The Kinetics of Nucleotide Binding to Isolated *Chlamydomonas* Axonemes Using UV-TIRF Microscopy

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ABSTRACT Cilia and flagella are long, slender organelles found in many eukaryotic cells, where they have sensory, developmental, and motile functions. All cilia and flagella contain a microtubule-based structure called the axoneme. In motile cilia and flagella, which drive cell locomotion and fluid transport, the axoneme contains, along most of its length, motor proteins from the axonemal dynein family. These motor proteins drive motility by using energy derived from the hydrolysis of ATP to generate a bending wave, which travels down the axoneme. As a first step toward visualizing the ATPase activity of the axonemal dyneins during bending, we have investigated the kinetics of nucleotide binding to axonemes. Using a specially built ultraviolet total internal reflection fluorescence microscope, we found that the fluorescent ATP analog methylanthraniloyl ATP (mantATP), which has been shown to support axonemal motility, binds all along isolated, immobilized axonemes. By studying the recovery of fluorescence after photobleaching, we found that there are three mantATP binding sites: one that bleaches rapidly (time constant ≈ 1.7 s) and recovers slowly (time constant = 44 s), one that bleaches with the same time constant but does not recover, and one that does not bleach. By reducing the dynein content in the axoneme using mutants and salt extraction, we provide evidence that the slow-recovering component, but not the other components, corresponds to axonemal dyneins. The recovery rate of this component, however, is too slow to be consistent with the activation of beating observed at higher mantATP concentrations; this indicates that the dyneins may be inhibited due to their immobilization at the surface. The development of this method is a first step toward direct observation of the traveling wave of dynein activity.

INTRODUCTION

Motile eukaryotic cilia and flagella are dynamic structures that move in a periodic wave-like pattern to propel cells through liquid and to move fluid across cell surfaces (1). At the core of cilia and flagella is the axoneme, a bundle of nine doublet microtubules surrounding a core of two single microtubules, together with hundreds of other associated proteins (2,3). Among these associated proteins are the axonemal dyneins, which are located between the doublets along the length of the axoneme (4). The dyneins use energy derived from the hydrolysis of ATP to slide microtubule doublets (5–7), and the sliding is converted to bending by constraints that prevent sliding at the base. The dyneins undergo an ATPase cycle: ATP binds and is hydrolyzed, and the products (ADP and organic phosphate) are released (8,9). The ATPase cycle is thought to be coupled to force...
generation through conformational changes of the dyneins (10–12). The similarity of the frequency of the axonemal bending wave to the ATPase rate of the axonemal dyneins suggests that each dynein ATP cycle is coupled to a cycle of doublet sliding and bending (13). In this way, dynein is thought to power the axonemal beat.

Dyneins are believed to generate sliding forces all along the length of the axoneme. First, the dyneins are present all along the length (14). And second, early modeling studies, which predated the discovery of dynein, argued that the bending wave was caused by a wave of activity that traveled all along the length: if motors were only active at the base, driving a whip-like motion of the axoneme, then the amplitude of the bending wave would decrease as it propagates (15), in contradiction to observations. The hypothesis that the axonemal bending wave is driven by a traveling wave of dynein activation forms the basis for many subsequent models, which can accurately predict the axonemal waveform (15–19). However, this hypothesized wave of dynein activity has never been directly measured or observed. Direct observation of a traveling wave of dynein activation is important not just to test the models but also to provide information about the extent of the modulation of the dynein activity, as well as its spatial and temporal properties.

One reason why it is difficult to observe directly the postulated traveling wave of dynein activation is that it is difficult to assay the activity of dynein. One experimental avenue is electron cryomicroscopy. A limitation of this approach, however, is that it is difficult to rapidly freeze a beating cilium or axoneme. For example, in a recent study (12), an active axoneme was rapidly frozen, but during the sample preparation, which took a few seconds, it is possible that beating stopped or was altered. Another avenue is to use fluorescence to monitor the binding and unbinding of ATP. However, this approach is difficult because dyneins have a higher specificity for nucleotides than other motors such as myosins and kinesins: only a small number of ATP analogs have been shown to support force generation by axonemal dyneins (20–22). Indeed, only the fluorescently labeled nucleotides 2′-(or 3′)-(N-methylanthraniloyl) ATP (mantATP) and 2′-(or 3′)-O-anthraniloyl ATP can support reactivation of isolated, demembranated axonemes, although at frequencies significantly reduced from that of ATP (21,23). MantATP also supports translocation of microtubules in gliding assays, in which outer-arm dyneins (24; https://www.qucosa.de/api/qucosa%3A34182/attachment/ATT-0/) or inner-arm dyneins (25) are bound to the glass surface of a perfusion chamber and microtubules introduced into the solution are observed to glide along the surface. Although the fluorescence of mantATP has been used in solution studies on axonemal dyneins (26), mantATP fluorescence has not been observed in the intact axoneme structure. This is partly due to the low quantum efficiency of the dye and partly due to the excitation and emission spectra of mantATP, which have peaks at 355 and 448 nm, respectively (https://www.jenabioscience.com/images/PDF/NU-202.0001.pdf), and make detection difficult because of the high background signal.

Recent developments in highly sensitive cameras for use with microscopy allow imaging of low-emitting dyes such as mantATP. Here, we report for the first time, to our knowledge, the direct observation of binding of mantATP to intact immobilized axonemes using ultraviolet (UV) total internal reflection fluorescence (TIRF) microscopy.

METHODS

Reagents

MantATP (trisodium salt) was purchased from Jena Bioscience (Jena, Germany). All other reagents were purchased from Sigma Aldrich (St. Louis, MO) unless stated otherwise.

Chlamydomonas cells

Chlamydomonas reinhardtii cells (CC-125 wild-type mt+/+ 137c) were grown in liquid tris-acetate-phosphate medium: 20 mM tris, 7 mM NH₄Cl, 0.40 mM MgSO₄, 0.34 mM CaCl₂, 2.5 mM PO₄³⁻, and 1000-fold diluted Hunter’s trace elements (14). The medium was titrated to pH 7.0 with glacial acetic acid.

Axonemes

Axonemes were purified using the method described in Alper et al. (27). Chlamydomonas cells were harvested by centrifugation at 900 × g for 5 min. They were then deflagellated by incubation with 4.2 mM dibucaine-HCl for 90 s. The flagella were separated from the cell bodies by centrifugation at 24,000 × g for 20 min on a 30% sucrose cushion. Flagella were then concentrated by resuspending the pellet in 10 mL of HMDE buffer (30 mM HEPES, 5 mM MgSO₄, 1 mM dithiothreitol, and 1 mM EGTA, titrated to pH 7.4 with KOH) with addition of 0.4 mM Pefabloc. The flagella were then demembranated by adding 0.2% IGEPAL CA-630 and washed in HMDE buffer. Salt extraction of axonemes was performed by incubating with the indicated amount of KCl (0.3–1 M) + HMDE for 20 min; the extracted axonemes were then centrifuged at 31,000 × g for 20 min to remove the salt and extracted protein. The axonemes were re-suspended in HMDE buffer.

Measurement of dynein content

Serial dilutions of each kind of axoneme were loaded on sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) gels, 4–15% precast Mini-PROTExAN TGX gels, tris/tricine/SDS running buffer, and SDS sample buffer (BioRad, Hercules, CA). Gels were then stained with Coomassie brilliant blue (Life Technologies, Carlsbad, CA). Images of gels were obtained by scanning the gel on an Epson V700 document scanner (Epson, Suwa, Nagano, Japan). The gels were analyzed in Fiji image analysis software (16). The amount of dynein and tubulin was measured by integrating the density of the respective bands in the scanned gels. The dynein content was defined as the dynein signal divided by the tubulin signal, the latter serving as a loading control.

Imaging

Imaging was performed using a Zeiss Axiovert 200M (Zeiss, Oberkochen, Germany) microscope with a Zeiss Alpha Plan-Fluar 100×/1.45 oil
objective and a home-built laser-TIRF line (Fig. 1, Table 1). A UV diode-pumped laser (model Zouk 05-01, 355 nm, maximal power 10 mW (Cobolt, Solna, Sweden)) was used as the light source. Images were recorded with an iXon 887 electron-multiplying CCD back-illuminated camera (Andor, Belfast, UK).

For imaging, axonemes in HMDEKP buffer (30 mM HEPES, 5 mM MgSO₄, 1 mM dithiothreitol, and 1 mM EGTA, 50 mM K-acetate, 1% w/v polyethylene glycol, titrated to pH 7.4 with KOH) were infused into an experimental chamber of depth 0.1 mm formed by a coverslip and a micropipette. We observed a fluorescent signal along the entire axoneme after the laser is shuttered (Fig. 2). The bleaching phase was fitted with the exponential functions using nonlinear least squares (MATLAB, “fit” function).

RESULTS

MantATP binds to the axoneme

We incubated flow cells with solutions containing isolated, demembranated *Chlamydomonas* axonemes and waited a few minutes until several had bound to the surface in each field of view (82 × 82 μm). We then infused a solution containing 1 μM mantATP. Using a TIRF microscope with long-wavelength UV illumination (Fig. 1, 355 nm laser), we observed a fluorescent signal along the entire axoneme (Fig. 2 A, TIRF image in upper panel, differential interference contrast image in lower panel). The same assay was performed in the presence of hog arc lamp Mercury arc lamp HBO 100 Zeiss

Hg arc lamp Mercury arc lamp HBO 100 Zeiss

Electronic controller Ludl Electronic Products MAC 2002 LEP

Computer Dell Precision T1700 workstation Dell

M1–M4 25.4 mm mirror, UV-enhanced aluminum ThorLabs

M5 Dichroic mirror, cutoff 400 nm long pass Chroma

L1 Fused silica plano-concave lens ThorLabs

L2 Fused silica plano-convex lens ThorLabs

L3 Fused silica plano-convex lens ThorLabs

Filter wheel Six-position motorized filter wheel, for 25.4 mm optics Fluoroview 102C ThorLabs

Shutter 25 mm optical shutter VS25 Uniblitz

Microscope Axiovert 200M microscope with differential interference contrast/phase capable condenser Zeiss

Objective Alpha Plan-Fluar 100× NA = 1.45 oil objective Zeiss

Camera iXon 887 electron-multiplying CCD back-illuminated camera Andor

Filter cube Interchangeable filter cube (in the microscope turret). For 355 nm laser TIRF: excitation filter 355/10 nm; emission filter 460/50 nm; mirror 400 nm long pass.

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**Table 1.** List of Parts for the UV-TIRF Microscope Schematic

| Abbreviation | Component Description | Manufacturer |
|--------------|-----------------------|--------------|
| Microscope   | Zeiss Axiovert 200M    | Zeiss        |
| Laser        | UV diode-pumped laser, model Zouk 05-01, 355 nm, maximal power 10 mW (Cobolt, Solna, Sweden) | Zeiss |
| Hg arc lamp  | Mercury arc lamp HBO 100 | Zeiss        |
| Electronic controller | Ludl Electronic Products MAC 2002 | LEP          |
| Computer     | Dell Precision T1700 workstation | Dell |
| L1           | Fused silica plano-concave lens | ThorLabs |
| L2           | Fused silica plano-convex lens | ThorLabs |
| L3           | Fused silica plano-convex lens | ThorLabs |
| Filter wheel | Six-position motorized filter wheel, for 25.4 mm optics Fluoroview 102C | ThorLabs |
| Shutter      | 25 mm optical shutter VS25 | Uniblitz |
| Microscope   | Axiovert 200M microscope with differential interference contrast/phase capable condenser | Zeiss |
| Objective    | Alpha Plan-Fluar 100× NA = 1.45 oil objective | Zeiss |
| Camera       | iXon 887 electron-multiplying CCD back-illuminated camera | Andor |
| Filter cube  | Interchangeable filter cube (in the microscope turret). For 355 nm laser TIRF: excitation filter 355/10 nm; emission filter 460/50 nm; mirror 400 nm long pass. | Zeiss/Chroma |
performed in the absence of mantATP did not produce a fluorescent signal from the axonemes. Thus, there are mantATP binding sites all the along the axoneme, though we cannot exclude the possibility that the fluorescence is reduced within a micrometer from the ends.

To investigate the kinetics of the binding and unbinding of mantATP to the axoneme, we used a bleaching and recovery protocol. In this assay, the fluorescence was bleached by intense, continuous UV illumination, and the recovery of fluorescence was monitored by intermittent brief light pulses. A typical trace is shown in Fig. 2 B, in which there is partial bleaching followed by partial recovery of fluorescence. During the bleaching phase, the fluorescent signal decreased over a few seconds from its initial value \( F_0 \) toward a steady-state value \( F_i \). During the recovery phase, the fluorescent signal approached the recovery value \( F_r \) over approximately 1 min. See Table 2 for the bleaching and recovery rates, together with the amplitudes \( F_0 \), \( F_i \), and \( F_r \). We identified two general features of the bleaching and recovery curves: 1) the bleaching rate was much higher than the recovery rate, and 2) the recovery was always incomplete (\( F_r < F_0 \)).

**The bleaching recovery data indicate that there are at least three nucleotide-binding sites**

To understand the bleaching and recovery curves, we first attempted to fit the time courses with the one-binding-site model shown in Fig. 3. The idea is that mantATP* binds to a site and is hydrolyzed to mantADP*, and the mantADP* unbinds at a slow rate. The (*) refers to the fluorescent nucleotide. If hydrolysis (\( h \)) is faster than bleaching (\( b \))—justified by data on the ATPase activity of Tetrahymena outer-arm dynein (29)—then the bleaching phase corresponds to the unbinding of mantATP* to mantADP\( \dagger \), where (\( \dagger \)) denotes the bleached nucleotide. In this scheme, recovery is limited by the slow unbinding of mantADP\( \dagger \). Usually, we will omit the (\( \dagger \)).

This one-binding-site model cannot account for the bleaching and recovery curves shown in Fig. 2 (see Appendix for the mathematical arguments). First, there should be complete recovery. Instead, the partial recovery indicates that there is a site that bleaches irreversibly. We call this the nonrecovering component. Second, the slow-recovering component should exhibit almost complete bleaching because the bleaching rate (\( k_1 = b + u \), in the scheme) is much faster than the recovery rate (\( k_2 = u \)), which corresponds to the unbinding of mantADP. Formally, \( (F_i/F_0) \approx (k_2/k_1) = (u/b + u) \ll 1 \) (see Appendix). Instead, the bleaching is only partial (\( F_i/F_0 \approx 0.4 \)). This indicates that there is a site that does not bleach; we call this the nonbleaching component. Therefore, the bleaching curves indicate that there are at least three components: a slow-recovering component, a nonbleaching component, and a nonrecovering component.

The three-component model provides a good fit to the experimental data (Fig. 4). In Fig. 4 B, the experimental data are shown in red circles; the fitted sum of the three components is shown with a solid blue curve; and the slow-recovering, nonbleaching, and nonrecovering components

**TABLE 2 Measured Parameters**

| Parameter               | Mean ± Standard Error |
|-------------------------|-----------------------|
| **Timescales**          |                       |
| Bleaching rate          | \( k_1 \) 0.70 ± 0.06 s\(^{-1} \) |
| Recovery rate           | \( k_2 \) 0.027 ± 0.004 s\(^{-1} \) |
| **Relative fluorescence intensities** |                 |
| Initial                 | \( F_0 \) 0.068 ± 0.008 |
| Intermediate            | \( F_i \) 0.027 ± 0.003 |
| Recovery                | \( F_r \) 0.045 ± 0.003 |

Parameters were measured from the bleaching and recovery curves like those shown in Fig. 2 for 10 separate experiments at 1 \( \mu \)M mantATP, with 50–150 axonemes analyzed in each experiment. The value of the intermediate level obtained by fitting the bleaching curves was similar to the value obtained by fitting the recovery curves (bleaching: 0.027 ± 0.003, recovery: 0.027 ± 0.005; mean ± Standard Error, \( n = 10 \) experiments).
Reducing dynein in axonemes using mutants and salt extraction

To determine whether axonemal dynein is contributing to one of the three binding sites, we used axonemes with different dynein contents. One way to reduce the number of dyneins is to use mutants lacking some of the dynein arms. Another way is use axonemes whose dyneins have been extracted with KCl (14). We compared wild-type axonemes, oda1 mutant axonemes (which are missing outer dynein arms), and salt-extracted oda1 axonemes to further reduce the number of dyneins. The amount of dynein in each type of axoneme was measured by scanning SDS-PAGE gels. Oda1 axonemes have less dynein than wild-type axonemes. Increasing the KCl salt concentration in the axonemes, consistent with earlier findings (see Introduction).

Nonrecovering

Nonbleaching

Slow-recovering

The parameters from Table 3 were used to infer the properties of the three components. The nonrecovering component has relative amplitude \( F_s - F_i/F_o \). The slow-recovering component has amplitude \( (F_s - F_i)/F_o \). The nonbleaching component is the rest and is approximately equal to \( F_i/F_o \).

In summary, it is likely that mantATP binds to inner-arm dyneins, consistent with earlier findings (see Introduction).
It is also possible that mantATP binds to outer-arm dynein; however, the large variability of the rate for wild-type axonemes precluded us from making a definitive statement one way or the other. By contrast, the nonbleaching component is not dynein because its amplitude was not diminished in salt-extracted oda1 axonemes. The nonrecovering component, which irreversibly bleaches during the experiment, is also not dynein because it was still present in the salt-extracted oda1 axonemes. Thus, only the slow-recovering component is likely to be dynein.

The bleaching and recovery rates, as well as the amplitudes of the three components, showed little dependence on mantATP concentration

To test whether the slow component is limited by the availability of mantATP, we varied the concentration of mantATP in the assay. Imaging conditions allowed us to vary the fluorescent nucleotide concentration only over a range of 0.1–30 μM. At the lowest concentration, the axonemes were so faint as to be barely visible against the background; at the highest concentration, the background was so bright that the axoneme could be barely discerned.

The recovery rate constant showed little dependence on the mantATP concentration (Fig. 6 C). This is expected because the unbinding of the bleached mantATP is not expected to be influenced by the mantATP in solution (Eq. 12). The bleaching rate was also little affected by mantATP (Fig. 6 D). For the nonrecovering site, this is consistent with its very slow release of bleached mantATP (>100 s). For the slow-recovering site, the lack of dependence on mantATP implies that even at 0.1 μM, most of the binding sites must be occupied (Eq. 10), implying a high affinity $K_D = k_r/k_c \leq 0.1$ μM. These data suggest that both the nonrecovering and slow-recovering sites are high affinity, consistent with the relative amplitudes of the respective components showing little dependence on mantATP concentration (Fig. 6, G and H). The relative amplitude of the nonbleaching component increases somewhat with mantATP (Fig. 6 F), as does its absolute amplitude; this might indicate that there is a low-affinity nonbleaching component. In summary, the bleaching and recovery curves show little dependence on the mantATP concentration.

DISCUSSION

Using UV-TIRF, we visualized mantATP binding to the axoneme and found nucleotide-binding sites distributed approximately uniformly along the axoneme. The bleaching and recovery experiments indicate that there are at least three components: nonrecovering, nonbleaching, and slow-recovering.

We suggest the following interpretations of these components, which have approximately equal amplitudes.

1) The slow-recovering component is a mantATP binding site on axonemal dynein that hydrolyzes mantATP and releases the product mantADP very slowly (on the timescale of the experiment; $\tau_u > 300$ s) so that once bleached, there is no measurable recovery. This explains the absence of a dependence of the bleaching rate and the amplitude of this component on the mantATP concentration.

2) The nonrecovering component could correspond to a high-affinity binding site that unbinds mantATP or mantADP very slowly (on the timescale of the experiment; $\tau_b > 300$ s). Alternatively, this component could correspond to a rapidly exchanging, nonhydrolyzing site with fast unbinding and binding rates compared to the sampling rate of 0.1 s.

If the slow-recovering component corresponds to axonemal dynein and the slow recovery is due to the unbinding of mantADP, this could account for the slow beating of Chlamydomonas axonemes in mantATP, as previously
proposed (21). The hydrophobic methylanthraniloyl substituent on the ribose may make the affinity for the analog to the hydrophobic dynein binding pocket higher, reducing the off rate. We found that isolated demembranated oda1 axonemes do not beat at mantATP concentrations at or below 60 μM (24). At 100 μM mantATP, they reactivate and have a beat frequency of 0.4 Hz (24). Note that the mantATP requirement for *Chlamydomonas* is much higher than previously reported for sea-urchin sperm, which has a beat frequency of 0.3 Hz at 2.5 μM mantATP (23). Unfortunately, we could not study the binding and bleaching dynamics of *Chlamydomonas* axonemes at high enough concentrations to reactivate the beat because of the poor signal/noise ratios. If one mantATP is hydrolyzed per dynein per beat cycle, at 100 μM mantATP, we expect a dynein turnover rate of 0.4 s⁻¹, and this must be faster than all rate constants between different states within the cycle, including mantADP unbinding. However, the recovery rate in our measurements was ~0.02 s⁻¹. If this rate corresponds to mantADP release, then release would need to be accelerated severalfold for beating at 0.4 Hz to occur. Therefore, we propose (assuming that our measured recovery rate corresponds to mantADP

![Figure 6 Dependencies of the fast and slow components on dynein content and mantATP concentration.](image)
release) that either beating itself accelerates mantADP release or that preventing beating through immobilization on the surface inhibits mantADP release. Thus, our observations are consistent with axonemal dyneins showing force-dependent inhibition. This is consistent with an earlier observation that dynein motors in the axoneme continue generating force even when the axoneme is prevented from beating, perhaps because of inhibition of dynein detachment by force (30).

We used a mixture of 2'- and 3'-mantATP. These isomers can have different binding characteristics (31); however, we only observed one distinct recovery rate for dynein in our assay, so we cannot distinguish between the two isomers. It is known that dynein has multiple nucleotide-binding sites with distinct nucleotide-binding properties, of which one is responsible for motility (26,32–34). Previous work has shown that mantATP binds to sea urchin outer-arm dynein with dissociation constants 4 and 60 μM (26). We observed binding of mantATP to the slow-recovering site (presumed to be dynein) even at 0.1 μM mantATP. The higher affinity in our experiments might be due to the dynein being in a different chemomechanical environment while it is in the intact, immobilized axoneme.

Directly observing labeled nucleotides bound to the axoneme is a promising method for characterizing dynein activity in the intact axoneme. If the fluorescence could be resolved in space and time, the traveling wave of activation of dynein could, in principle, be measured. The biggest difficulty currently is that dynein is very selective for which nucleotides it can utilize. We found that mantATP gives very low signals. Development of new fluorophores and conjugation methods is ongoing as demand grows for new and more stable fluorescent dyes. This gives us hope that a fluorescently labeled ATP suitable for dynein, with better optical characteristics, may be available in the future.

The use of TIRF microscopy assisted us in reducing the background of diffusing labeled ATP in solution by illuminating only ~200 nm from the surface of the cover glass. The method works very well for observing axonemes that are attached to the surface. Once the axoneme is beating, it will likely be too far from the surface of the glass chamber for this method. It must also be noted that the beat of the axoneme is not planar. It is more suitable to use epifluorescence microscopy for observations of fluorescence in the beating axoneme. To minimize background noise from diffusing fluorophore, a spiking assay, using a mixture of labeled and unlabeled ATP, could be used. This will require using a fluorescent dye that allows near-single-molecule dye resolution.

There are other important aspects to consider if one is to perform spatiotemporally resolved observation of dynein activity in a beating axoneme. The dynein proteins may be active all the time. In the case that the motors are at least downregulated when not producing force, there would still be a possibility of resolving the wave of force. The motors are also likely to be alternately active on different sides of the axoneme, making it harder to resolve their activity.

Careful separation of contributions from dynein and other proteins is also important, as well as being aware of the existence of regulatory sites. Further studies on mutant axonemes with various nondenyein components missing might shed light on the source of the nondenyein components and possible kinase or nondenyein motor activity in the axoneme. We hope that with this investigation, we lay the groundwork for future successful observation of dynein in the intact beating axoneme.

APPENDIX

In this section, we solve the kinetic scheme shown in Fig. 3 under simplifying assumptions. The total concentration of binding states is

\[ [M_{tot}] = [M] + [M^*] + [MT] + [MD^*] + [MD], \]

where M corresponds to the nucleotide-binding protein in the apo state, T* is fluorescent mantADP, D* is fluorescent mantADP, T is bleached mantATP, and D is bleached mantADP. \( k_b \) is the second-order rate constant for mantATP binding to M, and \( k_u \) is the unbinding rate. For *Teirahyomena* outer-arm dynein, Holzbaur and Johnson (29) found \( k_{u,ATP} = 4 \text{μM}^{-1} \text{s}^{-1} \) and \( k_{u,ADP} = 0.15 \text{μs}^{-1} \) for ATP binding and unbinding, respectively. We assume that \([T^*]\) is constant because the TIRF illumination depth (~0.2 μm) is only a small fraction of the 100-μm depth of the flow cell, and so bleaching of the mantATP in solution will be ~500 times slower than that in the TIRF field. As a shorthand, we write \( k = k_{u[T^*]}\), which is 4 s⁻¹ at 1 μM ATP. The rate constant \( h \) combines hydrolysis and phosphate (P) release and is irreversible because the phosphate concentration in solution is negligible. For the axonemal dynein ATPase, hydrolysis is fast and reversible (rates of 100 and 30 s⁻¹, respectively), and phosphate release is also fast (80 s⁻¹) (29); we therefore denote \( MT^* \) and \( MT \) as combined ATP and ADP ⋅ P, states with \((100/(100 + 30 + 80))\times80\), \( u \) is the unbinding rate of the mantADP (\( D^* \) and D), assumed not to depend on whether mantADP is bleached or not; the reverse rate is zero because the concentrations of \( D^* \) and D in solution are negligible. \( u_{ADP} \approx 4 \text{s}^{-1} \) is the rate-limiting step for the dynein ATPase and is increased up to fivefold in the presence of microtubules (29). \( b \) is the bleaching rate, which is irreversible.

We now make the simplifying assumption that \( h \gg k_u, b, u \). This assumption is true for ATP (given that the bleaching rate is \( b = 0.6 \text{s}^{-1} \). Table 3), and will be true for mantATP provided that hydrolysis is not disproportionally decreased. With this assumption, the kinetic scheme reduces to

\[
\begin{align*}
M + T^* & \rightarrow MD^* \rightarrow M + D \\
\downarrow b & \\
MD & \rightarrow M + D
\end{align*}
\]

This has a simple solution. The steady-state fluorescence in the absence of bleaching is

\[
F_0 = \frac{[MD^*]}{[M_{tot}]} = \frac{k}{k + u}
\]
and in the presence of bleaching

\[ F_i = \frac{u}{b+u} \frac{k}{k+u} \approx \frac{u}{b+u} F_0 \]

There are two rate constants, \( k + u \) and \( b + u \), taking \( k \) large, we obtain the rate constants \( k_1 = b + u \) in the presence of bleaching and \( k_2 = u \) in the absence of bleaching (i.e., during recovery). Thus,

\[ F_i \approx \frac{k_2}{k_1} F_0, \]

as stated in the main text after Fig. 3.

**AUTHOR CONTRIBUTIONS**

M.F. and J.H. designed the experiments. M.F. preformed the experiments and image and data analysis. J.H. and M.F. constructed the model. M.M. and M.F. built the microscope setup. M.F. and J.H. designed the experiments. M.F. preformed the experiments as stated in the main text afterFig. 3.

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**REFERENCES**

1. Carvalho-Santos, Z., J. Azimzadeh, .., M. Bettencourt-Dias. 2011. Evolution: tracing the origins of centrioles, cilia, and flagella. *J. Cell Biol.* 194:165–175.
2. Nicastro, D., C. Schwartz, .., J. R. McIntosh. 2006. The molecular architecture of axonemes revealed by cryoelectron tomography. *Science.* 313:944–948.
3. Li, J. B., J. M. Gerdes, .., S. K. Dutcher. 2004. Comparative genomics identifies a flagellar and basal body proteome that includes the BBSS human disease gene. *Cell.* 117:541–552.
4. Gibbons, I. R., and A. J. Rowe. 1965. Dynein: a protein with adenosine triphosphatase activity from cilia. *Science.* 149:424–426.
5. Summers, K. E., and I. R. Gibbons. 1971. Adenosine triphosphate-induced sliding of tubules in trypsin-treated flagella of sea-urchin sperm. *Proc. Natl. Acad. Sci. USA.* 68:3092–3096.
6. Johnson, K. A. 1985. Pathway of the microtubule-dynein ATPase and the structure of dynein: a comparison with actomyosin. *Annu. Rev. Biophys. Biophys. Chem.* 14:161–188.
7. Chen, D. T. N., M. Heymann, .., Z. Dogic. 2015. ATP consumption of eukaryotic flagella measured at a single-cell level. *Biophys. J.* 109:2562–2573.
8. Brokaw, C. J. 1991. Microtubule sliding in swimming sperm flagella: direct and indirect measurements on sea urchin and tunicate spermatozoa. *J. Cell Biol.* 114:1201–1215.
9. Johnson, K. A., S. P. Marchese-Ragona, .., T. J. Chilcote. 1986. Dynein structure and function. *J. Cell Sci. Suppl.* 5:189–196.
10. Ueno, H., K. H. Bui, .., T. Ishikawa. 2014. Structure of dimeric axonemal dynein in cilia suggests an alternative mechanism for force generation. *Cytoskeleton (Hoboken).* 71:412–422.
11. Sakakibara, H., and K. Oiwa. 2011. Molecular organization and force-generating mechanism of dynein. *FEBS J.* 278:2964–2979.
12. Lin, J., and D. Nicastro. 2018. Asymmetric distribution and spatial switching of dynein activity generates ciliary motility. *Science.* 360:eaar1968.
13. Yokota, E., and I. Mabuchi. 1994. CA/dynein isolated from sea urchin sperm flagellar axonemes. Enzymatic properties and interaction with microtubules. *J. Cell Sci.* 107:353–361.
14. Bui, K. H., T. Yagi, .., T. Ishikawa. 2012. Polarity and asymmetry in the arrangement of dynein and related structures in the Chlamydomonas axoneme. *J. Cell Biol.* 198:913–925.
15. Machin, K. E. 1958. Wave propagation along flagella. *J. Exp. Biol.* 35:796–806.
16. Sartori, P., V. F. Geyer, .., J. Howard. 2016. Dynamic curvature regulation accounts for the symmetric and asymmetric beats of Chlamydomonas flagella. *ELife.* 5:e13258.
17. Lindemann, C. B. 1994. A model of flagellar and ciliary functioning which uses the forces transverse to the axoneme as the regulator of dynein activation. *Cell Motil. Cytoskeleton.* 29:141–154.
18. Brokaw, C. J. 1975. Molecular mechanism for oscillation in flagella and muscle. *Proc. Natl. Acad. Sci. USA.* 72:3102–3106.
19. Riedel-Kruse, I. H., A. Hilfinger, .., F. Jülicher. 2007. How molecular motors shape the flagellar beat. *HFSP J.* 1:192–208.
20. Inoue, Y., and C. Shingyoji. 2004. The roles of noncatalytic ATP binding and ADP binding in the regulation of dynein motile activity in flagella. *Cell Motil. Cytoskeleton.* 64:690–704.
21. Lark, E., and C. K. Omoto. 1994. Axonemes paralyzed by the presence of dyneins unable to use ribose-modified ATP. *Cell Motil. Cytoskeleton.* 27:161–168.
22. Shimizu, T., K. Furusawa, .., R. D. Vale. 1991. Nucleotide specificity of the enzymatic and motile activities of dynein, kinesin, and heavy meromyosin. *J. Cell Biol.* 110:1189–1197.
23. Omoto, C. K. 1992. Sea urchin axonemal motion supported by fluorescent, ribose-modified analogues of ATP. *J. Muscle Res. Cell Motil.* 13:635–639.
24. Feožlova, M. 2016. Probing dynein motor activity in the intact chlamydomonas axoneme. PhD thesis. Technische Universität Dresden.
25. Kikushima, K. 2009. Central pair apparatus enhances outer-arm dynein activities through regulation of inner-arm dyneins. *Cell Motil. Cytoskeleton.* 66:272–280.
26. Mocz, G., M. K. Helms, .., I. R. Gibbons. 1998. Probing the nucleotide binding sites of axonomal dynein with the fluorescent nucleotide analogue 2(3’)-O-((N-Methylanthraniloyl)-adenosine 5’-triphosphate. *Biochemistry.* 37:9862–9869.
27. Alper, J. D., M. Tovar, and J. Howard. 2013. Displacement-weighted velocity analysis of gliding assays reveals that Chlamydomonas axonemal dynein preferentially moves conspecific microtubules. *Biophys. J.* 104:1989–1998.
28. Alper, J., V. Geyer, .., J. Howard. 2013. Reconstitution of flagellar sliding. *Methods Enzymol.* 524:343–369.
29. Holzhauser, E. L., and K. A. Johnson. 1989. Microtubules accelerate ADP release by dynein. *Biochemistry.* 28:7010–7016.
30. Schmitz, K. A., D. L. Holcomb-Wygle, .., C. B. Lindemann. 2000. Measurement of the force produced by an intact bull sperm flagellum in isometric arrest and estimation of the dynein stall force. *Biophys. J.* 79:468–478.
31. Oiwa, K., J. F. Eccleston, .., D. R. Trentham. 2000. Comparative single-molecule and ensemble myosin enzymology: sulfoindocyanine ATP and ADP derivatives. *Biophys. J.* 78:3048–3071.
32. Gibbons, I. R., B. H. Gibbons, .., D. J. Asai. 1991. Multiple nucleotide-binding sites in the sequence of dynein beta heavy chain. *Nature.* 352:640–643.
33. Ogawa, K. 1991. Four ATP-binding sites in the midregion of the beta heavy chain of dynein. *Nature.* 352:643–645.
34. Tani, T., and S. Kamimura. 1999. Dynein-ADP as a force-generating intermediate revealed by a rapid reactivation of flagellar axoneme. *Biophys. J.* 77:1518–1527.