid, with l_{\parallel} defined parallel to the swimming direction. We assume a two dimensional flow, and assume that the self-propelled tracer swims at a constant speed V_0 relative to the fluid without any noise or internally-imposed change in swimming direction (e.g., tumbling for a microbe). With these assumptions, the motion of the tracer is described by the following set of differential equations [36, 37, 38, 33]:

$$\dot{x} = u_x + V_0 \cos \theta, \qquad (2a)$$

$$\dot{y} = u_y + V_0 \sin \theta, \qquad (2b)$$

$$\theta = (1 + \alpha) (\omega/2) - \alpha (2u_{x,x} \cos \theta \sin \theta) -u_{x,y} \cos^2 \theta + u_{y,x} \sin^2 \theta), \qquad (2c)$$

where (*x*, *y*) are the coordinates of the tracer, θ is the angle of the tracer's swimming relative to the horizontal, and $\omega = u_{y,x} - u_{x,y}$ is





the vorticity of the flow. The parameter $\alpha = (\gamma^2 - 1)/(\gamma^2 + 1)$ denotes the shape and swimming direction of the swimmer (**Figure 2**). **Eq. 2a**, **Eq. 2b**, **Eq. 2c** assume that the motion of the swimmers does not alter the fluid flow.

For the zero-vorticity hyperbolic flow Eq. 1a, Eq. 1b, Eq. 2a, Eq. 2b, Eq. 2c simplify [33]:

$$\dot{x} = Ax + V_0 \cos \theta, \tag{3a}$$

$$\dot{y} = -Ay + V_0 \sin \theta, \tag{3b}$$

$$\theta = -\alpha A \sin\left(2\theta\right) \tag{3c}$$

If the self-propelled tracer swims smoothly with no internallydriven changes in swimming direction or external noise, there is no mechanism for the swimmer to rotate through angles $\theta = 0$, $\pi/2$, π or $3\pi/2$. In fact, swimming orientations of $\theta = 0$ and π are stable attractors for any tracer with positive α , and swimming orientations of $\theta = \pi/2$ and $3\pi/2$ are stable for swimmers with negative α .

We can rewrite **Eq. 3a**, **Eq. 3b**, **Eq. 3c** in non-dimensional form by scaling distances by the distance V_0/A and scaling time by *A*. Dimensionless coordinates are then defined as

$$\tilde{x} = xA/V_0, \tag{4a}$$

$$\tilde{y} = yA/V_0, \tag{4b}$$

$$\tilde{t} = tA.$$
 (4c)

With these definitions, dimensionless versions of Eq. 3a, Eq. 3b, Eq. 3c are:

$$\tilde{\tilde{x}} = \tilde{x} + \cos\theta, \tag{5a}$$

$$\dot{\tilde{y}} = -\tilde{y} + \sin\theta, \tag{5b}$$

$$\dot{\theta} = -\alpha \sin{(2\theta)}.$$
(5c)

There are *swimming fixed points* of Eq. 5 where \hat{x} , \hat{y} and $\hat{\theta}$ are all zero. As is the case for passive mixing (**Section 2.1**), these swimming fixed points have stable and unstable manifolds, referred to as *swimming invariant manifolds* (or *SwIMs*). Like the manifolds for passive mixing, SwIMs act as barriers. In the three-dimensional $(\tilde{x}, \tilde{y}, \theta)$ space, the swimming invariant manifolds form curved sheets, as shown in **Figure 3** [34]. For the hyperbolic flow, the stable manifold of the inward-blocking fixed points (blue sheet in **Figure 3A**) is independent of the *y*-coordinate and therefore projects into (\tilde{x}, θ) space as a 1D curve (**Figure 3B**). This projection separates all left-escaping from all right-escaping trajectories in (\tilde{x}, θ) space. Similarly, the unstable manifold of the outward-blocking fixed points (red sheet in **Figure 3A**) projects into (\tilde{y}, θ) space as a 1D curve (**Figure 3A**) projects into (\tilde{y}, θ) space as a 1D curve (**Figure 3A**) projects into (\tilde{y}, θ) space as a 1D curve (**Figure 3A**) projects into (\tilde{y}, θ) space as a 1D curve (**Figure 3A**) projects into (\tilde{y}, θ) space as a 1D curve (**Figure 3C**) that separates trajectories coming from above and below.

Projections (called "SwIM edges") of the edges of the SwIMs into physical (\tilde{x}, \tilde{y}) space are one-way barriers that block the motion of swimmers in one direction but allow them to pass in the other direction. If swimmers are released at the passive fixed point at the center of the hyperbolic flow (**Figure 1C**), they can swim outward (±*x*-direction) with the flow; i.e., in the unstable direction. But they can also swim up and down (±*y*) against the incoming flow (along the stable direction) until they reach swimming fixed points (colored red in **Figure 1C**) where the incoming flow has a speed V_0 , equal and opposite the swimmers. Thus, the red swimming fixed points at $(\tilde{x}, \tilde{y}) = (0, \pm 1)$ and the red SwIM edges at $\tilde{y} = \pm 1$ in **Figure 1C** block any outgoing $\pm y$ swimmers.









However, if a swimmer moves *inward* along the *y*-axis, the red SwIM edge along its path no longer blocks the swimmer, since the direction of the flow and the swimming are the same. The red SwIM edges are one-way barriers, blocking outward—but not inward—swimming tracers. Similarly, the blue swimming fixed points at $(\tilde{x}, \tilde{y}) = (\pm 1, 0)$ and SwIM edges at $\tilde{x} \pm 1$ in **Figure 1C** are one-way barriers, but these block *inward-swimming* (and not outward-swimming) tracers.

One-dimensional swimming invariant manifolds for $\alpha = -1$ (which coincide with BIMs that block reaction fronts in the same flow) have been shown theoretically [34] to block *all* swimmers, even those that tumble or have rotational diffusion or added rotational noise, as long as their swimming speed remains constant. The location of the swimming fixed points and

SwIM edges are independent of α for the hyperbolic flow and coincide with the BIMs. Consequently, the SwIM edges in the hyperbolic flow are predicted theoretically to be barriers even for tumbling swimmers.

3 EXPERIMENTAL METHODS

3.1 Microbes

The microbes (**Figure 4**) used in these studies are two types of motile algae—euglena and tetraselmis. Euglena gracilis (**Figure 4A**) are freshwater algae that measure approximately 40–50 μ m in length with a width of around 8–12 μ m with aspect ratios $\gamma \sim 3-5$ and $\alpha \sim 0.8$ –0.9. (The euglena are not rigid and

occasionally undulate in shape.) The euglena are propelled by a trailing flagellum that push the body forward. Tetraselmis (**Figure 4D**) are green marine algae that are almost circular with a typical diameter $10-15 \mu m$; we estimate α to be approximately 0.3–0.4. Tetraselmis are *pullers* with four flagella at the front that beat in a breast-stroke pattern to pull the body forward.

The estimates of α for both euglena and tetraselmis are crude. First, neither of them are perfect ellipsoids. Second, the shape varies from one organism to the next, especially for the euglena which are not even rigid. Furthermore, we neglect any effects of the extended flagella on α . But the difference in α is quite significant between the two organisms, with euglena being a reasonable approximation of an organism with $\alpha \approx 1$ and tetraselmis being a reasonable approximation of an organism with $\alpha \approx 0$.

The euglena and tetraselmis used in these experiments are standard, commercially-available samples obtained from Carolina Biological Supply without any effort at purifying the strain. The euglena are stored in the shipping tubes with continuous fluorescent illumination. Even a few months after shipment, euglena stored in this manner still swim actively. For use in an experimental run, the euglena samples from these tubes are diluted in soil-water medium (Carolina Biological Supply catalog #153785) at a concentration of 15–20% euglena sample by volume. For the tetraselmis, portions of the sample obtained from Carolina are cultured in Alga-Gro[®] Seawater medium (catalog #FAM_153754) with stirring and fluorescent illumination for 2 weeks, then transferred to 12 ml centrifuge tubes. Samples from these tubes are then diluted 1:1 with fresh Alga-Gro Seawater immediately before each experimental run.

3.2 PDMS Cells, Flow Generation and Measurement

The experiments are conducted in cross-channel cells (**Figure 1**) made from polydimethylsiloxane (PDMS); the channels are 4.0 mm wide and 2.0 mm deep. The flow is produced with two syringe pumps, one with two syringes attached via tubing to inlets/outlets at the top and bottom of the cross, and a second syringe pump with two syringes connected to inlets/outlets at the left and right ends of the cross. For a particular run, one syringe pump infuses fluid from two 1-ml syringes into the cell while the other syringe pump withdraws fluid at the same flow rate. The cell is mounted on an inverted microscope and imaged with a 4X objective. With this magnification, the visible region at the intersection of the two channels in the cross has a width and height of 3 mm.

For velocimetry measurements, $5.0 \,\mu\text{m}$ dyed (blue) polystyrene microspheres are used as passive tracers. In all experiments (both with passive and with swimming particles), the tracers are imaged only in a region near the mid-height of the cell to avoid slow-down in the flow due to the no-slip boundary conditions at the top and bottom.

Images are acquired using a CCD video camera. A background image determined by averaging frames over several seconds is subtracted from each image, and the resulting image is thresholded. These thresholded images are analyzed in IDL with a package developed by John Crocker and Eric Weeks [39] that determines centroid coordinates for clusters of pixels (with each cluster identified as a tracer), and then links coordinates in successive frames to determine trajectories. For the passive tracers, these trajectories are then used to determine local velocities. With a sufficient density, a velocity field is constructed, as shown in Figure 5A. (The region of interest is not centered on the flow, but that has no bearing on the analysis.) Linear regressions of slices of this velocity (Figure 5B) are used to determine the parameter A that denotes the strength of the velocity field of Eq. 1a, Eq. 1b. Deviations of the experimental velocity field from a true hyperbolic flow are shown in Figure 5C. Not surprisingly, the linear growth in the velocity is limited to a finite distance from the fixed point, since the fluid velocity has a finite peak at the inlet and outlet channels. This shows up in Figure 5C by the difference vectors which are opposite the flow far from the fixed point.

For runs with swimming microbes, we fit x – and y – components of the trajectories separately to sliding parabolas and take derivatives to determine the x – and y – components of the total velocity of the trajectory $\vec{V}_{tot} = \vec{V}_{swim} + \vec{u}_{flow}$. Subtracting the measured fluid velocity \vec{u}_{flow} Eq. 1a, Eq. 1b from \vec{V}_{tot} gives us the swimming velocity \vec{V}_{swim} from which we can determine both the swimming speed and direction $\theta = \arctan(V_{swim,v}/V_{swim,x})$ for each individual microbe. The average swimming speed $\langle V_0 \rangle$ is determined for each trajectory. We classify each trajectory as "tumbling" or "non-tumbling" by comparing the swimming orientation $\theta(t)$ at different times along the trajectory, as shown in **Figure 6**. The angles $\theta(t)$ are "unwrapped" so as not to be confined to the range 0 to 2π . For example, if a microbe is swimming with an orientation $\theta = 6.1$ radians and then rotates another +0.4 radians, the orientation θ = 6.5 radians with no wrap-around past zero. A microbe is deemed to "tumble" if $\Delta \theta \ge \pi$ between any two points in the trajectory, e.g., red and green curves in Figure 6. This is somewhat arbitrary-ideally, no smooth-swimming microbe should ever rotate beyond an angle of $\pi/2$ in a hyperbolic flow; however, periodic rocking motion of the swimming algae microbes (visible in the plots of $\theta(t)$ in Figure 6 and discussed in Section 4.1 below) can produce deviations in θ by as much as a radian, so a threshold $\Delta \theta$ of $\pi/2$ would result in a significant number of trajectories being labelled incorrectly as tumblers. The trajectories labelled non-tumblers (e.g., black curve in Figure 6) by this approach are not perfect smooth swimmers, partially because of the rocking motion of the swimming and also because of noise due to rotational diffusion (which is likely minimal considering the size of the microbes used in these studies) and other irregularities in the swimming motion.

A series of experimental runs for swimming microbes is obtained using the following protocol. After loading the tubing and PDMS cell with microbe-laden fluid, 1 ml syringes are loaded with new fluid and then data is taken with the flow in the "forward" direction (fluid pumped in along the vertical channel and out along the sides). After 1.0 ml of fluid is pumped (the capacity of the syringe), the flow is stopped. Data is then taken for a minute or two of the microbes



FIGURE 5 (A) Experimental velocity field, determined with passive tracers; flow rate 200 μ/min , A = 3.6 s⁻¹. The red circle shows the region of interest used for all of the statistical analyses of microbe data. (B) Slices of the velocity field; u_x along a horizontal line through the fixed point (blue), and u_y along a vertical line through the fixed point (red). These curves are fit to a line to determine A in Eq. 1a, Eq. 1b. (C) Vector field showing the deviation of the experimentally-measured velocity field from an ideal hyperbolic velocity field Eq. 1a, Eq. 1b; i.e., $\Delta \vec{v} = \vec{v}_{exp} - \vec{v}_{ideal}$. The arrows are scaled up by a factor of five larger than those for Figure 5A. The color map shows the magnitude $|\Delta \vec{v}|/\langle v \rangle$ of the velocity deviations, normalized by the average flow speed.



swimming without a flow. The flow is then reversed and data is taken with the fluid pumped in along the horizontal and out along the vertical. Another no-flow run is taken at the end of the "reverse" run, after which another forward run is obtained. This process of forward/no-flow/reverse/no-flow/forward/no-flow ... continues for 2–3 h for each set of data or until the microbe swimming is diminished.

4 RESULTS

4.1 No Flow

The euglena used in these studies swim fairly smoothly with only occasional tumbling. Even in the absence of tumbling, though, there is still variation in the swimming direction due to rocking back and forth in the swimming (a two-dimensional projection of helical swimming) and noise in the swimming behavior. **Figure 4B** shows a streak image of euglena swimming in the cross-channel cell with no imposed flow. (Movies of this are available online in **Supplementary Material**) Close inspection of some of the streaks in **Figure 4B** shows periodic wiggling due to the rocking motion. Virtually every euglena remains motile throughout the course of a sequence of experiments, which continue for 2–3 h.

The swimming behavior of the tetraselmis is more varied. Not all of the tetraselmis in a sample are motile; during any run, some of the tetraselmis remain sessile with the fraction of sessile microbes increasing over the course of a sequence of experimental runs. A streak image of tetraselmis in the channel with no flow is shown in **Figure 4E**. Similar to the euglena, some of the tetraselmis tumble noticeably and others swim more smoothly, although there is clear rocking (wiggling of some of the trajectories in **Figure 4E**) and noise in the orientations even for the tetraselmis that are not undergoing classic run-and-tumble motion.

Probability distributions of average swimming speeds in the absence of a flow are shown for euglena and tetraselmis in **Figures 4C,F**, respectively. (For the tetraselmis data in **Figure 4F**, the sessile microbes are excluded.). The mean swimming speed (with no flow) is 90 and 120 μ m/s for the euglena and tetraselmis, respectively. (The imposed flows in these experiments have no significant measurable effect on these mean swimming speeds.) Unlike the bacteria in the previous study [33], the swimming speed for an individual microbe is not necessarily constant in time. In fact, when tetraselmis tumble, they slow down to a stop before changing orientation.

4.2 Euglena in Hyperbolic Flow

A streak image of euglena moving in the hyperbolic flow is shown in **Figure 7A**. (A movie of the euglena swimming in the same flow can be found in online **Supplementary Material**) The effects of the swimming on the motion are clear in this figure, signified by



frequent crossing of the trajectories. As can be seen in the movie and the plot of the orientation for a typical non-tumbling microbe (black curve in **Figure 6**), there is a strong preference for the euglena to be oriented horizontally ($\theta = 0$ or π radians), particularly near the unstable passive manifold of the flow's fixed point. This is consistent with **Eq. 3c**, which indicates that $\dot{\theta}$ should cause all swimmers with $\alpha > 0$ in a hyperbolic flow to orient toward one of these two angles.

Figure 6 shows the swimming orientation θ for three different types of trajectories observed in these experiments. Non-tumbling ("smooth") trajectories often look like the black curve in **Figure 6**. There are two ways in which a tracer can show tumbling behavior. In some cases, similar to the red curve in **Figure 6**, the swimming is mostly smooth, with occasional and well-defined tumbling events where the orientation θ quickly changes. In many cases, though, the tumbling is more continuous, similar to the green curve in **Figure 6**. For the rest of the data analysis, we consider only the trajectories that are classified as non-tumbling.

Figure 7B shows a small sampling of trajectories determined by tracking the euglena; this figure shows the longest-duration, tracked, non-tumbling trajectories that initially enter from the bottom. It is difficult to interpret this plot vis-a-vis SwIM theory; since different euglena microbes swim with different average speed $\langle V_0 \rangle$, the location of the swimming fixed points and manifolds differ between each of the trajectories. To compare the trajectories with the swimming invariant manifold theory, we need to take into account the fact that each microbe has a different swimming speed. We use the dimensionless coordinates \tilde{x} and \tilde{y} defined in Eq. 4, approximating the swimming speed V_0 with the average speed $\langle V_0 \rangle$ over a trajectory. When plotted using these dimensionless variables (Figure 7C), we can see that the trajectories from Figure 7B all pass very close to the predicted inward-blocking swimming fixed points at $(\tilde{x}, \tilde{y}) = (\pm 1, 0)$. This is, in fact, the reason why these are the longest-duration trajectories. The nature of these swimming fixed points can be seen clearly in Figure 7C, as all of the trajectories come in vertically along the $\tilde{x} = \pm 1$ line, then make a sharp turn and move off either left or right horizontally from the swimming fixed points.

The blocking behavior of the SwIMs can be seen by plotting the trajectories in groupings sorted by the directions from which they enter and to which they leave the hyperbolic flow region. **Figure 8A** shows all of the microbes that enter from the top and depart the hyperbolic flow to the left. Virtually none of these trajectories appear at any point to the right of the inwardblocking SwIM edge at $\tilde{x} = 1$. Similar behavior can be seen in **Figures 8B–D**. Virtually none of the trajectories that leave to the left ever have coordinate $\tilde{x} > 1$ (i.e., to the right of the left-blocking SwIM edge at $\tilde{x} = 1$), and virtually none of the trajectories that leave to the right ever have \tilde{x} to the left of the right-blocking SwIM edge at $\tilde{x} = -1$.

In earlier studies with bacteria [33] (and with more than an order of magnitude fewer trajectories), the barriers due to the SwIM edges at $\tilde{x} = \pm 1$ were never violated. For the euglena, these barriers are not perfect; there are a small number of crossings visible in Figure 8. The blocking behavior can be quantified by considering the probability p_{left} (or p_{right}) for a microbe to escape to the left (right), given its initial starting location. In calculating these probabilities, we truncate all of the raw trajectories, excluding any motion outside a circle drawn around the flow's fixed point with the largest radius that keeps all of the circle within the microscope's field of view (red circle in Figures 5A,C). To account for any deviations in the left-right orientation symmetry in the sample of trajectories, we weight the *left-going* probability p_{lg} for a microbe eventually to move left (i.e., $\tilde{x}(t_{end}) < \tilde{x}(0)$) by the *left-facing* probabilities *p*_{*lf*} for tracers initially to be *facing* with their swimming direction to the left (i.e., with $\theta(0)$ between $\pi/$ 2 and $3\pi/2$):

$$p_{left} = \frac{\frac{p_{lg}}{p_{lf}}}{\frac{p_{lg}}{p_{lf}} + \frac{p_{rg}}{p_{rf}}}$$







For instance, since right-facing microbes move more slowly near the $(\tilde{x}, \tilde{y}) = (-1, 0)$ swimming fixed point, right-facing microbes can be oversampled in that region. Defined as above, p_{left} is the probability that a swimmer will end up moving left, assuming an equal number of left- and right-facing microbes at all points in the flow.

Plots of p_{left} for euglena are shown in **Figure 9A**. Theoretically, any microbe with $\tilde{x} < -1$ should have $p_{left} = 1$ since no microbe can cross over the right-blocking SwIM edge at $\tilde{x} = -1$. Similarly, any microbe with $\tilde{x} > 1$ should have $p_{left} = 0$. The slight crossing of the tails of this distribution beyond $\tilde{x} = \pm 1$ is indicative of the slight leakage in the SwIM edges seen in **Figure 8**.

The slight leakage of the SwIMs as one-way barriers can be understood by considering the swimming speeds. Unlike the bacteria studied previously [33], the euglena do not always swim at a constant speed. The speed $\langle V_0 \rangle$ used to nondimensionalize the coordinates is an average of the swimming speed during a trajectory. If the microbe is momentarily swimming with a speed larger than $\langle V_0 \rangle$, it can cross over what would be the barrier for a microbe swimming at the lower average speed.

The functional form of $p_{left}(\tilde{x})$ between $\tilde{x} = -1$ and 1 depends on several factors [34]. For perfectly smoothswimming microbes with $\alpha = 1$, inspection of the relative areas of left- and right-going regions in Figure 3B indicates that if microbe orientations θ are left-right symmetric, roughly half would escape to the left for most values of \tilde{x} between -0.5 and 0.5 transitioning up to 1 approaching $\tilde{x} = -1$ and down to 0 approaching $\tilde{x} = +1$. Qualitatively, this is consistent with the shape of the curve in Figure 9A. As discussed in Ref. [34], the presence of tumbling and rotational noise in the swimming modifies this curve, with the transition of p_{left} from 1 down to 0 approaching and even passing a straight diagonal line (dashed line in Figure 9A) with more tumbling and rotational noise. The lack of an appreciable change in the experimental curve for a variation in A by a factor of 4 indicates that these data are close to the limit of low noise. This is further indicated by the good agreement between the experimental data and the theoretical prediction (in green, from Ref. [34]) for the case with no noise and no tumbling. (Note, however, that the prediction assumes a uniform initial distribution of angles, whereas the calculated p_{left} is weighted only for left-right symmetry in the initial angles.)

8



Looking again at **Figure 8**, there is very little penetration of the euglena past the horizontal passive manifold at $\tilde{y} = 0$. This is consistent with the observation that most of the euglena are oriented horizontally ($\theta = 0$ or π) as they are moving in the vicinity of the unstable passive manifold of the hyperbolic flow. Without a significant vertical component in their swimming orientation, there is no mechanism for them to swim past the passive manifold. The microbes that *do* pass $\tilde{y} = 0$ are likely those that tumbled before the beginning of the tracked trajectory.

We make a 3D histogram in $(\tilde{x}, \tilde{y}, \theta)$ space from all of the "non-tumbling" trajectories that are also confined to the circular region around the flow's fixed point (see **Figure 5A**). Projections of this histogram for the euglena are shown in **Figures 10A–C**. A few features are notable in the (\tilde{x}, θ) and (\tilde{y}, θ) projections. First, there is a clear preference for horizontal orientations, shown by the dark bands at $\theta = 0$ and π in **Figures 10B,C**. Second, there is a low-density band in the region between $\tilde{x} = -1$ and 1 in the (\tilde{x}, θ) projection (**Figure 10B**) that matches up well with the projection of the SwIM (dashed blue) from **Figure 3B**. This is potentially a manifestation of the manifold as being repelling in (\tilde{x}, θ) space. As shown in Ref. [33], this SwIM projection also very effectively divides the left- and right-going trajectories.

There are significant density variations in the (\tilde{y}, θ) projections (**Figure 10C**). The euglena enter the field of view (away from \tilde{y} near 0) with a broad angle distribution, but near the centerline $\tilde{y} = 0$, the distribution is peaked heavily at $\theta = 0$ and π , as expected from the earlier discussion about the preference of horizontal orientations for the almost rod-shaped euglena with α near 1. It is also intriguing that there is a noticeable edge in the distribution that is close to—but not identical to—the projection of the SwIM into (\tilde{y}, θ) space. A

slice of the (\tilde{y}, θ) projection at $\tilde{y} = 0$ can be normalized to obtain a probability function for the microbe orientations, as shown in **Figure 11A**. As was the case for the left- and right-going probabilities (**Figure 9A**), these orientation distributions are roughly the same for the different flow rates studied in the experiments.

The distributions in Figure 11A cannot be compared quantitatively with the theoretical predictions for orientation distributions in Ref. [34]. The theoretical predictions assume a flow that is hyperbolic over an infinite extent, which is clearly not the case for the experiments. (In fact, there are even deviations within the field of view of the camera, as seen in Figure 5C.) In the experiments, the fluid and microbes are pumped through long tubes and then through channels leading to the intersection of the cross, and only in this intersection is the flow well approximated as hyperbolic. Furthermore, in the theoretical predictions, the distribution is calculated over all values of \tilde{v} , not just for $\tilde{v} = 0$. Qualitatively, though, the strong peaks in the experimental distribution at $\theta = 0$ and π are consistent with similar features in the theoretical predictions for swimmers with α near 1. The theory predicts that these peaks broaden with additional noise or for weaker flows (smaller A). The peaks for the lower flow case (blue symbols) in Figure 11A are slightly broader than those for the higher flow case (red symbols). But the differences are minimal, a further indication that the euglena data are in or near the low-noise limit.

Slices of the $(\tilde{x}, \tilde{y}, \theta)$ histograms at different orientational angles θ (**Figures 10D–G**) also reveal features predicted by the theory. The hyperbolic structure of the trajectories at $\theta = 0$ (**Figure 10D**) and $\theta = \pi$ (**Figure 10F**) show the inward-blocking swimming fixed points q^{in} and their 1D manifolds.





Slices at $\theta = \pi/2$ (Figure 10E) and $\theta = 3\pi/2$ (Figure 10G) show faint ghosts of the outward-blocking swimming fixed points, although these are difficult to see because there simply are not many tracers that reach the vicinity of these swimming fixed points.

4.3 Tetraselmis in Hyperbolic Flow

Trajectories of tetraselmis swimming in a hyperbolic flow are shown in **Figure 12**. These plots do not include the sessile microbes (determined by a threshold in measured $\langle V_0 \rangle$); we also exclude trajectories that are undergoing classic run-and-tumble behavior (determined again by examining differences in orientation $\Delta \theta$). Once again, we plot the longest-duration trajectories in **Figure 12B,C**

There is a notable qualitative difference between these plots and those for the euglena (**Figures 7B,C**). Whereas the longestduration euglena trajectories are those that pass closest to the inward-blocking swimming fixed points at $(\tilde{x}, \tilde{y}) = (\pm 1, 0)$, the longest-duration trajectories for the tetraselmis that enter from the bottom are those that display hyperbolic behavior around the outward-blocking swimming fixed point at $(\tilde{x}, \tilde{y}) = (0, 1)$. Furthermore, whereas the smoothest-swimming euglena rarely penetrate the passive manifold at $\tilde{y} = 0$, it is not uncommon for non-tumbling tetraselmis to pass $\tilde{y} = 0$ and approach the outward-blocking SwIM edge at $\tilde{y} = 1$.

The ability of tetraselmis to penetrate the passive manifold at $\tilde{y} = 0$ can be seen by plotting all of the trajectories in four quadrants (**Figure 13**). In contrast to the euglena (**Figure 8**), the tetraselmis often swim vertically near—but rarely beyond—the outward-blocking SwIMs at $(\tilde{x}, \tilde{y}) = (0, \pm 1)$. Similar to the euglena, the inward-blocking SwIM edges mostly define the range of the left-going and right-going microbes. This is supported by a plot of the fraction $p_{left}(\tilde{x})$ of left-escaping tetraselmis (**Figure 9B**), which is 1 for $\tilde{x} < -1$ and 0 for $\tilde{x} > 1$ with only slight leakage. This leakage is again due to variations in swimming speed around the average $\langle V_0 \rangle$.

There are two main reasons for the differences between the behavior of the euglena and tetraselmis. First, since tetraselmis are more circular than euglena, there is less of a tendency for them to orient horizontally in the hyperbolic flow. Unlike with the euglena, the orientations are not readily visible in the microscopic images. We determine the orientations of the tetraselmis using the same technique used with the euglena, subtracting the flow velocity from the total trajectory



FIGURE 13 |: Trajectories of tetraselmis in hyperbolic flow with $A = 1.8 \text{ s}^{-1}$. (These data exclude trajectories with classic run-and-tumble events during a trajectory.) The blue and red lines show the inward- and outward-blocking SwIM edges, respectively. (A) All trajectories that enter from the top and leave to the left. (B) All trajectories that enter from the bottom and leave to the left. (C) All trajectories that enter from the bottom and leave to the right.



velocities. The (\tilde{x}, θ) and (\tilde{y}, θ) distributions are shown in **Figures 14A–D** for two different flows. For weaker flows, (**Figures 14A,B**), there is a broad distribution in angles, with a slight preference for vertical orientation, likely due to alignment of the tracers in the inflow tubing and channels. This broader angle distribution—especially for weaker flows—can be seen by plotting the probability distribution for angles at $\tilde{y} = 0$ (**Figure 11B**).

With swimming velocities that have non-zero \tilde{y} -components, there is significantly less tendency for the tetraselmis to explore the inward-blocking fixed points and a stronger tendency for them to cross the passive manifold at $\tilde{y} = 0$ and explore the outward-blocking fixed points. This can be seen in slices of the $(\tilde{x}, \tilde{y}, \theta)$ histograms (**Figures 14E–H**) for which the outward-blocking fixed points are much more clearly sampled (**Figures 14F,H**) than for the euglena (**Figures 10E,G**).

The swimming of the tetraselmis is also noisier than that of the euglena, even in the absence of classic run-and-tumble events. This can be seen qualitatively in the no-flow streak plots (**Figure 4**). The noisier motion results in a distribution of orientations (**Figure 11B**) that is much broader than that for euglena (**Figure 11A**). Furthermore, the distribution is significantly more broad for weaker flows, as seen by comparing the blue and red symbols in **Figure 11B**, qualitatively consistent with the theoretical predictions about the effects of noise on the angle distributions. Noise in the swimming orientations is another mechanism that causes non-zero *y*-components in the swimming that enable tracers to pass the passive manifold at $\tilde{y} = 0$ and explore the outward-blocking SwIM edges.

Plots of escape probability $p_{left}(\tilde{x})$ for tetraselmis differ significantly from those for euglena (compare **Figures 9A,B**). We plot the theoretical curves for smooth-swimmers with both $\alpha = 0$ (light green curve) and $\alpha = 0.3$ (dark green) which corresponds to the experimental estimate of α for tetraselmis. The experimental escape probability is strikingly similar to the theoretical predication for an ideal, circular ($\alpha = 0$), smooth-swimming microbe, although it is possible that orientational noise in the swimming of the tetraselmis is playing a role in the shape of the plots in **Figure 9B**.

5 DISCUSSION

Overall, the experimental results are mostly consistent with predictions [33, 34] of swimming invariant manifolds as oneway barriers that impede the motion of self-propelled tracers in a hyperbolic fluid flow. In fact, the theory is quite robust. The swimming of both euglena and tetraselmis is not perfectly smooth and not even necessarily constant in time. Furthermore, the euglena are not even rigid—their shapes can undulate. Nevertheless, the SwIM theory still effectively predicts the one-way barriers that limit the motion of these microbes.

An important question is how to extend and apply SwIM theory to more realistic swimmers and more realistic flows. On the first point, it may be possible to modify the theory to account for swimming with a varying swimming speed, either by using perturbation methods for small variations in swimming speed or by explicitly adding terms to the \dot{x} and \dot{y} equations to account for fluctuations in swimming speed. It would also be interesting to study the range of validity of the SwIM approach to larger organisms (e.g., fish, birds), including those with the ability to make decisions and change their swimming/flying behavior in response to various stimuli. On the second point, further experiments are needed to test the SwIM theory in different laminar flows and to explore the potential applicability of the theory to time-periodic and time aperiodic fluid flows. Presumably, theoretical methods similar to Lagrangian Coherent Structure (LCS) approaches used for passive transport can be adapted to the more general active mixing problem.

It would also be of interest to do experiments with selfpropelled particles with negative α . Swimmers like this do not typically exist in nature, although it should be possible to fabricate synthetic swimmers such as Janus particles [31] that swim in a direction perpendicular to their semi-major axis. Of particular interest is the case where $\alpha = -1$ where the system of equations **Eq. 2a**, **Eq. 2b**, **Eq. 2c** are the same as those for front propagation.

We are also interested in the importance of SwIMs on the simultaneous mixing of both passive and active tracers in the same flow [40], and the effects of chemotaxis on the nature of SwIMs as barriers for active mixing, especially in cases where a flow is mixing both the self-propelled tracers and also the nutrients toward which they exhibit chemotaxis. The effects of scattering of swimmers off physical boundaries [41] on the SwIMs is another important issue. Another problem of interest is the case where the behavior of the active tracers *does* alter the flow field (e.g., bacterial turbulence [42–44]), resulting in a self-consistent process whereby the swimmers generate a flow

which then imposes barriers due to swimming invariant manifolds.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusion of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

HY, JB, MD, and PJ developed the techniques that were used in collecting the data in these experiments. SB and KM provided frequent feedback on the analysis and comparison with theory; they also provided data for theoretical curves. TS ran the experiments, did most of the analysis and wrote the first draft of the manuscript. All authors contributed to the revision of the manuscript.

FUNDING

These experiments were supported by the National Science Foundation under grants DMR-1806355 and CMMI-1825379.

ACKNOWLEDGMENTS

The authors would like to thank Jack Raup for frequent assistance with the PDMS cells, Matthew Heintzelman for guidance on working with swimming microbes, and Brandon Vogel for initial assistance in working with PDMS.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fphy.2022.861616/full#supplementary-material

Supplementary Video S1 | Movie corresponding to Figure 4B; motion of euglena without an imposed flow, shown with streaks.

Supplementary Video S2 | Movie corresponding to Figure 4B; motion of euglena without an imposed flow.

Supplementary Video S3 | Movie corresponding to Figure 4E; motion of tetraselmis without an imposed flow.

Supplementary Video S4 | Movie corresponding to Figure 4E; motion of tetraselmis without an imposed flow, shown with streaks.

Supplementary Video S5 | Movie corresponding to Figure 7A; motion of euglena with an imposed hyperbolic flow.

Supplementary Video S6 | Movie corresponding to Figure 7A; motion of euglena with an imposed hyperbolic flow, shown with streaks.

Supplementary Video S7 | Movie corresponding to Figure 12A; motion of tetraselmis with an imposed hyperbolic flow.

Supplementary Video S8 | Movie corresponding to Figure 12A; motion of tetraselmis with an imposed hyperbolic flow, shown with streaks.

REFERENCES

- Aref H. Stirring by Chaotic Advection. J Fluid Mech (1984) 143:1–21. doi:10. 1017/s0022112084001233
- 2. Ottino JM. The Kinematics of Mixing: Stretching, Chaos and Transport. Cambridge: Cambridge University Press (1989).
- MacKay RS, Meiss JD, Percival IC. Transport in Hamiltonian Systems. *Physica D: Nonlinear Phenomena* (1984) 13:55–81. doi:10.1016/0167-2789(84)90270-7
- Rom-Kedar V, Wiggins S. Transport in Two-Dimensional Maps. Arch Rational Mech Anal (1990) 109:239–98. doi:10.1007/bf00375090
- 5. Wiggins S. Chaotic Transport in Dynamical Systems. New York: Springer-Verlag (1992).
- Camassa R, Wiggins S. Chaotic Advection in a Rayleigh-Bénard Flow. *Phys Rev* A (1991) 43:774–97. doi:10.1103/physreva.43.774
- Solomon TH, Gollub JP. Chaotic Particle Transport in Time-dependent Rayleigh-Bénard Convection. *Phys Rev A* (1988) 38:6280–6. doi:10.1103/ physreva.38.6280
- Solomon TH, Tomas S, Warner JL. Role of Lobes in Chaotic Mixing of Miscible and Immiscible Impurities. *Phys Rev Lett* (1996) 77:2682–5. doi:10. 1103/physrevlett.77.2682
- Haller G. Lagrangian Structures and the Rate of Strain in a Partition of Two-Dimensional Turbulence. *Phys Fluids* (2001) 13:3365–85. doi:10.1063/1. 1403336
- Voth GA, Haller G, Gollub JP. Experimental Measurements of Stretching fields in Fluid Mixing. *Phys Rev Lett* (2002) 88:254501. doi:10.1103/physrevlett.88.254501
- Mathur M, Haller G, Peacock T, Ruppert-Felsot JE, Swinney HL. Uncovering the Lagrangian Skeleton of Turbulence. *Phys Rev Lett* (2007) 98:144502. doi:10. 1103/physrevlett.98.144502
- Mezić I, Loire S, Fonoberov VA, Hogan P. A New Mixing Diagnostic and Gulf Oil Spill Movement. Science (2010) 330:486–9. doi:10.1126/science.1194607
- Budisic M, Mezic I. Geometry of the Ergodic Quotient Reveals Coherent Structures in Flows. *Physica D* (2012) 241:1255
- Haller G. Lagrangian Coherent Structures. Annu Rev Fluid Mech (2015) 47: 137–62. doi:10.1146/annurev-fluid-010313-141322
- Coulliette C, Lekien F, Paduan JD, Haller G, Marsden JE. Optimal Pollution Mitigation in Monterey bay Based on Coastal Radar Data and Nonlinear Dynamics. *Environ Sci Technol* (2007) 41:6562–72. doi:10.1021/es0630691
- Di Giannatale G, Bonfiglio D, Cappello S, Chacón L, Veranda M. Prediction of Temperature Barriers in Weakly Collisional Plasmas by a Lagrangian Coherent Structures Computational Tool. *Nucl Fusion* (2021) 61:076013. doi:10.1088/ 1741-4326/abfcdf
- Darwish A, Norouzi S, Labbio GD, Kadem L. Extracting Lagrangian Coherent Structures in Cardiovascular Flows Using Lagrangian Descriptors. *Phys Fluids* (2021) 33:11707. doi:10.1063/5.0064023
- Serra M, Streichan S, Chuai M, Weijer CJ, Mahadevan L. Dynamic Morphoskeletons in Development. Proc Natl Acad Sci U.S.A (2020) 117: 11444–9. doi:10.1073/pnas.1908803117
- Mahoney J, Bargteil D, Kingsbury M, Mitchell K, Solomon T. Invariant Barriers to Reactive Front Propagation in Fluid Flows. *Epl* (2012) 98:44005. doi:10.1209/0295-5075/98/44005
- Mahoney JR, Mitchell KA. Finite-time Barriers to Front Propagation in Two-Dimensional Fluid Flows. Chaos (2015) 25:087404. doi:10.1063/1.4922026
- Bargteil D, Solomon T. Barriers to Front Propagation in Ordered and Disordered Vortex Flows. Chaos (2012) 22:037103. doi:10.1063/1.4746764
- Megson PW, Najarian ML, Lilienthal KE, Solomon TH. Pinning of Reaction Fronts by Burning Invariant Manifolds in Extended Flows. *Phys Fluids* (2015) 27:023601. doi:10.1063/1.4913380
- Gowen S, Solomon T. Experimental Studies of Coherent Structures in an Advection-Reaction-Diffusion System. *Chaos* (2015) 25:087403. doi:10.1063/ 1.4918594
- Doan M, Simons JJ, Lilienthal K, Solomon T, Mitchell KA. Barriers to Front Propagation in Laminar, Three-Dimensional Fluid Flows. *Phys Rev E* (2018) 97:033111. doi:10.1103/PhysRevE.97.033111
- Berg HC, Anderson RA. Bacteria Swim by Rotating Their Flagellar Filaments. Nature (1973) 245:380–2. doi:10.1038/245380a0

- Stocker R, Durham WM. Tumbling for Stealth? Science (2009) 325:400–2. doi:10.1126/science.1177269
- 27. Rusconi R, Guasto JS, Stocker R. Bacterial Transport Suppressed by Fluid Shear. Nat Phys (2014) 10:212-7. doi:10.1038/nphys2883
- Dehkharghani A, Waisbord N, Dunkel J, Guasto JS. Bacterial Scattering in Microfluidic crystal Flows Reveals Giant Active taylor-aris Dispersion. Proc Natl Acad Sci U.S.A (2019) 116:11119–24. doi:10.1073/pnas.1819613116
- Brockmann D, Hufnagel L, Geisel T. The Scaling Laws of Human Travel. Nature (2006) 439:462–5. doi:10.1038/nature04292
- Ebbens SJ, Howse JR. In Pursuit of Propulsion at the Nanoscale. Soft Matter (2010) 6:726. doi:10.1039/b918598d
- Das S, Garg A, Campbell AI, Howse J, Sen A, Velegol D, et al. Boundaries Can Steer Active Janus Spheres. Nat Commun (2015) 6:8999. doi:10.1038/ncomms9999
- Katuri J, Uspal WE, Simmchen J, Miguel-López A, Sánchez S. Cross-stream Migration of Active Particles. *Sci Adv* (2018) 4:eaao1755. doi:10.1126/sciadv. aao1755
- Berman SA, Buggeln J, Brantley DA, Mitchell KA, Solomon TH. Transport Barriers to Self-Propelled Particles in Fluid Flows. *Phys Rev Fluids* (2021) 6: L012501. doi:10.1103/physrevfluids.6.1012501
- Berman SA, Mitchell KA. Swimming Dynamics in Externally Driven Flows: The Role of Noise. *Phys Rev Fluids* (2022) 7:014501. doi:10.1103/ PhysRevFluids.7.014501
- Rom-Kedar V. Homoclinic Tangles-Classification and Applications. Nonlinearity (1994) 7:441–73. doi:10.1088/0951-7715/7/2/008
- Torney C, Neufeld Z. Transport and Aggregation of Self-Propelled Particles in Fluid Flows. *Phys Rev Lett* (2007) 99:078101. doi:10.1103/PhysRevLett.99.078101
- Khurana N, Blawzdziewicz J, Ouellette NT. Reduced Transport of Swimming Particles in Chaotic Flow Due to Hydrodynamic Trapping. *Phys Rev Lett* (2011) 106:198104. doi:10.1103/physrevlett.106.198104
- Zöttl A, Stark H. Nonlinear Dynamics of a Microswimmer in Poiseuille Flow. *Phys Rev Lett* (2012) 108:218104. doi:10.1103/physrevlett.108.218104
- Crocker JC, Weeks ER. Particle Tracking Using IDL (2011). Available at: http://physics.emory.edu/faculty/weeks/idl/.
- 40. Ran R, Brosseau Q, Blackwell BC, Qin B, Winter RL, Arratia PE. Bacteria Hinder Large-Scale Transport and Enhance Small-Scale Mixing in Time-Periodic Flows. *Proc Natl Acad Sci U S A* (2021) 118:e2108548118. doi:10.1073/ pnas.2108548118
- Chen H, Thiffeault J-L. Shape Matters: a Brownian Microswimmer in a Channel. J Fluid Mech (2021) 916:A15. doi:10.1017/jfm.2021.144
- Dombrowski C, Cisneros L, Chatkaew S, Goldstein RE, Kessler JO. Selfconcentration and Large-Scale Coherence in Bacterial Dynamics. *Phys Rev Lett* (2004) 93:098103. doi:10.1103/PhysRevLett.93.098103
- Sokolov A, Aranson IS, Kessler JO, Goldstein RE. Concentration Dependence of the Collective Dynamics of Swimming Bacteria. *Phys Rev Lett* (2007) 98: 158102. doi:10.1103/physrevlett.98.158102
- Dunkel J, Heidenreich S, Drescher K, Wensink HH, Bär M, Goldstein RE. Fluid Dynamics of Bacterial Turbulence. *Phys Rev Lett* (2013) 110:228102. doi:10.1103/physrevlett.110.228102

Conflict of Interest: Author PJ is employed by Velico Medical inc.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Publisher's Note: All claims expressed in this article are solely those of the authors and do not necessarily represent those of their affiliated organizations, or those of the publisher, the editors and the reviewers. Any product that may be evaluated in this article, or claim that may be made by its manufacturer, is not guaranteed or endorsed by the publisher.

Copyright © 2022 Yoest, Buggeln, Doan, Johnson, Berman, Mitchell and Solomon. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.