of Physics

Deutsche Physikalische Gesellschaft 🕕 DPG

IOP Institute of Physics

Published in partnership

with: Deutsche Physikalische

Gesellschaft and the Institute

New Journal of Physics

The open access journal at the forefront of physics

CrossMark

OPEN ACCESS

RECEIVED 22 May 2017

REVISED 1 September 2017

ACCEPTED FOR PUBLICATION

18 September 2017 PUBLISHED

7 November 2017

Original content from this work may be used under the terms of the Creative Commons Attribution 3.0 licence.

Any further distribution of this work must maintain attribution to the author(s) and the title of the work, journal citation and DOI.



Active particles in periodic lattices

Alexander Chamolly¹, Takuji Ishikawa² and Eric Lauga^{1,3}

- Department of Applied Mathematics and Theoretical Physics, University of Cambridge, Wilberforce Road, Cambridge CB3 0WA, United Kingdom
- ² Department of Finemechanics, Graduate School of Engineering, Tohoku University, 6-6-01, Aoba, Aoba-ku, Sendai 980-8579, Japan
 ³ Author to whom any correspondence should be addressed

E-mail: ajc297@damtp.cam.ac.uk, ishikawa@pfsl.mech.tohoku.ac.jp and e.lauga@damtp.cam.ac.uk

Keywords: low-Reynolds number locomotion, active particles, complex environment, Stokesian dynamics, lubrication theory Supplementary material for this article is available online

Abstract

PAPER

Both natural and artificial small-scale swimmers may often self-propel in environments subject to complex geometrical constraints. While most past theoretical work on low-Reynolds number locomotion addressed idealised geometrical situations, not much is known on the motion of swimmers in heterogeneous environments. As a first theoretical model, we investigate numerically the behaviour of a single spherical micro-swimmer located in an infinite, periodic body-centred cubic lattice consisting of rigid inert spheres of the same size as the swimmer. Running a large number of simulations we uncover the phase diagram of possible trajectories as a function of the strength of the swimming actuation and the packing density of the lattice. We then use hydrodynamic theory to rationalise our computational results and show in particular how the far-field nature of the swimmer (pusher versus puller) governs even the behaviour at high volume fractions.

1. Introduction

Swimming microorganisms live in a variety of natural and industrial environments, including the ocean, soil, intestinal tract and bioreactors, and they play diverse and important roles in environmental, agricultural and health issues [1–3]. Understanding the spreading of cells in various environments is therefore a fundamental problem, and this knowledge can be exploited for predicting and controlling the distribution of cells and their growth [4, 5].

From a physical point of view, the simplest environment for swimming cells is an unbounded fluid in the absence of background flow or the biological responses to stimulus, including chemotaxis, phototaxis or gravitaxis. The spreading of individual cells in such a simple environment has already been studied thoroughly. Early work investigated run and tumble motions of individual *Escherichia coli* (*E. coli*) bacteria and showed that the motion of the cells could be described by random walk models with the entire population displaying diffusive behaviour [6]. Similar results were also found for the unicellular alga *Chlamydomonas*, and cell populations displayed long-time diffusive behaviour [7–10].

The behaviour of swimming microorganisms in the presence of a planar wall has also been investigated extensively. Rothschild first reported a significant increase of cell density near a glass wall using bull spermatozoa [11]. Similar accumulation of cells near a wall was also found for human spermatozoa [12] and *E. coli* bacteria [13, 14]. The mechanism of far-field attraction to a wall may be in part explained by hydrodynamics [15]. When a swimming cell is a so-called pusher, i.e. its thrust is generated behind the cell body like spermatozoa or flagellated bacteria, the cell tends to direct towards the wall due to far-field hydrodynamic interactions. When a swimming cell is a puller, i.e. when its thrust is generated in front of the cell body, on the other hand, cells swimming parallel to the wall tend to be repelled by it hydrodynamically.

Some swimming microorganisms are seen to be trapped by planar walls. Swimming *E. coli* cells are drawn to a stable circular trajectory, and the physical mechanism underlying the entrapment has been explained by hydrodynamic and steric effects [16–19]. Spermatozoa can also be trapped by a planar wall, and the entrapment

mechanism is again explained by hydrodynamic and steric effects [20–22]. Such entrapment phenomena may reduce the spreading of cells near surfaces considerably.

Other types of surface-related behaviour have also been reported for both biological and artificial swimmers. *Chlamydomonas* algae cells scatter from a flat wall due to contact between its flagella and the surface and exhibit billard-like motion in polygonal geometries [23]. The collective behaviour of *Bacillus subtilis* swarms and their interactions has been shown to depend on the geometry of the enclosing cavities [24]. Chemically propelled rods tend to move on a surface along large circles with stochastic changes in the sign of the orbit curvature [25] and may be captured around solid spheres of various radii and materials [26]. Colloidal particles rolling along a planar wall displayed a transition to macroscopic directed motion for large populations [27]. Swimming *E. coli* cells were seen to be entrapped along convex walls provided their curvature was sufficiently low [28]. Theoretically, an analysis of the mechanism of entrapment showed that a swimmer approaching a spherical colloid can be captured hydrodynamically when the colloid is larger than a critical size [29].

In nature many microorganisms live in environments subject to complex geometrical constraints. Bacteria in soil, for instance, swim in heterogeneous granular matter, and their habitat is influenced by physical parameters including granular size and shape [30–34]. While locomotion of organisms in granular media with particle size smaller than the body length has been investigated widely [35, 36], only a few studies have addressed the spreading of microswimmers in complex geometries [37]. A study on the motion of Janus particles in a two-dimensional (2D) periodic pattern of ellipsoidal pillars arranged in a triangular lattice showed that microswimmers can steer even perpendicularly to an applied force in the periodic pattern [38]. Swimming Janus particles and *E. coli* cells in a 2D colloidal crystal showed that artificial particles orbit individual colloids with occasional hops while the circular orbits bacteria along planar walls were rectified into long, straight runs [39].

Given the potential environmental complexity encountered by both natural and artificial swimmers, a lot remains to be explored. In particular, the role played by both the type of swimmer and its environment in dictating how the swimmers spread on average is a fundamental quantity of interest. To this end, we consider in this paper a model system of swimming in a periodic environment. The swimmer is a spherical squirmer [40, 41], one of the simplest models of low-Reynolds number swimmers that allows us to compute the hydrodynamics precisely. The swimmer is then located in an infinite three-dimensional body-centred cubic (BCC) lattice of passive, rigid spheres of the same radius and its trajectory is computed using Stokesian dynamics [42]. Running a large number of simulations, we are able to address three main questions in our study: (1) which qualitative types of trajectories arise in periodic lattices, (2) how these depend on the strength of the swimmer and the packing density of the lattice and (3) how to explain these results from hydrodynamic arguments.

2. Model and simulations

Our computations are carried using a modified version of the Stokesian dynamics code used and validated in [42]. We consider a single spherical squirmer of radius 1 and 64 identical rigid spheres of the same radius 1 arranged as a rigid stationary BCC lattice. The flow satisfies the no-slip boundary condition on their surface. A unit lattice cell has side length *l* and contains the equivalent to two lattice spheres, while the computational cell consists of $4 \times 4 \times 2$ unit cells, thus including the equivalent to 64 lattice spheres, plus the squirmer (see figure 1). At the boundary of the computational cell, periodic boundary conditions for the flow are applied. The lattice volume fraction, or packing density, ϕ is related to l by $\phi = 8\pi/3l^3$.

The squirmer induces an effective slip velocity tangential to its surface in order to propel forward. Specifically, in spherical coordinates measured from its centre, the velocity boundary conditions applied by the squirmer in the swimming frame are given by [40, 41]

$$u_r(1,\theta) = u_\phi(1,\theta) = 0, \quad u_\theta(1,\theta) = \frac{3}{2}\sin\theta(1+\beta\cos\theta), \tag{1}$$

where θ is the azimuthal angle from the squirmer orientation **e** and β is the so-called squirming parameter dictating the nature of the far-field flow. A swimmer with $\beta > 0$ is a puller, like the alga *Chlamydomonas*, whereas one with $\beta < 0$ is a pusher, similar to most flagellated bacteria and spermatozoa.

In absence of the lattice, the boundary conditions in equation (1) have been chosen such that an individual squirmer would swim with unit speed [41]. The characteristic time used to non-dimensionalise the equations is thus the time scale it takes for one swimmer to swim is own radius. With this scaling, the simulations uses a timestep of 10^{-3} and a total of 10^{6} steps, thus is equivalent to 1000 dimensionless time units. The viscosity of the fluid, η , is also scaled to unity.

In our simulations the squirmer is initialised at the centre of a face of a unit lattice cell, with either random or prescribed orientation. Note that through the periodic boundary conditions our model corresponds to an infinite lattice with an infinite number of squirmers. However, the separation between them is large enough to



make any hydrodynamic interactions negligible compared to those with the surrounding lattice spheres, giving an appropriate model for a single swimmer in an infinite lattice.

In what follows, we examine volume fractions between 1% and 13%. The theoretical maximum to allow space for a squirmer to pass through the lattice along one of its axes (i.e. such that $l \ge 4$) is $\phi = \pi/24 \approx 13.1\%$, so the examined parameter range includes the very dilute limit up to almost the densest possible lattices that the swimmer could theoretically traverse. We also vary the strength of the squirming parameter by considering the range $-3 \le \beta \le 2$ and thus include both pullers an pushers.

3. Computational results

3.1. Zoology of trajectories

We ran a large number of simulations in our parameter range $-3 \le \beta \le 2$, $1\% \le \phi \le 13\%$ with random initial orientations. We observe qualitatively different features of the resulting trajectories originating from a complex interplay of both far-field and near-field interactions with the lattice (see illustration in figure 3 and details below). In order to classify the various different kinds of behaviour rigorously and systematically, we first introduce some definitions.

3.1.1. Definition #1: turns versus straight

A squirmer is said to make a turn every time its orientation changes by more than 60° from when it last made a turn (or from the initial orientation in case of the first turn). If a squirmer makes no turn during the entire simulation, or more than 500 time units elapse between two subsequent turns, then we say the trajectory is straight. The latter condition is added to discount transients occurring at the beginning of the simulation and to account for trajectories with very long straight segments, and the results are not sensitive to the exact value of the threshold, which has merely been chosen to be much larger than the time scale of scattering from the lattice. The angular threshold can be motivated by the typical behaviour observed in our numerical simulations: the swimmers do not perform U-turns, but often tend to wiggle around an otherwise straight path. The particular angular value we have chosen reflects these observations and has proven to capture the intuitive idea of a turn (as opposed to a wiggle) correctly in all cases considered. Note that the translational velocity direction and the orientation of the squirmer need not coincide and we may also have chosen this as a criterion for the definition (although it did perform not as effectively as the angular criterion). Note furthermore that our criterion does not distinguish between a sharp turn and a long curved trajectory; however the latter does not occur in our

3.1.2. Definition #2: random walks

When swimmers are able to spread in the lattice without being trapped, but do not display straight trajectories, their behaviour is akin to that of a random walk with periods of straight swimming followed by reorientations after collisions with lattice spheres. When a swimmer follows a random walk, we measure the total number of turns undergone by the swimmer over the entire course of the simulation (see below).



3.1.3. Definition #3: trapped

Writing the position of the centre of the squirmer as $\mathbf{r}(t)$, we define a squirmer to be trapped if there is a time t^* such that $|\mathbf{r}(t^* + t) - \mathbf{r}(t^*)| \leq 5$ for all times 0 < t < 100. This means that a swimmer is considered trapped if it does not move further than 5 radii in 20 times the time it would require to traverse that distance at full speed. The spatial threshold here is chosen to identify squirmers trapped by (and so remaining in contact with) a single lattice sphere, in which case the displacement is limited to slightly more than 4 radii. The temporal threshold is chosen to be much larger than the time required for the free case, and is not sensitive to the precise value as squirmers tend to get caught early in the simulations.

3.1.4. Definition #4: stuck versus orbits

If a swimmer remains in the proximity of a single lattice sphere, we introduce two further definitions to distinguish swimmers that stay in place from those that move on confined periodic or chaotic trajectories. A trapped trajectory is called stuck if it remains straight, and orbiting if it does not.

3.2. Phase diagram

With these definitions, we can now summarise our computational results in a phase diagram. This is shown in figure 2 in the $\beta-\phi$ plane for squirmers with random initial orientations. Four different behaviours can be distinguished: straight trajectories; random walks (with a colour scheme capturing the number of turns occurring in total during the full computation); orbiting swimmers; and stuck swimmers. The red line around $\beta \approx -1.8$ to -1.6 provide the boundary between trapped swimmers (strong pushers) and those which are not trapped (pullers and weak pushers).

3.2.1. Pullers

All pullers considered in our simulations ($\beta > 0$) move through the lattice in nearly straight paths at almost unit speed. These paths are not limited to perfectly straight lines parallel to the lattice axes but includes also diagonal paths with small wiggles around obstacles. A simple far-field theoretical approach in section 4 will allow us to rationalise these observations.



Figure 5. Sample trajectories for a swimmer moving in a BCC lattice for various values of the squirming parameter (β) and the lattice volume fraction (ϕ): (a) weak pusher ($\beta = -1.0$) undergoing straight swimming ($\phi = 11\%$); (b) weak pusher ($\beta = -1.0$) undergoing a random walk in the lattice ($\phi = 5\%$); (c) strong pusher ($\beta = -2.4$) orbiting around of the spheres in the lattice ($\phi = 5\%$); (d) strong pusher ($\beta = -2.4$) stuck in the lattice ($\phi = 11\%$). See the supplementary material available online at stacks. iop.org/NJP/19/115001/mmedia for animated versions of these and for pullers at $\beta = 1.0$.

3.2.2. Pushers

For weak pushers $(-1.6 \leq \beta < 0)$, similar straight trajectories are observed at small volume fractions, albeit with larger wiggles (figure 3(a)). However, when the volume fraction becomes larger the active particles are deflected through interactions with the lattice, leading to random walks (figure 3(b)). Most of these random walks are very weak, with a maximum of ≈ 20 turns over the course of the simulations, corresponding to a turn only every couple of lattice cells (see coloured cells in figure 2). Further we observe that the number of turns in the random walks increase with the volume fraction of the lattice (ϕ) and with the strength of the flow imposed by the swimmer (β), as might be expected from intuition. In contrast, for very small values of ϕ the interactions become weaker and the number of turns reduces to a small non-zero value for a wide range of β values. Interestingly, for intermediate-strength pushers, the random trajectories return to straight swimming for very high values of ϕ , as illustrated in figure 4.

When the pusher becomes stronger than a critical value, found to be approximately $\beta \leq -1.6$ in our simulations (and closer to $\beta \leq -1.8$ in the very dilute lattice limit) we observe a sharp transition to a trapped state in a manner which appears to be largely independent of the volume fraction in the lattice. In the trapped region, at small volume fractions, a swimmer orbits a single lattice sphere (figure 3(c)) but becomes stuck for larger values of ϕ (figure 3(d)). The range of values of β leading to a stuck swimmer shrinks as β is decreased further until all trajectories are orbits at $\beta \leq -3$. Interestingly, these orbits are nearly periodic circles for small volume fractions, but become chaotic for larger values of ϕ , apparently through constraints by the surrounding lattice environment. In all cases, orbiting squirmers are not oriented tangentially to the lattice sphere, but always tend to be at a small acute angle with it that increases with $|\beta|$, effectively 'sliding' around their captivators.



Figure 4. Long-time trajectories (blue solid lines) of intermediate-strength pushers ($\beta = -1$) transitioning from random walks at low volume fraction ((a), $\phi = 5\%$) to straight swimming at high volume fractions ((b), $\phi = 11\%$). Shadows on the *x*-*y*, *x*-*z* and *y*-*z* planes are drawn to help the eye (red dashed–dotted lines), and arrows indicate the direction of motion (length scales are scaled by the squirmer radius).



3.2.3. Dependence on initial orientation

In order to analyse the dependence of the random walks on the volume fraction of the lattice we performed multiple runs for selected values of β and ϕ using different initial orientations of the swimmer. The space of orientations is suitable for Lebedev quadrature so that expected values can be calculated for a randomly initialised squirmer [43]. We chose 17th-degree quadrature which ordinarily requires 110 sample orientations but this can be reduced to 12 by exploiting the symmetries of the lattice, reducing computational effort substantially. Recognising that the trajectory depends very sensitively on the initial orientation, we emphasise that the analysis of these results should be interpreted heuristically.

In figure 5 we plot for a given weak pusher ($\beta = -1.0$) the expected number of turns (blue solid line) and the probability that a trajectory is straight (dashed red line). Strikingly, the probability to obtain a straight trajectory rises sharply to unity at volume fractions greater or equal to 10%, along with a corresponding sharp decrease in the number turns (that need not be zero, see section 3.1.1). Likewise, we see that trajectories are on average more straight at very low volume fractions, which is what one would expect for the case of free swimmer. Similar computations for values of β in the trapped regime (strong pusher, not shown) also reveal that the average distance travelled by swimmers through the lattice decreases as either the volume fraction and or the squirming parameter increase.

4. Theory

Our simulations reveal a number of remarkable features of the locomotion of an active particle in a lattice. In this section, we provide theoretical arguments in order to rationalise these observations. We focus below on three main questions arising from the phase diagram in figure 2: (1) Why are pullers able to swim straight while weak pushers tend to undergo random walks? (2) Why is there a threshold for pushers to be trapped? (3) How can we understand the transition, in the trapped regime, between an orbiting and a stuck state as a function of the volume fraction of the lattice?

4.1. Straight swimming versus random walks

In order to understand why pullers swim straight though the lattice while weak pushers tend to undergo random walks, we use a far-field analysis of the hydrodynamic interactions between the swimming spheres and the stationary spheres in the lattice. In this limit, this can be accomplished using the method of reflections [44, 45]. Below we show analytically that squirmers swimming through a channel of spheres are stabilised and destabilised along their straight trajectory in approximately equal intervals but with varying strength and in a manner that changes sign with the squirming parameter.

The leading-order flow induced by the squirmer is that of a stresslet (force-dipole) which can be written as [46, 47]

$$\mathbf{u}(\mathbf{r}) = -\frac{3}{4}\beta \left(3\frac{\mathbf{r}(\mathbf{e}\cdot\mathbf{r})^2}{r^5} - \frac{\mathbf{r}}{r^3}\right),\tag{2}$$

where the squirmer is instantaneously centred at the origin and has orientation **e**. The leading-order hydrodynamic response to this flow by a lattice sphere located at \mathbf{r}_0 is a Stokeslet (point force) that can be calculated using Faxèn relations [45]. The response of the squirmer to this Stokeslet is then a perturbation, $\mathbf{u}_{\text{Stokes}}$, to its swimming velocity as

$$\mathbf{u}_{\text{Stokes}} = \frac{9}{8}\beta(3\cos^2\theta - 1)\frac{\mathbf{r}_0}{r_0^4} + \mathcal{O}(r_0^{-4}), \quad \cos\theta \coloneqq \mathbf{e} \cdot \hat{\mathbf{r}}_0, \tag{3}$$

with higher-order terms due to further flow singularities and their responses. Importantly, the sign of this velocity perturbation depends both on the squirming parameter, β , and on the angle that **e** makes with the displacement vector to the lattice sphere; this is a consequence of the dipolar nature of the flow in equation (2). From this, it is clear that the geometry of the lattice and the position and orientation of the squirmer within it determine its behaviour in a very non-trivial fashion.

Further progress can be made however by considering a simplified scenario, illustrated in figure 6. Consider a squirmer confined to a 2D plane moving through a 'channel' of lattice spheres positioned at $\mathbf{r}_i^{\pm} = il\hat{\mathbf{x}} \pm l/2\hat{\mathbf{y}}$ where *i* is an integer. Such a channel is a common feature of regular lattices such as cubic, BCC and FCC, which can all be thought of as assembled from interlaced channels at different orientations. In particular, by linearity of the flow in the dilute limit the results obtained from this model will give qualitative predictions for the threedimensional case at sufficiently low volume fractions.

Since the Stokeslet responses are directed towards the centre of the squirmer, they do not exert any net torque on it. For this reason the rate of change of \mathbf{e} is with $\mathcal{O}(l^{-5})$ at least two orders of magnitude smaller than the rate of change of ϵ . For the purpose of the argument below, we may therefore assume that the orientation of a swimmer, \mathbf{e} , remains unchanged. We focus on the case $\mathbf{e} = \hat{\mathbf{x}}$ and quantify the role of hydrodynamic interactions on the motion of the sphere. The squirmer position is denoted by $\mathbf{r} = (x(t)l, \epsilon(t)l)$ where $|\epsilon| \ll 1$. Neglecting all but the leading-order interactions we obtain the dynamical system for (x, ϵ) as

$$\frac{\mathrm{d}x}{\mathrm{d}\tau} = 1 + \mathcal{O}(\epsilon), \quad \frac{\mathrm{d}\epsilon}{\mathrm{d}\tau} \approx \frac{9\beta}{8l^3} \sum_{i,\pm} \frac{3\cos^2\theta_i^{\pm} - 1}{|\mathbf{r}_i^{\pm} - \mathbf{r}|^3} \sin\theta_i^{\pm}, \tag{4}$$

where $\tau = t/l$ and θ_i^{\pm} is the angle between the vectors **e** and $\mathbf{r}_i^{\pm} - \mathbf{r}$. Upon linearising in ϵ the terms in the sum we obtain the differential equation for the off-axis location of the swimmer as

$$\frac{\mathrm{d}\epsilon}{\mathrm{d}x} = -\frac{36\beta}{l^3} \left\{ \sum_{n=-\infty}^{\infty} \frac{3 - 52(n-x)^2 + 32(n-x)^4}{[1 + 4(n-x)^2]^4} \right\} \epsilon + \mathcal{O}(\epsilon^2), \tag{5}$$

which integrates to

$$\ln\left(\frac{\epsilon}{\epsilon_0}\right) \approx -\frac{36\beta}{l^3} \sum_{n=-\infty}^{\infty} \left\{ \frac{11(x-n) + 16(x-n)^5}{4(1+4(x-n)^2)^3} + \frac{1}{8} \tan^{-1}(2(x-n)) \right\}.$$
 (6)

Each term in the sum represents a kink in $\ln \epsilon$ centred around x = n of size $\pi/8$ and width O(1). Taking the average over unit intervals we can therefore deduce that







Figure 7. Instantaneous growth/decay of a perturbation ϵ away from a trajectory through the middle of a two-dimensional channel of rigid spheres. Results are shown in the case of a puller ($\beta > 0$); for a pusher the stability properties are reversed via a change of sign. The contribution from a single \pm -pair of spheres is displayed for illustration purposes (blue dashed line) while the total effect of all spheres is a superposition of such pair effects (red solid line).

$$\langle \ln(\epsilon/\epsilon_0) \rangle \approx -\frac{9\pi\beta}{2l^3} x = -\frac{27}{16}\beta\phi\tau \quad \Rightarrow \quad \epsilon(\tau) \sim e^{-(27/16)\beta\phi\tau}.$$
 (7)

From this we see that puller trajectories ($\beta > 0$) are stable, whereas pushers ($\beta < 0$) are not.

In figure 7 we plot the effect of a single pair of spheres at $y = \pm l/2$, corresponding to one term in the series (n = 0), on a puller squirmer ($\beta > 0$) depending on its position relative to them (dashed blue line). We also plot the influence of all spheres in the channel (solid red line). We note that any contributions from spheres more than $\approx 0.75l$ away from the swimmer are vanishingly small compared to the closer ones, a result self-consistent with our neglecting of any sphere above and below the channel, which thus allows us to generalise our heuristic argument to the full lattice. As the swimmer moves through the lattice, the perturbation of its position away from the central axis of the channel, ϵ , undergoes alternating periods of growth (positive regions in figure 7) and decay (negative regions). We observe that the perturbations grow in one direction when the swimmer is roughly between a \pm -pair of spheres and grow in the other when it is roughly in the middle between two such pairs. Importantly, the rate of change is considerably greater in the decay region than in the growth region. This is illustrated in figure 8 where we plot the spatial variation of the perturbation, ϵ , as the swimmer makes its way



Figure 8. Decay of a perturbation for the off-axis position of the swimmer in a two-dimensional channel of rigid spheres (setup in figure 6) in the case of puller ($\beta > 0$) and $\beta \phi = 0.1$ over 10 unit lattice lengths (blue solid line). The shape of the curve is generic for a puller. While the deviation of the swimmer away from the axis undergoes periods of both growth and decay, overall the latter dominates the dynamics, leading to exponential decay of the perturbation on average (equation (7), red dashed line).

through the lattice, as well as the log-average from equation (7). As a consequence, the perturbation undergoes an oscillating decay for a puller and the straight trajectory is therefore stable, consistent with our numerical simulations. In contrast, for a pusher the decay/growth regions in figure 7 are reversed and the swimmer is ultimately unstable, as observed in our simulations at low volume fractions. Furthermore we note that the force with which the squirmer is stabilised or destabilised in this dilute model scales as $\sim \beta \phi$. Therefore the effect becomes very small at low volume fractions; in particular, when the orientation of the squirmer is not aligned with an axis of the lattice, trajectories become more complex and are impacted by scattering from lattice spheres.

Conversely, as ϕ is increased for pushers, there is less space for them to turn and reorient, forcing them on an effectively straight trajectory. This is what is observed in figure 3(a). The clear jump in the probability that pushers go straight in figure 5 can therefore be explained by the existence of a critical volume fraction ϕ_c increasing with $|\beta|$, above which the squirmers cannot reorient quickly enough to leave an effectively straight path through the lattice.

4.2. Unbounded versus bounded trajectories

Similar far-field considerations may be used to obtain an estimate for the threshold in pusher strength beyond which all swimmers are trapped. For this, we make use of a recent paper which analysed the swimming of force dipoles near rigid spheres using the method of hydrodynamic images [29]. In that study, it was shown that for a pusher swimmer of unit radius creating a dipolar flow of strength α , trapping would occur along colloids whose radius, *A*, exceed a critical value A_c given by

$$A_c = \frac{64}{9\alpha^2}.$$
(8)

Swimmers near colloids for which $A < A_c$ will continue swimming (possibly with some scattering) while in the case $A > A_c$ pushers would all be predicted to be trapped.

Since the passive spheres of the lattice are of radius 1, this means that trapping would be predicted to occur for strong pushers, $|\alpha| > |\alpha_c|$, with

$$\alpha_c = \left(\frac{64}{9}\right)^{1/2} = \frac{8}{3}.$$
(9)

In our model, equation (2) indicates that the strength of the dipolar flow is given by

$$\alpha = -\frac{3}{4}\beta,\tag{10}$$

and thus trapping is predicted to occur for this far-field model for strong pushers satisfying

$$\beta < \beta_c, \quad \beta_c = -\frac{32}{9} \approx -3.56. \tag{11}$$

This number is off by about a factor of two from the results of our numerical simulations ($\beta_{c,num} \approx -1.8$). Since the theoretical prediction in [29] was derived in the case of a unique swimmer-sphere pair in the far field, and thus neglects hydrodynamic interactions with other spheres and all near-field effects, it is striking that the prediction remains robust and may also be used to predict the swimming behaviour in a lattice.





4.3. Stuck swimmer versus orbiting

So far we provided theoretical arguments for two of the main numerical observations, namely the qualitative difference between pushers and pullers as well as the transition, for the case of pushers, between a trapped and non-trapped regime. We now address the difference, within the trapped region, between swimmers which effectively remain stuck and those which continuously orbit around of the lattice spheres. In this case, the dynamics can only be explained through an interplay between generic far field hydrodynamic interactions with the lattice (as outlined above) and near-field hydrodynamic forces which necessarily depend on the details of the squirmer actuation.

In order to capture near-field effects, we now derive an argument based on lubrication theory (i.e. the equations of hydrodynamics valid in thin geometries [48]) to derive an expression for the torque experienced by a squirmer near an inert sphere as a function of its orientation (assuming that we are beyond the threshold for bounded dynamics). The distinct stuck versus orbiting behaviour would then arise depending on whether lubrication (high volume fraction) or far-field effects (low volume fraction) dominate.

To simplify calculations, we consider the 2D case of two cylinders ('spheres'), which although they might exhibit different scalings for the forces and torques, would capture the same flow physics (see sketch in figure 9). Because the dynamics are Stokesian there is no inertia, and hence a squirmer sliding around a sphere at constant velocity is completely equivalent to having a stationary squirmer next to a uniformly rotating sphere. We work in this frame in order to directly apply the squirming boundary condition, equation (1). We next assume that the squirmer remains at a constant distance h_0 from the inert 'sphere' surface and that its swimming direction, \mathbf{e} , makes an acute angle α with the normal to it. The gap width is thus written as

$$h(\varphi) = h_0 + 2(1 - \cos\varphi), \tag{12}$$

where φ is the angle between the normal and a point on the squirmer. We assume that the speed at which the surface is rotating is constant and write it as $u_{\text{slide}} = -\gamma \sin \alpha$ where γ is a constant between 0 and 1 that reflects the fact that it is slowed down by friction compared to the unit speed of an unobstructed swimmer.

From equation (1) we have that the tangential velocity on the squirmer is given by

$$u_{\text{Squirmer}} = -3/2\sin(\alpha - \varphi)(1 + \beta\cos(\alpha - \varphi)), \tag{13}$$

in the direction of increasing φ (see figure 9). In the lubrication limit the flow in the gap is a pure shear flow and therefore the total tangential stress on the squirmer is given by

$$\Gamma = -\int_{-\varphi_0}^{\varphi_0} \frac{u_{\text{Squirmer}} - u_{\text{slide}}}{h} \cos^2 \varphi \, \mathrm{d}\varphi \tag{14}$$

$$= -\int_{-\varphi_0}^{\varphi_0} \left(\frac{-\frac{3}{2}\sin(\alpha - \varphi)(1 + \beta\cos(\alpha - \varphi)) - (-\gamma\sin\alpha)}{h} \right) \cos^2\varphi \, \mathrm{d}\varphi, \tag{15}$$

where φ_0 is a cut-off angle for the region in which lubrication theory is valid ($\sqrt{h} \ll 1$) and the factor of $\cos^2 \varphi$ arises from the need to project integrand and integration measure into the direction perpendicular to the gap. Carrying out the integration leads to

$$\tau = \sin \alpha \frac{3\pi}{2\sqrt{h_0}} \left(1 - \frac{2}{3}\gamma + \beta \cos \alpha \right),\tag{16}$$

at leading order in h_0 as $h_0 \to 0$. We note that $\tau \sim h_0^{-1/2}$ in this 2D setting while for spheres one finds $\tau \sim \ln h_0^{-1}$ [49, 50].

This traction results in a torque on the squirmer and we are interested in the consequences of this torque on the dynamics of trapped swimmers. Analysing equation (16) for equilibrium points in α we find that $\alpha = 0$ is unstable for $\beta > \beta_c$ and stable for $\beta < \beta_c$ where

$$\beta_c = -1 + \frac{2}{3}\gamma,\tag{17}$$

and $0 < \gamma < 1$ implies $-1 < \beta_c < -1/3$. Compared with equation (11) we see that the threshold for entrapment predicted from considering near field effects only is much lower than that expected from far field effects. As the lubrication forces scale rather weakly with h_0 in three-dimensions and higher order effects from the global geometry will matter, we conclude that a theoretical prediction for the critical squirmer strength should lie between those two extremes, as is the case with our numerical observed value of $\beta_c \approx -1.8$.

We propose that the sliding behaviour can be understood as the consequence of a balance between far- and near-field hydrodynamic effects, shifting α to a non-zero equilibrium value that increases with the squirmer strength (i.e. with $|\beta|$). For strong pushers, the far-field hydrodynamic interactions dominates, explaining why there are more orbiting and fewer stuck trajectories. Additionally, at high volume fractions interactions with other lattice spheres start to become important. The net effect of these appears to be repulsion once a squirmer has been trapped. When the strength of the swimmer is not too strong and the volume fraction of the lattice is sufficiently large, the repulsion is strong enough to reduce the equilibrium value of α to a very small value, resulting in the behaviour classified as 'stuck'. In contrast, when the squirming strength is sufficiently large, the swimmer gets deflected from its near-circular trajectory around its captivator, giving rise to the observed 'orbiting' behaviour.

5. Conclusion

Using extensive numerical simulations we computed the trajectories of spherical squirmers through BCC lattices for a broad range of both the squirming parameter, β , and the lattice packing density, ϕ . We found a rich array of different qualitative behaviours, ranging from straight trajectories over weakly diffusive random walks to instant entrapment by the lattice.

We further demonstrated that much of this behaviour can be explained, at least qualitatively, using a combination of simple hydrodynamic near-field and far-field arguments for the squirmer and its interactions with individual spheres in the lattice. Specifically, pullers tend to be stabilised and pushers destabilised on trajectories far from lattice elements. Close to lattice spheres, near-field and far-field hydrodynamic torques compete. There is a clear threshold which is almost independent of the lattice volume fraction and a strong pusher above it is trapped, whereas a weak pusher below it is deflected but ultimately released back into the lattice. This may then result either in a trajectory resembling a weak random walk or, if too constrained by the lattice packing density, again a straight path. For a given squirming strength, there appears to be a clear threshold in packing density above which all trajectories are straight regardless of initial conditions.

In the parameter range for which the swimmer performs a random walk, it would be desirable to quantify the effective diffusive motion of the active particles. However, because reorientation events occur very infrequently, it is currently too computationally expensive to quantify it accurately from our simulations. A recent study has investigated this for a random Lorentz gas [37], and future work with more powerful numerical algorithms may shed more light on this issue taking hydrodynamics into account. Similarly, the effects of lattice geometry warrant further investigation: while test runs in a cubic lattice reveal the same generic behaviour of squirmers in the far field, the question to what extent the stuck-orbiting and trapped-free thresholds depend on the lattice geometry remains to be answered.

Future studies could address how additional properties of the squirmer can affect the dynamics, for example potential phoretic effects. In a setup with a chemically-patterned lattice, a passive particle may propel through phoretic interactions, or in the case of a weakly charged lattice a charged swimmer may be affected through electromagnetic effects. Practical applications in which our results may be put into use include the design of porous filters for swimmers of different types and strengths.

Acknowledgments

This work was supported by a Rouse Ball Studentship by Trinity College to AC, a JSPS Grant-in-Aid for Scientific Research (A) (17H00853) to TI and an ERC Grant PhyMeBa (682754) to EL.

References

- [1] Ingraham J L and Ingraham CA 2003 Introduction to Microbiology (Pacific Grove, CA: Brooks/Cole Publishing)
- [2] Wall D H, Nielsen U N and Six J 2015 Soil biodiversity and human health Nature 528 69-76
- [3] Chown SL, Clarke A, Fraser CI, Cary SC, Moon KL and McGeoch MA 2015 The changing form of antarctic biodiversity Nature 522 431–8
- [4] Pedley T J and Kessler J O 1992 Hydrodynamic phenomena in suspensions of swimming microorganisms Annu. Rev. Fluid Mech. 24 313–58
- [5] Guasto J S, Rusconi R and Stocker R 2012 Fluid mechanics of planktonic microorganisms Annu. Rev. Fluid Mech. 44 373-400
- [6] Berg H C 2004 Escherichia coli in Motion (New York: Springer)
- [7] Hegemann P and Bruck B 1989 Light-induced stop response in Chlamydomonas reinhardtii: occurrence and adaptation phenomena Cell Motil. 14 501–15
- [8] Hill N A and H\u00e4der D-P 1997 A biased random walk model for the trajectories of swimming micro-organisms J. Theor. Biol. 186 503–26
- [9] Vladimirov V A, Wu M S C, Pedley T J, Denissenko P V and Zakhidova S G 2004 Measurement of cell velocity distributions in populations of motile algae J. Exp. Biol. 207 1203–16
- [10] Polin M, Tuval I, Drescher K, Gollub J P and Goldstein R E 2009 Chlamydomonas swims with two gears in a eukaryotic version of runand-tumble locomotion Science 325 487–90
- [11] Rothschild L 1963 Non-random distribution of bull spermatozoa in a drop of sperm suspension Nature 198 1221-2
- [12] Winet H, Bernstein G S and Head J 1984 Observations on the response of human spermatozoa to gravity, boundaries and fluid shear J. Reprod. Fert. 70 511–23
- [13] Frymier P D, Ford R M, Berg H C and Cummings P T 1995 Three-dimensional tracking of motile bacteria near a solid planar surface Proc. Natl Acad. Sci. USA 92 6195–9
- [14] Berke A P, Turner L, Berg H C and Lauga E 2008 Hydrodynamic attraction of swimming microorganisms by surfaces *Phys. Rev. Lett.* 101 038102
- [15] Lauga E and Powers T R 2009 The hydrodynamics of swimming microorganisms Rep. Prog. Phys. 72 096601
- [16] Vigeant M A S, Ford R M, Wagner M and Tamm L K 2002 Reversible and irreversible adhesion of motile Escherichia coli cells analyzed by total internal reflection aqueous fluorescence microscopy Appl. Environ. Microbiol. 68 2794–801
- [17] Lauga E, DiLuzio W R, Whitesides G M and Stone H 2006 Swimming in circles: motion of bacteria near solid boundaries Biophys. J. 90 400–12
- [18] Giacche D, Ishikawa T and Yamaguchi T 2010 Hydrodynamic entrapment of bacteria swimming near a solid surface Phys. Rev. E 82 056309
- [19] Drescher K, Dunkel J, Cisneros L H, Ganguly S and Goldstein R E 2011 Fluid dynamics and noise in bacterial cell-cell and cell-surface scattering Proc. Natl Acad. Sci. USA 108 10940–5
- [20] Fauci L J and McDonald A 1995 Sperm motility in the presence of boundaries Bull. Math. Biol. 57 679-99
- [21] Smith D J, Gaffney E A, Blake J R and Kirkman-Brown J C 2009 Human sperm accumulation near surfaces: a simulation study *J. Fluid Mech.* 621 289–320
- [22] Kantsler V, Dunkel J, Polin M and Goldstein R E 2013 Ciliary contact interactions dominate surface scattering of swimming eukaryotes Proc. Natl Acad. Sci. USA 110 1187–92
- [23] Spagnolie S E, Wahl C, Lukasik J and Thiffeault J-L 2017 Microorganism billiards Physica D 341 33-44
- [24] Wioland H, Woodhouse F G, Dunkel J and Goldstein R E 2016 Ferromagnetic and antiferromagnetic order in bacterial vortex lattices Nat. Phys. 12 341–5
- [25] Takagi D, Braunschweig A B, Zhang J and Shelley M J 2013 Dispersion of self-propelled rods undergoing fluctuation-driven flips Phys. Rev. Lett. 110 038301
- [26] Takagi D, Palacci J, Braunschweig A B, Shelley M J and Zhang J 2014 Hydrodynamic capture of microswimmers into sphere-bound orbits Soft Matter 10 1784–9
- [27] Bricard A, Caussin J-B, Desreumaux N, Dauchot O and Bartolo D 2013 Emergence of macroscopic directed motion in populations of motile colloids Nature 503 95–8
- [28] Sipos O, Nagy K, Leonardo R D and Galajda P 2015 Hydrodynamic trapping of swimming bacteria by convex walls Phys. Rev. Lett. 114 258104
- [29] Spagnolie S E, M-Flores G R, Bartolo D and Lauga E 2015 Geometric capture and escape of a microswimmer colliding with an obstacle Soft Matter 11 3396–411
- [30] Hassink J, Bouwman L A, Zwart K B and Brussaard L 1993 Relationships between habitable pore and ce, soil biota and mineralization rates in grassland soils Soil Biol. Biochem. 25 47–55
- [31] Young I M and Crawford J W 2004 Interactions and self-organization in the soil-microbe complex *Science* **304** 1634–7
- [32] Or D, Smets B F, Wraith J M, Dechesne A and Friedman S P 2007 Physical constraints affecting bacterial habitats and activity in unsaturated porous media—a review Adv. Water. Resour. 30 1505–27
- [33] Carson J K, Gonzalez-Quinones V, Murphy D V, Hinz C, Shaw J A and Gleeson D B 2010 Low pore connectivity increases bacterial diversity in soil Appl. Environ. Microbiol. 76 3936–42
- [34] Vos M, Wolf A B, Jennings S J and Kowalchuk G A 2013 Micro-scale determinants of bacterial diversity in soil FEMS Microbiol. Rev. 37 936–54
- [35] Hosoi A E and Goldman D I 2015 Beneath our feet: strategies for locomotion in granular media Annu. Rev. Fluid Mech. 47 431-53
- [36] Majmudar T, Keaveny E E, Zhang J and Shelley M J 2012 Experiments and theory of undulatory locomotion in a simple structured medium J. R. Soc. Interface 9 1809–23
- [37] Morin A, Cardozo D L, Chikkadi V and Bartolo D 2017 Self-propulsion through disorder: localization transition in active Lorentz gases arXiv:1702.07655

- [38] Volpe G, Buttinoni I, Vogt D, Kummerer H-J and Bechinger C 2011 Microswimmers in patterned environments Soft Matter 7 8810–5
- [39] Brown A T, Vladescu I D, Dawson A, Vissers T, Schwarz-Linek J, Lintuvuori J S and Poon W C K 2016 Swimming in a crystal Soft Matter 12 131–40
- [40] Lighthill M J 1952 On the squirming motion of nearly spherical deformable bodies through liquids at very small reynolds numbers Comm. Pure Appl. Math. 5 109–18
- [41] Blake J R 1971 A spherical envelope approach to ciliary propulsion J. Fluid Mech. 46 199-208
- [42] Ishikawa T, Locsei J T and Pedley T J 2008 Development of coherent structures in concentrated suspensions of swimming model micro-organisms J. Fluid Mech. 615 401–31
- [43] Lebedev V I and Laikov D N 1999 A quadrature formula for the sphere of the 131st algebraic order of accuracy Doklady Mathematics 59 477–81
- [44] Happel J and Brenner H 1983 Low Reynolds Number Hydrodynamics (Dordrecht: Martinus Nijhoff)
- [45] Kim S and Karrila S J 2005 Microhydrodynamics: Principles and Selected Applications (Butterworth–Heinemann Series in Chemical Engineering) (New York: Dover)
- [46] Batchelor G K 1970 The stress system in a suspension of force-free particles J. Fluid Mech. 41 545–70
- [47] Blake J R 1971 A spherical envelope approach to ciliary propulsion J. Fluid Mech. 46 199–208
- [48] Batchelor G K 1967 An Introduction to Fluid Dynamics (Cambridge: Cambridge University Press)
- [49] Goldman A J, Cox R G and Brenner H 1967 Slow viscous motion of a sphere parallel to a plane walli motion through a quiescent fluid Chem. Eng. Sci. 22 637–51
- [50] Ishikawa T, Simmonds M P and Pedley T J 2006 Hydrodynamic interaction of two swimming model micro-organisms J. Fluid Mech. 568 119–60