Hydrodynamic interaction of microswimmers near a wall

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The hydrodynamics of an archetypal low-Reynolds number swimmer, called “squirmer,” near a wall has been numerically studied. For a single squirmer, depending on the swimming mechanism, three different modes are distinguished: (a) the squirmer escaping from the wall, (b) the squirmer swimming along the wall at a constant distance and orientation angle, and (c) the squirmer swimming near the wall in a periodic trajectory. The role of inertial effects on the near-wall motion of the squirmer is quantified. The dynamics of multiple squirmers swimming between two walls is found to be very different from a single squirmer. Near-wall accumulation of squirmers is observed. At a relatively small concentration $c = 0.1$, around 60–80\% of the squirmers are accumulated near the walls and attraction of pushers and pullers toward the wall is stronger than neutral squirmers. Near-wall squirmers orient normal to the wall, while in the bulk region, the squirmers are mostly oriented parallel to the wall. At a high concentration $c = 0.4$, the percentage of the near-wall squirmers is around 40\%. The orientation angle of squirmers in the bulk region is more uniformly distributed at high concentrations. In the near-wall region, pushers repel each other, while pullers are attracted to each other and form clusters.

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I. INTRODUCTION

Organisms swimming at a low-Reynolds-number regime near a no-slip wall is a subject of growing interest in recent years for its importance in many health and environmental problems [1,2]. For example, accumulation of bacteria near the surface and the formation of biofilms are closely related to many types of microbial infections [3]. Spermatozoa swimming in the female reproductive tract is another common example [4]. Studies have demonstrated that the presence of boundaries significantly affects the swimming speed and energetic properties of microswimmers [5–7]. Furthermore, recent experimental studies have observed many unexpected interesting behaviors for swimming organisms near a boundary, including the circular orbit of \textit{Escherichia coli} above a flat surface [8], wall-induced accumulation of organisms [9–12] and the “dancing” motion of pairs of \textit{Volvox} near a solid boundary [13].

The inertial effects are important for many planktomic swimmers in marine environments, such as larvae and 	extit{Pleurobrachia}. Small organisms use inertia to change their swimming direction, attack a prey, or escape from a predator [14,15]. Wang and Ardekani [16,17] analytically quantified the inertial effects by deriving the fundamental equation of motion for small organisms swimming in an unbounded quiescent fluid environment. The role of boundaries on the swimming of small organisms in a nonzero-Reynolds-number regime is of great importance. For example, the accumulation of larvae near the surface has a significant influence on the metamorphosis and survival of larvae [18].

Simplified swimming models have been widely used to investigate the dynamics of confined swimmers in a zero-Reynolds-number regime. The work of Zargar et al. [19] analyzed the influence of a wall on the dynamics of a three-sphere swimmer in the close proximity of a boundary. It is shown that the swimmer can swim toward or escape from the wall, depending on the initial angle and distance from the wall. The work of Or et al. [20] studied the dynamics of swimmers formed by small networks of rotating spheres attached by rigid rods. In their simulation, mobility matrices for multiple interacting spherical particles in the presence of a flat plane were approximately calculated by accounting for the far-field hydrodynamic interactions. Their results show that the swimmer may have a marginally stable motion or a periodic motion depending on the arrangement of the spheres. Similar swimming dynamics was reported in the work of Crowdy and Or [21] for a two-dimensional treadmilling swimmer model near a no-slip wall. The treadmilling swimmer was also utilized to study the swimming dynamics near a wall with a gap. Stable equilibrium points for the swimmer in the gap region, as well as Hopf bifurcations and periodic bound states were observed [22]. The work of Dunstan et al. [23] modeled a bacterial cell by using two spheres of different radii at a constant distance. The effect of the flagella was modeled by imposing a force on the tail sphere as well as adding equal and opposite torques on the two spheres. Their results show three different swimming behaviors depending on the initial condition: swimming in circles in contact with the wall, swimming in circles at a finite distance from the wall, and swimming away from the wall. Spagnolie and Lauga [24] provided a detailed comparison of the hydrodynamics of microswimmers near a boundary using far-field approximations and numerical solutions of the Stokes equation. They showed that the far-field approximation is surprisingly accurate for spherical swimmers when the distance between the center of the swimmer and the wall is larger than its diameter. However, the fundamental singularities of the Stokes equation cannot accurately account for the near-field effects when the swimmer comes close to the boundary. In this work, Navier-Stokes equations are directly solved to capture the hydrodynamic interaction of a single swimmer and a suspension of swimmers near a wall in a small, but nonzero, Reynolds-number regime.

One of the widely used swimmer models called “squirmer” is introduced by Lighthill [25] and Blake [26] to model the
ciliated microorganisms such as Volvox and Paramecium. Other studies have extended the model to general shape twodimensional [27], porous prolate-spheroidal [28], and smallamplitude deformable squirmers [29]. Here, we consider a spherical squirmer with a tangential surface deformation given as

\[ u_t = \left( e \cdot r \middle| r \mid - e \right) \sum_{n \geq 1} \frac{2}{n(n+1)} B_n P_n \left( e \cdot r \middle| r \mid \right), \] (1)

where \( e \) is the orientation vector of the squirmer, \( B_n \) is the \( n \)th mode of the tangential squirming velocity [26], \( P_n \) is the \( n \)th Legendre polynomial, \( P_n^\prime \) is its derivative, and \( r \) is the position vector. Similar to most previous studies [30–37], we only consider the first two squirming modes and consequently the magnitude of the tangential velocity is written as

\[ u_t^\prime(\theta) = B_1 \sin \theta + B_2 \sin \theta \cos \theta, \] (2)

where \( \theta \) is the polar angle measured from the swimming direction, and \( B_1 \) and \( B_2 \) are the first two squirming modes. The parameter \( \beta = B_2/B_1 \) distinguishes three kinds of swimmers: \( \beta > 0 \), corresponds to a puller generating thrust in front of the body such as Chlamydomonas; \( \beta < 0 \), corresponds to a pusher generating thrust behind the body such as most bacteria; \( \beta = 0 \) corresponds to a neutral squirmer, generating a symmetric flow field, such as Volvox. In the Stokes regime, the swimming speed of a squirmer in an unbounded domain is \( U_0 = 2B_1/3 \). The squirmer model has been widely used to study the dynamics of a single microswimmer, nutrient uptake [30,38], biomixing [39], unsteady propulsion of small organisms [17], as well as the effects of density stratification on the self-propulsion [31]. Recently, Zhu et al. [32] studied the locomotion of a squirmer in a capillary tube and found that a neutral squirmer generally follows a helical trajectory; a puller displays a stable locomotion along the tube, while a pusher crashes into the wall. Ishikawa, Pedley, and coworkers have studied the dynamics of multiple squirmers in an unbounded fluid in the Stokes regime, including the hydrodynamic interaction between two squirmers [33], rheology and diffusion of a suspension of squirmers [34,35], collective behavior [36], and vertical dispersion of squirmers in a shear flow [37].

Microorganisms are usually observed in high concentrations, where the interaction between microorganisms are important and they often exhibit collective behaviors resulting in swarms and vortices of large scales [40]. Previous experiments show that a suspension of microorganisms results in the variation of fluid viscosity [41], superdiffusion in short times [42], and enhanced mixing [41]. Hydrodynamic interaction of spherical particles near a wall or between two walls in a zero-Reynolds-number regime has been extensively studied using multipole expansion method and Stokesian dynamics [43–45]. The algorithm developed by Bhattacharya et al. accurately calculates the many-particle friction matrix by using spherical and Cartesian representation of Stokes flow to capture the interaction of the fluid with the particles and walls, respectively [45]. This method can be successfully applied to evaluate the friction matrix for a single sphere, a pair of spheres, and linear chains of spheres between two walls [46]. Based on the dumbbell swimmer model of two beads connected by a rigid rod, Underhill et al. [47] studied the diffusion and spatial correlation of swimmers as well as the correlations of stress and velocity [48]. Using the same model, Hernández-Ortiz et al. [49,50] studied the dynamics of suspension of swimmers between two walls and found that the swimmers aggregate near the walls at low concentrations, while at high concentrations, this distribution is disrupted by large-scale coherent motions. The diffusion and spatial correlation of swimmers are also affected by the confined geometry. Their model treated the swimmers as point-force dipoles and the swimmer-swimmer and swimmer-wall interactions are approximated using an excluded volume force. Wensink and Löwen [51] studied the suspension of self-propelled colloidal rods in a channel using two-dimensional Brownian dynamics simulation. Their results show that the aggregated self-propelled rods near the channel wall form “hedgehog”-shaped clusters with most of the rods pointing toward the wall.

In this paper, we investigate the dynamics of a single and a collection of squirmers(s) near a no-slip wall in a low-Reynolds-number regime. By directly solving the Navier-Stokes equation, the inertial effects are included and the motion of small swimming organisms near a solid surface is accurately captured.

\section{II. Numerical Method}

The distributed Lagrange-multiplier-based finite-volume method is used in this study and details of the method can be found in Refs. [52,53]. The Navier-Stokes equations for an incompressible Newtonian fluid are solved in the entire computational domain

\begin{align}
\rho \frac{Du}{Dt} &= \nabla \cdot (-p\mathbf{I} + \tau) + \mathbf{f}, \quad (3a) \\
\nabla \cdot \mathbf{u} &= 0, \quad (3b)
\end{align} \]

where \( \mathbf{u} \) is the velocity, \( \rho \) is the pressure, \( \mathbf{I} \) is the identity tensor, \( \tau = \mu(\nabla \mathbf{u} + (\nabla \mathbf{u})^T) \) is the viscous stress, and \( \mu \) is the viscosity. The density \( \rho \) is equal to the particle density \( \rho_p \) inside the squirmer and equal to the fluid density \( \rho_f \) in the fluid domain. In this study, we set \( \rho_p = \rho_f \), since the density of microorganisms is usually close to the background fluid. A rigid particle and a self-propelled particle can be modeled by adding a source term to the Navier-Stokes equation. The forcing term in each iteration is calculated as

\[ f = f^* + \alpha \frac{\phi \rho}{\Delta t} (\mathbf{U} + \omega_p \times \mathbf{r} + \mathbf{u}_i - \mathbf{u}), \] (4)

where \( \mathbf{u} \) is the translational velocity of the particle (inert or self-propelled), \( \omega_p \) is the particle angular velocity, \( \mathbf{u}_i \) is the imposed velocity causing the self-propulsion, \( f^* \) is the force calculated in the previous iteration, \( \alpha \) is a dimensionless factor whose value affects the convergence rate but not the solution, \( \phi \) is the volume fraction occupied by the particle in each computational cell (\( \phi = 1 \) inside, \( \phi = 0 \) outside, and \( 0 < \phi < 1 \) for the cells at the surface of the particle). The velocity field \( \mathbf{u} \) is obtained by solving Eq. (3). To recover the tangential velocity \( u_t^\prime \) on the surface of the squirmer [given in Eq. (2)], we impose the
following solenoidal velocity \( \mathbf{u}_i \) inside the squirmer,
\[
\mathbf{u}_i = \left[ \left( \frac{r}{\alpha} \right)^m - \left( \frac{r}{\alpha} \right)^{m+1} \right] \left( u_0^i \cot \theta + \frac{d u_0}{d \theta} \right) \mathbf{e}_r,
\]
\[
+ \left[ (m + 3) \left( \frac{r}{\alpha} \right)^{m+1} - (m + 2) \left( \frac{r}{\alpha} \right)^{m} \right] u_0^i \mathbf{e}_\theta,
\]
where \( \alpha \) is the radius of the squirmer, \( \mathbf{e}_r \) and \( \mathbf{e}_\theta \) are the unit vectors along \( r \) and \( \theta \) directions, \( m \) is an arbitrary positive integer, where the simulation results are independent of its value. It should be noted that \( \mathbf{u}_i \) is zero for an inert particle. The particle translational and angular velocities are calculated as
\[
U = \frac{1}{M_p} \int_{V_p} \rho_p (\mathbf{u} - \mathbf{u}_i) \, dV, \quad (6)
\]
\[
I_p \omega_p = \int_{V_p} \rho_p \mathbf{r} \times (\mathbf{u} - \mathbf{u}_i) \, dV, \quad (7)
\]
where \( V_p, M_p, \) and \( I_p \) are the volume, mass, and moment of inertia of the particle, respectively. The imposed velocity leads to a zero translational and rotational velocity for the particle (i.e., \( \int_{V_p} \mathbf{u}_i \, dV = 0 \) and \( \int_{V_p} \mathbf{r} \times \mathbf{u}_i \, dV = 0 \)). Iterations are repeated until the maximum of Euclidean norm of \( (f - f^*)/f \) and the normalized residual falls below the specified tolerance of \( 10^{-3} \). The normalized residual is defined as
\[
\int_{V_p} \frac{|U + \omega_p \times \mathbf{r} + \mathbf{u}_i - \mathbf{u}|}{U_0 V_p} \, dV. \quad (8)
\]
The velocity field inside the particle for the converged solution is \( \mathbf{u} = \mathbf{u} + \omega_p \times \mathbf{r} + \mathbf{u}_i \). It is straightforward to demonstrate that the converged solution is the equivalent of the particle equation of motion:
\[
M_p \frac{d^2 \mathbf{x}_p}{dt^2} = \int_{V_p} (-p \mathbf{1} + \mathbf{r}) \cdot n \, dS, \quad (9)
\]
where \( dS \) is the surface differential element. Combining Eqs. (4)–(7), one can show that \( \int_{V_p} f \, dV = 0 \). Integrating Eq. (3a) over the particle volume and using the Reynolds transport theorem leads to
\[
\frac{d}{dt} \int_{V_p} \rho (U + \omega_p \times \mathbf{r} + \mathbf{u}_i) \, dV = \int_{V_p} \nabla \cdot (-p \mathbf{1} + \mathbf{r}) \, dV, \quad (10)
\]
which is equivalent to Eq. (9). This method has been extensively used for the motion of inert particles in viscous fluids and verified in our previous publications [53–57].

Simulations are conducted using a finite-volume method on a fixed staggered grid. A conventional operator-splitting method is applied to enforce the continuity equation. The second-order total variation diminishing (TVD) Runge-Kutta method is used for time marching. The spatial derivatives in the convection term are evaluated using the quadratic upstream interpolation for convective kinetics (QUICK) scheme and the diffusion terms are discretized using the central difference scheme. The results are normalized by the characteristic length \( a \), velocity \( U_0 = 2B_1/3 \), and time \( a/U_0 \).

We first validate the motion of a single squirmer in an unbounded fluid at a finite Reynolds number defined as \( Re = \rho U_0 a/\mu \). The comparison between the present numerical results for the steady swimming speed and the analytical results, obtained using perturbation theory [17,58], is illustrated in Fig. 1. The first-order solution for the swimming speed of a squirmer in an unbounded domain is \( U \approx 1 - 0.15 \beta Re \) [17]. The swimming speed \( U \) of the neutral squirmer does not change with the Reynolds number. The swimming speed of a pusher increases with Re and our results are in good agreement with the analytical results at small values of Reynolds number. The swimming speed of a puller decreases with Re. At \( Re > 0.2 \), the first-order solution of Wang and Ardekani [17] and the second-order solution of Khair and Chisholm [58] starts to fail. Our results agree with Eq. (41) of Ref. [58].

In this study, we focus on the motion of a single squirmer near a wall and a suspension of squirmers between two walls. The schematic of a single squirmer near a wall is shown in Fig. 2. The distance between the center of the squirmer and the wall is \( h \) and the orientation angle of the squirmer from the horizontal direction is \( \alpha \). The wall is located at \( y = 0 \) and the squirmer moves in the \( x-y \) plane. For the cases of multiple squirmers, there are two walls at \( y = 0 \) and \( y = L \) and \( \alpha \) corresponds to the orientation of the squirmer relative to the nearest wall.

![FIG. 1. (Color online) Comparison of the results for the steady swimming of a squirmer in an unbounded domain at finite Re. Symbols, present numerical results; dashed lines, Eq. (39) from Ref. [17] (first-order); dashdot lines, Eq. (40) of Ref. [58] (second-order); solid lines, Eq. (41) of Ref. [58]. The domain size for the computational results is 40 × 40 × 40.](image1)

![FIG. 2. Schematic of a squirmer near a wall.](image2)
When the squirmer approaches the wall or another squirmer, the high pressure in the thin film between the squirmers and the wall prevents any unphysical overlaps. However, a very small grid resolution is needed to properly capture this dynamic process and consequently it is computationally expensive. As mentioned in Ref. [24], a short-range repulsive force [59] is necessary to prevent squirmer-wall or squirmer-squirmer overlaps. The repulsive force during the squirmer-wall or squirmer-squirmer collision is defined as

\[ f_r = C_m \frac{d - d_{min} - dr}{\varepsilon} e_r, \]

(11)

where \( C_m = \frac{M_g U_0^2}{a} \) is the characteristic force, \( \varepsilon = 10^{-4} \) is a small positive number, \( d \) is the distance between the center of the squirmer and the wall or the distance between two squirmers, \( d_{min} = a \) or \( 2a \) is the corresponding minimum possible distance, \( dr \) is the force range and is usually set to be the smallest grid size \( \Delta \) in the computational domain [59]. The direction of the repulsive force \( e_r \) is normal to the wall or along the line of center of the two squirmers. The above-mentioned repulsive force has been widely used to handle the collision between particles and walls [57,59]. Besides the formula used here, another form of repulsive force based on soft-sphere collision model is used in the numerical simulation of nutrient uptake of a suspension of squirmers in a thin film [30].

To check the convergence of the results, simulations of the near-wall motion of a neutral squirmer and a puller are conducted for different values of mesh size, time step, and magnitude of repulsive force. The size of computational domain is \( 64 \times 25.6 \times 12.8 \). The no-slip boundary condition is used on the wall and the far-field boundary condition is used at other boundaries of the domain. The squirmer is initially located at \( h_0 = 2, \alpha_0 = -\pi/4 \), and \( Re = 1 \). As shown in Fig. 3, the computed results are independent of the grid size, time step, and magnitude of the repulsive force. Consequently, for the calculations presented in this work, 20 grid points are used across the squirmer diameter and the time step is set to \( \Delta t = 1 \times 10^{-3} \).

III. RESULTS AND DISCUSSION

A. Swimming dynamics of a single squirmer near a wall

Figure 4 shows the trajectory as well as the time history of orientation angle \( \alpha \) and swimming speed \( U \) of the squirmer for different values of \( \beta \). The squirmer collides with the wall at \( t \approx 0.8 \), then it swims along the wall for a certain period of time referred to as the “contact time” and finally swims away from the wall. The contact time decreases as \( \beta \) increases. When squirmers detached from the wall, three different swimming modes are observed depending on the values of \( \beta \): (1) Squirmers of \( \beta \leq 1 \) will swim away from the wall with a positive angle. (2) Squirmers of \( 2 \leq \beta \leq 5 \) oscillate near the wall and eventually swim along the wall with a constant distance and a negative angle. It should be noted that the damping ratio for the cases of \( \beta = 4 \) and 5 is small and their oscillations are not fully damped during the simulation time. (3) Squirmers of \( \beta \geq 7 \) will swim in a cyclic motion, bouncing on the wall. The amplitude of the cyclic motion is small compared to the wavelength. Similar steady and periodic swimming modes along the wall have been observed in Ref. [20] using a three-sphere swimmer model. For the squirmers swimming away from the wall, the swimming speed is recovered to its value for a squirmer in an unbounded domain. For \( \beta \leq 1 \), the squirmers escape from the wall and their steady swimming speed decreases as \( \beta \) increases, which is consistent with analytical results of inertial squirmer in an unbounded domain [17]. On the other hand, pullers of \( \beta > 2 \) are trapped near the wall and their swimming speed increases with \( \beta \).

The effects of the initial angle and the initial height are also studied. The effects of the wall on the squirmer weakens for \( \alpha_0 > 0 \), since the squirmer swims away from the wall. Therefore, we mainly focus on the cases of \( \alpha_0 \leq 0 \). For the neutral squirmers at \( h_0 = 2 \) and \( Re = 1 \) in Fig. 5, the trajectories are affected by the initial angle while the swimming mode remains the same. For angles \( \alpha_0 \geq -\pi/12 \), the squirmer rotates away from the wall before getting to a close contact with the wall, whereas for initial angles \( \alpha_0 \leq -\pi/6 \), the squirmer collides with the wall and rotates away from the wall in a near-wall contact region. The squirmer eventually escapes the wall with a same final angle of \( \alpha_f \approx 0.4 \). In the work of Ref. [24], final angle \( \alpha_f \approx 0.4 \) and 0.35 were derived from
FIG. 4. (Color online) (a) Trajectory and (b) temporal evolution of distance away from the wall $y$, (c) orientation angle $\alpha$ and (d) swimming speed $U$ for squirmers of various $\beta$ at $Re = 1$, initially located at $h_0 = 2$ and $\alpha_0 = -\pi/4$.

The effects of the initial height of the squirmers are plotted in Fig. 7; the initial height affects the collision time between the squirmers and the wall but not the swimming mode.

In Fig. 8, we compare the trajectories of squirmers of $\beta = -3, 0, 3,$ and $7$ at $\alpha_0 = -\pi/4$ and $h_0 = 2$ for two different values of Reynolds number ($Re = 0.1$ and 1). The initial contact time decreases with Reynolds number. The pusher and neutral swimmer escape from the wall for both values of Reynolds number, whereas puller of $\beta = 3$ escapes from the wall at $Re = 0.1$ and is entrapped near the wall at $Re = 1$. At larger values of $\beta$, the puller is entrapped for both values of Reynolds number, but the one at $Re = 1$ has a larger bouncing frequency. In summary, the inertial effect decreases the initial wall contact time independent of the squirming type, but it leads to a stronger attraction toward the wall for the puller.

B. Swimming dynamics of multiple squirmers between two walls

We examine the swimming dynamics of a suspension of squirmers between two walls. The simulation is conducted in a cubic domain of $[0, 13.89] \times [0, 13.89] \times [0, 13.89]$ with two no-slip walls at $y = 0$ and $y = 13.89$. Periodic conditions are used along $x$ and $z$ directions. Without any overlap,
squirmers are initially randomly placed in a fluid otherwise at rest and their orientations are also randomly initialized. Three cases of $\beta = 0$, 3, and −3 at volume concentration $c = (4\pi N)/(3L^3) = 0.1$ are simulated. The case of $\beta = 0$ at $c = 0.4$ is also studied to consider the role of concentration. $N = 64$ and 256 squirmers are modeled for the two concentrations, respectively. Across the diameter of the squirmer, there are around 20 grid points for cases of $c = 0.1$ and 10 grids for $c = 0.4$, the time step is $\Delta t = 1 \times 10^{-3}$. The Reynolds number is $Re = 1$ for all cases.

Figure 9 shows the spatial distribution of squirmers at $t = 100$ at which the system has reached a statistical steady state. White and black points show the front and trailing ends of squirmers, respectively. At concentration $c = 0.1$, squirmers are accumulated near the walls for all values of $\beta$. Aggregation of pushers and pullers near the wall is stronger compared to the neutral squirmers. At high concentrations, squirmers are very close to each other and the accumulation near the walls is not obvious from Fig. 9. Another interesting phenomena is that there is a strong tendency for squirmers to orient toward the walls in the near-wall region.

We now quantitatively characterize the hydrodynamic interaction between the squirmers and the walls. We first define the percentage of near-wall squirmers $\varphi$ [see Fig. 10(a)] and the number of the collisions between squirmers normalized by the total number of squirmers $\varphi_c$ [see Fig. 10(b)]. Squirmer $n$ is considered to be near the wall when its distance from the wall $d_n = \min[y_n(t), L - y_n(t)]$ is smaller than a cutoff distance $d_{cut}$. Squirmer-averaged properties are plotted in Figs. 10(c)–10(f): the squirmer-averaged swimming speed $\overline{U} = \sum_{n=1}^{N} |u_n|/N$, the squirmer-averaged distance from the wall $\overline{d} = \sum_{n=1}^{N} d_n/N$, and the squirmer-averaged orientation angle $\overline{\varphi} = \sum_{n=1}^{N} \alpha_n/N$, where $\alpha_n$ is the orientation angle of the squirmer $n$ from the nearest wall. All the cases reach a statistical state at $t \simeq 20$. In the case of $c = 0.1$, around $60 \sim 80\%$ of squirmers swim within the distance of one diameter away from the wall, and most of them are actually in contact with the wall. In the suspension of squirmers at $c = 0.4$, the percentage of the near-wall squirmers is lower due to the dense packing and dominance of the steric interactions between squirmers. The percentage of squirmers undergoing collisions is shown in Fig. 10(b). There are more collisions among pushers compared to pullers and neutral squirmers at $c = 0.1$, which can be due to the larger swimming speed of pushers at $Re = 1$. At a high concentration, $c = 0.4$, $\varphi_c$ increases dramatically, reaching to about 2.

Figure 10(c) shows that the average swimming speed $\overline{U}$ decreases and reaches a statistically steady state at $t \simeq 20$. At small concentrations, $\overline{U}$ is approximately the same for different types of squirmers. At a high concentration (i.e., $c = 0.4$), the squirmer-averaged swimming speed $\overline{U}$ is reduced. Figure 10(d) shows the time evolution of average swimming speed for the squirmers in the near-wall region and in the bulk region. In the near-wall region, pullers swim faster; while in the bulk region, pushers swim faster. These results are consistent
with the results of a single squirmer shown in Fig. 4(d). We use the data of $t > 20$ to calculate the time-averaged speed over the total numbers of squirmers: $\langle U \rangle = \int_{0}^{T} U dt / T$. For $\beta = 0$, $-3$, and $3$ at $c = 0.1$, $\langle U \rangle$ is 0.44, 0.46, and 0.51, respectively, and 0.27 for $\beta = 0$ at $c = 0.4$. Recall from Fig. 1, the steady swimming speed of a single squirmer in an unbounded domain at $\text{Re} = 1$ is 1, 1.35, and 0.79 for $\beta = 0$, $-3$, and $3$, respectively. When near the wall, the swimming speed of a pusher decreases, while it increases for a puller as seen in Fig. 4(d). For a wall-trapped puller of $\beta = 3$ swimming along the wall, the steady speed is around 0.86.

In summary, in all our cases, the average swimming speed of suspension of squirmers decreases to around 30–60% of the speed of a single squirmer due to the effects of the walls and collision between squirmers.

Figures 10(e) and 10(f), show that both squirmer-averaged distance from the wall $d$ and squirmer-averaged orientation angle $\tilde{\alpha}$ decrease with time, indicating the tendency of squirmers to accumulate near the wall and to orient toward the wall. The distribution of squirmers is furthermore quantified by calculating the probability distribution function $f(\phi)$, which is defined as

$$f(\phi) = \frac{1}{N \Delta \phi} \sum_{n=1}^{N} (\delta(\phi_n - \phi)),$$  \hspace{1cm} (12)

and

$$\delta(\phi_n - \phi) = \begin{cases} 1, & \phi - \frac{\Delta \phi}{2} < \phi_n < \phi + \frac{\Delta \phi}{2}, \\ 0, & \text{otherwise}, \end{cases}$$  \hspace{1cm} (13)

where $\phi$ is the vertical position $y$ and orientation angle $\alpha$ in Figs. 11(a)–11(f), and $\Delta \phi$ is the interval of $\phi$ which is set to $\Delta y = 1.389$, $\Delta \alpha = 0.05\pi$, and $\Delta u = \Delta v = \Delta w = 0.1$. The choice of $\Delta \phi$ does not qualitatively change the results. The symbol $\langle \rangle$ represents a time-averaged quantity.

Figure 11(a) shows the probability distribution function of the vertical position of squirmers. The squirmers are accumulated near the walls and the probability distribution function rapidly falls away from the walls. When close to the wall, pushers and pullers have stronger tendencies to accumulate near the wall compared to neutral squirmers. This result is predictable since the stresslet (strongest far-field interaction) is absent in the case of the neutral squirmer [24]. At high concentrations, the peaks of $f(y)$ near the two walls are lower than the cases of $c = 0.1$ because the layer of the squirmers close to the wall is nearly saturated and the concentration in the middle of the channel grows. Similar results have been reported in Refs. [49,50]. In Fig. 12, we quantitatively compare our simulation results of pusher of $\beta = -3$ and $c = 0.1$ with published experimental and analytical results. The volume concentration of the cells in the experiments were reported in the range of $c = 0.01–0.1$ and the ratio between the channel width and the cell radius is around 40 for bull spermatozoa [9], 20–80 for E. coli [10], 66 for Caulobacter crescentus [11,12], and 6.6 for C.
FIG. 10. (Color online) Time evolution of (a) percentage of near-wall squirmers \( \varphi \), (b) percentage of colliding squirmers normalized by the total number of squirmers \( \varphi_c \), (c) squirmer-averaged swimming speed \( U \), (d) squirmer-averaged swimming speed for squirmers inside and outside the near-wall region distinguished by \( d_{cut} = 2 \), (e) squirmer-averaged distance from the wall \( d \), and (f) squirmer-averaged orientation angle \( \alpha \).

crescens [12], respectively. All the results show an increase in concentration of microorganisms near the wall, and the wall attraction is stronger for the smaller channel width. Our results show good agreements with the results of \textit{C. crescentus}, which is performed at a small channel width [12]. The analytical results solely based on the dipole interaction with the wall [10] overestimates the probability distribution of swimmers near the wall compared to the results of direct numerical simulation.

Figure 11(b) shows the probability distribution function of the orientation angle of the swimmers \( f(\alpha) \). A peak near \( \alpha = -\pi/2 \) is observed for all the cases. For squirmers of \( \beta = 0 \) at \( c = 0.1 \), another peak occurs near \( \alpha = 0 \). To better visualize the results, the probability distribution function of the orientation angle \( f(\alpha) \) is plotted for the squirmers near the wall and in the bulk region in Figs. 11(c) and 11(d), respectively. The near-wall squirmers are strongly oriented normal to the wall (see Fig. 9). Interestingly, for squirmers away from the wall, two different types of behavior are observed: for cases of \( \beta = 0 \) and \( 3 \) at \( c = 0.1 \), \( \alpha \) is mostly between \( -\pi/4 \) and \( \pi/4 \) and there is a clear peak at \( \alpha = 0 \); and for cases of \( \beta = -3, c = 0.1 \) and \( \beta = 0, c = 0.4 \), \( f(\alpha) \) is almost uniformly distributed over \( -\pi/2 \) to \( \pi/2 \). From the results of a single squirmer in Fig. 4(b), after the initial wall contact, all the squirmers detach from the wall with the final angle between \( -\pi/4 \) and \( \pi/4 \) (0.56, 0.41, and \( -0.24 \) for \( \beta = -3 \), 0, and 3, respectively). We have also shown that the final angles are independent of the initial impact angle of the squimmer if a close contact between the squimmer and the wall occurs. Therefore, when squirmers swim into the bulk region, their orientation angle is mainly between \( -\pi/4 \) and \( \pi/4 \). However, for cases of \( \beta = -3, c = 0.1 \) and \( \beta = 0, c = 0.4 \), squirmers collide with each other more frequently, which leads to a more uniform
distribution in the orientation angle. As shown in Fig. 11(e),
the distribution of vertical velocity component \( v \) dominates at
0, which also indicates the high percentage of the near-wall
squirmers. Similar results can also be found in Fig. 11(f) for \( u \)
and \( w \) components of velocity.

Previous studies have shown that the hydrodynamic inter-
action between squirmers generates large scale flows and leads
to a collective motion, with a range and duration depending
on the volume fraction and the swimming type \( \beta \) [36,60]. The
spatial correlation function \( g(r) \) can be used to quantify this interaction

\[
g(r) = \frac{L^3}{N(N-1)V} \sum_{m=1}^{N} \sum_{n=1}^{N} \langle \delta(r_{m,n} - r) \rangle,
\]

\[
\delta(r_{m,n} - r) = \begin{cases} 
1, & r - \frac{\Delta r}{2} \leq r_{m,n} < r + \frac{\Delta r}{2} \\
0, & \text{otherwise},
\end{cases}
\]

FIG. 12. (Color online) Comparison of vertical distribution of
microorganisms between present simulation results and previous
experimental and analytical studies.
where \( r_{m,n} \) is the distance between squirmers \( m \) and \( n \), 
\[ V(r_{m,n}) = \left(\frac{4\pi}{3}\right)\left(\frac{r_{m,n} + \Delta r/2}{\Delta r}\right)^3 - \left(\frac{r_{m,n} - \Delta r/2}{\Delta r}\right)^3 \]
is the volume of spherical shell of radius \( r_{m,n} \) and thickness \( \Delta r \).

Figures 13(a) and 13(b) show the pair distribution function for squirmers close to the wall and in the bulk region, respectively. For the squirmers near the wall, \( g(r) \) peaks at around \( r = 2 \) and 4, referring to the cluster formation of squirmers in the near-wall region. The curve of \( \beta = -3 \) has the highest peak at \( r \simeq 2 \), meaning that a larger number of pushers are in close contact [see Fig. 9(c)]. On the contrary, pullers are more scattered. This can be explained by the side-by-side interactions between two pushers, which is an attractive force, and two pullers, which is a repulsive force [2]. All curves have similar distributions for the bulk squirmers, the pair distribution function has a peak near \( r = 2 \), corresponding to the squirmers in close contact. Similar results were reported for the suspension of passive particles [61], bubbles [62], and two-dimensional swimming particles [60] in an unbounded domain.

The flow field near the bottom wall (\( y = 0 \)) generated by pushers (\( \beta = -3 \)) and pullers (\( \beta = 3 \)) is plotted in Fig. 14. The contourplot of the vertical velocity component is shown in the plane of \( y = 2.5 \). For clarity, the squirmers above the plane are not shown here. Near-wall pushers accumulate near each other and form large-scale coherent structures. Previous studies have shown that these coherent structures can increase the mass transport [41,42]. On the contrary, near-wall pullers are more scattered due to the side-by-side repelling force between them.

**IV. CONCLUSION**

We have studied the dynamics of a single and multiple low-Reynolds-number swimming organism(s) near a wall by conducting a three-dimensional direct numerical simulation. Each swimmer is modeled as a squirmer, which consists of a spherical body that propels itself by a tangential velocity distribution on its surface.

When a single squirmer is initially oriented toward the wall, three different modes are observed for \( \text{Re} = 1 \): (a) squirmers of \( \beta \leq 1 \) escapes from the wall, (b) squirmers of \( 2 \leq \beta \leq 5 \) oscillate near the wall and eventually swim along the wall keeping a constant distance and orientation angle, and (c) squirmers of \( \beta \geq 7 \) bounce on the wall. At a smaller Reynolds number, \( \text{Re} = 0.1 \), the initial wall contact time is increased independent of the swimming type, but a weaker attraction toward the wall is observed for the puller. The dynamics of suspension of squirmers between two walls is also studied. Similar to the observation of previous experiments [9–12], we found that the squirmers are strongly attracted to the walls. At a relatively small concentration of \( c = 0.1 \), around 60 ~ 80% of the squirmers are accumulated near the wall, the attraction of pushers and pullers
is stronger than neutral squirmers. At a high concentration, \( c = 0.4 \), around 40\% of the squirmers are near the wall. In the near-wall region, the squirmers mostly orient normal to the walls, while in the bulk region, the orientation angle of squirmers is more uniformly distributed or they orient in the direction parallel to the wall. The wall leads to the decrease of the average swimming speed of the squirmers. The pair distribution function shows that suspensions of pushers form large-scale clusters near a wall, which is not the case for pullers.

It is interesting to extend the results of this paper by including higher-order squirming modes which will affect the near-wall motion of the squirmer. In our simulations, the squirmer model is used to reduce the complexities of real microorganisms. The first two squirming modes capture the most important features of pusher- and puller-type microswimmers. Previous experiments of \( E. coli \) near a surface \([10,63]\) show that a stresslet and its image singularities, included to satisfy the no-slip boundary condition on the wall, describes the measured flow field around a bacterial cell near a wall with good accuracy. Since higher-order squirming modes will extensively expand the parameter space, our simulation as well as most of the previous studies \([30–37]\) only considered the first two squirming modes.

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APPENDIX

In addition to the validation test included in the main text, we provide two additional validation tests.

1. A squirmer colliding with a wall

Here, we compare the results of our numerical code against the published results of Spagnolie and Lauga \([24]\) for a neutral squirmer swimming near a wall. The computational domain is \([-16, 16] \times [0, 25.6] \times [-6.4, 6.4]\) with a uniform mesh size \( \Delta = 0.1 \) and the time step is set to \( \Delta t = 1 \times 10^{-3} \). The wall boundary is located at \( y = 0 \) and the squirmer is initially placed above the wall at \( h_0 = 2 \). Figure \( 15 \) shows the final steady angle of the squirmer of \( \beta = 0 \) with various initial angles \( \alpha_0 \). The results show good agreement with the simulation results of Spagnolie and Lauga \([24]\) at \( \text{Re} = 0 \) using a boundary-integral method based on regularized Stokeslets.

2. A particle moving between two parallel walls

Here, we compare the results of our numerical code against the published results of Bhattacharya \textit{et al.} \([46]\) for an inert particle moving with velocity \( U \) along \( x \) direction between two parallel walls and evaluate the drag force acting on the particle for different values of distance from the lower wall. The particle is initially located at \((0, h, 0)\). In order to facilitate the computation, we consider the frame of reference moving with the particle. In this frame, the particle is fixed and the walls, located at \( y = 0 \) and \( y = L \), move with velocity \( U \) along \( x \) direction. Inlet velocity \( U \) is imposed at \( x = -50 \). Simulations are conducted in a rectangular domain of \([-50, 50] \times [0, L] \times [-50, 50] \). The time step is set to \( \Delta t = 1 \times 10^{-3} \), and stretched grids are used where the mesh resolution around the particle is \( \Delta = 0.1 \). The distance between the particle and the lower wall is defined as \( \epsilon = h - 1 \) and the Reynolds number is set to \( \text{Re} = 0.01 \). We calculate the normalized drag force coefficient \( C_D = F_D / 6 \pi \mu U a \), where \( F_D \) is the drag force acting on the sphere. The comparison between our results and the semi-analytical results of Bhattacharya \textit{et al.} \([46]\) using a modified Stokesian dynamics method is shown in Fig. \( 16 \). Our results start to deviate from the results of Bhattacharya \textit{et al.} \([46]\) at \( \epsilon \approx 0.05 \), which corresponds to half a mesh size. Similar accuracy is also achieved by Lambert \textit{et al.} \([30]\) when simulating two spherical particles in shearing or squeezing motion.

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FIG. 15. (Color online) Comparison of the final angle of a neutral squirmer for various initial angles and a constant initial height of \( h_0 = 2 \) with the results of Ref. \([24]\).

FIG. 16. (Color online) Comparison of the results for a single translating particle between two walls. Symbols, present numerical results at \( \text{Re} = 0.01 \); lines, simulation results of Ref. \([46]\).