A fully three-dimensional model of the interaction of driven elastic filaments in a Stokes flow with applications to sperm motility

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ABSTRACT

In many animals, sperm flagella exhibit primarily planar waveforms. An isolated sperm with a planar flagellar beat in a three-dimensional unbounded fluid domain would remain in a plane. However, because sperm must navigate through complex, three-dimensional confined spaces along with other sperm, forces that bend or move the flagellum out of its current beat plane develop. Here we present an extension of previous models of an elastic sperm flagellar filament whose shape change is driven by the pursuit of a preferred curvature wave. In particular, we extend the energy of the generalized elastica to include a term that penalizes out-of-plane motion. We are now able to study the interaction of free-swimmers in a 3D Stokes flow that do not start out beating in the same plane. We demonstrate the three-dimensional nature of swimming behavior as neighboring sperm swim close to each other and affect each others’ trajectories via fluid-structure coupling.

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1. Introduction

For successful fertilization, sperm must be able to navigate complex environments to reach the egg. Thus, sperm motility patterns are vital components of sexual reproduction. In many species, sperm flagellar waveforms are planar or near-planar, though helical waveforms are also stable (Vernon and Woolley, 2004; Woolley and Vernon, 2001). Planar waveforms of an isolated beating flagellum result in trajectories that are characterized by either linear or circular paths within the plane, depending on the symmetry of the beat pattern (Phillips, 1972; Yanagimachi, 1970). However, forces exerted on the planar swimmer by proximal surfaces or nearby swimmers can easily bend or pull part or all of the flagellum out of its current beat plane.

Three-dimensional behavior of sperm flagella has been experimentally observed (Woolley and Osborn, 1984) and some species exhibit “Fig. 8” or helical deviations of the waveform out of the plane (Cosson et al., 2003; Woolley and Vernon, 2001). More recent experimental advances in tracking sperm at a high resolution have exposed the three-dimensional nature of swimming trajectories of sperm in populations, providing experimental evidence for helical or “chiral ribbon” trajectories (Corkidi et al., 2008; Su et al., 2012, 2013). The recent review of Guerrero et al. (2011) highlights the importance of understanding sperm motility in fully three-dimensional settings in order to correctly characterize and assess fertility in sperm.

The hydrodynamics of sperm motility is characterized by low Reynolds number, viscous fluid mechanics, and has been the inspiration of many classic studies in biological fluid dynamics (Fauci and Dillon, 2006), including Taylor’s swimming sheet (Taylor, 1951) and the resistive force theory of Gray and Hancock (1955). In these early models, the kinematics of the flagellar waveform were imposed, and the resulting swimming speed of the flagellum was calculated. Recognizing that a sperm flagellum is an elastic structure whose real behavior is determined by its coupling to the surrounding viscous fluid, models have been developed that treat the flagellum as a generalized Euler elastica, whose shape change is driven by the pursuit of a preferred curvature wave (Fauci and Peskin, 1988). Such flagellar models, essentially of swimming sheets, have been coupled to a two-dimensional fluid and have been used to study synchronization of finite swimmers and their attraction to walls (Fauci and McDonald, 1995), and the effect of viscoelasticity on flagellar swimming (Chisnell et al., 2013; Taran et al., 2010). More recently, these driven-elasctics have been coupled to a model for three-dimensional Stokes flow and have been used to study the emergence of a hyperactivated waveform due to calcium dynamics in a single, isolated flagellar filament (Olson et al., 2011), and the elastic attachment of a planar flagellar filament to an orthogonal planar wall (Simons et al., 2014). While the dynamics of these filaments were coupled to a three-dimensional domain of fluid, the symmetry imposed by a single flagellum confined to a plane in free-space or one beating perpendicular to an infinite planar wall does not result in any out-of-plane component of force on the flagellum.
Motivated by the understanding sperm–sperm and sperm–
boundary interactions in more general settings, even for driven
filaments that pursue a planar waveform, here we extend the
generalized Euler elastica energy formulation of Faucci and Peskin
(1988) to include a penalty for out-of-plane motion. The native
flagellar plane will evolve in time. The energy that drives this new
model is rotation- and translation-invariant, therefore the filament
will be a free-swimmer that generates zero total force and torque.

While several different mathematical approaches may be used
to couple flagellar forces to a fluid, here we use the 3D method of
regularized Stokelses (Cortez et al., 2005). In the following
sections, we present the mathematical formulation of the three-
dimensional model, how to incorporate multiple nearby sperm,
and then discuss the implications for three-dimensional trajec-
tories and behaviors in sperm populations.

2. Methods

2.1. Derivation of the three-dimensional model

We model the sperm flagellum as a three-dimensional curve \( \mathbf{X}(s, t) \) that exerts
forces along its length in a viscous fluid governed by the incompressible Stokes
equations:
\[
\nabla \cdot \mathbf{u} = 0 \\
\mu \nabla \mathbf{u} = -\nabla p + \nabla \cdot \mathbf{f}(\mathbf{X}(s, t), \mathbf{u}(\mathbf{X}(s, t))) ds
\]
(1)
where \( L \) is the total length of the flagellum, \( s \) is the arc length, \( t \) is time, \( \mu \) is the
dynamic viscosity, \( \mathbf{u} \) is the fluid velocity, and \( p \) is the pressure. The forces \( \mathbf{f}(\mathbf{X}, t) \)
exerted by the flagellum upon the surrounding fluid are regularized by a blob
function \( \phi_b \) defined by
\[
\phi_b(r) = \frac{15\delta^3}{8\pi r^4 + \epsilon^4}
\]
This is a regularized delta function that distributes the force in a small fluid volume
around the curve (Cortez, 2001; Cortez et al., 2005). This regularization eliminates
singularities in the velocity field and the parameter \( \epsilon \) is chosen to be on the order of the
diameter of the sperm flagellum. The flagellum will be considered an immersed
boundary that moves with the fluid velocity at all points along its length.

Our model assumes that sperm flagella have a nearly planar beat. Both helical
and planar flagellar beat forms may be stable. Planar waveforms may be more
prevalent near surfaces and observed helical waveforms may in fact be more aptly
described as “flattened helices” (Brokaw, 1965; Rinkenspoel, 1965; Woolley and
Vernon, 2001). The planar or quasi-planar nature of the sperm flagellum is most
likely controlled by the structure of the microtubule doublets and dynein bridges of
the axoneme (Gibbons et al., 1985; Inaguma et al., 1988; Lin et al., 2012; Lindemann
et al., 1992; Ono et al., 1990; Shingyoji et al., 1991). We do not seek to model the
underlying structure of the flagellum nor how a planar preference might arise, but
instead conceptualize the sperm flagellum as a thin filament that would naturally
have a planar shape in the absence of external forces.

This extended model is formulated so that a flagellum can accommodate three-
dimensional effects that cause deviations from planar configurations. These
effects could include surfaces, other sperm swimming nearby, or other external forces.
To accomplish this, we incorporate forces that restrict out-of-plane deviations of
the flagellum. Within the flagellar plane, we prescribe curvature-based forces along the
flagellum as in Faucci and Peskin (1988), which enable the sperm to move in a wave-
like manner.

The forces \( \mathbf{f} \) will be derived using a continuous energy formulation that is an
extension of the internally actuated filament model in Faucci and Peskin (1988). In
order to define the energy, we first define a local coordinate system in which the
flagellar plane is \( z = 0 \). This will be referred to as the flagellum frame of reference. In
this frame of reference, bending forces that modulate the swimming behavior of
the flagellum will occur in the plane \( z = 0 \) alone. Moreover, non-planar deviations can
be understood as deviations in the \( z \) component and forces restricting these
deviations can be found. For clarity, in the flagellum frame of reference, we refer to the
Cartesian coordinates as \( \hat{x}, \hat{y}, \hat{z} \) and \( \alpha \) and \( \beta \) and the coordinates of the flagellum itself as
\( X \).

Thus the flagellar plane is the plane \( z = 0 \). Using this notation, we define the total energy as
\[
\mathcal{E}(X, t) = \mathcal{E}_{\text{bend}}(X, t) + \mathcal{E}_{\text{tens}}(X, t) + \mathcal{E}_{\text{plasm}}(X, t)
\]
where \( \mathcal{E}_{\text{bend}} \) is a tensile energy term that enforces inextensibility, \( \mathcal{E}_{\text{tens}} \) is a time-
dependent bending energy that drives the beating behavior of the filament, and
\( \mathcal{E}_{\text{plasm}} \) is the energy arising from out-of-plane deviations. Each energy is defined as
\[
\mathcal{E}_{\text{bend}}(X, t) = \frac{1}{2} S \int_0^L \left( \left\| \frac{\partial \mathbf{X}}{\partial s} \right\| - 1 \right)^2 ds
\]
(2)
\[
\mathcal{E}_{\text{tens}}(X, t) = \frac{1}{2} S \int_0^L \left( \frac{\partial \mathbf{X}}{\partial s} \times \frac{\partial^2 \mathbf{X}}{\partial s^2} \right) \cdot \mathbf{e}_1 - C(\alpha, t) \right)^2 ds
\]
(3)
\[
\mathcal{E}_{\text{plasm}}(X, t) = \frac{1}{2} S \int_0^L \left( \frac{\partial \mathbf{X}}{\partial s} \right)^2 ds
\]
(4)
where \( S \) are stiffness coefficients, \( L \) is the length of the flagellum, and \( \mathbf{e}_1 = (0, 0, 1)^T \)
is the normal to the flagellar plane. The function \( C(\alpha, t) \) is the preferred curvature
that drives the swimming motion of the sperm. Since this function is restricted to the plane \( z = 0 \), the parameter \( \alpha \) refers to the projection of the filament arc length onto that plane, and \( \alpha \) is the total projected arc length in that plane. We take \( \alpha \) to be approximately constant because deviations from the plane are assumed to be small.

With these definitions, the energies (2) and (3) are the same energies described in
Faucci and Peskin (1988) and the energy (4) is the new addition to our model. The energy (4) is non-zero when there are local deviations of the flagellum in the \( \hat{z} \) direction (i.e. deviations from the flagellar plane). The total energy \( \mathcal{E} \) is translation- and rotation-invariant and forces in the flagellum frame of reference
\( f \) can be calculated as
\[
f = -\frac{\partial \mathcal{E}}{\partial \mathbf{X}}
\]
These resulting forces seek to minimize the total energy, where the tensile, bending and
planarity stiffness constants \( \{S, S_p, S_c\} \) determine how closely the preferred planar traveling wave is realized. In particular, the tensile stiffness \( S \) is chosen large enough so that the filament is effectively inextensible, the bending stiffness \( S_p \) reflects the bend modulus of the flexible filament, and the planarity stiffness \( S_c \) controls the penalty for deviation from the plane. Finding the forces \( f \) in the original frame of reference requires a transformation from the \( (\hat{x}, \hat{y}, \hat{z}) \) coordinates to the global Cartesian coordinates \((x, y, z)\).

For computational purposes, we will discretize the integral in (1) by defining
discrete points \( X_k \) along the flagellum (where \( k \) denotes the index of the point). Fig. 1
shows a schematic of the physical domain of the discretized flagellum for a preferred sinusoidal waveform with amplitude \( b \).

Using the method of regularized Stokelses described in Cortez et al. (2005), the
fluid velocity is given by
\[
\mathbf{u}(x) = \sum_{k} \frac{r_k^2 + 2x^2}{4(r_k^2 + x^2)^3/2} \mathbf{A}_k \cdot (\mathbf{X}_k - \mathbf{x}) / \left( \frac{r_k^2 + x^2 + \epsilon^2}{4} \right)^{3/2}
\]
where \( r_k = |\mathbf{X}_k - \mathbf{x}| \) and \( \epsilon \) is set to be close to the flagellum diameter. The points
will be discretized so that their spacing is less than \( \epsilon \). The discrete forces \( f_k \) define
the forces the flagellum exerts at points \( X_k \), and are calculated from discretized
versions of the energies (2)-(4). The motion of the flagellum points is described by the
following equation:
\[
\frac{\partial \mathbf{X}_k}{\partial t} = -\nabla \mathbf{u}(X_k, t)
\]
(5)
which is approximated using a forward Euler method. In the next several sections, we
describe the specific components of the three-dimensional preferred
curvature model.

2.2. Derivation of the discrete flagellar plane bending energy

We discretize the bending energy (3), as in Faucci and Peskin (1988):
\[
\mathcal{E}_{\text{bend}, \text{disc}} = \frac{1}{2} \sum_{k=2}^{N-1} \left( \frac{\hat{X}_{k-1} - \hat{X}_k}{\Delta t} \cdot \left( \hat{X}_{k+1} - \hat{X}_{k+2} - \hat{X}_{k-1} \right) \right)^2 / \Delta t
\]
(5)
where \( \hat{X}_k \) is the preferred curvature (in the work of Faucci and Peskin (1988))
and \( \mathcal{E}_{\text{bend}, \text{disc}} \) is the bending energy calculated using the preferred
sinusoidal waveform with amplitude \( b \).
where $S_b$ is the bending stiffness constant, $N$ is the total number of points along the flagellum, and $\Delta t$ is a constant equal to the desired spacing between the points $(x_k, y_k, 0)$. The preferred curvature $C_f(t)$ is a time-dependent sinusoidal function similar to the one used in Fauci and Peskin (1988):

$$C_f(t) = k^b \sin(\omega t - \alpha t)$$

(6)

where $\alpha t$ is the arc length distance along the flagellum (projected onto the plane $\hat{z} = 0$) from the head point $(\hat{x}, \hat{y}, 0)$ up to the point $(\hat{x}_k, \hat{y}_k, 0)$, and $k$ and $\omega$ are the wavenumber and frequency parameters, respectively. In this model, we choose $k = 2\pi/\Delta z$ so there is approximately one wavelength per flagellum length, but our results are not qualitatively sensitive to this choice.

Small out-of-plane deviations of the flagellum imply that $\Delta z$ may be smaller but not too different from $\Delta t$ (the desired spacing between points in 3D). In order not to overestimate the desired planar curvature forces for three-dimensional flagellar configurations, we consider the three-dimensional vector of length $\Delta s$ given by

$$r_{k+1} = \Delta s \hat{X}_{k+1} - \hat{X}_k$$

and define $\Delta t_{k+1}$ as the length of the projection of vector $r_{k+1}$ onto the flagellar plane:

$$\Delta t_{k+1} = |r_{k+1} - (e_1 \cdot r_{k+1}) e_1|$$

With this definition, the factors of $\Delta s^2$ and $\Delta t$ in Eq. (5) are replaced with

$$\Delta s^2 \to \Delta s \Delta t_{k+1} + \Delta t_{k+1} \Delta t_{k+1} \frac{\Delta t_{k+1} + \Delta s^2}{2}$$

$$\Delta t \to \Delta t_{k+1} + \Delta t_{k+1} \frac{\Delta s^2}{2}$$

(7)

We note that $\Delta s = \Delta t$ for all $k$ when the points are exactly in the plane $\hat{z} = 0$.

### 2.3. Plane restriction force

To address out-of-plane deviations, we discretize the energy (4) as

$$E_{plane} = \frac{1}{2} \sum_{k=1}^{N} \sum_{i=1}^{j} \left( \left[ \hat{z}_{k+1} - \hat{z}_k \right]^2 - \Delta s^2 \right).$$

This energy formulation restricts any non-planar deviations via forces in the $\hat{z}$-direction but is a local effect that seeks to minimize the total quadratic variation in $\hat{z}$, hence reducing the concavity of $z(k)$. The force derived from this energy acting upon an interior point $k \in (2, ..., N-1)$ would then be

$$f_z = S_p \Delta s \hat{e}_z$$

(8)

Because the energy function $E_{plane}$ like the bending and tensile energies, is rotationally and translationally invariant, the net force and the net torque due to $E_{plane}$ satisfy:

$$\sum_{k=1}^{N} \hat{X}_k \times f_z = 0.$$

This can be proven by direct calculation from (8) and is due to Noether’s Theorem (Arnold, 1989). These two conditions on the net force and net torque are constraints required for self-propelled organisms.

### 2.4. Incorporating asymmetry

Motility patterns in sperm are typically characterized as active (symmetric) or hyperactive (asymmetric with high amplitude). Because hyperactivity is required for successful fertilization of the egg (Suarez et al., 1991; Suarez, 2008; Vanagovich, 1970), we consider both asymmetric and symmetric preferred beat-forms. To incorporate asymmetric into this model, we use a simple amplitude modulation approach as in Olson et al. (2011) to set the preferred curvature function $C_f(t)$ in expression (6) along the flagellum for all times $t$. For our flagellum of length 100 $\mu$m, we let the amplitude $b$ of the curvature function (6) take the following values:

$$b = \begin{cases} b_1 & \text{if } \sin(\omega t - \alpha t) < 0 \\ b_2 & \text{if } \sin(\omega t - \alpha t) \geq 0 \end{cases}$$

(9)

In the case of a symmetric waveform, we choose $b_1 = b_2 = 10 \mu$m. To explore the effect of a weak asymmetry in curvature, we choose $b_1 = 10 \mu$m and $b_2 = 15 \mu$m. For a stronger, higher amplitude asymmetry we choose $b_1 = 10 \mu$m and $b_2 = 25 \mu$m. The stronger asymmetry parameters have been chosen such that the behavior is closer to the deeper bends characteristic of hyperactive sperm motility patterns (see previous models in Olson et al., 2011; Simons et al., 2014).

### 2.5. Defining the flagellum frame of reference

At each time step, we need to update the preferred flagellar plane and flagellar frame of reference. We use the method of least squares, as described in Eberly and Geometric Tools (2014), to define this plane. Given discrete points along the flagellum $X_k$, we find the center of mass of the flagellum, denoted by $\hat{X}$, as simply the average position of all points $X_k$. We define the flagellar plane as the plane that minimizes the sum of the squares of the orthogonal distances between the centered points $X_k - \hat{X}$ and $(y, z)$ and the plane. The plane is represented by the vector $\hat{v} = (a, b, c)$ corresponding to least squares fit plane $ax + by + cz = 0$. We note that because the points $(x, y, z)$ have zero average, this plane must go through the origin by construction.

The vector $\hat{v}$ defines the normal to the flagellar plane. This is equivalent to the $\hat{z}$ coordinate direction in the local (Lagrangian) flagellum frame of reference. Noting that a plane splits the space into two half-spaces, care is taken to ensure that $\hat{v}$ is always pointing towards the same half-space. This is important because the preferred curvature model in Fauci and Peskin (1988) requires the ability to discern the sign of the curvature. In other words, we must be able to choose between a plane normal vector pointing in the positive or negative $\hat{z}$ direction. Since our model is designed to represent a continuous process and we use a small numerical time step, the flagellum plane should not change significantly between time steps. Therefore, we add a check point in our algorithm that verifies that the normal vector from the previous time step and the normal vector from the current time step have a positive dot product.

Because forces within the flagellar plane are invariant under rotation and translation, we may choose any $\hat{x}$ and $\hat{y}$ directions that would result in an orthonormal triad with $\hat{v}$ with the appropriate “right-hand rule” relationship between the coordinate directions. Fig. 2 shows an example of a three-dimensional flagellar configuration, the least squares plane fit to the centered data, as well as the flagellum frame of reference.

### 2.6. Computational method

In all simulations, each flagellum is initialized with a non-collinear configuration at time step $j=0$ to ensure that a unique flagellar plane can be defined. The method for computing flagellar positions, forces, and velocities over time is as follows:

1. Calculate center of mass of flagellum points $X_k$ and translate data so they have a center of mass at the origin.
2. Using the method of least squares, find the flagellar plane for centered flagellar data. The normal to this plane is represented by the normal vector $\hat{v}$.
3. For time step $j > 0$, if $\hat{v} \cdot \hat{v}_{j-1} < 0$, let $\hat{v} = -\hat{v}$ (signed curvature checkpoint).
4. Define $X_k$ using the flagellum frame of reference determined by $\hat{v}$.
5. Find forces $f_k$ for the points $X_k$ based on the total energy $\epsilon$.
6. Rotate forces $f_k$ back to original frame of reference to obtain $f_k$.

![Image](http://dx.doi.org/10.1016/j.jbiomech.2015.01.050)
7. Solve for velocities $\mathbf{u}_k$ using the method of regularized Stokeslets and forces $\mathbf{f}_k$, as described in Olson et al. (2011).
8. Move all points $X_k$ by a forward Euler time step with velocities $\mathbf{u}_k$.
9. Let $j = j + 1$ and repeat.

### 2.7. Multiple sperm

We are interested in how sperm trajectories are influenced by nearby neighbors. The model proposed here can consider any three-dimensional flagellum configuration with multiple sperm present. However, for realistic scenarios with multiple sperm, the model must not allow flagellum points to overlap. To address this, we add a repulsive force that acts on points that come within a certain distance of each other.

The repulsive force acting on the point $X_k$ by the point $X_j$ on separate flagella will take the following form:

$$f_k^g = S \left( \frac{1}{d} \frac{1}{\min(r, d)} \right) (X_k - X_j) \quad (10)$$

where $r = |X_k - X_j|$ and $d$ is a fixed distance. We choose this form because it is continuous in $r$ and it tends to infinity as $r$ approaches zero, which ensures that points do not get too close to one another if the time step is small. The choice of parameter $S$ is arbitrary, but should be chosen so that the repulsion forces do not get too large too quickly (compared to the curvature forces) as $r$ falls below $d$. The forces $f_k^g$ will be added to the forces $f_k$ before solving for velocities in the computational method. Note that contributions from this repulsion force maintain force- and torque-free swimming.

### 3. Results

Here we explore the effects of extending the elastica model of driven filaments to include an energy contribution for out-of-plane motion. In particular, consider the cases of both a single planar flagellum swimming in 3D free-space, and a pair of coplanar flagella pursuing waves in that plane, whose initial configurations are all

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Value</th>
<th>Remark/reference(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\epsilon$</td>
<td>Regularization parameter</td>
<td>1.3 $\mu$m</td>
<td>See Cortez (2001)</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Diameter of sperm flagellum</td>
<td>0.5 $\mu$m</td>
<td>Serres et al. (1984)</td>
</tr>
<tr>
<td>$\Delta s$</td>
<td>Viscosity (water)</td>
<td>$10^{-3}$ kg m$^{-1}$ s$^{-1}$</td>
<td>Cummins and Woodall (1985)</td>
</tr>
<tr>
<td>$L$</td>
<td>Spatial (arc length) discretization</td>
<td>1 $\mu$m</td>
<td>Ohmuro and Ishijima (2006), Smith et al. (2009)</td>
</tr>
<tr>
<td>$K$</td>
<td>Flagellum length</td>
<td>100 $\mu$m</td>
<td>Ohmuro and Ishijima (2006), Smith et al. (2009)</td>
</tr>
<tr>
<td>$\omega$</td>
<td>Frequency</td>
<td>$2\pi$ (10 Hz)</td>
<td>Simons et al. (2014)</td>
</tr>
<tr>
<td>$S_t$</td>
<td>Tensile stiffness</td>
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<td>This work</td>
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<tr>
<td>$S_p$</td>
<td>Planar restriction stiffness</td>
<td>0.001–1 pN $\mu$m$^{-3}$</td>
<td>This work</td>
</tr>
<tr>
<td>$S_b$</td>
<td>Planar bending stiffness</td>
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<td>Olson et al. (2011)</td>
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<td>$d$</td>
<td>Repulsion length</td>
<td>3 $\mu$m</td>
<td>Set to be $&gt; 2 \delta$</td>
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Fig. 3. Initial configurations of sperm flagella. In simulations with a single sperm, the black curve in (a)–(c) is used, and has a fully three-dimensional configuration. For two sperm coplanar simulations, we consider the flagella in panels (a)–(c), and project the black curve in (a)–(c) onto the plane $z = 0$ so both sperm are in the plane $z = 0$. For two sperm simulations with a non-planar perturbation, we use the red and black curves as shown in panels (d)–(f). For two sperm with initial parallel flagellar planes, we use the configurations in panels (d)–(f). All units are in $\mu$m. These small amplitude initial configurations are chosen for simplicity, and the sperm shape at later times is a sinusoidal form with an amplitude near 10 $\mu$m, coming from the energies defined in Section 2.1. (For interpretation of the references to color in this figure caption, the reader is referred to the web version of this article.)
planar. In these cases, the energy contribution for out-of-plane motion is always zero, and the trajectories of the swimmers remain in the initial plane. However, a pair of flagella each pursuing planar waveforms in parallel planes will interact with each other and generate out-of-plane motion. Below we consider these cases, and also explore the consequences of out-of-plane perturbations of initial flagellar configurations in both an isolated flagellum and pairs of interacting flagella. All simulations use the model parameter values listed in Table 1 and were initialized with the flagellar configurations shown in Fig. 3. The top row (Fig. 3a–c) is used for coplanar or perturbed coplanar initial conditions, and the bottom row (Fig. 3d–f) is used for parallel plane initial configurations. From these small amplitude initial configurations (chosen for simplicity), the sperm flagellum evolves into its natural waveform over time. In all two sperm simulations presented here, the preferred curvature waves are specified to be in-phase. For further description of the method of regularized Stokeslets and scalings used for the fluid equations, please see Cortez (2001) and Simons et al. (2014).

3.1. Single sperm

We first consider a single flagellum that is pursuing a symmetric, planar wave, but whose initial small amplitude configuration is perturbed slightly out-of-plane. The flagellum was started with a primarily x–y sinusoidal planar conformation, with the head of the sperm slightly raised from the x–y plane (shown in black in Fig. 3). We examine the dynamics of relaxation of this sperm flagellum to its native planar form from its initial three-dimensional configuration for various planarity stiffnesses $S_p$. The higher the value of this stiffness parameter, the faster one would expect the flagellum to relax towards its native plane. The relaxation to the native flagellar plane can be characterized by considering the maximum value of $| z_k |$ over all flagellum points. Fig. 4 shows the relaxation behavior of this quantity over time for values of $S_p$ between 0.001 and 1 pN μm$^{-3}$. We note that a value of 1 is of the same order of magnitude as the inextensibility parameter connecting the flagellar points together. With this parameter value, the flagellum approaches its native plane within $10^{-4}$ s (or $10^{-3}$ beats). For symmetric waveforms, a single sperm would relax to a planar configuration and swim linearly forward within that plane. For an asymmetric waveform, the sperm would also relax to a planar configuration and swim in a circular motion within its native plane, as shown in Fig. 9a and Movie M1.

3.2. Two sperm

We examine the cases of two identical flagella pursuing in-phase planar waves in (a) the same plane and (b) parallel planes. The initial configurations are taken to be perfectly planar, with no perturbations (Fig. 3a–c with no perturbation from $z = 0$, and Fig. 3d–f). In the coplanar case, the sperm flagella attract and swim side by side, never leaving the plane $z = 0$. The trajectories for this scenario are shown in Fig. 5a and Movie M2. Woolley et al. (2009) demonstrated this attraction between bull spermatozoa in experiments where the swimming was confined to a glass surface. Numerical simulations using multi-particle collision dynamics in two dimensions by Yang et al. (2008) and bead models in three dimensions by Llopis et al. (2013) also demonstrate the attraction of nearby flagella.

In contrast, when two flagella are initialized in parallel planes, the flagella repel over time. This is shown by the trajectories in Fig. 5b and the Movie M3, where the two sperm are initially placed half a flagellum length (50 μm) apart. Below, we will see that when they are placed closer together initially, the flagella may attract before ultimately repelling and swimming away from each other.

The velocities achieved by two identical interacting symmetric swimmers, when compared to an identically actuated single swimmer, depend on relative distance and orientations of the flagella. The velocities versus minimum distance between the two flagella as a function of time are plotted in Fig. 6 for four different initial configurations of two swimmers that are pursuing identical symmetric waveforms, and whose planarity stiffness constant is $S_p = 1$. Whereas a single swimmer has an average path velocity (i.e. the speed of the sperm over one beat period) of approximately 62 μm/s, the coplanar swimmers (Fig. 5a) that are attracted towards each other have a decreased velocity of 50 μm/s after they reach a steady swimming state (see Fig. 6c). The swimmers that start 50 μm apart in parallel planes (see Fig. 5b) swim slower at all times, and approach a single swimmer’s speed as they swim away from each other, as shown in Fig. 6a. If the distance between initially parallel swimmers is reduced from 50 μm to 5 μm, the swimmers actually see a transient increase in velocity and attract towards each other. They eventually do swim away from each other, however, even without a repelling force, as demonstrated in
The initial attraction of parallel swimmers, rotating their native plane and leading to an effective repulsion at longer times, is consistent with the observations of Llopis et al. (2013).

Now we are going to consider perturbed coplanar initial configurations, as depicted in Fig. 3a–c. Figs. 7–9 and Movies M4–M7 show examples of two sperm swimming near each other with variations in the symmetry of the preferred curvature waveform. In particular, Fig. 7 shows the relative motions of the sperm bodies for the two symmetric swimmers over a period of 10 s. Initially, the two sperm are swimming side-by-side in slightly different planes (labeled by $t=0.1$ s). As time progresses, the sperm are attracted towards each other (labeled by $t=5.1$ s) and then move away from each other (labeled by $t=10.1$ s). The trajectories of the heads of these two swimmers are shown in Fig. 8a. While near each other, the fluid-structure coupling of the two sperm causes each filament to impact the trajectory (and preferred flagellar plane) of the other. The supplementary Movie M4 shows the swimming behavior of this pair of perturbed coplanar swimmers.

Fig. 6b. The initial attraction of parallel swimmers, rotating their native plane and leading to an effective repulsion at longer times, is consistent with the observations of Llopis et al. (2013).

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over time.

The end of trajectories, starting position for sperm heads is at $x = 0$. Units are $\mu$m. Initial configurations are shown in Fig. 3. The Movies M4 and M5 show the swimming behavior of the trajectories in (a) and (b), respectively, over time.

Although two perfectly coplanar swimmers pursuing a planar waveform would remain in that plane, this simulation shows that a small initial perturbation from that plane is never recovered. As is noted in Llopis et al. (2013), coplanar swimming is an unstable configuration. Fig. 8 shows that even with symmetric motility patterns, fully three-dimensional trajectories may appear. We also see that these trajectories depend upon the intrinsic properties of the actuated swimmer, since different planarity stiffness constants result in different trajectories. For the stiffness parameter values shown, it appears that a weaker planar restriction stiffness results in longer-term cooperativity, as the flagella remain close to each other for a longer period of time. This period of close swimming shows that symmetrically beating sperm in different planes may move in wave-like patterns, weaving around each other before eventually swimming away. Velocities for these swimmers decrease as they get closer to each other, but eventually increase towards that of a single swimmer as they swim apart (see Fig. 6d). It appears that for transient periods of time, these swimmers may swim faster than a single swimmer due to their evolving relative orientation.

For a single flagellum, an asymmetric waveform acting upon a planar flagellum would result in a circular trajectory within the flagellar plane alone. Fig. 9 shows that for two sperm, weak asymmetry in the waveform can lead to more complex three-dimensional trajectories. The flagellar plane of each individual sperm is rotated due to the presence of a nearby neighbor. Thus, spiral or pseudo-helical (slinky-like) trajectories can evolve from asymmetric waveforms. Movie M6 shows this swimming behavior over time. Movie M7 shows high amplitude, strongly asymmetric waveforms that would characterize hyperactivity. Such high asymmetry and amplitude cause neighboring sperm to swim in more ball-like trajectories.

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hydrodynamics of sperm attachment and detachment from epithelia by allowing for forces due to dynamic elastic linkages between the actuated elastic flagellum and a planar wall. We will use the 3D extension described here, along with these dynamic linkages, to capture the complex binding interactions observed in some species (Fisher and Hoekstra, 2010; Johnston et al., 2007; Moore et al., 2002; Moore and Taggart, 1995).

Our current model is a simplistic model chosen to compare with previous planar model results. For instance, the effect of a sperm head has not been included in our model, which would introduce drag at the base of the flagellum and could affect motility patterns particularly near surfaces (see the recent work of Ishimoto and Gaffney, 2014). Moreover, some species have been shown experimentally to have three-dimensional sperm configurations that cannot be represented by simple planar forces described in our model. In fact, there is evidence that sperm flagella exhibit helical or non-planar waveforms in some settings (Bishop, 1962; Phillips, 1972; Rikmenspoel, 1965; Woolley and Vernon, 2001). In Woolley and Osborn (1984), the three-dimensional nature of hamster spermatozoa was determined to give rise to primarily clockwise torsional component, indicating that this came not from randomly applied (or external) forces but some structural feature of the flagellum itself. Additionally, the sea urchin behavior observed in some studies (Cosson et al., 2003; Woolley and Vernon, 2001) is thought to demonstrate a natural three-dimensional beat pattern which only reduces to a near-planar motion near surfaces. For these intrinsic three-dimensional waveforms, the method of regularized Stokeslets coupled to a Kirchhoff rod model (Olson et al., 2013) should be explored.

Conflict of interest statement

The authors have no conflict of interests to disclose.

Acknowledgments

The work of J.S., R.C. and L.F. was supported, in part, by the National Science Foundation Grant DMS-104626.

Appendix A. Supplementary material

The following are the supplementary data to this paper:
Video S1 Video S2 Video S3 Video S4 Video S5 Video S6 Video S7

Video S1. A weakly asymmetric swimmer starting with a perturbed planar initial configuration (see black curve in Fig. 3a–c, with planar stiffness constant $S_p = 1$. The sperm travels in a circular path, recovering from its non-planar configuration quickly. Gray curve shows the projection (shadow) of the flagellum on the plane $z = 100$. Units are in microns. Length of simulation is 25 s. A video clip is available online. Supplementary material related to this paper can be found online at http://dx.doi.org/10.1016/j.jbiomech.2015.01.050.
Video S2. Two symmetric swimmers with a coplanar initial configuration (see Fig. 3a–c), with planar stiffness constant $S_p = 1$. The two swimmers attract over time. Light shaded curves show the projections (shadows) of the flagella on the plane $z = -100$. Units are in microns. Length of simulation is 25 s. A video clip is available online. Supplementary material related to this paper can be found online at http://dx.doi.org/10.1016/j.jbiomech.2015.01.050.

Video S3. Two symmetric swimmers with a parallel planar initial configuration (see Fig. 3d–f), with planar stiffness constant $S_p = 1$. The two swimmers repel over time. Light shaded curves show the projections (shadows) of the flagella on the plane $z = -100$. Units are in microns. Length of simulation is 25 s. A video clip is available online. Supplementary material related to this paper can be found online at http://dx.doi.org/10.1016/j.jbiomech.2015.01.050.
Video S4. Two symmetric swimmers with a perturbed planar initial configuration (see Fig. 3a–c), with planar stiffness constant $S_p = 1$. The two swimmers “weave” near each other before swimming apart. Light shaded curves show the projections (shadows) of the flagella on the plane $z = -150$. Units are in microns. Length of simulation is 25 s. A video clip is available online. Supplementary material related to this paper can be found online at http://dx.doi.org/10.1016/j.jbiomech.2015.01.050.

Video S5. Two symmetric swimmers with a perturbed planar initial configuration (see Fig. 3a–c), with planar stiffness constant $S_p = 0.01$. The two swimmers “weave” near each other for a more prolonged period than in Movie M4 because of a weaker stiffness constant. Light shaded curves show the projections (shadows) of the flagella on the plane $z = -100$. Units are in microns. Length of simulation is 25 s. A video clip is available online. Supplementary material related to this paper can be found online at http://dx.doi.org/10.1016/j.jbiomech.2015.01.050.
Video S6. Two weakly asymmetric swimmers with a perturbed planar initial configuration (see Fig. 3a–c), with planar stiffness constant $S_p = 1$. The two swimmers move in curved “slinky-like” trajectories over time. Light shaded curves show the projections (shadows) of the flagella on the plane $z = -250$. Units are in microns. Length of simulation is 25 s. A video clip is available online. Supplementary material related to this paper can be found online at http://dx.doi.org/10.1016/j.jbiomech.2015.01.050.
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