Kinematics of the swimming of *Spiroplasma*

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*Spiroplasma* swimming is studied with a simple model based on resistive-force theory. Specifically, we consider a bacterium shaped in the form of a helix that propagates traveling-wave distortions which flip the handedness of the helical cell body. We treat cell length, pitch angle, kink velocity, and distance between kinks as parameters and calculate the swimming velocity that arises due to the distortions. We find that, for a fixed pitch angle, scaling collapses the swimming velocity (and the swimming efficiency) to a universal curve that depends only on the ratio of the distance between kinks to the cell length. Simultaneously optimizing the swimming efficiency with respect to inter-kink length and pitch angle, we find that the optimal pitch angle is 35.5° and the optimal inter-kink length ratio is 0.338, values in good agreement with experimental observations.

Helices are profoundly involved in the swimming of many types of bacteria. For example, the classical picture of bacterial swimming involves long helical filaments extending out from the inner membrane of the cell [1], driving the bacterium forward by their rotation. But helices are also used in other more ingenious ways. In *Treponema pallidum*, the spirochetal bacterium responsible for syphilis, rotation of helical filaments encased between the bacterial cell wall and the outer membrane leads to periodic undulations that drive this cell through water and other viscous fluids. (This is not, presumably, how it crossed the Atlantic Ocean [2].) The *Lep-tospirotaeaceae* rely on a helical body plan and rotation of tightly-coiled filaments to drive their motility [2]. Generally speaking, it is the dynamics and shape together, the “swimming strategy”, that is the key to understanding the motility of helical bacteria. For the flagella-less family of plant pathogens represented by *Spiroplasma*, the precise propulsive strategy was shrouded in mystery until careful observations [3, 4] revealed that these tiny organisms processively flip the chirality of regions of their helical bodies, generating pairs of kinks moving down the cell body (Fig. 1(a)). The internal propulsive mechanism underlying this peculiar swimming strategy is still unclear, as is the quantification of its motility.

*Spiroplasma* swimming is realized by the propagation of a pair of kinks along the body axis of the cell, and the bacterium is propelled by the hydrodynamic force as the fluid associated with the kinks moves rearward with the “body wave”. Kinks start at the same end of the cell (front), and travel toward the other end (back). As the kink propagates, the cell changes direction through an angle related to the pitch angle of the helix. Because of the unbalanced viscous drag between different portions of the cell body separated by kinks, eventually the cell swims in a zigzag path. The helical transformation requires a cell to rotate about its body axis during kink propagation [3, 4, 5].

In this Letter we use the experimentally-observed geometry of *Spiroplasma* to explore its swimming behavior for a range of kinematic deformations. We consider a helically-shaped cell of fixed pitch angle and helix diameter. To swim, the bacterium flips the handedness of its helix beginning at one end. This change in helicity propagates toward the back of the cell with a uniform velocity, \( U_k = 10.5 \pm 0.3 \, \text{µm/s} \) [6]. After a time \( t_k \), the front of the cell reverts to its original handedness and this change in helicity also propagates toward the back of the cell with the same kink velocity. The distance between the kinks is \( L_k = U_k t_k \), and only an integer number of turns is considered. We treat \( U_k \), \( L_k \), and the cell length, \( L_c \), as parameters and use resistive-force theory (RFT) to calculate the swimming speed and trajectory as a function of these parameters. Our model is complementary to the work by Wada and Netz [8] where slender-body hydrodynamics [9] was used to explore the dependence of swimming speed on the pitch angle, \( \psi \). By varying the distance between kinks, we examine both the geometry and kinematics of the swimming process. It was found experimentally that the kink velocity \( U_k \) [4, 5] and the inter-kink distance \( L_k \) [6] show a remarkable consistency — what determines this preference? Our results demon-

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strate that the maximum propulsive efficiency is achieved when the pitch angle is around 35.5° (in agreement with [8]) and the ratio between inter-kink length and cell-body length is around 0.338, which is consistent with most experimental observations and suggests that the kinematics of *Spiroplasma* has been evolutionarily tuned.

Any helix can be described by its pitch, $p$, and helix radius, $r$. One helical repeat in the arclength $s$ is $\ell = \sqrt{p^2 + 4\pi^2 r^2}$. When two helices of differing pitch are concatenated, such that the tangent and normal vectors are continuous, a kink is formed [6, 10]. The bend angle $\alpha$ between the two axes in this construction is $\alpha = \pi - 2\psi$ (Fig. 1(b)). The parameters employed in the calculations are $p = 0.62 \mu m$, $r = D/2 = 0.069 \mu m$, and $\psi = 35^\circ$ [4, 5]. For the numerical integration, the cell body is discretized into straight-line elements. According to RFT [11, 12], the resistance force acting on a unit length of rod is given by $f = -[2\pi\mu/\ln(2/\epsilon)] (2I - \lambda \lambda) \cdot \nabla + O \left[\ln^{-2}(2/\epsilon)\right]$, where $\nabla$ is the local velocity, $I$ is the unit tensor, $\epsilon$ is the slenderness ratio of rod diameter to length, $\lambda$ is the unit tangent vector, and $\mu$ is viscosity. The total force and momentum on the unrestrained cell body is zero, i.e., $\int_0^L f(s) ds = 0$, $\int_0^L M(s) ds = 0$. The translational and rotational velocities of the cell body can be solved for from the resulting system of linear equations. To verify that the implementation of the model has been properly made, the translational velocity of a rigid helix with angular velocity $\omega$ around its body axis is calculated with the same numerical code. Since the total force on the rotating helix is zero, i.e., $\int_0^L f(s) ds = 0$ ($L$ is the total length of the rigid helix), the translational velocity of the helix is $U = B\ell \omega$, where $B = \sin^2 \psi \cos \psi / [2\pi(1 + \sin^2 \psi)]$ [8]. The numerical results match this resistive-force solution with relative errors less than 0.05% under our typical spatial discretization. A slender-body theory (SBT) calculation, using the straight line elements, was also performed to account for hydrodynamic interactions. Based on the aforementioned model, a number of dynamical questions on the swimming of *Spiroplasma* can be formulated and addressed.

Typical experimental and simulated displacement curves during one swimming cycle of *Spiroplasma* are shown in Fig. 2(a). Here the displacement is defined as the projected center-of-mass position. For comparison with the experimental data, our numerical simulation results are also smoothed with a 175-ms boxcar filter [8, 13]. Using the parameters given above, our simulations agree well with the corresponding experimental results [14], giving an average velocity of about 3 $\mu m/s$, close to the observed cell velocity, 3.3±0.2 $\mu m/s$ [5].

It is observed experimentally that the speed is decreased when two kinks are present as compared with when only one kink is present. This key feature of the displacement curve is reproduced in the numerical simulations (Fig. 2(a)). Our simulations show that the cell velocity increases linearly with kink velocity, and that the ratio of cell velocity to kink velocity is constant for fixed inter-kink length, with the proportionality constant varying with inter-kink length, consistent with the experimental reports [4, 5].

In order to delineate the effect of dynamically changing geometry on *Spiroplasma* motility, two dimensionless parameters are introduced. One is the ratio of cell velocity $U_c$ to kink velocity $U_k$; the other is the ratio of inter-kink length $L_k$ to cell body length $L_c$. We find that all the data fall on a single curve after the results from different cell-body lengths and inter-kink lengths are taken together (Fig. 2(b)). The curve describes a velocity ratio that increases with length ratio at small values, and then begins to decrease with length ratio after reaching a peak value. From Fig. 2(b), the maximum velocity ratio of the data-fitted curve is around 0.323 when the length ratio is about 0.32. The fact that the velocity ratios and length ratios observed in the experiment fall close to the aforementioned value suggests that wildtype *Spiroplasmas* have optimized their geometry to achieve the “fastest state” in the course of evolution. The study of Wada and Netz [8] suggests an optimized pitch angle as well, the treatment of which goes beyond the reach of RFT for pitch angles greater than roughly 35° [13, 16]. The full optimization over both length ratio and pitch angle (Fig. 3) using our SBT code gives a maximum velocity ratio at $L_k/L_c = 0.338$ and $\psi = 35.5^\circ$. These values are
consistent with the experimental values 0.303±0.051 an
34.8±1.4° respectively [3].

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FIG. 3: Velocity ratio as a function of kink length ratio and
pitch angle. Contour lines are velocity-ratio isolines. Exper-
imental observations are centered within the error ellipse.

The velocity ratio can be calculated in RFT in the
limit that the length ratio approaches one. As the length
ratio goes to one, there is effectively only one kink prop-
gagating along the cell body at any given time. In this
case, Spiroplasma can be treated as a filament consisting
of two helical sections, one right handed and the other
left handed. The two helical sections rotate around their
respective axes with the opposite sense. The total trans-
lational velocity is the vector sum of these two com-
ponents with an interior angle 2ψ. From geometrical con-
siderations, the kink velocity U_k can be represented as
a function of rotational velocity ω, i.e., U_k = 2ωr/ tan ψ.
Using the previously-mentioned translation velocity of a
rigid helix (Bℓω) the total velocity can be estimated to
be 2Bℓω cos ψ, and the corresponding velocity ratio re-
duces to 2πB = sin^2 ψ cos ψ / (1 + sin^2 ψ). For a pitch
angle of 35°, this ratio is around 0.202, not far from the
simulation result of 0.24. At a length ratio of one, the
kinematic asymmetry between the left-handed inter-kink
length and the right-handed extra-kink length vanishes,
and a chirality variable χ ≡ (Lc − Lk) / (Lc + Lk) can be
used to study the velocity ratio on both sides of χ = 0
[17]. We note in passing that the entire velocity-ratio
curve is in excellent empirical agreement with a quartic
polynomial [18]

U_c/U_k ≈ 0.24 + 0.73φ^2 − 1.61φ^4 \quad (1)

(the solid curve in Fig. 2(b)) where φ = tan^{-1} χ.

Not only translation, but also rotation of the cell body
must satisfy the dynamic balance, i.e., zero force and
torque. At the beginning of a kink cycle, one end of the
cell-body axis traces out a circular arc (see Fig. 3). The
maximum angle swept out, we term the envelope angle.
We use this concept in Fig. 2(b), where the dashed curve
shows that the envelope angle increases with length ratio.
Fig. 2(b) also shows that the velocity ratio achieves its

Experiments on Spiroplasma do not observe more than
two kinks along the cell body at one time. It is there-
fore interesting to consider the swimming efficiency as
a function of the number of kinks per cycle in order to
determine whether a two-kink cycle is optimal. A stan-
ard definition of swimming efficiency compares the en-
ergy dissipated by dragging the appropriate rigid cylin-
der through fluid to the power dissipated by the actual
swimmer [8, 12, 19]. This definition compares a swim-
ning object to a non-swimmer and also does not account
for the energy dissipated to create the swimming motion.
Since an unknown mechanism is responsible for produc-
ing the traveling kinks in Spiroplasma, we assume that
there is a fixed net energy, δE, required to produce a pair
of kinks [21] and that this mechanism sets a uniform kink
velocity. Under these assumptions, we hypothesize that
biology tries to maximize the distance traveled per unit
energy expended. Therefore, we consider the fuel mileage
γ = distance traveled per energy consumed (nδE + the
energy dissipated by the fluid, where n is the number
of kink pairs per cycle), which we compute in terms of
nanometers traveled per ATP burned [21]. First, ignor-
ing the energy to create the kinks (δE ≃ 0), we calculate
γ as a function of the length ratio (solid curve in Fig. 5).
The fuel mileage from different cell lengths can all be
collapsed onto one curve by constructing a scaling fac-
tor from a power of the slenderness ratio ε (= d/Lc) as
ε^{-δ}γ(L_k, L_c) with δ ≃ 2. In both RFT and SBT, we
find that the fuel mileage is maximized at a length ratio
of around 0.3, although the methods differ. The vehicle
different velocities lead to shifted γ values: γ_{RFT}/γ_{SBT} ≃ 3 − 3.4. Using
RFT, we find that a single double kink cycle requires
50-100 ATPs worth of energy and the cell moves about
26 nm/ATP, whereas SBT gives 150-340 ATPs per cycle
and the cell moves 8 nm/ATP. Interestingly, the latter is
the fuel mileage of a single kinesin molecule [22]. When
δE is small, we find that the fuel mileage for a 4-kink cy-
cle is larger than for a 2-kink cycle. However, when δE
is greater than 40 ATPs, then a 2-kink cycle is optimal.
Therefore, we expect that the mechanism producing the

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FIG. 4: The trace swept out by the cell-body axis at eight
different inter-kink lengths 2-9 ℓ (for cell length 10 ℓ). Three
snapshots of the cell body are superposed in the leftmost trace
(L_k = 2ℓ). Yellow lines indicate the trajectories of the two
kinks. The envelope angle is indicated.
double kink must require at least 40 ATPs per cycle.

In this Letter, we investigated via a numerical study the swimming of *Spiroplasma* with a model based on resistive-force theory and the kinematic deformations of *Spiroplasma*. The key features of *Spiroplasma* swimming are well reproduced with this model, and the simulation results agree quantitatively with the observed experimental data. The important effects of dynamic geometry on the motility of *Spiroplasma* are explored, and the motility efficiency is shown to assume the highest possible value given the kinematics observed in experiments. One implication is the optimization of the geometry and kinematics of *Spiroplasma* during the course of evolution.

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[13] The boxcar filter uses $S_{BC}(t) = \frac{1}{\sigma^2} \int_{t-\sigma}^{t+\sigma} S(t') dt'$, where $S$ is the displacement before filtering, $S_{BC}$ is the displacement after filtering, and $\sigma$ is the half width.

[14] The RFT results fit best with experimental data obtained in a high-viscosity medium rather than water. This suggests a possible explanation for the phenomena that *Spiroplasma* swims with higher speed in more viscous media. At low viscosity, Brownian motion randomly alters the orientations along the cell body, which could reduce the net propulsive force generated by the traveling kinks. The effect of these fluctuations on swimming speed may be reduced in higher-viscosity media.

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[16] A more tightly-wound helix means a more dominant hydrodynamic interaction. Jung et al. [15] found that RFT breaks down for values $2\pi R/P > 0.7$ ($R$ = superhelical radius, $P$ = superhelical pitch), which can be mapped onto our system as follows: superhelical turns $\rightarrow$ kinks, $\arctan(2\pi R/P) \rightarrow (\pi - \alpha)/2 \approx \psi$.

[17] Though $L_k > L_c$ is physically nonsensical per se, the solutions of the Stokes equations behave smoothly as inter-kink and extra-kink labels are exchanged at $L_k = L_c$, i.e. under the mapping $\chi \rightarrow -\chi$. This sign change is equivalent to a spatial inversion, under which the symmetries of the Stokes equations imply that the cell velocity is unchanged. Hence, the velocity ratio is a smooth, even function in $\chi$, and about $\chi = 0$ a best fit gives $U_c/U_k = 0.24 + 0.73x^2 - 2.04x^4 + O(x^6)$.

[18] The corresponding velocity ratio from SBT has a similar fit $U_c/U_k \approx 0.165 + 0.73x^2 - 1.82x^4$, to leading order a shift of 0.075 from Eq. (1).

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