

Kinesin Processivity Is Determined by a Kinetic Race from a Vulnerable One-Head-Bound State

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ABSTRACT Kinesin processivity, defined as the average number of steps that occur per interaction with a microtubule, is an important biophysical determinant of the motor's intracellular capabilities. Despite its fundamental importance to the diversity of tasks that kinesins carry out in cells, no existing quantitative model fully explains how structural differences between kinesins alter kinetic rates in the ATPase cycle to produce functional changes in processivity. Here we use high-resolution single-mole-cule microscopy to directly observe the stepping behavior of kinesin-1 and -2 family motors with different length neck-linker domains. We characterize a one-head-bound posthydrolysis vulnerable state where a kinetic race occurs between attachment of the tethered head to its next binding site and detachment of the bound head from the microtubule. We find that greater processivity is correlated with faster attachment of the tethered head from this vulnerable state. In compliment, we show that slowing detachment from this vulnerable state by strengthening motor-microtubule electrostatic interactions also increases processivity. Furthermore, we provide evidence that attachment of the tethered head is irreversible, suggesting a first passage model for exit from the vulnerable state. Overall, our results provide a kinetic framework for explaining kinesin processivity and for mapping structural differences to functional differences in diverse kinesin isoforms.

INTRODUCTION

Kinesin motor proteins drive many active processes in the cell, including vesicle transport (1-3), DNA and organelle repositioning (4,5), intraflagellar transport (6,7), microtubule dynamics control (8-10), and mitotic spindle organization (11-13). There are 45 kinesin genes in the human genome, each of which encodes an isoform that is optimized to drive some processes, but is incapable of driving others (14,15). Understanding the nature of this functional specialization is critical to elucidating the molecular bases of Charcot-Marie-Tooth disease (16), hereditary spastic paraplegia (17,18), Alzheimer's disease (19,20), and the various cancers (21,22) associated with either kinesin dysfunction or overactivity. Part of kinesin's functional diversity can be understood by major structural differences between isoforms. For example, kinesin-5 acts as a tetramer instead of the typical dimer (23), and kinesin-14 has its motor domain at its C-terminus rather than its N-terminus (24). However, most kinesins have a similar structure, and their functional specialization thus comes from the tuning of motility parameters such as velocity, processivity, and force sensi-

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tivity. Here we focus on processivity, or the number of steps kinesin takes per interaction with a microtubule. Processivity values for different kinesins vary over multiple orders of magnitude from just a few steps (25) to a thousand steps or higher (26,27), in part enabling them to perform their different tasks in the cell. Despite its fundamental importance, a consensus quantitative model that explains how processivity is tuned between kinesin isoforms is absent from the literature. This is in part due to the inability to measure kinetic intermediates in the ATPase cycle from which processivity can be controlled. Here we apply new high-resolution tracking technology (25,28–31) to fill this gap in the literature.

The simplest model for kinesin processivity is the kinetic race model, which posits that kinesin must proceed through a single vulnerable state each turnover of its ATPase cycle (32). In our previous work, we identified a one-head-bound posthydrolysis state as a candidate for the vulnerable state (28,33). This result suggested that processivity could be controlled by a kinetic race between tethered head attachment and bound head detachment from the one-head-bound intermediate. In this study, we examine these rate constants with high-resolution single-molecule microscopy as we implement four methods to alter processivity: changing the motor domain from kinesin-1 to -2, altering



the neck-linker (NL) length, changing the solvent ionic strength, and using slowly dissociating ATP analogs. We find that any method for increasing processivity leads to either the tethered head finding the next microtubule binding site more quickly or the bound head dissociating from the current microtubule binding site more slowly. Thus, we find that the kinetic race model is sufficient for quantitatively explaining processivity under physiological conditions, and that the intracellular capabilities of a given motor in part stem from the tuning of the two rate constants in the race.

MATERIALS AND METHODS

Constructs and protein preparation

All kinesin proteins used were expressed in BL21(DE3) bacteria (New England Biolabs, Ipswich, MA) with a C-terminal 6× His tag and purified by affinity chromatography followed by buffer exchange as reported in Mickolajczyk et al. (28) and Chen et al. (34). The kinesin-1 construct used was Drosophila KHC truncated at amino acid (aa) 559, and the elongated NL K117 construct was made by adding the 3-aa DAL at position 345 just preceding α 7, consistent with previous works (35–37). The kinesin-2 construct used was human KIF3A motor and NL homodimerized using the kinesin-1 coiled-coil (345-559) (36), and the K214 construct was made by deleting the DAL sequence at the end of the NL and replacing the kinked proline at positon 355 with an alanine (36). The GFP constructs included a C-terminal eGFP directly preceding the His tag (36-38). The N-terminal avi-tag constructs were described in Mickolajczyk et al. (28). For the kinesin-1 and -2 constructs, the linkers GG and GGAGG, respectively, were added directly downstream of the avi-tag. Biotin was added after induction, typically 0.5-2 h before cell lysis to empirically control the degree of biotinylation of the homodimers. Biotinylation was quantified by comparing the biotin concentration as measured by the colorimetric HABA assay (Sigma-Aldrich, St. Louis, MO) to total protein concentration as indicated by absorbance at 280 nm, and all motors prepared had <0.2 mol of bound biotin per mole of dimer motor. Kinesins used for biochemical assays were truncated at position 406 rather than 560 to enable higher yields (28,34). Insertions and deletions were made either using Q5 or Gibson Assembly (New England Biolabs).

Single-molecule experiments

Gold nanoparticles were imaged using a custom-built total internal reflection dark field microscope, employing a Sapphire-LP 532 nm laser (5-10 mW at sample; Coherent, Bloomfield, CT). Images were recorded using a Basler Ace acA640-750um CMOS camera (1000 frames per s, 945 μ s exposure; Basler, Ahrensburg, Germany) accessed by custom LabVIEW software (National Instruments, Cos Cob, CT). Microtubules were adhered to cleaned coverslips as in Mickolajczyk et al. (28). All assays, unless otherwise noted, were carried out in imaging solution: 0.5 mg/mL casein, 10 μ M taxol, 20 mM glucose, 20 µg/mL glucose oxidase, 8 µg/mL catalase, 0.2 mg/mL BSA, 1:200 β-mercaptoethanol, and 2 mM MgATP in BRB80 (80 mM K-PIPES, 1 mM EGTA, 1 mM MgCl₂, pH 6.8). Biotinylated motors were incubated with a stoichiometric excess of streptavidin-coated 30-nm gold nanoparticles (BBI Solutions, Cardiff, Wales, UK) on ice for 30 min, then spun down at 20,000 g for 4 min and resuspended at working levels (100-300 pM gold) in imaging solution. Point spread functions were fit using the software FIESTA (39) to obtain (X,Y,t) data.

GFP was imaged by total internal reflection fluorescence microscopy using a model No. TE2000 inverted microscope (Nikon, Melville, NY) as in Mickolajczyk et al. (28) and Shastry and Hancock (36,37). Frame rates were set to 3 and 20 frames per s for ATP and ADP experiments, respectively. GFP motors were used at 10–300 pM. ATP movies were analyzed using FIESTA (39), and velocity and run length were determined from the returned distance over time trace. Population run lengths were determined by fitting the cumulative density function (after removing runs <5 pixels, 71 nm/pixel) with an exponential with an *X* offset. Mant run lengths were measured in imaging solution with 0.1 mM mantATP. ADP durations were measured manually from kymographs drawn at positions coincident with Cy5-labeled microtubules in the software ImageJ (National Institutes of Health; http://imagej.nih.gov/ij/) (34). All experiments were performed at 22–23°C.

Stopped-flow spectrofluorometry and steadystate biochemistry

ATP half-site experiments were performed on an Applied Photophysics SX20 spectrofluorometer (Applied Photophysics, Leatherhead, Surrey, UK) (34). Data acquisition and fitting were performed in Pro-Data SX software (Applied Photophysics). One syringe was filled with 2-6 µM microtubules, 10 µM taxol, 500 nM mantADP, and 200-600 nM motor in BRB80 buffer (1:10 molar ratio motor dimer to microtubule). For kinesin-2, 500 nM mantGDP was used instead of mantADP due to the higher affinity of the motor for mantADP (34). The second syringe was filled with 4 mM ATP in BRB80. Excitation was set to 356 nm for the mantADP nucleotide and 450 nm emission was collected using a HQ480SP emission filter (Chroma Technology, Bellows Falls, VT). An integration time of 1 ms was set for the photomultiplier tube. Each experiment contained 1200 μ L split into 60 μ L shots, and the ensemble average fluorescence transient generated upon mixing was fit with a double exponential in the range of 1-500 ms. The faster rate constant was reported. The experiment was repeated at least nine times for each motor, and data sets were taken on at least two separate days for each motor. Solution exchange of cold versus mantADP to determine relative affinity was done as previously (34). ATPase experiments were carried out as in Chen et al. (34) with 5-10 nM motor. ATP turnover rates were determined from the Michaelis-Menton fit to the microtubule dependence of the observed reaction rate, corrected for initial kinesin concentration as determined by mant-ADP exchange (25). All experiments were performed at 22-23°C.

For details on the Hidden Markov Model, two-dimensional step-finding algorithm, and calculation of rates for the first passage model, see Supporting Material.

RESULTS

High-resolution single-molecule microscopy enables direct measurement of a one-headbound intermediate in the stepping cycle of kinesins-1 and -2

To test the hypothesis that kinesin processivity is set by a kinetic race between attachment of the tethered head and detachment of the bound head when the motor is in the one-head-bound (1HB) posthydrolysis state (Fig. 1 *A*, *state 3*), we performed high-resolution single-molecule tracking to measure the 1HB duration for multiple kinesins with different processivities. The motors used were wild-type kinesin-1 (K1₁₄), kinesin-1 with the NL extended to 17 aa (K1₁₇), wild-type kinesin-2 (K2₁₇), and kinesin-2 with the NL shortened to 14 aa (K2₁₄). It was previously shown that extending the NL reduces processivity independent of the motor domain (35–37). For all motors, a 30-nm-diameter gold nanoparticle was bound to the motor domain



FIGURE 1 Extending the NL reduces processivity by slowing the tethered head attachment rate. (*A*) Five-state mechanochemical cycle established in Mickolajczyk et al. (28) and Chen et al. (34), in which processivity is set by a kinetic race between attachment of the tethered (*blue*) head and detachment of the bound (*green*) head in the 1HB posthydrolysis state (state 3). Here *T* denotes ATP, *D* denotes ADP, *DP* denotes ADP plus phosphate, and φ denotes no nucleotide. (*B*) Example 1000 frames/s traces of kinesin-1 (K1) and kinesin-2 (K2) motors with 14- and 17-aa NL domains stepping in 2 mM ATP. Horizontal lines denote inferred microtubule binding sites of the nanoparticle-labeled head. Long states on the microtubule (*black points*) and short states off the microtubule (*colored points* corresponding to motor type) were determined by fitting to the (*X*,*Y*,*t*) data (details in Supporting Material). Each data point represents 1 ms. (*C*) Two-state mechanical model in which each 16.4-nm step is separated into 1HB and 2HB states. Long and short states correspond to black and colored points, respectively, in (*B*). Note: the terms "long" and "short" refer to duration, not molecular configuration. (*D*) Distributions of N = 370 long states and N = 347 short states for K1₁₄; see Fig. S2 for other motor distributions. (*E*) 1HB and 2HB durations showing that for both kinesin-1 and kinesin-2, lengthening the neck linker from 14 to 17 aa increased the 1HB duration and did not change the 2HB duration. Values were calculated from means of long and short state distributions. Error bars represent propagated error from the SE of long and short state distributions. (*F*) Run length is negatively correlated with 1HB duration for both kinesin-1 and -2, indicating that motors with long NL have reduced processivity due to a reduced k_{Attach} . Colors denoting motor type correspond to (*B*). To see this figure in color, go online.

using biotin-streptavidin (see Materials and Methods), a technique that has been shown both experimentally (28,30) and theoretically (40) to have little influence on stepping. The motors were observed in an in vitro single-molecule stepping assay under total internal reflection dark field microscopy at 1000 frames per s in 2 mM ATP. Example position versus time traces for the four kinesin constructs, obtained by fitting point spread functions of gold nanoparticles moving along microtubules (39,41), are shown in Fig. 1 *B*. Kinesin-1 with a single motor domain tagged has previously been shown to take 16.4 nm hand-over-hand steps (42–44), and we recently established that

at saturating ATP, a one-head-bound intermediate can be measured in which the tethered head is displaced ~ 8 nm from its previous binding site (28). Here we clearly observe this intermediate state for all kinesin-1 and -2 constructs tested (Fig. 1 *B*, colored points). Because only one head is labeled, this short duration intermediate ("short state"; Fig. 1, *C* and *D*) represents a single 1HB state. The longer duration intermediate ("long state"), where the labeled motor domain is bound to the microtubule (Fig. 1 *B*, black points), represents the two-head-bound (2HB) state of the labeled head plus the 1HB and 2HB states of the unlabeled head (Fig. 1 *C*). To quantify the duration the motors spent in the long and short states, the (X,Y,t) data were fit using a hidden Markov model (see *state space diagram* in Fig. S1; and see Materials and Methods and Supporting Material for details). From the stepping traces, distributions of at least 300 determinations of each long and short state were built up for each of the four motors studied (K1₁₄ distributions shown in Fig. 1 *D*; see Fig. S2 for all distributions). 1HB and 2HB durations were then calculated from long and short distribution sample means (Fig. 1 *E*). The key result was that for both kinesin-1 and kinesin-2, extending the NL from 14 to 17 aa increased the 1HB duration, while having little effect on the 2HB duration.

Altering the neck-linker length modulates the processivity of kinesin-1 and -2 by tuning the tethered head attachment rate

We next measured the run lengths (RL) of the four motors using GFP constructs (Fig. 1 *F*; distributions in Fig. S3). Consistent with previous measurements (35-37), constructs with the same NL length had similar RLs independent of motor domain. Importantly, a strong negative correlation was observed between the 1HB duration and the run length when comparing the 14 to the 17-aa NL construct of each motor type (Fig. 1 *F*). From the established mechanochemical cycle (Fig. 1 *A*), the 1HB state includes both a pre- and posthydrolysis state with detachment occurring from the posthydrolysis state (28,33). Therefore, the total 1HB duration (Fig. 1 *A* states 2 and 3) is:

$$au_{1\mathrm{HB}} = rac{1}{k_{\mathrm{Hydrolysis}}} + rac{1}{k_{\mathrm{Attach}} + k_{\mathrm{Detach}}}.$$

Assuming that the motor rarely returns backward to the 1HB state from the 2HB state (Fig. 1 *A* state $4 \rightarrow 3$ transition; discussed in detail below), we can derive the probability of stepping as a simple kinetic race between attachment and detachment:

$$P(\text{step}) = \frac{k_{\text{Attach}}}{k_{\text{Attach}} + k_{\text{Detach}}}$$

Because the probability of stepping is proportional to the run length, and the run length is >50 steps for all the motors investigated here, it follows that $k_{\text{Attach}} >> k_{\text{Detach}}$. Moreover, the existing evidence (45) indicates that $k_{\text{Hydrolysis}}$ is not significantly affected by NL length. Thus, the difference in the $\tau_{1\text{HB}}$ values for kinesins with the same motor domain but different NL lengths is as follows:

$$\Delta au_{1\mathrm{HB}} = au_{1\mathrm{HB}}^{17} - au_{1\mathrm{HB}}^{14} = rac{1}{k_{\mathrm{Attach}}^{17}} - rac{1}{k_{\mathrm{Attach}}^{14}}$$

Therefore, the differences in 1HB duration in Fig. 1, E and F are best understood as resulting from differences in k_{Attach} . It

follows that for both kinesin-1 and -2, extending the NL from 14 to 17 aa reduces processivity by decreasing k_{Attach} . This result shows that for kinesin-1 and -2, k_{Attach} is primarily determined by the NL domain and not by the motor domain.

Pre-steady-state kinetics independently support 1HB duration differences

To measure the effect of NL length on 1HB duration in an independent experimental platform, we performed ATP half-site release experiments for all four motor constructs using stopped-flow. Motors were incubated with microtubules and low concentrations of fluorescent 2'(3')-O-(Nmethylanthraniloyl) nucleoside disphosphate (mantNDP) to create a 1HB ATP waiting state with the mantNDP in the tethered head (46–48). The motor-microtubule complexes were then flushed against ATP, enabling the motors to proceed through their 1HB states, and a drop in fluorescence intensity was detected when they released their mantNDP into solution (Fig. 2 A). In the scheme of Fig. 2 A, if a high ATP concentration (fast k_{On}^{ATP}) and a low-affinity mantNDP (fast k_{Off}^{D*}) are used, the rate-limiting step in the half-site release process is $k_{1\text{HB}}$. Importantly, extending the NL was recently shown not to affect the mantNDP release rate from the front head in the 2HB state (Fig. 2 A k_{Off}^{D*} ; see Figs. 5E and 7D in (34)). Thus, by



FIGURE 2 Extending the neck linker increases the 1HB duration. (*A*) The sequence of states in the ATP half-site experiment; D^* denotes a fluorescent nucleotide. (*B*) Half-site release rates at 2 mM ATP, measured by fitting fluorescence transients (Fig. S4). Open circles represent 9–11 replicates for each motor, with mean value shown as a black bar. (*C*) Comparison of differences in the 1HB durations ($\Delta \tau_{1HB}$) between 14- and 17-residue neck-linker constructs for kinesin-1 and kinesin-2 measured by single-molecule (SM) experiments (Fig. 1 *E*) and biochemical (BC) assays (Fig. 2 *B*). Error bars denote propagated error from the SE of short state distributions for single-molecule experiments, and from the SE of half-site values for biochemical assays. To see this figure in color, go online.

comparing the half-site release rate of 14- to 17-aa NL constructs, differences in 1HB duration ($\Delta \tau_{1HB}$) can be calculated. Results for half-site release for the four motor constructs are shown in Fig. 2 *B* (see example transients in Fig. S4). In these experiments mantADP was used for kinesin-1, but due to the high mantADP affinity of kinesin-2 (34), mantGDP was used instead (investigated in detail below). Consistent with previous measurements using different motor constructs (49), elongating the NL was seen to decrease the half-site release rate. The $\Delta \tau_{1HB}$ measured by half-site was in excellent agreement with the $\Delta \tau_{1HB}$ measured by single-molecule (Fig. 2 *C*), providing a second independent line of evidence that extending the NL reduces processivity by reducing the tethered head attachment rate.

Altering ionic strength modulates the processivity of kinesin-1 and -2 by tuning the bound head detachment rate

We next investigated the effect of reducing the ionic strength on the velocity and run length of all four motor constructs. Each motor was measured in BRB80 (as above in Figs. 1 and 2), and in buffers containing 40, 20, and 12 mM PIPES. Consistent with previous kinesin-1 measurements (36), reducing the ionic strength was seen to slightly decrease the velocity of all motors (Fig. 3 *A*) and to increase the run lengths of all motors to different extents (Fig. 3 *B*; distributions in Fig. S3). Under the model that processivity is controlled by a kinetic race between tethered head attachment and bound head detachment in the 1HB state, an increase in processivity due to an increase in k_{Attach} must lead to a faster, not a slower velocity. Thus, the measured increase in processivity at low ionic strength more likely stems from a decrease in $k_{\text{Detach.}}$ To test this claim, we measured the microtubule dwell time of all four motors at saturating ADP as a function of ionic strength (Fig. 3 C; see distributions and example kymographs in Fig. S5). Consistent with the prediction that low ionic strength reduces detachment rates, the dwell time increased as the ionic strength was reduced for all four motors. Strikingly, as highlighted when the run lengths (Fig. 3 D) and ADP dwell times (Fig. 3 E) were normalized to their values in BRB80, increases in both parameters were shared by each motor type independent of NL length: K1₁₄ and K1₁₇ both had an \sim 2.5-fold increase in RL and ADP dwell time, whereas $K2_{14}$ and $K2_{17}$ both had an \sim 4–6-fold increase in RL and ADP dwell time. Thus, k_{Detach} can be adjusted by altering the electrostatic interactions between the motor domain and the microtubule, and tuning k_{Detach} is a means of controlling RL in an approximately proportional fashion. The fact that elongating the NL did not affect the ADP dwell time bolsters the claim that changes in RL due to differences in NL length stem from changes in k_{Attach} .

Slowing the ADP off-rate modulates processivity by shunting the motor backward into the 1HB state

The data in Figs. 1–3 establish that kinesin-1 and-2 processivity can be understood as a kinetic race between k_{Attach}



FIGURE 3 Ionic strength alters processivity by tuning the detachment rate from the 1HB vulnerable state. (*A*) Velocities in 2 mM ATP, (*B*) run lengths in 2 mM ATP, and (*C*) dwell times in 2 mM ADP for GFP motors in BRB12, BRB20, BRB40, and BRB80 buffers. Velocities are shown as mean \pm SE with a 10% error added for ~1°C variability in temperature (61). Run lengths and dwell times were determined by distribution fitting with error bars determined by bootstrapping (52). (*D*) Run lengths and (*E*) ADP dwell times normalized to BRB80 values (with propagated error) to emphasize the pairing by motor domain rather than by neck-linker length. All panels are color-coded by legend in (*B*), consistent with Figs. 1 and 2. To see this figure in color, go online.

and k_{Detach} when the motor is in the vulnerable 1HB state (Fig. 4 A, state 2). However, completion of a forward step also involves ADP release to lock in the gains made by the attachment step. Hence, there is a second kinetic race after tethered head attachment-either ADP is released to complete the step, or the attachment step is reversed and the motor returns to the vulnerable 1HB state (Fig. 4 A, state $3 \rightarrow 4$ versus $3 \rightarrow 2$ transition). Depending on the relative kinetic parameters, this process can be described in two ways. In a "first passage model", kAttach is relatively slow and ADP release follows rapidly such that the motor rarely reverts to the 1HB state. In this sense, tethered head attachment is the irreversible ratcheting step. Alternatively, in a "rapid equilibrium model", both k_{Attach} and the reverse rate $k_{\text{Detach}}^{\text{FH}}$ are very fast, and the equilibrium favors the 1HB state sufficiently that it is detected in single-molecule tracking (28). Here, the rare outpacing of $k_{\text{Detach}}^{\text{FH}}$ by $k_{\text{Off}}^{\text{ADP}}$ locks the motor into the 2HB state (Fig. 4 A, equilibrium between states 2 and 3 until the $3 \rightarrow 4$ transition occurs), and ADP release is thus the irreversible ratcheting step. To test between these models, we measured the run lengths of our four GFP constructs in mantATP. Although the off-rates for mant- and cold ADP are similar for kinesin-1 (Fig. 4 B; Fig. S6 A) (28), kinesin-2 has a 15-fold lower off-rate for mantADP than cold ADP (Fig. 4 B) (34). Thus, under the first passage model the kinesin-2 RL should drop only slightly in mantADP, whereas under the rapid equilibrium model the kinesin-2 RL should drop \sim 15-fold in mantATP. We found that in mantATP the RL of $K2_{14}$ and $K2_{17}$ only dropped roughly twofold, thus ruling out the rapid equilibrium model (Fig. 4 C, distributions in Fig. S6 B). As expected from the similarity in their ADP and mantADP off-rates, the



 $K1_{14}$ and $K1_{17}$ RLs were not measurably different in mantATP versus cold ATP.

To ensure quantitative consistency of the first passage model, we modeled the six-state process shown in Fig. 4 A as a Markov chain. We solved for k_{Attach} and k_{Detach} using our cold ATP RL and $\Delta \tau_{1\text{HB}}$ values (Figs. 1 A and 2 C), used literature values for $k_{\text{off}}^{\text{ADP}}$ or $k_{\text{off}}^{\text{mantADP}}$ (28,34), and used measured values for $k_{\text{Detach}}^{\text{ADP}}$ (Fig. 3 C; and see Table S1 and Supporting Material for details). Using transition probabilities calculated from the rate constants, we determined the run length by calculating the average number of times the motor entered the ATP waiting state before detaching (Fig. 4 A state 6). Comparing the modeled mantATP versus cold ATP RL (changing only k_{off}^{ADP} to $k_{off}^{mantADP}$ as in Fig. 4 B), we were able to nearly perfectly replicate the experimental data (Fig. 4 C), validating the first passage model. Hence in physiological ATP, tethered head attachment is the ratcheting step of the cycle and locking-in steps via ADP release occurs with nearly 100% probability.

DISCUSSION

By changing the identity of the motor domain, the length of the NL domain, the ionic strength of the solvent, and the nucleotide off-rate, we demonstrate here that kinesin processivity can be quantitatively described as a kinetic race between tethered head attachment and bound head detachment. This model should generalize to N-terminal kinesins beyond kinesin-1 and -2, and it provides a quantitative framework for understanding how small structural differences between isoforms can tune rates in the ATPase cycle to yield functional differences in motor behavior.

> FIGURE 4 Tethered head attachment follows a first passage model. (A) Mechanochemical model based on Fig. 1 A, showing allowed transitions. When ADP release (state $3 \rightarrow 4$) is significantly slowed, front-head (FH) or rear-head (RH) detachment from state 3 may occur, forcing kinetic flux into vulnerable states (2,5). In the first passage model, $k_{\text{Detach}}^{\text{FH}}$ is small relative to $k_{\text{Off}}^{\text{ADP}}$, whereas in the rapid equilibrium model, $k_{\text{Detach}}^{\text{FH}}$ is very fast relative to k_{Off}^{ADP} (*red arrows*). (*B*) Previous determinations of k_{Off}^{ADP} by front-head nucleotide exchange experiments for kinesin-1 (28) and kinesin-2 (34). These rates are independent of neck-linker length (34). (C) Run lengths in mantATP relative to run lengths in unlabeled (cold) ATP for each motor. Consistency of experiment and model (black squares) supports first passage model for k_{Attach} . To see this figure in color, go online.

Elongating the neck-linker tunes processivity by changing the tethered head attachment rate

Elongating the NL has been previously shown to reduce processivity (36,37,43,49), but the underlying mechanism has remained mysterious. Using high-resolution tracking and stopped-flow, we show here that NL mutants alter the vulnerable state duration, and that independent of motor domain there is a strong negative correlation between vulnerable state duration and processivity (Fig. 1 F). Recent gold-nanoparticle tracking results from Isojima et al. (30) showed that human kinesin-1, with cys-lite modifications in 12 mM PIPES buffer containing saturating ATP, had an increased 1HB duration when the NL was shortened to 13 aa, and slightly decreased duration when the NL was increased to 20 aa (six-glycine insert). However, kinesins with a 13-aa NL have a reduced run length (37) and Cys-lite mutants with various 20-aa NLs have a surprisingly increased run length (50). Hence, their results follow the same negative correlation between vulnerable state duration and processivity. Additionally, our 17-aa NL constructs did not show the unbinding events, sideways steps, or futile hydrolysis (Fig. 1; Fig. S7) that 20-aa NL cys-lite did (30,43), indicating that not all elongated NL constructs should be lumped together for comparison.

One possible explanation for why elongating the NL increases the 1HB duration is that the longer NL increases the conformational search volume of the tethered head. However, an argument against this model is that Drosophila kinesin-1 constructs with NLs from 15-20 residues all have roughly the same RL (27,36). A second possibility is that elongating the NL alters the interaction of the tethered head with the bound head while in the 1HB state. Alonso et al. (51) used biochemical and structural data to argue that in the ATP waiting state, the tethered head is nestled next to the bound head, and chemical events in the bound head release the tethered head to step to the next binding site. If extending the NL disrupted the ability of the bound head to free the tethered head, this would increase the measured 1HB duration and time spent in the vulnerable state. A third possibility is that a NL of 14 aa provides the ideal geometry for the tethered head to find its next binding site, perhaps because the docked NL is rigid and any lengthening or shortening of this domain causes misregistration between the tethered head and the next binding site, thereby reducing the attachment rate.

Electrostatic interactions stabilize the kinesinmicrotubule complex in the vulnerable state

We found that adding or subtracting the 3-aa DAL to the NL affected the tethered head attachment rate but not the bound head detachment rate. Thus, kinesins were paired by NL length when observing attachment (Fig. 1) and paired by motor domain when observing detachment (Fig. 3). The boundhead detachment rate was evaluated by proxy using the dwell time in saturating ADP, an approach justified by the close

agreement of the measured $k_{\text{Detach}}^{\text{ADP}}$ values (Fig. 3 C) with the k_{Detach} values calculated from measured RLs and Δau_{1HB} (Table S1). The effect most likely comes from electrostatic interactions between positively charged residues in kinesin and negatively charged residues in the microtubule, particularly in the tubulin C-terminal tail (52). Consistent with this, cleaving off the C-terminal tail with subtilisin has been shown to decrease kinesin processivity (52–54). It is notable that k_{Detach} can also be affected by adding positive residues outside the motor domain. Adding lysines to the neck coil has been shown to increase processivity (52). Adding lysines when elongating the kinesin-1 NL has been shown to decrease the run length in BRB80 but not in BRB12 (36,43), a result best understood by offsetting changes in kAttach (longer NL) and kDetach (ionic strength dependence). We verified this by showing that $K1_{17}$ with a lysine (KAL inserted) had a stronger ionic strength dependence of run length and dwell time in ADP than K1₁₇ without a lysine (DAL inserted; Fig. S8).

Slowing ADP release reduces processivity by reversing tethered head attachment

When mantATP is used to power kinesin-2 stepping, the slow mantADP release rate increases the probability that the newly bound front head fails to complete its step by releasing its ADP, and instead reverts to the 1HB vulnerable state. This effect only decreases run length approximately twofold (Fig. 4 C), which is surprisingly mild and suggests a first passage model: attachment of the tethered head is the irreversible ratcheting step, and rapid ADP release subsequently locks-in the 1HB to 2HB transition. Our results and model predictions show that kinesin-1 and -2 both rectify steps with nearly 100% efficiency in cold ATP, meaning that the vulnerable state is only entered once per cycle the majority of the time. In Fig. 4, the efficiency of capturing the forward step was compromised chemically by using a high-affinity nucleotide analog. It can also be compromised mechanically by increasing intramolecular tension: Isojima et al. (30) showed that kinesin-1 with a 13-residue NL occasionally returned to the 1HB state after tapping down on the microtubule-binding site, indicating an increased $k_{\text{Detach}}^{\text{FH}}$. Multiple isoforms of kinesin have been shown to undergo one-dimensional diffusion when forced into the ADP state (26,55-58), and some low-processivity kinesins such as MCAK utilize diffusion in the ADP or ADP-Pi state in the cell to perform their role in mitosis (58). It follows that these kinesins have a greatly slowed ADP off-rate (59), such that the efficiency of locking-in steps is reduced and kinetic flux is forced into the 1HB ADP-Pi and ADP states.

Assisting loads decrease run length by enhancing detachment

This work also provides insights into how external loads alter kinesin processivity. Optical trapping experiments on

wild-type kinesin-1 have shown that hindering loads reduce motor velocity and run length, whereas assisting loads have no effect on velocity but strongly reduce run length (27,33,60). Our results show that either k_{Attach} or k_{Detach} can affect run length, but of these two only k_{Attach} affects velocity. Thus, the decrease in run length under hindering load may result from changes to k_{Attach} and k_{Detach} , but the decrease in run length under assisting load must result solely from changes to k_{Detach} . This makes intuitive sense, as it not clear how an assisting load would decrease kAttach, and assisting loads have been shown to rescue the irregular stepping of mutants (43). It also means that assisting loads must speed k_{Detach} to a greater degree than do hindering loads. External loads may also increase the front head or rear head detachment rates from the 2HB state (Fig. 4 A, states $3 \rightarrow 2$ and $3 \rightarrow 5$ transitions, respectively), and thus increase the flux into vulnerable states in a fashion similar to mantATP (Fig. 4).

Here we show that processivity, an important biophysical determinant of intracellular function, is set by a kinetic race between attachment of the tethered head and detachment of the bound head when kinesin is in a vulnerable one-headbound state. This simple and general model provides a link between kinesin structure and function that should apply across the kinesin superfamily in both healthy and disease states.

SUPPORTING MATERIAL

Supporting Materials and Methods, eight figures, and one table are available at http://www.biophysj.org/biophysj/supplemental/S0006-3495(17) 30511-8.

AUTHOR CONTRIBUTIONS

K.J.M. built the total internal reflection dark field microscope, designed experiments, performed experiments, analyzed data, and wrote the article. W.O.H. designed experiments and wrote the article.

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SUPPORTING CITATIONS

References (62,63) appear in the Supporting Material.

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