Comparison of explicit and mean-field models of cytoskeletal filaments with crosslinking motors

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Abstract

In cells, cytoskeletal filament networks are responsible for cell movement, growth, and division. Filaments in the cytoskeleton are driven and organized by crosslinking molecular motors. In reconstituted cytoskeletal systems, motor activity is responsible for far-from-equilibrium phenomena such as active stress, self-organized flow, and spontaneous nematic defect generation. How microscopic interactions between motors and filaments lead to larger-scale dynamics remains incompletely understood. To build from motor–filament interactions to predict bulk behavior of cytoskeletal systems, more computationally efficient techniques for modeling motor–filament interactions are needed. Here, we derive a coarse-graining hierarchy of explicit and continuum models for crosslinking motors that bind to and walk on filament pairs. We compare the steady-state motor distribution and motor-induced filament motion for the different models and analyze their computational cost. All three models agree well in the limit of fast motor binding kinetics. Evolving a truncated moment expansion of motor density speeds the computation by \(10^3\)–\(10^6\) compared to the explicit or continuous-density simulations, suggesting an approach for more efficient simulation of large networks. These tools facilitate further study of motor–filament networks on micrometer to millimeter length scales.

1 Introduction

The cytoskeleton generates force and reorganizes to perform important cellular processes [1], including cell motility [2, 3], cytokinesis [4], and chromosome segregation in mitosis [5]. The cytoskeleton is made of polymer filaments, molecular motors, and associated proteins. The two best-studied cytoskeletal filaments are actin and microtubules [1]. It remains incompletely understood how diverse cytoskeletal structures dynamically assemble and generate force of pN to nN [1, 2].

Force generation and reorganization in the cytoskeleton depend on the activity of crosslinking motor proteins that align and slide pairs of filaments (Fig. 1). Reorganization of actin networks by myosin motors is important for muscle contraction [6–8], cell crawling and shape change [9–11], and cytokinesis [4, 12]. Microtubule sliding by crosslinking kinesins and dyneins contributes to mitotic spindle assembly [5, 13–16], chromosome segregation [17–20], cytoplasmic stirring in Drosophila oocytes [21], and beating of cilia and flagella [22–24].

Filament–motor interactions produce diverse cellular structures and dynamics, but linking molecular properties of motors to larger-scale assembly behavior remains challenging. Crosslinking motors vary in binding affinity, speed, processivity, and force–velocity relation. These same ingredients can be reconstituted and show dynamic self-organization into asters or contractile bundles [25–27], active liquid crystals [28–31], or other structures [32–34]. Even in reconstituted systems, our ability to predict and control dynamics and self-organization is limited.

Improved theory and simulation of cytoskeletal assemblies with crosslinking motors would allow better prediction of both cellular and reconstituted systems. Currently, few mesoscale modeling methods for filament–motor systems are available between explicit particle simulations and continuum hydrodynamic theory. Explicit motor simulations have several existing software tools, including Cytosim [35], MEDYAN [36], and AFINES [37], and others [38]. Explicit motor simulations are straightforward to extend to include, for example, a new force–velocity relation or motor cooperativity. However, the cost of explicit particle simulations scales linearly or quadratically with the number of particles (depending on the type of interactions),
making simulation of large systems challenging. Continuum models of coarse-grained fields can be computationally tractable and predict macroscopic behavior [39–46]. Current continuum models invoke symmetry considerations to determine the structure of the model without reference to an underlying microscopic mechanisms [39,47–49], or simplify a microscopic model by making assumptions about the physics of motors [43,50–57]. Furthermore, previous continuum theories have coarse-grained the filament distribution, with simplifying assumptions about the motor distribution. This presents an opportunity to better understand how the distribution of motors evolves and affects filament motion. Further development of mesoscale modeling techniques focusing on crosslinking motors could help bridge the gap between detailed explicit particle models and continuum theories.

To develop mesoscale modeling tools, we focus on the fundamental unit of a crosslinked filament network: two filaments with crosslinking motors that translate and rotate the filaments. We study three different model representations in a coarse-graining hierarchy and compare computational cost and accuracy. For explicit motors, we extend previous work that uses Brownian dynamics and kinetic Monte Carlo simulation to handle filament motion and binding kinetics [43,56,58–62]. At the first level of coarse-graining, we average over discrete bound motors to compute the continuum mean-field motor density (MFMD) between filaments, and evolve this density according to a first-order Fokker–Planck equation [58]. This requires computing the solution to a single partial differential equation (PDE) for each filament pair, rather than separately tracking each individual motor. The MFMD determines the force and torque on each filament needed to evolve its position and orientation. At the second level of coarse-graining, we expand the MFMD in moments to derive a system of ordinary differential equations (ODEs) for the time evolution of the moments. While the moment expansion does not close, an approximate treatment of filament motion can be modeled by low-order moments. To compare these three model implementations, we consider test cases of parallel, antiparallel, and perpendicular filaments. Under the same initial conditions, the three model implementations give similar results on average. Remarkably, the reduced moment expansion achieves a computational cost that is \(10^3\)–\(10^6\) lower than the other models, suggesting a route to computationally tractable large-scale simulations.

2 Model overview

We consider a pair of rigid, inextensible filaments that move and reorient under the force and torque applied by crosslinking motors. Filaments move in three dimensions, experience viscous drag, and are constrained to prevent overlaps. Motors bind to and unbind from the filaments consistent with detailed balance in binding. Crosslinking motors walk with a force-dependent velocity toward filament plus ends and unbind when they reach the ends. We investigate models at three levels: an explicit motor model where motors are represented with a discrete density, a continuum mean-field motor density (MFMD) model, and a moment expansion model.

2.1 Filaments

We model filament motion using Brownian dynamics, balancing the force applied by motors against viscous drag and constraint forces. Because the force that induces Brownian motion is typically smaller than that due to motors, we neglect Brownian noise [65]. Filaments translate according to the force–balance equation

\[
\dot{\mathbf{r}}_i = \mathbf{M}_i \left( \sum_n \mathbf{F}_{n,i} \right),
\]

where \(\mathbf{r}_i\) is the center of filament \(i\) with mobility matrix \(\mathbf{M}_i\) acting on by forces \(\mathbf{F}_{n,i}\). The mobility matrix for a perfectly rigid rod in a viscous medium is

\[
\mathbf{M}_i = \left( \left( \gamma_{\parallel,i} - \gamma_{\perp,i} \right) \hat{u}_i \hat{u}_i + \gamma_{\perp,i} \mathbf{I} \right)^{-1},
\]

where \(\mathbf{I}\) is the identity matrix and \(\gamma_{\parallel,i}\) and \(\gamma_{\perp,i}\) are the parallel and perpendicular drag coefficients with respect to the filament orientation \(\hat{u}_i\). Cytoskeletal filaments with length \(L_i\) and diameter \(D_{fil}\) typically have a large aspect ratio \(L_i/D_{fil} \gg 1\), so we approximate the drag coefficients using slender body theory [66].

The torque–balance equation is

\[
\dot{\hat{u}}_i = \frac{1}{\gamma_{\theta,i}} \left( \sum_n \mathbf{T}_{n,i} \right) \times \hat{u}_i,
\]

where \(\mathbf{T}_{n,i}\) are the torques acting on filament \(i\) and \(\gamma_{\theta,i}\) is the rotational drag coefficient about the center of filament \(i\).

The force and torque exerted by crosslinking motors depend on where motors are attached, the motor tether extension, and the relative position and orientation of filaments. Given the crosslinking motor distribution along the filaments \(\psi_{i,j}(s_i, s_j)\), where \(s_i\) is the bound motor head position on filament \(i\), the total crosslinking force and torque exerted by filament \(i\) on filament \(j\) are

\[
\mathbf{F}_{i,j} = \int_{L_i} \int_{L_j} \mathbf{f}_{i,j}(s_i, s_j) \psi_{i,j}(s_i, s_j) ds_i ds_j,
\]

\[
\mathbf{T}_{i,j} = \int_{L_i} \int_{L_j} s_j \hat{u}_j \times \mathbf{f}_{i,j}(s_i, s_j) \psi_{i,j}(s_i, s_j) ds_i ds_j.
\]

where \(\mathbf{f}_{i,j}(s_i, s_j)\) is the force exerted on filament \(j\) by the crosslinking motor attached at \(s_i\) and \(s_j\) (Fig. 1E).
Fig. 1  Experimental systems of cytoskeletal filaments with crosslinking motors and overview of the model. a–c Fluorescence microscopy images of cytoskeletal networks. a Mitotic spindle showing microtubules (green), chromosomes (blue), and spindle-pole component TPX2 (red) [63]. b Reconstituted active gel of microtubules (white) driven by crosslinking kinesin motor clusters with local flow field shown (yellow arrows) [28]. Scale bar: 80 μm. c Reconstituted active network of actin (magenta) and myosin II (green) [64]. Scale bar: 50 μm. d Schematic of filament–motor network with green filaments and red motors. e Schematic of filament pair (green) crosslinked by a motor (red) with model variables position of filament $i$’s center $r_i$, orientation vector of filament $i$ $\hat{u}_i$, vector between filament centers $r_{i,j} = r_j - r_i$, vector between motor heads $h_{i,j}$, motor tether extension $|h_{i,j}|$, and motor speed on filament $i$ while attached to filament $j$.

For brevity, we use subscripts on variables such as $f_{i,j}$ to indicate that these are functions of the relative position and orientation of filaments $i$ and $j$. Our three model implementations all models use Eqs. (4) and (5) to compute the force and torque that evolve filament position and orientation but differ in how the computation of $\psi_{i,j}$.

We constrain the motion of filaments to prevent overlap, which avoids numerical instabilities introduced by a hard potential between filaments. To implement the constraint, we construct a vector $\hat{u}_{\text{min}}$ that is perpendicular to both infinite carrier lines defined by $\hat{u}_i$ and $\hat{u}_j$ and parallel to the vector of closest approach between these lines. The vector $\hat{u}_{\text{min}}$ is used to define two normal planes that confine the filaments, leading to the modified force and torque

$$\tilde{F}_{i,j} = F_{i,j} - (F_{i,j} \cdot \hat{u}_{\text{min}})\hat{u}_{\text{min}}$$

$$\tilde{T}_{i,j} = (T_{i,j} \cdot \hat{u}_{\text{min}})\hat{u}_{\text{min}}.$$  

Note that for filaments lying in the same confining plane and $|\hat{u}_i \cdot \hat{u}_j| < 1$, $\hat{u}_{\text{min}} = 0$ and our constraints break down. However, if only the first condition is satisfied, i.e., (anti)parallel filaments, $\tilde{T}_{i,j} = 0$ and $\tilde{F}_{i,j}$ is parallel to $\hat{u}_i$ and $\hat{u}_j$. After computing the force and torque, we numerically integrate Eqs. (1) and (3) to update filament position and orientation.

2.2 Motors

In our model motors bind and unbind, crosslink between two filaments, exert force and torque when crosslinking, and walk with a force-dependent velocity. Typically motor proteins diffuse in solution until they are near a filament, then stochastically bind to that filament. Once one head binds, the other head can bind to a second filament, forming a crosslink, or the motor can unbind. Crosslinking motors can unbind to a state with one head bound, or can unbind completely from both filaments. We consider an infinite reservoir of unbound motor proteins. The diffusion of motors in solution is fast relative to the motion of filaments, so we assume the motor reservoir has uniform, constant concentration. We neglect steric interactions between motors. This approximation holds for filaments sparsely pop-
ulated with motors and motors that do not cluster on filaments or in solution.

Motors crosslinking filaments have a potential energy $U_{i,j}(s_i, s_j)$ (Fig. 1). The energy depends on the motor head separation vector $\mathbf{h}_{i,j}(s_i, s_j) = \mathbf{r}_j + s_j \hat{u}_j - (\mathbf{r}_i + s_i \hat{u}_i)$ that gives the motor tether extension

$$k_{i,j}(s_i, s_j) = \sqrt{r_{i,j}^2 + s_i^2 + s_j^2 + 2r_{i,j} \cdot (s_j \hat{u}_j - s_i \hat{u}_i) - 2s_is_j(\hat{u}_i \cdot \hat{u}_j)},$$

where $\mathbf{r}_{i,j} = \mathbf{r}_j - \mathbf{r}_i$, and $r_{i,j}^2 = \mathbf{r}_{i,j} \cdot \mathbf{r}_{i,j}$ (Fig. 1c).

The bound motor heads walk with a speed $v_{i,j}$ that depends on the force component on the motor head parallel to the walking direction, $\hat{u}_i \cdot f_{j,i}$ [67]. This projected force is used to determine the motor speed via the force-velocity relation, as discussed below. This model is based on processive microtubule motors such as kinesin and dynein, but a similar model has been used for myosin minifilaments [36,37].

### 3 Explicit motor model

In the explicit motor model individual bound motors are modeled, allowing fluctuations in bound motor number and binding kinetics that recover the correct equilibrium distribution of crosslinking proteins in the limit of no motor walking (Fig. 2a) [43,56,59–62].

#### 3.1 Binding kinetics and stepping

A motor diffuses in solution until one of its heads bind to a filament; we model this by an infinite reservoir of unbound motors with a uniform and constant concentration $c_o$. Filaments have a linear binding site density $c$, and the binding site has an association constant $K_a$ (units of $\mu M^{-1}$). First motor head binding has rate

$$k_{on,S} = K_a c_o \mu L_{tot} k_{o,S},$$

where $L_{tot} = \sum_i L_i$ is the total length of filaments and $k_{o,S}$ is the bare (force-independent) unbinding rate for singly bound heads. All binding locations have equal binding probability. Singly bound motors unbind at rate $k_{off,S} = k_{o,S}$.

A motor with one head bound crosslinks to another filament, which may stretch or compress its tether. This makes crosslinking kinetics force dependent; our models satisfy detailed balance in binding, so we recover the thermal equilibrium Boltzmann distribution in the limit of passive crosslinkers. Motor motion shifts the crosslinking distribution away from equilibrium. Motor unbinding rate can depend on the force applied to bound heads [68–73]. Previous work shows how this force dependence can be included while maintaining detailed balance in binding [62,74,75]. For simplicity, here we include the force dependence in the binding rate only and discuss possible implications below. With one head bound to filament $i$ at position $s_i$, the free motor head binds to filament $j$ at position $s_j$ with a probability proportional to a Boltzmann factor of binding energy

$$P_{S \rightarrow C} \propto \exp(-\beta U_{i,j})$$

with $\beta = (k_B T)^{-1}$ (Fig. 3a). Here, $S \rightarrow C$ denotes the motor’s transition from a single head bound (S) to crosslinking (C). The total binding rate is computed by integration over all binding positions on filament $j$

$$k_{on,C} = \frac{\epsilon K_p k_{o,C}}{V_{bind}} \int_{L_j} e^{-\beta U_{i,j}} ds_j,$$

where $k_{o,C}$ is the bare (force-independent) unbinding rate for a crosslinking motor, $K_p$ is the crosslinking association constant. The unbound motor head explores a volume $V_{bind}$ centered about the bound head, computed as the integral of the unbound head’s position weighted by the Boltzmann factor

$$V_{bind} = \int e^{-\beta U_{i,j}} dr^3 = 4\pi \int_{0}^{R_{cut,C}} e^{-\beta U_{i,j}} r^2 dr.$$

Beyond the cutoff radius $R_{cut,C}$, the integrand becomes small, enabling the use of a lookup table (Appendix B). The probability distribution of binding position depends on the Boltzmann factor. We recover the proper binding distribution through inverse transformation sampling of Eq. (11) (Appendix B.2).

As discussed above, the unbinding rate of a single head of a motor crosslinking two filaments is assumed to be force-independent,

$$k_{off,C} = k_{o,C}.$$

Force-dependent unbinding affects the density of motor proteins most when stretched [70]; larger motor stretch occurs when external force is applied against the force generated by motors. Therefore, sliding filaments slowed only by drag, like those in active nematics, will be less affected by force-dependent unbinding than stationary filaments or jammed filaments like microtubules in mitotic spindles. We can include force-dependent unbinding in the explicit motor and MFMD model but not in the moment expansion model (Sect. 5). We chose the time step small enough that individual motors undergo only one transition per time step (Appendix A).
Fig. 2  Comparison of motor representations in three hierarchical models with schematics on the left and 2D motor distributions on the right. 

**A** Explicit motor model with t

1 head bound

Unbound

Crosslinking

 filament $j$

filament $i$

$\psi_{i,j}(s_i, s_j) = \left\langle \sum_n \delta(s_i - s_n) \delta(s_j - s'_n) \right\rangle$

$\mu_{i,j}^{k,l} = \int_{L_i} \int_{L_j} s_i^k s_j^l \psi_{i,j} ds_i ds_j$

$\nu_{i,j} = \nu_0 \left( 1 + \frac{\hat{u}_i \cdot f_{j,i}}{f_{\text{stall}}} \right)$, $0 < \hat{u}_i \cdot f_{j,i}$

$\nu_0 \left( 1 + \frac{\hat{u}_i \cdot f_{j,i}}{f_{\text{stall}}} \right)$, $-f_{\text{stall}} < \hat{u}_i \cdot f_{j,i} < 0$

$\nu_0 \left( 1 + \frac{\hat{u}_i \cdot f_{j,i}}{f_{\text{stall}}} \right)$, $\hat{u}_i \cdot f_{j,i} < -f_{\text{stall}},$

where $f_{\text{stall}}$ is the motor stall force (Fig. 3b).

3.2 Distribution of explicitly modeled motors

The bound motor distribution is

$\psi_{i,j}(s_i, s_j, t) = \sum_{n=1}^{N_{i,j}(t)} \delta(s_i - s_n(t)) \delta(s_j - s'_n(t))$,  

(15)
Choice for motor tether potential, force–velocity relation, and filament initial configurations. 

- **a** Plot of the normalized potential energy in motor tether as a function of motor extension (blue) and equivalent zero-length tether potential (orange line). Both potentials have identical slope at the distance $f_{\text{stall}}/k_{\text{cl}}$ (red dashed line) where motors stall.

- **b** Plot of normalized motor speed as a function of force (blue) and its linear approximation (dashed orange).

- **c** Chosen initial configurations of pairs of 1μm filaments. Filament centers are separated by $D_{\text{fil}} = 25$ nm perpendicular to both filament orientation vectors.

where $\delta(s_i)$ is the Dirac delta function and $N_{i,j}$ is the total number of motors crosslinking filaments $i$ and $j$. Here, $s_i$ and $s'_n$ are the attached positions of the heads of the $n$th crosslinking motor. Motors with one head bound to filament $i$ have a distribution

$$
\chi_i(s_i, t) = \sum_{n=1}^{N_i(t)} \delta(s_i - s_n(t)),
$$

where $N_i$ is the number of one head bound motors on filament $i$. Only motors crosslinking exert forces between filament pairs, but $\chi_i$ and $\chi_j$ are needed to calculate the evolution of $\psi_{i,j}$.

### 4 Mean-field motor density model

Under typical experimental conditions, there can be tens to thousands of crosslinking motors between a filament pair. Motor force and torque fluctuations occur because of stochastic motor binding and unbinding. As the number of motors increases, the standard deviation relative to the mean decreases as $1/\sqrt{N}$. For our explicit motor model, antiparallel filaments with an average of 14 motors bound show a standard deviation in bound motor number of 27% of the mean. This shows that the fluctuations are quite significant for order 10 motors. The $1/\sqrt{N}$ scaling predicts that for an average of 1000 motors, the standard deviation would be only 3.2% of the mean. The force and torque scale similarly. Therefore, for large motor number, we may use the average motor distribution to derive a mean-field motor density (MFMD) to accurately describe force and torque on filaments by motors. We can then evolve the MFMD instead of explicit motors (Fig. 2b). We previously showed that the average steady-state density of crosslinking motors between stationary parallel filaments agreed well with a solution to a multi-dimensional Fokker–Planck equation (FPE) [58]. Here, we expand this approach to model crosslinking motor density between filaments in three dimensions, allow filament motion, and study time-dependent behavior of coupled systems of motors and filaments.
For a one-step binding model, the MFMD evolves according to
\[
\frac{\partial \psi_{i,j}(s_i,s_j,t)}{\partial t} = -\frac{\partial (v_{i,j}\psi_{i,j})}{\partial s_i} - \frac{\partial (v_{i,j}\psi_{i,j})}{\partial s_j} + k_{on} - k_{off}\psi_{i,j},
\]
with motor velocity \(v_{i,j}\), motor crosslinking rate \(k_{on}\), and unbinding rate \(k_{off}\). To satisfy detailed balance in binding, we use the rates \(k_{on} = 2k_0ce^{-\beta U_{i,j}(s_i,s_j)}\) and \(k_{off} = 2k_0\), with the effective concentration \(c\) (units \(\text{nm}^{-2}\)) [58]. The factors of two occur because there are two ways a motor can crosslink. To numerically solve the hyperbolic Eq. (17), we use a first-order accurate upwind difference method (Appendix C).

The mean-field motor density model differs from the explicit model in that motors with one head bound are not modeled explicitly. To properly compare the different binding models, we establish a mapping of parameters between these two models (Appendix D), which gives
\[
c = \frac{c^2 K_a K_E}{V_{\text{bind}}} c_o.
\]

Some model parameters are difficult to measure directly. For example, the association constant \(K_E\) may differ from \(K_a\) if proteins change their molecular conformation when bound. We discuss an approach to estimate such parameters in Appendix E.

### 4.1 Steady-state solution for MFMD on antiparallel filaments

If filaments move slowly compared to the timescale of motor rearrangement, then a quasi-steady-state approximation can be used. In the quasi-steady limit, the force and torque on filaments are computed from the steady-state MFMD [61]. The quasi-steady approximation is computationally efficient compared to numerical integration of the time-dependent PDE. A steady-state solution also provides a convenient route to compare our model implementations.

At steady state, Eq. (17) becomes
\[
\psi_{i,j}\frac{\partial v_{i,j}}{\partial s_i} + v_{i,j}\frac{\partial \psi_{i,j}}{\partial s_i} + \psi_{i,j}\frac{\partial v_{j,i}}{\partial s_j} + v_{j,i}\frac{\partial \psi_{i,j}}{\partial s_j} + 2k_0\psi_{i,j} = 2k_0ce^{-\beta U(s_i,s_j)}.
\]

Here, we choose functional forms of \(U_{i,j}\) and \(v_{i,j}\) consistent with previous models [36,37,58,60,61]. Motors have a potential energy \(U_{i,j} = \frac{k_t}{2} (h_{i,j} - h_{cl})^2\) determined by the tether spring constant \(k_t\) and tether length \(h_{cl}\) (Fig. 3a), which implies a motor crosslinking filaments \(i\) and \(j\) exerts a force on \(f_{i,j} = -k_{cl}(1 - \frac{h_{i,j}}{h_{cl}}) h_{i,j}\) on filament \(j\). The force–velocity relation of a motor head attached to filament \(i\) while the other head is bound to \(j\) follows Eq. (14). Here, we assume that motors that reach filament ends walk off, i.e., no end pausing.

A semi-analytic steady-state solution can be derived for antiparallel filaments when motor tethers have zero length \(h_{cl} = 0\) because the FPE is symmetric under the transformation \(i \leftrightarrow j\). For zero-tether-length motors to mimic their non-zero-length counterparts, we modify the zero-length motor’s spring constant so both types of motors stall at the same extension \(h_{i,j} = h_{\text{stall}}\). This implies \(k'_{cl}h_{\text{stall}} = k_{cl}(h_{\text{stall}} - h_{cl}) = f_{\text{stall}}\) with the solution
\[
k'_{cl} = \frac{k_{cl}f_{\text{stall}}}{f_{\text{stall}} + k_{cl}h_{cl}},
\]
where \(h_{\text{stall}} = f_{\text{stall}}/k'_{cl}\). Note this choice changes the binding dynamics, because the potential energy is now larger for larger motor extension (Fig. 3A).

To find the steady-state solution, note that \(\hat{u}_i, r_{i,j} : \hat{u}_j = 0\) and \(\hat{u}_i \cdot \hat{u}_j = -1\) for antiparallel filaments with centers aligned. Therefore, \(h_{i,j} = \sqrt{r_{i,j}^2 + (s_i + s_j)^2}\) and \(\hat{u}_i \cdot f_{i,j} = \hat{u}_j \cdot f_{j,i} = -k'_{cl}(s_i + s_j)\).

Since \(U_{i,j}, v_{i,j}\), and \(v_{j,i}\) depend exclusively on the sum of \(s_i\) and \(s_j\), we make the change of variables \(\xi = s_i + s_j\) in Eq. (19) to find
\[
(v_{i,j} + v_{j,i})\frac{\partial \psi_{i,j}}{\partial \xi} + (\frac{\partial v_{i,j}}{\partial \xi} + v_{j,i} + v_{i,j} + 2k_0) \psi_{i,j} = 2k_0ce^{-\beta U(\xi)}.
\]

There are three regions of solution determined by the force–velocity relation Eq. (14): \(\xi \leq 0\), \(0 \leq \xi \leq h_{\text{stall}}\), and \(h_{\text{stall}} < \xi\). For \(\xi \leq 0\), \(v_{i,j} = v_{j,i} = v_0\) and Eq. (21) becomes
\[
\frac{\partial \psi_{i,j}}{\partial \xi} + \frac{\psi_{i,j}}{l_o} = \frac{c}{l_o} e^{-\beta U(\xi)},
\]
where \(l_o = v_o/k_o\) is the motor run length. This is solved with an integrating factor, giving
\[
\psi_{i,j}(\xi) = e^{-\xi/l_o} \psi_{i,j}(-L) + \frac{c}{l_o} e^{-\xi/l_o} \int_{-L}^{\xi} e^{\xi/l_o} e^{-\beta U(\xi)} dx.
\]

Applying the boundary condition \(\psi_{i,j}(-\frac{L}{2}, -\frac{L}{2}) = \psi_{i,j}(-L) = 0\), we remove the last term in Eq. (23) and rewrite the Gaussian integral as
\[
\psi_{i,j}(\xi) = \frac{c}{l_o} \sqrt{\frac{\pi}{2|k'_{cl}|}} \exp \left( \frac{1}{2|k'_{cl}|} - \frac{\xi}{l_o} \right) \left[ \text{erf} \left( \frac{\beta k'_{cl} f_{\text{stall}} - 1}{l_o \sqrt{2|k'_{cl}|}} \right) \right]_{\xi = -L}^{\xi = L}.
\]
For $0 \leq \xi \leq h_{\text{stall}}$, the velocity $v_{i,j} = v_{j,i} = 1 - \frac{\xi}{h_{\text{stall}}}$. Equation (21) becomes

$$
(h_{\text{stall}} - \xi) \frac{\partial \psi_{i,j}}{\partial \xi} + \left( \frac{h_{\text{stall}}}{l_o} - 1 \right) \psi_{i,j} = \frac{h_{\text{stall}}}{l_o} ce^{-\beta U(\xi)}. \tag{25}
$$

Solving with an integrating factor, we find

$$
\psi_{i,j}(\xi) = \psi_{i,j}(0) \left( \frac{h_{\text{stall}}}{h_{\text{stall}} + \xi} \right)^1 - \frac{h_{\text{stall}}}{l_o} ce^{-\beta U(\xi)} + \frac{l_o}{l_o (h_{\text{stall}} - \xi)^{1 - \frac{h_{\text{stall}}}{l_o}}} \int_0^\xi (h_{\text{stall}} - x) - \frac{h_{\text{stall}}}{l_o} ce^{-\frac{1}{2} (r_{i,j}^2 + x^2)} dx.
$$

We match the solution for $\psi_{i,j}(0)$ to Eq. (23) to enforce continuity. The exponential term in Eq. (26) can be approximated by a series expansion or integrated numerically. Here, we use numerical integration.

For $\xi > h_{\text{stall}}$, the velocity and velocity derivatives are zero, so

$$
\psi_{i,j}(\xi) = ce^{-\frac{1}{2} (r_{i,j}^2 + \xi^2)}. \tag{27}
$$

Since the motor velocity is zero at $\xi = h_{\text{stall}}$, motors do not walk from $\xi < h_{\text{stall}}$ to $\xi > h_{\text{stall}}$. A nonzero MFMD exists for $\xi > h_{\text{stall}}$ only if motors bind at these lengths. This appears as an integrable discontinuity at $\xi = h_{\text{stall}}$.

### 5 MFMD moment expansion

A series expansion or reduced representation of a continuous distribution can lower the computational cost of solving a system’s time evolution [57, 76, 77]. Here, we use low-order moments of the MFMD to calculate motor number, mean and standard deviations of motor head distribution, and filament motion.

The moments of $\psi_{i,j}$ are

$$
\mu_{i,j}^{k,l}(t) = \int_{L_i} \int_{L_j} s_i^k s_j^l \psi_{i,j} ds_i ds_j, \tag{28}
$$

where $k, l$ are nonnegative integers. The moments are symmetric under exchange of both filaments and powers so that $\mu_{i,j}^{k,l} = \mu_{j,i}^{l,k}$. The zeroth moment $\mu_{i,j}^{0,0} = N_{i,j}$ is the total number of motors bound to the two filaments, and the first moments $\mu_{i,j}^{1,0}, \mu_{i,j}^{0,1}$ are proportional to the mean motor head position along each filament $P_i = \frac{\mu_{i,j}^{1,0}}{N_{i,j}}, P_j = \frac{\mu_{i,j}^{0,1}}{N_{i,j}}$. The first two second moments determine the standard deviation of motor head density

$$
\sigma_i = \sqrt{\mu_{i,j}^{2,0} N_{i,j} - \mu_{i,j}^{1,0} \mu_{i,j}^{1,0}}. \tag{29}
$$

The symmetric second moment term $\mu_{i,j}^{1,1}$ determines the covariance of motor head position

$$
V_{i,j} = \frac{\mu_{i,j}^{1,1}}{N_{i,j}} - P_i P_j. \tag{30}
$$

The positional means, standard deviations, and covariance are used to reconstruct an approximate MFMD for visualization using a bivariate normal distribution (Fig. 2c, Videos 1–6).

Using the approximation of zero-length tethers as in Sect. (4.1) above, $f_{i,j}$ is a linear function of $s_i$ and $s_j$. In this case, filament motion can be computed from low-order moments using Eqs. (4) and (5):

$$
F_{i,j} = -k c_l \int_{L_i} \int_{L_j} (r_{i,j} + s_j \ddot{u}_j - s_i \ddot{u}_i) \psi_{i,j} ds_i ds_j
$$

$$
= -k c_l \left( \mu_{i,j}^{0,0} r_{i,j} + \mu_{i,j}^{0,1} \ddot{u}_j - \mu_{i,j}^{1,0} \ddot{u}_i \right) \tag{31}
$$

and

$$
T_{i,j} = -k c_l \int_{L_i} \int_{L_j} s_j \ddot{u}_j \times (r_{i,j} + s_j \ddot{u}_j - s_i \ddot{u}_i) \psi_{i,j} ds_i ds_j
$$

$$
= -k c_l \ddot{u}_j \times \left( \mu_{i,j}^{0,1} r_{i,j} - \mu_{i,j}^{1,1} \ddot{u}_i \right) \tag{32}
$$

Substituting Eqs. (31) and (32) into Eq. (1) and (3) shows that only moments up to second order are needed to compute filament motion from crosslinking motors. Thus, motor and filament evolution can be written as a system of ODEs that depend on the dynamical evolution of the moments. This dynamical evolution is computed by taking the time derivative of Eq. (28) and substituting in the FPE (17)

$$
\frac{\partial \mu_{i,j}^{k,l}}{\partial t} = \int_{L_i} \int_{L_j} s_i^k s_j^l \frac{\partial \psi_{i,j}}{\partial t} ds_i ds_j. \tag{33}
$$

However, this coupled system of equations for the moment time evolution does not close. Because the piecewise motor force–velocity relation is not linear, moments depend on higher-order moments recursively. Also, filament ends introduce boundary terms that prevent closure. Despite this, in certain limits a truncated moment expansion shows good agreement with the explicit and MFMD models.
We first introduce a linear approximation to the force–velocity relation (Fig. 3b)

$$v_{i,j} \approx v_o \left(1 + \frac{\hat{u}_i \cdot f_{i,j}}{f_{\text{stall}}}\right)$$

$$= v_o \left(1 + \frac{k \kappa_{i,j}}{f_{\text{stall}}} (r_{i,j} \cdot \hat{u}_i + \hat{u}_i \cdot \hat{u}_j s_j - s_i)\right).$$

(34)

This approximation is valid for $h_{\text{stall}} \gg 1/k\alpha$, in which case motors do not bind beyond their stall stretch. We also require that $v_o \gg 2k_o \sqrt{1/k\alpha}$, ensuring that motors pulled towards the plus ends with $\hat{u}_i \cdot f_{i,j} > 0$ move quickly into a regime -- $f_{\text{stall}} < \hat{u}_i \cdot f_{i,j} < 0$, where the linear and piecewise force–velocity functions agree.

We substitute the linearized force–velocity function from Eq. (34) into the MFMD Eq. (17) to obtain

$$\frac{\partial \psi_{i,j}}{\partial t} = 2k_o c e^{-\beta\psi_{i,j}} + (2\kappa - 2k_o) \psi_{i,j}$$

$$- (v_o + \kappa (r_{i,j} \cdot \hat{u}_i + \hat{u}_i \cdot \hat{u}_j s_j - s_i)) \frac{\partial \psi_{i,j}}{\partial s_i}$$

$$- (v_o + \kappa (r_{j,i} \cdot \hat{u}_j + \hat{u}_j \cdot \hat{u}_i s_i - s_j)) \frac{\partial \psi_{i,j}}{\partial s_j},$$

(35)

where $\kappa = v_o k\alpha / f_{\text{stall}}$ is the rate at which motors reach their stall force. Integrating Eq. (35) directly returns the zeroth-moment equation

$$\frac{\partial \mu_{i,j}}{\partial t} = 2k_o q_{i,j}^{0,0} - 2k_o q_{i,j}^{0,0}$$

$$+ \left[(-v_o - \kappa r_{i,j} \cdot \hat{u}_i + \hat{u}_j s_j) B_{i,j}^1 - \kappa \hat{u}_i \cdot \hat{u}_j B_{i,j}^1\right]_{\partial L_i}$$

$$+ \left[(-v_o - \kappa r_{j,i} \cdot \hat{u}_j + \hat{u}_i s_i) B_{i,j}^0 - \kappa \hat{u}_i \cdot \hat{u}_j B_{i,j}^1\right]_{\partial L_j},$$

(36)

where we have defined $q_{i,j}^{k,l}$ are analytically integrated in one dimension and then numerically integrated using the quad method also from sciprintegrate (Appendix F).

The boundary terms in square brackets contain moments and $B_{i,j}^l$ an order higher than $\partial \mu_{i,j}/\partial t$. In Appendix G, we write the analogous time evolution for the $B_{i,j}^l$, and show that it does not close. Therefore, the moment evolution equations do not close.

To close the system of equations, we set the boundary terms to zero. Physically, this means we neglect motor unbinding from filament plus ends. If motors pause at plus ends, this approximation will lead to significant error. However, if motor unbinding is relatively rapid (including at filament plus ends), this is a good approximation. To explore the impact of not including these boundary terms, below we quantify the discrepancy between this model and the explicit motor and MFMD models. Neglecting boundary terms truncates the system of equations at second order, because only terms up to second order are needed to calculate force and torque on filaments.

We evolve equations (1, 3, 38) using solver_jvp in the scipy.integrate library [78]. This code uses the LSODA integrator, an Adams/BDF integration method that automatically detects stiffness, from the Fortran ODEPACK library [79]. The source terms $q_{i,j}^{k,l}$ are analytically integrated in one dimension and then numerically integrated using the quad method also from scipy.integrate (Appendix F).

6 Results

To test the degree of agreement between explicit motor and mean-field models, we first selected parameters based on microtubules and kinesin-5 motor proteins because they are relatively well-studied cytoskeletal proteins [81, 82, 84, 85] (Table 1). We studied three characteristic sets of initial filament pair position and orientation: antiparallel, parallel, and perpendicular (Fig. 3, Video 1–3), and compared both stationary and moving filaments. We choose an initial condition with no motors bound to filaments, in order to observe the effects of time evolution of the motor density. For stationary filaments, we found good agreement for all three models. For moving filaments, we found qualitative agreement but fluctuations in motor dynamics and different end boundary conditions contributed to quantitative differences in filament motion. We measured the computational cost for stationary antiparallel filaments.
Table 1 Model parameters for MTs and kinesin-5 for explicit motor distribution and MFMD calculations

| Parameter                        | Symbol | Value               | Notes                                      |
|----------------------------------|--------|---------------------|--------------------------------------------|
| Total time                       | $N_t$  | 20 sec              | Chosen                                     |
| Explicit motor time step size    | $\Delta t_{\text{Explicit}}$ | 0.0001 sec         | Chosen for numerical stability             |
| MFMD time step size              | $\Delta t_{\text{MFMD}}$ | 0.001 sec          | Chosen for numerical stability             |
| MFMD grid spacing                | $\Delta s$ | 1 nm               | Chosen for numerical stability             |
| Viscosity                        | $\eta$ | $10^{-6}$ pN sec nm$^{-2}$ | Chosen (viscosity of cytoplasm) |
| Filament length                  | $L$    | 1 $\mu$m            | Chosen                                     |
| Filament diameter                | $D_{\text{fil}}$ | 25 nm              | Diameter of microtubules [80]             |
| Explicit motor concentration     | $c_o$  | 11 nM               | Chosen                                     |
| MFMD effective concentration     | $c$    | 0.0093 nm$^{-2}$    | Calculated (Sect. D)                      |
| Modified tether length           | $h_{\text{cl}}$ | 0 nm               | Chosen (Sect. 2.2)                        |
| Effective tether spring constant | $k_{\text{cl}}$ | 0.037 pN nm$^{-1}$ | Calculated (Sect. 2.2), spring constant [81], tether length [82] |
| Filament binding site density    | $\epsilon$ | 0.25 nm$^{-1}$  | Estimated, one site every four nanometers |
| Inverse temperature              | $\beta = \frac{1}{k_B T}$ | 0.2433 pN$^{-1}$ nm$^{-1}$ | Room temperature                         |
| Motor speed                      | $v_o$  | 50 nm sec$^{-1}$    | [83]                                       |
| Motor stall force                | $f_{\text{stall}}$ | 2 pN              | [84]                                       |
| Association constant             | $K_a$  | 0.005 nM$^{-1}$     | [85]                                       |
| Association constant (unbound→one head bound) | $K_{\epsilon}^*$ | 2.56 | Calculated (Sect. D), [86] |
| Multi-step bare off rate         | $k_{o,S}$ | 0.77 sec$^{-1}$ | [85]                                       |
| Multi-step bare off rate (unbound→one head bound) | $k_{o,C}$ | 0.77 sec$^{-1}$ | Chosen to match $k_{o,S}$ |
| One-step bare off rate           | $k_o$  | 0.77 sec$^{-1}$     | Chosen to match $k_{o,S}$                 |

and found that the moment expansion model can give a dramatic improvement in performance.

6.1 Stationary filament pairs

When filaments are held stationary, motor density reaches or fluctuates around a steady-state solution (Fig. 4). To compare with the mean-field models, we averaged 48 realizations of each explicit motor simulation; the results agreed within error with the mean-field models (Fig. 4b–d). This agreement between models demonstrates that the mean-field models capture the average behavior of our explicit model.

Beyond the steady state, we characterize the evolution of motor number, force, and torque (Fig. 4e–m). In all configurations, the crosslinking motor number in the explicit motor model lags that of the MFMD and moment expansion models (Fig. 4e–g). The crosslinking rate in the two-step binding algorithm depends on the density of motors with one head bound, resulting in a slower approach to steady state.

For antiparallel filaments, force generation increases with crosslinking motor number (Fig. 4h, Video 1) because motors walk in opposite directions, causing the motor tether to stretch and generate force. If free to move, these antiparallel filaments would slide. No average sliding would occur for parallel filaments, and the small number of crosslinking proteins for perpendicular filaments results in small relative force (Fig. 4i, j). The average explicit motor motor torque in the $\hat{z}$-direction shows significant fluctuations about the mean (Fig. 4k–m). Because motor torque increases for motors
Fig. 4 Comparison of model results for three different stationary filament configurations. a Schematic of the three different filament configurations and legend for following plots. b Plot of total crosslink motor numbers at steady state. c Plot of steady-state motor force components from filament i on filament j. d Bar graph of steady-state torque in the \( \hat{z} \)-direction from filament i on filament j. Explicit motor model error bars in (b–d) indicate the Standard Error of the Mean (SEM) of the last 30 seconds of 40 second long simulations (\( n = 48 \)). e–g Bound motor number versus time. Purple and blue solid lines are the average of 48 individual explicit motor simulations (translucent lines) for one head bound and crosslinking motors. h–j Motor force in the \( \hat{x} \)-direction and \( \hat{y} \)-direction. Individual explicit motor runs are represented as blue for both directions. k–m Motor torque in the \( \hat{z} \)-direction. Full explicit motor model range not shown to better see average. n–p Steady-state motor probability density as a function of motor extension for semi-analytic (black), explicit motor, and MFMD models. Motor minimum extension is set by the separation of filaments at closest point of approach, 25 nm.
further from the filament centers, the torque fluctuations increase with filament length.

We compared the steady-state distribution of motor extension for both explicit motor and MFMD models (Fig. 4n–p). (Note that the moment expansion loses this information in coarse-graining.) The distribution of motors crosslinking antiparallel filaments has two peaks (Fig. 4n). The larger peak represents the most probable binding distance \( \Delta y \), and the second peak corresponds to motors near their stall extension \( h = \sqrt{\Delta y^2 + h^2_{\text{stall}}} \). The shape of the distribution results from motor kinetics, walking, and stalling. Motors on parallel filaments show a peak at \( \Delta y \) (Fig. 4o, Video 2), but no second peak because the motor heads walk in the same direction with similar speed. For motors crosslinking perpendicular filaments, the extension distribution is singly peaked and broader than for parallel filaments (Fig. 4p, Video 3). This occurs because the parallel force component on perpendicular filaments increases more gradually as the motors extend, causing a more gradual decrease in motor speed. This broad distribution indicates a larger average force per motor for perpendicular filaments compared to aligned filaments.

6.2 Dynamical evolution of filament pairs

Here, we consider the same three filament starting configurations and allow filament motion (Fig. 5, Videos 4–6). The final filament position and orientation are comparable for the explicit motor and MFMD models, while the moment expansion model overestimates the range of filament translation and rotation (Fig. 5b, c; note that filament rotation only occurs for the perpendicular initial configuration).

To compare motor activity between models over the whole simulation, we calculated the total work done by motors. We numerically integrate both filaments using the trapezoid rule [87],

\[
W_{\text{tot}} = W_{\text{lin}} + W_{\text{rot}} = \sum_{i \neq j} \int \mathbf{F}_{i,j} \cdot d\mathbf{r}_j + \sum_{i \neq j} \int \mathbf{T}_{i,j} \cdot d\theta_j, \tag{39}
\]

where \( \theta_j \) is the angle the vector \( \mathbf{u}_j \) rotates through over the simulation. The infinitesimal vector \( d\mathbf{\theta}_i = \hat{\theta}_i d\theta_i \) where

\[
\hat{\theta}_i = \frac{\mathbf{u}_i \times \dot{\mathbf{u}}_i}{|\mathbf{u}_i \times \dot{\mathbf{u}}_i|}. \tag{40}
\]

Total work computed for the mean-field models is within error of the explicit motor model (Fig. 5d). We note that the explicit motor model produces greater total work because fluctuations in motor binding cause fluctuations in sliding direction which generate larger work. Motors generate rotational work only for initially perpendicular filaments, due to the constraints. The magnitude of the rotational work is relatively small because filaments rotate slowly (due to high rotational drag and low motor torque), and this slower velocity produces less work in the overdamped limit.

The crosslinking motor number depends on the filament overlap length, which changes as filaments move (Fig. 5e–j). The crosslinking motor number in the explicit motor model lags the mean-field models initially due to differences in binding, but becomes comparable after the initial transient. As antiparallel filaments slide apart, their overlap decreases so fewer motors crosslink, while crosslinking motors continue to unbind at a constant rate. However, the overlap length has little effect on the number of motors with one head bound (Fig. 5e, h). The dynamics of motor number for parallel stationary and moving filaments are nearly identical because there is negligible sliding. (Fig. 5f, i). Moving perpendicular filaments maintain a similar overlap length to stationary perpendicular filaments, leading to an approximately constant motor number, until the plus-ends move close together (Fig. 5g, j). Then, motors continue to bind but immediately walk off, producing little force or torque.

The motor force between antiparallel filaments rapidly reaches a force plateau which persists until the antiparallel overlap length is small enough that motor binding is negligible (Fig. 5k). The nearly constant force implies that motor extension decreases as the number of crosslinking motors increases to give a constant sliding speed (Fig. 5m, Video 4). This steady-state force is an order of magnitude smaller than the stall force (Table 1). The moment expansion model shows a slower decrease in force as the overlap approaches zero compared to the MFMD model (Fig. 5h). This is a consequence of our neglect of boundary terms, which physically means neglecting motor dissociation at filament ends. This unphysical slow force decrease drives filaments beyond the zero overlap configuration to larger than expected separation (Fig. 5b).

Parallel filaments remain with their centers aligned on average because sampling the full distribution of motor crosslinking extension generates restoring force for any fluctuations away from full overlap (Eqs. 11, 13). Neither the MFMD nor the moment expansion models produce a net force, but in the explicit motor model fluctuations in motor number and binding lead to force and position fluctuations (Fig. 5f, i, l, Video 5). For perpendicular filaments, the small number of crosslinking motors results in large force fluctuations in the explicit motor model (Fig. 5j). The mean-field models show a rapid increase to half the maximum force of the antiparallel configuration followed by a decrease as the filaments align parallel (Fig. 5k, m). The lag caused by the two-step binding model is more apparent here because the explicit lower motor number means filaments move more slowly into the parallel configuration where binding is favored (Video 6).

6.3 Computational cost and accuracy

To compare the accuracy and computational cost of our models, we focus on stationary antiparallel fil-
Fig. 5 Comparison of model results for three different initial filament configurations evolved with constrained motion. **a** Schematic of initial and final filament configurations. **b** Plot of final filament center separations. **c** Plot of change in angle between filaments starting in a perpendicular configuration. Data shown is final configuration after 100 seconds for the explicit motor model and 20 seconds for MFMD and moment expansion model. **d** Plot of translational (solid bars) and rotational (hatch bars) work done by motors on filaments during simulation. **e–g** Plots of filament centers separation as a function of time. **h–j** Plots of motor number versus time. Purple and blue solid lines are the average of 48 explicit motor simulations (translucent lines) for one head bound and crosslinking motors. **k–m** Plots of motor force in the \( \hat{x} \)-direction with individual explicit motor runs (translucent blue lines) and average (solid blue). Full explicit motor model range not shown to better see average. Model error bars in (b–d) indicate the SEM of simulation realizations (\( n = 48 \)).
Fig. 6 Comparison of computational cost and accuracy of models for stationary filaments in an antiparallel configuration. Error compared against steady-state solution. a Plot of error (triangle) and CPU time (circle) vs time step $\Delta t$ for explicit motor and MFMD models. Each for explicit motor model data point consists of 48 parameter set realizations. MFMD simulations were run 3 times to ensure consistency of time scaling. Standard error of the mean (SEM) of CPU time plotted but not visible. b Plot of error and CPU time vs $\Delta s$ for MFMD model. Simulations were run 3 times to ensure consistency. SEM of CPU time plotted but not visible. c, d Plot of CPU time vs unbound motor concentration $c_o$ and filament length $L$ for the three models. Explicit motor simulation data points in C and D consist of 24 parameter set realizations while MFMD and moment expansion data points consist of 3 runs for concentration and 4 runs for filament length. SEM of CPU time plotted but not visible.

The solution error is the average magnitude of the deviation of the steady-state numerical solution from $\psi_{i,j}$ of Eqs. (23), (26), and (27),

$$\text{Error} = \frac{1}{L_i \times L_j} \int_{L_i} \int_{L_j} \left| \psi_{i,j} - \bar{\psi}_{i,j} \right| ds_i ds_j$$

$$\approx \sum_{m,n} \left| \bar{\psi}_{i,j}(m \Delta s_i, n \Delta s_j) - \psi_{i,j}(m \Delta s_i, n \Delta s_j) \right| \Delta s_i \Delta s_j,$$

(41)
results closer to those of the explicit model because stochastic fluctuations average out. The explicit motor model has a cost linear in filament length (due to the larger number of bound motors on longer filaments), while for the MFMD model it is quadratic (Fig. 6d). The cost of the moment expansion model is length independent.

7 Discussion

To improve modeling methods for cytoskeletal filaments crosslinked by motors (Fig. 1), we studied crosslinked filament pairs and compared an explicit motor model to two levels of coarse-grained mean-field motor models (Fig. 2). The explicit motor model uses Brownian dynamics and kinetic Monte Carlo to describe individual motor binding and unbinding, motion, and force generation. In the first level of coarse graining, we average over individual motors and solve a PDE for the mean-field motor density (MFMD). To further coarse grain, we compute a moment expansion of the MFMD model and solve a system of ODEs for the motor moments and filament motion.

We compared the model implementations for filaments that are initially antiparallel, parallel, or perpendicular (Fig. 3). When filaments are held stationary, the motor distribution reaches a steady state with similar average motor distribution, force, and torque for the three implementations (Fig. 4). The explicit motor simulations showed significant fluctuations that by construction are not present in the mean-field models. Interestingly, we found that a significant portion of crosslinking motors on antiparallel filaments do not reach their stall force for our parameter set.

When filaments move, the final filament separation is similar for the explicit motor and MFMD models, although the moment expansion model overestimates the range of displacement and reorientation as a result of neglecting boundary terms (Fig. 5). The dynamics of bound motor number, force, and torque were similar for the MFMD and moment expansion models. Motor fluctuations in the explicit motor model lead to greater overall work done by motors.

To compare computational cost across the model implementations, we studied stationary filaments and motors at steady state (Fig. 6). Both mean-field models have a simulation time independent of motor concentration, potentially making them faster than explicit models for systems with many motors. The moment expansion model’s CPU time is also independent of filament length, which could make it particularly efficient for systems with long filaments. Overall, the moment expansion model was $10^3 - 10^6$ faster than the other models. This method could therefore be useful for simulating bulk active filament networks.

Future work could address the simplifying assumptions and approximations made in the moment expansion model. An improved treatment of boundary terms may improve the computation of filament motion. Incorporating additional motor physics into the moment expansion model, such as non-zero-length motors, force-dependent detachment, and steric interactions between motors could improve its ability to simulate microscopic motor behavior at the mesoscale, bridging current explicit motor and continuous active network theories. Implementing the moment expansion model in systems of many filaments is of interest for testing whether the improvements in computational cost we identify are present in larger systems.

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Data Availability Statement This manuscript has associated data in a data repository. [Authors’ comment: The datasets generated and analysed for this manuscript are stored in a data repository. Data is available from the corresponding author upon request.]

Author contribution statement ARL, MAG, and MDB formulated the explicit motor binding model and kinetic Monte-Carlo algorithm. ARL and JMM wrote the Brownian dynamics simulation code. ARL, FF, MAG, MJS, and MDB derived the MFMD model and numerical integration scheme. ARL, MAG, and MDB derived the moment expansion model. ARL ran the simulations. ARL, MAG, and MDB analyzed and compared results. ARL and MJS drafted the manuscript. ARL, MAG, MJS, and MDB edited the manuscript.

Appendices

A Determining the time-step for binding

Our kinetic Monte Carlo algorithm assumes that multiple binding/unbinding events do not occur in the same time step $\Delta t$. As $\Delta t$ becomes large relative to the kinetic rates, this approximation fails. A time step is appropriate if the maximum probability of two events occurring in $\Delta t$ satisfies

$$\max\{P(C(\Delta t) \cup B(t'))|A(0))\} < \delta$$

for a tolerance $\delta$, where $A$, $B$, and $C$ denote motor bound states (including unbound, single head bound, and crosslinking) at time $\Delta t > t'$, $P(C(\Delta t) \cup B(t'))|A(0)) = P(C(\Delta t)|B(t'))P(B(t')|A(0))$ and each individual state change follows a single event Poisson process with $P(B(t)|A(0)) = 1 - \exp[-k_{A \rightarrow B} t]$. The maximum probabili-
ity for a double event occurs at $t' = t'_\text{max}$ found by solving

$$
\frac{dP(C(\Delta t) \cup B(t'))|A(0))}{dt'} = 0
$$

$$
k_{A\rightarrow B} \left(e^{k_{B\rightarrow C}(\Delta t - t')} - 1\right) - k_{B\rightarrow C} \left(e^{k_{A\rightarrow B}t'} - 1\right) = 0.
$$

(43)

While no analytic solution exists, $t'_\text{max}$ can be numerically computed.

There are four unique processes that must be considered with a two-step binding process with unbound (U), single head bound (S), and crosslinking (C) states: $U \rightarrow S \rightarrow U$, $U \rightarrow S \rightarrow C$, $S \rightarrow C \rightarrow S$, and $C \rightarrow S \rightarrow U$. The process $C \rightarrow S \rightarrow C$ has the same probability as $S \rightarrow C \rightarrow S$, similarly, $S \rightarrow U \rightarrow S$ has the same probability as $U \rightarrow S \rightarrow U$. If modeling filament motion with some force- or energy-dependent unbinding, $k_{\text{off}}$ may be large. This means that in the limit of large unbinding rate the probabilities $P(C \rightarrow S \rightarrow C) \rightarrow P(S \rightarrow C)$ and $P(C \rightarrow S \rightarrow U) \rightarrow P(S \rightarrow U)$.

B Lookup table for kinetic Monte Carlo binding

Equation (11) gives the transition probability of a singly bound motor crossing a filament as an integral of a Boltzmann factor. If $h_{\text{cl}} = 0$, $k_{\text{on},C}$ is functionally similar to an error function. However, to model non-zero-length tethers, we numerically integrate Eq. (11). Rather than directly numerically integrating at each time step, we precompute a lookup table.

The cumulative distribution function (CDF) of Eq. (11), is a function $h_{i,j}$. All other variables in the integral are constant for a given motor species. We reduce the CDF dimensionality by considering the lab position of each bound motor head and an infinite carrier line defined by the position and orientation of the unbound filament. Binding is then determined by the minimum distance $r_\perp$ between the bound motor head position and the filament ends $[s_-, s_+]$ on the carrier line.

The carrier line CDF is

$$
\text{CDF}(r_\perp, s) = \int_{-\infty}^{s} e^{-\beta U(r_\perp, s')} ds',
$$

allowing us to write the crosslinking rate as

$$
k_{\text{on},C}(r_\perp, s_+, s_-) = k_0 d e K_E [\text{CDF}(r_\perp, s_+) - \text{CDF}(r_\perp, s_-)].
$$

(45)

We notice that $e^{-\beta U_{i,j}}$ is symmetric in $s$, so CDF($r_\perp, s$) \text{- CDF($r_\perp, 0$)} is anti-symmetric. Therefore, instead of integrating from negative infinity, we use

$$
\text{CDF}'(r_\perp, s) = \text{sgn}(s) \int_{0}^{s} e^{-\beta U(r_\perp, s')} ds'
$$

(46)

and (45) to find the crosslinking rate.

We find the values of Eq. (46) by Gauss–Konrad integration. The accuracy desired sets the maximum values for $s$ and $r_\perp$. The integrand is always positive for real values of $s$ and $r_\perp$, so the CDF asymptotes for large values of either variable. The maximum of the integral is the point when the Boltzmann factor drops to the accuracy limit $\delta$. Therefore, the lookup table domain is

$$
s, r_\perp \in \left[0, \sqrt{-\frac{2 \ln(\delta)}{\beta h_{\text{cl}}} + h_{\text{cl}}} \right].
$$

(47)

Given a specified grid spacing $\Delta s$, $\Delta r$, the memory required for the lookup table scales as $(s_{\text{max}}/\Delta s) \times (r_{\perp, \text{max}}/\Delta r)$.

B.1 Interpolation of lookup table values

Since the lookup table is not a continuous function, we interpolate values between discrete grid points. The 2D linear interpolation for input values of $r_\perp$ and $s$ is

$$
\text{CDF}(r_\perp, s) \approx \left(1 + m - \frac{r_\perp}{\Delta r}\right) \left(1 + n - \frac{s}{\Delta s}\right)
$$

$$
\text{CDF}_{m,n} + \left(\frac{r_\perp}{\Delta r} - m\right) \left(1 + n - \frac{s}{\Delta s}\right) \text{CDF}_{m+1,n} + \left(1 + m - \frac{r_\perp}{\Delta r}\right) \left(\frac{s}{\Delta s} - n\right) \text{CDF}_{m,n+1} + \left(\frac{r_\perp}{\Delta r} - m\right) \left(\frac{s}{\Delta s} - n\right) \text{CDF}_{m+1,n+1},
$$

(48)

where CDF$_{m,n} =$ CDF$(m \Delta r, n \Delta s)$ are the lookup table values at $m$ and $n$ if $r_\perp$ lies within $m \Delta r$ and $(m+1) \Delta r$ and $s$ lies within $n \Delta s$ and $(n+1) \Delta s$.

B.2 Reverse lookup algorithm

When a motor head binds to a filament, the binding position probability distribution function (PDF) is defined by the Boltzmann factor. We sample the PDF by using the lookup table. To transform a uniform random variable $X$ to random variable $Y$ with an arbitrary PDF $f_Y$, $X$ is inserted into the inverted CDF of $Y$

$$
Y = \text{CDF}_Y^{-1}(X).
$$

(49)

Since the lookup table holds the CDF values and given a random number from a uniform distribution, we apply a combination of search and interpolation to quickly find the corresponding random number from the PDF. The algorithm is as follows
1. Sample a uniform random number $X \in [0, \text{CDF}_{\text{max}}]$. Note that the maximum value does not need to be 1.
2. Given $r_\perp$, locate index $m$ such that $m\Delta r \leq r_\perp \leq (m + 1)\Delta r$
3. Use $m$ to find the set of indices $\{n_-\}$ such that $\text{CDF}_{m,n_-} \leq X \leq \text{CDF}_{m,n_-+1}$ and $\text{CDF}_{m+1,n_-} \leq X \leq \text{CDF}_{m+1,n_-+1}$.
4. Use the CDF values to interpolate the binding locations $s_-, s_+$ corresponding to the perpendicular distances $r_- = m\Delta r$ and $r_+ = (m + 1)\Delta r$. For example, if the lookup table has the form of Eq. 7 and a
5. Find $s$ by interpolating the across the lookup table grid with respect to $r_\perp$

$$s \approx (s_- - s_+)(r_\perp - r_-)\Delta r + s_-$$

While this algorithm succeeds in most circumstance, the low slope of the CDF at large values of $s$ can cause errors. For example, if the lookup table has the form of Fig. 7 and a protein is located at a perpendicular distance of $r_\perp = 35$ nm, given a random number of $X = 10^3$, no value for $s_-$ will be found since CDF$(30, s_{\text{max}}) < 10^3$. To correct for this, we solve for $s_-$ using a binary search algorithm.

The binary search algorithm is as follows
1. Determine if $\text{CDF}_{m,n_-}$ or $\text{CDF}_{m+1,n_-}$ is less than $X$. If $\text{CDF}_{m,n_-} < X$, set $s_- = s_{\text{max}}$. If $\text{CDF}_{m+1,n_-} < X$, set $s_+ = s_{\text{max}}$.
2. Find other $s_\pm$ using the inverted lookup table and Eq. (50) or (51).
3. Find the average of $s_- \text{ and } s_+$
4. Use the lookup table interpolation algorithm to find the CDF($r_\perp, s_{\text{avg}}$).
5. If CDF($r_\perp, s_{\text{avg}}$) > $X$ set the larger of the two $s_\pm$ values to $s_{\text{avg}}$. Otherwise, set the smaller of the two to $s_{\text{avg}}$.
6. Repeat steps 3-5 until CDF($r_\perp, s_{\text{avg}}$) - $X$ < $\delta$ for some desired tolerance $\delta$.

This process converges at a rate $O(\log_2(\delta s_{\text{max}}))$.

C Numerical integration of the MFMD equation

We approximate the solution $\psi_{i,j}(s_i, s_j, t)$ by discretizing the solution in time and space

$$\psi_{i,j}(s_i, s_j, t) \approx \psi_{i,j}^{m,n,k} = \psi_{i,j}(m\Delta s, n\Delta s, k\Delta t)$$

for $\psi_{i,j}^{m,n,k} \in \mathbb{R}^{(M_i+1)\times(M_j+1)\times k}$, where $M_i$ is the number of discretized points along filament $i$. Additional boundary points for $m, n = 0$ are added.

We use forward Euler time-stepping so our discrete differential operator for time is

$$\frac{\partial \psi_{i,j}}{\partial t} \approx \frac{1}{\Delta t}(\psi_{i,j}^{m,n,k} - \psi_{i,j}^{m,n,k-1}).$$

To solve the hyperbolic FPE (17), we use a first-order accurate upwind method [88].

$$\frac{\partial \psi_{i,j}}{\partial s_i} \to \frac{1}{\Delta s}(\psi_{i,j}^{m,n,k} - \psi_{i,j}^{m-1,n,k}).$$

Note this only holds for the indices $0 < m$ and $0 < n$. The matrix representation for Eq. (55) is

$$\frac{1}{\Delta s} (\psi_{i,j}^{0,n,k} - \psi_{i,j}^{1,n,k}) = \psi_{M_i-1,n,k}^{i,j}$$

where $c_m$ and $d_m$ are chosen to satisfy the boundary conditions. We choose the notation $>m,n$ for this matrix. To differentiate along $s_j$, we use the identity $\psi_{i,j}^{m,n,k} = \psi_{i,j}^{n,m,k}$, apply $>m,n$ on the matrix, and then convert back,

$$\frac{\partial \psi_{i,j}}{\partial s_i} \to \left( \sum_{a} >a,m \psi_{a,j}^{m,n,k} \right)^T,$$

which in index notation is $\psi_{i,j}^{m,a,n} > a \psi_{i,j}^{m,n,a}$. For brevity, we use the notation $\psi_{i,j}^{m,a,n} > a \psi_{i,j}^{m,n,a}$.

The discretized Fokker–Planck Eq. (17) is then

$$\psi_{i,j}^{m,n,k+1} = \Delta t \left( - >m,n \psi_{i,j}^{a,n,k} \psi_{i,j}^{a,n,k} + c_{m,n,k} - 2k_o \psi_{i,j}^{m,n,k} \right)$$

where $U_{i,j}^{m,n,k}$ and $v_{i,j}^{m,n,k}$ are the discretized potential and velocity at time $k\Delta t$. Note that $U_{i,j}^{m,n,k} = U_j^{n,m,k}$, but $v_{i,j}^{m,n,k} = v_{i,j}^{n,m,k}$.

In cases where the flux of the motors $\frac{\partial \psi_{i,j}}{\partial s_i}$ is known at the boundaries, we construct $> a$ to satisfy the requirements. When filaments are in solution, there is zero flux from the minus ends, so all $c_m = 0$. In our simulations, motors walk of filament ends with out pausing, so $d_{M_i-1} = -1$ and $d_M = 1$ with all other $d_m = 0$. Although not modeled in this paper, some biological motors end pause at filament plus ends. To model this, $d_{M_i-1} = -1$ and every other $d_m = 0$.

D Conversion of binding parameters from an explicit to mean-field motor density model

To relate binding parameters of the one-step and multi-step binding models, we use that at steady state, the motor distribution $\psi_{i,j}$ should be equivalent for both models. Since we only compare binding kinetics, we simplify the Fokker–Planck equation to keep only the binding terms: in Eq. (17), we set $v_{i,j} = v_{j,i} = 0$,

$$\frac{\partial \psi_{i,j}}{\partial t} = 2k_o \psi_{i,j} - 2k_o \psi_{i,j}.$$
The steady-state solution is
\[ \psi_{i,j} = e^{-\beta U_{i,j}}, \tag{60} \]
which is a Boltzmann factor multiplied by an effective concentration.

The multi-step binding model can be written
\[ \frac{\partial \psi_{i,j}(s_i, s_j)}{\partial t} = \epsilon K'_{E} k_o, c (\chi_i + \chi_j) e^{-\beta U_{i,j}} - 2k_o, c \psi_{i,j}, \tag{61} \]
\[ \frac{\partial \chi_i(s_i)}{\partial t} = c_o K_a e \chi_i - k_o, S \chi_i \]
\[ + \int_{L_i} \left( k_o, c \psi_{i,j} - \epsilon K'_{E} k_o, c \chi_i e^{-\beta U_{i,j}} \right) ds_j, \tag{62} \]
\[ \frac{\partial \chi_j(s_j)}{\partial t} = c_o K_a e \chi_j - k_o, S \chi_j \]
\[ + \int_{L_i} \left( k_o, c \psi_{i,j} - \epsilon K'_{E} k_o, c \chi_j e^{-\beta U_{i,j}} \right) ds_i, \tag{63} \]
where \( \chi_i \) is the mean-field density of motors with one head bound to filament \( i \) (cf. Eq. 16). We define \( K'_E = K_E/V_{\text{bind}} \) and solve for the steady state, giving
\[ \psi_{i,j} = \frac{\epsilon K'_{E} k_o, c}{2} (\chi_i + \chi_j) e^{-\beta U_{i,j}}, \tag{64} \]
\[ \chi_i = \frac{\epsilon K'_{E} k_o, c}{2k_o, S} \left( \int_{L_i} (\chi_j - \chi_i) e^{-\beta U_{i,j}} ds_j \right) + \epsilon K_a c_o, \tag{65} \]
\[ \chi_j = \frac{\epsilon K'_{E} k_o, c}{2k_o, S} \left( \int_{L_i} (\chi_i - \chi_j) e^{-\beta U_{i,j}} ds_i \right) + \epsilon K_a c_o. \tag{66} \]

The equations for \( \chi_i \) and \( \chi_j \) have the forms
\[ X(s) = C \int_a^b (Y(t) - X(s)) K(s, t) dt + D, \tag{67} \]
\[ Y(t) = C \int_c^d (X(s) - Y(t)) K(s, t) ds + D, \tag{68} \]
where \( t \in [a, b] \) and \( s \in [c, d] \). Distributing the integrals, we can rewrite
\[ X(s) = C \int_a^b Y(t) K(s, t) dt - C X(s) F(s) + D, \tag{69} \]
\[ Y(t) = C \int_c^d X(s) K(s, t) ds - C Y(t) G(t) + D, \tag{70} \]
where \( F(s) = \int_a^b K(s, t) dt \) and \( G(t) = \int_c^d K(s, t) ds \). Solving for \( X(s) \) and \( Y(t) \) gives
\[ X(s) = \frac{D}{1 + C F(s)} + \frac{C}{1 + C F(s)} \int_a^b Y(t) K(s, t) dt, \tag{71} \]
\[ Y(t) = \frac{D}{1 + C G(t)} + \frac{C}{1 + C G(t)} \int_c^d X(s) K(s, t) ds, \tag{72} \]
After plugging Eq. (71) into (72), we find
\[ Y(t) = \frac{D}{1 + C G(t)} + \frac{CD}{1 + C G(t)} \int_a^b K(s, t) ds \]
\[ + C^2 \int_a^b \int_c^d Y(t') K(s, t) K(s, t') ds dt. \tag{73} \]
This can be rearranged into the form
\[ Y(t) = A(t) + \int_a^b Y(t') B(t, t') dt', \tag{74} \]
which implies that \( Y(t) \) and \( X(s) \) each satisfy a Fredholm equation of the second kind. Both \( A \) and \( B \) are continuous given \( K(s, t) = e^{-\beta U_{i,j}(s, t)} \), so the Fredholm equations of the second kind have unique solutions. By inspection, the solution to Eqs. (67) and (68) is \( X(s) = Y(t) = D \). When we substitute this solution in Eqs. (65) and (66), we find \( \chi_i = \chi_j = e \epsilon K_a c_o \) and
\[ \psi_{i,j} = \frac{\epsilon^2 K_a K'_E c_o e^{-\beta U_{i,j}}}{V_{\text{bind}}} . \tag{75} \]

Setting Eq. (60) equal to (75) gives
\[ c = \frac{\epsilon^2 K_a K'_E c_o}{V_{\text{bind}}}. \tag{76} \]

E Calculating binding parameters from experiments

The experimental parameters for motor binding are not always independently measured. If all but one binding parameter are known, then the unknown parameter can be found from Eq. (18) and the ratio of the number of motors with one head bound and number of motors crosslinking.

As an example, suppose we wish to find \( K_E \). The number of motors with one head bound is \( N_S = c_o K_a e L \), where \( L \) is the filament length. In vitro experiments [86] can measure the crosslinking motors number \( N_d \). Integrating Eq. (75), we obtain the model prediction for the number of crosslinking motors as
\[ N_C = c_o \epsilon^2 K_a K'_E \int_{L_i} \int_{L_j} e^{-\beta U_{i,j}} ds ds_j. \tag{77} \]
For fully parallel or antiparallel filaments of the same length with adjacent centers, the total number of motors in Eq. (77) is proportional to \( L \). If \( L \gg \sqrt[2]{2/\beta k_{cl}} \), the Gaussian integral \( \approx L \sqrt{2\pi/\beta k_{cl}} e^{-\beta k_{cl} r_{\perp}^2} \), where \( r_{\perp} \) is the center-to-center separation between filaments. The ratio of the number of crosslinking motors relative to the number motors with one head bound is
\[ \rho = \frac{N_C}{N_S} = \epsilon K'_E \sqrt{\frac{2\pi}{\beta k_{cl}}} e^{-\beta k_{cl} r_{\perp}^2}, \tag{78} \]
allowing us to estimate \( K_E = \epsilon \sqrt{\frac{2\pi}{\beta k_{cl}}} e^{\beta k_{cl} r_{\perp}^2} \).
F Gaussian integrals in the moment expansion

The source terms in the moment expansion require a double integral over two filaments. To lower the numerical integral’s computational cost, we find an analytic solution for either the semi-integrated term $Q_j^i(s_i)$ or the fully integrated term $q_{k,j}^i$.

The integrated source terms are
\[
q_{k,j}^i = c e^{-\frac{\beta}{2\pi}} \int_{L_j} s_i \exp \left\{ -\frac{s_j^2 - 2 s_j r_{k,j} - \hat{u}_i - (r_{j,i} + \hat{u}_i + \hat{u}_j s_j)^2}{\alpha^2} \right\} ds_j,
\]
\[
\int_{L_j} s_j \exp \left\{ -\frac{(s_j - r_{j,i} - \hat{u}_j - \hat{u}_i s_j)^2}{\alpha^2} \right\} ds_j.
\]

where $\alpha = \sqrt{\frac{\pi}{2\beta c_1}}$. We define the quantity $A = -r_{j,i} \cdot \hat{u}_j - \hat{u}_i \cdot \hat{u}_j s_j$ so that the integral over $s_j$ becomes
\[
Q_j^i(s_i) = \int_{L_j} s_j^\alpha (\frac{s_j + A}{\alpha})^2 ds_j.
\]

This integral has an analytic form in terms of error functions, which can be rapidly computed. For $l = 0, 1, 2, 3$, we find
\[
\begin{align*}
Q_0^i(s_i) &= \frac{\alpha \sqrt{\pi}}{2} \left[ \text{erf} \left( \frac{s_i + A}{\alpha} \right) \right] \partial L_j , \\
Q_1^i(s_i) &= -\alpha \left[ \text{erf} \left( \frac{s_i + A}{\alpha} \right) \right] \partial L_j , \\
Q_2^i(s_i) &= \alpha \left[ 2 \alpha (A - s_j) \text{erf} \left( \frac{s_j + A}{\alpha} \right) \right] \partial L_j , \\
Q_3^i(s_i) &= \alpha \left[ 2 \alpha (A^2 + s_j^2 + 3 A^2) \text{erf} \left( \frac{s_j + A}{\alpha} \right) \right] \partial L_j ,
\end{align*}
\]

G Moment expansion boundary terms

To generally define boundary conditions, instead of integrating over both $s_i$ and $s_i$, we integrate over just one variable. This makes the boundary condition a function of a single filament attachment position. For example, the boundary terms for the first filament are
\[
B_j^i(s_i) = \int_{L_j} \left( 2 k_o e^{-\beta u_{i,j}} + (2 \kappa - 2 k_o) \psi_{i,j} \right)
- (v_o + \kappa(r_{j,i} \cdot \hat{u}_j + \hat{u}_i \cdot \hat{u}_j s_j - s_j)) \frac{\partial \psi_{i,j}}{\partial s_j}
- (v_o + \kappa(r_{j,i} \cdot \hat{u}_j + \hat{u}_i \cdot \hat{u}_j s_j - s_j)) \frac{\partial \psi_{i,j}}{\partial s_j},
\]

These boundary terms are evaluated at $-L_j/2$ and $L_j/2$. We derive a recursion relation by integrating Eq. (85) over $s_j$ and using the definition in Eq. (37)
\[
B_j^i(s_i) = 2 k_o c Q_j^i(s_i) + l (v_o + \kappa(r_{j,i} \cdot \hat{u}_j + \hat{u}_i \cdot \hat{u}_j s_j)) B_j^i - (2 k_o + \kappa(l - 1)) B_j^i \\
- (v_o + \kappa(r_{j,i} \cdot \hat{u}_j + \hat{u}_i \cdot \hat{u}_j s_j)) \frac{\partial B_j^i}{\partial s_j} - \kappa \hat{u}_i \cdot \hat{u}_j \frac{\partial B_j^i}{\partial s_j} \\
- \left[ s_j (v_o + \kappa(r_{j,i} \cdot \hat{u}_j + \hat{u}_i \cdot \hat{u}_j s_j - s_j)) \psi_{i,j} (s_i, s_j) \right] \partial L_j ,
\]

Solving this equation requires finding the time evolution of the boundary term spatial derivatives, which solve
\[
\begin{align*}
\frac{\partial B_j^i(s_i)}{\partial s_j} &= 2 k_o c Q_j^i(s_i) + l (v_o + \kappa(r_{j,i} \cdot \hat{u}_j + \hat{u}_i \cdot \hat{u}_j s_j)) B_j^i - (2 k_o + \kappa(l - 2)) \frac{\partial B_j^i}{\partial s_j} \\
- (v_o + \kappa(r_{j,i} \cdot \hat{u}_j + \hat{u}_i \cdot \hat{u}_j s_j - s_j)) \frac{\partial^2 B_j^i}{\partial s_j^2} + \kappa \hat{u}_i \cdot \hat{u}_j \frac{\partial^2 B_j^i}{\partial s_j^2} \\
+ \text{corner terms}.
\end{align*}
\]

This shows that the boundary terms do not close. However, if the higher-order terms or their coefficients are small compared to the moments $\mu_{k,j}^i$, we may take a zeroth-order approximation. We consider this approximation in Sect. 6.

References

1. Jonathon Howard, Mechanics of Motor Proteins and the Cytoskeleton (Sinauer Associates Publishers, Sunderlund, Mass, 2001)
2. Dennis Bray, Cell Movements: From Molecules to Motility, 2nd edn. (Garland Pub, New York, 2001)
3. Laurent Blanchin, Rajaa Boujemaa-Paterski, Cécile Sykes, Julie Plastino, Actin dynamics, architecture, and mechanics in cell motility. Physiol. Rev. 94(1), 235–263 (2014)
4. Thomas D. Pollard, John A. Cooper, Actin, a central player in cell shape and movement. Science 326(5957), 1208–1212 (2009)
5. J.Richard McIntosh, Maxim I Molodtsov, Fazly I Ataullakhanov, Biophysics of mitosis. Q. Rev. Biophys. 45(2), 147–207 (2012)
6. A.F. Huxley, Muscle structure and theories of contraction. Prog. Biophys. Biophys. Chem 7, 255–318 (1957)
7. A.F. Huxley, S. Tideswell, Filament compliance and tension transients in muscle. J. Muscle Res. Cell Motil. 17(4), 507–511 (1996)
8. A.F. Huxley, S. Tideswell, Rapid regeneration of power stroke in contracting muscle by attachment of second myosin head. J. Muscle Res. Cell Motil. 18(1), 111–114 (1997)
9. Stephanie L. Gupton, Clare M. Waterman-Storer, Spatiotemporal feedback between actomyosin and Focal-Adhesion systems optimizes rapid cell migration. Cell 125(7), 1361–1374 (2006)
10. Maxime F. Fournier, Roger Sauser, Davide Ambrosi, Jean-Jacques Meister, Alexander B. Verkhovsky, Force transmission in migrating cells. J. Cell Biol. 188(2), 287–297 (2010)

11. Erín L. Barnhart, Kun-Chun Lee, Kinneret Keren, Alex Mogilner, Julie A. Theriot, An adhesion-dependent switch between mechanisms that determine motile cell shape. PLOS Biol. 9(5), e1000159 (2011)

12. G. Laevsky, Cross-linking of actin filaments by myosin II is a major contributor to cortical integrity and cell motility in restrictive environments. J. Cell Sci. 116(18), 3761–3770 (2003)

13. Iain Hagan, Mitsuhiro Yanagida, Kinesin-related cut 7 protein associates with mitotic and meiotic spindles in fission yeast. Nature 356(6364), 74 (1992)

14. W.S. Saunders, D. Koshland, D. Eshel, I.R. Gibbons, M.A. Hoyt, Saccharomyces cerevisiae kinesin- and dynein-related proteins required for anaphase chromosome segregation. J. Cell Biol. 128(4), 617–624 (1994)

15. Tarun M. Kapoor, Thomas U. Mayer, Margaret L. Coughlin, Timothy J. Mitchison, Probing spindle assembly mechanisms with monastral, a small molecule inhibitor of the mitotic kinesin, Eg5. J. Cell Biol. 150(5), 975–988 (2000)

16. Shang Cai, Lesley N. Weaver, Stephanie C. Emms-McClung, Claire E. Walczak, Kinesin-14 family proteins HSET/XCTK2 control spindle length by cross-linking and sliding microtubules. Mol. Biol. Cell 20(5), 1348–1359 (2008)

17. R. Wollman, G. Civelekoglu-Scholey, J.M. Scholey, A. Mogilner, Reverse engineering of force integration during mitosis in the Drosophila embryo. Mol. Syst. Biol. 4(1), 195 (2008)

18. Gul Civelekoglu-Scholey, Jonathan M. Scholey, Mitotic force generators and chromosome segregation. Cell. Mol. Life Sci. 67(13), 2231–2250 (2010)

19. Zhen-Yu. She, Wan-Xi Yang, Molecular mechanisms of kinesin-14 motors in spindle assembly and chromosome segregation. J. Cell Sci. 130(13), 2097–2110 (2017)

20. Krumo Vukusić, Renata Buda, Agneza Bosilj, Ana Milas, Nenad Pavin, Iva M. Tolić, Microtubule sliding within the bridging fiber pushes kinetochore fibers apart to segregate chromosomes. Dev. Cell 45(1), 11–23.e6 (2017)

21. Stijn Ganguy, Lucy S. Williams, Isabel M. Palacios, Raymond E. Goldstein, Cytoplasmic streaming in Drosophila oocytes varies with kinesin activity and correlates with the microtubule cytoskeleton architecture. Proc. Nat. Acad. Sci. 109(38), 15109–15114 (2012)

22. Peter Satir, Studies on cilia. J. Cell Biol. 39(1), 77–94 (1968)

23. Keith E. Summers, I.R. Gibbons, Adenosine triphosphate-induced sliding of tubules in trypsin-treated flagella of sea-urchin sperm. Proc. Nat. Acad. Sci. U.S.A. 68(12), 3092–3096 (1971)

24. Stephen M. King, Turning dyneins off bends cilia. Cytoskeleton 75(6), 372–381 (2018)

25. F.J. Nedelec, T. Surrey, A.C. Maggs, S. Leibler, Self-organization of microtubules and motors. Nature 389(6648), 305–308 (1997)

26. Thomas Surrey, François Nédélec, Stanislas Leibler, Eric Karsenti, Physical properties determining self-organization of motors and microtubules. Science 292(5519), 1167–1171 (2001)

27. F. Backouche, L. Haviv, D. Groswater, A. Bernheim-Groswater, Active gels: dynamics of patterning and self-organization. Phys. Biol. 3(4), 264–273 (2006)

28. T. Sanchez, D.T.N. Chen, S.J. DeCamp, M. Heymann, Z. Dogic, Spontaneous motion in hierarchically assembled active matter. Nature 491(7421), 431–434 (2012)

29. Amin Doostmohammadi, Jordi Ignés-Mullol, Julia M. Yeomans, Francesc Sagüés, Active nematic. Nat. Commun. 9(1), 3246 (2018)

30. Linnea M. Leima, Stephen J. DeCamp, Zhihong You, Luca Giomi, Zvonimir Dogic, Statistical properties of autonomous flows in 2D active nematics. Soft Matter 15(15), 3264–3272 (2019)

31. G. Duclos, R. Adkins, D. Banerjee, M.S.E. Peterson, M. Varghese, I. Kolvin, A. Baskaran, R.A. Pelcovits, T.R. Powers, A. Baskaran, F. Toschi, M.F. Hagan, S.J. Streichan, V. Vitelli, D.A. Beller, Z. Dogic, Topological structure and dynamics of three-dimensional active nematics. Science 367(6482), 1120–1124 (2020)

32. Jan Brugués, Valeria Nuzzo, Eric Mazur, Daniel J. Needleman, Nucleation and transport organize microtubules in metaphase spindles. Cell 149(3), 554–564 (2012)

33. Johanna Roostalu, Jamie Rickman, Claire Thomas, François Nédélec, Thomas Surrey, Determinants of Polar versus nematic organization in networks of dynamic microtubules and mitotic motors. Cell 175(3), 796–808.e14 (2018)

34. Kimberly L. Weirich, Kinjal Dusibiswas, Thomas A. Witten, Suriyanarayanan Vaikuntanathan, Margaret L. Gardel, Self-organizing motors divide active liquid droplets. Proc. Nat. Acad. Sci. 116(23), 11125–11130 (2019)

35. Francois Nedelec, Dietrich Foethke, Collective langevin dynamics of flexible cytoskeletal fibers. New J. Phys. 9(11), 427 (2007)

36. Konstantin Popov, James Komianos, Garegin A. Papoian, MEDYAN: mechanochemical simulations of contraction and polarity alignment in actomyosin networks. PLOS Comput. Biol. 12(4), e1004877 (2016)

37. Simon L. Freedman, Shiladitya Banerjee, Glen M. Hocky, Aaron R. Dinner, A versatile framework for simulating the dynamic mechanical structure of cytoskeletal networks. Biophys. J. 113(2), 448–460 (2017)

38. D.A. Head, W.J. Briers, Gerhard Gymmer, Nonequilibrium structure and dynamics in a microscopic model of thin-film active gels. Phys. Rev. E 89(3), 032705 (2014)

39. Igor S. Aranson, Lev S. Tsimring, Pattern formation of microtubules and motors: inelastic interaction of polar rods. Phys. Rev. E 71(5), 050901 (2005)

40. K. Kruse, J.F. Joanny, F. Jülicher, J. Prost, K. Sekimoto, Generic theory of active polar gels: a paradigm for cytoskeletal dynamics. Eur. Phys. J. E 16(1), 5–16 (2005)

41. David Saintillan, Michael J. Shelley, Instabilities and pattern formation in active particle suspensions: kinetic theory and continuum simulations. Phys. Rev. Lett. 100(17), 178103 (2008)

42. Luca Giomi, Mark J. Bowick, Xu Ma, M. Cristina Marchetti, Defect annihilation and proliferation in active nematics. Phys. Rev. Lett. 110(22), 228101 (2013)

43. Tong Gao, Robert Blackwell, Matthew A. Glaser, M.D. Betterton, Michael J. Shelley, Multiscale model-
75. Robert Blackwell, Oliver Sweezy-Schindler, Christopher Edelmaier, Zachary R. Gergely, Patrick J. Flynn, Salvador Montes, Ammon Crapo, J. Alireza Doostan, Richard McIntosh, Matthew A. Glaser, Meredith D. Betterton, Contributions of microtubule dynamic instability and rotational diffusion to kinetochore capture. Biophys. J. 112(3), 552–563 (2017)

76. Shenshen Wang, Peter G. Wolynes, On the spontaneous collective motion of active matter. Proc. Nat. Acad. Sci. 108(37), 15184–15189 (2011)

77. A.J.T.M. Mathijssen, A. Doostmohammadi, J.M. Yeomans, T.N. Shendruk, Hydrodynamics of microswimmers in films. J. Fluid Mech. 806, 35–70 (2016)

78. Fundamental algorithms for scientific computing in python, P. Virtanen, R. Gommers, T.E. Oliphant, M. Haberland, T. Reddy, D. Cournapeau, E. Burovski, P. Peterson, W. Weckesser, J. Bright, S.J. van der Walt, M. Brett, J. Wilson, K.J. Millman, N. Mayorov, A.R.J. Nelson, E. Jones, R. Kern, E. Larson, C.J. Carey, L. Polat, Y. Feng, E.W. Moore, J. VanderPlas, L. Axalade, J. Perktold, R. Cimrman, I. Henriksen, E.A. Quintero, C. R. Harris, A.M. Archibald, A.H. Ribeiro, F. Pedregosa, P. van Mulbregt, and SciPy 1.0. Contributers. Scipy 1.0. Nature Methods 17, 261–272 (2020)

79. C. Alan, Hindmarsh, ODEPACK, a Systematized Collection of ODE Solvers, in Scientific Computing, ed. by R.S. Stepleman, et al. (Science and Education Publishing, North-Holland, Amsterdam, 1983), pp. 55–64

80. Alberts. Molecular biology of the cell, 5th edition by B. Alberts, A. Johnson, J. Lewis, M. Raff, K. Roberts, and P. Walter, Biochem. Mol. Biol. Edu. 36(4), 317–318 (2008)

81. K. Kawaguchi, S. Ishitawa, Nucleotide-dependent single- to double-headed binding of kinesin. Science 291(5504), 667–669 (2001)

82. A.S. Kashina, R.J. Baskin, D.G. Cole, K.P. Wedaman, W.M. Saxton, J.M. Scholey, A bipolar kinesin. Nature 379(6562), 270–272 (1996)

83. Adina Gerson-Gurwirtz, Christina Thiede, Natalia Movshovich, Vladimir Fridman, Maria Podolskaya, Tsali Danieli, Stefan Lakämper, Dieter R. Kloppeinstein, Christoph F. Schmidt, Larisa Gheber, Directionality of individual kinesin-5 Cin8 motors is modulated by loop 8, ionic strength and microtubule geometry. EMBO J. 30(24), 4942–4954 (2011)

84. Megan T. Valentine, Polly M. Fordyce, Troy C. Krzysiatk, Susan P. Gilbert, Steven M. Block, Individual dimers of the mitotic kinesin motor Eg5 step progressively and support substantial loads in vitro. Nat. Cell Biol. 8(5), 470–476 (2006)

85. J.C. Cochran, Kinesin motor enzymology: chemistry, structure, and physics of nanoscale molecular machines. Biophys. Rev. 7(3), 269–299 (2015)

86. Yuta Shimamoto, Scott Forth, Tarun M. Kapoor, Measuring pushing and braking forces generated by ensembles of kinesin-5 crosslinking two microtubules. Dev. Cell 34(6), 669–681 (2015)

87. H.I. Siyyam, M.I. Syam, The modified trapezoidal rule for line integrals. J. Comput. Appl. Math. 84(1), 1–14 (1997)

88. Randall J. LeVeque, Finite Difference Methods for Ordinary and Partial Differential Equations: Steady-State and Time-Dependent Problems (Society for Industrial and Applied Mathematics, Philadelphia, PA, 2007)