

Quantitative Determination of the Probability of Multiple-Motor Transport in Bead-Based Assays

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ABSTRACT With their longest dimension typically being less than 100 nm, molecular motors are significantly below the optical-resolution limit. Despite substantial advances in fluorescence-based imaging methodologies, labeling with beads remains critical for optical-trapping-based investigations of molecular motors. A key experimental challenge in bead-based assays is that the number of motors on a bead is not well defined. Particularly for single-molecule investigations, the probability of single-versus multiple-motor events has not been experimentally investigated. Here, we used bead travel distance as an indicator of multiple-motor transport and determined the lower-bound probability of bead transport by two or more motors. We limited the ATP concentration to increase our detection sensitivity for multiple- versus single-kinesin transport. Surprisingly, for all but the lowest motor number examined, our measurements exceeded estimations of a previous model by ≥ 2 -fold. To bridge this apparent gap between theory and experiment, we derived a closed-form expression for the probability of bead transport by multiple motors, and constrained the only free parameter in this model using our experimental measurements. Our data indicate that kinesin extends to ~57 nm during bead transport, suggesting that kinesin exploits its conformational flexibility to interact with microtubules at highly curved interfaces such as those present for vesicle transport in cells. To our knowledge, our findings provide the first experimental approach utilized here (limiting ATP concentration) may be generally applicable to studies in which molecular motors are labeled with cargos that are artificial or are purified from cellular extracts.

INTRODUCTION

Microtubule-based molecular motors are protein machines that drive long-range mechanical transport in cells (1). This transport process is critical for cellular function and survival. A quantitative understanding of molecular-motor function at the single-molecule level is crucial for deciphering the complexity of mechanical transport in cells and for engineering biomimetic machineries on the nano/microscale. Optical trapping has been instrumental in elucidating the mechanochemical functions of molecular motors, particularly at the single-molecule level (2–4).

Perhaps paradoxically, the single-molecule range in optical trapping experiments has yet to be defined in quantitative terms. Uncertainty arises because optical trapping studies use dielectric beads to label molecular motors. Each bead is typically decorated by a Poisson-distributed number of motors, rather than by a well-defined number of motors. The three-dimensional nature of the bead further

Submitted September 21, 2015, and accepted for publication May 10, 2016. *Correspondence: jxu8@ucmerced.edu Editor: David Warshaw. http://dx.doi.org/10.1016/j.bpj.2016.05.015

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complicates the problem, since not all motors on the bead can reach the microtubule at the same time. Although a reduction in the ratio of motors to beads can lower the probability of cargo transport by two or more motors, this reduction also limits the fraction of motor/bead complexes that are capable of interacting with the microtubule, and in turn significantly reduces the experimental throughput. Although force measurements can shed light on the number of motors engaged in transport, work by Furuta et al. (5) revealed that, depending on the motor type, the force generated by multiple motors is not necessarily sensitive to the motor number. Previously, Beeg et al. (6) determined the average number of simultaneously engaged motors on a bead for a range of motor concentrations, making it possible to evaluate the probability of multiple-motor events measured under the same experimental conditions. However, Beeg et al. (6) used a substantially smaller bead size (100 nm diameter) than that optimized for optical trapping (~500 nm diameter (4). Because a fivefold increase in bead size can impact both the density of motors on the bead surface and the fraction of bead surface available to the microtubule, it is nontrivial



to translate the experimental conditions from (6) to those suitable for optical trapping.

To date, only one theory model has provided a general guide for estimating the probability of multiple-motor transport at a given motor number in bead-based assays (3). However, the assumptions of this previous model have not been experimentally verified. Importantly, the motors are assumed to be fully extended and their motor domains effectively in contact with each other (Fig. S1 A in the Supporting Material). There is increasing consensus that motors should be able to bind different locations along the length of the microtubule (7,8) (Fig. S1 B). However, this updated geometry does not readily lend itself to theory predictions because only limited information is available regarding a key parameter in this updated geometry (i.e., how far the motor can extend during bead transport; Fig. S1 B). As a result, predictions from the previous model (3) have remained the only guide for estimating the probability of multiple-motor transport in optical trapping studies (9–13).

Here, we sought to develop a quantitative guide to understanding the probability of multiple-motor events in the dilute motor range. We focused on experimental details that are typically used in optical trapping studies (500-nmdiameter spherical beads and randomly distributed motors on the bead surface) and carried out our experiments using the major microtubule-based motor, kinesin-1 (conventional kinesin). We used the motile fraction (the probability of beads exhibiting motility along microtubules) as an experimental readout for the average number of active motors on the bead (2,3) (Supporting Material, Text 1; Fig. S2). We used the bead travel distance to identify bead transport by multiple kinesins for a range of motile fractions, and used a limiting ATP concentration to increase our detection sensitivity for these multiple-motor events (14–18). Surprisingly, for all but the lowest motile fractions examined, we detected substantially higher probabilities of multiple-motor transport than that predicted by the previous theory model (3). We therefore applied both theory and simulation to unravel this discrepancy.

MATERIALS AND METHODS

Proteins and reagents

Bovine brain tubulin was purified over a phosphocellulose column as previously described (19). Conventional kinesin was purified from bovine brain as previously described (20), except that 9S kinesin was eluted from the Mono-Q resin using customized salt gradients to separate kinesin from other polypeptides in the 9S sucrose fractions (21). Chemicals were purchased from Sigma Aldrich (St. Louis, MO).

Microtubule preparation

To assemble taxol-stabilized microtubules, purified tubulin (40 μ M) was supplemented with 0.5 mM GTP and incubated for 20 min at 37°C. Assembled microtubules were mixed with an equal volume of PM buffer (100 mM PIPES, 1 mM MgSO₄, 2 mM EGTA, pH 6.9) supplemented with 40 μ M taxol and incubated for 20 min at 37°C. Microtubules were then kept at room temperature in a dark box and used within 4 days of preparation.

Motor/bead complex preparation

Kinesin was incubated with carboxylated polystyrene beads (500 nm diameter; Polysciences, Warrington, PA) in motility buffer (67 mM PIPES, 50 mM CH₃CO₂K, 3 mM MgSO₄, 1 mM dithiothreitol, 0.84 mM EGTA, 10 μ M taxol, pH 6.9) for 10 min at room temperature. Before motility measurements, this solution was supplemented with an oxygen-scavenging solution (250 μ g/mL glucose oxidase, 30 μ g/mL catalase, 4.6 mg/mL glucose) and ATP (1 mM or 0.01 mM, as indicated). The bead concentration was kept constant at 3.6 × 10⁵ particles/ μ L and the concentration of kinesin was varied to give rise to a range of motile fractions (Fig. S2).

In vitro optical trapping

Optical trapping was carried out in flow cells, imaged via differential interference microscopy, and video-recorded at 30 Hz as previously described (16). For all studies presented here, we limited the trap power to <20 mW (at fiber output), such that the trap positioned individual beads but was not sufficient to stall beads carried by a single kinesin (stall force \sim 4.5 pN (22)). To measure the motile fraction, we used the optical trap to position individual beads in the vicinity of the microtubule; a motile event was scored if and only if the bead demonstrated directed motion away from the center of the trap. We observed occasional events in which the bead appeared to bind (as indicated by reduced Brownian motion of the bead) but did not move processively away from the trap center (data not shown). We did not score these events for our motile fraction measurements, as they constituted neither motile events nor nonmotile events. The beads in these events typically detached shortly after we turned off the optical trap, without demonstrating any clear movement along the microtubule (data not shown). For motile beads, upon observation of directed bead motion along the microtubule, we turned off the optical trap to enable cargo transport in the absence of external load.

Data analysis

Video recordings of bead motion were particle-tracked to 10 nm resolution (1/3 pixel) using a template-matching algorithm as previously described (16,23). The travel distance for each bead was determined as the net displacement of the bead along the microtubule axis upon the bead's binding to and then detaching from the microtubule. The distribution of travel distances for each experimental condition (motile fraction and ATP concentration) was fitted to a single-exponential decay (2). The mean travel distance and the associated standard error of the mean for each distribution were determined from the fitted decay constant and uncertainty, respectively. To account for the human reaction time to manually shut off the optical trap, only trajectories > 0.3 μ m were used to determine the distribution of travel distances at each motile fraction.

Best fitting of motile fraction measurements to a one-motor Poisson curve was carried out in OriginPro9.1 (OriginLab, Northampton, MA). We used least- χ^2 fitting to constrain our theory model using our experimental measurements via a custom routine in MATLAB (The MathWorks, Natick, MA).

Simulation

Our simulation randomly distributes a mean number of active motors (n) on a 51 \times 51 lattice with periodic boundaries (a torus lattice), identifies the location of a lattice site that is occupied by one motor, and evaluates the number of motors in a closed patch surrounding this particular lattice site. The size of the closed path is determined as the area of the square

lattice multiplied by α , the probability that any two randomly distributed motors on the bead will be in close enough proximity to be in simultaneous reach of the microtubule. We repeated the simulation 10,000 times to determine the probability of counting two or more motors for each set of *n*- and α -values tested. We evaluated the motile fraction corresponding to each *n* using the following relationship: motile fraction = $1 - e^{-n}$ (Fig. S2) (2,3).

Statistical analysis

Standard errors for binomially distributed data were calculated as $\sqrt{P(1-P)/N}$, where *P* is the measured fraction and *N* is the sample size. We did not observe a substantial difference between this error calculation and that determined using a 68.3% confidence interval for a binomial distribution (data not shown). In case of a zero-fraction measurement (at

motile fraction = 0.2; Fig. 1 *B*), the standard error was estimated as the upper level of the 68.3% confidence interval for a binomial distribution.

RESULTS

Lower-bound measurements of the probability of multiple-motor transport

We used the bead travel distance to identify multiple-motor events versus single-motor events. The average travel distance for a single kinesin is ~1 μ m (2,24). When a bead is transported by a single kinesin, there is a <0.1% probability that the travel distance will be >6.9 μ m (Supporting Material, Text 2). Thus, when we observed a motile event longer



FIGURE 1 Measurements of bead travel distances for different motile fractions. (*A*) Distribution of travel distances for three motile fractions measured at 1 mM ATP (*top*) and 0.01 mM ATP (*bottom*). Solid line, best fit to a single exponential decay for travel distances of $\leq 6.9 \ \mu\text{m}$. Blue bar, cumulative counts of events with travel distance of >6.9 $\ \mu\text{m}$. D (\pm standard error), fitted mean travel distance for measurements shorter than our travel threshold; $F_{>6.9\ \mu\text{m}}$ (\pm standard error), fraction of motile events longer than our travel threshold; *N*, sample size. (*B*) Fraction of motile beads with travel distance of >6.9 $\ \mu\text{m}$, measured at 0.01 mM ATP (*blue scatter*). Error bars, standard error. Magenta line, predicted fraction of motile events transported by two or more motors, based on a previous theory (3) (see Supporting Material, Text 3). To see this figure in color, go online.

than our travel threshold, we excluded it from the set of single-motor events with high confidence and identified it as a multiple-motor event. Since not all multiple-motor events exceed this 6.9 μ m threshold, our measurements of long-travel events represent lower-bound values for the probability of multiple-motor transport.

We used ATP as an experimental handle to increase the likelihood that a multiple-motor event would travel >6.9 μ m (*blue bars*, Fig. 1 A). Previous studies demonstrated that the distance traveled by multiple kinesins is inversely tuned by the ATP concentration (0.01–1 mM) (14–18). Importantly, the single-kinesin travel distance remains constant over the same ATP range (15,24). We thus anticipated that a limiting ATP concentration (0.01 mM) would increase our detection sensitivity for multiple-motor events.

We varied the input kinesin concentration to experimentally access a range of motile fractions (or motor numbers) (Fig. S2 A). For each motile fraction, we measured 50–150 motile trajectories and determined the corresponding distribution of travel distances (Fig. 1 A). For both ATP concentrations examined, we found that the fraction of long-travel events increased with increasing motile fraction (*blue bars*, Fig. 1 A). At the lowest motile fraction tested (0.2), we did not observe any long-travel events for either ATP concentration. At the highest motile fraction tested (0.8), the fraction of long-travel events was substantially larger than zero for both 0.01 mM and 1 mM ATP. This observation is consistent with the increased presence of motors on the bead for the higher motile fractions (Fig. S2 B).

Importantly, for the higher motile fractions (>0.2), we observed substantially higher fractions of long-travel events at the lower ATP concentration ($F_{>6.9\mu m}$; Fig. 1 A). For example, at a motile fraction of 0.8, the fraction of longtravel events was 23% at 0.01 mM ATP, ~4-fold larger than the 6% at 1 mM ATP. Since we used the same trap power for measurements at both ATP concentrations, the increase in the fraction of long-travel events was due to the change in ATP concentration, not experiment artifacts from optical trapping. Further, for measurements below our travel threshold (gray shading, Fig. 1 A), the mean travel distances measured at 1 mM ATP approximately doubled as the motile fraction increased from 0.2 to 0.8, whereas the mean travel distances measured at 0.01 mM ATP remained approximately constant at the single-kinesin level $(\sim 1 \ \mu m \ (2,24))$ over the same motile fraction range. This observation indicates that only the travel distance, but not the likelihood, of multiple-motor events was amplified by the lower ATP concentration. Given the increase in multiple-motor travel distance at 0.01 mM ATP, multiple-motor events are less likely to influence the distribution of travel distances shorter than 6.9 μ m. Together, these data demonstrate that our approach of limiting ATP is effective in uncovering the presence of multiple-motor transport in bead-based assays.

We measured the fraction of long-travel events for a range of motile fractions at 0.01 mM ATP (*blue scatters*, Fig. 1 *B*). We detected a substantial difference between these lowerbound measurements and the predictions of a previous model (3) (Supporting Material, Text 3; Fig. 1 *B*, *blue scatters* versus *magenta line*). For motile fractions > 0.4, our lower-bound measurements were substantially larger than predictions from the previous model. For example, at a motile fraction of 0.5, our lower-bound measurements indicate that at least 8% of motile trajectories are transported by multiple motors, which is more than twice the 3% predicted previously.

We examined the possibility that our measurements were influenced by application of the optical trap's force to the motors. To measure the bead travel distance, we used an optical trap to initially confine individual beads to the vicinity of a microtubule. Previous studies reported that the force exerted by an optical trap can promote synergistic cooperation between multiple kinesins (25,26). It is therefore possible that our use of an optical trap may have biased the multiple-motor travel distance toward higher values by influencing how many motors were engaged in transport at the start of a multiple-motor event (before the trap was turned off). We speculate that the magnitude of this effect in our experiments is small, since the trap's effect is more pronounced when the trap's force is larger than or comparable to the force produced by a single kinesin (25,26). In this study, we used a trap with substantially lower power. Beads carried by approximately one motor moved processively away from the trap center without detaching prematurely (for example, motile fraction = 0.2; Fig. 1 A), which is difficult to achieve when the trap's force is comparable to the motor's force (26-28). Importantly, regardless of the magnitude of the trap's effect on multiple-motor travel distance, the trap's force cannot increase the distance traveled by a single motor under any circumstance (26–28). Thus, while the trap's force may improve the sensitivity for detecting a multiple-motor event, it cannot lead to false identification of a single-motor event as a multiple-motor event.

To understand the discrepancy between our measurements and the previous model (Fig. 1 B), we next derived a general theory model for the probability of transport by multiple motors (Eqs. 1 and 2; Fig. 2). We then used our lower-bound measurements to constrain the free parameter in our model and to gain insight into the geometry of multiple-motor transport in bead-based assays (Fig. 3).

An exact theory expression for the probability of multiple-motor transport

Using the same framework as developed in the previous model (3), we described the probability of bead transport by two or more motors as the weighted sum $P(\ge 2) = \sum_{k=2}^{\infty} p(k | n) \cdot g(2 | k)$, where *n* is the average number of motors on the bead, p(k | n) is the Poisson probability that there



FIGURE 2 Probability that a bead is transported by two or more motors, $P(\ge 2)$, evaluated for three values of α (the probability that two randomly attached motors are within simultaneous reach of the microtubule). Error bars in the simulation results indicate standard error (sample size N = 10,000 per simulation condition). To see this figure in color, go online.

are exactly k motors on the bead, and g(2 | k) is the probability that at least two of the k motors on the bead are available for transport. Whereas the Poisson probability $p(k | n) = n^k e^{-n}/k!$ only concerns the total number of motors on the bead and does not differ between experiments, the weighting factor g(2 | k) is sensitive to experimental details (bead size) and assumptions of multiple-motor transport geometry (Fig. S1). As detailed below, the previous model (*magenta line*, Fig. 1 B) used an approximation for this weighting factor. We hypothesized that incorporating the exact expression of g(2 | k) would help bridge the gap between experiment and theory in Fig. 1 B.

To arrive at an exact expression for g(2 | k), we used α to denote the probability that any two randomly attached motors are within simultaneous reach of the microtubule. The value of α is important because it critically impacts the probability of multiple-motor transport for a given motor number *n*. Intuitively, the larger the value of α , the more likely it is that two or more motors will engage in transport at a given motor number. The evaluation of α is difficult, however, because it depends sensitively on bead size as well as assumptions of two-motor transport geometry (Fig. S1). The experimental measurements in this study (Fig. 1 *B*) provided the first opportunity, to our knowledge, to constrain the value of α for kinesin without any geometry assumptions.

The exact expression for the weighting factor is $g(2|k) = 1 - (1 - \alpha)^{k-1}$. For any one motor present on the bead, the probability that none of the remaining k - 1 motors will be within simultaneous reach of the microtubule is $(1 - \alpha)^{k-1}$. This exact expression differs from the approximated form used in a previous model, $g(2|k) = \alpha$ (3). The previous approximation $(g(2|k) = \alpha)$ underestimates the weighting factor for all k > 2. For example, when there are 20 motors on the bead, the probability that two or more motors are within simultaneous reach of the microtubule is larger than that obtained when only two motors are present on the bead.

Using $g(2|k) = 1 - (1 - \alpha)^{k-1}$, we arrived at a closed-form description for the probability of multiple-motor



FIGURE 3 Quantitative comparison between experiment and theory. (A) Lower-bound probability of a bead transported by two or more motors. Scatter, experimental measurements. Error bars, standard error. Solid line and shaded area, least- χ^2 fit and standard error, respectively, for Eq. 2. (B) Updated geometry for two-motor transport (not to scale). $h_{\text{best-fit}}$ extension length of the kinesin motor, corresponding to the best-fit α -value determined in (A). To see this figure in color, go online.

transport (for a detailed derivation, see Supporting Material, Text 4):

$$P(\geq 2) = 1 + e^{-n} \left(\frac{\alpha}{1-\alpha}\right) - \frac{e^{-n\alpha}}{1-\alpha},$$
 (1)

where *n* indicates the average number of active motors present on the bead. Since we absorbed geometry considerations into the probability α , this closed-form expression (Eq. 1) is general and does not depend on details of multiple-motor geometry (for example, Fig. S1 A versus S1 B). We then recast the dependence on the mean motor number (*n*) as that on the experimental measurable, motile fraction, using the following relationship: motile fraction = $1 - e^{-n}$ (Fig. S2) (2,3).

We used a numerical simulation to test the validity of Eq. 1 (Fig. 2). Our numerical simulation captured the three-dimensional nature of the cargo via the use of a torus lattice, and we also included one free parameter to reflect the probability α (Materials and Methods). The predictions from Eq. 1 were in excellent agreement with the results of our numerical simulations for all α -values tested (black line versus scatter, Fig. 2). On the other hand, when we approximated the probability of multiple-motor transport using $g(2 | k) = \alpha$ as in the previous model, we consistently obtained a lower value than that returned by our numerical simulation (magenta line versus black scatter, Fig. 2). The extent of underestimation was more pronounced for small values of α and diminished as α approached one. This scenario is expected, since in the limit of $\alpha = 1$, both the exact and approximate forms of the weighting factor also converge to one. These data support our hypothesis that the underestimation in g(2 | k) contributes to the gap between experiment and theory in Fig. 1 B.

Interestingly, correcting g(2 | k) alone was not sufficient to bridge the gap between experiment and theory. When evaluated using the previously estimated value for α (0.099), the probability of multiple-motor transport (as determined by Eq. 1 and simulation; *black line* and *scatters*, Fig. 2) remained substantially smaller than our lower-bound measurements (blue scatters, Fig. 1 B). For example, at a motile fraction of 0.8, we anticipate that $\sim 10\%$ of the motile events are transported by two or more motors ($\alpha = 0.1$, Fig. 2). This value is less than half of the 23% observed experimentally (Fig. 1). Thus, our theory study indicates that the value of α is substantially higher than previously anticipated. This finding is perhaps not surprising, since the geometry of two-motor transport used in the previous estimation was itself an approximation (Fig. S1 A). Importantly, in the updated geometry for two-motor transport, the value of α becomes larger than its previous estimated value (0.099) when kinesin extends beyond 16% of its contour length (80 nm (29,30) (Fig. S1 B). Since kinesin has been found to extend to at least 28% of its contour length during active transport (31), it is conceivable that the α -value may be larger than previously estimated.

Quantitative comparison between experiment and theory

To enable a direct comparison between theory and experiments, we recast Eq. 1 to reflect the lower-bound nature of our experimental measurements. Our experimental measurements represent lower-bound values, since our distance threshold excludes the population of multiple-motor transport events that travel $\leq 6.9 \ \mu m$ (Fig. 1). Imposing the same threshold, we derived (Supporting Material, Text 5) the expected lower-bound probability of multiple-motor transport as

$$P_{lower-bound}(\geq 2) = P(\geq 2) - f \cdot \left(\frac{\alpha \cdot e^{-n\alpha}}{(1-\alpha)^2} - \frac{\alpha \cdot n \cdot e^{-n}}{1-\alpha} - \frac{\alpha \cdot e^{-n}}{(1-\alpha)^2}\right),$$

$$(2)$$

where $P(\ge 2)$ is as described in Eq. 1 and $f = 0.556 \pm 0.096$, as we previously measured for the experimental condition used in this investigation (0.01 mM ATP) (15). Similarly to Eq. 1, this description is a function of the experimental measureable motile fraction and the free parameter, α .

Constraining Eq. 2 using our experimental measurements, we obtained a best-fit value of $\alpha = 0.405$ (Fig. 3 *A*). Note that since our theory expressions describe the probability of multiple-motor transport for all beads (including those without active motors), we scaled our measurements in Fig. 1 *B* by their associated motile fractions to obtain the lower-bound measurements on the same probability (*blue scatters*, Fig. 3 *A*). Since this best-fit value of α is 4-fold larger than that estimated using the previous model (0.099; Supporting Material, Text 3), our data indicate that the geometry assumed in the previous model (Fig. S1 A) is unlikely to occur during bead transport.

Using an updated two-motor geometry (7,8) (Fig. S1 B) and our best-fit value of $\alpha = 0.405$ (Fig. 3 A), we arrived at a mean extension length of 57 nm for kinesin (Fig. 3 B). This mean extension length and its associated upper and lower limits (79 nm and 40 nm, respectively) are reasonable as they are within kinesin's contour length (80 nm (29,30)). These data support the updated geometry and address the question of kinesin's extension length for the typical bead size used in optical trapping (500 nm diameter).

Estimated fraction of motile beads transported by multiple motors

Based on our findings (Fig. 3 and Eq. 1), we evaluated the probability that a motile bead is transported by two or more motors (Fig. 4). To do so, we normalized our



FIGURE 4 Fraction of motile beads transported by multiple motors as a function of motile fraction, estimated for two major microtubule-based molecular motors. Magenta lines, a previous model (3). Black lines and gray area, this study. α , the probability that two randomly attached motors are within simultaneous reach of the microtubule. h, the extension length of the motor while bound to a bead. Left: solid black line and gray area, mean ± standard error based on our experimentally constrained α -value for kinesin (Fig. 3). Right: dashed black line and solid black line, minimum and maximum values based on the updated two-motor geometry (Fig. S1 B) and the motor's extension length, h, supported by previous structural studies of dynein (32-34). To see this figure in color, go online.

expression in Eq. 1 by the associated motile fraction. Since only motile events can contribute to transport measurements, the normalized probability directly reflects the impact of multiple-motor events on the measured motility parameters.

Using the best-fit α -value for kinesin (Fig. 3), we evaluated the fraction of motile beads transported by multiple kinesins (Fig. 4, *left*). For all motile fractions measured, our experimentally constrained estimations were >4-fold larger than previous estimations (*black line* versus *magenta line*, Fig. 4, *left*). We estimate that for motile fractions below 0.387, \geq 90% of motile events are due to the action of a single kinesin (Fig. 4, *left*).

Using the updated two-motor geometry supported by our study (Fig. 3 *B*), we extended our evaluation to another major microtubule-based motor, dynein (Fig. 4, *right*). Here, we could not constrain the extension length of dynein as we did for kinesin, since there is currently no experimental handle to amplify the difference between single- and multiple-dynein travel distances. Instead, we referred to previous structural studies (32-34) to estimate the range for dynein's motor extension length. This approach is reasonable because the flexibility of dynein is expected to be substantially more limited than that of kinesin (1).

We used the size of dynein's motor domain to place a lower bound on its extension length (26 nm (34)) and obtained a minimum α -value of 0.197 (Fig. S1 B). This lower limit may be more relevant for studies that use minimal dynein constructs containing only the motor domains. Estimations using this lower limit remained >5-fold larger than previous predictions for dynein (*black dashed line* versus *magenta line*, Fig. 4, *right*). We estimate that the motile fraction can be as high as 0.609 while still ensuring that <10% of the motile events are transported by multiple dyneins (*dashed black line*, Fig. 4, *right*).

We used dynein's contour length as the maximum extension length (50 nm) (32,33), corresponding to a maximum α -value of 0.36 (Fig. S1 B). As expected, the probability of multiple-motor transport increased at the higher α -value. Using this upper-bound value, we estimate that <10% of the motile measurements reflect transport by multiple dyneins for motile fractions below 0.420 (*solid black line*, Fig. 4, *right*).

DISCUSSION

Here, we combined experiments and theory to determine the probability of multiple-motor transport in bead-based assays. In the experimental portion of our study, we increased our detection sensitivity for multiple- versus single-kinesin transport using the ATP concentration as an experimental handle to tune the multiple-kinesin travel distance (Fig. 1). We verified that the increased likelihood of long-travel events was directly associated with lower ATP levels (Fig. 1 A). In our theory work, we derived an exact and closed-form expression for the probability of multiple-motor transport that contains only one free parameter, α (Eq. 1). Our numerical simulation validated the predictions of our expression for a range of α -values (Fig. 2). We then recast our expression to reflect the lower-bound nature of our experimental measurements (Eq. 2). We constrained the one free parameter, α , in our theory model using our experimental measurements (Fig. 3A). The resulting predictions constitute a set of quantitative guides for estimating the probability of multiple-motor contributions in single-molecule investigations using optical trapping (Fig. 4). To our knowledge, this is the first set of such guides that has been experimentally constrained.

In this study, we focused on experimental details used in typical optical trapping studies; specifically, individual motors were uniformly distributed atop spherical beads (500 nm diameter). Under these conditions, we found that it was not necessary to limit single-molecule investigations to motile fractions substantially below 0.2 (such as 0.07 in (9)), where <5% of motile events were due to multiple motors for both kinesin and dynein (Fig. 4). Our study also supports the use of a motile fraction of 0.3 for single-molecule investigations (10–13), since we estimated that

 $7\% \pm 2\%$ motile events were due to multiple kinesins (<7% for multiple dyneins; Fig. 4). Depending on the expected cooperativity of motors functioning in groups and specific requirements for measurement precision, experiments may be carried out at somewhat higher motile fractions to boost experimental throughput.

In general, the results shown in Fig. 4 apply only to studies in which the motors are randomly distributed on the bead surface, since we assumed a single parameter, α , across the bead surface. Despite this restriction, our model is appropriate for studies that use intermediate binding sites (antibodies (5,15) and/or DNA scaffolds (5,11)) to group motors together locally on the bead surface. In these special cases, the intermediate binding sites are randomly distributed atop the bead surface. Motors bound to individual intermediate sites may be considered to constitute a single "supermotor" complex, and evaluation of Eq. 1 yields the probability that a bead is transported by multiple supermotor complexes (11,15).

Further studies are necessary to understand how changing the bead size would impact the results shown in Fig. 4. Our study supports an updated geometry for two-motor transport in which the motors bind at different locations along the length of the microtubule (Fig. 3). In this updated geometry, the α -value depends sensitively on both the bead size and the motor extension length (Fig. S1 B). For a given extension length, the larger the bead, the less likely it is that two randomly attached motors will be within simultaneous reach of the same microtubule, thus reducing α . Therefore, if the motor is relatively rigid (as is likely the case for dynein (1)), an increase in bead size (for example, to the ~1 μ m diameter used in some optical trapping studies) has the potential to decrease the probability of multiple-motor transport at a particular motile fraction. For flexible kinesin, however, the impact of bead size on the α -value will depend on how the motor's extension length varies with bead size. We are currently investigating the effect(s) of bead size on kinesin's extension length.

For 500-nm-diameter beads, our data indicate that kinesin extends to 57 nm (~71% of its contour length) in beadbased assays (Fig. 3 B). This finding supports the general assumption that kinesin is in an extended conformation during bead transport (for example, see (3)). However, an extended conformation for kinesin is also surprising, since kinesin was previously found to adopt a more compact conformation in microtubule-gliding assays, extending to only $\sim 28\%$ of its contour length (31). The key difference between bead-based assays and microtubule-gliding assays may be that the interface between microtubules and the cargo surface is highly curved in the former but flat in the latter. Our study raises the intriguing possibility that kinesin can exploit its conformational flexibility to seek and interact with microtubules at highly curved interfaces, such as those that occur during vesicle transport in cells.

The experimental approach used here harnesses the increased kinesin/microtubule association time at lower ATP concentrations (14,15). Thus, this approach is not limited to a specific cargo geometry, motor/cargo recruitment method, or measurement approach (optical trapping or fluorescence). It is, however, specific to studies of kinesin, for which the inverse correlation between multiplemotor travel and ATP level has been directly demonstrated (14–18). Future investigations expanding our ability to tune multiple-motor travel distance for other molecular motors (such as dynein) will critically empower this experimental approach. Despite this limitation, when combined with selective small-molecule inhibitors of other motors (such as dynein (35,36)), the ATP-based strategy may complement current photobleaching- (37) and force-based (38) methods to shed light on the number of kinesin motors that are involved in the transport of cargos purified from cellular extracts.

SUPPORTING MATERIAL

Supporting Materials and Methods and two figures are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(16)30297-1.

AUTHOR CONTRIBUTIONS

J.X. designed the investigation, Q.L. performed the experiments, A.G. and J.X. carried out the theory and simulation work, and S.J.K. purified proteins. All authors analyzed the data. J.X., S.J.K., and A.G. wrote the manuscript.

ACKNOWLEDGMENTS

We thank our reviewers for helpful comments, Steven P. Gross for helpful discussions, and Tiffany J. Vora for manuscript editing.

This work was supported by grants from the UC Merced Health Sciences Research Institute (Biomedical Seed Grant to J.X.), the UC Merced Academic Senate Committee on Research (to J.X. and A.G.), the National Institutes of Health (NS048501 to S.J.K.), the National Science Foundation (EF-1038697 to A.G.), and the James S. McDonnell Foundation (to A.G.).

REFERENCES

- Vale, R. D. 2003. The molecular motor toolbox for intracellular transport. *Cell*. 112:467–480.
- Block, S. M., L. S. Goldstein, and B. J. Schnapp. 1990. Bead movement by single kinesin molecules studied with optical tweezers. *Nature*. 348:348–352.
- Svoboda, K., and S. M. Block. 1994. Force and velocity measured for single kinesin molecules. *Cell*. 77:773–784.
- Neuman, K. C., and S. M. Block. 2004. Optical trapping. *Rev. Sci. Instrum.* 75:2787–2809.
- Furuta, K., A. Furuta, ..., H. Kojima. 2013. Measuring collective transport by defined numbers of processive and nonprocessive kinesin motors. *Proc. Natl. Acad. Sci. USA*. 110:501–506.
- Beeg, J., S. Klumpp, ..., R. Lipowsky. 2008. Transport of beads by several kinesin motors. *Biophys. J.* 94:532–541.

- Walcott, S., P. M. Fagnant, ..., D. M. Warshaw. 2009. Smooth muscle heavy meromyosin phosphorylated on one of its two heads supports force and motion. J. Biol. Chem. 284:18244–18251.
- Korn, C. B., S. Klumpp, ..., U. S. Schwarz. 2009. Stochastic simulations of cargo transport by processive molecular motors. *J. Chem. Phys.* 131:245107.
- 9. King, S. J., and T. A. Schroer. 2000. Dynactin increases the processivity of the cytoplasmic dynein motor. *Nat. Cell Biol.* 2:20–24.
- Ori-McKenney, K. M., J. Xu, ..., R. B. Vallee. 2010. A cytoplasmic dynein tail mutation impairs motor processivity. *Nat. Cell Biol.* 12:1228–1234.
- Jamison, D. K., J. W. Driver, ..., M. R. Diehl. 2010. Two kinesins transport cargo primarily via the action of one motor: implications for intracellular transport. *Biophys. J.* 99:2967–2977.
- Jannasch, A., V. Bormuth, ..., E. Schäffer. 2013. Kinesin-8 is a lowforce motor protein with a weakly bound slip state. *Biophys. J.* 104:2456–2464.
- Nicholas, M. P., L. Rao, and A. Gennerich. 2014. An improved optical tweezers assay for measuring the force generation of single kinesin molecules. *Methods Mol. Biol.* 1136:171–246.
- Klumpp, S., and R. Lipowsky. 2005. Cooperative cargo transport by several molecular motors. *Proc. Natl. Acad. Sci. USA*. 102:17284– 17289.
- Xu, J., Z. Shu, ..., S. P. Gross. 2012. Tuning multiple motor travel via single motor velocity. *Traffic*. 13:1198–1205.
- Xu, J., S. J. King, ..., B. Nemec. 2013. Interplay between velocity and travel distance of kinesin-based transport in the presence of tau. *Biophys. J.* 105:L23–L25.
- Norris, S. R., V. Soppina, ..., K. J. Verhey. 2014. A method for multiprotein assembly in cells reveals independent action of kinesins in complex. J. Cell Biol. 207:393–406.
- Ando, D., M. K. Mattson, ..., A. Gopinathan. 2014. Cooperative protofilament switching emerges from inter-motor interference in multiple-motor transport. *Sci. Rep.* 4:7255.
- Sloboda, R. D., and J. L. Rosenbaum. 1982. Purification and assay of microtubule-associated proteins (MAPs). *Methods Enzymol.* (85 Pt B):409–416.
- Schroer, T. A., and M. P. Sheetz. 1991. Two activators of microtubulebased vesicle transport. J. Cell Biol. 115:1309–1318.
- Vershinin, M., B. C. Carter, ..., S. P. Gross. 2007. Multiple-motor based transport and its regulation by Tau. *Proc. Natl. Acad. Sci. USA*. 104:87–92.
- Xu, J., B. J. Reddy, ..., S. P. Gross. 2012. Casein kinase 2 reverses tailindependent inactivation of kinesin-1. *Nat. Commun.* 3:754.

- Carter, B. C., G. T. Shubeita, and S. P. Gross. 2005. Tracking single particles: a user-friendly quantitative evaluation. *Phys. Biol.* 2:60–72.
- Yajima, J., M. C. Alonso, ..., Y. Y. Toyoshima. 2002. Direct long-term observation of kinesin processivity at low load. *Curr. Biol.* 12:301–306.
- Driver, J. W., D. K. Jamison, ..., M. R. Diehl. 2011. Productive cooperation among processive motors depends inversely on their mechanochemical efficiency. *Biophys. J.* 101:386–395.
- Jamison, D. K., J. W. Driver, and M. R. Diehl. 2012. Cooperative responses of multiple kinesins to variable and constant loads. *J. Biol. Chem.* 287:3357–3365.
- Schnitzer, M. J., K. Visscher, and S. M. Block. 2000. Force production by single kinesin motors. *Nat. Cell Biol.* 2:718–723.
- Andreasson, J. O., B. Milic, ..., S. M. Block. 2015. Examining kinesin processivity within a general gating framework. *eLife*. 4:e07403.
- Hirokawa, N., K. K. Pfister, ..., G. S. Bloom. 1989. Submolecular domains of bovine brain kinesin identified by electron microscopy and monoclonal antibody decoration. *Cell*. 56:867–878.
- Scholey, J. M., J. Heuser, ..., L. S. Goldstein. 1989. Identification of globular mechanochemical heads of kinesin. *Nature*. 338:355–357.
- Kerssemakers, J., J. Howard, ..., S. Diez. 2006. The distance that kinesin-1 holds its cargo from the microtubule surface measured by fluorescence interference contrast microscopy. *Proc. Natl. Acad. Sci. USA*. 103:15812–15817.
- Vallee, R. B., J. S. Wall, ..., H. S. Shpetner. 1988. Microtubule-associated protein 1C from brain is a two-headed cytosolic dynein. *Nature*. 332:561–563.
- Sakakibara, H., H. Kojima, ..., K. Oiwa. 1999. Inner-arm dynein c of Chlamydomonas flagella is a single-headed processive motor. *Nature*. 400:586–590.
- Kon, T., K. Sutoh, and G. Kurisu. 2011. X-ray structure of a functional full-length dynein motor domain. *Nat. Struct. Mol. Biol.* 18:638–642.
- Kobayashi, T., T. Martensen, ..., M. Flavin. 1978. Inhibition of dynein ATPase by vanadate, and its possible use as a probe for the role of dynein in cytoplasmic motility. *Biochem. Biophys. Res. Commun.* 81:1313–1318.
- Firestone, A. J., J. S. Weinger, ..., J. K. Chen. 2012. Small-molecule inhibitors of the AAA+ ATPase motor cytoplasmic dynein. *Nature*. 484:125–129.
- Hendricks, A. G., E. Perlson, ..., E. L. Holzbaur. 2010. Motor coordination via a tug-of-war mechanism drives bidirectional vesicle transport. *Curr. Biol.* 20:697–702.
- Barak, P., A. Rai, ..., R. Mallik. 2013. Quantitative optical trapping on single organelles in cell extract. *Nat. Methods.* 10:68–70.