Motor Protein Accumulation on Antiparallel Microtubule Overlaps

Hui-Shun Kuan1,2 and Meredith D. Betterton1,*

1Department of Physics and 2Department of Chemistry and Biochemistry, University of Colorado at Boulder, Boulder, Colorado

ABSTRACT Biopolymers serve as one-dimensional tracks on which motor proteins move to perform their biological roles. Motor protein phenomena have inspired theoretical models of one-dimensional transport, crowding, and jamming. Experiments studying the motion of Xklp1 motors on reconstituted antiparallel microtubule overlaps demonstrated that motors recruited to the overlap walk toward the plus end of individual microtubules and frequently switch between filaments. We study a model of this system that couples the totally asymmetric simple exclusion process for motor motion with switches between antiparallel filaments and binding kinetics. We determine steady-state motor density profiles for fixed-length overlaps using exact and approximate solutions of the continuum differential equations and compare to kinetic Monte Carlo simulations. Overlap motor density profiles and motor trajectories resemble experimental measurements. The phase diagram of the model is similar to the single-filament case for low switching rate, while for high switching rate we find a new (to our knowledge) low density-high density-low density-high density phase. The overlap center region, far from the overlap ends, has a constant motor density as one would naively expect. However, rather than following a simple binding equilibrium, the center motor density depends on total overlap length, motor speed, and motor switching rate. The size of the crowded boundary layer near the overlap ends is also dependent on the overlap length and switching rate in addition to the motor speed and bulk concentration. The antiparallel microtubule overlap geometry may offer a previously unrecognized mechanism for biological regulation of protein concentration and consequent activity.

INTRODUCTION

The motion of motor proteins on biopolymers is important for diverse biological processes (1). Actin, microtubules, and nucleic acids can serve as one-dimensional tracks on which motor proteins (including myosins, kinesins, helicases, and ribosomes) move (2,3). Motors must accumulate on filaments in sufficient density to perform their biological roles. This motor accumulation along a biopolymer is affected by two key effects: directional walking of motors, and motor binding to/unbinding from the filament.

Motor accumulation on filaments is related to extensive theoretical work on asymmetric simple exclusion processes (ASEP) (4). In ASEP models, particles move on a one-dimensional lattice by biased hopping and experience excluded-volume interactions with other particles. These models have been applied to diverse examples of one-dimensional nonequilibrium transport ranging from molecular motors to vehicular and pedestrian traffic. Active particle motion leads to nonzero flux of particle density and nontrivial flux and density profiles. To compare to experimental studies of molecular motors, ASEP models have been extended to incorporate important biophysical ingredients such as binding kinetics. A model extending the totally asymmetric simple exclusion process (TASEP) to include motor binding and unbinding (Langmuir kinetics) predicted motor density profiles along single fixed-length filaments (5). Experimental work measured kinesin-8 motor protein traffic jams on microtubules and found good agreement with the predicted density profiles (6).

Motor motion on cytoskeletal filaments is important for biological length regulation, including regulation of the length of the polymer (7), lengths of overlap regions between filaments (8), and the length of cytoskeletal assemblies such as the mitotic spindle (9,10) and even whole cells (11,12). The case of regulation of microtubule (MT) length has seen the most work. MTs undergo nonequilibrium polymerization dynamics characterized by switching between distinct growing and shrinking states. While this dynamic instability alone leads to a broad distribution of MT lengths (13), numerous proteins targeted to MT ends modify their dynamics and can dramatically alter the length...
distribution (14). Single MTs can have their length regulated, for example, by kinesin-8 motors that walk with directional bias and shorten the MT from its end (7,15,16). Theoretical work on TASEP-like models has described how length-dependent depolymerization affects otherwise static filaments (7,17,18), filaments with simplified polymerization kinetics (19–22), and dynamic MTs (23–25).

Because the mitotic spindle includes arrays of overlapping antiparallel MTs at the spindle midzone, regulation of MT overlaps is important for mitotic spindle function and cytokinesis. The MT crosslinking protein PRC1/Ase1/MAP65 and kinesin-4 motors (chromokinesins) play roles in maintenance of the spindle midzone in anaphase (26–28), along with other motors and MAPs (29). Direct binding interactions of PRC1 and kinesin 4 can cause length-dependent accumulation at plus ends of single MTs (30). Bieling et al. (8) reconstituted a minimal system of stable antiparallel MT overlaps in which PRC1 bound preferentially to overlapping regions of antiparallel MTs (8) (see also Subramanian et al. (31)). PRC1 recruited the kinesin-4 motor Xklp1 to the overlap. Xklp1 motors could bind to and unbind from the MTs, walk toward the plus end of each MT, and switch between the two MTs at a relatively high rate (8). Motors present near the MT plus ends slowed the polymerization speed, consistent with earlier work showing that Xklp1 inhibits dynamic instability (32) and affects spindle MT mass (33). As a result, antiparallel MT overlaps reached a constant length that depended on the bulk concentration of motors. This work demonstrated that motor-dependent regulation of dynamics and length can occur not just for single MTs, but also for overlapping MT pairs.

Here we model the density profiles of motor proteins on antiparallel MT overlaps of fixed lengths. We do not explicitly consider MT length regulation in this system, as the combination of binding kinetics and coupled switching of the motors between the two antiparallel filaments is sufficient to produce a rich phenomenology. Our work is an extension of the TASEP with binding kinetics on a single filament (34) to include two antiparallel filaments coupled by switching; it is also an extension of TASEP models of two antiparallel lanes with lane switching (35,36) to incorporate binding and unbinding kinetics.

We first develop the model and show that measured and estimated parameters can give overlap motor density profiles and motor trajectories qualitatively similar to those found in the experiments of Bieling et al. (8). We then develop an analytic solution to the mean-field steady-state equations, and use it to determine the phase diagram of the model. For high motor switching rate between filaments, we find a new (to our knowledge) low density-high density-low density-high density phase. We then study the model for the reference parameter set in more detail. Because the motor density profiles are controlled by both boundary conditions and a nonlocal total binding constraint, the density in the center of the overlap depends not just on motor binding kinetics but also on motor speed and overlap length. We find an analytical approximation that describes the overlap center density and the size of the motor-dense boundary layer near the overlap ends. The degree of motor accumulation near the overlap ends depends on the overlap length and motor switching rate in addition to the motor speed and bulk concentration. The coupling of motor motion, binding, and switching kinetics on antiparallel MT overlaps may therefore offer a previously unrecognized mechanism for the control of one-dimensional motor density profiles.

MATERIALS AND METHODS

Model

In this section, we develop a mathematical model of motor density on antiparallel filament overlaps, inspired by the experiments of Bieling et al. (8). In the experiments of Bieling et al. (8), the cross-linking protein PRC1 binds preferentially to overlapping regions of antiparallel MTs and recruits the kinesin motor protein Xklp1 to the overlap region. Because PRC1 and therefore the motors are present at much higher concentrations in the overlap, in the model we consider the overlap region only and model the regions of single filaments as sources or sinks of motor proteins (Fig. 1 A). While previous work has shown that PRC1/Ase1 alone can develop density inhomogeneities and exert forces on sliding MTs (37,38), in the experiments of Bieling et al. (8), little to no MT sliding occurred and the PRC1 distribution was uniform in the overlaps (8). Therefore we do not explicitly model the PRC1 molecules or their spatial distribution, but assume they are uniformly distributed so that each lattice site in the overlap is identical with a binding affinity that would correspond to the density-weighted average of the PRC1 and bare tubulin affinities. We treat each MT as a single track and neglect the multiple-protofilament structure of the MT, consistent with previous theoretical work (7,17–22,24). In our model, motors can bind to and unbind from each of the MTs, walk toward the plus end of a MT, and switch between the two MTs (Fig. 1 A).

There are two competing processes in this model: motor stepping (TASEP); and motor binding, unbinding, and MT switching (Langmuir kinetics). The Langmuir kinetics dominate the system behavior if the overlap length is sufficiently large, $N > 1$, so that each bound motor can only walk for a short fraction of the overlap length before it unbinds. Thus, the competition between TASEP and Langmuir kinetics occurs only if the overall binding rate to one MT in the overlap $K_{on} = Nk_{on}$ (where $c$ is the bulk motor concentration), and unbinding rate $K_{off} = Nk_{off}$, are of similar magnitude to the motor speed $v$ (5).

In the next section, we develop the discrete microscopic model and corresponding stochastic simulation, then derive the mean-field continuum description.

Discrete model

We consider two overlapping antiparallel MTs of fixed length, so the number of sites $N$ is fixed (Fig. 1 A). At each site, motor binding or unbinding can occur with binding rate $k_{on}c$, where $k_{on}$ is the binding rate constant per site and $c$ is the bulk motor concentration; the unbinding rate is $k_{off}$. Each bound motor steps at rate $v$ to the next site toward the MT plus end (if the next site is unoccupied), and switches at rate $s$ to the site on the adjacent MT (if that site is unoccupied). Note that Xklp1 is plus-end directed, and we therefore assume that the motion on a single MT is unidirectional.

The occupation number $i$ is 1 if site $i$ is occupied or 0 if site $i$ is empty. Then the equations for interior sites ($2 < i < N - 1$) on MTs with plus ends to the right ($R$) and left ($L$) are
Simulations used to generate these images used the reference parameter set (Table 1) and the indicated bulk motor concentrations. (C) Simulated kymograph made from our kMC model with motor spatial position on the horizontal axis and time increasing downwards. Horizontal scale bar, 10 μm. Vertical scale bar, 5 μm.

At steady state, we can derive a total binding constraint on the equations by summing all of the sites on both filaments. The bulk flux terms of the form $n_{i-1}(t) - \tilde{n}_i(t)$ sum to zero and only the binding and boundary terms remain:

$$\sum_{i=2}^{N-1} [2k_{on}c - (k_{on}c + k_{off}) (\tilde{n}_{R,i} + \tilde{n}_{L,i})] + v\alpha [2 - \tilde{n}_{R,1}(t) - \tilde{n}_{L,N}(t)] - v\beta [\tilde{n}_{R,N}(t) + \tilde{n}_{L,1}(t)] = 0.$$  

(7)

We find a binding constraint on the total motor binding

$$\sum_{i=1}^{N} \tilde{n}_{R,i} + \tilde{n}_{L,i} = 2(N-2) \frac{k_{on}c}{k_{on}c + k_{off}} + \frac{2v(\alpha(1-\alpha) - \beta(1-\beta))}{k_{on}c + k_{off}}$$

(8)

where we have defined the Langmuir density $\rho_0 = k_{on}c/(k_{on}c + k_{off})$. Therefore, at steady state, an equilibrium involving binding, unbinding, and the filament-end boundary conditions must be reached on average for the entire overlap. This is related to the zero-current condition found in previous work on the antiparallel TASEP without binding kinetics (35,36). We did not find an analytical solution to the discrete equations. Instead, we performed kinetic Monte Carlo (kMC) simulations of the discrete model (Supporting Material) and compared it to solutions in the continuum limit.

**Mean-field continuum model**

We derive the mean-field continuum model as in Parmeggiani et al. (5) by taking the stationary average $\langle \tilde{n}_i \rangle \equiv \rho_i$, and applying the random phase approximation $\langle \tilde{n}_{i+1} \rangle \approx \rho_i \langle \tilde{n}_i \rangle$. We also assume motor commutation during track switching, $(\tilde{n}_{R,i} \tilde{n}_{L,j}) = \langle \tilde{n}_{L,j} \tilde{n}_{R,i} \rangle$, to give discrete mean-field equations with linear switching terms (Supporting Material). We take the continuum limit and nondimensionalize the parameters and variables by choosing the length of the overlap, $L$, as the unit of length and $LV$ as the unit of time. Capital letters denote the nondimensionalized parameters.
Motors on Antiparallel MT Overlaps

(S = sL/v and so on; Supporting Material). We choose x = 0 as the center of the overlap, and the boundary conditions become ρ_R(−0.5) = ρ_L(0.5) = α and ρ_R(0.5) = ρ_L(−0.5) = 1 − β. The steady-state continuum mean-field equations are

\[ 0 = (2ρ_R - 1) \frac{\partial ρ_R}{\partial x} + K_{oc}c(1 - ρ_R) - K_{off}ρ_R - Sρ_R + Sρ_L; \]  

\[ (9) \]

\[ 0 = (1 - 2ρ_L) \frac{\partial ρ_L}{\partial x} + K_{oc}c(1 - ρ_L) - K_{off}ρ_L + Sρ_R - Sρ_L. \]  

\[ (10) \]

In the continuum mean-field model, the total binding constraint (Eq. 8) is

\[ \int_{-1/2}^{1/2} dx \rho_R = \int_{-1/2}^{1/2} dx \rho_L = \frac{K_{oc}c}{K_{oc}c + K_{off}} \left( \frac{α(1 - α) - β(1 - β)}{K_{oc}c + K_{off}} \right) = ρ_0 + \frac{α(1 - α) - β(1 - β)}{K_{oc}c + K_{off}}. \]  

\[ (11) \]

In the continuum mean-field model, the total binding constraint (Eq. 8) is

\[ \int_{-1/2}^{1/2} dx \rho_R = \int_{-1/2}^{1/2} dx \rho_L = \frac{K_{oc}c}{K_{oc}c + K_{off}} \left( \frac{α(1 - α) - β(1 - β)}{K_{oc}c + K_{off}} \right) = ρ_0 + \frac{α(1 - α) - β(1 - β)}{K_{oc}c + K_{off}}. \]  

\[ (11) \]

**RESULTS**

**Motor density and trajectories in overlap**

To study whether our model can qualitatively describe the motor density in antiparallel MT overlaps observed experimentally, we determined a reference parameter set corresponding to the experiments of Bieling et al. (8) (Table 1). Most parameters were directly measured by Bieling et al. We estimated the motor on rate constant k_{on} by comparison to the single-molecule imaging of low-density Xklp1 in an overlap (8). To estimate the minus-end boundary condition α, note that Bieling et al. found that Xklp1 binding was greatly increased on overlaps due to recruitment by PRC1. Low ionic strength (which favors motor-MT binding) was needed to observe significant Xklp1 binding to single MTs outside of overlaps. Therefore, we assume that the flux of motors into the overlap from outside is negligible, and set α = 0 in our reference parameters. The plus-end boundary condition β is related to the motor unbinding rate at MT plus ends. While this end off-rate was not directly measured by Bieling et al. (8), kinesin motors typically pause at MT plus ends and have an end unbinding rate smaller than the unbinding rate in the bulk. We therefore expect that β lies between 0 (no end unbinding) and 2.7 × 10^{-3} (the value of β corresponding to an unbinding rate equal to k_{off}, the motor unbinding rate in the bulk). We found that β = 2.7 × 10^{-3} is so small that the motor density profiles were indistinguishable from those with β = 0 (Fig. S1 in the Supporting Material). Therefore we used β = 0 in our reference parameter set.

With these parameters, we studied kMC simulations of our model with varying motor concentration and overlap lengths corresponding to the steady-state values measured by Bieling et al. (8). We made simulated images that represent how our model’s motor distributions would appear in an experiment with fluorescently tagged motors (green) and antiparallel MT overlap region (red; Fig. 1 B, and Supporting Material). Motors decorate the overlap, with greater density and end accumulation for higher bulk motor concentration. The simulated images are qualitatively similar to the experimental images (see Fig. 4 in Bieling et al. (8)). We also made simulated kymographs (Fig. 1 C, and Supporting Material), which show directed motor motion with direction reversal, qualitatively similar to the experimental results (see Figs. 6 and S5 in Bieling et al. (8)). We note that the experimental kymographs show a larger population of paused/immobile motors than occurs in our model.

**Analytic solution of the steady-state continuum equations**

To further study the behavior of our model, we determined analytic steady-state solutions of the continuum mean-field

---

**TABLE 1 Parameter Values for the Reference Parameter Set**

| Symbol | Parameter Description | Reference Value | Notes |
|--------|-----------------------|-----------------|-------|
| v      | motor speed           | 0.5 μm s⁻¹      | measured by Bieling et al. (8); varied up to 8 μm s⁻¹ to study effect of varying speed on density profiles |
| k_{on} | binding rate constant | 2.7 × 10⁻⁴ nM⁻¹ s⁻¹ | estimated based on motor density profiles and kymographs to give affinity of 6.3 nM, ~15× higher than measured for motor on single MT by Bieling et al. (42) |
| c      | bulk motor concentration | 1–200 nM       | varied by Bieling et al. (8) |
| k_{off} | unbinding rate       | 0.169 s⁻¹      | measured by Bieling et al. (8) |
| s      | switching rate        | 0.44 s⁻¹       | measured by Bieling et al. (8) |
| α      | motor flux constant into overlap from MT minus end | 0 | motors bind primarily inside the overlap; see discussion in the Results; we varied α between 0 and 1 to determine the model phase diagram |
| β      | motor flux constant out of overlap from MT plus end | 0 | an upper bound on the end motor unbinding rate is β = 2.7 × 10⁻³; see discussion in the Results; we varied β between 0 and 1 to determine the model phase diagram |
| N      | number of sites       | 120–2500        | varied to study overlaps of length 1–20 μm |
| δ      | length of a single site | 8 nm           | length of an α-β tubulin dimer; see Bray (1) |
equations. One solution to Eqs. 9 and 10 is the constant Langmuir density set by binding/unbinding equilibrium, \( \rho_0 = K_{onc}/(K_{onc} + K_{off}) \). To find spatially varying solutions, we first define \( \sigma_{RL} = \rho_{RL} - 1/2 \), the difference of the motor occupancies from 1/2. The rate combinations are \( k = K_{onc} + K_{off} + S \) and \( \gamma = K_{onc} - K_{off} \). Then the equations become

\[
\frac{d\sigma_R}{dx} = \frac{k}{2} - \frac{\gamma}{4\sigma_R} - \frac{S\sigma_L}{2\sigma_R} \tag{12}
\]

\[
\frac{d\sigma_L}{dx} = -\frac{k}{2} + \frac{\gamma}{4\sigma_L} + \frac{S\sigma_R}{2\sigma_L} \tag{13}
\]

Equations 12 and 13 are well defined for \( \sigma_{RL} \neq 0 \). This allows solution of the differential equation relating \( \sigma_R + \sigma_L \) and \( \sigma_R - \sigma_L \) with one integration constant (Supporting Material). Our kMC simulation results for motor density in the overlap differ substantially from this analytic solution (Fig. S2), because the analytic solution (Eq. S18 in the Supporting Material) does not satisfy the total binding constraint (Eq. 11). Therefore, the full density profiles must be determined by matching continuum solutions corresponding to different integration constants. This mathematical result implies several remarkable properties of the density profiles, as discussed below.

**Phase diagram**

We determined the phase diagram of the model as a function of the boundary condition parameters \( \alpha \) and \( \beta \), as well as how the phase diagram changes with the other model parameters. We find four phases previously observed for the single-filament case (5), the low density (L), high density (H), low density-high density (LH), and Meissner (M) phases (Figs. 2 and S3). We distinguish two different classes of L, H, and LH phase on overlaps by whether they have a local maximum (labeled \( x \)) or a local minimum (labeled \( n \)) of the total motor density at the overlap center. In addition, for high switching rate, we observe a new (to our knowledge) low density-high density-low density-high density (LHLH) phase (Fig. 2).

We studied flows in the \( \sigma_L - \sigma_R \) phase plane to determine the motor concentration profiles (Fig. S4) and therefore the phase diagram, as discussed in the Supporting Material and a followup article (39). Representative phase diagrams for small and large switching rates are shown in Fig. 3. We note that the new (to our knowledge) LHLH phase occurs when \( \alpha < 0.5 \) and the switching frequency is sufficiently high, conditions that could occur experimentally. This phase can appear for high switching rate when two analytic solutions to the steady-state continuum equations intersect, causing transition points in the phase plane where the density is ill defined (Figs. S4 and S5). Our LHLH phase is reminiscent of multiphase coexistence, as found by Pierobon et al. (40). However, in this previous work the multiphase coexistence occurs due to a point defect in the lane, while here it occurs for spatially uniform dynamics.

The phase boundaries change with parameters of the model. Increases (decreases) to the Langmuir density (through changes in bulk motor concentration or binding/unbinding rate constants) shift the lower boundary of the LH phase closer to (farther from) the line where \( 1 - \beta = 0.5 \) when \( \alpha > \rho_0 \) and closer to (farther from) the line \( \alpha = \beta \) when \( \alpha < \rho_0 \). Similar shifts occur for the upper boundary of the LH phase. The motor speed affects the width and the shape of the LH phase: for higher motor speed...
speed, the LH phase region becomes narrower in width and the LH boundaries become more flat.

### Density profiles for reference parameters

The motor density profiles that we determine by kMC simulation are qualitatively consistent with those observed experimentally (Figs. 1 and 4). Here we illustrate our results for the reference parameter set (Table 1); in the Supporting Material we also discuss a large-S parameter set chosen to approximate the limit of large switching rate (Table S1; Fig. S6). This parameter regime typically shows the LH phase, where the density profiles have three regions separated by domain walls. For motors moving to the right, there is a boundary layer on the left with the motor density increasing from zero, then a region of approximately linearly varying density, then a sharp transition to another boundary layer of linearly increasing density that approaches 1 on the right boundary (Fig. 4). If we fix all other parameters, the boundary layer regions increase in size as the bulk motor concentration increases. The total motor density in the overlap is the sum of the motor densities on the two filaments, which have the symmetry \( r_L(x) = r_R(-x) \). The linear density variation near the overlap ends can be seen in the continuum mean-field equations (Supporting Material): the slope is \( K_{\text{on}} + S \) at the minus end and \( K_{\text{off}} + S \) at the plus end. This approximation agrees well with simulation results near the overlap ends (Figs. S2 and S7).

### Control of center density by the total binding constraint

Motor density profiles must satisfy the total binding constraint of Eq. 11, which requires that when \( \alpha = \beta = 0 \), the integral of the density on a single MT must equal \( r_0 \), the Langmuir density determined by binding/unbinding equilibrium. We verified that the total binding constraint is satisfied in our simulations by determining the integrated motor density and comparing it to \( r_0 \) (Fig. S8). However, as noted above, the analytic solutions we found for the steady-state motor density (Eqs. S18, S23, and S24) do not in general satisfy the total binding constraint. As a result, the motor concentration in the center of the filaments does not necessarily approach the Langmuir density \( r_0 = K_{\text{on}} c / (K_{\text{on}} c + K_{\text{off}}) \) (Fig. 4). This is a significant difference between the model of microtubule overlaps we study and the single-filament model (5).

The total binding constraint also controls the length of the boundary layer regions near the overlap ends. To derive an analytic approximation for this length, we approximate the motor densities as piecewise linear, and assume the...
domain walls are infinitely thin so that we can neglect them in integrating the concentration (Fig. S9). A filament is divided into three regions: boundary layers near the plus- and minus-ends (\(x_{bl}\)) and the central region. We can then determine the motor density profile (Eq. S28), the boundary layer ends (Eq. S30), and the motor density in the center of the overlap, \(\rho_c\) (Eq. S31).

The motor density at the center of the overlap is \(\rho_c = \rho_0\), even when the overlap is long. Instead, the center density depends on the motor speed and filament switching rate in addition to binding parameters (Fig. 4). In Fig. 5, we compare simulation results and the prediction of Eq. S31, showing that our approximation is in good agreement with simulations. (Note that our approximation of the density as linearly varying becomes less exact for slow motor speeds, leading to a discrepancy between simulation results and our analytic approximation.) For comparison, we show the Langmuir density \(\rho_0\) for the same parameters. Varying motor speed has no effect on \(\rho_0\) (right panel), but a significant effect on \(\rho_c\). To further illustrate the dependence of the center density on motor kinetic parameters, we show in Fig. 6 examples of motor density profiles from kMC simulations with varying motor speed. Changing motor kinetic parameters can have a large effect on the center density in the overlap, even for systems with identical binding kinetic parameters and \(\rho_0\).

**Control of end accumulation by the total binding constraint**

The length of the boundary layer at the overlap ends depends on motor binding and kinetic parameters as well as the overlap length (Eq. S30). We compared the analytic approximation to simulation results and found good agreement (Fig. S10), allowing us to predict how varying experimental control parameters will alter the degree of motor accumulation near the overlap ends (Fig. 7). We note that although the boundary layer lengths predicted for typical experimental parameters are below the optical resolution limit of \(\approx 250\) nm, they could be distinguished by fits of fluorescent motor intensity profiles to model predictions (compare to Figs. 1, 4, and 6), similar to how fits to a Gaussian fluorescence intensity profile allow subresolution localization of single molecules (41).

The total binding constraint means the boundary layer length is determined as fraction of the overlap length. In the single-filament model, the motor density profile in the MT minus-end boundary layer is independent of MT length (17, 24), with important consequences: MT length regulation by the kinesin-8 motor Kip3 (7, 16) depends on a motor density profile that maintains the same functional form near the minus end as MT length changes. By contrast, on antiparallel MT overlaps, the density profile, boundary-layer length, and extent of motor accumulation at the overlap-ends change with overlap length (Fig. 7). This means that a length-regulation mechanism identical to that of Kip3 cannot occur on antiparallel MT overlaps.

**CONCLUSIONS**

We studied a model of motor motion on antiparallel MT overlaps that incorporates motor binding and unbinding, plus-end directed motor motion, and switching between filaments (Fig. 1). Our model is inspired by the experiments of Bieling et al. (8) on the motion of the kinesin-4 motor Xklp1 on antiparallel MT overlaps. Our model is an extension of previous theory that studied motor motion on a single MT with binding kinetics (5, 34), or motor motion on two antiparallel filaments with lane switching, but no binding kinetics (35, 36). To our knowledge, this is the first theoretical study of a two-lane TASEP model with oppositely oriented lanes, switching, and binding/unbinding.

To compare to the experiments of Bieling et al. (8), we used measured or estimated parameters (Table 1). Because PRC1 recruits motors directly into the overlap and motor binding to single MTs is much weaker, we neglect motor binding to MTs outside of the overlap and set the minus-end flux parameter \(\alpha = 0\). Using the bulk motor unbinding rate \(K_{off}\) as an upper bound on the plus-end unbinding rate, we find \(\beta \leq 2.7 \times 10^{-3}\). This value is sufficiently small that it gives motor density profiles indistinguishable from those with \(\beta = 0\) (Fig. S1). Therefore no-flux boundary conditions with \(\alpha = \beta = 0\) are used to model the experiments. Simulated images of the motor distribution in an overlap and simulated kymographs of motor trajectories are similar to those found experimentally (Fig. 1).

**FIGURE 5** Motor density at the center of the overlap. (Left) Variation with bulk motor concentration; (right) variation with motor speed. Points indicate simulation results and dashed lines theoretical prediction from Eq. S31. The red line shows the value of the Langmuir density \(\rho_0\) for the same parameters. To see this figure in color, go online.
We derived analytical and approximate solutions of the continuum steady-state equations and compared them to kMC simulation results (Fig. S2). Focusing on the symmetric case where the boundary conditions $a$ and $b$ are the same for both filaments in the overlap, we used both kMC simulations and phase plane flows (Figs. S3–S5) to determine the nonequilibrium phases possible in our model (Fig. 2). For low rate of motor switching between filaments in the overlap, we find the low-density, high-density, low-density/high-density, and Meissner phases previously studied for the single-lane case (5). In addition, for high switching rate we find a novel (to our knowledge) low density-high density-low density-high density phase with three domain walls. We determined representative phase diagrams for small and large switching rates (Fig. 3). We discuss determination of the phase diagram in more detail in a longer follow-up article (39).

We then studied the model in more detail both for the reference parameters and a high-switching-rate parameter set (Table S1). Results of kMC simulations (Figs. 4 and S6) agree well with exact and approximate solutions of the continuum steady-state equations (Figs. S2 and S7). In contrast to systems in which motors move on single filaments (7,17–22,24), we find that antiparallel overlaps with no-flux boundary conditions have zero total current. This leads to a total binding constraint that the integral of the total motor density on a single filament must equal $\rho_0$, the motor density set by binding/unbinding equilibrium (Fig. S8).

For the experimentally relevant low-density/high-density coexistence phase, the density profiles are approximately piecewise linear (Fig. S9). This motivates an analytic approximation to determine the density profiles consistent with the total binding constraint and gives analytic expressions for the overlap center motor density and the length of the boundary layer in which motors accumulate near the overlap ends. We find that as a result of the total binding constraint, the motor density at the center of the overlap is not determined solely by the motor binding equilibrium, but is also controlled by the overlap length, motor speed, and filament switching rate (Figs. 5 and 6). These same parameters control the length of the boundary layer at the overlap ends where motors accumulate (Fig. S10).

The mitotic spindle contains arrays of overlapping antiparallel MTs to which multiple motors and cross linkers bind (29). The surprising differences in motor density profiles between single filaments and the antiparallel overlaps we study here are therefore of interest in the study of the spindle midzone. For antiparallel overlaps, both the motor density far from the overlap ends and the number of motors near the overlap ends can be tuned not just by motor binding kinetics but also by motor speed, filament switching rate, and overlap length (Fig. 7). The antiparallel filament geometry gives biological systems additional handles to control motor density, its spatial distribution, and therefore motor function. Motor density can affect recruitment of other proteins and MT dynamics. Therefore, this previously undescribed mechanism of regulation of motor density along MTs may offer advantages to the control of motor activity.

For single MTs, length regulation by the kinesin-8 motor Kip3 depends on a functional form of motor density that is independent of MT length (7,16). The physics we describe here means that the motor density and extent of motor

![FIGURE 6 Dependence of density profiles on motor speed. (Left) Motor density $p_R(x)$ on MT with rightward-moving motors. (Right) Total motor density $p_R(x) + p_L(x)$ on both MTs in the overlap. Varying motor speed can significantly alter the motor density at the center of the overlap. These simulations use the large-$S$ parameter set of Table S1 with bulk motor concentration $c = 200$ nM and the motor speeds indicated in the legend. To see this figure in color, go online.](image1)

![FIGURE 7 Control of boundary layer length. (Left) Boundary layer length as a fraction of total overlap length, shown as a function of bulk motor concentration and motor speed. (Right) Boundary layer length as a function of overlap length for the reference parameter set and varying bulk motor concentration. The boundary layer length is the distance at each end of the overlap where significant motor accumulation occurs and is determined by Eq. S30. To see this figure in color, go online.](image2)
accumulation at the overlap ends depends on overlap length. Therefore, length regulation of antiparallel MT overlaps must occur differently. In future work, it will be of interest to understand how the motor density profiles on fixed-length overlaps can be used to understand the length regulation of dynamic overlaps.

SUPPORTING MATERIAL

Supporting Materials and Methods, ten figures, and one table are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(16)30153-9.

AUTHOR CONTRIBUTIONS

H.-S.K. and M.D.B. designed research, performed research, contributed analytic tools, analyzed data, and wrote the article.

ACKNOWLEDGMENTS

We thank Robert Blackwell, Matthew Glaser, and Loren Hough for useful discussions.

This work was supported by National Science Foundation grant No. DMR-0847685 and National Institutes of Health No. K25 GM110486 to M.D.B., a fellowship to H.-S.K. provided by matching funds from the National Institutes of Health/University of Colorado Biophysics Training Program, and facilities of the Soft Materials Research Center under the National Science Foundation Materials Research Science and Engineering Centers grants No. DMR-0820579 and No. DMR-1420736. This work used code from the Danuser and Jaqaman labs, provided through the Computational Image Analysis in Cellular and Developmental Biology course at MBL, and funded by National Institutes of Health No. R25 GM103792.

REFERENCES

1. Bray, D. 2000. Cell Movements: from Molecules to Motility. Routledge, New York.
2. Kolomeisky, A. B. 2015. Motor Proteins and Molecular Motors. CRC Press, Boca Raton, FL.
3. Chowdhury, D. 2013. Modeling stochastic kinetics of molecular machines at multiple levels: from molecules to modules. *Biophys. J.* 104:2331–2341.
4. Helbing, D. 2001. Traffic and related self-driven many-particle systems. *Rev. Mod. Phys.* 73:1067–1141.
5. Parmeggiani, A., T. Franosch, and E. Frey. 2004. Totally asymmetric simple exclusion process with Langmuir kinetics. *Phys. Rev. E Stat. Nonlin. Soft Matter Phys.* 70:046101.
6. Leduc, C., K. Padberg-Gehle, …, J. Howard. 2012. Molecular crowding creates traffic jams of kinesin motors on microtubules. *Proc. Natl. Acad. Sci. USA.* 109:6100–6105.
7. Varga, V., C. Leduc, …, J. Howard. 2009. Kinesin-8 motors act cooperatively to mediate length-dependent microtubule depolymerization. *Cell.* 138:1174–1183.
8. Biegler, P. I. A. Telley, and T. Surrey. 2010. A minimal midzone protein module controls formation and length of antiparallel microtubule overlaps. *Cell.* 142:420–432.
9. Goshima, G., R. Wollman, …, R. D. Vale. 2005. Length control of the metaphase spindle. *Curr. Biol.* 15:1979–1988.
10. Walczak, C. E., T. J. Mitchison, and A. Desai. 1996. XKCM1: a *Xenopus* kinesin-related protein that regulates microtubule dynamics during mitotic spindle assembly. *Cell.* 84:37–47.
11. Rivero, F. B. Köppel, …, A. A. Noegel. 1996. The role of the cortical cytoskeleton: F-actin crosslinking proteins protect against osmotic stress, ensure cell size, cell shape and motility, and contribute to phagocytosis and development. *J. Cell Sci.* 109:2679–2691.
12. Revenu, C., R. Athman, …, D. Louvard. 2004. The co-workers of actin filaments: from cell structures to signals. *Nat. Rev. Mol. Cell Biol.* 5:635–646.
13. Dogterom, M., and S. Leibler. 1993. Physical aspects of the growth and regulation of microtubule structures. *Phys. Rev. Lett.* 70:1347–1350.
14. Drummond, D. R. 2011. Regulation of microtubule dynamics by kinesins. *Semin. Cell Dev. Biol.* 22:927–934.
15. Gupta, M. L., Jr., P. Carvalho, …, D. Pellman. 2006. Plus end-specific dynein activity of KiP3, a kinesin-8 protein, explains its role in positioning the yeast mitotic spindle. *Nat. Cell Biol.* 8:913–923.
16. Varga, V., J. Helenius, …, J. Howard. 2006. Yeast kinesin-8 depolymerizes microtubules in a length-dependent manner. *Nat. Cell Biol.* 8:957–962.
17. Hough, L. E., A. Schwabe, …, M. D. Betterton. 2009. Microtubule depolymerization by the Kinesin-8 motor KiP3p: a mathematical model. *Biophys. J.* 96:3050–3064.
18. Reese, L., A. Melbinger, and E. Frey. 2011. Crowding of molecular motors determines microtubule depolymerization. *Biophys. J.* 101:2190–2200.
19. Govindan, B. S., M. Gopalakrishnan, and D. Chowdhury. 2008. Length control of microtubules by depolymerizing motor proteins. *Europhys. Lett.* 83:40006.
20. Johann, D., C. Erlenkämper, and K. Kruse. 2012. Length regulation of active biopolymers by molecular motors. *Phys. Rev. Lett.* 108:258103.
21. Melbinger, A., L. Reese, and E. Frey. 2012. Microtubule length regulation by molecular motors. *Phys. Rev. Lett.* 108:258104.
22. Reese, L., A. Melbinger, and E. Frey. 2014. Molecular mechanisms for microtubule length regulation by kinesin-8 and XMAP215 proteins. *Interface Focus.* 4:20140031.
23. Tischer, C., P. R. Ten Wolde, and M. Dogterom. 2010. Providing positional information with active transport on dynamic microtubules. *Biophys. J.* 99:726–735.
24. Kuan, H.-S., and M. D. Betterton. 2013. Biophysics of filament length regulation by molecular motors. *Phys. Biol.* 10:036004.
25. Glunčić, M., N. Magelli, …, I. M. Tolić. 2015. Kinesin-8 motors improve nuclear centering by promoting microtubule catastrophe. *Phys. Rev. Lett.* 114:078103.
26. Kurasawa, Y., W. C. Earnshaw, …, K. Kodokoro. 2004. Essential roles of KIF4 and its binding partner PRC1 in organized central spindle midzone formation. *EMBO J.* 23:3237–3248.
27. Zhu, C., and W. Jiang. 2005. Cell cycle-dependent translocation of PRC1 on the spindle by Kif4 is essential for midzone formation and cytokinesis. *Proc. Natl. Acad. Sci. USA.* 102:343–348.
28. Khmelinskii, A., C. Lawrence, …, E. Schiebel. 2007. Cdc14-regulated midzone assembly controls anaphase B. *J. Cell Biol.* 177:981–993.
29. Fededa, J. P., and D. W. Gerlich. 2012. Molecular control of animal cell cytokinesis. *Nat. Cell Biol.* 14:440–447.
30. Subramanian, R., S.-C. Ti, …, T. M. Kapoor. 2013. Marking and measuring single microtubules by PRC1 and kinesin-4. *Cell.* 154:377–390.
31. Subramanian, R., E. M. Wilson-Kubalek, …, T. M. Kapoor. 2010. Insights into antiparallel microtubule crosslinking by PRC1, a conserved nonmotor microtubule binding protein. *Cell.* 142:433–443.
32. Bringmann, H., G. Skiniotis, …, T. Surrey. 2004. A kinesin-like motor inhibits microtubule dynamic instability. *Science.* 303:1519–1522.
33. Castoldi, M., and I. Vernos. 2006. Chromokinesin Xklp1 contributes to the regulation of microtubule density and organization during spindle assembly. *Mol. Biol. Cell.* 17:1451–1460.
34. Parmeggiani, A., T. Franosch, and E. Frey. 2003. Phase coexistence in driven one-dimensional transport. *Phys. Rev. Lett.* 90:086601.
35. Juhasz, R. 2007. Weakly coupled, antiparallel, totally asymmetric simple exclusion processes. *Phys. Rev. E Stat. Nonlin. Soft Matter Phys.* 76:021117.

36. Ashwin, P., C. Lin, and G. Steinberg. 2010. Queueing induced by bidirectional motor motion near the end of a microtubule. *Phys. Rev. E Stat. Nonlin. Soft Matter Phys.* 82:051907.

37. Braun, M., Z. Lansky, ..., M. E. Janson. 2011. Adaptive braking by Ase1 prevents overlapping microtubules from sliding completely apart. *Nat. Cell Biol.* 13:1259–1264.

38. Lansky, Z., M. Braun, ..., S. Diez. 2015. Diffusible crosslinkers generate directed forces in microtubule networks. *Cell.* 160:1159–1168.

39. Kuan, H.-S., and M. D. Betterton. 2016. Phase-plane analysis of the totally asymmetric simple exclusion process with binding kinetics and switching between antiparallel lanes. *ArXiv:1604.00690 [physics.bio-ph]* 7 March 2016.

40. Pierobon, P., M. Mobilia, ..., E. Frey. 2006. Bottleneck-induced transitions in a minimal model for intracellular transport. *Phys. Rev. E Stat. Nonlin. Soft Matter Phys.* 74:031906.

41. Yildiz, A., J. N. Forkey, ..., P. R. Selvin. 2003. Myosin V walks hand-over-hand: single fluorophore imaging with 1.5-nm localization. *Science.* 300:2061–2065.

42. Bieling, P., I. Kronja, and T. Surrey. 2010. Microtubule motility on reconstituted meiotic chromatin. *Curr. Biol.* 20:763–769.