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Hotspots of boundary accumulation: Dynamics and statistics of micro-swimmers in flowing films

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Biological flows over surfaces and interfaces can result in accumulation hotspots or depleted voids of microorganisms in natural environments. Apprehending the mechanisms that lead to such distributions is essential for understanding biofilm initiation. Using a systematic framework we resolve the dynamics and statistics of swimming microbes within flowing films, considering the impact of confinement through steric and hydrodynamic interactions, flow, and motility, along with Brownian and run-tumble fluctuations. Micro-swimmers can be peeled off the solid wall above a critical flow strength. However, the interplay of flow and fluctuations causes organisms to migrate back towards the wall above a secondary critical value. Hence, faster flows may not always be the most efficacious strategy to discourage biofilm initiation. Moreover, we find run-tumble dynamics commonly used by flagellated microbes to be an intrinsically more successful strategy to escape from boundaries than equivalent levels of enhanced Brownian noise in ciliated organisms.

I. INTRODUCTION

Surfaces, interfaces and confinements are ubiquitous in microbial environments.^{1,2} Living near surfaces can provide a variety of benefits to microbes over life in bulk fluids. Whereas interfaces may be the source of oxygen and sunlight, solid surfaces accumulate sediments including nutrients and offer anchoring points for the formation of extracellular matrices and biofilms. Moreover, noslip surfaces locally slow flows and may provide peaceful respite from violent mixing. However, an unavoidable consequence of no-slip surfaces embedded in flows is shearing and, though microbes at surfaces may not be subject to large velocities, they are commonly subjected to non-negligible shears.³

Confinement within films, between a rigid substrate and an interface, is particularly fascinating with respect to microbial life since it offers the immediate presence of two surfaces with differing properties. Intense biological activities and accumulations of swimming cells are often associated with films.^{4–6} Indeed, countless species of bacteria secrete their own extracellular polymeric substances in order to form biofilms.^{7,8} Films on passive or living substrates allow pathogens to swim or even swarm in order to colonise a wide variety of surfaces including soil, plant leaves, animal tracts or skin.^{9–11} Liquid-air interfaces have recently attracted interest because of bioremediation of oil consuming bacteria at oil spill sites.^{12–14}

The majority of previous research on swimming in confined environments has focused on investigating the effect of a single solid boundary on swimming dynamics. In particular, the interactions between organisms and surfaces have been studied in detail, both theoretically^{15–28} and in experiments.^{29–36} Moreover, biological traits such as nutrient uptake in a quiescent fluid have been numerically investigated for a suspension of spherical model



FIG. 1. A micro-swimmer immersed in a liquid film. The film flows in the x direction, indicated by the arrows and the parabolic velocity profile. The micro-swimmer is advected and rotated by this flow, and interacts with the film surfaces, sterically, and hydrodynamically through its self-generated flow field. Angles $-\pi/2 < \phi < \pi/2$ indicate upstream orientation.

swimmers (squirmers) constrained between a wall and a free surface.³⁷ The interplay of a flowing fluid and swimming cells has been studied recently for Newtonian^{38–46} and non-Newtonian fluids.^{47–51} Additionally, biological systems often feature large fluctuations that affect the dynamics of living cells.^{52–59} Together, these works emphasise the pivotal roles of confinement, the background flow and noise as determinants of swimmer trajectories in micro-environments. However, despite the widespread implications the combined effects of motility, external flows, hydrodynamics and biological non-determinism remain obscure due to the complexity of the dynamics.

Here, we present a comprehensive description of the dynamics and statistics of swimming microbes in flowing films between a solid boundary and a free surface. We

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focus on two model micro-organism classes: flagellated swimmers, like E. coli bacteria, and ciliated swimmers, such as *Volvox carteri* microphytes (**§IV B**). To model swimming trajectories and the distribution of microswimmers within films, we consider the various contributions to a swimmer's equation of motion, first separately then concurrently. We begin by considering only the effects of external flow and steric interactions with the film boundaries (**§III**) before including detailed hvdrodynamic interactions with the two surfaces ($\S IV$). Additionally, stochastic effects must also be accounted for. We scrutinise the swimmer distributions that will arise in flowing films because swimmers are subjected to thermal noise $(\S V A)$. It is well known that the primary source of biological stochasticity in many flagellated microbes is run-tumble dynamics, whereas ciliated organisms are subject to enhanced Brownian fluctuations, e.g. due to cilia beating out of synchrony. Run-tumble dynamics are seen to prevent boundary accumulation more successfully than equivalent levels of enhanced Brownian noise $(\S V B)$. Our results have implications for the control of cell distributions within flowing films, and highlight the cellular swimming strategies against which defouling schemes must contend.

II. SWIMMER DYNAMICS IN A FILM

We consider a motile swimming microbe within a flowing film, with a bottom no-slip wall at z = 0 and a noshear top interface at height z = H. The micro-swimmer is modelled as an ellipsoid of semi-major and minor axes a and b at a position $\mathbf{r} = (x, y, z)$ and orientation \mathbf{p} (**Fig.** 1). It should be noted that we do not account separately for the swimmer's body and flagella, but we account for the swimmer's shape by modelling it as an ellipsoid with an aspect ratio that includes the flagellar length. Following examples from the literature, 25,39,42 we describe the microbe's position and orientation with the equations of motion

$$\dot{\boldsymbol{r}} = \boldsymbol{v}^{\mathrm{S}} + \boldsymbol{v}^{\mathrm{F}} + \boldsymbol{v}^{\mathrm{ST}} + \boldsymbol{v}^{\mathrm{HI}} + \boldsymbol{v}^{k_{\mathrm{B}}T}, \qquad (1)$$

$$\dot{\boldsymbol{p}} = \left(\boldsymbol{\Omega}^{\mathrm{F}} + \boldsymbol{\Omega}^{\mathrm{ST}} + \boldsymbol{\Omega}^{\mathrm{HI}} + \boldsymbol{\Omega}^{k_{\mathrm{B}}T} + \boldsymbol{\Omega}^{\mathrm{RT}}\right) \times \boldsymbol{p}.$$
 (2)

The contributions to the motion come from selfpropulsion (a swimming velocity $\boldsymbol{v}^{\mathrm{S}} = v_{\mathrm{s}} \boldsymbol{p}$), background flow (characterised by velocity $\boldsymbol{v}^{\mathrm{F}}$ and angular velocity $\Omega^{\rm F}$), steric interactions with the substrate and the inter-face in a film ($v^{\rm ST}$ and $\Omega^{\rm ST}$), hydrodynamic interactions with the surfaces ($v^{\rm HI}$ and $\Omega^{\rm HI}$), and stochastic dynam-ics due to thermal noise ($v^{k_{\rm B}T}$ and $\Omega^{k_{\rm B}T}$) or run-tumble dynamics (controlled by Ω^{RT}). We shall consider each of these in turn.

The translational invariance of Eqs. (1-2) along the directions parallel to the surfaces allows us to consider motion of swimmers in the y = 0 plane where we take the flow along the x direction. The swimmer orientation is represented in cylindrical coordinates as p =

 $-(\cos\phi, 0, \sin\phi)$, where $\phi \in [-\pi, \pi]$ is the angle in the x-z plane, where upstream orientation corresponds to $\phi = 0$ (Fig. 1). The dynamics of the system can be calculated by solving two coupled equations,³⁹ $\dot{z} = \dot{z}(z, \phi)$ and $\dot{\phi} = \dot{\phi}(z, \phi)$, followed by integrating two further uncoupled equations, $\dot{x} = \dot{x}(z, \phi)$ and $\dot{y} = \dot{y}(z, \phi)$.

III. SWIMMING CELL TRAJECTORIES WITH STERIC INTERACTIONS

We begin by considering the simplified picture that microbes swim in a flowing film where their interactions with the film boundaries are purely steric. Several recent studies have considered pure kinematic interaction of swimmers with solid boundaries in the search for non-hydrodynamic mechanisms of boundary accumulation.^{60,61} Neglecting hydrodynamic interactions is expected to give qualitatively relevant conclusions but not quantitative results. In \S{IV} and \S{V} we characterise the roles of hydrodynamic interactions and noise, respectively, but until then we only have Eq. 1 and Eq. 2 with $\left\{ \boldsymbol{v}^{\text{HI}}, \boldsymbol{v}^{k_{\text{B}}T}, \boldsymbol{\Omega}^{\text{HI}}, \boldsymbol{\Omega}^{k_{\text{B}}T}, \boldsymbol{\Omega}^{\text{RT}} \right\} = \boldsymbol{0}.$ The background fluid flow in the *x* direction is pre-

scribed in the form of a half-parabolic profile,⁶²

$$\boldsymbol{u}^{\mathrm{F}}(\boldsymbol{r}) = v_{\mathrm{max}} \left[2\frac{z}{H} - \frac{z^{2}}{H^{2}} \right] \boldsymbol{\hat{e}}_{x}, \qquad (3)$$

which has a maximum speed v_{max} at z = H. The flow u^{F} enacts a drag on a swimmer at position r, which induces the velocity v^{F} and angular velocity Ω^{F} . The flow-induced translational and rotational velocities of the force-free and torque-free swimmer are

$$\boldsymbol{v}^{\mathrm{F}} = v_{\mathrm{max}} \left[2\frac{z}{H} - \frac{z^2}{H^2} \right] \boldsymbol{\hat{e}}_x, \tag{4}$$

$$\mathbf{\Omega}^{\mathrm{F}} = v_{\mathrm{max}} \left[\frac{H-z}{H^2} \left(1 - G \cos 2\phi \right) \right] \hat{\boldsymbol{e}}_y, \tag{5}$$

where the geometry factor $G = \frac{\gamma^2 - 1}{\gamma^2 + 1} \in [0, 1)$ is a function of the aspect ratio $\gamma = a/b$ of an elongated swimmer.

Steric interactions with the surfaces are modelled by a repulsive force and torque, resulting in the velocity and angular velocity

$$\boldsymbol{v}^{\text{ST}} = v_{\text{s}} A(\phi)^{12} \left(\frac{1}{z^{12}} - \frac{1}{(H-z)^{12}} \right) \hat{\boldsymbol{e}}_{z},$$
 (6)

$$\mathbf{\Omega}^{\rm ST} = a^2 v_{\rm st} \xi^{-1} G \sin\left(2\phi\right) \hat{\boldsymbol{e}}_y,\tag{7}$$

so that a microbe facing a surface would be at equilibrium at the distance of closest approach between the swimmer's body and the boundary.³⁸ For an elongated swimmer, that distance is $A(\phi) = \sqrt{a^2 + (b^2 - a^2)\cos^2 \phi}$. The steric torque is derived from the potential⁶³ $U \propto \cos^2 \phi$ so that $\dot{\phi}^{ST} = -\frac{1}{\xi} \frac{\partial U}{\partial \phi}$, where ξ is the rotational drag coefficient with units of volume and $v_{\rm st} = |\boldsymbol{v}^{\rm ST}|$.

The trajectories of a microbial swimmer in a flowing film (without hydrodynamic interactions or noise) are shown in Fig. 2. The vorticity of the background flow



FIG. 2. Deterministic trajectories of a micro-swimmer in a liquid film flowing from left to right, with steric interactions but without hydrodynamic interactions. Three trajectories are shown with initial orientation $\phi = 0.1$, x = 0 and z/H = 0.1, 0.5, 0.9 in solid red, dashed blue and dotted green lines, respectively. The background flow is (a,b) slow: $v_{\text{max}} = 2v_s$ and (c,d) fast: $v_{\text{max}} = 10v_s$. The swimmer body shape is (a,c) spherical (like *Volvox*) with a = b = H/20 and (b,d) elongated (like *E. coli*) with aspect ratio $\gamma = 3$, size a = 3b = 3H/20 and rotational drag coefficient $\xi = 10a^3$. (Main panels) Dynamics in real x - z space. Motion is from left to right, along with the background flow, and arrows indicate the direction of swimmer orientation. (Side panels) Corresponding dynamics in $\phi - z$ phase space. Background colours indicate the sign of the z component of the swimmer velocity. The white regions are inaccessible for the swimmer due to the steric interactions.

is strongest at the bottom wall but vanishes at the top, turning an upstream-oriented swimmer up towards the film interface. In slowly flowing films, spherical swimmers, such as *Volvox*, remain there whilst being advected downstream and slowly rotated, until their orientation is sufficiently changed to swim down into the film, where the vorticity is larger. Hence, the swimmer 'dips' down but quickly finds itself returned towards the free interface once again, only for the process to repeat (**Fig. 2**a).

Elongated swimmers, such as E. coli, tend to reside at the top surface longer due to their Jeffery orbits⁴¹ (Fig. 2b). Furthermore, the steric interactions reorient the long bodies parallel to the surface. When an elongated swimmer is oriented in the upstream direction at the top interface, the vorticity and steric torque due to motility counter each other, leading to a stable fixed point in phase space at $(\phi = 0, z = H - b)$ (red and green trajectories). The downstream-oriented fixed point is unstable at the top of the film, because the vorticity and steric torque cooperate, so that the swimmer can move away from the top interface (blue trajectory). Conversely, at the bottom wall the (upsteam-) downstreamorientation fixed point is (unstable) stable, as shown by the red trajectory. Therefore, even without hydrodynamics, steric interactions can cause trapping of elongated swimmers at surfaces, oriented downstream at the bottom wall and upstream at the interface, in agreement with simulations²² and experiments.³¹

At sufficiently large flow speeds, an additional type of trajectory can be observed: A fast rotation ('spinning') of swimmers. Spinning trajectories are confined to the lower regions of the film where, due to the strong vorticity of the flow, the swimmer is rotated rapidly. Hence, its motility averages to zero, leaving the swimmer vorticitytrapped. For spherical *Volvox*, we see spinning in the lower region of the film without trapping at the bottom wall (Fig. 2c; red trajectory). In the top region, swimmers start 'dipping' but interact sterically with the top surface. Therefore, they are forced to follow the trajectory exactly between dipping and spinning in the middle region (blue and green trajectories). This special trajectory is called the separatrix, in which swimmers dip from the top, down to a critical height z^* . Elongated E. coli also feature dipping and spinning in the middle region (Fig. 2d; blue trajectory), but in the top and bottom regions they can get trapped close to the surfaces (red and green trajectories). The chance of encountering a surface is smaller when the motility is averaged to zero in rapid rotation, thus the probability of spinning increases with flow speed.

The transition between dipping and spinning is defined by the separatrix in phase space. Dipping trajectories are above this line and spinning trajectories below the separatrix. The lowest point in the film that this special trajectory can reach is the critical height,

$$z^* = \left(1 - \frac{2}{\sqrt{v_{\text{max}}/v_{\text{s}}}}\right)H,\tag{8}$$

both for spherical *Volvox* and elongated *E. coli*. Therefore, the separatrix touches the bottom wall at a critical flow rate of $v_{\text{max}} = 4v_{\text{s}}$. Spinning trajectories do not exist below that critical flow. The larger the flow rate, the larger the fraction of phase space taken up by spinning trajectories, and therefore the probability of finding these trajectories in the lower regions of the film.

In summary, these results show that in the absence of hydrodynamic interactions and noise, the background flow and steric interactions of a micro-swimmer with the film boundaries lead to distinct swimmer trajectories depending critically on background flow speed. Above the critical flow speed $v_{\rm max} > 4v_{\rm s}$ a swimmer is likely to be observed spinning near the bottom boundary due to the trapping in the high-vorticity region, whereas if $v_{\rm max} < 4v_{\rm s}$ swimmers tend to reside near the top surface.

IV. TRAJECTORIES OF SWIMMING CELLS WITH HYDRODYNAMIC INTERACTIONS

In addition to differing size and shape, *Volvox* and *E.* coli have pronouncedly different swimming strategies and consequently have very different hydrodynamic interactions with the bounding surfaces of the film through the swimmer-generated flow field.⁶⁴ In order to derive the hydrodynamic interactions ($v^{\rm HI}$ and $\Omega^{\rm HI}$ in Eq. 1 and Eq. 2), we first compute the swimmer-generated flow field in a film (§IV A). We then use our results to describe the specific hydrodynamic effects on trajectories of our example swimmers (*E. coli* and *Volvox*; §IV B) and study the opposing effects of hydrodynamic boundary accumulation versus peeling of swimmers from the bottom wall by flow (§IV C).

A. Hydrodynamic interactions with surfaces

To find the flow at the position \boldsymbol{x} produced by a microorganism at $\boldsymbol{r} = (x, y, z)$ we first require the flow solution of a point force (Stokeslet) in a film. Once the Stokeslet solution is established, we use a multipole expansion to find the flow field generated by a micro-swimmer. Full details can be found in Ref.⁶⁵. This flow field then allows us to compute the hydrodynamic interactions of the swimmer with the surfaces and other swimmers. For simplicity, we consider organisms in the dilute limit, and therefore do not include swimmer-swimmer interactions.

In unbounded fluids, the fluid velocity due to a point force \boldsymbol{f} is

$$\boldsymbol{u}^{\mathrm{S}\infty}(\boldsymbol{x},\boldsymbol{r},\boldsymbol{f}) = \left(\frac{\mathbb{I}}{\overline{d}} + \frac{\boldsymbol{d}\boldsymbol{d}}{\boldsymbol{d}^3}\right) \cdot \frac{\boldsymbol{f}}{8\pi\mu},\tag{9}$$

where d = x - r and $\underline{\mathbb{I}}$ is the identity matrix. We account for the presence of the wall and the interface by including auxiliary flow fields, written in terms of image systems. Surfaces are mathematically replaced with a hydrodynamic image system, analogous to the method of images in classical electrostatics. To account for the no-slip wall, we consider an image swimmer located at the position $\mathcal{R}^{(0)} = (x, y, -z)$ and producing an additional flow field that is given by the Blake tensor.⁶⁶ The

no-shear interface located at z = H also requires a hydrodynamic image at the position $\mathcal{R}^{(-1)} = (x, y, 2H - z)$ and the flow field produced by this image is a simple direct reflection of the Stokeslet flow (**Eq. 9**).

However, a single image for each surface is not sufficient: Just as two parallel mirrors create an infinite series of images, so too do two parallel surfaces. These hydrodynamic images are found at the positions $\mathbf{r}^{(n)} = (x, y, z - 2nH)$ and $\mathcal{R}^{(n)} = (x, y, -z - 2nH)$, where $n = 0, \pm 1, \pm 2, \ldots$ and $\mathbf{r} = \mathbf{r}^{(0)}$. The flows produced by each image can be determined by successively repeating reflection operations to produce the appropriate hydrodynamic images for the bottom wall or top interface. The final flow at the point \mathbf{x} due to a Stokeslet at \mathbf{r} in the film is given by

$$\boldsymbol{u}^{\mathrm{S}}(\boldsymbol{x},\boldsymbol{r},\boldsymbol{f}) = \underline{\underline{\mathcal{F}}} \cdot \boldsymbol{f} / (8\pi\mu), \tag{10}$$

$$\underline{\underline{\mathcal{F}}}(\boldsymbol{x},\boldsymbol{r}) = \sum_{n=-\infty}^{\infty} \left[\underline{\underline{\mathcal{G}}}(\boldsymbol{x},\boldsymbol{r}^{(n)}) + \underline{\underline{\mathcal{G}}}(\boldsymbol{x},\boldsymbol{\mathcal{R}}^{(n)})\right], \quad (11)$$

where the image tensors $\underline{\underline{\mathcal{G}}}(x, r^{(n)})$ and $\underline{\underline{\mathcal{G}}}(x, \mathcal{R}^{(n)})$ are given explicitly in Ref.⁶⁵ In practice, the series converges rapidly and is safely truncated after only a few images on each side of the film. In this work we use four images on each side of the film in all presented text and figures, unless explicitly stated otherwise.

Knowing the analytic form of the Stokeslet in a film allows us to find the flow field generated by a motile microbe through the use of a multipole expansion.⁶⁷ Neutrally buoyant micro-swimmers are force-free and so do not subject their surrounding fluid to a net force but rather a force dipole and higher order moments. Hence, the flow generated by the swimmer is

$$\boldsymbol{u}(\boldsymbol{x},\boldsymbol{r},\boldsymbol{p}) = \boldsymbol{u}^{\mathrm{D}} + \boldsymbol{u}^{\mathrm{Q}} + \boldsymbol{u}^{\mathrm{SD}} + \boldsymbol{u}^{\mathrm{RD}} + \dots$$
 (12)

where each term represents a multipole term in the swimmer-generated flow field: u^{D} is the Stokes dipole, u^{Q} the quadrupole, u^{SD} the source doublet, and u^{RD} the rotlet doublet flow. They are related to Eq. 11 by

$$\boldsymbol{u}^{\mathrm{D}}(\boldsymbol{x},\boldsymbol{r},\boldsymbol{p}) = \kappa \ (\boldsymbol{p} \cdot \tilde{\boldsymbol{\nabla}})(\underline{\boldsymbol{\mathcal{F}}} \cdot \boldsymbol{p}), \tag{13}$$

$$\boldsymbol{u}^{\mathrm{Q}}(\boldsymbol{x},\boldsymbol{r},\boldsymbol{p}) = -\frac{1}{2}\nu \ (\boldsymbol{p}\cdot\tilde{\boldsymbol{\nabla}})^{2}(\underline{\boldsymbol{\mathcal{F}}}\cdot\boldsymbol{p}), \tag{14}$$

$$\boldsymbol{u}^{\mathrm{SD}}(\boldsymbol{x},\boldsymbol{r},\boldsymbol{p}) = -\frac{1}{2}\sigma \; \tilde{\nabla}^2(\underline{\boldsymbol{\mathcal{F}}}\cdot\boldsymbol{p}),$$
 (15)

$$\boldsymbol{u}^{\mathrm{RD}}(\boldsymbol{x},\boldsymbol{r},\boldsymbol{p}) = -\frac{1}{2}\tau \ (\boldsymbol{p}\cdot\tilde{\boldsymbol{\nabla}})\tilde{\boldsymbol{\nabla}}\times(\underline{\boldsymbol{\mathcal{F}}}\cdot\boldsymbol{p}), \qquad (16)$$

where the derivatives act on \boldsymbol{r} . The multipole coefficient κ has units $[\mu \text{m}^3/\text{s}]$ and ν, σ, τ have units $[\mu \text{m}^4/\text{s}]$. Note that the infinite image series and the multipole expansion can be used in conjunction because of the linearity of the Stokes equations.

Eqs. 13-16 account for the generic attributes of microswimmers. The dipole $\boldsymbol{u}^{\mathrm{D}}$ models the opposing propulsion and drag forces on the fluid. Pusher-type swimmers drive fluid out along the swimming axis and $\kappa > 0$, while pullers have $\kappa < 0$. The quadrupole term $\boldsymbol{u}^{\mathrm{Q}}$ represents the fore-aft asymmetry of the microorganism with $\nu > 0$



FIG. 3. Typical deterministic swimming trajectories in a quiescent film for two different organism types: (a) *E. coli* bacteria as an example of flagellated swimmers. $H = 10\mu$ m, $a = 1.5\mu$ m, $b = 0.5\mu$ m, $v_s = 50\mu$ m/s, $\kappa = 30\mu$ m³/s, $\sigma = -25\mu$ m⁴/s, $\nu = 25\mu$ m⁴/s, $\tau = -8.3\mu$ m⁴/s and $\xi = 1\mu$ m³. Axes units are in μ m. (b) *Volvox* microphytes as an example of ciliated swimmers. H = 4mm, $a = b = 200\mu$ m, $v_s = 100\mu$ m/s, $\kappa = 10^6\mu$ m³/s, $\sigma = 10^9\mu$ m⁴/s and $\nu = \tau = 0\mu$ m⁴/s. Axes units are in mm. (a,b) All other multipole moments are set to zero. Trajectories result from numerical integration of Eqs. 1-2. Colour indicates time passing, ranging from violet (t = 0) to red (t = 2s for (a) and t = 4s for (b)). Inset images show typical examples of flagellated and ciliated organisms.

expected for flagellated bacteria, such as *E. coli*. The finite hydrodynamic size of the swimmer is included via the source doublet u^{SD} . Ciliated organisms, like *Volvox*, possess positive σ values, while for non-ciliated swimmers $\sigma < 0$. The rotlet doublet u^{RD} represents opposing rotation of a swimmer's head and tail.

The Faxén relations couple the translation and rotation of the swimmer $(v^{\rm HI} \text{ and } \Omega^{\rm HI})$ with the flow (u). By solving these relations for the force-free and torquefree swimmer, the surface-induced translational and rotational velocities are found from Eq. 12 as a function of swimmer position and orientation.⁶⁵ It must, however, be noted that near-wall swimming might strongly deviate from the predictions of a multipole expansion due to contact interactions.^{34,35}

B. Two specific examples: E. coli and Volvox

Here we apply our model to two specific example organisms of different classes: the flagellated *E. coli* bacterium and the ciliated *Volvox carteri* microphyte. Many of the micro-swimmers occurring in nature can be classified in one of these two categories. For a detailed list of species, we refer to e.g. Lighthill.⁶⁸ Using experimental data we estimate the swimming parameters including the multipole coefficients and swimming speed, and use these to calculate the swimming trajectories.

For *E. coli*-like organisms, we estimate the dimensions as $a = 1.5\mu$ m and $b = 0.5\mu$ m. The dipole strength has been measured as $\kappa \approx 30\mu$ m³/s.³⁴ The source doublet coefficient is expected to be negative for a nonciliated organism and can be estimated from an approximate hydrodynamic size $a_{\rm H} \sim 1\mu$ m and swimming speed $v_{\rm s} \approx 50\mu$ m/s,⁶⁹ leading to $\sigma \approx -\frac{1}{2}a_{\rm H}^3 v_{\rm s} \approx -25\mu$ m⁴/s. Similarly, we estimate the quadrupolar coefficient $\nu \approx$ 25μ m⁴/s for a flagellated bacterium. The rotlet dipole coefficient τ can be estimated from experimental measurements of the radius of curvature of swimming trajectories near surfaces.^{29,70–73} Using a radius of curvature $R \approx 20\mu \text{m}$, aspect ratio $\gamma = 3$ and swimming height $z = b \approx 0.5\mu \text{m}$ gives $\tau \approx \frac{32z^4 v_{\text{s}}}{3R(1-G)} \approx -8.3\mu \text{m}^4/\text{s}$, where $G = \frac{\gamma^2 - 1}{\gamma^2 + 1}$ and the minus sign corresponds to swimming in the clock-wise direction above the no-slip wall.⁶⁵ To emphasise the effects of a relatively thin film, we choose $H = 20b = 10\mu \text{m}$.

For Volvox-like organisms, we estimate the dimensions as $a = b = 200\mu m$, and for the film height we choose H = 20b = 4mm. The primary multipole coefficients have been measured as $\kappa \approx 10^6 \mu m^3/s$ and $\sigma \approx 10^9 \mu m^4/s$ as has the swimming velocity $v_s \approx 100 \mu m/s$.⁷⁴ This type of organism is fore-aft symmetric and so $\nu \approx 0$, and it does not have rotating flagella so we also expect $\tau \approx 0$. Volvox is not always neutrally buoyant and therefore a Stokeslet flow field should be present, but we neglect the effects of gravity here, assuming the sedimentation speed is smaller than the swimming speed. This model would, however, be even more appropriate for the many smaller ciliated micro-organisms in nature⁶⁸ of which the multipole coefficients have not yet been measured directly.

Fig. 3 shows typical trajectories for the two example organisms in the absence of flow. Random initial positions and orientations were chosen to show the diversity of the dynamics. E. coli-like organisms tend to accumulate at the surfaces (Fig. 3a), with a small bias towards the bottom wall because of the dipolar hydrodynamic interactions, and they remain tightly bound there because of quadrupolar hydrodynamics, in agreement with earlier calculations.⁶⁵ At both surfaces the swimmers move in clockwise circles, but the radius of curvature of the trajectories at the bottom wall is larger than at the top interface by a factor of $\gamma^2 + \mathcal{O}(b^2/H^2) \approx 9$ due to the body elongation. Volvox-like organisms, in contrast, turn away from the bottom wall but assemble at the top interface (Fig. 3b). The hydrodynamic binding to the top interface is not so strong as that of the bacteria, because of the positive source doublet moment. With a small fluctuation this allows for motion away from the interface into the film, before returning back to the top again.

The resulting trajectories for these different swimmer types could offer important biological implications. Micro-organisms such as *E. coli* bacteria might have developed into slender and fairly fore-aft symmetric bodies that tend to accumulate at boundaries, e.g. to facilitate biofilm formation. Likewise, sperm cells could benefit from the ability to swim upstream along the walls of the female reproductive tract. On the other hand, ciliated organisms like the *Paramecium* protozoa or *Volvox* tend to accrue at the top interface only, which could be beneficial to collect oxygen or to facilitate photosynthesis.

C. Flow-induced peeling

Next, we re-introduce the external flow to the dynamics of the swimming microbes, in addition to steric and hydrodynamic interactions with the surfaces. The trajectories for swimmers including hydrodynamic interactions (**Fig. 3**) are substantially modified because of the additional vorticity of the background flow. They more closely resemble the swimming paths of **Fig. 2** far from the surfaces. However, close to the surfaces, and particularly near the bottom wall where the vorticity is largest, there is a competition between the external flow field and the hydrodynamic interactions with the wall.

The external flow can prevent hydrodynamic interaction-induced boundary accumulation by peeling swimmers off the bottom wall. This detachment from surfaces by imposed flows has been demonstrated recently in experiments.⁴⁵ In **Fig. 4** the minimum flow strength required to detach a swimmer from the bottom wall is shown as a function of dipole moment κ and source doublet moment σ . Notice the asymmetry – a pusher can be washed off more easily than its equivalent puller. Recall that both *E. coli* and *Volvox* are pushers with $\kappa > 0$. Neutral swimmers ($\kappa = 0$; steric interactions only) can accumulate at the surfaces in strong confinement, but can also be detached with small background flows.

We theoretically estimate the critical flow speed required to detach swimmers from the bottom wall by equating the angular velocity of the dipolar interactions $(\Omega^{D}$ given in Ref.⁶⁵) and of the flow (Eq. 5). Considering only two images and spherical swimmers, we obtain

$$\frac{v_{\max}}{v_{s}}\Big|_{\rm crit} = \frac{3}{16} \frac{H |\kappa|}{a^{3} v_{s}} \left|\sin 2\phi_{c}\right|,\tag{17}$$

where ϕ_c is the critical angle that the swimmer must rotate through to overcome the hydrodynamic barrier. Note that this expression scales linearly with the film height, because the local vorticity decreases with increasing *H*. This equation is straightforwardly generalised to account for elongated swimmers, such as *E. coli*, by replacing the radius *a* with the distance of closest approach between the ellipse and the wall, $A(\phi) =$



FIG. 4. Minimum flow strength required to peel microswimmers off the bottom wall of a liquid film, for (a) A swimmer with only a variable dipole moment $\kappa \ [\mu m^3/s]$. (b) A swimmer with variable dipole and source doublet moment $\sigma \ [\mu m^4/s]$. Lines in (a) show theoretical predictions for a puller (red, solid) and pusher (blue, dashed) from Eq. 17. Blue dots in (a) and contours in (b) result from numerical integration of Eqs. 1-2. Included are motility, external flow, steric and hydrodynamic interactions with the walls, but no noise. All other multipole coefficients are zero. Swimmers are released facing the bottom wall ($z = a, \phi = \pi/2$). Film height $H = 10\mu$ m, swimmer dimensions $a = b = 0.5\mu$ m and swimming speed $v_s = 50\mu$ m/s.

 $\sqrt{a^2 + (b^2 - a^2) \cos^2 \phi}$. Furthermore, by defining a dimensionless number $\lambda = a^3 v_{\text{max}}/\kappa H$ that characterises the degree of detachment by external flows (**Table I**), **Eq. 17** can be recast as $\lambda_{\text{crit}} = 3 |\sin 2\phi_c| / 16$, which is independent of the swimming speed. Here $\lambda < \lambda_{\text{crit}}$ corresponds to attachment and $\lambda > \lambda_{\text{crit}}$ to detachment.

For pullers, the equilibrium orientation without flow is pointing towards the wall, $\phi = \pi/2$. Hence, pullers must turn upwards over the hydrodynamic barrier from $\phi_c = \pi/2$ to the orientation parallel to the wall, $\phi = (0, \pi)$, in order to swim away. Inserting this angle ϕ_c and the figure parameters into **Eq. 17** gives a prediction for the minimum flow strength required for pullers (red line in **Fig. 4**a).

Pushers are oriented parallel to the wall at equilibrium, but they must also overcome the attraction towards the bottom wall. Therefore, the angle at which the pushers can swim away, using two image systems, is given by the requirement that

$$-\sin\phi > \frac{3}{16}\frac{\kappa}{v_{\rm s}a^2}(3\cos 2\phi - 1).$$
 (18)

Inserting the resulting angle ϕ_c into Eq. 17 gives a prediction for the minimum flow strength required for pushers (blue dashed line in Fig. 4a). Since the angle is smaller for pushers than for pullers, the former can escape from the surface more easily.

The inclusion of higher-order multipoles in the hydrodynamic interactions with the surfaces changes the critical flow speed (**Fig. 4**b). Flagellated swimmers with source doublet moment $\sigma < 0$ are more tightly bound to the wall, thus requiring a stronger flow to peel them off. Pushers such as *E. coli* in **Fig. 3**a with $\kappa = 30 \mu \text{m}^3/s$ and $\sigma = -25 \mu \text{m}^4/s$ require $v_{\text{max}} \approx 25 v_{\text{s}}$ compared to $\approx 2.5 v_{\text{s}}$ when $\sigma = 0$. Non-ciliated pullers such as *Chlamy*- domonas reinhardtii with both $\kappa, \sigma < 0$ are even more tightly bound. Ciliated swimmers with $\sigma > 0$, on the other hand, are much less strongly attracted to the wall (**Fig. 3**b) and hardly need any flow for detachment.

V. SWIMMER DISTRIBUTIONS

Swimmer trajectories in real microbial environments are subject to stochastic fluctuations. Having established the deterministic features of the swimmer trajectories in the presence of flow, steric and hydrodynamic interactions, we now examine the distributions of microswimmers if noise, originating from both thermal fluctuations and run-tumble dynamics, is added. This is important because since measuring distributions of swimming cells is easier in many experimental setups than following individual trajectories. For simplicity, in the following we examine archetypal spherical swimmers with a dominant contribution from dipolar hydrodynamic interactions.

We model the thermal/Brownian noise as drawn from a Gaussian distribution so that the mean squared displacement and angular displacement are $\langle |\mathbf{r}(t) - \mathbf{r}(0)|^2 \rangle = 6Dt$ and $\langle |\phi(t) - \phi(0)|^2 \rangle = 2D_r t$ in the large-time limit, where D and D_r are the translational and rotational Brownian diffusion coefficients, respectively. For typical micronsized swimmers we approximate the thermal diffusion coefficients as $D \approx 0.25 \mu \text{m}^2/s$ and $D_r = 0.2 \text{rad}^2/\text{s}$. We assume that these diffusion coefficients are isotropic and remain constant as a function of film height. To be concrete, we consider a narrow film of height $H = 10\mu\text{m}$ and organisms of dimensions $a = b = 0.5\mu\text{m}$ and swimming velocity $v_s = 50\mu\text{m/s}$.

When only thermal noise is considered, the swimmers' trajectories are deterministic over a timescale of crossing the film. Once they reach a surface, there is a competition between the noise, flow and hydrodynamic interactions. Using only the dipolar contribution from the first image system ($v^{\rm D}, \Omega^{\rm D}$ given in Ref.⁶⁵), we define translational and rotational Hydrodynamic Interactions Péclet numbers to be

$$\mathrm{P\acute{e}^{HI}} \equiv \frac{v^{\mathrm{HI}}a}{D} = \frac{3\kappa}{8b^2} \frac{a}{D},$$
 (19)

$$\mathbf{P}\mathbf{\acute{e}_r^{HI}} \equiv \frac{\Omega^{\mathrm{HI}}}{D_r} = \frac{3\kappa}{16b^3} \frac{1}{D_r}.$$
 (20)

With a dipole strength of $\kappa=30\mu\mathrm{m}^3/\mathrm{s}$, we estimate $\mathrm{P\acute{e}^{HI}}\sim100$ and $\mathrm{P\acute{e}^{HI}}_\mathrm{r}\sim200$. Since both $\mathrm{P\acute{e}^{HI}}$, $\mathrm{P\acute{e}^{HI}}_\mathrm{r}\gg1$, thermal noise is not sufficient to allow swimmers to overcome hydrodynamic attraction to the surfaces, in the absence of a background flow. Experiments have shown that flagellated bacteria have smaller passive rotational diffusion coefficients than unflagellated cells, 34,56 so $\mathrm{P\acute{e}^{HI}}_\mathrm{r}\sim200$ is in fact a conservative estimate. Consequently, other mechanisms must be employed by swimming cells to prevent boundary accumulation within films.

In §V A, we first study the combined effect of Brownian noise and external flow on distributions of archetypal micro-swimmers, i.e. *neutral* swimmers, *pushers* and *pullers*. In §V B, we add non-thermal fluctuations and investigate the impact of the nature of this noise by comparing swimmers subject to run-and-tumble dynamics (*tumblers*) and enhanced Brownian noise (*drifters*).

A. Prevention of boundary accumulation by external flow

§III describes the effect of flow on steric swimmers without noise. In **§IV** we include hydrodynamic interactions with the surfaces, but still exclude noise. In this section, we investigate the combined effects of steric and hydrodynamic interaction, film flow, and thermal noise.

The accumulation of swimmers at the surfaces is evaluated using the swimmer density distribution $\rho(z, \phi)$, normalised such that $(2\pi H)^{-1} \int_0^H \int_{-\pi}^{\pi} \rho(z, \phi) d\phi dz = 1$. This distribution is established by solving **Eqs. (1-2)** numerically for a population of 10^4 swimmers that are initially distributed at random positions $z \in [0, H]$ and orientations $\phi \in [-\pi, \pi]$. The simulation is continued until the distribution $\rho(z, \phi)$ reaches a steady state, which is after ~ 30 seconds. Subsequently, the spatial distribution is found by integrating over the angular coordinates, f(z) = $(2\pi)^{-1} \int_{-\pi}^{\pi} \rho(z, \phi) d\phi$, and similarly for the orientational distribution, $g(\phi) = H^{-1} \int_0^H \rho(z, \phi) dz$. The fraction of swimmers at the bottom wall and top interface are evaluated as $f_w = H^{-1} \int_0^{\epsilon} f(z) dz$ and $f_i = H^{-1} \int_{H-\epsilon}^H f(z) dz$, where $\epsilon = 1.1a$ to allow for small fluctuations.

Fig. 5 shows distributions of swimmer positions and orientations as a function of the background flow for *neu*tral swimmers ($\kappa = 0$), pushers ($\kappa > 0$) and pullers $(\kappa < 0)$, while subject to thermal diffusion. In the absence of a background flow, the swimmers accumulate at the two surfaces due to steric and hydrodynamic interactions, with a small bias towards the bottom wall for the pushers and pullers due to the stronger hydrodynamic dipolar interactions near no-slip surfaces (Fig. 5a; first window). Once trapped, the equilibrium orientation for pushers is parallel to both surfaces (**Fig. 5**b, orange peaks at $\phi = 0, \pm \pi$). Pullers are bound more strongly since their equilibrium orientation is perpendicular to the surface (blue peaks at $\pm \pi/2$). Indeed, Schaar *et al.*⁷⁵ have shown that the detention times of pullers in the absence of external flow can be several orders of magnitudes larger than those of pusher or source doublet swimmers.

Introduction of a small flow is enough to drive the majority of neutral swimmers towards the top surface (**Fig. 5**a; second window). The transition to free interface accumulation is a general trend for all swimmer types, but occurs at different flow speeds for each. The fraction of neutral swimmers at the bottom wall (f_w) drops rapidly to zero around $v_{\text{max}} \approx 0.5v_{\text{s}}$ and at the top interface the fraction (f_i) rises to unity (**Fig. 5**c; green circles). Pushers start to get detached from the bottom



FIG. 5. Prevention of boundary accumulation of micro-swimmers due to an external flow in a film. Statistics are shown for an ensemble of 10^4 swimmers, 30 seconds after release from a random position and orientation, when distributions have reached steady-state. (a) Distributions of swimmer position across the film for various background flow rates. (b) Distributions of swimmer orientation. (c) Fraction of swimmers at the bottom wall (f_w ; filled symbols) and top interface (f_i ; empty symbols). (d) Schematic of the observed dynamics. Colours indicate *neutral* swimmers (green circles), *pushers* (orange squares) and *pullers* (blue diamonds). Parameters used are film height $H = 10\mu$ m, swimmer dimensions $a = b = 0.5\mu$ m, swimming speed $v_s = 50\mu$ m/s, thermal diffusion coefficients $D = 0.25\mu$ m²/s and $D_r = 0.2$ rad²/s, dipole strength $\kappa = 30\mu$ m³/s and all other multipole coefficients are set to zero. Flow rates are normalised with respect to the swimming speed, v_s .

wall around $v_{\text{max}} \approx 2v_{\text{s}}$ (orange squares), which is a bit smaller than the value predicted in **§IV C** for deterministic swimmers (**Fig. 4**a; $v_{\text{max}} \approx 2.5v_{\text{s}}$ at $\kappa = 30\mu\text{m}^3/\text{s}$). This is because the noise occasionally kicks microbes away from the wall. By $v_{\text{max}} \approx 4v_{\text{s}}$ (**Fig. 5**a; third window), the majority of pushers have accumulated at the top interface. Pullers remain attached until a flow of strength $v_{\text{max}} \approx 8v_{\text{s}}$ is applied (blue diamonds), also a bit smaller than the deterministic equivalent $\approx 9v_{\text{s}}$. The distributions across the film show this in more detail (**Fig. 5**a,b; windows 2, 3 & 4 for the three swimmer types respectively).

Interestingly, by further increasing the flow speed, swimmers can again get trapped near the bottom wall. This is not due to hydrodynamic interactions but rather is directly related to the 'spinning' trajectories in highvorticity flows that were described in **§III**. Consequently, at high flow rates swimmers do not accumulate in the regions of smaller vorticity near the top interface. This is consistent with earlier findings of shear-trapping by Rusconi et al.⁴² It occurs because, whereas the vorticity alone is too small at the top surface to reinject swimmers into the film at a significant rate, small thermal fluctuations can lead swimmers to the high-vorticity region where they get trapped. Therefore, the number density of swimmers in the film is again larger near the bottom wall for flow rates $v_{\text{max}} > 8v_{\text{s}}$, and the accumulation at the top surface is suppressed entirely at large flow rates (**Fig. 5**c and **Fig. 5**a,b; last two windows).

In summary, the results show that in the absence of the background flow the swimmers accumulate at the top and bottom surfaces. Introducing flow and thermal Brownian noise can substantially modify distributions of swimming microbes in the film (**Fig. 5**d). For moderate flow rates swimmers are peeled off the bottom wall and accumulate at the top interface. However, for large flow rates the swimmers can get vorticity-trapped near the bottom wall again. This counter-intuitive result suggests that faster flow rates are not always the best strategy to discourage wall accumulation and the consequent initiation of biofilms. Instead, there is a finite range of intermediate flows within which the top-accumulation is optimised.

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FIG. 6. Prevention of boundary accumulation of micro-swimmers due to (a) run-and-tumble dynamics and (b) Gaussiandistributed enhanced Brownian noise, evaluated at the same rotational HI Péclet number. Statistics are shown for an ensemble of 10^4 swimmers, after 10 seconds of release from a random position and orientation, when the distributions have reached steadystate. Shown is the fraction of swimmers at the bottom wall (f_w ; solid lines) and top interface (f_i ; dotted lines). Colours indicate pushers (orange), pullers (blue) and neutral swimmers (green). Parameters used are film height $H = 100\mu$ m, swimmer dimensions $a = b = 0.5\mu$ m, swimming speed $v_s = 50\mu$ m/s, dipole strength $\kappa = 30\mu$ m³/s, all other multipole coefficients are set to zero and there is no background flow.

B. Prevention of boundary accumulation by run-tumble dynamics

Typical microbial swimmers do not rely solely on thermal noise to prevent boundary accumulation — the HI Péclet numbers (Eqs. 19-20) are much too large. Many micro-organisms have developed different mechanisms to actively change their orientation at a given moment. This could be an effective enhanced rotational Brownian noise, e.g. due to cilia which temporarily beat out of synchrony 57,58,76-78 or to inhomogeneous external influences.⁵⁹ Moreover, many microbial species make use of a 'tumbling' mechanism that allows them to suddenly change their orientation, rather than slowly decorrelating over time.⁷⁹ Various tumbling mechanisms have been observed, including ones based on flagellar unbundling^{80,81} and flagellar buckling instabilities⁸² for bacteria. Such strategies are not limited to flagellated bacteria but are also employed by ciliated organisms such as Paramecium that suddenly eject trichocysts.⁸³

In this section, we measure the extent to which such mechanisms cause boundary detachment by comparing swimmers with run-tumble dynamics (tumblers) to swimmers subject to a Gaussian-distributed enhanced Brownian noise (drifters) of the same effective rotational diffusion coefficient.

Fig. 6 shows the fraction of tumblers and drifters at the film surfaces for a given run time between tumble events, $\tau_{\rm r} \in [0.01, 100]$ s. This range corresponds to ${\rm P\acute{e}}_{\rm r}^{\rm HI} \in [0.1, 1000]$, since the effective rotational diffusion coefficient is $D_r^{\rm eff} = \pi^2/(2\tau_{\rm r})$. In agreement with the expectations, microbes of all swimmer types are seen to accumulate at the two surfaces at large HI Péclet numbers.

Tumblers can detach fairly easily from the surfaces when the HI Péclet number is reduced (**Fig. 6**a), because sudden decorrelating tumbling events are momentarily sufficient to overcome the hydrodynamic attraction. With increasing tumbling rate, there is a gradual crossover from surface accumulation to residence in the bulk of the film. Interestingly, we find that the hydrodynamic swimmer-type has little effect on this crossover. This is an important observation as it indicates that hydrodynamic interactions are not a dominant factor when run-and-tumble dynamics are present, as in *E. coli*.

Neutral drifters (swimmers subject to enhanced Brownian noise) can also escape from the surfaces with ease. However, drifters with hydrodynamic interactions remain attached to the surfaces at much lower HI Péclet numbers (**Fig. 6**b). Moreover, the escape crossover is much sharper. This difference arises because the slowly accumulating Brownian noise is continuously countered by hydrodynamic and steric interactions, and therefore a certain threshold must be exceeded before swimmers can escape from a surface. The level of critical noise is Pé_r^{HI*} \approx 8 for pusher drifters and \approx 2 for puller drifters. Also notice that the attraction towards the bottom wall is a factor of 3/2 larger for pushers,⁶⁵ and therefore the fraction of pushers at the bottom wall peaks before decaying with decreasing HI Péclet number.

This crossover from boundary accumulation to detachment is particularly relevant in many typical microenvironments. For *E. coli*-like with $\tau_{\rm r} \approx 0.9 {\rm s} \, ({\rm P\acute{e}}_{\rm r}^{\rm HI} \approx 8)$, the total fraction at both surfaces is $f_{\rm s} = f_{\rm w} + f_{\rm i} \approx 0.3$. However, for longer run times near surfaces, as observed in experiments,³⁶ $\tau_{\rm r} \approx 2 {\rm s} \, ({\rm P\acute{e}}_{\rm r}^{\rm HI} \approx 20)$, the fraction almost doubles to $f_{\rm s} \approx 0.55$.

In summary, the tumbling mechanism is more effective in preventing surface accumulation than Brownian noise of equivalent strength, particularly at the HI Péclet numbers relevant in nature. Flagellated organisms like *E. coli* bacteria could employ this to their advantage, using hydrodynamic interactions to get close to surfaces ($\S IV$) and then tumbling to escape them. We have demonstrated that hydrodynamic interactions with confining walls break the equivalence between active Brownian motion and run-tumble dynamics. While it is commonly true that active Brownian motion and run-tumble dynamics are equivalent this is not always the case.⁵³ Recent research that did not account for hydrodynamic interactions has shown that run-and-tumble dynamics lead to different density distributions near surfaces than simple active Brownian particles.⁵² Our results support this previous finding and further demonstrate that runtumble dynamics dominate over hydrodynamic interactions that can likely be safely neglected in future studies of bacteria.

VI. CONCLUSIONS AND DISCUSSION

We have presented a comprehensive description of the dynamics and statistics of swimming cells in flowing films, demonstrating that swimmer trajectories show distinct behaviours depending on the swimming mechanism of the organism. We focus on two classes of model organisms, *E. coli*-like and *Volvox*-like swimmers, with parameters tuned to experiments. Flagellated swimmers accumulate at surfaces due to hydrodynamic interactions, whereas ciliated organisms can avoid surfaces.

We have shown that swimmers can be detached from the bottom wall above a critical flow strength, which we predict analytically and obtain numerically for dipolar and higher-order hydrodynamic interactions. Conversely, we see that steric interactions with the surfaces and background film flow favours wall (interface) accumulation above (below) a critical flow speed, due to a vorticity-trapping mechanism. Therefore, boundary accumulation is not always prevented by imposing stronger flows. Instead, there is a finite range of intermediate flows for which microbes do not accumulate at the bottom wall. Our work predicts the extent of this range for both pusher and puller swimmers in terms of three dimensionless numbers (**Table I**). The flow-controlled crossover from surface accumulation to interface accumulation may have important implications for biofouling.

In addition, we demonstrate that run-and-tumble dynamics can act as a mechanism that organisms, such as $E.\ coli$, could employ to prevent boundary accumulation at surfaces, whereas enhanced Brownian noise requires much smaller Péclet numbers (greater noise) to achieve this goal. Whereas hydrodynamic interactions are important for systems with enhanced Brownian noise, we find that they have little impact on the swimmer distributions for the run-tumble dynamics expected for microorganisms such as $E.\ coli$. This is a conclusion has ramifications for future theoretical and computational investigations as it implies that computationally expensive and theoretically cumbersome hydrodynamic interactions may be neglected in biologically relevant scenarios if experimen-

Symbol	Definition	Name
λ	$a^3 v_{\rm max} / \kappa H$	Detachment number
Pé ^{HI}	$3a\kappa/8b^2D$	Translational HI Péclet number
$P\acute{e}_{r}^{HI}$	$3\kappa/16b^3D_{ m r}$	Rotational HI Péclet number

TABLE I. Summary of dimensionless number definitions.

tally significant run-tumble noise is properly accounted for.

Our results provide a number of testable predictions with implications for biofilm initiation. Genetic modification of *E. coli* can alter run-and-tumble dynamics.⁷⁹ *E. coli* modified in this way is predicted to have markedly different distributions within flowing films compared to unmutated samples. Likewise, different motile microbes, such as *Volvox*, are expected to reside at different points in the flowing film due to their different swimming strategies. The fraction of swimmers at the no-slip wall compared to the number at the no-shear interface could be measured. Measuring such fractions as a function of flow rate would provide direct experimental verification of the predictions made here. The rate of biofilm initiation at the solid surface is expected to correlate with the fraction of swimmers at the wall and so our work suggests a non-monotonic dependence of the initiation rate on the film flow velocity, with a maximum at moderate flow strengths.

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REFERENCES

- ¹J. Elgeti, R. G. Winkler, and G. Gompper, "Physics of microswimmerssingle particle motion and collective behavior: A review," Rep. Progr. Phys. **78**, 056601 (2015).
- ²R. M. Harshey, "Bacterial motility on a surface: Many ways to a common goal," Annu. Rev. Microbiol. 57, 249–273 (2003).
- ³K. Miki and D. E. Clapham, "Rheotaxis guides mammalian sperm," Curr. Biol. 23, 443 – 452 (2013).
- ⁴L. H. Cisneros, R. Cortez, C. Dombrowski, R. E. Goldstein, and J. O. Kessler, "Fluid dynamics of self-propelled microorganisms, from individuals to concentrated populations," Experim. Fluids 43, 737–753 (2007).
- ⁵I. S. Aranson, A. Sokolov, J. O. Kessler, and R. E. Goldstein, "Model for dynamical coherence in thin films of self-propelled microorganisms," Phys. Rev. E **75**, 040901 (2007).
- ⁶A. Doostmohammadi, R. Stocker, and A. M. Ardekani, "Low-Reynolds-number swimming at pycnoclines," Proc. Nat. Acad. Sci. **109**, 3856–3861 (2012).
- ⁷L. Hall-Stoodley, J. W. Costerton, and P. Stoodley, "Bacterial biofilms: From the natural environment to infectious diseases," Nat. Rev. Microbiol. 2, 95–108 (2004).
- ⁸J. C. Conrad, "Physics of bacterial near-surface motility using flagella and type IV pili: Implications for biofilm formation," Res. Microbiol. **163**, 619 629 (2012).

- ⁹M. A. Bees, P. Andresen, E. Mosekilde, and M. Givskov, "The interaction of thin-film flow, bacterial swarming and cell differentiation in colonies of *Serratia liquefaciens*," J. Math. Biol. 40, 27–63 (2000).
- ¹⁰J. Hill, O. Kalkanci, J. L. McMurry, and H. Koser, "Hydrodynamic surface interactions enable *Escherichia coli* to seek efficient routes to swim upstream," Phys. Rev. Lett. **98**, 068101 (2007).
- ¹¹B. Quiñones, G. Dulla, and S. E. Lindow, "Quorum sensing regulates exopolysaccharide production, motility, and virulence in *Pseudomonas syringae*," Mol. Plant-Microbe Interact. **18**, 682– 693 (2005).
- ¹²A. Karimi, D. Karig, A. Kumar, and A. Ardekani, "Interplay of physical mechanisms and biofilm processes: Review of microfluidic methods," Lab Chip **15**, 23–42 (2015).
- ¹³L. Vaccari, D. Allan, N. Sharifi-Mood, A. Singh, R. Leheny, and K. Stebe, "Films of bacteria at interfaces: Three stages of behaviour," Soft Matter **11**, 6062–6074 (2015).
- ¹⁴H. A. Stone and H. Masoud, "Mobility of membrane-trapped particles," J. Fluid Mech. **781**, 494–505 (2015).
- ¹⁵T. J. Pedley and J. O. Kessler, "The orientation of spheroidal microorganisms swimming in a flow field," Proc. Roy. Soc. Lond. B **231**, 47–70 (1987).
- ¹⁶E. Lauga, W. R. DiLuzio, G. M. Whitesides, and H. A. Stone, "Swimming in circles: Motion of bacteria near solid boundaries," Biophys. J. **90**, 400 – 412 (2006).
- ¹⁷A. P. Berke, L. Turner, H. C. Berg, and E. Lauga, "Hydrodynamic attraction of swimming microorganisms by surfaces," Phys. Rev. Lett. **101**, 038102 (2008).
- ¹⁸Y. Or and R. M. Murray, "Dynamics and stability of a class of low Reynolds number swimmers near a wall," Phys. Rev. E **79**, 045302 (2009).
- ¹⁹D. G. Crowdy and Y. Or, "Two-dimensional point singularity model of a low-Reynolds-number swimmer near a wall," Phys. Rev. E 81, 036313 (2010).
- ²⁰G. Li and J. X. Tang, "Accumulation of microswimmers near a surface mediated by collision and rotational Brownian motion," Phys. Rev. Lett. **103**, 078101 (2009).
- ²¹K. Drescher, K. C. Leptos, I. Tuval, T. Ishikawa, T. J. Pedley, and R. E. Goldstein, "Dancing Volvox: Hydrodynamic bound states of swimming algae," Phys. Rev. Lett. **102**, 168101 (2009).
- ²²J. Elgeti and G. Gompper, "Self-propelled rods near surfaces," EPL 85, 38002 (2009).
- ²³H. Shum, E. A. Gaffney, and D. J. Smith, "Modelling bacterial behaviour close to a no-slip plane boundary: The influence of bacterial geometry," Proc. Roy. Soc. Lond. A (2010), 10.1098/rspa.2009.0520.
- ²⁴G.-J. Li, A. Karimi, and A. Ardekani, "Effect of solid boundaries on swimming dynamics of microorganisms in a viscoelastic fluid," Rheologica Acta 53, 911–926 (2014).
- ²⁵S. E. Spagnolie and E. Lauga, "Hydrodynamics of self-propulsion near a boundary: Predictions and accuracy of far-field approximations," J. Fluid Mech. **700**, 105–147 (2012).
- ²⁶K. Martens, L. Angelani, R. Di Leonardo, and L. Bocquet, "Probability distributions for the run-and-tumble bacterial dynamics: An analogy to the Lorentz model," Eur. Phys. J. E 35, 84 (2012).
- ²⁷J. Elgeti, U. B. Kaupp, and G. Gompper, "Hydrodynamics of sperm cells near surfaces," Biophys. J. **99**, 1018 – 1026 (2010).
- ²⁸A. Costanzo, J. Elgeti, T. Auth, G. Gompper, and M. Ripoll, "Motility-sorting of self-propelled particles in microchannels," EPL **107**, 36003 (2014).
- ²⁹P. D. Frymier, R. M. Ford, H. C. Berg, and P. T. Cummings, "Three-dimensional tracking of motile bacteria near a solid planar surface," Proc. Nat. Acad. Sci. **92**, 6195–6199 (1995).
- ³⁰A. P. Berke, L. Turner, H. C. Berg, and E. Lauga, "Hydrodynamic attraction of swimming microorganisms by surfaces," Phys. Rev. Lett. **101**, 038102 (2008).
- ³¹G. Li and J. X. Tang, "Accumulation of microswimmers near a surface mediated by collision and rotational brownian motion,"

Phys. Rev. Lett. 103, 078101 (2009).

- ³²G. Li, J. Bensson, L. Nisimova, D. Munger, P. Mahautmr, J. X. Tang, M. R. Maxey, and Y. V. Brun, "Accumulation of swimming bacteria near a solid surface," Phys. Rev. E 84, 041932 (2011).
- ³³J. Leach, H. Mushfique, S. Keen, R. Di Leonardo, G. Ruocco, J. M. Cooper, and M. J. Padgett, "Comparison of Faxén's correction for a microsphere translating or rotating near a surface," Phys. Rev. E **79**, 026301 (2009).
- ³⁴K. Drescher, J. Dunkel, L. H. Cisneros, S. Ganguly, and R. E. Goldstein, "Fluid dynamics and noise in bacterial cell-cell and cell-surface scattering," Proc. Nat. Acad. Sci. **108**, 10940–10945 (2011).
- ³⁵V. Kantsler, J. Dunkel, M. Polin, and R. E. Goldstein, "Ciliary contact interactions dominate surface scattering of swimming eukaryotes," Proc. Nat. Acad. Sci. **110**, 1187–1192 (2013).
- ³⁶M. Molaei, M. Barry, R. Stocker, and J. Sheng, "Failed escape: Solid surfaces prevent tumbling of *Escherichia coli*," Phys. Rev. Lett. **113**, 068103 (2014).
- ³⁷R. A. Lambert, F. Picano, W.-P. Breugem, and L. Brandt, "Active suspensions in thin films: Nutrient uptake and swimmer motion," J. Fluid Mech. **733**, 528–557 (2013).
- ³⁸A. Costanzo, R. Di Leonardo, G. Ruocco, and L. Angelani, "Transport of self-propelling bacteria in micro-channel flow," J. Phys.: Condens. Matter 24, 065101 (2012).
- ³⁹A. Zöttl and H. Stark, "Nonlinear dynamics of a microswimmer in Poiseuille flow," Phys. Rev. Lett. **108**, 218104 (2012).
- ⁴⁰E. Altshuler, G. Mino, C. Perez-Penichet, L. d. Rio, A. Lindner, A. Rousselet, and E. Clément, "Flow-controlled densification and anomalous dispersion of *E. coli* through a constriction," Soft Matter **9**, 1864–1870 (2013).
- ⁴¹A. Zöttl and H. Stark, "Periodic and quasiperiodic motion of an elongated microswimmer in Poiseuille flow," Eur. Phys. J. E 36, 4 (2013).
- ⁴²R. Rusconi, J. S. Guasto, and R. Stocker, "Bacterial transport suppressed by fluid shear," Nat. Phys. **10**, 212–217 (2014).
- ⁴³H. Masoud and H. A. Stone, "A reciprocal theorem for marangoni propulsion," J. Fluid Mech. **741**, R4 (2014).
- ⁴⁴V. Kantsler, J. Dunkel, M. Blayney, and R. E. Goldstein, "Rheotaxis facilitates upstream navigation of mammalian sperm cells," eLife **3**, e02403 (2014).
- ⁴⁵N. Figueroa-Morales, G. Miño, A. Rivera, R. Caballero, E. Clément, E. Altshuler, and A. Lindner, "Living on the edge: Transfer and traffic of *E. coli* in a confined flow," Soft Matter **11**, 6284–6293 (2015).
- ⁴⁶H. Masoud, H. A. Stone, and M. J. Shelley, "On the rotation of porous ellipsoids in simple shear flows," J. Fluid Mech. **733**, R6 (2013).
- ⁴⁷A. J. T. M. Mathijssen, T. N. Shendruk, J. M. Yeomans, and A. Doostmohammadi, "Upstream swimming in microbiological flows," arXiv:1507.00962 preprint (2015).
- ⁴⁸J. Gachelin, G. Miño, H. Berthet, A. Lindner, A. Rousselet, and É. Clément, "Non-Newtonian viscosity of *Escherichia coli* suspensions," Phys. Rev. Lett. **110**, 268103 (2013).
- ⁴⁹A. Karimi, S. Yazdi, and A. M. Ardekani, "Hydrodynamic mechanisms of cell and particle trapping in microfluidics," Biomicrofluidics 7, 021501 (2013).
- ⁵⁰H. M. López, J. Gachelin, C. Douarche, H. Auradou, and E. Clément, "Turning bacteria suspensions into superfluids," Phys. Rev. Lett. **115**, 028301 (2015).
- ⁵¹A. Ardekani and E. Gore, "Emergence of a limit cycle for swimming microorganisms in a vortical flow of a viscoelastic fluid," Phys. Rev. E 85, 056309 (2012).
- ⁵²J. Elgeti and G. Gompper, "Run-and-tumble dynamics of selfpropelled particles in confinement," EPL **109**, 58003 (2015).
- ⁵³M. E. Cates and J. Tailleur, "When are active Brownian particles and run-and-tumble particles equivalent? Consequences for motility-induced phase separation," EPL **101**, 20010 (2013).
- ⁵⁴J. Saragosti, V. Calvez, N. Bournaveas, B. Perthame, A. Buguin, and P. Silberzan, "Directional persistence of chemotactic bacteria

in a traveling concentration wave," Proc. Nat. Acad. Sci. 108, 16235–16240 (2011).

- ⁵⁵J. Saragosti, P. Silberzan, and A. Buguin, "Modeling *E. coli* tumbles by rotational diffusion. Implications for chemotaxis," PloS one 7, e35412 (2012).
- ⁵⁶S. Tavaddod, M. Charsooghi, F. Abdi, H. Khalesifard, and R. Golestanian, "Probing passive diffusion of flagellated and deflagellated *Escherichia coli*," Eur. Phys. J. E **34**, 16 (2011).
- ⁵⁷K. Y. Wan and R. E. Goldstein, "Rhythmicity, recurrence, and recovery of flagellar beating," Phys. Rev. Lett. **113**, 238103 (2014).
- ⁵⁸R. Ma, G. S. Klindt, I. H. Riedel-Kruse, F. Jülicher, and B. M. Friedrich, "Active phase and amplitude fluctuations of flagellar beating," Phys. Rev. Lett. **113**, 048101 (2014).
- ⁵⁹M. Jabbarzadeh, Y. Hyon, and H. C. Fu, "Swimming fluctuations of micro-organisms due to heterogeneous microstructure," Phys. Rev. E **90**, 043021 (2014).
- ⁶⁰J. Elgeti and G. Gompper, "Wall accumulation of self-propelled spheres," Europhys. Lett. **101**, 48003 (2013).
- ⁶¹B. Ezhilan, R. Alonso-Matilla, and D. Saintillan, "On the distribution and swim pressure of run-and-tumble particles in confinement," J. Fluid Mech. **781**, R4 (2015).
- ⁶²S. J. Weinstein and K. J. Ruschak, "Coating flows," Annu. Rev. Fluid Mech. **36**, 29–53 (2004).
- ⁶³F. Peruani, A. Deutsch, and M. Bär, "A mean-field theory for self-propelled particles interacting by velocity alignment mechanisms," Eur. Phys. J. Spec. Top. **157**, 111–122 (2008).
- ⁶⁴A. P. Berke, L. Turner, H. C. Berg, and E. Lauga, "Hydrodynamic attraction of swimming microorganisms by surfaces," Phys. Rev. Lett. **101**, 038102 (2008).
- ⁶⁵A. J. T. M. Mathijssen, A. Doostmohammadi, J. M. Yeomans, and T. N. Shendruk, "Hydrodynamics of microswimmers in films," Under review (2015).
- ⁶⁶J. R. Blake, "A note on the image system for a Stokeslet in a no-slip boundary," Math. Proc. Camb. Phil. Soc **70**, 303 – 310 (1971).
- ⁶⁷A. J. T. M. Mathijssen, D. O. Pushkin, and J. M. Yeomans, "Tracer trajectories and displacement due to a micro-swimmer near a surface," J. Fluid Mech. **773**, 498—519 (2015).
- ⁶⁸J. Lighthill, "Flagellar hydrodynamics," SIAM Rev. 18, 161–230 (1976).
- ⁶⁹G. Lowe, M. Meister, and H. C. Berg, "Rapid rotation of flagellar bundles in swimming bacteria," Nature **325**, 637–640 (1987).

- ⁷⁰H. C. Berg and L. Turner, "Chemotaxis of bacteria in glass capillary arrays. *Escherichia coli*, motility, microchannel plate, and light scattering." Biophys. J. 58, 919 (1990).
- ⁷¹E. Lauga, W. R. DiLuzio, G. M. Whitesides, and H. A. Stone, "Swimming in circles: Motion of bacteria near solid boundaries," Biophys. J. **90**, 400–412 (2006).
- ⁷²R. Di Leonardo, D. Dell'Arciprete, L. Angelani, and V. Iebba, "Swimming with an image," Phys. Rev. Lett. **106**, 038101 (2011).
- ⁷³A. S. Utada, R. R. Bennett, J. C. Fong, M. L. Gibiansky, F. H. Yildiz, R. Golestanian, and G. C. Wong, "Vibrio cholerae use pili and flagella synergistically to effect motility switching and conditional surface attachment," Nat. Comm. 5 (2014), 10.1038/ncomms5913.
- ⁷⁴K. Drescher, R. E. Goldstein, N. Michel, M. Polin, and I. Tuval, "Direct measurement of the flow field around swimming microorganisms," Phys. Rev. Lett. **105** (2010), 10.1103/Phys-RevLett.105.168101.
- ⁷⁵K. Schaar, A. Zöttl, and H. Stark, "Detention times of microswimmers close to surfaces: Influence of hydrodynamic interactions and noise," arXiv:1412.6435 preprint (2014).
- ⁷⁶K. A. Wemmer and W. F. Marshall, "Flagellar motility: All pull together," Curr. Biol. **14**, R992–R993 (2004).
- ⁷⁷V. F. Geyer, F. Jülicher, J. Howard, and B. M. Friedrich, "Cellbody rocking is a dominant mechanism for flagellar synchronization in a swimming alga," Proc. Nat. Acad. Sci. **110**, 18058– 18063 (2013).
- ⁷⁸K. Y. Wan, K. C. Leptos, and R. E. Goldstein, "Lag, lock, sync, slip: the many 'phases' of coupled flagella," J. R. Soc. Interface **11**, 20131160 (2014).
- ⁷⁹H. C. Berg and D. A. Brown, "Chemotaxis in *Escherichia coli* analysed by three-dimensional tracking," Nature **239**, 500–504 (1972).
- ⁸⁰R. M. Macnab, "Bacterial flagella rotating in bundles: A study in helical geometry," Proc. Nat. Acad. Sci. **74**, 221–225 (1977).
- ⁸¹L. Turner, W. S. Ryu, and H. C. Berg, "Real-time imaging of fluorescent flagellar filaments," J. Bacteriology **182**, 2793–2801 (2000).
- ⁸²K. Son, J. S. Guasto, and R. Stocker, "Bacteria can exploit a flagellar buckling instability to change direction," Nat. Phys. 9, 494–498 (2013).
- ⁸³A. Hamel, C. Fisch, L. Combettes, P. Dupuis-Williams, and C. N. Baroud, "Transitions between three swimming gaits in *Paramecium* escape," Proc. Nat. Acad. Sci. **108**, 7290–7295 (2011).