

Allosteric control of kinesin's motor domain by tubulin: a molecular dynamics study†

 Aliaksei Krukau,^{ab} Volker Knecht^{ac} and Reinhard Lipowsky^{*a}

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Molecular motors such as kinesin are essential for many biological processes. These motors have two motor domains, which bind to tubulin filaments, hydrolyze ATP, and transduce the released chemical energy into directed movements. The general principles of this chemomechanical coupling are now well-established but the underlying molecular mechanisms remain elusive because small conformational changes within large proteins are difficult to detect experimentally. Here, we use atomistic molecular dynamics simulations to monitor such changes within a single motor domain of KIF1A, which belongs to the kinesin-3 motor family. The nucleotide binding pocket of this domain can be empty or occupied by ATP or ADP. For these three nucleotide states, we determine the mobility of the backbone of the protein, both in solution and attached to tubulin. Only one subdomain of the motor domain is found to exhibit a strongly increased mobility upon binding to tubulin: the neck linker that presumably acts as a mechanical transmitter to the other motor domain in dimeric kinesin-3 motors. Furthermore, upon binding to tubulin, the neck linker mobility becomes sensitive to the bound nucleotide and is highly increased after phosphate release, which implies undocking of this linker from the core of the motor domain. These simulation results are consistent with experimental data from EPR spectroscopy, FRET, and cryo-electron microscopy. A detailed analysis of our simulation data also reveals that the undocking of the neck linker in the ADP-kinesin-tubulin state arises from allosteric interactions between the nucleotide and tubulin and that the β -sheet core undergoes a twist both during phosphate release and ATP binding. The computational approach used here can be applied to other motor domains and mechanoenzymes in order to identify allosteric interactions between the subdomains of these proteins.

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1 Introduction

Molecular motors such as kinesin that interact with cytoskeletal filaments are essential for many biological processes including intracellular cargo transport, cell motility, and cell division. These cytoskeletal motors are intriguing because they involve a wide range of scales, from the cleavage of a single covalent bond during ATP hydrolysis up to the transport over mesoscopic or even macroscopic scales. In order to elucidate the connection between these different scales, a wide variety of different experimental methods has been applied. The molecular structure of kinesin motors has been studied using X-ray crystallography^{1–5} and cryo-electron microscopy (Cryo-EM).^{4,6,7} The conformational changes within these motor proteins have been probed by

electron paramagnetic resonance (EPR) spectroscopy,^{6,8,9} Förster or fluorescence resonance energy transfer (FRET),⁶ mutagenesis,^{9,10} and biochemical methods.¹¹ The mechanical stepping of single motors can be controlled by laser traps and resolved by optical microscopy.¹² The latter methods are also useful in order to determine how multiple motors work together in the environment of the cell.¹³

From the theoretical point of view, the transport properties of single motors¹⁴ and small teams of motors^{15,16} can be understood in terms of stochastic processes that describe the motors as mechanoenzymes, which undergo both chemical transitions and mechanical steps. These theories have clarified the general principles that govern the chemomechanical coupling of the motors as well as the dependence of their transport properties on intrinsic motor parameters and external control parameters. However, these stochastic theories cannot determine the underlying molecular mechanisms, which are related to relatively small conformational changes within the large motor proteins. Here, we address these molecular mechanisms *via* atomistic molecular dynamics (MD) simulations using the motor domain of KIF1A mouse kinesin, which belongs to the kinesin-3 family, as an instructive example. In contrast to previous simulation studies,^{17–23}

^a *Theory & Bio-Systems, Max Planck Institute of Colloids and Interfaces, 14424 Potsdam, Germany. E-mail: lipowsky@mpikg.mpg.de; Fax: +49 331 5679602; Tel: +49 331 5679600*

^b *Leibniz-Rechenzentrum, Bavarian Academy of Sciences, 85748 Garching, Germany*

^c *Biomolecular Dynamics, University of Freiburg, 79104 Freiburg, Germany*

† Electronic supplementary information (ESI) available: Appendix on the choice of the Cartesian coordinate system with two Fig. S1 and S2 as well as Table S1 with neck linker displacements. See DOI: 10.1039/c3cp53367k



we will explicitly study the allosteric role of tubulin by simulating three nucleotide states of the motor domain both in solution and attached to tubulin. A quantitative comparison between these six different states reveals that, when the motor domain is bound to tubulin, the phosphate release from the nucleotide binding pocket leads to a strong increase in the mobility of the neck linker. The associated undocking of this linker from the core of the motor domain, which is essential for the motility of the kinesin-1 and related constructs,^{6,7,8} may also play an important role for the motility of both monomeric^{24,25} and dimeric kinesin-3.^{26–28} To the best of our knowledge, we report here the first simulation study that provides evidence for allosteric interactions between kinesin and tubulin or, more generally, between a molecular motor and the associated filament.

Many different kinesin motors have been identified that form a large superfamily of proteins.^{27,29} All members of this superfamily possess a motor domain. Here, we study the motor domain of KIF1A mouse kinesin as an example for a kinesin-3 motor. The latter motors can exist as monomers or homodimers.²⁷ In fact, both monomeric^{24,25} and dimeric^{26–28} KIF1A can move progressively along microtubules.

High-resolution X-ray crystal structures of kinesin's motor domain are now available for several members of the kinesin superfamily.^{1–5} The high-resolution structure of the kinesin-3 motor KIF1A is displayed in Fig. 1, based on X-ray data from ref. 5. As shown in this figure, the motor domain contains several important subdomains: the nucleotide binding pocket, the microtubule binding site, and the neck linker together with the $\alpha 6$ -helix.

The *nucleotide binding pocket* can be empty or occupied by ATP or ADP. As shown in Fig. 1, this pocket consists of several regions: the P-loop, which interacts with the phosphoryl groups of the bound nucleotide, as well as the L9-loop, which is directly connected to the $\alpha 3$ -helix; the latter two structural elements form the so-called switch-I region, which is located close to the

γ -phosphate and is involved in the cleavage of this phosphate from ATP.¹¹

The *microtubule binding site* or switch-II region is responsible for the interactions of the motor domain with the filament, and changes its affinity to tubulin upon release of phosphate or ADP from the nucleotide binding pocket. The microtubule binding site contains the L11- and L12-loops, as well as the $\alpha 4$ -helix, see Fig. 1. The L11-loop (or switch-II-loop) is located very close to the L9-loop (or switch-I-loop) of the nucleotide binding pocket, and these two loops are believed to be crucial for the cross-talk between the nucleotide binding pocket and the microtubule binding site.⁵

The *neck linker* is a short peptide segment of about 10 residues, see NL in Fig. 1, and provides a connection, together with the $\alpha 6$ -helix, between the motor domain and the neck region of dimeric kinesin-3. Mutagenesis and biochemical experiments indicate that the neck-linker determines the direction of movement of the kinesin motors along tubulin filaments.¹⁰ The conformation and mobility of the neck linker as probed by EPR spectroscopy, FRET, and Cryo-EM were found to be sensitive to the nucleotide state when kinesin-1 and related constructs were attached to tubulin but not for these motors in solution.^{6,8,9} Because of its nucleotide-dependent mobility, the neck linker should be essential for force generation^{6,9} even though the molecular displacements are expected to be small.¹²

Some atomistic molecular dynamics (MD) simulations have been performed previously in order to address the conformational changes of kinesin's motor domain in solution¹⁷ and attached to tubulin.¹⁹ The latter simulations were restricted to very short runs of only 1 ns. More recently, atomistic MD simulations were performed applying secondary structure restraints as deduced from Cryo-EM maps.²² The corresponding constrained MD trajectories had an overall run time of 42 ns. Atomistic^{20,21} and coarse-grained¹⁸ MD simulations were used to study force generation by the neck linker. Coarse-grained MD was also applied recently to the diffusion of kinesin along filaments.²³

In our MD study, both the motor domain and the tubulin dimer were modelled with atomic resolution and fully immersed in about 15 000 and 45 000 water molecules, see Methods section below. We distinguished three different nucleotide states, denoted by *ATP*, *ADP* or *Empty*, of the motor domain. Our *ATP* structures are based on AMP-PNP structures from the protein database. We first validated our simulation code by calculating the crystal structure of the motor domain in the *ATP* state and demonstrating that this computed structure is in good agreement with the crystal structure as obtained from X-ray diffraction. We then studied the different nucleotide states of the motor domain both in solution and attached to tubulin. Each of these six states was simulated for 300 ns. For the *ATP* state of the motor domain, the initial configuration of our simulations was provided by the corresponding X-ray structure, see Fig. 1. For the *ADP* and *Empty* states, we modified this initial configuration by removing the γ -phosphate and the ATP molecule, respectively, from the motor domain. Comparing the conformations of the *Empty* state with those of the *ATP* state, we can determine the conformational changes during ATP binding. Likewise, a comparison of the *ATP*

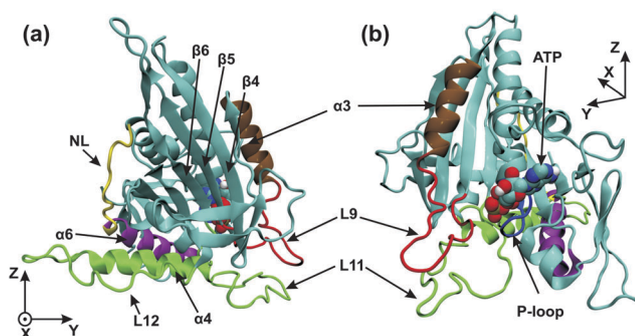


Fig. 1 (a) Front and (b) back view of the motor domain in the *ATP* state. The image is generated from a structural model based on X-ray data (PDB 1vfw) for KIF1A mouse kinesin,⁵ a kinesin-3 motor, with a docked neck linker using VMD.³⁰ The microtubule binding site (green) consists of the L11-loop, the L12-loop, and the $\alpha 4$ -helix. The nucleotide binding pocket consists of the P-loop (blue) and the L9-loop (red), which is directly connected to the $\alpha 3$ -helix (ochre). The last subdomain at the C-terminal end of the motor domain is the neck linker (NL, yellow), which is preceded by the $\alpha 6$ -helix (purple). The C-terminus of the $\alpha 6$ -helix is in close proximity to the C-terminus of the $\alpha 4$ -helix. The coordinates x , y , and z are used to describe the conformational changes of the motor domain.



state with the *ADP* state provides insight into the conformational changes associated with phosphate release.

Using this MD set-up, we first studied the mobility of the 361 C_{α} -atoms of KIF1A's motor domain by calculating the positional fluctuations of these atoms. Essentially, all of these fluctuations are found to remain unchanged or become reduced when the motor domain binds to tubulin, with one remarkable exception: the neck linker, which exhibits an essentially constant mobility both in solution and in the *Empty* and *ATP* states of the kinesin-3-tubulin complex but a strongly *enhanced* mobility in the *ADP*/kinesin-3/tubulin state. This enhanced mobility of the neck linker implies an undocking of this linker from the core of the motor domain, in agreement with experimental data from EPR spectroscopy, FRET, and Cryo-EM.^{6–8} Furthermore, a detailed analysis of our simulation data also shows that the enhanced mobility of the neck linker arises from rotational movements of the $\alpha 4$ -helix (or switch-II-helix), which belongs to the microtubule binding site, and the $\alpha 6$ -helix, which is directly connected to the neck linker. These helix rotations are caused by L11–L9 loop interactions, which are modified because the L9-loop (or switch-I-loop) moves away from the nucleotide during phosphate release. In this way, our simulations provide direct evidence for an allosteric coupling mechanism that leads from phosphate release to neck linker undocking *via* a sequence of conformational changes involving several subdomains, namely the L9 loop, the L11 loop, the $\alpha 4$ -helix, and the $\alpha 6$ -helix.

Our paper is organized as follows. In the “Results” section, we first describe the mobility of the different subdomains for the three nucleotide states of the motor domain, both in solution and attached to tubulin. We show that the neck linker of *ADP*-kinesin-3 exhibits a strongly enhanced mobility upon binding to tubulin. Next, we determine the rotations of the $\alpha 6$ - and $\alpha 4$ -helices in the six different states of the motor domain and show that these rotations are quite different for the motor domain in solution and attached to tubulin. We also find a tilt of the strand orientations within the β -sheet core, both during phosphate release and *ATP* binding. Finally, we study the interactions between the L9- and L11-loops for the kinesin-3-tubulin complex and find that the L9-loop moves away from the nucleotide during phosphate release. In the “Discussion” section, we compare our results with available experimental data and argue that our simulation results provide direct evidence for tubulin's allosteric control of the chemo-mechanical coupling between phosphate release and neck linker undocking. The Methods section at the end contains a detailed description of our computational approach.

2 Results

2.1 Subdomain mobilities of kinesin-3 in solution and attached to tubulin

The motor domain of KIF1A mouse kinesin (hereafter called ‘kinesin-3’) consists of 361 residues and, thus, of 361 C_{α} -atoms that define the protein's backbone. In this subsection, we will first discuss the mobility of these C_{α} -atoms by calculating the standard deviations or root-mean-square-fluctuations (rmsf) of

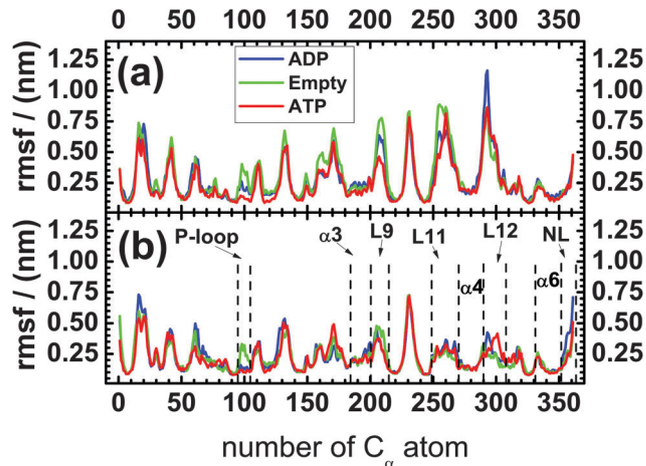


Fig. 2 Standard deviations or root-mean-square-fluctuations (rmsf) of the 361 C_{α} -atoms of the kinesin's backbone (a) in solution and (b) attached to a tubulin dimer for the three nucleotide states *ADP* (blue), *Empty* (green), and *ATP* (red). The x-axis is labeled by the numbers of the C_{α} -atoms: the first C_{α} -atom with number 1 corresponds to the N-terminus of the motor domain, the last C_{α} -atom with number 361 represents the C-terminus of this domain and of the neck linker. The different loops (P-loop, L9, L11, L12) and helices ($\alpha 3$, $\alpha 4$, $\alpha 6$) as well as the neck linker (NL) have been introduced in Fig. 1.

their coordinates. These fluctuations are displayed in Fig. 2(a) and (b) for kinesin-3 in solution and attached to tubulin, respectively. Inspection of this figure reveals that the overall fluctuations of the backbone of kinesin-3 are strongly reduced upon binding to tubulin. The largest decrease is observed for the microtubule binding site, which is located between the C_{α} -numbers 250 and 310. The mobility of the L11- and L12-loops of kinesin-3 attached to tubulin is about half of that in solution, and the mobility of the $\alpha 4$ -helix drops by 25% from 0.2 to 0.15 nm upon attachment to tubulin.

2.1.1 Mobility of the nucleotide binding pocket. Another important subdomain, for which fluctuations of the backbone depend both on the tubulin binding and on the nucleotide state, is the nucleotide binding pocket. The L9-loop of kinesin-3 in the *Empty* nucleotide state reduces its mobility from 0.8 to 0.45 nm upon attachment to tubulin. A large decrease in mobility from 0.65 to 0.45 nm is observed for this loop when kinesin binds to tubulin with an *ADP* molecule in its binding pocket. In contrast, the mobility of the L9-loop undergoes only relatively small changes from 0.45 to 0.4 nm upon binding to tubulin when kinesin is in the *ATP* nucleotide state. The fluctuations of the $\alpha 3$ -helix do not depend on the nucleotide but on the tubulin binding state. Indeed, the mobility of the $\alpha 3$ -helix is reduced by 20% from 0.25 to 0.2 nm when the motor domain attaches to tubulin. Fluctuations of the P-loop, which also belongs to the nucleotide binding pocket, depend both on the binding state and on the nucleotide state. Upon binding to tubulin, the mobility of the P-loop in the *Empty* and *ADP* states is reduced from 0.45 to 0.35 nm and from 0.25 to 0.15 nm, respectively. In contrast, the mobility of the P-loop remains the same and equal to 0.15 nm both in solution and attached to tubulin when kinesin contains *ATP* in its nucleotide binding pocket.



2.1.2 Mobility of neck linker. The mobility of the neck linker as observed in the simulations behaves in an unexpected manner. Indeed, the standard deviation of the neck linker has the constant value of 0.45 nm (i) for all nucleotide states of kinesin in solution and (ii) for tubulin-bound kinesin both in the *ATP* and in the *Empty* state, whereas the mobility of the neck linker is *increased* to 0.75 nm in the *ADP* nucleotide state of tubulin-bound kinesin. In contrast, all other subdomains exhibit a mobility that remains essentially unchanged or is *reduced* when the motor domain binds to tubulin.

In order to corroborate this peculiar behavior of the neck linker, we also performed a principal component analysis of its mobility. The outcome of this analysis for the C_α -atoms of the neck linker is shown in Fig. 3. In solution, we found small differences in the first eigenvalue, but these differences are smaller than the statistical error, see Fig. 3(a). Therefore the mobility of the neck linker in solution is essentially independent of the nucleotide state. In contrast, the fluctuations of the neck linker depend strongly on the nucleotide state when the motor domain is attached to tubulin as shown in Fig. 3(b). The first eigenvalue for the *ADP* state is now 2.3 and 3.7 times larger than for the *Empty* and *ATP* nucleotide states, respectively. In fact, all first four eigenvalues are 2–3 times larger for the *ADP* compared to the *ATP* or *Empty* states. Upon binding to tubulin, the first eigenvalue is increased by a factor 2.1 and 1.4 for the *ADP* and the *Empty* states and decreased by a factor 1.4 for the *ATP* state, as follows from a comparison of the values in Fig. 3(a) and (b).

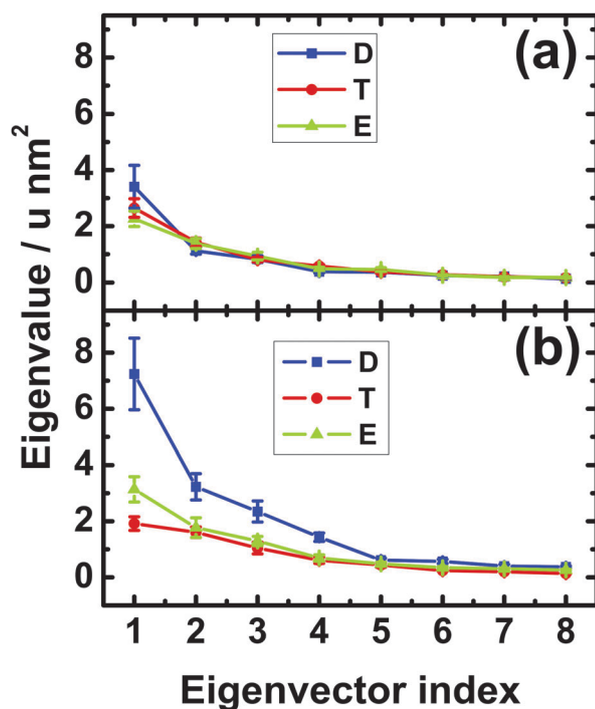


Fig. 3 Eigenvalues of the covariance matrix for the C_α atoms of the neck linker region, residues 351–361, for the motor domain (a) in solution and (b) attached to a tubulin dimer. The eigenvalues for the *ADP*, *ATP*, and *Empty* nucleotide states are displayed in blue (D), red (T), and green (E), respectively. The parameter u denotes the atomic mass unit.

Therefore, for the motor domain bound to tubulin, both the root-mean-square-fluctuations in Fig. 2(b) and the eigenvalues of the principal component analysis in Fig. 3(b) imply that the mobility of the neck linker is hardly changed during ATP binding whereas this mobility is strongly increased during phosphate release. The latter increase corresponds to an undocking of the neck linker from the core of the motor domain.

For the motor domains of kinesin-1^{20,21} and kinesin-5,³¹ the neck linker interacts with the N-terminal cover strand, located between the C_α -numbers 0 and 9, and forms a cover neck bundle upon ATP binding. We did not analyze this structural feature in the present study but, for kinesin-3 attached to tubulin, the mobility of the N-terminal cover strand is strongly reduced for the *ATP* compared to the *Empty* state; see Fig. 2(b), as one would expect if a cover neck bundle is formed during ATP binding.

2.2 Rotations of $\alpha 4$ - and $\alpha 6$ -helices in solution and attached to tubulin

As shown in Fig. 1, the last subdomain of kinesin-3 that precedes the neck linker is the $\alpha 6$ -helix which is in close contact with the $\alpha 4$ -helix, the latter being part of the microtubule binding site. Therefore, we have also analyzed the conformation of the $\alpha 4$ - and the $\alpha 6$ -helices to study possible interactions^{4,5} between the microtubule binding site and the neck linker.

In order to visualize the dependence of the molecular structure of kinesin-3 on its nucleotide state, we have compared the dominant conformations of the *ATP* and *ADP* states as well as those of the *ATP* and *Empty* states. For a given pair of states, this comparison was performed by superimposing their β -sheet cores and identifying the rotations of the $\alpha 4$ - and $\alpha 6$ -helices. In order to describe these rotations, we used the Cartesian coordinate system (x, y, z) as shown in Fig. 1 and described in more detail in the Appendix and Fig. S1 and S2, see the ESI.†

2.2.1 Kinesin-3 in solution. The dominant conformations of the $\alpha 4$ - and $\alpha 6$ -helices as obtained from our simulations for kinesin's motor domain in solution are shown in Fig. 4, where the red, blue, and green color correspond to the *ATP*, *ADP*, and *Empty* states, respectively.

Relative to its orientation in the *ATP* state (red), the $\alpha 4$ -helix in the *ADP* state (blue) is rotated by 14° around the x -axis as shown in Fig. 4(a). Thus, the latter rotation is observed within the yz -plane. In contrast to the $\alpha 4$ -helix, the $\alpha 6$ -helix does not undergo any significant rotation in the yz -plane during the transition from the *ATP* to the *ADP* state. Furthermore, when projected onto the xy -plane, neither the $\alpha 4$ - nor the $\alpha 6$ -helices change their orientation, see Fig. 4(b).

In Fig. 4(c) and (d), the orientations of the $\alpha 4$ - and $\alpha 6$ -helices for the *Empty* state (in green) are shown together with those for the *ATP* state (in red). When the motor domain binds ATP, the $\alpha 4$ -helix rotates by about 13° within the yz -plane, see Fig. 4(c), whereas both the $\alpha 4$ - and the $\alpha 6$ -helices remain essentially unchanged when projected onto the xy -plane, see Fig. 4(d). Furthermore, inspection of all four subfigures Fig. 4(a)–(d) shows that the helix orientations in the *Empty* state closely resemble those in the *ADP* state.

2.2.2 Kinesin-3 attached to tubulin. The dominant conformations of the $\alpha 4$ - and $\alpha 6$ -helices of kinesin-3 attached to tubulin are



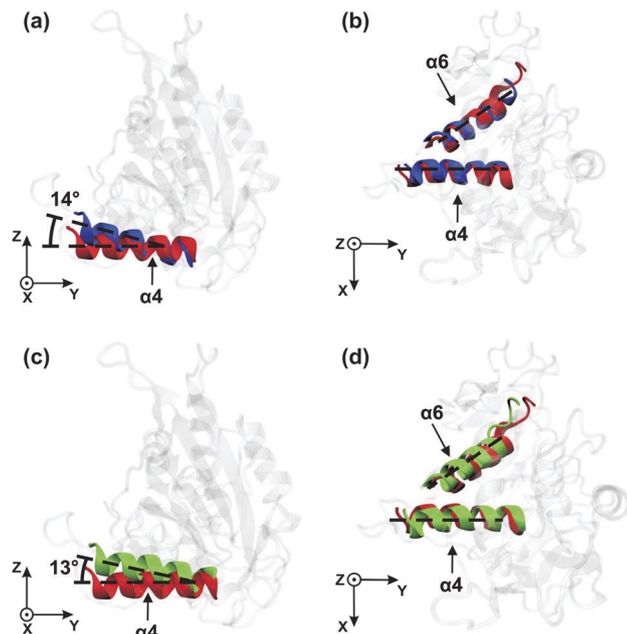


Fig. 4 Kinesin-3 in solution: dominant orientations of the $\alpha 4$ - and $\alpha 6$ -helices for the three nucleotide states ATP (red), ADP (blue), and Empty (green) when projected onto the yz - and the xy -plane: (a and b) ATP cleavage and phosphate release (red \rightarrow blue) and (c and d) ATP binding (green \rightarrow red). In both processes, the helices undergo pronounced rotations parallel to the yz -plane as depicted in (a) and (c).

shown in Fig. 5. The comparison of the ATP state (red) with the ADP state (blue) in Fig. 5(a) and (b) now reveals that, in contrast to the solution case, the $\alpha 4$ -helix is no longer rotated within the yz -plane, but both the $\alpha 4$ - and the $\alpha 6$ -helices are now rotated within the xy -plane, which is tangential to the surface of the microtubule. More precisely, the $\alpha 4$ - and the $\alpha 6$ -helices are rotated by 13° and 15° , respectively, in the ADP state relative to the corresponding orientation in the ATP state, see Fig. 5(b).

In Fig. 5(c) and (d), the orientations of the helices are compared for the Empty state (green) with those for the ATP state (red). Upon binding of ATP, the $\alpha 4$ -helix undergoes relatively small rotations in the yz -plane and in the xy -plane, respectively, while the $\alpha 6$ -helix undergoes no appreciable rotation in the yz -plane and a rotation by 10° in the xy -plane. In order to directly determine the effect of these helix rotations onto the neck linker, we have also measured the neck linker's average displacement and its root-mean-square-fluctuations in the different nucleotide states, see Table S1 in the ESI.†

Very recently, it has been proposed^{32,33} that the β -sheet core of kinesin-1 may undergo a twist during nucleotide release, similar to myosin motors.³⁴ In order to find out whether such twists may occur in kinesin-3, we have studied the relative orientations of neighboring β -strands within the β -sheet core of this motor, as explained in the Methods section. For kinesin-3 attached to tubulin, we find that the β_4 -strand is rotated by about 10° relative to its neighboring strands β_3 and β_5 , both during phosphate release and during ATP binding, see Table 1.

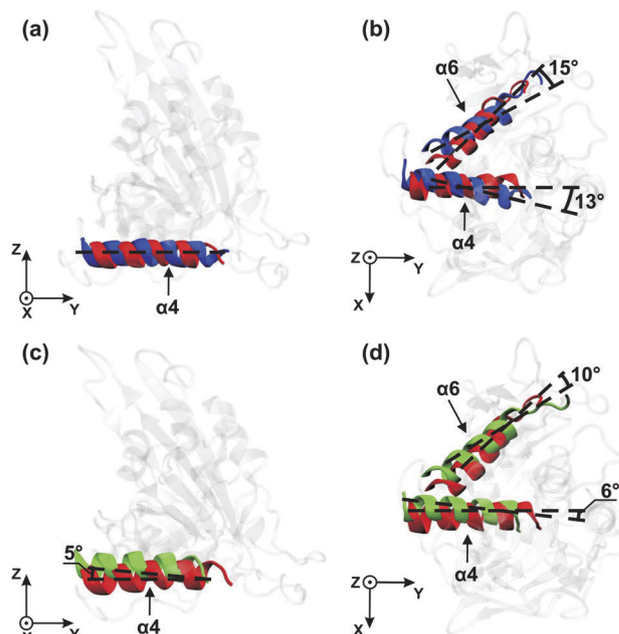


Fig. 5 Kinesin-3 attached to tubulin: dominant orientations of the $\alpha 4$ - and $\alpha 6$ -helices for the three nucleotide states ATP (red), ADP (blue), and Empty (green) when projected onto the yz - and xy -planes: (a and b) ATP cleavage and phosphate release (red \rightarrow blue) and (c and d) ATP binding (green \rightarrow red). In both cases, the helices undergo pronounced rotations parallel to the xy -plane as depicted in (b) and (d).

Table 1 Average tilt angles ϕ_{ij} between neighboring strands β_i and β_j of the β -sheet core for kinesin-3 attached to tubulin in the three nucleotide states ATP, ADP, and Empty, together with the associated standard errors. The tilt angles ϕ_{34} and ϕ_{45} change by about 10° , both during phosphate release and ATP binding, whereas the tilt angle ϕ_{56} is essentially independent of the nucleotide state

| State | ϕ_{34} ($^\circ$) | ϕ_{45} ($^\circ$) | ϕ_{56} ($^\circ$) |
|-------|--------------------------|--------------------------|--------------------------|
| ATP | 20 ± 1.0 | 10 ± 0.7 | 16 ± 1.3 |
| ADP | 8.6 ± 0.5 | 18 ± 1.0 | 17 ± 1.2 |
| Empty | 9.3 ± 2.0 | 19 ± 1.8 | 15 ± 1.8 |

2.3 Interactions of L9- and L11-loops in kinesin-3 attached to tubulin

One possible pathway for the mechanical transmission of conformational changes from the nucleotide binding pocket to the microtubule binding site is provided by the interactions between the L9-loop, which is located in the vicinity of the γ -phosphate of the ATP, and the L11-loop, which is attached to the N-terminus of the $\alpha 4$ -helix, the latter being part of the microtubule binding site, see Fig. 1. The dominant conformations of the L9- and L11-loop in the ATP and the ADP nucleotide state are displayed in Fig. 6(a) and (b), respectively. In the ATP nucleotide state, essentially the complete L9-loop is found to be close to the γ -phosphate and probably participates in the ATP hydrolysis. In addition, the central part of the L9-loop has an α -helical conformation and interacts with the N-terminal part of the L11-loop. At the same time, the C-terminal part of the L9-loop interacts with the N-terminal part of the $\alpha 4$ -helix, see Fig. 6(a). In the ADP nucleotide state, on the other hand, the



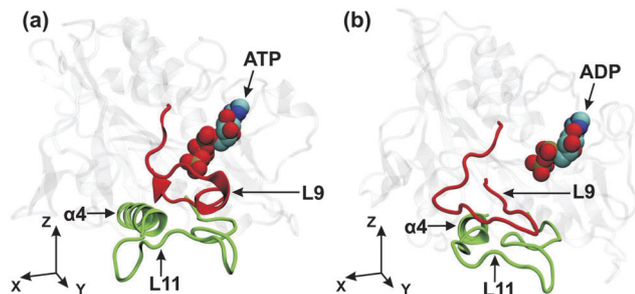


Fig. 6 Kinesin-3 attached to tubulin: dominant conformations of the L9-loop (red) and the L11-loop (green) for (a) the *ATP* and (b) the *ADP* states of the motor domain. As before, the (x,y) -plane is tangential to the tubulin surface and the y -axis is parallel to the long principal axis of the $\alpha 4$ -helix in the *ATP* state. Comparison of the two conformations reveals that the γ -phosphate release from the nucleotide binding pocket causes the L9-loop to move away from the nucleotide, to change its interaction pattern with the L11-loop, see the corresponding atomic separations as described in Fig. 7 below, and to induce a rotation of the $\alpha 4$ -helix within the (x,y) -plane as in Fig. 5(b).

L9-loop changes its conformation, moving away from the nucleotide, and the central part of this loop starts to interact with the central part of the L11-loop, which induces the rotation of the $\alpha 4$ -helix in the xy -plane, see Fig. 6(b).

To describe the observed conformational changes between the *ATP* and the *ADP* state in a more quantitative manner, we calculated the separations of different atoms within these two regions of kinesin-3 attached to tubulin, see Fig. 7. In the *ATP* nucleotide state, contacts are found between the peptide segments Ala²⁰⁶–Arg²¹⁶ of L9 and Ala²⁵⁵–Lys²⁶¹ of L11, between the segments Thr²¹³–Arg²¹⁶ of L9 and Ala²⁴⁸–Ala²⁵⁵ of L11, as well as between the segments Ser²¹⁵–Arg²¹⁶ of L9 and Asn²⁷²–Thr²⁷⁶ of the $\alpha 4$ -helix, as indicated by the purple circles in Fig. 7(a). In addition, several very stable hydrogen bonds were observed

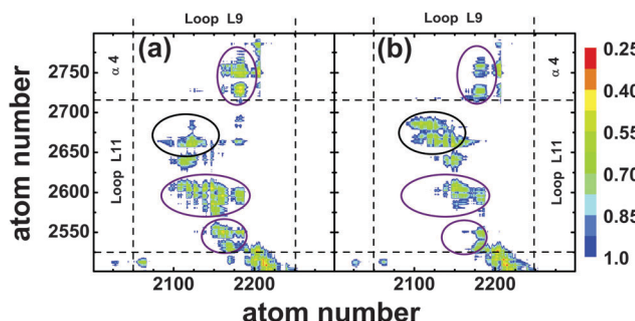


Fig. 7 Kinesin-3 attached to tubulin: distance between the atoms of the L9-loop, which belongs to the nucleotide binding pocket, and the L11-loop, which is part of the microtubule binding site, for (a) the *ATP* and (b) the *ADP* nucleotide states. The color code represents the average distance between two atoms in nm. In the *ATP* state, the central part of the L9-loop interacts with the N-terminal part of the L11-loop (lower purple ellipses) and the C-terminal part of the L9-loop interacts with the N-terminal part of the $\alpha 4$ -helix (upper purple ellipses). All of these interactions are strongly reduced in the *ADP* state, which exhibits, however, enhanced interactions between the inner part of the L9-loop and the C-terminal part of the L11-loop (black ellipses).

between L9-Thr²¹³ and L11-Ala²⁵⁵ (existence probability 0.861), L9-Ser²¹⁴ and L11-Glu²⁵³ (0.962), as well as L9-Arg²¹⁶ and L11-Asn²⁷² (0.721). The last hydrogen bond connects the C-terminus of the L9-loop to the N-terminus of the $\alpha 4$ -helix, and thus may stabilize the orientation of the helix. When ATP is replaced by ADP in the binding pocket, almost all contacts present in the *ATP* state disappear, see purple circles in Fig. 7(b). Furthermore, all hydrogen bonds that were formed in the *ATP* state are destroyed as well. However, a region of stable contact is observed between the peptide segments Ala²⁰⁷–Thr²¹³ of L9 and Arg²⁶⁴–Ala²⁶⁹ of L11, see red circle in Fig. 7(b). In addition, two stable hydrogen bonds are found between L9-Asn²⁰⁹ and L11-Glu²⁶⁷ (0.95) as well as L9-Met²¹⁰ and L11-Glu²⁶⁷ (0.883).

Thus, after the release of the γ -phosphate, conformational changes in the L9-loop induce the breaking of several hydrogen bonds between the loops, destabilizing the *ATP* state orientation of the $\alpha 4$ -helix. At the same time, the appearance of contacts between the central segments of the loops facilitate rotation of the $\alpha 4$ -helix, and hydrogen bonds between residues in these segments stabilize the new orientation of this helix. The complete removal of the ATP from the binding pocket leads to a three-fold decrease in the stability of the hydrogen bonds between the termini of the L9- and L11-loops. However, for this *Empty* state, the appearance of new contacts between any segments of the loops is not observed. Therefore, the destabilization of hydrogen bonds between the termini of the L9- and L11-loops in the *Empty* state, without the creation of contacts between their central parts, leads to a rotation of the $\alpha 4$ -helix with a magnitude only half of that in the *ADP* state.

3 Conclusions

In the molecular dynamics simulations reported here, we studied the motor domain of KIF1A mouse kinesin and focussed on those subdomains that should play an important role for free energy transduction and motility of kinesin-3 motors: the nucleotide binding pocket, the microtubule binding site, and the neck linker. First, we determined the mobility of these subdomains as described by the positional fluctuations of the C_{α} -atoms of the protein's backbone.

For the motor domain in solution, we do not observe significant differences in the mobility of the neck linker for the three nucleotide states, see Fig. 2(a) and Fig. 3(a). This independence of the mobility on the nucleotide state agrees with the results of EPR⁶ and FRET measurements.^{6,8} Comparing our simulation data on the mobility of the neck linker in solution and attached to tubulin, see the data for NL in Fig. 2(a) and (b), we find that fluctuations of the neck linker in the *ADP* nucleotide state almost double in amplitude upon binding to tubulin. In contrast, the fluctuations of the neck linker in the *ATP* state are slightly reduced when the motor domain is attached to tubulin. The increase in the mobility of the neck linker in the *ADP* relative to the *ATP* state, when the motor domain is attached to tubulin, again agrees with Cryo-EM and EPR data.^{6,9} In fact, in our simulations, the neck linker is found



to be the *only* region within the motor domain, for which tubulin binding leads to an *increased* mobility, see Fig. 2.

Furthermore, when the motor domain is attached to tubulin, the mobility of the neck linker is slightly higher in the *Empty* state compared to the *ATP* nucleotide state, see Fig. 3(b). This result correlates well with Cryo-EM experiments,⁷ which show that the neck linker was docked onto the motor head in the *ATP* state, but did not have any pronounced structure in the *Empty* state.

One important result of our simulation study is the remarkable difference between the conformational changes of the motor domain in solution and attached to tubulin. In solution, the $\alpha 4$ -helix is found to rotate in the (y , z)-plane during γ -phosphate release, *i.e.*, during the transition from the *ATP* to the *ADP* state, see Fig. 4(a), which agrees with experiment as previously mentioned. However, inspection of Fig. 4(b) and (d) also shows that we do not observe any rotation of the $\alpha 6$ -helix. Therefore, in solution, we find no evidence for a direct coupling between the $\alpha 4$ -helix, which belongs to the microtubule binding site, and the $\alpha 6$ -helix, which is directly connected to the neck linker.

However, when the motor domain is attached to tubulin, the helices are found to behave in a completely different way. First, during phosphate release, the $\alpha 4$ -helix no longer rotates in the yz -plane, but rather in the xy -plane, which is tangential to the tubulin surface. Most likely, the rotation of the $\alpha 4$ -helix within the yz -plane is suppressed by the steric hindrance provided by tubulin. Second, and even more importantly, we now observe a large rotation of the $\alpha 6$ -helix during phosphate release, see Fig. 5(b). In this way, we observe direct crosstalk between the microtubule binding site and the neck linker when the motor domain is attached to tubulin.

We also analyzed the distance between atoms of the L9-loop, which belongs to the nucleotide binding pocket, and those of the L11-loop, which is part of the microtubule binding site. When the motor domain is attached to tubulin, the interactions between these loops are found to depend on the nucleotide within the binding pocket.

In the *ATP* state, the central and the C-terminal parts of the L9-loop are in contact with the N-terminal part of the L11-loop and the $\alpha 4$ -helix, respectively, as shown in Fig. 6(a). The loops are connected by several stable hydrogen bonds. However, the release of the γ -phosphate from the nucleotide binding pocket induces a change in the conformation of L9, the disappearance of the contacts between the L9-loop and the termini of the L11-loop and the $\alpha 4$ -helix, as well as the destruction of the previously existing hydrogen bonds between the loops, see Fig. 6(b). These changes lead to an increased mobility of the L11-loop and the $\alpha 4$ -helix. At the same time, the appearance of contacts between the central regions of the L9- and L11-loops, as described in Fig. 7(b), facilitates the large rotation of the $\alpha 4$ -helix, as observed in the (xy)-plane tangential to the tubulin surface.

The release of the complete *ATP* molecule from the binding pocket, on the other hand, leads only to a destabilization of the hydrogen bonds between the termini of the two loops and to a small rotation of the $\alpha 4$ -helix in the *Empty* state, see Fig. 5(c) and (d). This rotation increases the mobility of the neck linker

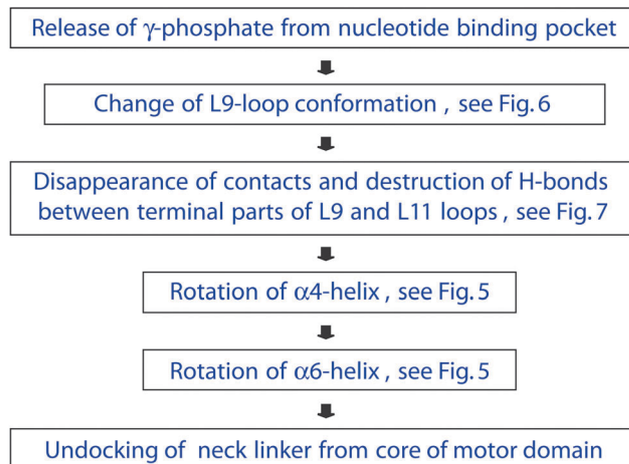


Fig. 8 Chemomechanical coupling between phosphate release and neck linker undocking for kinesin-3 attached to tubulin as mediated by allosteric interactions between the motor's subdomains. First, phosphate release affects the L9-loop conformation, which moves away from the nucleotide and loses many contacts with the L11-loop. These reduced interactions between the two loops rotate the $\alpha 4$ -helix and, in this way, modify the molecular interactions between the $\alpha 4$ - and the $\alpha 6$ -helices. As a result, the $\alpha 6$ -helix undergoes a substantial rotation as well and undocks the neck linker from the motor domain's core. These allosteric interactions are absent for the motor domain in solution, see Fig. 4.

only slightly, compared to the *ADP* state, and does not lead to its complete undocking from the core of the motor domain.

The conformational changes of kinesin-3 during phosphate release as deduced from our simulations of the *ATP* and *ADP* state are consistent with the scheme shown in Fig. 8. First, phosphate release leads to a conformational change of the L9-loop, which moves further away from the bound nucleotide, see Fig. 6, and modifies the interactions of this loop with the L11-loop. The latter loop is directly connected to the $\alpha 4$ -helix and then acts to rotate this helix. Furthermore, because the C-terminus of the $\alpha 4$ -helix is in close proximity to the C-terminus of the $\alpha 6$ -helix, the rotation of $\alpha 4$ leads to a rotation of $\alpha 6$. The latter rotation generates a force on the N-terminus of the neck linker, which is then pulled away from its docked position, see the measured displacements in Table S1 (ESI[†]) and increases the neck linker's mobility, see Fig. 2.

It is interesting to note that our chemomechanical coupling scheme as summarized in Fig. 8 predicts that the rotation of the $\alpha 4$ helix *precedes* the undocking of the neck linker. The rotation of the $\alpha 4$ helix, which is part of the microtubule binding site, may act to release the motor domain from its strongly bound *ATP* state. Such a sequence of conformational transitions seems to be consistent with the motility model for monomeric KIF1A as reviewed in ref. 25. In the latter model, the KIF1A motor becomes loosely bound to the microtubule *before* it undergoes a conformational change that biases its diffusion towards the plus end of the filament.

We also determined the strand orientations within the β -sheet core for the different nucleotide states. For kinesin-3 attached to tubulin, we found a twist within this core, both during phosphate release and during *ATP* binding, see Table 1.



Since the members of the kinesin-3 family have motor domains that are close structural homologs, the allosteric coupling mechanism found here for KIF1A mouse kinesin should apply to other kinesin-3 motors as well. In addition, the recent observation^{26–28} that KIF1A motors can form dimers *in vivo* indicates that a similar allosteric coupling mechanism may even apply to other members of the kinesin superfamily but this remains to be studied. Apart from such an extension to different kinesins, our computational approach can be applied to other motor domains and mechanoenzymes as well because it is based on a relatively simple strategy: consider the mechanoenzyme in different states that are distinguished by their ligands or binding partners and determine, for these different states, the mobility of important subdomains within the protein. Ligand-induced changes in the mobilities will then reveal allosteric interactions between these subdomains and the ligands. Other possible targets for future applications of our computational approach include the rotary motor F_1 -ATPase³⁵ and the actin-based stepping motor myosin V.³⁶

4 Methods

4.1 Crystal structure of KIF1A's motor domain and code validation

The motor domain of KIF1A mouse kinesin consisting of 361 residues was studied in explicit water using MD simulations with atomic resolution. In order to validate our computational method, we first simulated the crystal structure of this protein in the *ATP* state and compared it with the corresponding X-ray crystal structure,⁵ protein data bank id 1VFW. The missing loops, L11 and L12, of KIF1A corresponding to residues 252–266 and 287–300, respectively, were modeled using the ModLoop server.^{37,38} The complete unit cell of the crystal, which contains four copies of the motor domain and 5139 water molecules, see Fig. 9, was constructed using the Computational Crystallography Toolbox (cctbx) plug-in for PyMOL.³⁹ Using the X-ray crystal structure as the initial configuration, we performed 10 simulation runs with a 4 fs time step in an NPT ensemble at temperature $T = 350$ K and pressure $P = 1$ bar. Each run took 30 ns, which implies a total simulation time of 300 ns.

In order to estimate the overall accuracy of our simulations, we saved the simulated configurations of the crystal every 2 ps which leads to a total number of 15×10^4 configurations. Using these configurations, we calculated the root-mean-square deviation (RMSD) of the simulated crystal from the X-ray crystal structure, averaging over all C_α atoms of the protein. This RMSD value was found to be 0.25 ± 0.02 nm, where the standard deviation was calculated using the 40 protein copies used in the 10 simulation runs. Thus, within the accuracy of this computation, the RMSD value of the simulated crystal is compatible with the spatial resolution of 0.23 nm as estimated for the X-ray crystal structure.⁵

For the motor domain in aqueous solution and attached to tubulin, we found characteristic rotations of the α_4 and α_6 helices. In order to estimate the accuracy of these simulation results, we also determined the RMSD values for the ends of

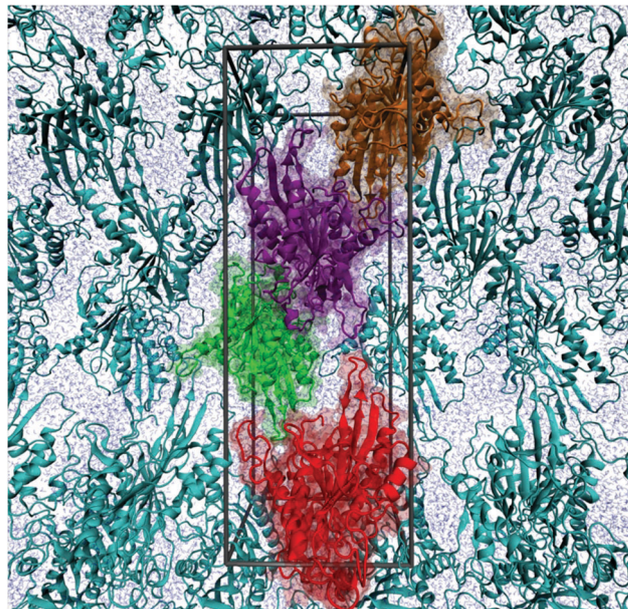


Fig. 9 Crystal structure of KIF1A mouse kinesin, protein data bank id 1VFW. The simulation box (black frame) is identical with the unit cell, which is a rectangular cuboid and contains four copies of the protein. The unit cell has the dimensions 4.26 nm \times 5.55 nm \times 15.67 nm. All protein molecules are displayed in ribbon representation. In addition, the four copies within the simulation box are depicted in transparent spacefilling representation and distinguished by four different colors. The blue background is provided by water molecules.

these helices from the crystal structure simulations. The N- and C-termini of the α_4 -helix were found to have the RMSD values of 0.26 ± 0.07 nm and 0.32 ± 0.1 nm, respectively. Likewise, the RMSD values for the N- and C-termini of the α_6 -helix were found to be 0.19 ± 0.04 nm and 0.16 ± 0.03 nm. Thus, the RMSD values of the ends of these two helices are also compatible with the spatial resolution of the X-ray crystal structure.

We also analyzed the orientations of the α_4 - and α_6 -helices in our crystal simulations. We found that the α_4 - and α_6 -helices of the simulated crystal structure were rotated by $4^\circ \pm 1.6^\circ$ and $4.6^\circ \pm 1.5^\circ$, respectively, compared to the X-ray crystal structure. Such a rotation angle is again compatible with the spatial resolution of 0.23 nm for the X-ray structure. Indeed, if we rotate the α_4 -helix, which has a length of 2.15 nm, around one of its ends by 4° , the other end of the helix is displaced by 0.15 nm. The same consideration applies to the α_6 -helix with a length of 2.18 nm.

4.2 Motor domain in solution

In solution, we studied the *ATP*, *ADP*, and *Empty* states of the motor domain, corresponding to bound ATP, bound ADP, and no bound nucleotide. As an initial configuration for the *ATP* state, we used again the X-ray structure of KIF1A mouse kinesin in the *ATP* state,⁵ protein data bank id 1VFW. As in the crystal structure simulations, the missing loops, L11 and L12, of the motor domain corresponding to residues 252–266 and 287–300, respectively, were modeled using the ModLoop server.^{37,38} The protein was placed in a dodecahedral box of initial edge length



9 nm containing 15 263 simple point charge (SPC) water molecules⁴⁰ and 4 Cl⁻ ions to ensure electroneutrality of the system. The simulations were performed using periodic boundary conditions. The structure of the missing loops was energy minimized for 10⁶ steps *via* steepest descent and equilibrated for 5 ns with a 1 fs time step in an NPT ensemble at temperature $T = 300$ K and pressure $P = 1$ bar. Afterwards the system was simulated at $T = 350$ K for 60 ns and a structure cluster analysis based on the Daura algorithm⁴¹ was performed to find the dominant conformation of the missing loops.

The ADP and Empty nucleotide states of the motor head were created from its ATP state by removing the γ -phosphate and ATP, respectively, from the nucleotide binding pocket. Both ATP and ADP were complexed with Mg²⁺ ions. Depending on the nucleotide state, three to five Cl⁻ ions were placed in the boxes to ensure electroneutrality of the systems. These systems were energy minimized for 10⁶ steps *via* steepest descent and then equilibrated for 10 ns with a 1 fs time step in an NPT ensemble at $T = 300$ K and $P = 1$ bar. To achieve better sampling of conformational space, each system was simulated 10 times using different initial velocities. Every production run continued 30 ns with a 4 fs time step and configurations were saved every 10 ps. So each system was simulated for a total time of 300 ns and 3×10^4 configurations were available for analysis. The production runs were conducted in an NPT ensemble at $T = 350$ K and $P = 1$ bar. The GROMOS43a1 force field was used for the motor domains and the ions. The GROMOS43a1 parameter set was also used to model ATP and ADP, apart from the partial charges which were taken from.⁴² A spherical cutoff of 1 nm was used for the short-range Lennard-Jones and electrostatic interactions; long range electrostatic interactions were taken into account by particle mesh Ewald summation.⁴³ The Berendsen scheme for temperature and pressure control⁴⁴ with the relaxation times of 0.1 ps and 1 ps, respectively, was used, and all simulations were performed using the GROMACS software package.⁴⁵

4.3 Motor domain attached to tubulin

One motor domain of KIF1A mouse kinesin in the ATP state attached to a tubulin dimer,⁴ protein data bank id 2hxf, was taken as the initial configuration for the simulations. The missing loops L11 and L12 of the motor domain, and the missing regions of the E-hook of tubulins, corresponding to residues 440–451 and 438–455 for α - and β -tubulin, respectively, were modeled using the scheme described above. One example for the resulting configuration of the kinesin-3-tubulin complex is shown in Fig. S1 in the ESI.† The Empty and ADP states of the motor domain were created as in solution. The protein complexes were placed in dodecahedral boxes of initial edge length 13 nm with 45 732 SPC water molecules in every box. Depending on the nucleotide state 48–50 Na⁺ ions were added to obtain electrically neutral systems, which were energy minimized, equilibrated and simulated using the protocol described above. Tubulin, GTP, and GDP were modeled using the GROMOS43a1 force field.

4.4 Analysis methods

Dominant conformations were obtained from a structure cluster analysis using the algorithm by Daura *et al.*⁴¹ The largest clusters were considered to provide the dominant contributions and were represented by the corresponding central configurations. Angles between regions of the motor domain in the different states were calculated using VMD.³⁰ A principal component analysis was performed for the C $_{\alpha}$ atoms of the region of interest using the analysis tools of GROMACS.

Two residues were considered to be hydrogen bonded when the distance between the donor X₁H and the acceptor X₂ did not exceed 0.35 nm and the angle X₁HX₂ exceeded 130°; here X₁ represents a nitrogen, oxygen or sulfur atom, and X₂ represents a nitrogen or oxygen atom.

4.5 Strand orientations within the β -sheet core

The β -sheet core of kinesin-3 consists of eight strands, β_1 to β_8 , corresponding to the residues 54–57 for β_1 , 7–12 for β_2 , 324–330 for β_3 , 91–97 for β_4 , 237–248 for β_5 , 218–230 for β_6 , 138–150 for β_7 , and 153–156 for β_8 , in the spatial order of the strands and according to PDB ID 1VFW.

For the β -sheet core of myosin, a twist has been found between neighboring β -strands.³⁴ In order to investigate the possibility that analogous twists occur in the β -sheet core of kinesin-3, as recently proposed for kinesin-1 in ref. 32 and 33 the orientation of the strand β_i extending from residue n_i to residue $n_i + l_i$ was characterized by the unit vector \hat{e}_i that points from the C $_{\alpha}$ -atom of residue $n_i + 1$ to the C $_{\alpha}$ -atom of residue $n_i + l_i - 1$.

The average tilt angle ϕ_{ij} between two neighboring strands β_i and β_j was evaluated *via* $\phi_{ij} = \min(\langle \psi_{ij} \rangle, 180^\circ - \langle \psi_{ij} \rangle)$ with $\langle \psi_{ij} \rangle \equiv \langle \arccos(\hat{e}_i \cdot \hat{e}_j) \rangle$. For each state, the angle $\langle \psi \rangle$ represents an average over all configurations. The corresponding standard error was evaluated by averaging over the individual trajectories. As a result, the tilt angle ϕ_{34} between the strands β_3 and β_4 as well as the tilt angle ϕ_{45} between the strands β_4 and β_5 are found to change by about 10°, both during phosphate release and during ATP binding, whereas the tilt angle ϕ_{56} is found to be essentially independent of the nucleotide state, see Table 1 in the main text.

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