Processive Movement by a Kinesin Heterodimer with an Inactivating Mutation in One Head†

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ABSTRACT: A single molecule of the motor enzyme kinesin-1 keeps a tight grip on its microtubule track, making tens or hundreds of discrete, unidirectional 8 nm steps before dissociating. This high duty ratio processive movement is thought to require a mechanism in which alternating stepping of the two head domains of the kinesin dimer is driven by alternating, overlapped cycles of ATP hydrolysis by the two heads. The R210K point mutation in Drosophila kinesin heavy chain was reported to disrupt the ability of the enzyme active site to catalyze ATP P→O bond cleavage. We expressed R210K homodimers as well as isolated R210K heads and confirmed that both are essentially inactive. We then coexpressed tagged R210K subunits with untagged wild-type subunits and affinity purified R210K/wild-type heterodimers together with the inactive R210K homodimers. In contrast to the R210K head or homodimer, the heterodimer was a highly active (>50% of wild-type) microtubule-stimulated ATPase, and the heterodimer displayed high duty ratio processive movement in single-molecule motility experiments. Thus, dimerization of a subunit containing the inactivating mutation with a functional subunit can complement the mutation; this must occur either by lowering or by bypassing kinetic barriers in the ATPase or mechanical cycles of the mutant head. The observations provide support for kinesin-1 gating mechanisms in which one head stimulates the rate of essential processes in the other.

Kinesin-1 and myosin V are the best characterized examples of motor enzyme molecules that move unidirectionally and continuously along the cytoskeletal filaments (microtubules and F-actin, respectively) that serve as their tracks (1–3). Each enzyme maintains a tight association with its track, undergoing tens or hundreds of catalytic turnovers without detaching from or sliding freely relative to it. Such “processive high duty ratio” movement allows individual isolated enzyme molecules to make forward progress even when opposed by substantial elastic loads. This property may be essential to the biological function of these enzymes as engines that propel subcellular organelles through the viscoelastic cytoplasm. Processive high duty ratio movement is thought to require that the actions of the two identical motor or “head” domains of the enzyme be coordinated so that at least one head is attached to the track at any given instant during movement. This head coordination hypothesis is consistent with observations that one-headed kinesin-1 constructs lack the ability to move processively (4–7) and with high duty ratio (8). The hypothesis is also consistent with demonstrations in two-headed molecules that catalytic turnover of the two heads is not independent, at least under some conditions (9–11).

The combined results from a series of studies (10, 12–17) show that kinesin heads move by an alternating-sites hand-over-hand mechanism, in which the two heads alternately hydrolyze ATP and each head moves forward 16 nm along the microtubule in every other hydrolysis cycle. This results in net movement of the enzyme as a whole by 8 nm per cycle. Curiously, the two heads do not behave quite identically; the data suggest that even- and odd-numbered steps begin with different conformations of the domain that links the heads to the C-terminal coiled-coil and that even and odd cycles have different $k_{cat}$ values under some circumstances (15, 17).

In addition to defining the sequence of domain movements that underlie kinesin motility, the alternating-sites hand-over-hand mechanism provides an attractive hypothesis to explain the head coordination that is thought to be responsible for the high duty ratio processive property of kinesin movement. In particular, it is widely accepted that such head coordination, and hence processive movement, is a consequence of the alternating ATP hydrolysis between the two heads (9, 18–20). Alternation of the ATPase cycles is enforced by the mechanical linkage of the two subunits through the neck linker domains. Specifically, the linkage is thought to mediate gating processes in which a step in the mechanochemical cycle of one head is slowed or blocked by the presence of the other head in particular states. Such inhibitory gating processes include (i) blocking of ADP release by the “tethered” head until nucleoside triphosphate binds to the “bound” head (9, 11, 21) and (ii) slowing of ATP binding to the leading head when the neck linker in the trailing head is docked (22, 23). Conversely, acceleration of trailing head detachment by mechanical strain transmitted from the leading head may represent an unusual stimulatory gating process,
but there are disagreements concerning the magnitude and mechanistic significance of this process (4, 23, 24).

To investigate possible additional stimulatory gating processes in kinesin, we prepared by coexpression a heterodimeric kinesin derivative in which one head has the wild-type sequence and the other has a single point mutation. Unlike previous studies using heterodimers (4, 10, 25, 26), we chose to focus on an active site mutation known to specifically target the chemical step of ATP hydrolysis. The Drosophila kinesin-1 R210K mutation almost completely blocks the ATP phosphoanhydride bond hydrolysis step while having little or no effect on other head functions (27). We here show that the R210K/wild-type heterodimer has novel properties distinct from those of previously characterized heterodimers. Pairing the inactive R210K subunit with a wild-type subunit produces a dimer with essentially the full mechanochemical function of homodimeric wild-type kinesin.

MATERIALS AND METHODS

Plasmids. Plasmid pWC2 encodes wild-type subunit K401-BIO-H6, which consists of the first 401 residues of Drosophila kinesin heavy chain fused to a biotin acceptor domain (this fusion is K401-BIO of ref 5) with the C-terminal extension (one-letter amino acid code) LSETSGHHHHHH added to permit Ni$^{2+}$ chelation affinity purification, pWC2 is identical to pEY4 (ref 5 and GenBank AY621072) except for the insertion 5′-TATCTGAGACTAGTGGGCACCACCATCACCATCTGTA-3′, which encodes the extension, at position 1476. pEY4 is identical to pSKA (28), except that the coding region is that given in GenBank AY621072. Plasmid pTTR210K encoding the mutant subunit K401-BIO-H6 R210K was made from pWC2 by directed mutagenesis (QuikChange, Stratagene) of the R210 CGA codon to AAA. pTTSTOP, which encodes untagged wild-type K401, was generated by directed mutagenesis of codon 402 (CGC) of pEY4 to the stop codon TGA. Coexpression experiments used pTTSTOP2, which was prepared by ligation of the BsrBI/HindIII coding region fragment of pTTSTOP with the BsaHI/HindIII replication origin fragment of pACYC184 (29), pTTR210K340, which encodes the tagged mutant kinesin head K340-BIO-H6 R210K, was constructed by ligating the 374 bp PstI/EcoRI fragment of pKA2 (28) to the 3390 bp PstI/EcoRI fragment of pTTR210K. The entire coding regions of pTTSTOP, pTTSTOP2 and pTTR210K were confirmed by sequencing.

Protein Expression and Purification. K401-BIO-H6 R210K homodimers were expressed from strain BL21(DE3) pLYsS pTTR210K and purified as described (5), except that a Ni$^{2+}$ chelation column (Qiagen Ni-NTA Superflow) was used in place of the avidin column. The resin was incubated with lysate overnight, and then washed for 2 h with 20 mM imidazole-Cl$^{-}$ pH 7.2, 4 mM MgCl$_2$, 10 mM 2-mercaptoethanol. Tagged kinesin was eluted with 500 mM imidazole-Cl$^{-}$ pH 7.2, 4 mM MgCl$_2$, 10 mM 2-mercaptoethanol and dialyzed against 50 mM imidazole-Cl$^{-}$ pH 6.7, 4 mM MgCl$_2$, 10 mM 2-mercaptoethanol prior to ion-exchange chromatography. The same procedure was used to purify heterodimers from BL21(DE3) pTTR210K pTTSTOP2, wild-type homodimers from BL21(DE3) pLYsS pWC2 and monomeric mutant heads from BL21(DE3) pLYsS pTTR210K340. Untagged wild-type K401 homodimer was expressed using BL21(DE3) pLYsS pTTSTOP, and BL21(DE3) pLYsS pWC2 was used to express K401-BIO-H6 homodimer.

Sedimentation. Protein diffusion coefficients were measured by fitting (SVEDBERG; John Philo Software) absorbance profiles obtained in a synthetic boundary experiment at low speed (8,000 rpm) in an analytical ultracentrifuge (XL-A with An60Ti rotor; Beckman-Coulter) in 20 mM imidazole-Cl$^{-}$ pH 7.2, 4 mM MgCl$_2$, 0.1 mM tris(2-carboxyethyl)phosphine at 20 °C. Absorbance of the sample before centrifugation (OD$_{280}$ = 1.2; 1.0 cm path length) corresponded closely with that measured in the centrifuge (OD$_{280}$ = 1.4; 1.2 cm), demonstrating the absence of significant quantities of protein aggregates large enough to be pelleted. Profiles from a conventional meniscus depletion experiment (44,000 rpm) were then analyzed by differential methods and fit to determine the sedimentation coefficient (30). Separate peaks from the heterodimer and mutant homodimer were not resolved in sedimentation experiments because the differences between the molecular weights and shapes of these species are small.

ATPase and Motility Assays. Steady-state ATPase activity was measured as described (28), except that the tubulin concentration was 1.2 mg/mL. Motility assays and bead tracking were conducted essentially as described (5) except that the beads were 150 nm diameter and the ATP regeneration system was omitted. A bead movement was included in the data set only when the complete sequence of bead binding to the microtubule, processive movement, and release was observed, and only movements longer than a minimum length needed for reliable detection (225 and 500 nm for heterodimer and wild-type, respectively) were used for subsequent quantitative analysis. To measure the characteristic run length of a given enzyme species, individual movement length measurements were binned (100 and 500 nm bin width for heterodimer and wild-type, respectively) and normalized, and the resulting histogram was fit with a single exponential probability density function.

RESULTS

Properties of R210K Homodimers and Monomers. In the mutant enzyme studied here, arginine substitutes for lysine at position 210 (R210K) in the well-conserved switch I loop (31) adjacent to the kinesin active site. Despite the conservative nature of this mutation, Klumpp et al. (27) showed that a homodimeric R210K construct is almost completely catalytically inactive; its ATPase $k_{cat}$ of 0.22/dimer/s is ~200-fold reduced from the corresponding wild-type construct. In contrast, microtubule association kinetics and nucleotide affinities of the mutant homodimer were similar to wild-type. Furthermore, nucleotide-stimulated ADP release was unimpaired in the R210K homodimer, evidence that machinery for head-head communication remains functional. To confirm and extend these earlier studies, we expressed and purified a similar mutant kinesin consisting of the N-terminal 401 amino acids of Drosophila kinesin with C-terminal biotin and six-histidine tags (K401-BIO-H6 R210K). K401 constructs lack the complete C-terminus of intact kinesin heavy chain, but contain at least 40 amino acids of the α-helical coiled-coil neck domain, a length sufficient to induce formation of stable dimers (32–34). We also
prepared a K340-BIO-H6 R210K construct that lacks this dimerization domain; that protein was homogeneously monomeric as expected (Figure 1). The ATPase specific activities of both the mutant monomer and homodimer constructs were dramatically reduced relative to wild-type dimers (Table 1, lines 4 and 2 vs line 1). We further tested the ability of the mutant homodimers to move streptavidin-coated 150 nm diameter polystyrene beads along immobilized microtubules, using an experimental design in which wild-type homodimers show robust activity (5). No bead movement was detected at enzyme:bead mole ratios of 1:1, 10:1, or 100:1 (Table 1 and data not shown). At all ratios, bead-labeled mutant homodimers remained bound at fixed positions on microtubules in 1 mM ATP or in controls with no added nucleotide, but did not bind in controls with 1 mM ADP. These findings are consistent with observations in bulk experiments (27) that linkage between nucleotide and microtubule binding is retained in the mutant.

**Wild-Type/R210K Heterodimers.** Having confirmed that R210K inactivates kinesin both in homodimers and in isolated heads, we made preparations containing heterodimers with one wild-type and one mutant subunit. Tagged mutant subunits (K401-BIO-H6 R210K) and untagged wild-type subunits (K401) were coexpressed and proteins that contained at least one tagged subunit were purified by affinity and ion exchange chromatography, yielding mutant/wild-type heterodimers mixed with the inactive mutant homodimers. Denaturing electrophoresis of the purified protein reveals two bands at the expected sizes of the tagged and untagged subunits (Figure 2). The tagged:untagged molar concentration ratio estimated by densitometry was 1.6:1. That is close to the 2:1 ratio expected if the two types of subunits are expressed equally, subunits associate randomly, and untagged wild-type homodimers are fully removed during purification. Sedimentation velocity measurements on the heterodimer sample demonstrate that it is uniformly dimeric, with sedimentation and diffusion coefficients indistinguishable from those of wild-type homodimers (Figure 1). Thus, the

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**Table 1: ATPase Activity and Motility of Kinesin Derivatives**

| Enzyme Species | ATPase Activity (mean ± S.D.) (s⁻¹) | Movement Velocity (mean ± S.D.) (μm s⁻¹) | Run Frequency (mean ± S.E.) (min⁻¹) | Run Length (± S.E.) (μm) |
|---------------|-----------------------------------|-----------------------------------|-----------------------------------|------------------|
|               | 24.3 ± 0.5 (N = 6)                | 0.8 ± 0.2 (N = 79)                | 1.0 ± 0.2 (N = 25)                | ≥1.5 ± 0.1 \(^d\) (N = 79) |
|               | 0.19 ± 0.02 (N = 6)               | <0.00010 (N = 8)                 | <0.04 *                         | No motility |
|               | 10.0 ± 0.1 (N = 3)                | 0.09 ± 0.12 (N = 50)             | 0.4 ± 0.1 (N = 10)               | 0.31 ± 0.06 (N = 50) |
|               | 0.18 ± 0.02 (N = 3)               | N.D. \(^f\)                      | N.D. \(^f\)                     | N.D. \(^f\) |

\(^{a}\) Schematic diagrams of enzyme species present, using the same symbols as in Figure 1. \(^{b}\) Microtubule-stimulated ATPase specific activity per dimer (rows 1–3) or per monomer (row 4) in 1 mM ATP, 1.2 mg/mL microtubules, 25 °C. \(^{c}\) Data set included some runs where motor ran off the microtubule end. \(^{d}\) No motility detected. Value given is the detection limit. \(^{e}\) Implies that heterodimer specific activity is ∼15 s⁻¹, since the preparation is estimated by gel densitometry to contain ∼35% mutant homodimers, which have negligible activity. \(^{f}\) Not determined.

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**Figure 1:** Population distributions \(g(s^\ast)\) of apparent sedimentation coefficients \(s^\ast\) in sedimentation velocity experiments on the heterodimer (blue) and mutant monomer (purple) samples (left). Circles are measurements; lines are theoretical non-interacting single-species distributions using the sedimentation and diffusion coefficients (right, rows 1 and 2) determined by fitting (see Materials and Methods). Cartoons show the domain structures of protein species in the samples: red, wild-type head; yellow, R210K mutant head; thick black, neck coiled-coil; green, biotin and six-histidine tags; white, biotin tag. Wild-type data (right, rows 3 and 4) from ref 5 are included for comparison.
R210K mutation does not affect the oligomeric state of the construct nor significantly alter its overall shape, consistent with previous conclusions (27) that R210K causes a local structural alteration at the catalytic site, not a global folding defect.

If the kinesin mechanism requires that the heads alternate catalysis (9, 18–20), a mutation which blocks ATP hydrolysis in one head of a mutant/wild-type heterodimer would be expected to prevent steady-state catalytic turnover of the wild-type head. Despite the fact that R210K heads are essentially inactive in both monomer and homodimer contexts, the heterodimer sample microtubule-stimulated ATPase specific activity is only ∼2-fold reduced relative to that of wild-type homodimers (Table 1). The observed activity was not from the mutant homodimers present in the sample; that species is inactive (Table 1). Nor can the activity be explained by postulating contamination of the heterodimer sample by wild-type homodimers, because the wild-type homodimer and heterodimer sample ATPase activities were observed to have quite different salt dependencies: the wild-type homodimer ATPase decreased ∼58-fold upon addition of 150 mM NaCl whereas heterodimer ATPase decreased by only ∼8-fold.

Motility of Heterodimer Sample. Can ATP hydrolysis by the mutant/wild-type heterodimer drive processive movement of a single enzyme molecule along a microtubule? The heterodimer sample was mixed with streptavidin-coated beads at a 1:1 mole ratio of protein dimers to beads; in the motility experiment these beads moved processively and unidirectionally along microtubules, at a velocity ∼9-fold slower than that of wild-type homodimers (Table 1; Video S1 and Video S2, Supporting Information).

At the 1:1 dimer:bead ratio, most beads had ≤1 enzyme molecule. Nonetheless, the frequency of heterodimer bead movements was roughly similar to that observed with wild-type homodimers (Table 1). This is strong evidence that processive movement can be generated by single heterodimers; it is not restricted to the extremely small subpopulation of beads with >1 enzyme molecule simultaneously interacting with the microtubule (2). The small (∼2-fold) reduction in movement frequency in the heterodimer sample relative to wild-type homodimers is explained in large part by the fact that a significant fraction of molecules in the former are inactive mutant homodimers.

In principle, the heterodimer preparation might contain wild-type homodimers due to incomplete removal in the purification, homologous recombination between the plasmids, or subunit exchange after purification. Two lines of evidence confirm that the observed motility is not from wild-type homodimers. First, no motility was seen in control experiments in which the streptavidin-coated beads were pretreated with excess biotin before adding enzyme. This implies that the movements observed in the heterodimer sample arise from enzyme molecules specifically attached to the beads by a biotin–streptavidin linkage; the wild-type homodimers should lack the biotin tag. Second, movement in the heterodimer samples has different properties from that driven by wild-type homodimers, suggesting that the two arise from distinct enzyme species. Velocity distributions observed for wild-type homodimers and the heterodimers are nearly disjoint (Figure 3). Also, the characteristic distance moved by each enzyme molecule (“run length”) was significantly lower for heterodimers, and the mean duration of the run (i.e., run length divided by mean velocity) was substantially larger (Table 1).

In control experiments, we examined motility from equimolar mixtures of wild-type K401-BIO-H6 with the mutant/wild-type heterodimer sample or with mutant homodimers. In the former, the bead velocity distribution had two peaks, one at the wild-type velocity (∼800 nm/s) and one at the heterodimer velocity (∼95 nm/s). The latter had only a single peak at the wild-type velocity. These experiments demonstrate that movement of wild-type homodimers is not slowed by the simultaneous presence of heterodimers or mutant homodimers, confirming that contaminating wild-type homodimers do not cause the slow motility observed in the heterodimer sample.

Nanometer-scale tracking of single molecules (35) was used to determine whether beads driven by the mutant/wild-type heterodimers moved unidirectionally. As is the case with wild-type homodimers, clear backward displacements were rare (Figure 4). For each molecule tracked, the distribution of bead positions normal to the microtubule axis was narrow and movement was parallel to the axis. These observations
The properties of the R210K/wild-type heterodimer do not conform to this prediction. We observed that this species moved rapidly along microtubules, and it hydrolyzed ATP at rates >50-fold faster than the R210K homodimer. The high activity of the heterodimer is a striking contrast to previously reported processive mutant/wild-type kinesin heterodimers, none of which has steady-state activity significantly greater than twice that of the corresponding mutant homodimer (10, 26). We found single R210K/wild-type heterodimers to be processive, high duty ratio motors like wild-type homodimers. Additional experiments confirmed that the activities attributed to heterodimers are not due to contamination by wild-type homodimers or mutant homodimers and are not a result of aggregation or higher-order oligomerization of heterodimers.

Intradimer Complementation of the Motility Defect in the R210K Subunit. High duty ratio processive motility requires two heads; it is absent in single-headed kinesin-1 constructs (4–7). The data presented here show that motility is restored when a wild-type subunit is associated with an R210K subunit that in other contexts (monomer or homodimer) is nonmotile and almost completely inactive. Analogous effects have been seen with other motor enzymes: for both myosin II and Rep helicase, dimer-specific functions can be restored when a wild-type subunit associates with a nonfunctional mutant subunit (38, 39). Since the R210K/wild-type kinesin heterodimer takes 8 nm steps, tracks microtubule protofilaments, and appears to limp, there is a strong likelihood that it moves by the same pattern of alternating hand-over-hand head movements as wild-type homodimers.

**DISCUSSION**

**Association of the Inactive R210K Subunit with a Wild-Type Subunit Produces a Dimer with Near-Wild-Type ATPase and Motor Activity.** Kaseda et al. (10) hypothesized that the activity of some kinesin mutant/wild-type heterodimers can be explained by a hand-over-hand mechanism in which the mutation slows a step in every other catalytic cycle, leading to the alternating long and short dwell times that they observed between successive 8 nm steps. In its simplest form, this hypothesis predicts that movement speed and ATPase $K_{cat}$ of a wild-type/mutant heterodimer will be no more than twice those of the corresponding mutant/mutant homodimer, since the slow step occurs in every cycle of the latter and in every other cycle of the former.

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**Notes:**

1. A potential exception is the loop 12/helix 5 microtubule binding site triple mutant Y274A/R278A/K281A. However, this mutant homodimer binds microtubules so weakly that it was not possible to measure the ATPase $K_{cat}$ or single-molecule movement velocity (10, 37). Thus, homo- and heterodimer activities for this species cannot be compared.
In the most widely accepted view of the kinesin catalytic cycle (see refs 23, 40), the enzyme hydrolyzes ATP in the trailing head in a configuration in which the leading head is empty and both heads are bound to the microtubule (Figure 5A, uppermost species). Klumpp et al. (27) showed that the hydrolysis step is blocked by the R210K mutation. How is this kinetic barrier overcome in the R210K/wild-type heterodimer? In purely formal terms, the presence of the wild-type subunit can complement the kinetic defect in the R210K subunit either by acting directly to lower the kinetic barrier (Figure 5, orange) or by bypassing the barrier through an alternative movement mechanism (for example, Figure 5, violet). In the schemes shown, both possibilities result in hand-over-hand movement coupled to ATP hydrolysis. The key difference is that the former hydrolyzes one ATP per 8 nm step (Figure 5B), like wild-type homodimers, while the latter hydrolyzes one ATP per two steps (Figure 5C). These two possibilities could in principle be distinguished by comparing the ATPase specific activity in bulk with the single-molecule movement velocity measured under the same conditions (35, 41). However, approaches based on specific activity will not be reliable until the heterodimer mechanism is understood in enough detail that active site titrations methods can be used to make accurate measurements of the concentration of active heterodimer molecules (35, 41), and the extent to which heterodimer ATPase might be partially uncoupled from motility is assessed. Without such additional data, both barrier lowering and barrier bypass pathways remain as plausible explanations for the complementation of the R210K kinetic defect by heterodimerization with a wild-type subunit.

Precedents for Overcoming the R210K Kinetic Barrier in Heterodimers. All kinesin and myosin motor domains have an invariant arginine residue in the switch I motif at the position corresponding to R210 in Drosophila kinesin (31, 42). For myosin, there is extensive evidence (reviewed in ref 42) that the arginine side chain, together with an adjacent conserved glutamic acid side chain to which it forms a salt bridge when the nucleotide binding pocket is closed, plays an essential role in nucleoside triphosphate hydrolysis by precisely positioning a water molecule adjacent to the γ-phosphate of ATP. A similar role has been proposed for R210 in kinesin (27). In both families of enzymes, single point mutations at these positions abolish or greatly reduce microtubule- or actin-stimulated ATPase activity of isolated monomer heads (this report and refs 27, 39, 42-45). Thus, available data suggest that mutations at R210 impair ATPase function primarily by local disruption of the active site geometry needed for catalysis. This view is supported by structural modeling which shows that the conservative R210K substitution is expected merely to subtly alter the network of H-bonds in the ATP-bound active site (27).

If R210K causes only a subtle localized alteration in active site conformation, the barrier lowering pathway is a reasonable explanation for its near wild-type function. Specifically, one might hypothesize that when the mutant head is trailing (yellow head in the topmost state of Figure 5A), mechanical strain from the bound leading head transmitted through the neck linker might be sufficient to rearrange the structure of the active site and restore the hydrolytic activity that is absent in the R210K monomer, which lacks the internal mechanical strain of the two head bound state. There is ample precedent for mechanical strain transmitted through the neck linkers altering the kinetics of chemical processes within the heads; for example, this is thought to be the mechanism that prevents premature binding of ATP to the lead head (22, 46).

Conversely, the essential steps of the barrier bypass pathway shown in Figure 5A are also supported by precedents in the literature. This pathway (violet arrow) postulates detachment of the trailing mutant head (yellow) while it is in the strongly microtubule binding ATP state. This process is expected to be slower than detachment of an ADP bound trailing head as occurs in the wild-type kinesin mechanism (Figure 5A, green box), but inhibitor studies suggest that it can occur at a substantial rate (23, 47). The feasibility of the bypass pathway is supported by a computational model of R210K/wild-type heterodimer function (48). Also, the idea that a kinesin-1 heterodimer can function with only one catalytically active head parallels the recent revelation that the heterodimeric kinesin-14 Kar3/Vik1 functions as a motor despite the absence of catalytic activity in the Vik1 head (49). However, the barrier bypass pathway proposed here, in which the inactive R210K head binds ATP only to release it later in the cycle, is probably not directly applicable to Kar3/Vik1 since the structure of an isolated Vik1 head shows that the nucleotide binding pocket is absent in this protein (49).

Failure To Overcome the Kinetic Barrier in R210K Homodimers. In the hypothesized scheme for heterodimer function, 8 nm steps occurring through the barrier lowering or bypass pathway alternate with steps occurring by the wild-type pathway (Figure 5B, C). The implication is that since no such alternation can take place in the R210K homodimer, reactions of that enzyme should be restricted to the altered pathway in every step. Thus, the bypass pathway predicts no ATPase activity and no ATP-driven motility, in good agreement with the observed properties of the homodimer (this study and ref 27). In contrast, the barrier lowering pathway predicts that the homodimer will be functional as long as it can successfully enter the two head bound state shown at the top of Figure 5A. Therefore, to explain the low activity of the homodimer in the barrier lowering scheme, it is necessary to make the additional assumption that the enzyme becomes trapped in a kinetically refractory state (e.g., an off-cycle one-head-bound species) from which it is unable to efficiently enter the catalytic cycle. This scenario is consistent with available data, but there is no independent evidence for such a state.

Implications for the Mechanism of Wild-Type Kinesin-1. High duty ratio processive movement by kinesin-1 requires coordination of the cycles of the two head domains. Most gating processes that have been demonstrated or proposed to enforce this coordination are inhibitory in nature. Both in the slowing by internal mechanical strain of lead head ATP binding (22) and in the blockage of ADP release from the tethered head (9), a reaction step that occurs readily in one-headed constructs is inhibited (in some parts of the mechanoochemical cycle) as a consequence of the head→head linkage in the dimer. It is only in the hypothesized strain-induced dissociation of the trailing head (4, 24) that the partner head is proposed to have a stimulatory rather than inhibitory effect. The observation that the presence of the wild-type head in the R210K/wild-type heterodimer complements the kinetic defect in the mutant head provides a new example of such
FIGURE 5: Scheme for the mechanochemical function of R210K/wild-type heterodimers. Microtubule protofilaments with alternating αβ-tubulin dimers (gray/white) are shown with plus ends to the right; T, ATP; D, ADP; P_i, inorganic phosphate. (A) Overcoming the kinetic barrier when the R210K head is in the trailing position. In the single turnover cycle for wild-type kinesin homodimers (analogous to that shown in the green box, but with two identical heads), ATP is hydrolyzed in the trailing head from a state (top) in which both heads are bound to the microtubule. In the R210K/wild-type heterodimer, individual cycles with the mutant head in the leading position (not shown) alternate with individual cycles (green box) in which the wild-type head (red) is leading and the mutant head (yellow) is trailing. In these cycles where the R210K head is trailing, its pairing with a wild-type head in the heterodimer must enable it to evade the kinetic barrier to ATP hydrolysis (red “X”) present in isolated R210K monomers. Evasion could be accomplished by lowering the barrier (orange arrow), allowing the trailing mutant head to hydrolyze ATP much more rapidly in the heterodimer context than in the monomer context, or by bypassing the barrier via a pathway in which the trailing mutant head steps forward without accompanying hydrolysis of ATP. The mechanism of this putative bypass pathway might consist of ATP binding to the leading head, followed by stepping, followed by release of unhydrolyzed ATP from the (now leading) mutant head (violet arrow), but other arrangements of steps are also possible. (A possible alternative is that ATP remains continuously bound to the mutant head, as in the short runs seen in mixtures of ATP and an inhibitor in ref 47.) In the wild-type kinesin mechanism, the position of ADP-associated heads has been variously reported as microtubule-bound, tethered, or docked to the leading head (see discussion and citations in refs 40, 50); for simplicity, only the first of these possibilities is illustrated. Docked and undocked neck linker strands in the trailing head are represented with straight and curved lines, respectively. (B and C) Summary of predicted reaction stoichiometries for the barrier lowering (B) and barrier bypass (C) mechanisms. Each box shows the net stoichiometry of substrate binding and product release for two consecutive 8 nm steps; the step with the mutant head (yellow) trailing is shown first, followed by the step with the wild-type head (red) trailing. In these schemes, the first 8 nm step is assumed to proceed by the pathways shown as orange (B) or violet (C) arrows in panel A. The second step is assumed to proceed by the wild-type mechanism since the functions performed by the leading head (binding ATP and stimulating nucleotide release from the other head) are essentially unaffected by the R210K mutation (27).

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SUPPORTING INFORMATION AVAILABLE

Differential interference contrast microscopy video recording of polystyrene bead-labeled wild-type kinesin homodimers (Video S1) or wild-type/R210K heterodimers (Video S2) moving along stationary microtubules at 1 mM ATP. The microtubule was near vertical in S1 and was diagonal (upper right to the lower left) in S2. Frame width ∼2 µm in both S1 and S2. This material is available free of charge via the Internet at http://pubs.acs.org.

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