EPR Spectroscopy Shows a Microtubule-Dependent Conformational Change in the Kinesin Switch 1 Domain

Nariman Naber,* Sarah Rice,† Marija Matuska,* Ronald D. Vale,† Roger Cooke,*‡ and Edward Pate§
*Department of Biochemistry and Biophysics, †Department of Cellular and Molecular Pharmacology, and §Department of Mathematics, Washington State University, Pullman, Washington 99164

ABSTRACT We have used site-directed spin-labeling and electron paramagnetic resonance spectroscopy to monitor a conformational change at the nucleotide site of kinesin. Cys-lite kinesin (K349 monomer) with the mutation S188C was spin labeled with MSL or MTSL. This residue is at the junction between the switch 1 region (which is a structure known to be sensitive to bound nucleotide in the G-proteins) and the α3-helix, adjacent to the nucleotide site. The spectra showed two or more components of mobility, which were independent of nucleotide in the absence of microtubules (MTs). The spectra of both labels showed a change of mobility upon binding to MTs. A more mobile spectral component became enhanced for all triphosphate analogs examined, AMPPNP, ADP·AlF4, or ADP·BeF4, in the presence of MTs, although the magnitude of the new component and the degree of mobility varied with nucleotide analog. The ADP state showed a much-reduced spectral change with a small shift to the more immobilized component in the presence of MTs. For kinesin·ADP·MT, a van’t Hoff plot gave ∆H° = −96 kJ/mol implying that the conformational change was extensive. We conclude there is a conformational change in the switch 1 region of kinesins when kinesin binds to MTs.

INTRODUCTION

One fundamental goal in the field of motility is to define the conformational changes at the nucleotide site associated with the interaction of the motor with the physiological substrate, and how the conformational changes that accompany nucleotide hydrolysis are translated into movement. Kinesin-family motors, along with myosin motors and the guanosine triphosphate binding proteins, are members of the larger G-protein superfamily (Vale, 1996; Kull et al., 1998). In this superfamily, the triphosphate-binding domain is composed of three conserved amino acid sequences: the P-loop, switch 1, and switch 2. A comparison of crystal structures shows that the conformation of switch 1 can vary. In the initial structures of kinesin-family motors, the phosphate-binding portion of the nucleotide site formed an open track, with the triphosphates exposed to solvent (Gulick et al., 1998; Kull et al., 1996; Sablin et al., 1996; Sack et al., 1997) and the switch 1 region (amino acids 190–204 in human kinesin) displaced away from the nucleotide. In all myosin motor x-ray structures, the switch 1 region is found to be adjacent to the nucleotide, forming in conjunction with the P-loop and switch 2 a closed phosphate tube (Yount et al., 1995) in which the phosphates are tightly bound. In G-protein x-ray structures, switch 1 is sensitive to both the bound nucleotide and the G-protein hydrolysis are translated into movement. Kinesin-family motors (Naber et al., 2002), have reported conformational changes in the mobility of nucleotide-analog, and EPR probes bound at the active site of kinesin-family motors (Naber et al., 2002), have reported a conformational change at the active site upon binding to microtubules (MTs). Switch 1 was not directly monitored in the studies, but the conformational change observed was compatible with the closed switch 1 conformation proposed by Minehardt et al. (2001). This fully closed structure has been proposed to be essential for both nucleotide hydrolysis and for the correct coordination of the conserved glycine in switch 2 to the γ-phosphate of the nucleotide. By analogy with nucleotide-site conformational changes in both myosin and the G-proteins, a popular hypothesis is that this latter coordination is an essential element for motility (Vale and Milligan, 2000; Smith and Rayment, 1996; Kull and Endow, 2002; Woehlke, 2001; Holmes and Geeves, 2000; Cooke, 1997; Schief and Howard, 2001).

EPR spectroscopy has been a useful tool in monitoring conformational changes in motor proteins (Baumann et al.,
Covalent attachment of an EPR probe at an endogenous cysteine suffers from two major deficiencies. First, the cysteine may not be at the precise location in the protein that one wishes to investigate. Second, the presence of multiple cysteines can result in multiple labels per protein, each with their own distinct spectral signature, dramatically complicating interpretation of spectral changes. Thus the use of endogenous residues has been augmented by site-directed spin labeling (SDSL) approaches (Serag et al., 2002; Isas et al., 2002; Hubbell et al., 1996). Endogenous, reactive cysteines are replaced with nonreactive amino acids ("Cys-lite" proteins (Rice et al., 1999)), and a single cysteine is then introduced at a specific desired location. The presence of a single reactive element simplifies spectral analysis and allows for more aggressive conditions for covalent labeling, enhancing EPR signal strength. SDSL has proved to be particularly powerful when x-ray crystal structures are available, and suggests new approaches to the study of conformational changes in kinesin. An EPR spin probe can be introduced onto the motor protein by covalent attachment to a cysteine residue in the protein. The motion of the attached probe is restricted by the adjacent protein surface. Conformational changes can thus be detected via changes in probe mobility, monitored as changes in the EPR spectrum.

In kinesin, the switch 1 region is located between a \( \beta \)-ribbon (\( \beta_5 \)) that forms part of the central, core \( \beta \)-sheet of the protein and a four-turn \( \alpha \)-helix (\( \alpha_3 \)), part of which forms one edge of the nucleotide-binding pocket (Fig. 1). Ser-188 is a surface-exposed residue located at the junction of the switch 1 region and the \( \alpha_3 \)-helix, adjacent to the nucleotide site in kinesin (Fig. 1). The residue is not conserved in kinesin-family motors. To further probe conformational changes in the switch 1 region, we have made use of the conservative mutation S188C for SDSL in a previously studied kinesin construct containing no other reactive cysteines (Rice et al., 1999). Previous use of this construct has been limited to labeling with undecagold to locate the position of the catalytic domain of kinesin on the microtubule, and a single fluorescence resonance energy transfer distance observation from kinesin already complexed with microtubules (Rice et al., 1999). There have been no previous spectroscopic studies directly monitoring conformational changes in the switch 1 region upon interaction with MTs. Here we demonstrate that EPR probes at the Cys-188 site show a conformational change in the switch 1 region upon binding to MTs, and that the conformational change differs in the di- and triphosphate states.

![FIGURE 1 Ribbon diagram of S188C kinesin. Human kinesin (Kull et al., 1996) with Cys-188 (yellow) at the junction of the \( \alpha_3 \)-helix (cyan, aa 176–189) with the switch 1 region (magenta, aa 190–213). The side chain of Ser-188 projects out into the aqueous environment in the crystal structure. The nucleotide-binding domain contains the P-loop (blue), switch 1, and switch 2 (green) homologous to myosin-family motors and the G-proteins. The \( \beta_5 \)-ribbon, aa 205–216, is shown in orange.]

### METHODS

The mutation S188C was introduced into monomeric Cys-lite K349 kinesin, and the protein was bacterially expressed and purified as previously described (Rice et al., 1999); amino acids 1–349. Kinesin was initially dialyzed into buffer A containing 25 mM PIPES, 100 mM NaCl, 2 mM MgCl₂, 1 mM EGTA, 50 mM ATP, pH 7.0, 4°C. The protein was spin labeled by reacting overnight at 4°C after addition of a twofold molar excess of 4-maleimido-2, 2, 6, 6-tetramethyl-1-piperidinyloxy (MSL) or 1-oxyl-2, 2, 5, 5-tetramethyl-\( \alpha \)-pyrrolin-3-methyl methane thiosulfonate (MTSL). Repetitive steps of protein concentration via centrifugation through a 10 kDa cutoff sizing filter, and subsequent dilution with buffer A, were used to remove excess spin label. The final protein concentration was ~100 \( \mu \)M. Protein concentration was determined by the method of Bradford (1976) using bovine serum albumin as a reference standard. The molecular weight of kinesin was taken as 39,700.

Tubulin was prepared from bovine brain tissue following the protocols of Ma and Taylor (1995) and stored in a buffer containing 100 mM PIPES, 1 mM MgCl₂, 1 mM EGTA, 1 mM Na₂GTP, –80°C. For microtubule polymerization, thawed tubulin was spun at 100,000 \( g \) for 20 min, 4°C, to remove aggregated material, and 1 mM Na₂GTP 10% DMSO and 2 mM MgCl₂ were then added to the supernatant. Polymerization was induced by 30 min incubation at 37°C. Taxol (20 \( \mu \)M) was added and the microtubules were spun down at 100,000 \( g \) for 25 min, 25°C. The pellet was resuspended in a buffer containing 25 mM NaCl, 25 mM PIPES, 1 mM EGTA, 2 mM MgCl₂, 20 \( \mu \)M taxol, pH 7.0, 25°C. The final concentration of the tubulin \( \alpha \beta \)-dimer was 200–250 \( \mu \)M. The molecular weight of the dimer was taken to be 110,000.

The microtubule-activated ATPase rates were determined using 1 \( \mu \)M kinesin, 10 \( \mu \)M microtubules in buffer A containing an additional 2 mM Na₂ATP. Liberated phosphate was determined using the malachite green assay (Kodama et al., 1986) with ATPase time points taken every 30 s over a 150 s time period. The slope of a least-squares fit to the data gave the ATPase rate.

First derivative EPR spectra were accumulated with an ER200D spectrometer (IBM Instruments, Danbury, CT) with the following instrument settings: microwave power, 25 mW; center field, 345.5 mT; modulation, 0.1 mT at a frequency of 100 kHz; gain, 0.1–2.0 \( \times \) 10⁶. Between 10 and 25, 12.7 mT wide, 50 s sweeps were averaged for final data.
analysis. Kinesin (50 μM) in buffer A with added 5 mM Li₂AMPPNP, 5 mM Na₂ADP, or 5 mM Na₂ADP and either 2 mM AlCl₃ or 2 mM BeCl₂ and 10 mM NaF was mounted in a 50 μL glass capillary and placed in a TE₀₁₁ cavity. Temperature was maintained by passing cooled or heated air through the radiation slits in the cavity and monitored by a thermistor located close to the experimental sample. For spectra in the presence of microtubules, 30 μM kinesin and 90 μM MTs were added to ligand-modified buffer A. The mixture was spun at 100,000 g for 25 min, 25°C. The pellet was then mounted in a depression in a rexolite flat cell, surrounded by grease, covered with a glass coverslip to prevent dehydration, and placed in the TE₀₁₁ cavity for data accumulation.

Spectra for MSL-labeled protein in the presence of ADP were deconvolved by first obtaining the pure components. An almost pure immobilized spectrum was observed experimentally in the presence of MTs, 2°C, and an almost pure mobile component was observed experimentally in the absence of MTs, 30°C. The small amounts of the other component in each of these were deleted by subtraction, and the resulting spectra taken as an approximately pure component for further analysis. 

Taxol was obtained from Molecular Probes (Eugene, OR). MSL was obtained from Aldrich (Milwaukee, WI). MTSL was obtained from Toronto Research Chemicals (Toronto, Ontario, Canada). All other compounds (nucleotides, phosphate analogs, and buffer salts) were obtained from Sigma Chemical (St. Louis, MO).

RESULTS

To fix notation, KS188C will refer to the K349 Cys-lite kinesin monomer, with mutation S188C used in these studies. KS188C has a MT-activated ATPase rate (1 μM KS188C, 10 μM MT) of 15.4 ± 0.4 kinesin⁻¹ s⁻¹ (mean ± range, two observations). This is ~75% of the value previously observed for Cys-lite K349 (Rice et al., 1999) when differences in ionic strength are taken into account. With MSL-labeled protein, MSL-KS188C, the MT-activated ATPase decreases to 9.8 ± 0.4 kinesin⁻¹ s⁻¹ (two observations), and for MTSL-KS188C the MT-activated ATPase is 8.3 ± 0.4 kinesin⁻¹ s⁻¹ (two observations). These values imply that spin probe-labeled protein retains catalytic competence, although the modification of the protein at Cys-188 does affect the rates of some step(s) in the cycle, emphasizing the importance of this region.

Fig. 2 demonstrates our fundamental results. The spectra in the absence (presence) of MTs are shown by dashed (solid) lines for MSL-labeled kinesin at 20°C. In the absence of MTs, the spectra appear independent of the state of the bound nucleotide. The spectra show that when MSL-KS188C-triphosphate analog binds to MTs, there is a shift of EPR probes from a more immobilized to a more mobile component. This is demonstrated by the shift in spectral intensity from the low-field component, A, to an up-field component, B, and by the decrease in intensity at the high-field dip, component C, in the AMPPNP, ADP·AlFx, and ADP·BeFx spectra when the spectra in the presence of MTs are compared with those in the absence of MTs. Additionally, the fact that these spectral components appear to occur at the same positions in the magnetic field suggests that the probes in all three triphosphate-analog states are reporting the same conformational change. A small spectral change is observed with ADP at the nucleotide site. When MSL-KS188C·ADP binds to MTs, there is a small outward shift of spectral intensity at both the low field (D) and high field (E) regions of the spectra. This implies that a fraction of the probes in the MSL-KS188C·ADP state now become more immobilized upon binding to MTs. Fig. 3 shows the results from MSL-labeled protein when the temperature is lowered to 5°C. The spectral changes are more pronounced than at 20°C. For the ATP-analog states, there is a significant inward shift in spectral intensity in the low-field components (A → B) and a decrease in intensity of the high-field dip (C). This again implies an increase in mobility of a fraction of the probes upon binding to MTs. There is, however, a small outward shift of the high-field dip (C) in the presence of ADP·BeFx, indicating that the magnitude of the mobility increase may be slightly less than with the other two
In the presence of ADP, the spectral change is smaller with a very small decrease in the mobility of some of the probes. Upon binding to MTs, there is a slight outward shift in spectral intensity in the low-field component (E → D) as shown by the change in the ratio of peak heights, and a corresponding outward shift in the high-field dip (F). These imply a shift to a more immobilized probe orientation.

Figs. 4 and 5 show that the spectral changes obtained in the presence and absence of MTs when Cys-188 is labeled with MTSL are more complex. In the absence of MTs at 20°C, there is a broad low-field shoulder, component A, which decreases in intensity in the presence of MTs for the triphosphate analogs. There is little change in the presence of ADP (Fig. 4). For all of the nucleotides, there is also an intensity decrease in the right-most portion of the low-field shoulder (C), with the appearance of a new peak with greater splitting (component B). Using the quantitative analysis and order parameter of Griffith and Jost (1976), the 4.37 ± 0.025 mT (four observations) splitting of this component can be modeled as motion within a cone with a vertex angle of 130°. Cone angles of mobility for the other two, more (A) and less (C) immobilized components of the spectra, cannot be defined as precisely. Thus at a minimum, the spectra indicate the presence of three distinct spectral components and potentially more.

At 5°C (Fig. 5), in the presence of triphosphate analogs, MTSL-KS188C shows the same basic spectral change as observed at 20°C. For the low-field peak, there is a shift in spectral intensity from a more immobilized component (A) to a component (B) of intermediate immobilization. This is clearest for MTSL-KS188C, where the A → E and B → D splittings of 6.64 ± 0.05 mT (two observations) and 4.52 ± 0.04 mT (four observations), respectively, correspond to an increase in the cone angle of motion from 52° to 126°, with the larger splitting corresponding to the smaller cone angle.

For MTSL-KS188C•ADP, spectral differences in the presence and absence of MTs show only a very small shift to a more immobilized conformation in the presence of MTs.

### The effect of temperature

As noted, the spectra we have obtained are complex. Many are the sum of three or more distinct spectral components, suggesting transitions between three or more distinct protein states. These are difficult to deconvolute, confounding more quantitative analysis. However, for MSL-KS188C bound to MTs in the presence of ADP, the probe shows only two different spectral components. Investigating the relative fractions of these two components in the spectra of MT•MSL-KS188C•ADP as a function of temperature allows us to define better the thermodynamics associated with the transition between the two protein conformations. Fig. 6, top, shows the spectra taken as the immobile component (solid line), and the mobile component (dashed line), used in defining the spectral shift as temperature varied for MT•MSL-KS188C•ADP. Pure components were obtained as described in Methods. The spectra of MT•MSL-KS188C•ADP could then be expressed as a linear sum of fractions of the two spectra shown in Fig. 6, top. Fig. 6, middle (MT•MSL-KS188C•ADP, 15°C), shows such a linear summation. Fig. 6, bottom, shows the quality of the fit in Fig. 6, middle, giving the residual spectrum obtained by subtracting the fitted spectrum from the experimentally determined spectrum. Letting $K = [\text{immobile fraction}] / [\text{mobile fraction}]$ be the equilibrium constant between the two conformations in the spectra of MT•MSL-KS188C•ADP as a function of temperature, Fig. 7 gives $-\Delta Rn(K)$ for the titration data versus $1/T$. Using the van’ t Hoff relationship, the slope of a least-squares linear fit to the data over the temperature range 5–30°C ($r = 0.99$) yields a mean value for $\Delta H^0 = -96$ kJ/mol and a mean $\Delta G$ of 2.2 kJ/mol. This value for $\Delta H^0$ implies a significant conformational change involving more than just a few amino acids. By comparison,
the enthalpy of α-helix formation in water is \( \sim 5 \text{ kJ/mol per residue} \) (Hermans, 1966). It must be emphasized, however, that we do not know the precise secondary structure involved in the kinesin conformational change, and that the adjacent protein surface can also significantly influence the enthalpy. For the remaining spectra, a change of comparable magnitude appears to be occurring, but this cannot be quantitated due to the overlap of the spectral components.

**DISCUSSION**

We have examined the changes in mobility of two different EPR probes covalently bound to KS188C in triphosphate-analog and diphosphate states in the presence and absence of MTs. The resulting EPR spectra are complex in that some have three or more distinct components. To simplify the discussion of the data, we have assumed that a given spectral component represents a single probe conformation. However, we cannot rule out the possibility of multiple conformations giving similar spectral signatures. Thus we may be underestimating the number of distinct probe conformations involved. Nonetheless, taken in toto, the data unambiguously report a conformational change at Cys-188 when kinesin binds to MTs. Furthermore, the probes report different changes in mobility between ADP- and ATP-analog states. AMPPNP and ADP\textsubscript{BeFx} are ATP analogs with the γ-phosphorous oxygens or fluorines in a tetrahedral orientation. ADP\textsubscript{AlFx} is instead thought to mimic the γ-phosphate hydrolysis transition state intermediate. Both MTSL and MSL show a trend from more immobilized to less immobilized spectra upon binding to MTs for triphosphate analogs. Furthermore, for a given EPR probe at a given temperature, the triphosphate analogs report similar spectra irrespective of the type of coordination around the γ-phosphorous. For ADP the situation is reversed. Although small, the trend is from more mobile to less mobile probes.

**RELATIONSHIP TO OTHER STUDIES**

A number of crystal structures of kinesin-family motors have been solved and are compatible with the possibility of a conformational change at the Cys-188 position. Kinesin Ser-188 is at the junction of the α3-helix and loop 9. Loop 9 and the adjacent α3a-helix at the nucleotide site form the switch 1 region of kinesin-family motors (Fig. 1). Fig. 8 shows the α3-helix-loop 9 junction (Ca backbone only) from representative kinesin-family x-ray structures. The α3-helix is at the top; loop 9 is at the bottom. The residue homologous to kinesin Ser-188 is shown as a solid sphere. As is evident, the x-ray structures show considerable lateral motion at the Cys-188 position. In Kar3\textsubscript{ADP} (Yun et al., 2001)/nct\textsubscript{ADP} (Sablin et al., 1996), it is displaced closest to/furthest from the nucleotide. Other x-ray structures lie between these extremes. Additionally, in the x-ray structure of Kar3 with the mutation, R598A (red), the α3-helix becomes disordered one amino acid (Asn-582) to the N-terminus (α3-helix side) of the location homologous to kinesin Ser-188 (Lys-583). This disorder continues into loop 9.

The switch 1 region of kinesin has a structure homologous to similar regions in the G-proteins and in the myosin motors. This structural element was first identified in the G-proteins as having a conformation that is sensitive to the state of the nucleotide (Sprang, 1997). It is also sensitive to the binding of other proteins (Goldberg, 1998; Boriack-Sjodin...
Kinesin-family motors generate biologically useful movement only in the presence of the polymer roadway. A fundamental difference between these previous crystallographic and molecular dynamics studies, and our present observations, is that MTs were not present in these earlier studies. Here we have use EPR spectroscopy to additionally investigate the influence of the polymer, and show that it is responsible for a conformational change adjacent to the nucleotide-binding site. Thus our results support previous observations of MT-induced conformational changes at the nucleotide site (Naber et al., 2002). By influencing the structure of the switch 1 region, the MT could be acting as a nucleotide exchange factor or could be promoting nucleotide hydrolysis. The rates of both exchange and hydrolysis are known to be enhanced by the binding of MTs.

In summary, EPR spectroscopy shows a conformational change at the nucleotide site of kinesin upon binding to MTs. Thermodynamic analysis implies a significant number of amino acids are involved.

REFERENCES


