Kinesin-related proteins (KRPs)

Kinesin-related proteins (KRPs)1, also known as kinesin-like proteins (KLPs) or kinesin family proteins (KIFs), form a superfamily of microtubule-based mechanochemical motors. Kinesin itself, the archetypal member of the family, was discovered in squid axoplasm, suggesting a role in axonal transport (Brady, 1985; Vale et al., 1985). As expected for this role, kinesin heavy chain mutants in Drosophila have defective neuromuscular function (Saxton et al., 1991; Gho et al., 1992). Kinesin-related proteins have since been found in a wide range of organisms, including several fungi, Arabidopsis, Drosophila, Xenopus, and mammals, and are known to be involved in such diverse cellular processes as spindle pole separation, chromosome movement, and neuronal vesicle transport (for review see Goldstein, 1993; Bloom and Endow, 1994, 1995; Moore and Endow, 1996).

The question of how many different kinesin-related proteins there might be in a single organism has been addressed by in situ hybridization studies, using a probe generated by PCR amplification with primers to highly conserved sequences in the motor domain (Endow and Hatsumi, 1991; Stewart et al., 1991). This led to the unexpected conclusion that there are at least 30 members of the kinesin superfamily in Drosophila. It now appears that other higher eukaryotes have a similarly large number of kinesin-related proteins. A number of these have now been sequenced, and their specific functions and regulation are the subject of intense investigation.

Kinesin itself and a number of kinesin-related proteins are involved in the movement of membrane-bound organelles and vesicles. However, a large number of kinesin-related proteins have been found to be involved in various aspects of meiosis and mitosis. This is perhaps not surprising in view of the central role of the microtubule spindle in these processes. Known or postulated roles include all aspects of spindle structure and function, microtubule dynamics, chromatin structure and function, and chromosome movement (Vernos et al., 1995; Walczak et al., 1996; Boleti et al., 1996; for review see Goldstein, 1993; Bloom and Endow, 1995; Moore and Endow, 1996).

Kinesin and KRPs share a conserved motor domain of ~340 amino acids, which defines the superfamily of kinesin-related proteins. Similarities between the motor domains have been used to construct molecular phylogenies of the KRPs (Moore and Endow, 1996). KRPs with similar motor domains often have similar functions, but this is by no means always the case (see Discussion).

Outside of the motor domains the kinesin-related proteins differ from each other, suggesting that the different cellular roles of these proteins depend on the non-motor sequences. Like kinesin heavy chain, many kinesin-related proteins have a central α-helical region. This region is thought to mediate the homodimerization of the kinesin heavy chain in the native tetrameric protein. The “tail” region is thought to bind the “cargo” and/or accessory proteins. Little is known about the proteins that bind to KRPs, although they are clearly critical to force transduc-

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1. Abbreviations used in this paper: cdk, cyclin-dependent kinase; DAPI, 4',6-diamidino-2-phenylindole; KLP, kinesin-like protein; KRP, kinesin-related protein; ORF, open reading frame.
tion to the “cargo” and probably also to the regulation of the motor.

Kinesin itself is a tetramer, comprising two heavy chains and two light chains. The heavy chains contain the motor domain at the NH2 terminus and the light chains bind toward the COOH terminus. Another protein, kinectin, is thought to bind to kinesin (Toyoshima et al., 1992; Kumar et al., 1995). Molecular cloning and antibody studies suggest that kinectin is an integral membrane protein anchored in the ER (Futterer et al., 1995; Yu et al., 1995), and thus may function as a molecular link between motor and cargo. Very few associated proteins have been identified for kinesin-related proteins; candidates include Cik1 for Kar3 (Page et al., 1994) and a heterotrimeric sea urchin complex consisting of two different kinesin-related proteins and a non-kinesin protein (Cole et al., 1993).

Little is known of the associated proteins; regulation of motility and cargo binding of kinesin-related proteins is correspondingly poorly understood. Several reports have suggested that phosphorylation may play an important role. Both the heavy and light chains of kinesin are phosphorylated, as is kinectin (Hollenbeck, 1993; Lee and Hollenbeck, 1995). Membrane-associated kinesin heavy chain is more highly phosphorylated than soluble kinesin heavy chain, which suggests that membrane association is regulated by phosphorylation. Several different kinases have been implicated in the phosphorylation of kinesin and KRP. Pharmacological studies have shown that protein kinase A activation selectively inhibits anterograde axonal transport of vesicles, but not mitochondrial transport or retrograde transport (Okada et al., 1995). This activation induces the phosphorylation of several axonal proteins including kinesin. On the other hand, protein kinase A activation stimulates plus end–directed pigment granule movement in fish chromatophores (Rozdzial and Haimo, 1986; Rodionov et al., 1991; Sammak et al., 1992). The Xenopus kinesin-related protein Eg5 contains a predicted cyclin-dependent kinase (cdk) phosphorylation site in the COOH-terminal tail that is required for spindle association and function (Sawin and Mitchison, 1995; Blangy et al., 1995). Another kinase that may regulate a kinesin-related protein is polo-like kinase (Plk). Plk is a murine protein kinase related to the Drosophila polo, Saccharomyces cerevisiae CDC5, and Schizosaccharomyces pombe plo1 protein kinases, and colocalizes with the kinesin-related protein CHO1/MKLP-1, which it can phosphorylate in vitro (Lee et al., 1995).

Even less is known about the protein phosphatases that must act antagonistically to these kinases. Serine/threonine protein phosphatases are classified into four major classes: types 1, 2A, 2B, and 2C (PP1, PP2A, PP2B, PP2C; for review see Cohen, 1989; Bollen and Stalmans, 1992; Sheno-linar, 1994). Okadaic acid, a potent inhibitor of PP2A and PP1, stimulates kinesin motor activity at least twofold, apparently through the hyperphosphorylation of kinesin-associated proteins (McIlvan et al., 1994). Subsequent work has shown that maximal inhibition occurs at 500 nM okadaic acid, implicating PP1 rather than PP2A, which is completely inhibited at much lower concentrations of okadaic acid (McIlvan, J.M., Jr., L. Lindesmith, Y. Argon, and M.P. Sheetz, 1994. Mol. Cell. Biol. 5(Suppl.):31a). This shows that PP1 is present in semipurified motor fractions and regulates motor activity.

In this paper we describe the identification of a Drosophila kinesin-related protein, KLP38B, in a screen for proteins that associate with type 1 serine/threonine protein phosphatase (PP1) and show that KLP38B binds PP1 in vitro. KLP38B is expressed in all proliferating cells examined, and is required for normal chromosome condensation in mitosis and for male and female fertility. Incomplete lethality of KLP38B mutants implies a degree of functional redundancy, although KLP38B sequence is not closely related to other known Drosophila KRP.

**Materials and Methods**

**Two-Hybrid System**

Two-hybrid screening for proteins capable of binding to PP1 87B was carried out essentially as in Harper et al. (1993). The start codon of a PP1 87B cDNA (Dombrádi et al., 1989; Axton et al., 1990) was modified to an NdeI site and subcloned into the NdeI site of pAS2 to create pAS2-PP1. A Drosophila third instar larval cDNA library, constructed in pACT (Durfee et al., 1992), was a gift from Stephen Elledge (Baylor College of Medicine, Houston, TX), together with the yeast strains and plasmids described in Harper et al. (1993). This library was screened for cDNAs that interact with pAS2-PP1 in the yeast strain Y190 (Harper et al., 1993). 25 cDNAs were isolated that are strong HIS+ lacZ+ positives in combination with pAS2-PP1 but not with control plasmids (pAS-p53, pAS-cdc2, pAS-lamin). Three of these 25 positives are derived from KLP38B.

**Sequence Analysis**

Sequence of the 3,605-bp cDNA (see Fig. 1) was determined using cycle sequencing with fluorescent terminators and a semiautomated sequencer (ABI 377; Perkin-Elmer Corp., Norwalk, CT), according to the manufacturer’s instructions. Synthetic oligonucleotide sequencing primers were supplied by Hobolth DNA Syntese (Hillerød, Denmark). Other KLP38B cDNAs were sequenced by using a combination of Eco III deletions and primer walking. Sequence analysis and searches of sequence databases were performed using the Genetics Computer Group (GCG) programs (Devereux et al., 1984), PACTIO (Berger et al., 1995), and PEPCOIL (Lupas et al., 1991; Lupas, 1996).

**KLP38B cDNA Isolation**

cDNAs encoding part of KLP38B were obtained using the two-hybrid system (see above). Longer cDNAs were isolated from the Nick Brown 0–4-h embryonic cDNA library (Brown and Kaferos, 1988) by hybridization, using the longest two-hybrid cDNA as a probe. Two longer cDNAs were isolated, but neither contains the complete KLP38B open reading frame (ORF). Screening a testes cDNA library (kindly provided by T. Hazelrigg, Columbia University, New York) identified another seven cDNAs; again none of them contained the complete KLP38B ORF. Full-length cDNAs were obtained by using a novel PCR-based screening procedure developed for the purpose. XL-2 cells (Stratagene, La Jolla, CA) were transformed with 3 ng of the Nick Brown imaginal disk library. 100 cultures were observed from a single row pool and a single column pool, unambiguously identifying one of the original 100 cultures as containing a 3-kb KLP38B cDNA. This was purified by hybridization with the KLP38B-specific oligo (5′-CGGAAGTACGAGTCCACCGAGTGGG-3′) and a vector-specific oligo (5′-GCTCAGAATAAACGCTCAACTTTGGG-3′). The size of the amplified fragments was determined by gel electrophoresis. A 1.1-kb band was observed from a single row pool and a single column pool, unambiguously identifying one of the original 100 cultures as containing a 3.6-kb KLP38B cDNA. This was purified by hybridization with the KLP38B-specific oligo, and the identity of the purified clone was confirmed by PCR. This method is described in detail in Alphey (1997).

**KLP38B Cosmid Isolation**

A high density gridded filter from the European Genome
Identification of KLP38B Mutants

A large-scale screen has identified 2,711 lethal and semilethal P[lacW]-induced mutations on the second chromosome (Török et al., 1993). This modified P element contains a plasmid origin of replication and antibiotic resistance gene, allowing plasmid rescue of DNA flanking the insertion site. Plasmids representing 1,836 lines were independently rescued and pooled in batches of 10 and 100. Pools of 100 plasmids were screened by hybridization with genomic DNA from the KLP38B locus. Hybridizing pools were then narrowed down to single plasmids by screening their constituent subpools and finally individual plasmids, and corresponding mutant lines were obtained. This method is described in detail in Guo et al. (1996). Six pools of 100 plasmids showed cross-hybridization with the KLP38B probe, a 7-kb BamHI/EcoRI fragment from cosmid 164E4 (see Fig. 4). Subdivision of the positive pools identified six insertion lines: l(2)k08002, l(2)k03903, l(2)k09102, l(2)k04914, l(2)k04805, and l(2)k05702. Two additional alleles (KLP38B-1 and KLP38B-4) were gifts from Mike Gold-ettner (Cornell University, Ithaca, NY). Other stocks were from Bloomington Drosophila Stock Center (Indiana University, Bloomington, IN) and The European Drosophila Stock Center (University of Umeå, Umeå, Sweden).

\[ P[lacW] \] Reversion

Excision of P[lacW] elements to test for phenotypic reversion was conducted by crossing the P element-bearing lines to the [P(2-3)]9B transposase source (Robertson et al., 1988; Cooley et al., 1988). Sequences flanking the P element insertions were cloned by plasmid rescue, using the bacterial origin and \( \beta \)-lactamase gene engineered into the P element (Bier et al., 1989).

Complementation of KLP38B Mutants by a KLP38B cDNA

KLP38B cDNA was excised from the vector pNB40 (Brown and Kafatos, 1988) by digestion with HindIII and NotI. The Drosophila expression vector pWsg26 (Lopez et al., 1994) was linearized with EcoRI. These two molecules were end-filled and ligated together to place the KLP38B cDNA under the control of the hs26 promoter. This construct was then injected into embryos carrying the P[\( \Delta \)-3]9B transposase source (Robertson et al., 1988; Cooley et al., 1988). Transformed progeny of these injected individuals were identified by virtue of the \( w^+ \) marker in pWsg26. A third chromosome insertion was selected. Standard breeding strategies were used to construct flies homozygous or hemizygous for KLP38B mutants and also carrying a single copy of the hs26-KLP38B transgene. The phenotype of these flies was analyzed in comparison with flies of the same KLP38B genotype but lacking the transgene.

Cytological Preparations

Preparation, fixation, antigen detection, and 4',6-diamidino-2-phenylindole (DAPI) staining of ovaries, embryos, and larval brains was described in Gonzalez and Glover (1993). Embryos were incubated in 5 mM taxol for 30 s to stabilize microtubules, as recommended by these authors. The anti-tubulin antibody used was mouse monoclonal TAT-1 (Woods et al., 1989), a gift from Keith Gull (University of Manchester, UK). Texas red–conjugated secondary antibody was purchased from Sigma (Sigma-Aldrich Co., Dorset, UK). RNA in situ hybridization was performed according to Tautz and Peitlle (1989) and Gonzalez and Glover (1993).

SEM

Adult flies were anesthetized with CO\(_2\), mounted on a stub with carbon dag, ventral surface to the stub, and then placed in the low vacuum chamber of an ElectroScan Environmental Scanning Electron Microscope (Cambridge Instruments, Welwyn Garden City, UK). Specimens were observed at 20 kV. Digital images were recorded and manipulated using Adobe Photoshop (Adobe Systems, Inc., Mountain View, CA).

Microscopy

Fluorescence microscopy was performed using a DM IRBE microscope (Leitz, Wetzlar, Germany). Images were captured using a cooled slow-scan CCD camera (CH250; Photometrics, Ltd., Tucson, AZ), using IP Lab Spectrum software (Signal Analytics, Co., Vienna, VA). This software was also used for pixel counting. Deconvolution was performed using Power Hazebuster software (Vaytek, Inc., Fairfield, IA), and the images were assembled into figures using Adobe Photoshop. Confocal microscopy was performed using a Leitz DM IRBE microscope with Leica TCS 4D confocal system including UV laser.

Protein Expression and In Vitro Binding

A PPI 87B cDNA with the start codon mutated to an NdeI site (a gift from David Glover, Dundee University, Dundee, UK) was subcloned into pET-16b (Novagen, Madison, WI). After induction in the expression host cells BLR(DE3) (Novagen), pET-16b–PPI 87B expresses full-length PPI 87B protein with an N-terminal fusion encoding His\(_6\), allowing affinity purification on immobilized Ni\(^{2+}\) ions. PPI 87B with no insert was used as a control; this encodes a short (48-aa) protein including all the vector sequences present in the His\(_6\)-PPI 87B protein.

35S-labeled KLP38B and procollagen were synthesized by in vitro transcription–translation (Riboprobe and Flexi rabbit reticulocyte lysate kits; Promega, Madison, WI) from full-length KLP38B cDNA and polI(III)\( \alpha \) (Lees and Bulleid, 1994), respectively. Soluble extracts of BLR(DE3) containing oligo-histidine–tagged PPI 87B or control pET-16b protein were prepared and incubated with 10 \( \mu \)l Ni-NTA Sepharose in buffer A (50 mM Tris-HCl, \( \fi \), 100 mM KCl, 5 mM MgCl\(_2\), 0.1% Triton X-100, 20% glycerol, 10 mM imidazole) for 1 h at 4°C. The Ni-NTA beads were washed four times in this buffer, and then incubated with 1 \( \mu \)l reticulocyte lysate containing 35S-labeled KLP38B or procollagen in 100 \( \mu \)l buffer A for 30 min at room temperature. The beads were washed an additional four times in buffer A supplemented with imidazole to 50 mM, and then eluted with 20 \( \mu \)l of buffer A containing 200 mM imidazole. 10 \( \mu \)l of each eluate was analyzed by SDS-PAGE and phosphoimaging (Fuji Bas 2000; Fuji Photo Film Co., Tokyo, Japan). 0.5 \( \mu \)l of reticulocyte lysate was loaded for comparison (see Fig. 6, lanes I and 4).

Results

Isolation of KLP 38B

The PPI catalytic subunit will dephosphorylate a wide variety of substrates in vitro, but in vivo is found associated with a range of targeting and regulatory subunits that modulate its subcellular localization and substrate specificity (Hubbard and Cohen, 1993). To obtain potential PPI regulators, we used the yeast two-hybrid interaction trap (for review see Fields and Sternglanz, 1994). This system identifies cDNAs encoding polypeptides that will bind to a “bait” protein in yeast. We used a cDNA library from Drosophila third instar larval RNA (see Materials and Methods). Drosophila has four genes encoding PPI isoforms (Dombrádi et al., 1999, 1993), but 80% of the total PPI activity is contributed by the PPI 87B isoform (Axton et al., 1990). Using full-length PPI 87B cDNA fused to GAL4 as the bait, 25 independent cDNAs were identified that interacted with PPI 87B but not with control baits. Sequencing and in situ hybridization to polytene chromosomes revealed that these 25 cDNAs are derived from 16 genes. One of these genes shows strong sequence similarity to NIPP-1, a biochemically characterized mammalian PPI inhibitor (Van Eynde et al., 1995). Another, identified by three independent cDNAs from the two-hybrid screen, showed sequence similarity to kinesin-like proteins. These cDNAs hybridize to cytological position 38B on chromosome 2L. We refer to this gene as KLP38B (kinesin-like protein at 38B), following established convention (Goldstein, 1993). Using sequence information from these two-hybrid
KLP38B cDNAs, we used a PCR-based screen to isolate a 3.6-kb cDNA from an imaginal disk library (see Materials and Methods). This cDNA contains a single long ORF encoding a kinesin-related protein (Fig. 1). Based on translation from the first ATG, the sequence predicts a protein of 1,121 amino acids with a molecular mass of ~125 kD and an isoelectric point (pI) of 7.6. Sequence analysis suggests the existence of four distinct domains (Figs. 1 and 2). Amino acid residues 1–122 at the NH₂ terminus share no strong similarity with other known sequences. By comparison with other kinesin-related proteins, residues 123 to ~512 encode the motor domain (Goldstein, 1993), while similarity to most other KRPs ends at residue 528. COOH-terminal to the motor domain is a 210 region of sequence similarity to Unc-104, Kif1A, Kif1B, and OrfW, but not to other known KRPs (Fig. 2). Within this region these five proteins share 39% similarity; 27% of these residues are identical between KLP38B and Unc-104, and 29% between KLP38B and OrfW. The motor domain of KLP38B is also more similar to Unc-104-like KRPs than to others.

The remaining 300 residues at the COOH terminus show no significant similarity to other known proteins. On the basis of these comparisons, we place KLP38B in the Unc-104-like subfamily of kinesin-related proteins (Bloom and Endow, 1995; Moore and Endow, 1996; see Discussion). Comparisons of either the motor domain or the entire sequence do not indicate that KLP38B is more closely related to any one Unc-104-like KRP than to another.

Kinesin heavy chains fold into an α-helical coiled-coil in the stalk domain to form a dimer (Yang et al., 1989; de Cuevas et al., 1992). Most kinesin-related proteins are similarly thought to form homodimers, mediated by an extended coiled-coil region, although tetrameric, heterotrimeric, and monomeric kinesin-related proteins have been described (Cole et al., 1993, 1994; Bloom and Endow, 1995). We therefore used the PEPCOIL (Lupas et al., 1991; Lupas, 1996) and PAIRCOIL (Berger et al., 1995) programs to examine the KLP38B sequence for regions likely to form a coiled-coil. The PEPCOIL prediction is shown in Fig. 2 C. The PAIRCOIL output is essentially identical except that there is no predicted coiled-coil region near the COOH terminus (amino acids 1,061–1,077, see Fig. 2 C). This comparatively low level of coiled-coil is comparable to Kif1A (Okada et al., 1995) and Kif1B (Nangaku et al., 1994), which are murine KRPs related to KLP38B and Unc-104 (Fig. 2 A). Recombinant Kif1A and Kif1B exist predominantly as globular monomers, as judged by gel filtration, native PAGE, differential light scattering (Kif1A only), and EM. This suggests that KLP38B may similarly exist predominantly as a monomer.

Sequencing another four cDNAs revealed a number of single base changes relative to the sequence of the 3,605-bp cDNA. These could represent natural polymorphism or mutations induced in the process of cDNA synthesis. The majority of these variants are silent in terms of predicted polypeptide sequence, suggesting that these variants represent natural polymorphism. The single variant affecting the amino acid sequence of the motor domain (S454I)

Figure 1. Sequence of the KLP38B cDNA. The sequence of the coding strand of the longest available KLP38B cDNA is depicted with the predicted 1,121–amino acid sequence of its product, based on translation from the first ATG (see Fig. 2 and text). The putative motor domain is shown with a double underline, regions predicted to form coiled-coils by PAIRCOIL with a single underline (see Fig. 2 C). These sequence data are available from EMBL under accession number X99617.
potential cdk phosphorylation site is not related to the BimC/Eg5 subfamily. Comparison of the putative cdk phosphorylation site in KLP38B with that of BimC/Eg5-like kinesin-related proteins. Of the residues common to the BimC/Eg5 sequences, only the core cdk consensus (S/T P × K/R) and a leucine are also present in KLP38B. (C) Predicted coiled-coil formation by KLP38B. PEPCOIL output: probabilities of the KLP38B protein forming α-helical coiled-coils as predicted from the algorithms of Lupas et al. (1991), using a 28-residue window. The horizontal axis is position within the protein, and the NH2 terminus is at the origin; the vertical axis is probability of coiled-coil formation. Open boxes above the plot indicate the frame of the coiled-coil repeat. (D) Domain structure of KLP38B. The motor domain, region of extended sequence similarity with Unc-104–related KRPs, and the COOH-terminal PPI-binding region are shown at the same scale as in C. The PPI-binding region is defined by the shortest KLP38B clone from the two-hybrid screen.

maps to a nonconserved loop. Details of these variants have been deposited in the EMBL database, accession number X99617.

Inspection of the sequence shows a potential cdk phosphorylation site at residue 1,108, close to the COOH terminus. Cdk phosphorylation has been shown to be important for function in Eg5, where phosphorylation is required for spindle association (Sawin and Mitchison, 1995; Blangy et al., 1995; for review see Walczak and Mitchison, 1996). The COOH-terminal cdk phosphorylation site in KLP38B does not have the additional residues characteristic of the BimC/Eg5 subfamily. Unc-104, Kif1A, Kif1B, and OrfW also have potential cdk phosphorylation sites; these also share no extended similarity with the BimC/Eg5 subfamily or with each other (not shown).

**Transcription Pattern of KLP38B**

We have studied the expression pattern of KLP38B by RNA in situ hybridization to various *Drosophila* tissues and developmental stages (Fig. 3). KLP38B is expressed in proliferating cells at every developmental stage, as well as in the testis and ovary. In the ovarioles, KLP38B is expressed from the gerarium onward. Expression is much higher in the nurse cells at later stages, leading to a maternal contribution in the syncytial embryo (not shown). Later in development, KLP38B is expressed in the proliferating cells of the imaginal disks (e.g., eye disk, Fig. 3 B) and central nervous system (Fig. 3 A). In the testis, KLP38B transcripts accumulate during the 90-h growing stage and are degraded shortly after meiosis (Fig. 3 C; see Discussion).

**Isolation of KLP38B Mutants**

The major reason for using *Drosophila* as a model system in which to investigate KRPs and PPI is the advanced genetics and developmental biology. Gene function can be determined by analyzing the phenotype of mutants defective for the gene. We therefore isolated deficiencies and mutants for KLP38B as follows.

KLP38B was localized to cytological position 38B on chromosome 2L by in situ hybridization of the cDNA to wild-type polytene chromosomes (data not shown). This mapping was further refined by hybridization to deficiency stocks. This showed that KLP38B lies within Df(2L)pr-A20 (38A3-4 to 38B6-C1) and Df(2L)TW9 (37E2-F4 to 38A6-C1). This defines the cytological location as 38A3-38C1, which is consistent with the initial localization to 38B. A PCR-based in situ hybridization experiment aimed at estimating the number and cytogenetic location of genes encoding *Drosophila* kinesins and kinesin-related proteins had previously suggested that there is such a gene at 38B (Endow and Hatsumi, 1991).

Cosmid 164E4 from the European *Drosophila* Genome Project (Siden-Kiamos et al., 1990; Kafatos et al., 1991) was identified as containing the KLP38B gene by hybridization to a KLP38B-specific probe. The KLP38B cDNA was mapped onto the cosmid by restriction mapping and further
ther hybridization to determine the gene organization (Fig. 4). A 7-kb BamHI–EcoRI fragment from the middle of the gene was used to identify recessive lethal P element mutants with insertions in the KLP38B gene, using plasmid rescue from a large collection of P element–induced lethal and semilethal mutants (see Materials and Methods). Six such mutants were identified. Two further P element-induced alleles of \textit{KLP38B}, designated 24-O and 93-E, were generously provided by Mike Goldberg. We refer to these alleles as \textit{KLP38B}\textsuperscript{24-O} and \textit{KLP38B}\textsuperscript{93-E}. Consistent with the in situ mapping, these P element mutants are all semilethal over \textit{Df(2L)pr-A20}.

All \textit{KLP38B} alleles were tested in combination with each other and over \textit{Df(2L)pr-A20}, a deficiency that removes the \textit{KLP38B} gene. The observed phenotypes of all the \textit{KLP38B} alleles, and \textit{Df(2L)pr-A20}, are entirely recessive. Several of the mutant chromosomes are completely lethal when homozygous, but only semilethal or sterile over other alleles or over \textit{Df(2L)pr-A20}, indicating that they have one or more additional recessive lethal mutations on the same chromosome. The phenotype of \textit{KLP38B}\textsuperscript{24-O} homozygotes is indistinguishable from that of \textit{KLP38B}\textsuperscript{24-O}/\textit{Df(2L)pr-A20} hemizygotes, so we conclude that \textit{KLP38B}\textsuperscript{24-O} has little or no residual \textit{KLP38B} activity and that \textit{KLP38B} activity is not absolutely required for viability (see Discussion).

When using mutant chromosomes carrying a P element, there is a possibility that the observed phenotype is not due to the P element insertion, but rather to another mutation on the same chromosome. This can be tested by excising the P element and observing the new phenotype: complete reversion to wild type indicates that the mutant phenotype is entirely due to the P element insertion. Using the strongest mutant, \textit{KLP38B}\textsuperscript{24-O}, we found that excision of the P[\textit{lacW}] element resulted in complete phenotypic reversion, thus demonstrating that the phenotypes observed are due to the insertion of P[\textit{lacW}] at the \textit{KLP38B} locus.

Another possibility is that the P element insertions affect one or more genes other than \textit{KLP38B}. We therefore created transgenic flies carrying the \textit{KLP38B} cDNA under the control of the hsp26 promoter. Even without heat shock, this construct rescues the sterility of \textit{KLP38B}\textsuperscript{24-O} homozygotes and of \textit{l(2)k03903}/\textit{Df(2L)pr-A20} hemizygotes. Similarly, the visible phenotypes are much less se-

\textbf{Figure 4.} Molecular map of the \textit{KLP38B} region. The P[\textit{lacW}] element insertions in \textit{KLP38B} mutants are shown in relation to the \textit{KLP38B} exons and intron. The P[\textit{lacW}] insertion site of \textit{KLP38B}\textsuperscript{24-O} maps \textasciitilde 3 kb away from the other insertions, which are too close to separate on this scale. Of these other mutants, the insertion site in \textit{l(2)k03903} is the most 3’ relative to the \textit{KLP38B}, while \textit{l(2)k04805}, \textit{l(2)k04912}, \textit{l(2)k04914}, and \textit{l(2)k05702} have identical insertion sites 53 bp away and the \textit{KLP38B}\textsuperscript{93-E} insertion is another 242 bp further toward the 5’ end of \textit{KLP38B}. The central gray line represents genomic DNA, the \textit{KLP38B} exon map is below, and the P element insertion sites are marked above. Asterisks mark the ends of the BamHI–EcoRI fragment used to identify the mutants. B, BamHI; R, EcoRI; X, Xbal.
vere in these flies. We therefore conclude that the phenotypes we observe in the P element mutants are entirely due to disruption of the \textit{KLP38B} gene.

\textbf{The Molecular Basis of the KLP38B Mutations}

Plasmid rescue was repeated on each of the P element lines. Restriction maps of the recovered plasmids were compared with the genomic map from cosmid 164E4 to generate a molecular map of the \textit{KLP38B} locus and mutants. All of the P element insertions map to the \textit{KLP38B} intron. The recovered plasmids were sequenced using a P element–specific primer to determine the exact insertion sites of the P elements. These data are summarized in Fig. 4.

\textbf{KLP38B Is Required for Normal Cell Proliferation}

The range of phenotypes observed—rough eyes (Fig. 5, A–C), missing bristles (Fig. 5, D–F, and Table I), reduced viability, and male and female sterility—correlates with our RNA in situ data (Fig. 3; unpublished observations) and suggests a role for \textit{KLP38B} in cell proliferation and gametogenesis (see Discussion). We have also observed an “upheld wing” phenotype in adults, in which the wing blade is held vertically rather than horizontally. Newly eclosed flies (young adults) do not show this phenotype, rather it develops a few days after eclosion. Affected individuals cannot fly. The “upheld wing” phenotype has been observed in mutants affecting development of the indirect flight muscles (DeSimone et al., 1996), but examination of these muscles dissected from \textit{KLP38B}24-O homozygotes revealed no obvious abnormalities in muscle structure. Homozygous adults also have poor locomotive ability.

We examined the male sterility of \textit{KLP38B}24-O homozygotes in more detail. Squashed preparations of testes from \textit{KLP38B}24-O homozygous males were examined by phase-contrast microscopy. This method allows the detection of meiosis and cytokinesis defects by measuring the number of postmeiotic spermatids and the relative size and number of their nuclei and Nebenkerns, which are the mitochondrial derivatives (for examples see Alphey et al., 1992; Williams et al., 1995). No such gross meiotic defect was observed, but no motile sperm were present (not shown).

\textbf{KLP38B Is Required for Female Fertility and Follicle Cell Proliferation}

\textit{KLP38B}24-O homozygous females are completely sterile. Mutant ovaries are small, with egg chambers rarely developing past stage 5 of oogenesis (stages according to King, 1970; Mahowald and Kambysellis, 1980; see Spradling, 1993, for review of oogenesis in \textit{Drosophila}). Further investigation of the mutant ovaries revealed a failure of folli-

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\centering
\includegraphics[width=\textwidth]{figure5.png}
\caption{\textit{KLP38B} mutants show rough eyes and missing bristles. Scanning EM shows clear defects in the eye structure of \textit{KLP38B}24-O homozygotes (B and C) compared with the highly ordered wild-type eye (A). Facets are of uneven size and shape and many bristles are missing, consistent with aberrant cell proliferation. Similarly, the wild-type array of abdominal bristles (D) is disrupted in \textit{KLP38B}24-O homozygotes (E and F). C and F are details from B and E, respectively. Bars, 50 \textmu m.}
\end{figure}
The larger egg chambers are not completely covered by follicle cells (Fig. 6). In wild-type ovaries, follicle cells divide four or five times during stages 1–5, giving \(1,000\) follicle cells. These then cease cell division but increase in size by polyploidization. At stage 9 the majority of the cells migrate over the expanding oocyte and form a columnar epithelium, while \(50\) remain as a thin layer over the nurse cells and \(6–10\) migrate between the nurse cells to become the border cells. The majority of \(KLP38B^{24-O}\) egg chambers arrest at stage 5 or earlier. No eggs are laid by these females. DAPI staining clearly shows that these egg chambers have a much lower number of follicle cells than wild type (Fig. 6, A and B).

### KLP38B Is Required for Mitotic Chromatin Condensation

As described above, \(KLP38B^{24-O}\) homozygous females are completely sterile and produce no mature oocytes. A weaker allele, \(l(2)k03903\), does lay some eggs but these very rarely hatch. \(l(2)k03903\) also has somewhat less severe rough eyes and bristle defects than \(KLP38B^{24-O}\) and is male fertile. We investigated the basis of the maternal effect lethality in embryos derived from \(l(2)k03903\) females. For clarity, we will refer to these as k03903 embryos.

The majority of k03903 embryos fail to cellularize. Those that do cellularize appear to have uneven cell size and distribution compared with wild type, but gastrulate apparently normally even though they generally fail to hatch. This is not unexpected by comparison with the zygotic phenotype of \(string\) and other cell cycle genes, in which failure of mitosis leads to an embryo with far less cells than wild type, which nonetheless gastrulates and develops relatively normally but fails to hatch (Hartenstein and Posakony, 1990; Foe et al., 1993). k03903 embryos fail to form pole cells. They also have a high frequency of “drop-out” nuclei (not shown). These are nuclei that are eliminated from the surface blastoderm layer and fall into the interior, and are indicative of a cell cycle defect (Foe et al., 1993; see Discussion).

We have examined syncytial and newly cellularized k03903 embryos by DAPI staining and immunofluorescence with anti-tubulin antibodies and find metaphase and anaphase spindles associated with undercondensed chromatin (Fig. 7). This phenotype is clearly distinguishable from wild type and is 100% penetrant: every metaphase or anaphase figure from k03903 embryos has undercondensed chromater.

### Table I. Scutellar Bristle Defects in KLP38B Mutants

<table>
<thead>
<tr>
<th>Bristle No.</th>
<th>(KLP38B^{24-O})</th>
<th>l(2)k03903</th>
<th>l(2)k05702</th>
<th>Oregon R</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anterior</td>
<td>67</td>
<td>30</td>
<td>3</td>
<td>60</td>
</tr>
<tr>
<td>Posterior</td>
<td>76</td>
<td>24</td>
<td>0</td>
<td>82</td>
</tr>
<tr>
<td>Duplicated</td>
<td>0</td>
<td>3</td>
<td>97</td>
<td>0</td>
</tr>
<tr>
<td>Extra</td>
<td>0</td>
<td>3</td>
<td>97</td>
<td>0</td>
</tr>
</tbody>
</table>

Wild-type flies have four scutellar bristles, two anterior and two posterior. Disruption of this pattern is frequently observed in cell cycle mutants. Scutellar bristles were counted for 100 homozygous individuals of each of the three \(KLP38B\) mutant alleles. Oregon R is a standard wild-type strain. Duplicated bristles are two bristles from the same socket, or two bristles immediately adjacent to each other; extra bristles are bristles not in the characteristic location.
Figure 7. k03903 embryos undergo mitosis with undercondensed chromatin. Metaphase (A–H) and anaphase (I–P) figures from wild-type and k03903 embryos. k03903 embryos have undercondensed chromatin at both metaphase (F and H) and anaphase (N and P) relative to wild type (B and D, and J and L, respectively). k03903 spindles are generally similar to those of wild type, but a range of defects is occasionally seen including kinked spindles (G). Mitotic figures are shown in pairs, with tubulin staining on the left and DAPI on the right. Bar, 5 μm.
KLP38B Is Required for Normal Cell Cycle Progression

Cell cycle progression in Drosophila embryos is unusual in a number of respects (for review see Foe et al., 1993). Syncytial embryos have no detectable G1 or G2 phases, and of course have no cytokinesis. They appear to lack many of the checks and controls present in a normal somatic cell cycle. By contrast, neuroblasts in the brain of wandering third instar larvae follow a typical cell cycle of G1-S-G2-M. We therefore examined these cells from KLP38B mutants.

The mutant phenotype is characterized by an increased mitotic index (21% of 2,565 cells scored in acetic acid–orcein squashes of 14 KLP38B mutant embryos compared with 6.8% for wild-type brains). Approximately two-thirds of these mitotic figures are polyplody. The diploid cells have a metaphase/anaphase ratio not significantly different from wild type (2.6), whereas the polyploid cells have a metaphase/anaphase ratio of 46. Chromatin condensation varies from one cell to another (see Discussion).

We further investigated these abnormal mitotic figures by confocal microscopy of whole brains (Fig. 8). Many of these polyploid cells have chromosomes arranged on monopolar spindles, indicating a centrosome duplication or separation defect. Fig. 8, A–J, is a series of confocal sections through such a cell. Fig. 8, K and L, shows a comparable wild-type metaphase (note the difference in magnification between A–J and K–M). Fig. 8 M shows a field of mitotic cells, illustrating the elevated mitotic index.

KLP38B Associates with PP1

As described above, KLP38B was originally isolated as a cDNA encoding a protein capable of binding to PP1 87B, the major PP1 catalytic subunit of Drosophila. These cDNAs were isolated: they differ in length from each other and thus are independent and not derived from clonal amplification of the same original cDNA. All three encode the COOH-terminal region of KLP38B, encoding amino acids 587–1,121, 725–1,121, and 848–1,121, respectively. This demonstrates that the COOH-terminal 273 residues are sufficient to bind PP1.

While the two-hybrid system has now been successfully used many times to demonstrate protein–protein binding, it is also known to be capable of producing false positives. We therefore investigated the association of KLP38B and PP1 87B by a completely independent method. We expressed full-length PP1 87B in bacteria, with an oligo-histidine tag to allow affinity chromatography using immobilized Ni$^{2+}$ ions. We bound PP1 87B to Ni-NTA Sepharose, and then incubated these beads with $^{35}$S-labeled KLP38B from an in vitro transcription–translation reaction. After extensive washing we eluted the PP1 87B by increasing the imidazole concentration and analyzed the samples by SDS-PAGE followed by phosphoimaging. We found that 35% of the KLP38B remained bound to the column through the washes and coeluted with PP1 87B (Fig. 9, lane 6). No KLP38B (<0.5%) bound to control beads prepared using extracts from bacteria that cannot synthesize PP1 87B (Fig. 9, lane 5). We conclude that KLP38B can bind to PP1 87B in vitro. A procollagen control does not bind to PP1 under the same conditions. Interestingly, phosphorylation of KLP38B is not required for binding to PP1, as bacterially expressed KLP38B also binds to PP1 (not shown).

Discussion

KLP38B Structure and Function

Structural similarities between kinesin-related proteins in many cases reflect functional similarities. For example, the BimC-related kinesin-related proteins Bim5, KLP61F, Cin6/Kip1, and Cut7 are all involved in the formation of bipolar spindles (Enos and Morris, 1990; Hagan and Yana-gida, 1990; Hoyt et al., 1992; Heck et al., 1993; Kashina et al., 1996a,b). Loss-of-function mutations of these KRPks are able to duplicate their centrosomes/spindle pole bodies but fail to separate them. Of the Unc-104–like KRPs, Kif1A is a neuron-specific vesicle transporter and is thought to be a functional homologue of Unc-104 (Okada et al., 1995). Kif1B has been found to be associated with mitochondria and is capable of transporting mitochondria along microtubules in vitro (Nangaku et al., 1994). No function is known for OrfW, which was detected in a large-scale sequencing project (Nomura et al., 1994). Loss-of-function unc-104 mutants are paralyzed; they show correct axon formation, but few synapses and synaptic vesicles accumulate in the neuron cell body (Hall and Hedgcock, 1991). This suggests that Unc-104 transports synaptic vesicles along axonal microtubules. The functions ascribed to Kif1B and Unc-104 are clearly very different from the mitotic role we have demonstrated for KLP38B. Drosophila also has another KRP more closely related to Unc-104. For these reasons, we do not believe that KLP38B is a true functional homologue of Unc-104.

The Mitotic Role of KLP38B

The expression pattern of KLP38B in proliferating cells suggests a role for KLP38B in mitosis. This is confirmed by the phenotype of KLP38B mutants. The effect of cell cycle mutants is often seen only in late larval/pupal stages (Gatti and Baker, 1989), as most larval growth is by cell enlargement. Survivors typically show rough eyes and missing...
bristles, as a consequence of occasional failure of proliferation in the cells from which these structures are derived. Maternal effect lethality is another well-known phenotype of mutation in cell cycle genes: the mother has to provide the embryo with all the components necessary for 13 rapid nuclear divisions, so lack of any gene product required for mitosis will lead to nonviable embryos (Gatti and Baker, 1989; Sullivan et al., 1990; Foe et al., 1993; Gonzalez et al., 1994). KLP38B mutants show all of these phenotypes.

KLP38B 24-O escapers are female sterile: egg chambers do not develop beyond the early stages because of failure of follicle cell proliferation. Embryos derived from l(3)k03903 homozygous females (“k03903 embryos”) have abnormal mitotic figures, with undercondensed chromatin on a metaphase-like spindle. This implies a role for KLP38B in the establishment or maintenance of mitotic chromatin condensation. We cannot rule out the possibility of transient chromatin condensation in these embryos, but we have never observed properly condensed chromatin, so any such condensation cannot last long. Undercondensed chromatin on a metaphase-like spindle could be explained by a metaphase arrest or delay followed by partial decondensation of the chromatin, but we also find that all the anaphase-like figures have undercondensed chromatin, which suggests that these embryos go through a complete mitosis with undercondensed chromatin. These embryos also exhibit a low frequency of spindle abnormalities, which may be a secondary consequence of the abnormal chromatin. We also observe occasional chromatin bridges in cycle 14 (newly cellularized) embryos, as well as some cytokinesis defects, possibly as a consequence of the chromatin bridges.

The syncytial embryo may be incapable of arresting its mitotic cycles in response to problems in chromosome condensation or segregation and instead eliminates the defective nuclei from the blastoderm layer (Sullivan et al., 1990; Foe et al., 1993). Cell divisions later in development have more checkpoints and controls and may be able to

Figure 8. Mitotic defects in KLP38B 24-O neuroblasts. (A–J) Confocal sections through a KLP38B 24-O neuroblast. A–E are DAPI staining; F–J show the tubulin. Sections are at 1-μm intervals. A wild-type metaphase is shown for comparison K and L. M is a confocal section through a field of mitotic cells. Only a single anaphase is present; the remainder are metaphases in various orientations. Bars, 5 μm.
Differential control of spindle formation and chromatin condensation by cdc25 homologues has been previously observed (White Cooper et al., 1993; Gabrielli et al., 1996). Vernos et al. (1995) analyzed XKlp1 function by depletion experiments using antisense oligos or anti-XKlp1 antibodies. They concluded that XKlp1 is required for chromosome positioning and bipolar spindle stabilization. The delocalized chromosomes that they observe (Fig. 8 of Vernos et al., 1995) also appear to be unevenly condensed, which might imply a role for XKlp1 in chromatin condensation. Another precedent for KRP involvement in genome organization comes from fission yeast, where microtubule-based movement of chromosomes through attachment of telomeres to the spindle pole body is implicated in homologue recognition (for review see Kohli, 1994).

KLP38B transcripts accumulate in the growing stage of spermatogenesis and are degraded shortly after meiosis. This is very similar to the testis expression pattern of the cdc25 homologue twin (Alphey et al., 1992) and other cell cycle-regulatory genes required for meiosis (Alphey, L., unpublished observations). However, there is little or no postmeiotic transcription in the male germ line (Olivieri and Olivieri, 1965; Gould-Somero and Holland, 1974; for review see Fuller, 1993), so genes involved in postmeiotic differentiation are expected to follow a similar transcription pattern; the gene products are then stored as mRNA or protein until required. Since the male sterility seen in KLP38B mutants does not appear to be associated with severe meiotic defects, we conclude that KLP38B either has no role in male meiosis, or else its role is redundant with another kinesin-related protein.

Does KLP38B Have Nonmitotic Roles?

We have demonstrated a role for KLP38B in mitosis. This does not exclude the possibility that KLP38B may have additional roles. The observation that KLP38B mRNA is present in postembryonic tissues only in cells that are about to divide suggests that KLP38B acts only in mitosis and does not function in nonmitotic microtubule-based transport in these tissues. As noted above, we have observed that males hemizygous or homozygous mutant for KLP38B+4- are completely sterile. These flies produce no motile sperm but appear to complete meiosis and cytokinesis normally. We therefore deduce a role for KLP38B in postmeiotic spermatid differentiation, in addition to the mitotic role.

Redundancy of KLP38B with Other Kinesin-related Proteins

The phenotypes of KLP38B+4- /KLP38B+4- and KLP38B+4- / Df(2L)pri-A20 are indistinguishable, indicating that this allele has very little, if any, residual KLP38B activity. Despite this finding, none of our KLP38B alleles are completely lethal. This suggests a degree of functional redundancy with other Drosophila kinesin-related pro-
teins. Such redundancy has been observed before in the case of CIN8/KIP1, two redundant budding yeast kinesin-related proteins (Hoyt et al., 1992). Another possibility is perdurance of the maternal product, which has many precedents among Drosophila cell cycle genes (for review see Gonzalez et al., 1994). We have observed a reduction in the penetrance of the lethality and scutellar bristle phenotypes of KLP38B<sup>24-0</sup> in a continuously maintained culture over a 12-mo period, such that the viability is now considerably higher (30–50% relative to heterozygous siblings) and the penetrance of the scutellar bristle defects lower than a year ago (data not shown). This suggests that the stock can readily accumulate modifiers, perhaps subtle allelic variants of genes with overlapping functions. The complete sterility of KLP38B<sup>24-0</sup> homozygotes, both males and females, demonstrates an absolute requirement for the KLP38B gene product in gametogenesis. This is reminiscent of KLP3A mutants, which are sterile but viable, even though the KLP3A protein is present in proliferating somatic cells (Williams et al., 1995).

**KLP38B and PP1**

We have demonstrated binding of KLP38B to PP1 in vitro. Drosophila melanogaster has four genes encoding isoforms of PP1 (Dombrádi et al., 1990, 1993). Strong alleles of PP1<sup>87B</sup> show PP1 activity reduced to ~20% of wild type in extracts of larvae and adults, indicating that this gene is the major isoform at these stages (Baksa et al., 1993). These mutants have a strong cell cycle phenotype: neuroblasts are delayed in mitosis and show aberrant spindle organization and hypercondensed chromatin. Some alleles are also dominant suppressors of position effect variegation, indicating that PP1 has an interphase role in the regulation of chromatin structure as well as a mitotic role. Other roles of PP1 may be masked by redundancy between the isoenzymes and other related protein phosphatases.

If PP1<sup>87B</sup> and KLP38B associate in the fly, as suggested by our in vitro experiments, this suggests an explanation for their observed phenotypes: KLP38B is required for normal chromatin condensation and is inhibited by PP1<sup>87B</sup>. PP1 has been found in partially purified axonal kinesin, and inhibition of the phosphatase with okadaic acid stimulates kinesin activity (McIlvain et al., 1994a; McIlvain, J.M., Jr., L. Lindesmith, Y. Argon, and M.P. Sheetz. 1994. Mol. Cell. Biol. 5(Suppl.):31a). However, the true situation may not be so simple. Both PP1<sup>87B</sup> and KLP38B are pleiotropic; furthermore, they are both members of gene families with overlapping functions. Axton et al. (1990) suggest that the cell cycle effects of PP1<sup>87B</sup> mutants may be due in part to the presence of excess free regulatory subunits interfering with the function of another PP1 isoenzyme or with other proteins essential for mitosis.

Another attractive hypothesis is that KLP38B may be involved in localizing PP1. The PP1 catalytic subunit will dephosphorylate a wide range of substrates in vitro, but in vivo is found associated with targeting and regulatory proteins that modulate its subcellular location and substrate specificity, and it seems likely that there is one or more specific subunits for each cellular function of the enzyme (Hubbard and Cohen, 1993). Association with KLP38B may be involved in the regulation of PP1 regarding chromatin condensation or spindle function.

Regulation of the cell cycle, motor activity, spindle structure, and chromatin condensation by phosphorylation has been found in a variety of systems. We have identified a novel kinesin-related protein (KLP38B) in a screen for PP1-binding proteins and demonstrated for the first time a physical association between a kinesin-related protein and a protein phosphatase. Our phenotypic analysis of mutants defective for KLP38B demonstrates a requirement for this gene for mitotic chromatin condensation and cell proliferation.

We are grateful to the following colleagues who sent us reagents: Stephen Elledge for the yeast two-hybrid system, including an unpublished Drosophila cDNA library; Myles Axton and David Glover for PP1<sup>87B</sup> cDNA; Istvan Torok and Mike Goldberg for P element mutants; Richard Wilson and Neil Bulleid for procollagen cDNA and assistance with in vitro translation. Doug Ruden, Pedro Ripoll, and Sharyn Endow generously shared unpublished data. We thank Robert Saunders for help with cytogenetics, and Trevor Sherwin and Klaus Ernfled for help with image analysis. Computing facilities at SEQNET and HGMP were made available by the United Kingdom Biotechnology and Biological Sciences Research Council and Medical Research Council, respectively. We also thank Viki Allan, Doug Drummond, and Iain Hagan for helpful discussions and critical reading of the manuscript.

This work was principally supported by grant SP229001010 from the Cancer Research Campaign to L. Alphey, with additional funding from the Royal Society (to L. Alphey) and United Kingdom Biotechnology and Biological Sciences Research Council (to K. Kaiser). Confocal and fluorescence microscopes and semiautomated sequencing facilities were available through the generosity of the Wellcome Trust (grants 045183/Z/95/Z and 044327/Z/95/Z, respectively).

Received for publication 23 December 1996 and in revised form 14 April 1997.

**References**


