

Filament mechanics in a half-space via regularised Stokeslet segments

B. J. Walker^{1,†}, K. Ishimoto², H. Gadêlha³ and E. A. Gaffney¹

¹Wolfson Centre for Mathematical Biology, Mathematical Institute, University of Oxford, Oxford OX2 6GG, UK

²Graduate School of Mathematical Sciences, The University of Tokyo, Tokyo, 153-8914, Japan

³Department of Engineering Mathematics, University of Bristol, Bristol BS8 1UB, UK

(Received 4 April 2019; revised 17 July 2019; accepted 28 August 2019;
first published online 1 October 2019)

We present a generalisation of efficient numerical frameworks for modelling fluid–filament interactions via the discretisation of a recently developed, non-local integral equation formulation to incorporate regularised Stokeslets with half-space boundary conditions, as motivated by the importance of confining geometries in many applications. We proceed to utilise this framework to examine the drag on slender inextensible filaments moving near a boundary, firstly with a relatively simple example, evaluating the accuracy of resistive force theories near boundaries using regularised Stokeslet segments. This highlights that resistive force theories do not accurately quantify filament dynamics in a range of circumstances, even with analytical corrections for the boundary. However, there is the notable and important exception of movement in a plane parallel to the boundary, where accuracy is maintained. In particular, this justifies the judicious use of resistive force theories in examining the mechanics of filaments and monoflagellate microswimmers with planar flagellar patterns moving parallel to boundaries. We proceed to apply the numerical framework developed here to consider how filament elasto-hydrodynamics can impact drag near a boundary, analysing in detail the complex responses of a passive cantilevered filament to an oscillatory flow. In particular, we document the emergence of an asymmetric periodic beating in passive filaments in particular parameter regimes, which are remarkably similar to the power and reverse strokes exhibited by motile 9 + 2 cilia. Furthermore, these changes in the morphology of the filament beating, arising from the fluid–structure interactions, also induce a significant increase in the hydrodynamic drag of the filament.

Key words: micro-organism dynamics

1. Introduction

The mechanics of flexible filaments on the microscale underpin much of biology, from the propulsive flagella of motile bacteria and spermatozoa to nodal cilia, the

† Email address for correspondence: benjamin.walker@maths.ox.ac.uk

latter hypothesised to be responsible for the breaking of left–right symmetry in mammals (Gray 1928; Berg & Anderson 1973; Smith, Montenegro-Johnson & Lopes 2019). Furthermore, the dynamics of elastic filaments is of intense interest in the physics of microdevices and surface flows, including near-wall dynamics. For instance, the consideration of soft deformable sensors has already motivated extensive studies of attached filaments (Roper *et al.* 2006; Guglielmini *et al.* 2012), as has the characterisation of attached filament forces for understanding the drag induced by slender appendages (Pozrikidis 2011; Curtis *et al.* 2012; Simons *et al.* 2014). Such appendages range from the primary cilium to carbon nanotube mats, with an extensive review of the field presented by du Roure *et al.* (2019), which notes that both theoretical and numerical developments are very much still required in this field. Indeed, with advances in microscopy enabling ever more detailed quantification of kinematics, often with confining geometry such as a cover slip or substrate, the development of validated, and ideally simple, methodologies would be beneficial in estimating mechanics from kinematics.

Furthermore, the complex mechanics of fluid–structure interaction is an important problem and has been well studied, as illustrated by the slender body theory of Tornberg & Shelley (2004) and Liu *et al.* (2018). Numerical simulations that attempt to move past slender body theory are frequently plagued by extensive numerical stiffness, such as the regularised Stokeslet simulations of Olson, Lim & Cortez (2013) and Ishimoto & Gaffney (2018). The recent advance of Moreau, Giraldi & Gadêlha (2018) and its subsequent extension by Hall-McNair *et al.* (2019) have sought to address this stiffness, with their methodologies significantly reducing the computational cost associated with filament–fluid interactions via the use of integrated force and moment balance equations, but have not considered even the simplest confined geometry that is an infinite planar wall. Hence our first objective will be to generalise these improved frameworks to dynamics in a half-space bounded by a wall, adapting the recent regularised Stokeslet segment approach of Cortez (2018) for drag calculations and elasto-hydrodynamics.

We will validate in detail the computation of drag from kinematic data against the earlier work of Ramia, Tullock & Phan-Thien (1993), additionally validating our proposed elasto-hydrodynamic framework against the gold-standard boundary element method of Pozrikidis (2010). These validations demonstrate the high accuracy of our approach in capturing the mechanics of filaments, and thus it is of extensive use for understanding cellular swimming. For instance, resistive force theories are very popular due to their ease of use (Lauga *et al.* 2006; Gadêlha *et al.* 2010; Sznitman *et al.* 2010; Schulman *et al.* 2014; Utada *et al.* 2014; Moreau *et al.* 2018), and they have been shown to be of reasonable accuracy in free space for small-bodied cells such as spermatozoa (Johnson & Brokaw 1979). With freedom to choose the resistive coefficients, such local drag theories have been shown to perform very well even near a boundary (Friedrich *et al.* 2010), as supported by the boundary element method validation studies for straight rods of Ramia *et al.* (1993). This suggests that the refinements of Brenner (1962) and Katz, Blake & Paveri-Fontana (1975) to free-space resistive force theories, valid for straight filament motion parallel or perpendicular to a planar wall, may be useful for the analysis of microscopy data, and would be very popular if demonstrated to be accurate. In particular, in the common circumstance where subject cells are imaged swimming parallel to a coverslip, for example Riedel-Kruse & Hilfinger (2007) and Friedrich *et al.* (2010), and thus have unchanging boundary separation, we hypothesise that these refined resistive force theories may be sufficient to accurately capture the

hydrodynamic drag on a curved filament. Hence, as our second objective and first application, we use the regularised Stokeslet segment framework of Cortez (2018) to test this hypothesis in exemplar problems. In particular we will consider both the case when a filament is moving parallel to a boundary and the case when it is attached to the surface and moving extensively perpendicular to the wall. Further, the presented elastohydrodynamic framework is sufficiently flexible to enable sophisticated considerations of fluid–structure interactions, inherited from the approach of Moreau *et al.* (2018). Hence, as an application of non-local elastohydrodynamics in a half-space, our final objective is to proceed to demonstrate the methodology’s ease of use in the context of such complex physics by examining the mechanics of a flexible inextensible adhered filament in an oscillatory flow, generalising Pozrikidis’ study of carbon nanomat surface drag to time dependent flows (Pozrikidis 2011).

More generally, testing the boundary-corrected resistive force theories when applied to videomicroscopy data has not been considered in detail. Such evaluation is important for ascertaining whether these corrected resistive force theories may be reliably and accurately used to provide simple access to the viscous drag on filaments in the presence of boundaries, pertinent to the analysis of diverse videomicroscopy data, for example Ohmuro & Ishijima (2006) and Smith *et al.* (2009). Furthermore, the methodology presented is both impactful and a non-trivial extension of previous studies in that it will enable the efficient numerical study of the elastohydrodynamical problem of filament motion near a boundary, once more facilitating the mechanical investigation of a wealth of flagellated microswimmers such as the well-studied human spermatozoon (Fauci & McDonald 1995; Elgeti, Kaupp & Gompper 2010; Ishimoto & Gaffney 2015). These elastohydrodynamic methods also provide clear precursors for the application of the framework to simulations of oscillating systems in physiological and soft matter modelling, such as primary and motile ciliary systems respectively in the kidneys and lungs, in addition to numerical studies of active surfaces, for example Shum *et al.* (2013) and Balazs *et al.* (2014).

2. Methods

2.1. Piecewise-linear filaments

We consider a planar slender inextensible filament represented by N piecewise-linear segments of constant length and described by arclength parameter $s \in [0, L]$, where L is the filament length. The segment endpoints are taken to correspond to material points, with their positions being denoted $\mathbf{x}_1, \dots, \mathbf{x}_{N+1}$ and where \mathbf{x}_i and \mathbf{x}_{i+1} correspond to the i th segment. We will refer to \mathbf{x}_1 as the base of the filament and correspondingly \mathbf{x}_{N+1} as the tip, and denote the constant arclength associated with the material point \mathbf{x}_i as s_i . As the filament is assumed to be planar, valid in the absence of external symmetry-breaking features, without loss of generality we assume that it lies in a plane spanned by orthogonal unit vectors \mathbf{e}_x and \mathbf{e}_y , with \mathbf{e}_z completing the orthonormal right-handed triad $\mathbf{e}_x\mathbf{e}_y\mathbf{e}_z$, and we may write $\mathbf{x}_i = x_i\mathbf{e}_x + y_i\mathbf{e}_y$. Following Moreau *et al.* (2018) we note that, once the segment lengths v_1, \dots, v_N are prescribed, the filament may be completely described by the $N + 2$ scalars $x_1, y_1, \theta_1, \dots, \theta_N$, where θ_i is defined as the angle between the i th segment and the unit vector \mathbf{e}_x as shown in figure 1. Explicitly, from these $N + 2$ variables we may recover the filament endpoints as

$$\mathbf{x}_j = \mathbf{x}_1 + \sum_{i=1}^{j-1} (\cos \theta_i \mathbf{e}_x + \sin \theta_i \mathbf{e}_y) v_i, \quad (2.1)$$

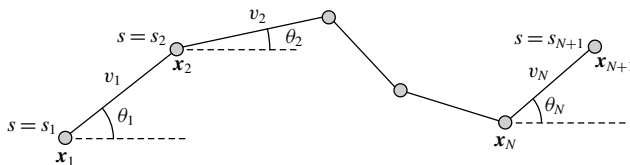


FIGURE 1. The two-dimensional piecewise-linear filament. We consider an inextensible filament composed of N straight segments each of length v_i , connected at material points \mathbf{x}_i with each segment making angle θ_i with the global \mathbf{e}_x axis. The endpoints \mathbf{x}_1 and \mathbf{x}_{N+1} shall be referred to as the base and tip respectively, and we note that, given segment lengths v_i , the entire filament may be described by the location of the base and the angles θ_i .

and, using dots to denote derivatives with respect to time, the velocity of each material point is given by

$$\dot{\mathbf{x}}_j = \dot{\mathbf{x}}_1 + \sum_{i=1}^{j-1} (-\sin\theta_i \mathbf{e}_x + \cos\theta_i \mathbf{e}_y) \dot{\theta}_i v_i, \tag{2.2}$$

again expressible using only the $N + 2$ variables due to the imposed geometrical constraints.

2.2. Force distributions via regularised Stokeslet segments

To describe the low Reynolds number fluid dynamics pertinent to the filament we utilise the recent work of Cortez (2018), namely the method of regularised Stokeslet segments (RSS). Here we briefly recapitulate the formulation of this methodology as presented by Cortez, relating the force density applied on the fluid by the filament to the fluid velocity via non-local hydrodynamics.

As in the popular method of regularised Stokeslets (Cortez 2001), we begin by considering solutions of the three-dimensional smoothly forced Stokes equations, which may be stated for viscosity μ and force \mathbf{f} applied on the fluid at the origin as

$$\mu \nabla^2 \mathbf{u} = \nabla p - \mathbf{f} \phi_\epsilon, \quad p, \mathbf{u} \rightarrow 0 \quad \text{as } |\mathbf{x}| \rightarrow \infty, \tag{2.3a,b}$$

where \mathbf{u} is the fluid velocity, p is the pressure and ϕ_ϵ is a smooth approximation to the Dirac delta distribution dependent on the small parameter ϵ . Following § 2 of Cortez (2018) we take

$$\phi_\epsilon(\mathbf{x}) = \frac{15\epsilon^4}{8\pi(|\mathbf{x}|^2 + \epsilon^2)^{7/2}}, \tag{2.4}$$

for which the solution to (2.3) is known and given by

$$8\pi\mu\mathbf{u}(\mathbf{x}) = \left[\left(\frac{1}{R} + \frac{\epsilon^2}{R^3} \right) \mathbf{I} + \frac{\mathbf{x}\mathbf{x}^\top}{R^3} \right] \mathbf{f}, \quad R(\mathbf{x}) = \sqrt{|\mathbf{x}|^2 + \epsilon^2}. \tag{2.5a,b}$$

By linearity of the Stokes equations the velocity at a point $\tilde{\mathbf{x}}$ due to a distribution of regularised Stokeslets along the filament with force density $\mathbf{f}(s)$ is therefore given

by

$$8\pi\mu\mathbf{u}(\tilde{\mathbf{x}}) = \int_0^L \left[\left(\frac{1}{R(\mathbf{r})} + \frac{\epsilon^2}{R(\mathbf{r})^3} \right) \mathbf{I} + \frac{\mathbf{r}\mathbf{r}^\top}{R(\mathbf{r})^3} \right] \mathbf{f}(s) \, ds, \quad \mathbf{r}(s) = \tilde{\mathbf{x}} - \mathbf{x}(s). \quad (2.6a,b)$$

Whilst the original method of regularised Stokeslets would entail approximating the force distribution by a finite number of smoothed point forces, the method of regularised Stokeslet segments instead considers a linear distribution of forces along the straight segments of the discretised filament. With the discretisation of the force distribution as a continuous piecewise-linear function along each of the segments, the fluid velocity is instead given by a sum of integrals over each of the segments, where each individual integral may be analytically evaluated. Parameterising the j th segment by $\alpha \in [0, 1]$ so that $\mathbf{x} = \mathbf{x}_j - \alpha\mathbf{v}$ for $\mathbf{v} = \mathbf{x}_j - \mathbf{x}_{j+1}$, and additionally writing the force density as $\mathbf{f} = \mathbf{f}_j + \alpha(\mathbf{f}_{j+1} - \mathbf{f}_j)$ for force densities $\mathbf{f}_j, \mathbf{f}_{j+1}$ at $\mathbf{x}_j, \mathbf{x}_{j+1}$ respectively, the integral along the j th segment is given by

$$v_j \int_0^1 \left[\left(\frac{1}{R} + \frac{\epsilon^2}{R^3} \right) \mathbf{I} + \frac{(\tilde{\mathbf{x}} - \mathbf{x}_j + \alpha\mathbf{v})(\tilde{\mathbf{x}} - \mathbf{x}_j + \alpha\mathbf{v})^\top}{R^3} \right] [\mathbf{f}_j + \alpha(\mathbf{f}_{j+1} - \mathbf{f}_j)] \, d\alpha, \quad (2.7)$$

where we have identified $|\mathbf{v}| = v_j$ and $R = \sqrt{|\tilde{\mathbf{x}} - \mathbf{x}_j + \alpha\mathbf{v}|^2 + \epsilon^2}$. As noted by Cortez (2018, § 2.2) this may be written as a sum of integrals of the form

$$T_{m,q} = \int_0^1 \alpha^m R^q \, d\alpha \quad (2.8)$$

for $(m, q) \in \{(0, -1), (1, -1), (0, -3), (1, -3), (2, -3), (3, -3)\}$, each of which is implicitly dependent on j and may be evaluated by explicit computation of $T_{0,-1}$ and $T_{0,-3}$ and subsequent application of the recurrence relation

$$v_j^2 T_{m+1,q} = -\frac{m}{(q+2)} T_{m-1,q+2} - (\tilde{\mathbf{x}} - \mathbf{x}_j) \cdot \mathbf{v} T_{m,q} + \frac{1}{q+2} \alpha^m R^{q+2} \Big|_0^1, \quad q \neq -2, \quad (2.9)$$

which follows from the definition of $T_{m,q}$ and consideration of $(d/d\alpha)(\alpha^m R^q)$.

The simple extension of this methodology to a half-space bounded by a no-slip planar wall via the image system of Ainley *et al.* (2008) (with the typographical error corrected Smith (2009)) is presented in detail by Cortez (2018, § 3.3). Exemplified in figure 2, we will consider two configurations of half-space, one in which the infinite planar boundary is parallel to the plane containing the filament, given by $z = 0$, and another where the boundary is situated perpendicular to this plane and given by $y = 0$. In both cases the hydrodynamics may be described by the same regularised singularity representation, each requiring computation of $T_{m,q}$ for the additional values $(m, q) \in \{(0, -5), (1, -5), (2, -5), (3, -5), (4, -5), (5, -5)\}$ in order to include the additional necessary regularised singularities. Omitted from the previous work of Cortez, we compute

$$T_{0,-5} = \frac{(B + \alpha C^2) (-B^2 + C^2[3A^2 + 4\alpha B + 2\alpha^2 C^2 + 3\epsilon^2])}{3R^3 (B^2 - C^2[A^2 + \epsilon^2])^2} \Big|_0^1, \quad (2.10)$$

where $A = |\tilde{\mathbf{x}} - \mathbf{x}_j|$, $B = (\tilde{\mathbf{x}} - \mathbf{x}_j) \cdot \mathbf{v}$ and $C = |\mathbf{v}| = v_j$. The remaining values of $T_{m,q}$ may be computed via the recurrence relation equation (2.9). Simple computation of

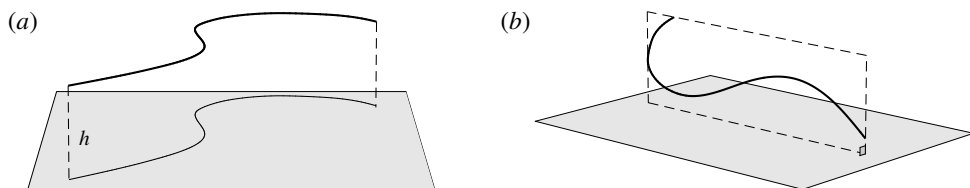


FIGURE 2. Schematic configurations of planar filaments near boundaries. (a) A filament moving parallel to the infinite planar boundary, at constant separation h . The projection of the filament onto the boundary is shown as a thin black line. (b) A filament contained in a plane perpendicular to the boundary, with the plane of motion shown dashed.

the coefficients of the $T_{m,q}$ results in a matricial form of the velocity contribution at $\tilde{\mathbf{x}}$ from the j th segment as a linear operator acting on \mathbf{f}_j and \mathbf{f}_{j+1} , the details of which are cumbersome and given in the supplementary material, available at <https://doi.org/10.1017/jfm.2019.723>. The overall velocity at the evaluation point $\tilde{\mathbf{x}}$ is then given by the sum of these contributions, resulting in a matricial equation of the form

$$\mathbf{u}(\tilde{\mathbf{x}}) = \mathbf{M}(\tilde{\mathbf{x}})\mathbf{F}, \tag{2.11}$$

where $\mathbf{F} = (f_{1,x}, f_{1,y}, \dots, f_{N+1,x}, f_{N+1,y})^\top$ herein denotes the composite vector of force densities applied on the fluid at the segment endpoints \mathbf{x}_j in the directions \mathbf{e}_x and \mathbf{e}_y , where we write $\mathbf{f}_j = f_{j,x}\mathbf{e}_x + f_{j,y}\mathbf{e}_y$. Taking the evaluation point $\tilde{\mathbf{x}}$ as each \mathbf{x}_i in turn yields a square system of linear equations in $2(N + 1)$ variables, relating the velocities $\mathbf{u}(\mathbf{x}_i)$ to the force distribution on the filament via non-local hydrodynamics. Application of the no-slip condition at the segment endpoints then gives a relation between the filament velocities and the forces applied on the fluid by the filament, which we write in brief as

$$\dot{\mathbf{X}} = \mathbf{A}\mathbf{F}, \tag{2.12}$$

for the square matrix \mathbf{A} composed row-wise of blocks $\mathbf{M}(\mathbf{x}_i)$ for $i = 1, \dots, N + 1$, and where $\dot{\mathbf{X}} = (\dot{x}_1, \dot{y}_1, \dots, \dot{x}_{N+1}, \dot{y}_{N+1})^\top$. Given kinematic data, this linear system may be readily solved to give the force densities applied on the fluid by the filament.

2.3. Extension to efficient solution of elastohydrodynamic equations

We now proceed to combine the work of Cortez (2018) and Moreau *et al.* (2018) to give an efficient scheme for solving non-local planar elastohydrodynamics, similar in concept to the piecewise-constant force density approach of Hall-McNair *et al.* (2019) but with continuous piecewise-linear force discretisation and the additional inclusion of an infinite planar boundary. As formulated by Moreau *et al.* (2018), we integrate the pointwise conditions of force and moment balance on the filament, given by

$$\mathbf{n}_s - \mathbf{f} = \mathbf{0}, \tag{2.13}$$

$$\mathbf{m}_s + \mathbf{x}_s \times \mathbf{n} = \mathbf{0} \tag{2.14}$$

for contact force and couple denoted \mathbf{n} , \mathbf{m} respectively and where a subscript of s denotes differentiation with respect to arclength, noting that we have assumed

slenderness of the filament. This yields the integrated equations

$$-\sum_{j=1}^N \int_{s_j}^{s_{j+1}} \mathbf{f}(s) \, ds = \mathbf{n}(0), \tag{2.15}$$

$$-\sum_{j=i}^N \int_{s_j}^{s_{j+1}} (\mathbf{x}(s) - \mathbf{x}_i) \times \mathbf{f}(s) \, ds = \mathbf{m}(s_i), \quad i = 1, \dots, N. \tag{2.16}$$

Here we have assumed conditions of zero contact force and couple at the tip of the filament, i.e. $\mathbf{n}(L) = \mathbf{m}(L) = \mathbf{0}$, retaining generality at the base for the time being. With the assumption of piecewise-linear distributions of force density \mathbf{f} over each segment, again with $\mathbf{f} = \mathbf{f}_j + \alpha(\mathbf{f}_{j+1} - \mathbf{f}_j)$ on the j th segment for $\alpha \in [0, 1]$, we may write these integrals as linear operators on \mathbf{F} , yielding the system

$$-\mathbf{B}\mathbf{F} = \mathbf{R}, \tag{2.17}$$

where $\mathbf{R} = (\mathbf{n}(s_1) \cdot \mathbf{e}_x, \mathbf{n}(s_1) \cdot \mathbf{e}_y, m(s_1), \dots, m(s_N))^T$ and we are writing $\mathbf{m}(s_i) = m(s_i)\mathbf{e}_z$ by planarity of the filament. The matrix \mathbf{B} is of dimension $(N + 2) \times 2(N + 1)$, which under the assumption of equispaced segment endpoints has rows \mathbf{B}_k given by

$$\mathbf{B}_1 = \frac{\Delta s}{2} [1, 0, 2, 0, 2, \dots, 2, 0, 1, 0], \tag{2.18}$$

$$\mathbf{B}_2 = \frac{\Delta s}{2} [0, 1, 0, 2, 0, 2, \dots, 2, 0, 1], \tag{2.19}$$

with the additional rows \mathbf{B}_{i+2} each encoding the integrated expression

$$\mathbf{e}_z \cdot \sum_{j=i}^N \left\{ \left[\frac{\Delta s}{2} (\mathbf{x}_j - \mathbf{x}_i) + \frac{\Delta s^2}{6} \mathbf{t}_j \right] \times \mathbf{f}_j + \left[\frac{\Delta s}{2} (\mathbf{x}_j - \mathbf{x}_i) + \frac{\Delta s^2}{3} \mathbf{t}_j \right] \times \mathbf{f}_{j+1} \right\}, \tag{2.20}$$

where $\Delta s = L/N$ is the length of each segment, $\mathbf{t}_j = \cos \theta_j \mathbf{e}_x + \sin \theta_j \mathbf{e}_y$ is the local unit tangent and $i = 1, \dots, N$. Thus we may write the coupled elasto-hydrodynamic problem as

$$-\mathbf{B}\mathbf{A}^{-1}\dot{\mathbf{X}} = \mathbf{R}, \tag{2.21}$$

where \mathbf{B} represents integration over segments and \mathbf{A} encodes the relationship between the forces on the surrounding fluid and the velocities of material points on the filament, here non-local and assumed to be invertible.

As noted in § 2.1, the material velocities $\dot{\mathbf{x}}_i$ may be expressed in terms of the time derivatives of the base point \mathbf{x}_1 and the segment angles $\theta_1, \dots, \theta_N$. Defining \mathbf{Q} to be the $2(N + 1) \times (N + 2)$ matrix such that

$$\mathbf{Q}\dot{\boldsymbol{\theta}} = \dot{\mathbf{X}}, \tag{2.22}$$

where $\boldsymbol{\theta} = (x_1, y_1, \theta_1, \dots, \theta_N)^T$, we have \mathbf{Q} given explicitly by

$$\mathbf{Q} = \left[\begin{array}{cc|cc} 1 & 0 & & \\ \vdots & \vdots & \mathbf{Q}_1 & \\ 1 & 0 & & \\ \hline 0 & 1 & & \\ \vdots & \vdots & \mathbf{Q}_2 & \\ 0 & 1 & & \end{array} \right]_P, \tag{2.23}$$

where the subscript P denotes that the i th row of \mathbf{Q} is to be permuted to

$$P(i) = \begin{cases} 2(i-1) + 1, & i = 1, \dots, N + 1, \\ 2(i - N - 1), & i = N + 2, \dots, 2N + 2. \end{cases} \quad (2.24)$$

This permutation of \mathbf{Q} allows us to define the blocks \mathbf{Q}_1 and \mathbf{Q}_2 as being strictly lower triangular matrices of dimension $(N + 1) \times N$ with entries

$$\mathbf{Q}_1^{ij} = \Delta s \begin{cases} -\sin \theta_j, & j < i, \\ 0, & j \geq i, \end{cases} \quad (2.25)$$

$$\mathbf{Q}_2^{ij} = \Delta s \begin{cases} +\cos \theta_j, & j < i, \\ 0, & j \geq i. \end{cases} \quad (2.26)$$

Given the matrix \mathbf{Q} we may rewrite the full coupled elasto-hydrodynamic problem simply as

$$-\mathbf{BA}^{-1}\mathbf{Q}\dot{\boldsymbol{\theta}} = \mathbf{R}, \quad (2.27)$$

with $\mathbf{BA}^{-1}\mathbf{Q}$ being a square matrix of dimension $(N + 2) \times (N + 2)$. Finally, and here for example assuming force and moment-free conditions at the filament base, the standard linear constitutive relation $m(s_i) = EI \theta_s(s_i) \approx EI (\theta_i - \theta_{i-1})/\Delta s$ for bending stiffness EI , valid for $i = 2, \dots, N$, gives \mathbf{R} explicitly in terms of $\boldsymbol{\theta}$, yielding a linear system of low dimension that may be readily solved for $\dot{\boldsymbol{\theta}}$, with the temporal dynamics then computable via existing ordinary differential equation (ODE) methods. As noted by Moreau *et al.*, this formulation is readily extensible to the imposition of a variety of boundary conditions, in particular that of a cantilevered filament base, achieved by replacing the overall zero-force and zero-moment equations with $\dot{x}_1 = \dot{y}_1 = \dot{\theta}_1 = 0$.

We non-dimensionalise this system by scaling spatial coordinates with filament length L , forces with EI/L^2 and time with some characteristic time scale T , yielding the non-dimensional system

$$-E_h \hat{\mathbf{B}} \hat{\mathbf{A}}^{-1} \hat{\mathbf{Q}} \hat{\dot{\boldsymbol{\theta}}} = \hat{\mathbf{R}}, \quad E_h = \frac{8\pi\mu L^4}{EIT} \quad (2.28)$$

for elasto-hydrodynamic number E_h , where the notation $\hat{\cdot}$ denotes non-dimensional quantities. The dimensional and non-dimensional quantities are related by

$$\mathbf{B} = L^2 \hat{\mathbf{B}}, \quad \mathbf{A} = \frac{1}{8\pi\mu} \hat{\mathbf{A}}, \quad \mathbf{Q}\dot{\boldsymbol{\theta}} = \frac{L}{T} \hat{\mathbf{Q}} \hat{\dot{\boldsymbol{\theta}}}, \quad \mathbf{R} = \frac{EI}{L} \hat{\mathbf{R}}, \quad (2.29a-d)$$

where we have multiplied the two force balance equations by $\Delta s = L/N$ and absorbed the dimensional scalings of x_1, y_1 into \mathbf{Q} for convenience, with $\hat{\boldsymbol{\theta}} = (\hat{x}_1, \hat{y}_1, \theta_1, \dots, \theta_N)^\top$. In order to pose the non-dimensional equations in terms of the commonly used sperm number Sp (Delmotte, Climent & Plouraboué 2015; Ishimoto & Gaffney 2018; Moreau *et al.* 2018), we proceed following Ishimoto & Gaffney (2018) to define

$$Sp^4 = \frac{\xi L^4}{EIT}, \quad \xi = \frac{4\pi\mu}{\log(2L/\epsilon)} \quad (2.30a,b)$$

for resistive force coefficient ξ and $L/\epsilon = 10^3$, giving the approximate relation $E_h \approx 15.2 Sp^4$.

2.4. Implementation, verification and parameter choice

Both the calculation of force densities from kinematic data and the solution of full elastohydrodynamics were implemented in MATLAB[®]. We utilise the inbuilt stiff ODE solver `ode15s` (Shampine & Reichelt 1997), making use of variable step sizes in order to satisfy configurable error tolerances that are typically set here at 10^{-6} , with the presented results insensitive to this choice. Prior to the recent work of Hall-McNair *et al.* (2019) the solution of non-local elastohydrodynamics has required significant computational work and minimal timestep for the solution of this stiff problem (Olson *et al.* 2013; Ishimoto & Gaffney 2018), and we replicate in our implementation the low computational cost associated with the integrated elasticity equations of Moreau *et al.* (2018). In particular, typical simulations of a cantilevered filament in background flow, explored in detail in §§ 3.4 and 3.5, have a typical runtime of 10 s, where $N = 40$ and we simulate over 10 periods of oscillation of the background flow on modest hardware (Intel[®] Core[™] i7-6920HQ CPU).

Verification was first performed on the reduction to an unbounded domain, with the full elastohydrodynamics being verified by comparison with the resistive force theory results of Moreau *et al.* (2018) for filament relaxation. Further drag calculations were compared against the work of Cortez (2018) for the case of uniform motion, with the degree to which the no-slip condition is numerically satisfied along the filament length having previously been investigated in detail (Cortez 2018, § 3.1). The regularised image system in a half-space was then verified by explicit evaluation of the fluid velocity on the no-slip boundary, with the numerical result being zero to machine precision, and further by noting that far-field results were seen to converge to those of the free-space system. Further qualitative checks were performed, an example being the successful reproduction of the intuitive result that relaxation time scales increase for filaments with reduced separation from the no-slip stationary boundary, along with numerical comparison against the previous works of Pozrikidis regarding the elastohydrodynamics of filaments in steady shear flow (Pozrikidis 2010, 2011), a brief example of which being presented in appendix B. Results of additional verification against the boundary element method of Ramia *et al.* (1993) are shown in figure 3(a). Sufficient accuracy is typically achieved with $N = 20$ segments, similar to the discretisation used by Cortez (2018), although we typically take $N = 40$ to enable the capturing of high-curvature filaments.

As noted by Cortez (2018), the use of regularised Stokeslet segments allows the regularisation parameter to represent the radius of the filament being modelled, verified here by comparison with the boundary element method of Ramia *et al.* (1993) in figure 3, subject to the phenomenological condition that ϵ be less than the length of the segments, which appears necessary for convergence. We will take $\epsilon/L = 10^{-3}$ unless otherwise stated, with typical slender filament aspect ratios yielding ϵ/L in the range $(10^{-2}, 10^{-3})$ (Yonekura, Maki-Yonekura & Namba 2003; Ishimoto & Gaffney 2016). Whilst the results that follow naturally depend on the size of the regularisation parameter, it being used as a proxy for the filament radius, this dependence appears to be predominantly quantitative, with the same qualitative conclusions holding across the range of physically relevant values of ϵ/L .

2.5. Endpoint effects

Inherent to the method of regularised Stokeslet segments as presented here is the presence of apparent large variations in computed force densities at the tips of filaments. This was initially observed by Cortez (2018, § 3.1, figure 4), who remarked

that these endpoint effects may be reduced when ϵ/L is small. Indeed, for ϵ/L in our range of physical interest these endpoint effects contribute minimally to overall drag calculations, in particular having little effect on the computed total drag exerted on a filament and the computed force densities away from the filament tip, whose presentation we focus on herein. Exploration of the effects of non-uniform segment lengths yields the observation that these oscillatory endpoint effects remain limited to approximately the 3 segments proximal to the ends of the filament, thus the integral contribution of these oscillations may be reduced by the clustering of segments near the filament tips. For our typical parameters of $N = 40$ and $\epsilon/L = 10^{-3}$ we observe a difference in integrated drag along of filaments of less than 2% between linear, quadratic and Chebyshev segment endpoints, hence we proceed with calculations utilising segments of uniform length.

2.6. Boundary-corrected resistive force theory

The leading-order approximation of slender body hydrodynamics that is resistive force theory (RFT) was first introduced by Gray and Hancock (Hancock 1953; Gray & Hancock 1955), relating the local drag on a body to its local tangential and normal velocities u_t, u_n via

$$f_t = -C_t u_t, \quad f_n = -C_n u_n. \tag{2.31a,b}$$

Here f_t, f_n are the local tangential and normal components of force applied on the fluid, with the constant resistive coefficients C_t, C_n typically being functions of filament aspect ratio. As RFT relates forces only to local velocities, calculations using this simple local drag theory are not able to account for the presence of a boundary. Brenner (1962) and Katz *et al.* (1975) posed corrections to resistive force theory for straight filaments in asymptotic regimes far from or near to an infinite no-slip boundary, with the latter having been verified against high-accuracy boundary element methods by Ramia *et al.* (1993). In the case of filament motion confined to a plane parallel to the boundary, as exemplified in figure 2(a), by this wall-corrected resistive force theory (W-RFT) a straight filament parallel to the boundary has resistive coefficients given by

$$C_t = \frac{2\pi\mu}{\log\left(\frac{2}{\epsilon}\right) - 0.807 - \frac{3L}{8h}}, \quad C_n = \frac{4\pi\mu}{\log\left(\frac{2}{\epsilon}\right) + 0.193 - \frac{3L}{4h}} \quad \text{if } L \ll h, \tag{2.32a,b}$$

$$C_t = \frac{2\pi\mu}{\log\left(\frac{2h}{\epsilon}\right)}, \quad C_n = \frac{4\pi\mu}{\log\left(\frac{2h}{\epsilon}\right)} \quad \text{if } L \gg h, \tag{2.33a,b}$$

where h is the boundary separation of the filament and ϵ has been taken as the filament radius. Similarly, and as in figure 2(b), when aligned parallel to the boundary and moving in a plane perpendicular to the surface the coefficients are given by

$$C_t = \frac{2\pi\mu}{\log\left(\frac{2}{\epsilon}\right) - 0.807 - \frac{3L}{8h}}, \quad C_n = \frac{4\pi\mu}{\log\left(\frac{2}{\epsilon}\right) + 0.193 - \frac{3L}{2h}} \quad \text{if } L \ll h, \tag{2.34a,b}$$

$$C_t = \frac{2\pi\mu}{\log\left(\frac{2h}{\epsilon}\right)}, \quad C_n = \frac{4\pi\mu}{\log\left(\frac{2h}{\epsilon}\right) - 1} \quad \text{if } L \gg h, \quad (2.35a,b)$$

where all coefficients are as summarised by Brennen & Winet (1977) and (2.35) and (2.33) additionally require $\epsilon \ll h$. In free space we will adopt the resistive force coefficients defined by the large- h limit of (2.32) and (2.34), and refer to this simpler theory as free-space resistive force theory (F-RFT). As given by Brenner and Katz *et al.*, the leading-order boundary corrections to C_t and C_n are $O(L/h)^3$ when $L \ll h$, and $O(h/L)$ when $L \gg h$, although the overall error in the resistive force approximation remains logarithmic in the filament aspect ratio.

3. Results and applications

3.1. Evaluation of wall-corrected resistive force theory for straight filaments

For the case of a straight uniform filament aligned parallel to a planar boundary, utilising the approach of § 2.2 we compute the hydrodynamic drag on the slender body as it moves parallel to the wall along its tangent at unit non-dimensional velocity, comparing the solutions given by the methods of regularised Stokeslet segments, free-space RFT and wall-corrected RFT, solving separately (2.12) and (2.31) for force density. Having normalised by the F-RFT solution, it being independent of the boundary separation h , we show the computed non-dimensional total drag in figure 3. Good agreement near the boundary can be seen between the W-RFT of Katz *et al.* and our implementation of the method of regularised Stokeslet segments, the former as previously validated in the limit $h/L \rightarrow 0$ with high-accuracy boundary element methods by Ramia *et al.* (1993). Indeed, direct comparison of the RSS solution with Ramia *et al.* (1993, figure 4c) provides additional verification of our methodology, where we note that for this comparison we are taking $\epsilon/L = 10^{-2}$ to ensure the filament radius matches that used in the boundary element computations. Further, far from the boundary the correction of Brenner (1962) lies within 10% of the RSS solution, evidencing good overall agreement between the two schemes as the normalised wall separation h/L increases. Analogous agreement between these methodologies can additionally be seen for total normal drag when moving normal to the boundary, thus we conclude that the corrections to resistive force theory of Brenner and Katz *et al.* agree closely with regularised Stokeslet segments for straight filaments in their respective asymptotic regimes. Hence, agreement in more generality may be reasonably hypothesised.

Expressions for corrected drag coefficients pertaining to filaments oriented perpendicular to a planar boundary are also available in the literature, and may be found summarised in the work of Brennen & Winet (1977). Repetition of the above validation, translating the filament along its axis and again comparing against the gold-standard boundary element method of Ramia *et al.* (1993), yields similar conclusions regarding the validity of near and far-field expressions for the resistive coefficients. The same good far-field agreement can be seen between the RSS solution and the corrected resistive force theory, however the near-boundary corrected theory has no dependence on boundary separation in the limit and thus does not capture near-boundary variations in drag. With significant experimental and theoretical study concerning filaments aligned approximately parallel to a boundary, for example the theoretical works of Elgeti *et al.* (2010) and typical videomicroscopy data of Ohmuro & Ishijima (2006) and Smith *et al.* (2009), herein we focus on the expressions given in § 2.6 and on the validity of resistive force theories in more generality.

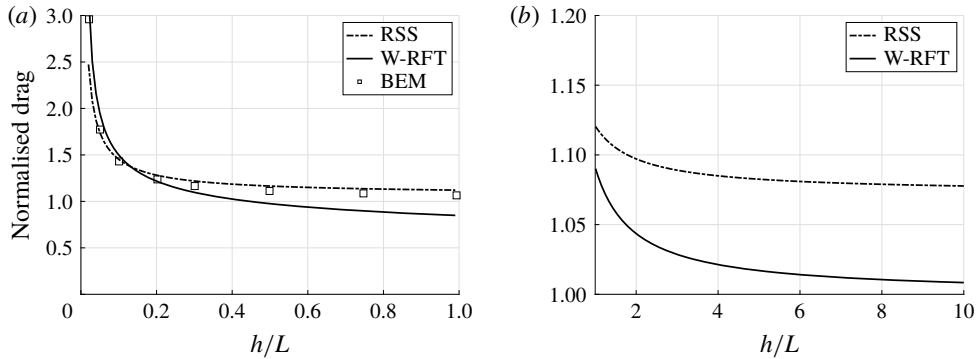


FIGURE 3. Computed total drag on a straight parallel filament moving along a boundary parallel to its tangent, here with $\epsilon/L = 10^{-2}$. (a) Near the boundary we see very good agreement between the regularised Stokeslet segment (RSS) solution (dot-dashed) and the wall-corrected resistive force theory (W-RFT) solution of Katz *et al.* (solid), with the latter corresponding to (2.33) and fair agreement maintained outside the region of validity of W-RFT as $h/L \sim 1$. The boundary element method (BEM) data of Ramia *et al.* are marked with squares, adapted from Ramia *et al.* (1993, figure 4c) and show good agreement with the regularised Stokeslet segments. (b) Similar agreement can be seen far from the boundary, with the asymptotic solution of Brenner given in (2.32) losing validity when $h/L \sim 1$ although remaining within 10% of the RSS solution.

3.2. Computing hydrodynamic drag on filaments parallel to boundaries

Following the established good agreement of W-RFT and RSS for straight filaments we proceed to evaluate the agreement for curved filaments moving in a plane parallel to the no-slip boundary, again solving (2.12) and (2.31). Motivated by the previous use of free-space resistive force theory for drag determination from captured kinematic data (Friedrich *et al.* 2010; Gaffney *et al.* 2011; Ishijima 2011; Ooi *et al.* 2014), we compute the forces on a free filament from the kinematic data of Ishimoto *et al.* (2017) corresponding to the flagellum of a human spermatozoon. These data, captured from free-swimming spermatozoa situated approximately 10–20 μm from a planar boundary, were smoothed by Ishimoto *et al.* (2017) and given in the form of an arclength parameterisation, with approximately 100 frames being recorded per beat cycle.

We first consider the filament moving in very close proximity to the boundary, with normalised separation $h/L = 0.01$, within the typical range of slithering motion for spermatozoa (Nosrati *et al.* 2015). A summary of the resulting drag calculations is shown in figure 4, from which we observe that the wall-corrected RFT of Katz *et al.* (1975) can demonstrate strong pointwise agreement with regularised Stokeslet segments over a single frame (figure 4a). However, calculations of the total drag force over the filament highlight a general trend of over-estimation of force densities by W-RFT, with the differences between W-RFT and RSS having a median of 37% over the 100 frame range shown here, with differences measured relative to the RSS value. Free-space resistive force theory appears to do the opposite, systematically underestimating the magnitude of total drag, with increased median deviation of 45% from the RSS solution. Most notable however is the tight clustering of the effective ratio of drag coefficients C_n/C_t , shown in figure 4(c), computed pointwise from the RSS solution and with an analogous distribution of the effective values of C_n/μ

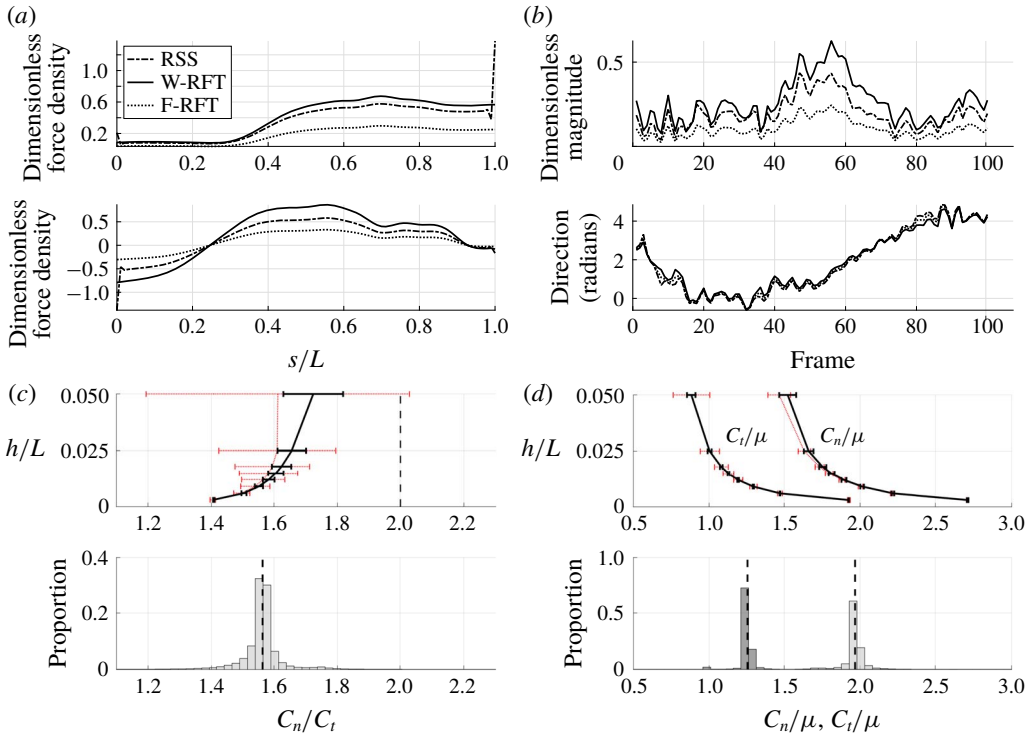


FIGURE 4. (Colour online) Drag calculations from kinematic data captured parallel to a planar boundary, with data from Ishimoto *et al.* (2017). (a) The tangential and normal components of the drag along the filament for a single frame (upper and lower panels respectively) at separation $h/L = 0.01$, showing good qualitative agreement between methodologies of drag computation and fair quantitative agreement between W-RFT and RSS. (b) The magnitude and direction of the total integrated force on the filament (upper and lower panels respectively) at separation $h/L = 0.01$, with direction measured relative to an arbitrary fixed axis. As in (a), agreement is fair and qualitative features are captured by all methodologies. Remarkably, the direction of the resultant force as computed using the resistive force theories strongly agrees with that of the non-local RSS. Endpoint effects are visible in the upper panel of (a), as discussed in detail in § 2.5. (c) Ratio of effective drag coefficients C_n/C_t , as computed via RSS, against boundary separation (upper), shown as the median over all frames and all material points (740 frames, 100 points per frame). The ratio as predicted by Katz *et al.*, 2, is shown as a dashed line for comparison. Error bars corresponding to half the interquartile range are shown, representative of dispersal about the median as the distributions do not appear significantly skewed. Modifying the kinematic data of Ishimoto *et al.* (2017) as described in the main text to simulate a pinned filament, analogous results are shown in red (dotted, thin), highlighting a reduction in validity of resistive force theories for pinned data but retaining notable accuracy when very close to the boundary. The lower panel is a histogram corresponding to $h/L = 0.01$, showing the distribution of effective coefficient ratios in the free-swimming case, highlighting tight grouping about the median. (d) Median values of effective resistive coefficients against boundary separation (upper), with error bars of half the interquartile range. Pinned data are shown in red (dotted, thin) for comparison. The lower panel is a histogram corresponding to $h/L = 0.01$, showing the tight distribution of dimensionless effective drag coefficients C_n/μ and C_t/μ , with C_t/μ shown darker. Medians are shown dashed.

and C_t/μ shown in figure 4(d). This suggests that an appropriate choice of resistive coefficient would enable the accurate approximation of local filament drag using only a local theory, in concurrence with the findings of Friedrich *et al.* (2010) though seen here much closer to the boundary and at a local level, rather than Friedrich *et al.*'s comparison to observed sperm behaviour, which necessarily averages over the cell. We verify this result using additional waveform data, modifying the kinematic data of Ishimoto *et al.* (2017) to crudely approximate a pinned spermatozoa by fixing both the endpoint and local tangent in space, along with the idealised pinned waveforms used by Curtis *et al.* (2012) given in appendix C. In particular, the tight clustering of effective coefficient ratios at low boundary separations is retained, though we note reduced agreement compared to that present for free-swimming data. Surprisingly, these coefficients and ratios do not align with the corrected coefficients of Katz *et al.* (1975), derived for straight filaments parallel to a boundary.

Additionally, for approximate separations $0.3 < h/L < 1$ we observe very strong agreement between resistive force theories and the non-local solution, with median differences in total drag between methodologies consistently less than 15%, in many cases being less than 5%. This is coupled with additional agreement concerning the direction of the resultant filament drag, which is retained even at reduced separations (figure 4b, lower). However, lost at such intermediate separations is any clear choice of effective resistive coefficient ratio, with the distribution of effective ratios significantly broadening above $h/L \approx 0.05$.

At a much greater distance from the boundary, with $h/L = 10$, as expected F-RFT and W-RFT give approximately equal estimates for the drag on the free-swimming filament, with the W-RFT solution having approached that of F-RFT. As in the case of near-boundary swimming, only small differences are present between the W-RFT and RSS solutions, with median differences between methods of approximately 6%, in this case with the magnitude of all computed drag forces having been reduced from their near-wall values by approximately a factor of two. Thus, in the medium and far field of a boundary resistive force theories appear remarkably accurate for determining the total drag on even curved filaments moving parallel to the boundary, although surprisingly at this increased boundary separation there is little grouping of the effective resistive coefficient ratios.

3.3. Hyperactivation-induced tugging of tethered spermatozoa

Explored initially by Curtis *et al.* (2012) using both free-space and wall-corrected resistive force theory, and reconsidered by Simons *et al.* (2014) using regularised Stokeslets, we re-evaluate the observation that the hyperactivation of mammalian spermatozoa aids in surface escape via a beat-induced tugging effect on the tether point of boundary-attached spermatozoa, and consider in detail the drag on the filament. Adopting the idealised beat patterns used by Curtis *et al.* and given in appendix C, appropriately non-dimensionalised, we position the base of the filament $0.01L$ from the boundary and compute both the total and local drag from this kinematic data for both hyperactivated and normal beating, noting that the plane of filament beating is perpendicular to the boundary. Mirroring the set-up of Curtis *et al.*, we assume that the filament is clamped at its base, implemented by rotating the kinematic data to align the basal tangent in each instant at some angle θ_0 to the boundary.

Figure 5 shows the results of the drag computation over a single beat period for both the hyperactivated and normal beat patterns for $\theta_0 = \pi/2$. We see reaffirmed

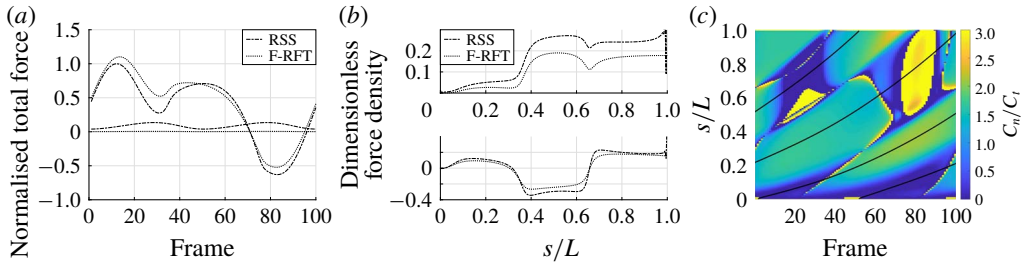


FIGURE 5. (Colour online) Drag computation for pinned spermatozoa with a base clamped perpendicular to a boundary over a single period of 100 frames. (a) Normalised total force on the boundary over a single beat period as computed by regularised Stokeslet segments (dot-dashed) and free-space resistive force theory (dotted), in the cases of hyperactive and normal beating patterns (high and low amplitudes respectively). Normal beating gives rise to a force on the boundary of unchanging sign, whilst hyperactivated beating generates a force whose sign changes over the beat period, corresponding to a so-called tugging effect. (b) Tangential (upper panel) and normal components (lower panel) of force density as a function of arclength for a single frame, shown for the hyperactivated beat pattern. In contrast to the fair agreement between methods seen in (a), with mean error of 24 % for the hyperactivated beat between F-RFT and RSS, tangential force density differs on average by 43 % between methods for the frame shown. (c) Ratio between effective drag coefficients computed using regularised Stokeslet segments, with contours of zero filament curvature superimposed as black curves, showing significant variation in the effective ratio.

by the method of regularised Stokeslet segments the conclusion of Curtis *et al.*, with the hyperactivated beat pattern exhibiting a change of sign in force component perpendicular to the boundary, whilst no such change is observed for the normally beating flagellum. Sampling $\theta_0 \in [\pi/4, \pi/2]$, we note that this observation holds across a range of basal orientations for these beat patterns, in agreement with the RFT-established conclusions of Curtis *et al.* Overall, figure 5(a) demonstrates fair agreement between the local free-space resistive force theory solution and that obtained using regularised Stokeslet segments, suggesting a surprising validity in using simple local theories in this circumstance, as concluded broadly by Simons *et al.* However, the pointwise values of the drag density highlight a stark disagreement between local and non-local theories, with deviations of up to 43 % for the tangential component of the drag for the frame of hyperactivated beating shown in figure 5(b). For reference, the integrated total drag along the filament on average gives differences of only 24 % between RSS and F-RFT, suggesting that a serendipitous cancellation occurs when integrating over the flagellum.

In figure 5(c) we show the ratio of effective tangential and normal drag coefficients for the hyperactivated beating, along with contours of zero curvature for reference. Significant variation in this ratio can be seen, in stark contrast to those computed for near-wall parallel swimming above. Thus the conclusion of Friedrich *et al.* (2010), that resistive force theory is accurate to high precision, does not hold for a bound spermatozoon, and hence the use of simple resistive force theories for near-boundary drag calculations is not reliable in general. This precludes the general use of the near-boundary correction of Katz *et al.* (1975) in this circumstance, supported by the poor agreement seen in figure 5, whilst the far-field result of Brenner (1962) is inappropriate in such close boundary proximity.

3.4. Morphological bifurcation of cantilevered filaments in an oscillatory flow

In the absence of a resistive force theory capable of accurately quantifying the details of pinned filament motion perpendicular to boundaries, we extensively utilise the non-local theory of regularised Stokeslet segments to consider the fully coupled elasto-hydrodynamics of a cantilevered filament in an oscillating background flow, as formulated in § 2.3. With a planar infinite no-slip boundary situated at $\hat{y} = 0$ and the filament base located at $(\hat{x}, \hat{y}) = (0, 0.01)$ to avoid numerical degeneracy, we consider the non-dimensional background flow with velocity $\hat{\mathbf{u}}_b = \hat{a} \sin(2\pi\hat{t} - \hat{t}_0)\hat{\mathbf{y}}\mathbf{e}_x$ for amplitude \hat{a} and phase \hat{t}_0 . Note that this flow preserves symmetry in the plane containing the filament, thus we may retain the assumption of filament planarity. Hence we have the modified non-dimensional system

$$-E_h \hat{\mathbf{B}}\hat{\mathbf{A}}^{-1} \hat{\mathbf{Q}}\hat{\boldsymbol{\theta}} = \hat{\mathbf{R}} - E_h \hat{\mathbf{B}}\hat{\mathbf{A}}^{-1} \hat{\mathbf{U}}_b, \tag{3.1}$$

where the components of $\hat{\mathbf{U}}_b$ are given by the background flow evaluated at the endpoints of filament segments, explicitly

$$\hat{\mathbf{U}}_b = \hat{a} \sin(2\pi\hat{t} - \hat{t}_0)(\hat{y}_1, 0, \hat{y}_2, \dots, \hat{y}_{N+1}, 0)^\top. \tag{3.2}$$

Noting that $\mathbf{n}(s_1)$ and $\mathbf{m}(s_1)$ are *a priori* unknown for a cantilevered filament, we impose the appropriate constraints of $\dot{x}_1 = \dot{y}_1 = \dot{\theta}_1 = 0$ in place of the total force and moment-balance equations, again yielding a square linear system. Under the assumption of a straight initial configuration with $\theta_i = \pi/2$ for $i = 1, \dots, N$, which we make throughout, there is freedom in the three non-dimensional parameters \hat{a} , \hat{t}_0 and E_h , the latter being equivalent to the sperm number Sp . In appendix A we consider in detail the effects on the dynamics of varying the phase \hat{t}_0 of the background flow, concluding that phase can effectively be neglected in long-time dynamics, and thus we will fix $\hat{t}_0 = 0$, noting that this breaks the inherent left–right symmetry of the initial condition.

For fixed \hat{a} we consider the beating of the filament as it is driven by the background flow with the sperm number being varied. With $\hat{a} = 2\pi$, we showcase in figure 6 the diverse range of observed behaviours. For low sperm numbers, in this case below a threshold value of around $Sp = 2.5$, we observe symmetric, low amplitude beating of the filament. Increasing the sperm number beyond this point results in the remarkable emergence of an asymmetric beating, shown in figure 6(b), strongly resembling the characteristic beating of a cilium. In particular, we see reproduced in this passive, flow-driven filament the defining power and recovery strokes of beating cilia (Brennen & Winet 1977).

Increasing the sperm number further results in the filament buckling and its beating regaining approximate symmetry, as can be seen in figure 6(c) for $Sp = 4.55$. For very high sperm numbers we see the appearance of buckling modes of higher order, resolved in figure 7 using $N = 150$ segments to capture filaments with such exceptionally high curvatures. With such high sperm numbers, we observe the collapse of modes onto those corresponding to lower-order buckling as time progresses, in addition to a greatly increased relaxation time to periodic beating in comparison to stiffer filaments.

In order to classify the mode of periodic beating we introduce the symmetry measure S , defined for our piecewise-linear filament with segment endpoint coordinates

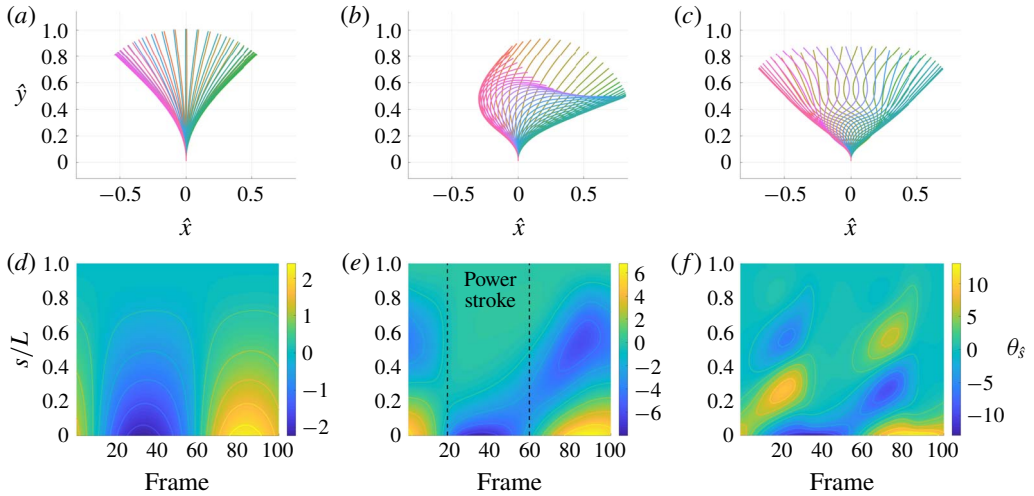


FIGURE 6. (Colour online) Periodic beating of a cantilevered filament in oscillatory background flow of amplitude $\hat{a} = 2\pi$, shown here over a single period of the background flow after convergence to periodic filament motion has occurred. (a) At low sperm numbers, here with $Sp = 1.14$, flow-driven beating is approximately symmetric. (b) For medium sperm numbers a remarkable ciliary-type asymmetric beating emerges, shown here for $Sp = 2.84$ and globally stable, with a clear distinction between effective forward and reverse strokes. (c) At higher sperm numbers the filament buckles significantly and regains a symmetric periodic beat, here with $Sp = 4.55$. (d–f) Plots of signed curvature corresponding to (a–c) above, sampled at 100 frames per period and each independently scaled for clarity, with select contours of curvature shown. Symmetry is clearly visible in (d) and (f), with the latter displaying a low-order buckling mode. (e) Exemplifies cilia-like beating, with the effective power stroke occurring approximately between frames 20 and 60 (shown black, dashed).

$\hat{\mathbf{x}}_i(\hat{t})$ as

$$S = 1 - \frac{1}{2} \int \frac{\sum_i |H\hat{\mathbf{x}}_i(\hat{t}) - \hat{\mathbf{x}}_i(\hat{t} + 1/2)|^2}{\sum_i |\hat{\mathbf{x}}_i(\hat{t})|^2} d\hat{t}, \tag{3.3}$$

where $H\hat{\mathbf{x}}_i$ denotes the reflection of $\hat{\mathbf{x}}_i$ in the plane $\hat{x} = 0$, and the integral over \hat{t} runs over half the period of the motion after convergence to periodic beating has occurred. We note that the left–right symmetric beats of figures 6(a) and 6(c) give $S = 1$, with motion that breaks left–right symmetry such as the distinct forward and reverse strokes of ciliary beating yielding reduced values of S . We find that values of S a small tolerance less than unity indeed correspond well to the ciliary-type beating of figure 6(b). We additionally distinguish between approximately straight and highly buckled beating by considering the number of filament regions with curvature of unchanging sign, classifying filaments with three or more such regions as buckled. Shown in figure 7(c) are the regions of the Sp – \hat{a} parameter space that approximately correspond to low-amplitude symmetric, ciliary and buckled modes of filament beating.

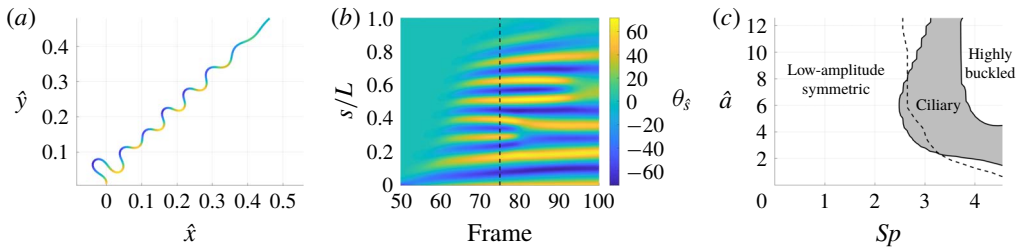


FIGURE 7. (Colour online) Bifurcation into high-order buckling modes for a filament with high sperm number, here with $Sp=16$ and sampled at 100 frames over the initial period of the background flow. (a) A single frame of the initial motion of the filament, where high-curvature regions can be seen to be resolved by the $N=150$ segments used here, with curvature represented by colour. (b) The curvature of the filament over the latter part of the initial period of the background flow, from which we may identify the collapse of high-order modes onto lower-order buckling modes. The frame shown in (a) is indicated on the curvature plot by a dashed line, with colour scalings consistent between plots. (c) A bifurcation diagram highlighting the regions of parameter space corresponding to low-amplitude symmetric, ciliary and buckled modes of beating. Buckled modes appear to the right of the dividing dashed line, as defined by the transition between less than three and three regions of constant curvature sign along the filament. Furthermore, ciliary beating occurs within contours delimiting where S is essentially unity, shown shaded. In the overlap region between ciliary and buckled beating we observe ciliary-type beats with buckled tips.

3.5. Time-averaged total drag on cantilevered filaments in oscillatory flow corresponds to distinct beating morphologies

Computation of the total hydrodynamic drag exerted on the filament by the background oscillatory flow over a single period reveals a bifurcation structure closely aligned with that of the morphology of beating. Shown in figure 8(a) and rescaled by flow amplitude, the normalised total drag in the direction parallel to the boundary is seen to be of constant sign, and minimal for the approximately symmetric beating present at low sperm numbers, with a region of non-trivial total drag appearing for $Sp \approx 1$ where elastic effects further distance the beating from reciprocal motion. Figure 8(c) demonstrates that such minimal total drag arises as the result of gross cancellation over the period of oscillation, consistent with the symmetry of the associated beating mode. Upon entering the region of parameter space consistent with ciliary-type beating there is a significant change in the magnitude of the experienced total drag, intuitively correlated with the emergence of the distinct forward and backward strokes of ciliary beating. With reference to the bifurcation diagram of figure 7(c), highly buckled beating at high values of Sp results in little total drag in the direction parallel to the boundary, consistent with the truly time-reversible motion obtained in the limit of (3.1) as $E_h \rightarrow \infty$. Notably, the direction of the time-averaged drag exerted on the filament over a period of the background flow points in the direction of the power stroke; with reference to the motion shown in figure 6(b), this corresponds to a resultant drag in the direction of increasing \hat{x} , opposite in direction to the resultant drag that might be expected to act on an actively beating cilium in otherwise stationary fluid.

Considering similarly the total force applied on the filament base in the direction normal to the boundary over a single beat period, we again observe that the total

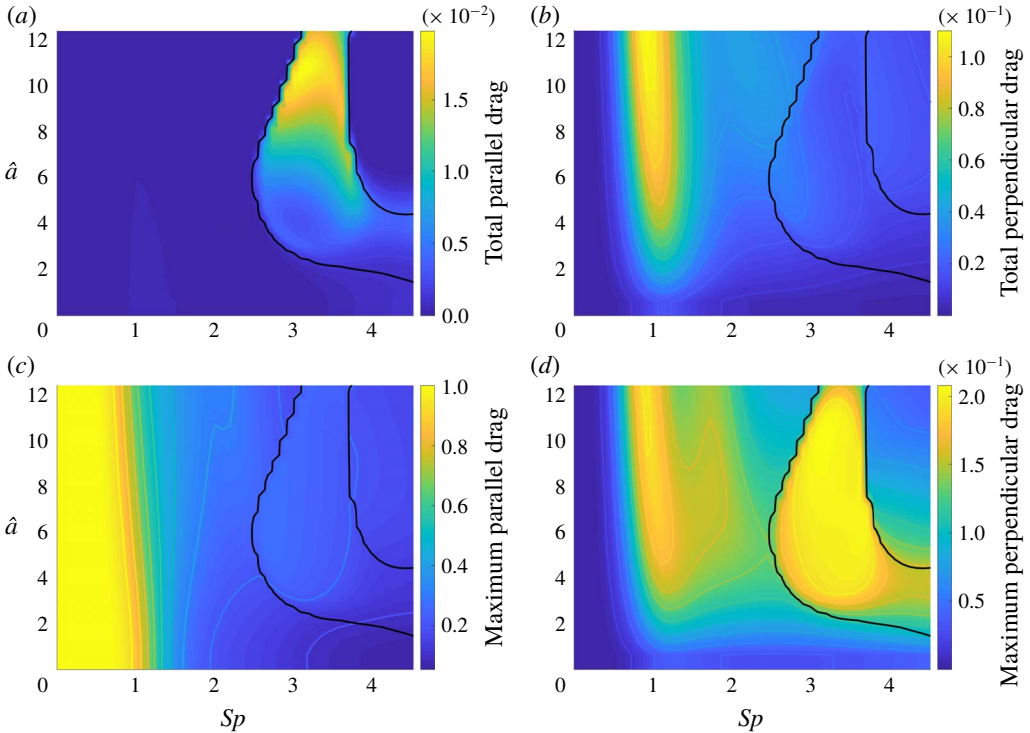


FIGURE 8. (Colour online) Hydrodynamic drag associated with cantilevered filaments in oscillating flow over a single period. (a) Total drag on the filament in the direction parallel to the boundary, averaged over a single period. A steep increase in total drag is noted to occur upon entering the parameter regime of ciliary beating, with a less-prominent local maximum present around $Sp = 1$. (b) Total pushing force applied on the base of the filament in the direction perpendicular to the boundary over a single period of the background flow. A clear maximum can be seen around $Sp = 1$, corresponding to the symmetric beating of stiff filaments, with the high sperm number border of the ciliary region being aligned with a sharp, albeit limited increase in total drag for $\hat{a} > 6$. (c) Maximum instantaneous total drag on the filament in the direction parallel to the boundary. Comparison with (a) highlights gross total drag cancellation occurring for low Sp , with reduced cancellation for the asymmetric ciliary-type beating. (d) Maximum absolute instantaneous force applied on the base of the filament in the direction perpendicular to the boundary. By comparison with (b) we observe limited cancellation of total drag in this direction in stiff filaments, with significant cancellation occurring for ciliary-type beating. Each panel is normalised by the maximum instantaneous total parallel drag and rescaled by flow amplitude to enable meaningful comparison. Select contours of total drag are shown and the region of ciliary-type beating is outlined in black, the latter determined only from the kinematic symmetry measure S .

force is of constant sign, corresponding to the filament pushing towards the boundary. Shown in figure 8(b), the maximal force occurs around $Sp = 1$ for the majority of sampled flow amplitudes, aligning with the local maximum of total parallel drag noted previously and pertaining to the non-reciprocal symmetric beating of stiff filaments. Comparison with figure 8(d) reveals in this case that significant total drag cancellation occurs for the asymmetric ciliary-type beating. In contrast, the total drag on stiffer

filaments in this direction sees limited cancellation, exemplifying the significance of beating morphology in filament drag calculations.

4. Discussion

In this work we have examined the accuracy of resistive force theories in quantifying the mechanics of planar filament motion in a half space via regularised Stokeslet segments, further exploring the coupled non-local elastohydrodynamics of cantilevered filaments in an oscillatory flow. We have seen that the corrections to free-space resistive force theory of Brenner (1962) and Katz *et al.* (1975) perform well both in the near and far field of planar boundaries when applied to straight filaments, in agreement with the verification of Ramia *et al.* (1993) and serving as additional validation of our methodology.

By considering the motion of curved filaments in planes parallel to a boundary, in line with our hypothesis we have found that these corrections capture quantitative filament dynamics to moderate accuracy in all but the most extreme boundary proximities. Remarkably however, when very close to the boundary we have nonetheless observed a tightly distributed ratio of effective drag coefficients, and indeed tightly distributed coefficients, suggesting that a resistive force theory with such coefficients could yield accurate estimates for hydrodynamic drag in this circumstance. With subjects imaged in the relative near field of a coverslip, the similar conclusion of Friedrich *et al.* (2010) is thus in part supported by our non-local calculations, albeit here at greatly reduced boundary separations, and suggests the future development of an accurate, empirical resistive force theory in the near field of boundaries. This has potential application to the study of a slithering mode of swimming, as reported to be prevalent in spermatozoa by Nosrati *et al.* (2015) although this mode is far from ubiquitous. Analytical exploration of this result would require consideration of the asymptotic limit where the separation of singularities from their images is much less than the length scale of the filament radius of curvature, with such detailed calculations expected to be a subject of significant future study.

We have seen that the agreement between methodologies for parallel-swimming filaments does not carry through to those moving in a plane perpendicular to the boundary, with no clear consensus on effective coefficient ratio for pinned filaments moving perpendicular to boundaries. Thus we conclude that constant-coefficient resistive force theories cannot be expected to give accuracy comparable to non-local methods for filaments moving perpendicular to boundaries. Despite this, here we have reverified the conclusion of Curtis *et al.* (2012), based originally on resistive force theory, although this appears to rely on serendipitous cancellations on integrating along the filament.

In particular, we have noted a potential lack of reliability in local drag theories when applied to pinned filaments, as evidenced by the loss of coherence in effective drag coefficient ratio when using artificially pinned kinematic data in § 3.2, supported further by the findings of § 3.3 for tethered spermatozoa. Accordingly, we have utilised the non-local hydrodynamics of Cortez (2018) to examine the response of a cantilevered elastic filament to an oscillatory background flow, with this application highlighting the flexibility in our treatment of the coupled elastohydrodynamics. Having established convergence to a single limiting periodic behaviour from a range of initial conditions for fixed filament and flow parameters, as detailed in appendix A, we have evidenced in passive flow-driven filaments the existence of a remarkable mode of beating typically characteristic of actively beating cilia. Notably, the fact

that the movement of our passive filaments appears qualitatively similar to that of actively beating cilia highlights that one cannot naively infer internal mechanics by simply observing filament motion alone.

The asymmetric ciliary-type mode of beating was found to be accompanied by two distinct highly symmetric beating modes. Most prevalent in the $Sp-\hat{a}$ parameter space was that of low-amplitude beating, with the pinned filament retaining low curvature throughout and generating the maximal time-averaged total normal force into the boundary amongst beating morphologies. Highly buckled modes corresponding to less-rigid filaments produced minimal time-averaged total drag over each period of the background flow, with the high-order buckling modes still resolved by our piecewise-linear formulation of the governing elastohydrodynamics. The near symmetry of each of these modes resulted in little time-averaged total drag in the direction parallel to the boundary. In contrast, entering the region of parameter space corresponding to ciliary beating is seen to strongly correlate with a sharp increase in time-averaged total parallel drag, intuitively the result of kinematic asymmetry arising from the distinct power and recovery strokes of ciliary beating. Hence, elastohydrodynamically induced morphological changes in general have a dramatic effect on filament drag mechanics. Furthermore, the efficiency of the presented methodology would facilitate in-depth mechanical studies of primary cilia with regards to transmitted force and mechanotransduction, noting that the interaction of bending modulus variations, effective boundary conditions at the ciliary base, changes in cross-section of the cilium and cilium length are reported to be under-explored even in the restricted context of renal physiology (Nag & Resnick 2017). Additionally, the methodology as presented here is readily extensible, with the potential to enable efficient multi-filament elastohydrodynamic simulations applicable to the study of a variety of physical and biological systems, such as actively driven ciliary carpets or carbon nanotube mats. An interesting additional direction, although requiring significant further modelling development, is generalising this framework to non-planar filament motions, as would occur with symmetry-breaking background flows.

In summary, we have considered in detail the validity of traditional and corrected resistive force theories for filaments in the presence of a planar boundary, concluding that, whilst in general these local drag theories may not be relied upon to perform well against non-local solutions for curved filaments, for filament motion parallel to a boundary the use of a resistive force theory may be remarkably accurate with appropriately chosen coefficients, and viable over a range of boundary separations. Verified against previous high-accuracy methods, we have additionally presented an efficient non-local formulation of the governing elastohydrodynamics in a half-space, exemplifying its flexibility by application to a cantilevered filament in oscillatory flow. This study of passive filaments revealed a surprising asymmetric beating morphology found typically in active cilia, highlighting a complex relationship between passive elastic fibres, internally forced filaments, the flows that they drive and the forces that they induce.

Acknowledgements

B.J.W. is supported by the UK Engineering and Physical Sciences Research Council (EPSRC), grant EP/N509711/1. K.I. is supported by JSPS-KAKENHI for Young Researchers (18K13456), JSPS Overseas Research Fellowship (29-0146) and MEXT Leading Initiative for Excellent Young Researchers (LEADER).

The research materials supporting this publication can be accessed at <http://dx.doi.org/10.5287/bodleian:MzdNe99JY>.

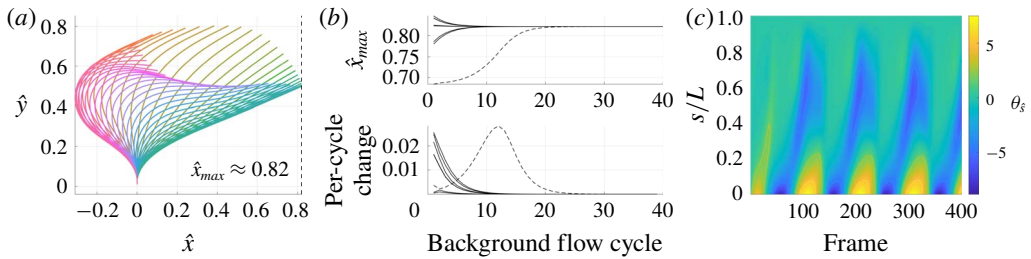


FIGURE 9. (Colour online) Evaluating the sensitivity of long-time behaviour to background flow phase. (a) An illustration of the measure \hat{x}_{max} used to determine periodicity and convergence of filament behaviour, equal to the maximal displacement of the filament tip from the centreline over one period of the background flow. Filament configurations throughout one period of motion are shown as coloured curves. (b) The evolution of \hat{x}_{max} over 40 periods of the background flow, for various choices of initial phase $\hat{t}_0 \in [0, \pi]$, here for $Sp = 3.13$, $\hat{a} = 2\pi$. In the upper panel we see that all choices of \hat{t}_0 lead to convergence to the same behaviour, with even the approximate initial symmetry of the $\hat{t}_0 = \pi/2$ instance (dashed) collapsing onto the common limiting behaviour. *A posteriori* validation of convergence to periodic motion from all phases is provided by consideration of the per cycle change in \hat{x}_{max} , as shown in the lower panel and highlighting eventual periodicity. Symmetric counterparts for $\hat{t}_0 \in [\pi, 2\pi]$ have been omitted. (c) Signed curvature of the filament during the first periods of the background flow, with select contours of curvature shown, and sampled at 100 frames per period for $\hat{t}_0 = 3\pi/8$, showing the transition to ciliary beating from a straight initial configuration.

Supplementary material

Supplementary material is available at <https://doi.org/10.1017/jfm.2019.723>.

Appendix A. Influence of flow phase on cantilevered filament dynamics

One might expect the phase \hat{t}_0 of the background oscillatory flow of § 3.4 to have significant impact on long-time filament dynamics, the most notable of such effects being the selection of left–right polarity in resulting behaviours. We examine the long-time dynamics for a range of phases, flow amplitudes and elasto-hydrodynamic numbers, tracking in particular the maximally attained displacement of the filament tip from the centreline $\hat{x} = 0$ throughout each period of the background flow, denoted \hat{x}_{max} and exemplified by figure 9(a). In particular, convergence of \hat{x}_{max} to a constant would serve to validate, *a posteriori*, the assumption of periodic motion. Sampling Sp and \hat{a} each from the range $[0.5, 12]$, as expected we observe a duality in solutions inherited from the left–right symmetry of the problem, and we additionally note the convergence of all solutions to a single periodic motion, where the period is aligned with that of the background flow to working precision. This is demonstrated by figure 9(b), which shows the evolution of \hat{x}_{max} over a number of cycles of the background flow for $Sp = 3.13$ and $\hat{a} = 2\pi$. We see convergence of all solutions to a common limit cycle of periodic behaviour, with even the approximately symmetric transient motion of $\hat{t}_0 = \pi/2$ (shown dashed) eventually collapsing onto the same mode of behaviour. Hence we resolve that exploration of the Sp – \hat{a} parameter space, holding \hat{t}_0 constant, is sufficient to capture the long-time behaviours of the cantilevered filament system.

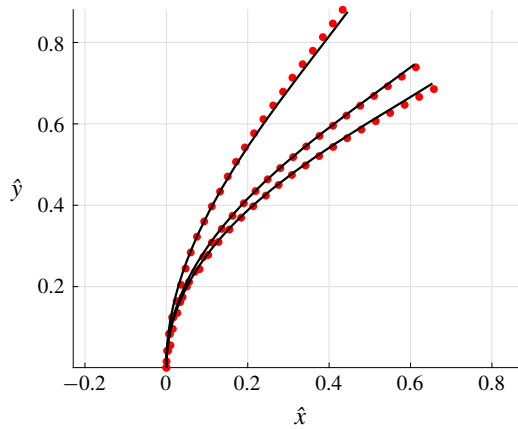


FIGURE 10. (Colour online) The limiting deflections of cantilevered filaments in background shear flows, with elastohydrodynamic numbers $E_h \approx 34, 68, 85$ (black, solid), compared with data adapted from Pozrikidis (2010, figure 11) and corresponding to $\hat{E}_B = 1.0, 0.5, 0.4$ from the publication of Pozrikidis. The latter data are shown as red circles, with good agreement between solutions, serving as validation of our treatment and implementation of the governing elastohydrodynamics. Here $N = 80$ and $\epsilon/L = 0.05$, the latter in line with the set-up of Pozrikidis (2010).

Appendix B. Validation of elastohydrodynamics in a half-space

Here we present an example verifying our implementation of our methodology for simulating filament elastohydrodynamics in a half-space via comparison to the boundary element computations of Pozrikidis, with data adapted from Pozrikidis (2010, figure 11). Given a stationary background shear flow of unit dimensionless shearing rate, so that $\hat{u}_b = \hat{y}e_x$ in the notation of § 3.4, we simulate the motion of a passive cantilevered filament attached to a planar boundary until convergence of the filament to a stationary solution. Taking $E_h \approx 34, 68, 85$ in turn, in figure 10 we show the resulting filament shapes against those found in the work of Pozrikidis (2010). The remarkable agreement between the boundary element computations of Pozrikidis and our solution for the filament shape serves as a further validation of our methodology, and in particular demonstrates the capability of our proposed scheme to accommodate non-trivial background flows and filament boundary conditions.

Appendix C. Kinematic data used in Curtis *et al.*

With \hat{x}, \hat{y} here denoting dimensionless coordinates, the flagellar waveforms of Curtis *et al.* (2012) used in §§ 3.2 and 3.3 may be parameterised as

$$\hat{x}(\xi, \hat{t}) = \xi, \tag{C 1}$$

$$\hat{y}(\xi, \hat{t}) = A_N \xi \sin(B_N \xi - \hat{t}), \tag{C 2}$$

for normal beating, and

$$\hat{x}(\xi, \hat{t}) = \xi, \tag{C 3}$$

$$\hat{y}(\xi, \hat{t}) = A_H \xi [1 - \cos(B_H \xi - \hat{t})], \tag{C 4}$$

in the case of hyperactive beating. Here $\xi \in [0, \xi^*]$, where ξ^* is taken to preserve a total dimensionless arclength of 1, and \hat{t} denotes dimensionless time. Following Curtis *et al.* (2012), where data from Ohmuro & Ishijima (2006) and Smith *et al.* (2009) have been considered, we have parameters $(A_N, B_N) = (0.1, 2.6\pi)$ and $(A_H, B_H) = (0.4, 4.24\pi)$.

REFERENCES

- AINLEY, J., DURKIN, S., EMBID, R., BOINDALA, P. & CORTEZ, R. 2008 The method of images for regularized Stokeslets. *J. Comput. Phys.* **227** (9), 4600–4616.
- BALAZS, A. C., BHATTACHARYA, A., TRIPATHI, A. & SHUM, H. 2014 Designing bioinspired artificial cilia to regulate particle–surface interactions. *J. Phys. Chem. Lett.* **5** (10), 1691–1700.
- BERG, H. C. & ANDERSON, R. A. 1973 Bacteria swim by rotating their flagellar filaments. *Nature* **245** (5425), 380–382.
- BRENNEN, C. & WINET, H. 1977 Fluid mechanics of propulsion by cilia and flagella. *Annu. Rev. Fluid Mech.* **9** (1), 339–398.
- BRENNER, H. 1962 Effect of finite boundaries on the Stokes resistance of an arbitrary particle. *J. Fluid Mech.* **12** (01), 35–48.
- CORTEZ, R. 2001 The method of regularized Stokeslets. *SIAM J. Sci. Comput.* **23** (4), 1204–1225.
- CORTEZ, R. 2018 Regularized Stokeslet segments. *J. Comput. Phys.* **375**, 783–796.
- CURTIS, M. P., KIRKMAN-BROWN, J. C., CONNOLLY, T. J. & GAFFNEY, E. A. 2012 Modelling a tethered mammalian sperm cell undergoing hyperactivation. *J. Theor. Biol.* **309**, 1–10.
- DELMOTTE, B., CLIMENT, E. & PLOURABOUÉ, F. 2015 A general formulation of Bead Models applied to flexible fibers and active filaments at low Reynolds number. *J. Comput. Phys.* **286**, 14–37.
- ELGETI, J., KAUPP, U. B. & GOMPPER, G. 2010 Hydrodynamics of sperm cells near surfaces. *Biophys. J.* **99** (4), 1018–1026.
- FAUCI, L. J. & MCDONALD, A. 1995 Sperm motility in the presence of boundaries. *Bull. Math. Biol.* **57** (5), 679–699.
- FRIEDRICH, B. M., RIEDEL-KRUSE, I. H., HOWARD, J. & JULICHER, F. 2010 High-precision tracking of sperm swimming fine structure provides strong test of resistive force theory. *J. Expl Biol.* **213** (8), 1226–1234.
- GADÉLHA, H., GAFFNEY, E. A., SMITH, D. J. & KIRKMAN-BROWN, J. C. 2010 Nonlinear instability in flagellar dynamics: a novel modulation mechanism in sperm migration? *J. R. Soc. Interface* **7** (53), 1689–1697.
- GAFFNEY, E. A., GADÉLHA, H., SMITH, D. J., BLAKE, J. R. & KIRKMAN-BROWN, J. C. 2011 Mammalian sperm motility: observation and theory. *Annu. Rev. Fluid Mech.* **43** (1), 501–528.
- GRAY, J. 1928 *Ciliary Movement*. Cambridge University Press.
- GRAY, J. & HANCOCK, G. J. 1955 The propulsion of sea-urchin spermatozoa. *J. Expl Biol.* **32** (4), 802–814.
- GUGLIELMINI, L., KUSHWAHA, A., SHAQFEH, E. S. G. & STONE, H. A. 2012 Buckling transitions of an elastic filament in a viscous stagnation point flow. *Phys. Fluids* **24** (12), 123601.
- HALL-MCNAIR, A. L., GALLAGHER, M. T., MONTENEGRO-JOHNSON, T. D., GADÉLHA, H. & SMITH, D. J. 2019 Efficient implementation of elastohydrodynamics via integral operators. 1–43; [arXiv:1903.03427](https://arxiv.org/abs/1903.03427).
- HANCOCK, G. J. 1953 The self-propulsion of microscopic organisms through liquids. *Proc. R. Soc. Lond. Ser. A* **217** (1128), 96–121.
- ISHIJIMA, S. 2011 Dynamics of flagellar force generated by a hyperactivated spermatozoon. *Reproduction* **142** (3), 409–415.
- ISHIMOTO, K., GADÉLHA, H., GAFFNEY, E. A., SMITH, D. J. & KIRKMAN-BROWN, J. 2017 Coarse-graining the fluid flow around a human sperm. *Phys. Rev. Lett.* **118** (12), 124501.
- ISHIMOTO, K. & GAFFNEY, E. A. 2015 Fluid flow and sperm guidance: a simulation study of hydrodynamic sperm rheotaxis. *J. R. Soc. Interface* **12** (106), 20150172.

- ISHIMOTO, K. & GAFFNEY, E. A. 2016 Mechanical tuning of mammalian sperm behaviour by hyperactivation, rheology and substrate adhesion: a numerical exploration. *J. R. Soc. Interface* **13** (124), 20160633.
- ISHIMOTO, K. & GAFFNEY, E. A. 2018 An elastohydrodynamical simulation study of filament and spermatozoan swimming driven by internal couples. *IMA J. Appl. Maths* **83** (4), 655–679.
- JOHNSON, R. E. & BROKAW, C. J. 1979 Flagellar hydrodynamics. A comparison between resistive-force theory and slender-body theory. *Biophys. J.* **25** (1), 113–127.
- KATZ, D. F., BLAKE, J. R. & PAVERI-FONTANA, S. L. 1975 On the movement of slender bodies near plane boundaries at low Reynolds number. *J. Fluid Mech.* **72** (03), 529–540.
- LAUGA, E., DILUZIO, W. R., WHITESIDES, G. M. & STONE, H. A. 2006 Swimming in circles: motion of bacteria near solid boundaries. *Biophys. J.* **90** (2), 400–412.
- LIU, Y., CHAKRABARTI, B., SAINTILLAN, D., LINDNER, A. & DU ROURE, O. 2018 Morphological transitions of elastic filaments in shear flow. *Proc. Natl Acad. Sci. USA* **115** (38), 9438–9443.
- MOREAU, C., GIRALDI, L. & GADÊLHA, H. 2018 The asymptotic coarse-graining formulation of slender-rods, bio-filaments and flagella. *J. R. Soc. Interface* **15** (144), 20180235.
- NAG, S. & RESNICK, A. 2017 Biophysics and biofluid dynamics of primary cilia: evidence for and against the flow-sensing function. *Am. J. Physiol.-Renal Physiol.* **313** (3), F706–F720.
- NOSRATI, R., DRIOUCHI, A., YIP, C. M. & SINTON, D. 2015 Two-dimensional slither swimming of sperm within a micrometre of a surface. *Nat. Commun.* **6** (1), 8703.
- OHMURO, J. & ISHIJIMA, S. 2006 Hyperactivation is the mode conversion from constant-curvature beating to constant-frequency beating under a constant rate of microtubule sliding. *Mol. Reproduction Dev.* **73** (11), 1412–1421.
- OLSON, S. D., LIM, S. & CORTEZ, R. 2013 Modeling the dynamics of an elastic rod with intrinsic curvature and twist using a regularized Stokes formulation. *J. Comput. Phys.* **238**, 169–187.
- OOI, E. H., SMITH, D. J., GADELHA, H., GAFFNEY, E. A. & KIRKMAN-BROWN, J. 2014 The mechanics of hyperactivation in adhered human sperm. *R. Soc. Open Sci.* **1** (2), 140230.
- POZRIKIDIS, C. 2010 Shear flow over cylindrical rods attached to a substrate. *J. Fluids Struct.* **26** (3), 393–405.
- POZRIKIDIS, C. 2011 Shear flow past slender elastic rods attached to a plane. *Intl J. Solids Struct.* **48** (1), 137–143.
- RAMIA, M., TULLOCK, D. L. & PHAN-THIEN, N. 1993 The role of hydrodynamic interaction in the locomotion of microorganisms. *Biophys. J.* **65** (2), 755–778.
- RIEDEL-KRUSE, I. H. & HILFINGER, A. 2007 How molecular motors shape the flagellar beat. *HFSP J.* **1** (3), 192–208.
- ROPER, M., DREYFUS, R., BAUDRY, J., FERMIGIER, M., BIBETTE, J. & STONE, H. A. 2006 On the dynamics of magnetically driven elastic filaments. *J. Fluid Mech.* **554**, 167–190.
- DU ROURE, O., LINDNER, A., NAZOCKDAST, E. N. & SHELLEY, M. J. 2019 Dynamics of flexible fibers in viscous flows and fluids. *Annu. Rev. Fluid Mech.* **51** (1), 539–572.
- SCHULMAN, R. D., BACKHOLM, M., RYU, W. S. & DALNOKI-VERESS, K. 2014 Undulatory microswimming near solid boundaries. *Phys. Fluids* **26** (10), 101902.
- SHAMPINE, L. F. & REICHEL, M. W. 1997 The MATLAB ODE Suite. *SIAM J. Sci. Comput.* **18** (1), 1–22.
- SHUM, H., TRIPATHI, A., YEOMANS, J. M. & BALAZS, A. C. 2013 Active ciliated surfaces expel model swimmers. *Langmuir* **29** (41), 12770–12776.
- SIMONS, J., OLSON, S., CORTEZ, R. & FAUCI, L. 2014 The dynamics of sperm detachment from epithelium in a coupled fluid-biochemical model of hyperactivated motility. *J. Theor. Biol.* **354**, 81–94.
- SMITH, D. J. 2009 A boundary element regularized Stokeslet method applied to cilia- and flagella-driven flow. *Proc. R. Soc. A* **465** (2112), 3605–3626.
- SMITH, D. J., GAFFNEY, E. A., BLAKE, J. R. & KIRKMAN-BROWN, J. C. 2009 Human sperm accumulation near surfaces: a simulation study. *J. Fluid Mech.* **621**, 289–320.
- SMITH, D. J., MONTENEGRO-JOHNSON, T. D. & LOPES, S. S. 2019 Symmetry-breaking cilia-driven flow in embryogenesis. *Annu. Rev. Fluid Mech.* **51** (1), 105–128.

- SZNITMAN, J., SHEN, X., SZNITMAN, R. & ARRATIA, P. E. 2010 Propulsive force measurements and flow behavior of undulatory swimmers at low Reynolds number. *Phys. Fluids* **22** (12), 121901.
- TORNBERG, A. K. & SHELLEY, M. J. 2004 Simulating the dynamics and interactions of flexible fibers in Stokes flows. *J. Comput. Phys.* **196** (1), 8–40.
- UTADA, A. S., BENNETT, R. R., FONG, J. C. N., GIBIANSKY, M. L., YILDIZ, F. H., GOLESTANIAN, R. & WONG, G. C. L. 2014 *Vibrio cholerae* use pili and flagella synergistically to effect motility switching and conditional surface attachment. *Nat. Commun.* **5** (1), 4913.
- YONEKURA, K., MAKI-YONEKURA, S. & NAMBA, K. 2003 Complete atomic model of the bacterial flagellar filament by electron cryomicroscopy. *Nature* **424** (6949), 643–650.