Autonomous Motility of Active Filaments due to Spontaneous Flow-Symmetry Breaking

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We simulate the nonlocal Stokesian hydrodynamics of an elastic filament which is active due a permanent distribution of stresslets along its contour. A bending instability of an initially straight filament spontaneously breaks flow symmetry and leads to autonomous filament motion which, depending on conformational symmetry, can be translational or rotational. At high ratios of activity to elasticity, the linear instability develops into nonlinear fluctuating states with large amplitude deformations. The dynamics of these states can be qualitatively understood as a superposition of translational and rotational motion associated with filament conformational modes of opposite symmetry. Our results can be tested in molecular-motor filament mixtures, synthetic chains of autocatalytic particles, or other linearly connected systems where chemical energy is converted to mechanical energy in a fluid environment.

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Components which convert chemical energy to mechanical energy internally are ubiquitous in biology. Common examples where this conversion leads to autonomous propulsion are molecular motors (at the subcellular level) and bacteria (at the cellular level) [1]. Recently, biomimetic elements which convert chemical energy into translational [2] or rotational [3] motion have been realized in the laboratory. While the detailed mechanisms leading to autonomous propulsion in these biological and soft matter systems show a wonderful variety [4], their collective behavior tends to be universal and can be understood by appealing to symmetries and conservation laws [5]. This realization has led to many studies of the collective properties of suspensions of hydrodynamically interacting autonomously motile particles [6].

There are ample instances in biology, however, where the conversion of chemical to mechanical energy is not confined to a particle-like element but is, instead, distributed over a linelike element. Such a situation arises, for example, in a microtubule with a row of molecular motors converting energy while walking on it. The mechanical energy thus obtained not only produces motion of the motors but also generates reaction forces on the microtubule, which can deform elastically in response. Hydrodynamic interactions between the motors and between segments of the microtubule must be taken into account since both are surrounded by a fluid. This combination of elasticity, autonomous motility through energy conversion, and hydrodynamics is found in biomimetic contexts as well. A recent example is provided by mixtures of motors which crosslink and walk on polymer bundles. A remarkable cilia-like beating phenomenon is observed in these systems [7]. A polymer in which the monomeric units are autocatalytic nanorods provides a nonbiological example of energy conversion on linear elastic elements. Though such elements are yet to be realized in the laboratory, active elements coupled to passive components through covalent bonds have been synthesized [2] and may lead to new kinds of nanomachines [3].

Motivated by these biological and biomimetic examples, we study, in this Letter, a semiflexible elastic filament immersed in a viscous fluid with energy converting active elements distributed along its length. We present an equation of motion for the filament that incorporates the effects of nonlinear elastic deformation, active processes, and nonlocal Stokesian hydrodynamic interactions. We use the lattice Boltzmann (LB) method to numerically solve the active filament equation of motion. Our simulations show that a straight active filament is linearly unstable to transverse perturbations. Depending on the symmetry of the perturbation, the hydrodynamic flows produce translational or rotational motion of the filament. Conformational symmetry provides a qualitative way of understanding the dynamics of the nonlinear fluctuating states that develop at high ratios of activity to elasticity. We describe these results and our model in detail below.

Model.—Our model for the active filament consists of \( N \) beads, with coordinates \( r_m \), interacting through a potential given by

\[
U(r_1, \ldots, r_N) = \sum_{m=1}^{N-1} U_S(b_m) + \sum_{m=1}^{N-2} U_B(b_m, b_{m+1}) + \frac{1}{2} \sum_{m,n=1}^{N} U_{LJ} (r_n - r_m).
\]

(1)

The two-body harmonic spring potential \( U_S(b_m) = \frac{1}{2} k (b_m - b_0)^2 \) penalizes departures of \( b_m \), the modulus of the bond vector \( b_m = |r_m - r_{m+1}| \), from its equilibrium value of \( b_0 \). The three-body bending potential \( U_B(b_m, b_{m+1}) = k (1 - \cos \phi_m) \) penalizes departures of
the angle $\phi_m$ between consecutive bond vectors from its equilibrium value of zero. The rigidity parameter $\kappa$ is related to the bending rigidity as $\kappa = b_0 \kappa$. The repulsive Lennard-Jones potential $U_{\text{LJ}}$ vanishes if the distance between beads $r_{nm} = |r_m - r_n|$ exceeds $\sigma_{\text{LJ}}$. The nth bead experiences a force $f_n = -\partial U/\partial r_n$ when the filament stretches or bends from its equilibrium position. With the above choice of potential the connected beads approximate an inextensible, semiflexible, self-avoiding filament.

Active nonequilibrium processes, such as those that convert chemical energy to mechanical energy, are internal to the fluid and hence cannot add net momentum to it. Then, the integral of the force density on a surface enclosing the active element must vanish. This is ensured if the active force density is the divergence of a stress. Since the active processes cannot add angular momentum to the fluid, the stress must be symmetric [8]. The most dominant Stokesian processes cannot add net momentum to it. Then,\[\text{Integral of force density on a surface enclosing the fluid and hence cannot add net momentum to it.}\]

The tangential stresses exerted by motors walking on microtubules [7], we choose the activity. The results of other choices of sign and orientation will be presented elsewhere.

Elastic forces and active stresses produce velocities in the fluid. In the Stokesian regime, the velocity in a threedimensional unbounded fluid at location $r$ produced by a force $f$ at the origin is $v_o(r) = O_{\alpha\beta}(r)f_\beta$, where $O_{\alpha\beta}(r) = (\delta_{\alpha\beta} + \hat{r}_\alpha \hat{r}_\beta)/8\pi \eta r$ is the Oseen tensor, Cartesian directions are indicated by Greek indices, $\eta$ is the fluid shear viscosity, and $\hat{r}_\alpha = r_\alpha/r$. Similarly, the velocity at location $r$ produced by a stresslet $\sigma$ at the origin is $v_o(r) = D_{\alpha\beta\gamma}(r)\sigma_{\gamma\beta}$, where $D_{\alpha\beta\gamma}(r) = (-\hat{r}_\alpha \delta_{\beta\gamma} + 3\hat{r}_\alpha \hat{r}_\beta \hat{r}_\gamma)/8\pi \eta r^2$ [10]. In the presence of rigid or periodic boundaries the tensors $O$ and $D$ must be replaced by the appropriate Green’s functions of Stokes flow that vanish at the boundaries or have the periodicity of the domain [10]. Similarly, two-dimensional Green’s functions must be used when studying the motion of filaments in planar flows [9]. The velocity of the nth bead is obtained by summing the force and activity contributions from all beads, including itself, to the fluid velocity at its location. An isolated spherical bead with a force $f$ acquires a velocity $\mu f$ where $\mu$ is its mobility. By symmetry, an isolated spherical bead with a stresslet $\sigma$ cannot acquire a velocity. This gives the following equation of motion for the active filament:

$$\dot{r}_n = \sum_{m=1}^{N} [O(r_n - r_m) \cdot f_m + D(r_n - r_m) \cdot \sigma_m],$$

where, for $m = n$, $O_{\alpha\beta} = \mu \delta_{\alpha\beta}$ and $D_{\alpha\beta\gamma} = 0$.

Equations (1)–(3) represent our model for the nonlocal Stokesian hydrodynamics of an active elastic filament. In the absence of bending rigidity and activity, our model reduces to Zimm dynamics of a polymer in a good solvent [11].

The ratio of the stresslet and Stokeslet terms in the equation of motion is a dimensionless measure of activity. Estimating the curvature elastic force as $\kappa/L^2$, where $L = (N - 1)b_0$ is the length of the filament, yields the activity number $\mathcal{A} = L\sigma_0/\kappa$. The rates of active and elastic relaxation are $\Gamma_\sigma = \sigma_0/\eta L^d$ and $\Gamma_\kappa = \kappa/\eta L^{d+1}$, respectively. Since $\mathcal{A} = \Gamma_\sigma/\Gamma_\kappa$, the activity number also

![FIG. 1 (color online). Spontaneous symmetry breaking due to a linear instability. (a) A straight filament produces a flow symmetric about both its axis and its midpoint. This flow can only produce an extensile motion of the filament. (b) The straight conformation, however, is linearly unstable to transverse perturbations. Here the perturbation is symmetric about an axis passing through the filament midpoint. This leads to a flow which is no longer symmetric about the initial axis. It enhances the instability and can produce center of mass motion of the filament. Bead positions are shown as yellow filled circles, while the fluid velocity is shown as streamlines with the background color indicating its magnitude.

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measures the ratio of time scales associated with active and elastic relaxation. As $A < 0$ the active time scale diverges and conformational changes occur only due to elastic forces. As $A > 1$ conformational changes due to activity are much more rapid than those due to elasticity.

Method.—We use the lattice Boltzmann method [12] to obtain the incident flow in Eq. (3), corresponding to terms with $m / C_{222} n$, and then integrate the equations using the forward Euler method. The method of obtaining the incident flow is described in detail in Refs. [13] and [14], and is related to methods used in Ref. [15]. The subtleties in using the lattice Boltzmann method to solve Eq. (3) are described in detail in Ref. [14]. We use lattice units in which both spatial and temporal discretization scales are unity. We choose $b_0 = 2$ such that there is less than 1% variation in contour length. We choose $\bar{\kappa}$ in the range 0.0 to 0.5, $\sigma_0$ in the range 0 to 0.05, and $N$ in the range 16 to 96. The initial filament conformation is a superposition of small amplitude transverse sinusoidal deformations of wavelengths a few integer multiples of $L$. The integration is carried out for several million time steps. Our results, unless otherwise stated, are for periodic planar lattices of size 128.

Results.—We summarize our results in Figs. (1 and 2) and the movies in Ref. [14]. Our simulations find a linear instability of an initially straight filament. On dimensional grounds, this instability occurs only when $L > l_3 \sim \kappa / \sigma_0$, where numerical prefactors can be obtained from a linear stability analysis of Eq. (3). In contrast, the elastic Euler instability of a filament under force $F$ occurs when $L > l_E \sim \sqrt{\kappa / F}$. A linear instability of passive filaments in an active medium without nonlocal hydrodynamics was reported in Ref. [16], while bow-shaped conformation for filaments driven by external forces were reported in Ref. [17].

FIG. 2 (color). Filament motion for high activity. (a) For an even initial conformation with $A = 250.7$, the motion is predominantly translational. (b) For an identical initial conformation with $A = 150.4$, the motion is translational as well as rotational due to the spontaneous appearance of conformations with odd symmetry. In both (a) and (b) the color of the trace corresponds to the center of mass speed $|R|$ normalized by its maximum value. (c) and (d) Time traces of the $x$ (top) and $y$ components (bottom) of $R$ (olive dashed line) and $K = -(\sigma_0 / 4 \pi \eta b_0) \langle \bar{\kappa} \rangle$ (black solid line), where $\langle \bar{\kappa} \rangle$ is the average curvature, corresponding to simulations in (a) and (b), respectively. The velocity and the curvature remain strongly correlated for various filament conformational modes and activity numbers. Times are in $10^3$ LB steps.
We show the nature of the linear instability, as $A \to \infty$ and $k \to 0$, in Fig. (1) and its accompanying movie [14]. The flow produced by a straight filament is symmetric about the filament center and the filament axis, as shown in Fig. 1(a). It can only produce an extensile motion in the filament which is transient, since it is balanced by the stretching elasticity. A spontaneous transverse perturbation breaks the flow symmetry about the initial axis, enhancing the perturbation, as shown in Fig. 1(b). The flow now develops an uncompensated directional component in which the filament can translate. Since the hydrodynamic drag on the filament is greater at its ends [18], a balance between elastic deformation, active propulsion, and drag ensues and the filament propels steadily in a deformed bow-shaped conformation [14].

In Fig. (1) the initial perturbation is symmetric about the filament midpoint. We call this an even mode. Initial perturbations which are antisymmetric about the midpoint are also linearly unstable. However, these odd modes produce flows of a completely different nature than the even modes. Instead of an uncompensated linear component, the flow develops a vorticity centered on the filament midpoint which results in pure rotational motion [14]. A generic initial perturbation is a superposition of both even and odd modes and, thus, both rotates and translates. At low $A$, there is little coupling between the conformational modes, due to which the filament has steady motion. However, with increasing $A$ greater elastic deformations are needed to balance the activity, due to which the filament develops nonlinear fluctuating states with large-amplitude deformations, as shown in Fig. (2) and the accompanying movies [14]. Conformational modes are now coupled, and modes absent from the initial perturbation can spontaneously appear. With a predominance of even modes, the motion is primarily translational as seen in Fig. 2(a), but when a spontaneously generated odd mode appears the motion is both translational and rotational as seen in Fig. 2(b). In cubic flows, we see similar linear instabilities and nonlinear fluctuating states [14]. It is surprising that such complex animate behavior can be generated by Eq. (3). Remarkably, its qualitative aspects can be understood from basic symmetry arguments relating conformations to the flows they produce. The role of symmetry in the motility of active droplets has been studied recently in Ref. [19].

In the free-draining approximation, it is possible to relate the center of mass velocity $\mathbf{R}$ to the mean curvature of the filament [14],

$$\mathbf{R} = -\frac{\sigma_0}{4\pi\eta b_0} \langle \chi \mathbf{n} \rangle,$$

where $\chi$ is the local curvature and $\mathbf{n}$ is the local unit normal vector. In Figs. 2(c) and 2(d) we plot components of this equation corresponding to simulations in Figs. 2(a) and 2(b), respectively. The agreement is good in both cases. Thus local hydrodynamics provides a good approximation for the translational velocity but nonlocal hydrodynamics is required to fully explain the conformational fluctuations. The interplay between nonlocal hydrodynamics and semiflexibility is necessary for the rotational motion of the filament, as has been noted earlier in a different context [17].

For a microtubule of size $L \sim 30 \mu m$, $\kappa \sim 50 \text{ pN} \cdot \text{m}^2$ with about 200 motors per micron exerting approximately 6 pN of force, we obtain $A \sim 60$. These parameters provide a translational speed of $\sim 1 \text{ mm s}^{-1}$ for a semicircular shape in water. The activity can be manipulated in motor-polymer bundle systems or in polymers of autonomously motile nanorods over a large range of $A$. These systems would be the best candidates for a verification of our results.

Discussion and conclusion.—Our model has several important variations. We argued that active processes cannot add linear or angular momentum to the fluid and, so, must be represented by Stokesian singularities with those properties. This ruled out the Stokeslet and the rotlet but allowed for higher singularities, of which the stresslet, being the most dominant, was retained. The stresslet, with a $C_0$ axis, is not forbidden by symmetry as a representation of a polar active element. If it is forbidden for non-symmetry reasons, we must use the potential dipole $d$ [10], the leading singularity with polar symmetry, whose velocity field is $v_\alpha(r) = G_{\alpha\beta}(r) d_\beta$, $G_{\alpha\beta} = (-\delta_{\alpha\beta} + 3\hat{r}_\alpha \hat{r}_\beta) / 8\pi\eta r^4$, in Eq. (3). The axis of the stresslet or the potential dipole can be oriented normally or obliquely to the local tangent of the filament and the stresslet can also be contractile, $\sigma_0 < 0$. The precise nature of the nonlinear steady states obtained from these various combination will be reported in future work. A generic equation of motion encompassing these specific cases is provided in Ref. [14].

Semiflexibility is crucially important in obtaining the results above. A rigid rod ($\kappa = \infty$, $A = 0$) is immune to the active instability. Since the uniaxial axes of the stresslets and the rod are aligned, it cannot, by symmetry, acquire any translational or rotational motion. It is only by the breaking of this symmetry, possible when $A \neq 0$, that the filament is able to acquire motion.

We have presented a model for an autonomously motile semiflexible filament which takes into account nonlocal hydrodynamic interactions. Our model opens up the possibility of studying the nonequilibrium dynamics of active filaments, for example the cilialike beating of motor-polymer bundles [7], as well as the collective properties of networks of active filaments, such as the cytoskeleton.

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