A “molecular guillotine” reveals an interphase function of Kinesin-5

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Abstract
Motor proteins are important for transport and force generation in a variety of cellular processes and morphogenesis. Here we design a general strategy for conditional motor mutants by inserting a protease cleavage site at the “neck” between the head domain and the stalk of the motor protein, making the protein susceptible to proteolytic cleavage at the neck by the corresponding protease. To demonstrate the feasibility of this approach, we inserted the cleavage site of TEV protease into the neck of the tetrameric motor Kinesin-5. Application of TEV protease led to a specific depletion and functional loss of Kinesin-5 in Drosophila embryos. By this, we revealed that Kinesin-5 stabilized the microtubule network during interphase in syncytial embryos. The “molecular guillotine” can potentially be applied to many motor proteins due to the conserved structures of kinesin, dynein and myosin with accessible necks.

Author summary
We design a general strategy for conditional motor mutants by inserting a protease cleavage site between head and stalk domain of the motor protein, making it susceptible to specific proteolytic cleavage. We demonstrate the feasibility of the approach with the motor Kinesin-5 and the protease TEV in Drosophila embryos. This approach can potentially be applied to motor proteins kinesin, dynein and myosin due to the conserved structures.

Introduction

Cytoskeletal motor proteins, including myosins, dyneins and kinesins, convert the chemical energy of ATP hydrolysis into mechanical work. Motor proteins are wildly involved in multiple fundamental cellular processes such as intracellular transport, cell division, cell shape change and migration [1]. The structure of motor proteins is conserved. They contain a motor domain, referred to as “head”, which catalyzes ATP and binds microtubules or F-actin. The catalytic cycle links ATP hydrolysis to a conformational change of the protein that translates into unidirectional movement of the motor protein on the filament. A second part of the protein, the stalk, links the head to the cargo binding site, contains coil-coiled structures for oligomerization or associates with other subunits. Head and stalk are parts of the same polypeptide, which is functionally relevant as a tight link of head and stalk is essential for transmission of mechanical force [2].

Genetic analysis of the physiological function of motor proteins is hampered, since many motor proteins fulfill an essential function for the cell or organism. For example, Kinesin-5 serves indispensable functions during mitosis, making an analysis of its function in interphase or in terminally differentiated cells difficult. Conditional mutations, such as temperature sensitive alleles, can overcome these limits of genetic analysis [3]. Gene knock down by RNAi
approaches relays on protein turnover, leading to insensitivity of stable
proteins. Pharmacological approaches with small molecules inhibitors or
specific antibodies provide an alternative and have been applied for motor
protein inhibition [4–6]. However, chemical approaches cannot be generalized,
and need to be developed case by case.

Kinesin-5 belongs to kinesin family member 11 (KIF11), with the motor domain
on N terminus, followed by a coiled-coil rod containing a central bipolar
assembly (BASS) domain. Forming bipolar homo tetramers, Kinesin-5 can
crosslink anti-parallel aligned microtubules. The motor activity enables filament
sliding, e.g. during formation and elongation of the mitotic spindle [7]. In
Drosophila syncytial embryos, Kinesin-5 is enriched at mitotic spindles and is
essential for spindle formation and chromosome segregation. Injection of
antibodies specific for Kinesin-5 into embryos leads to collapse of newly
formed spindle and the formation of mono-asters of microtubules [5,6].

Making proteins susceptible to proteolytic cleavage represents a generally
applicable strategy for generation of conditional alleles[8-10].

Here we apply this concept to motor proteins by inserting a proteolytic site
between the head and stalk region (“neck”). We designated this strategy a
“molecular guillotine” (Fig. 1A). We chose well-characterized Kinesin-5 in order
to demonstrate the feasibility of this approach. As a protease, we employ TEV,
which is highly specific. No match of TEV recognition motif within the
Drosophila proteome has been identified, and flies expressing TEV are viable
and fertile [10].

Results

Design of a “molecular guillotine”
We inserted three copies of the TEV recognition motif at one of two positions, G394 or Q499, into the stalk region. G394 and Q499 are located within conserved coiled-coil regions next to the head domain (Fig. 1B, C). In addition, we fused GFP to the C-terminus, which does not affect the function of Kinesin-5, as previously reported [11]. These constructs were expressed as transgenes in levels comparable to the endogenous allele with a ubiquitously active promoter, as assayed by western blot (Fig. 1D). Due to the C-terminal GFP moiety, the constructs showed a slower mobility in SDS-PAGE than wild type Kinesin-5. The TEV sites do not affect the functionality of Kinesin-5 as the construct with the insertion at G394 (Kin-5[G394tev]-GFP) complemented the lethality of a Kinesin-5 (Klp61f) mutation. For this, we recombined Kin-5[G394tev]-GFP with a Kinesin-5 mutation. The resulting flies only expressed Kin-5[G394tev]-GFP, were viable and fertile and can be kept as a homozygous stock. In embryos from this line, Kinesin-5 was detected only at the molecular weight corresponding to transgenic Kin-5[G394tev]-GFP, which confirms the absence of endogenous Kinesin-5 (Fig. 1E).

Kinesin-5 cleavage in vivo

We expressed TEV protease in stripes in embryos under the control of the engrailed promoter. Control embryos with no TEV expression showed uniform Kin-5[G394tev]-GFP expression. In contrast, the GFP signal was strongly depleted in stripes with TEV expression (Fig. 2A). Next we turned to syncytial embryos, which are characterized by their rapid and synchronous nuclear division cycles and the associated remodeling of the cytoskeleton. During mitosis, microtubules and their motors are important for formation and function of mitotic spindles and chromosome segregation, whereas they function in nuclear arrangement and stabilization of the nuclear array in interphase [12,13]. Kinesin-5 localizes to the mitotic spindle and is involved in
chromosome segregation during mitosis [5,6,11]. We microinjected TEV protease into syncytial embryos and recorded GFP fluorescence. Following injection of TEV protease but not buffer, GFP fluorescence rapidly dropped (Fig. 2B). Correspondingly, the specific staining pattern, such as labelling of mitotic spindles or cytoplasmic asters was lost in TEV injected embryos (Fig. 2D). Quantification of total GFP fluorescence provided an estimate for an approximate half life of about 30 min (Fig. 2C). Kinesin-5 was specifically cleaved, since the electrophoretic mobility of Kin-5[G394tev]-GFP was higher in TEV than buffer injected embryos (Fig. 2E). Kin-5[G394tev]-GFP embryos were lysed about 30 min after injection and extracts analyzed by western blot against the C-terminus of Kinesin-5. The observed difference in electrophoretic mobility was consistent with proteolytic cleavage at the TEV sites at the neck and corresponding loss of the head domain. As we detected a single band, proteolytic cleavage was close to complete under our experimental conditions (Fig. 2E).

Cleavage of Kinesin-5 leads the loss-of-function in mitosis

Next we analyzed the functional consequences of the Kinesin-5 cleavage. To track the nuclear cycles and behavior of chromosomes, we co-injected fluorescent labelled histone-1 and TEV protease into Kinesin-5 null embryos expressing the Kin-5[G394tev]-GFP transgene. Following TEV injection, we observed a failure of chromosome separation and monoastral spindles (Fig. 3). These phenotypes were observed in individual spindles interspersed between normally appearing spindles. These phenotypes were consistent with the previously reported mitotic defects following Kinesin-5 antibody injection [5].

Interphase function of Kinesin-5

An interphase function of Kinesin-5 has not been investigated, yet. In interphases of syncytial embryos, Kinesin-5-GFP is strongly enriched at the
centrosomes and associated asters. In addition, dynamic extended structures
between adjacent asters were detected (Fig. 4C). These transient signals may
represent microtubules coated with Kinesin-5 and possibly antiparallel aligned
microtubules.

As hypothesized previously [12,13], Kinesin-5 may be involved in nuclear
positioning and formation of the nuclear array in syncytial Drosophila embryos.
Kinesin-5 bound to anti-parallel aligned microtubules may push adjacent
asters away from each other and thus generate a repulsive force, which may
lead to uniform internuclear distances. In this model, Kinesin-5 would promote
movements of centrosome and their associated asters. Alternatively, Kinesin-5
may crosslink microtubules from adjacent asters and stabilize the syncytial
microtubule network. In this model Kinesin-5 would suppress movement of
centrosomes and associated asters (Fig. 4B). To distinguish these two models,
we recorded the dynamics of centrosomes in the scale of seconds [13]. From
the recorded tracks, fluctuations of the centrosomes were calculated as
previously reported [13]. These fluctuations have the dimension of a diffusion
constant and do not contain slow (minute-scale) drift movement. We recorded
centrosome dynamics in embryos with injected TEV protease and calculated
the second-scale fluctuations (Fig. 4D). We find that the fluctuations are
strongly increased to about 20x10⁻³ µm²/s as compared to about 6x10⁻³ µm²/s
in control embryos injected with buffer. Passive fluctuations as detected in
embryos depleted of ATP are in the range of 1.2x10⁻³ µm²/s [13] (Fig. 4E).
Since cleavage of Kinesin-5 leads to an increased centrosome mobility, we
conclude that functional Kinesin-5 stabilizes the dynamics of the microtubule
array.

Discussion
The function of Kinesin-5 for spindle formation and elongation during mitosis is well established [6]. Consistently, inhibition of Kinesin-5 by antibody injection induces defects in chromosome segregation in syncytial Drosophila embryos. Although expressed, a function of Kinesin-5 during interphase has been unknown, partly because such an interphase function was obscured by the mitotic defects in Kinesin-5 depleted embryos. The problem that one phenotype obscures other phenotypes is common to proteins with widespread functions, such as molecular motors. To circumvent this problem, we developed a method for conditionally inactivating Kinesin-5. In addition to Kinesin-5, this method is potentially suitable for other motor proteins, as well. With a “molecular guillotine”, we specifically inactivated Kinesin-5 by administration of TEV protease. In this way, we revealed an interphase function for the stabilization of the syncytial microtubule network. In syncytial embryos, the microtubule asters originating from centrosomes can directly interact with neighboring asters, since they are not physically separated by plasma membranes. These interactions lead to formation of an extended network covering the embryonic cortex. The phenotypic behavior of centrosomes and their associated nuclei reflect their intrinsic properties but also, as part of the network, the influences from the neighbors. Adjacent microtubule asters potentially interact via crosslinkers such as Feo/Ase1p, bundling proteins or motors with sliding activity, such as Kinesin-5. Here we tested the hypothesis that Kinesin-5 generates repulsive forces between adjacent astral microtubules in interphase. We expected that a loss of force generation would have led to a reduced mobility of the network and its nodes, the centrosomes. Using the fluctuations of centrosomes as an indicator of network dynamics, we rejected our hypothesis, because we measured an increased mobility of the centrosomes, when Kinesin-5 was inactivated. We interpret this data in that the in vivo function of Kinesin-5 as a crosslinker is
more dominant than its function for sliding of anti-parallel aligned microtubules
and thus pushing apart adjacent microtubule asters. The in vivo function of
Kinesin-5 is similar to Kinesin-1, which is enriched at the cortex and F-actin
and actin caps. Both may be involved in anchoring microtubule asters to the
cortex and in this way counteract fluctuation movements of centrosomes.

Having identified a suppressive function of Kinesin-5, the questions remains
about the origin of the forces driving centrosome fluctuations. Fluctuations are
due to an active component, since ATP depletion leads to loss of fluctuations.
The (−)-end directed motor Kinesin-14 may serve as a force generator [6].

“Molecular guillotine” can be used as a conditional mutant tool to study
the function of motor protein

The “molecular guillotine” is potentially a versatile method for conditional
inactivation of motor proteins. TEV protease has been used in inactivation of
cohesin in yeast [14] and in fly [9], as well as Drosophila claudin [10]. However,
this approach had not been used in motor proteins. The approach of a
“molecular guillotine” as reported in this study can be applied widely to
members of the motor protein families. Unlike using the small molecules
inhibitor [4,15], TEV protease can be specifically expressed using UAS-GAL4
system in any genetically tractable cell type, and thus decapitate the selected
motor protein in a tissue and developmental stage specific manner. Direct
cleavage by TEV potentially leads to a faster inactivation kinetics than by the
degron [16] or deGradFP systems [17], which rely on the ubiquitin-mediated
protein degradation machinery. In addition, inactivation of motor protein by the
“molecular guillotine” approach can be fine-tuned by titrating the TEV protease
concentration, which help to identify additional functions of the motor proteins.
In summary, the novel approach of a “molecular guillotine” enabled us to investigate a specific function of the motor protein Kinesin-5 in interphase. Potentially, the decapitation approach can be correspondingly applied to other kinesin motors as well as dyneins and myosins, as they have a related domain structure in common.

**Materials & Methods**

**Genetics**

Fly stocks (en-Gal4, Sas6-GFP, Klp61f^{07012}) [18,19] were obtained from the Bloomington Stock Center, if not otherwise noted. Transgenes of ubi-Kin5-tev-GFPQ499, ubi-Kin5-tev-GFPG394 and sqh-Kin5-GFP were generated by P element mediated random genome integration. We isolated multiple insertions on the third chromosome with varying expression. The ubi-Kin5-tev-GFPG394 line with strongest GFP fluorescence was recombined with an amorphic Kinesin-5 mutation (Klp61f^{07012}) and kept as a homozygous line. The transgenes spq-Kin5-GFP without TEV sites complemented the lethality of the Kinesin-5 mutation (Klp61f^{07012}) as well. TEV protease was expressed from a UAS-T-TEV transgene or injected as a purified recombinant protein.

**Cloning**

A sequence coding for three recognitions sites of TEV protease (PS ENLYFQG PR ENLYFQG GS ENLYFQG PR) was inserted behind the codons of G394 or Q499 of the Kinesin-5 cDNA. This sequence was cloned together with the ubiquitin promoter and eGFP into the multiple cloning site of a pUASt vector derivative lacking the UAS and hsp70TATA sites. Kinesin5-GFP was cloned by fusing eGFP coding sequence to the 3’ end of Kinesin-5 coding
sequence and transferred to transformation vector sGMCA [20]. Sequence information and details of the cloning procedure are available upon request.

**Western blotting, Kinesin antibody**

The *Kinesin5* coding sequence corresponding to the C-terminal tail (aa 600–1066) was cloned by PCR with SK-Klp61f (Drosophila genomic resource center, Bloomington) as template into a protein expression vector with a N-terminal 9xHis tag. The His9-Kinesin-5-C600 protein with an apparent molecular weight of about 70 kd in SDS polyacrylamide gel electrophoresis (SDS-PAGE) was purified under denaturing conditions (Trenzyme, Konstanz) and used for immunization of rabbits (BioGenes, Berlin). Embryonic extracts were analyzed by SDS-PAGE and immunoblotting as previously described [21]. Briefly, proteins were blotted to nitrocellulose filters by wet transfer (100 mA per mini gel, overnight). The blots were blocked with 5% fat-free milk in PBS, incubated with primary antibodies in PBT (PBS with 0.1% Tween20, Kinesin-5, rabbit, 1:5000, α-Tubulin, mouse, 1:100000, B512, Sigma, GFP, rabbit, 1:5000, Torrey Pines Biolabs) and fluorescently labelled secondary antibodies (LiCOR, 1:20000, 0.05 µg/ml in PBT) for each two hours at room temperature. The developed blots were recorded with a LICOR system.

**Microinjection**

Embryos were dechorionated and aligned on a coverslip, desiccated for 10 min, and covered with halocarbon oil (Voltalef 10S, Lehmann & Voss). We injected TEV protease at 10 µM purified from overexpressing E. coli (a gift from Dirk Görlich) or Histone1-Alexa-488 at 2 mg/ml (ThermoFisher).

**Microscopy**
Images were recorded with a Zeiss microscope equipped with a spinning disc (25x/NA0.7 multi immersion, 40x/NA1.3 oil). Centrosome movement was recorded in Sas6-GFP expressing embryos as previously described with a frame rate of 1 Hz [13]. Kin-5-GFP distribution in interphase was recorded with a confocal microscope (Zeiss LSM780 with airy scan unit, 63x/NA1.4/oil). Images were processed with Fiji/ImageJ.

**Fluctuation analysis**

The centrosomes tracking and measurement of fluctuation were carried out as previously described [13].

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**References**


**Figure legends**

**Figure 1. Design of a molecular guillotine for Kinesin-5.** (A) Schematic illustration of motor protein molecular guillotine by inserting a protease substrate site next to the head domain of a motor. (B) TEV cleavage site in three copies (3x) is inserted in the coiled-coil region in stalk domain at position G394 or Q499. Domain structure of Kinesin-5 (Drosophila, Klp61F) and secondary structure prediction are indicated (C) Sequence alignment of the insertions sites at G394 and Q499 in wild type and Kinesin-5 mutant background. (D, E) Western blots with embryonic extracts (0–4h) from wild type and flies with the Kinesin-5-tev-GFP transgene, probed as indicated with antibodies for Kinesin-5, GFP, α-tubulin. Apparent molecular weight in kilo Dalton.

**Figure 2. Kin-5[G394tev]-GFP is cleaved by TEV protease.** (A) Image of living embryos expressing Kin-5[G394tev]-GFP with or without TEV protease expressed in striped pattern. Scale bar 50 µm. Region marked by squares in yellow are shown in high magnification. Scale bar 10 µm. Quantification of GFP signal along the anterior-posterior body axis (line in green). (B–D) TEV protease or buffer was injected into syncytial embryos mutant for Kinesin-5 and expressing Kin-5[G394tev]-GFP. (B) Images from time lapse recording. Time in minute:second. (C) Quantification of GFP fluorescence. N, number or embryos. (D) Images of living embryos before and 30 min after injection. Scale bar 10 µm. (E) Western blot with extracts from embryos 30 min after injection with TEV or buffer probed with Kinesin-5 and α-Tubulin antibodies. Apparent molecular weight in kilo Dalton.

**Figure 3. Phenotype of Kin-5[G394tev]-GFP cleavage by TEV protease.**
(A, B) Images from time lapse recording of embryos mutant for Kinesin-5 expressing the Kin-5[G394tev]-GFP transgene and injected with fluorescent labeled Histone H1. (B) Coinjection of TEV protease. Arrow head in yellow points to defective mitotic figure. Schematic drawing of the mitotic figures. Scale bar: 10 µm.

Figure 4. Interphase function of Kinesin-5

(A) Projected image of an embryo expressing Histone 2Av from selective plane illumination microscopy in side view and cross section (position indicated by lines in blue). Magnified section illustrate the interactions between the nuclei and between nuclei and cortex. Dots in yellow indicate centrosome pairs. (B) Illustration of microtubule asters with overlapping microtubules in anti-parallel orientation. Kinesin-5 may slide microtubules apart (Model 1) or crosslink adjacent asters (Model 2). (C) Image of living embryo expressing Kinesin-5-GFP (apical position) Scale bar 5 µm. (D) Images from living embryo mutant for Kinesin-5 expressing Kin-5[G394tev]-GFP and SAS6-GFP and injection with TEV protease or buffer. Trajectories of centrosomes over 220 s on an image from time lapse recoding. Scale bar 5 µm. (E) Box plot displaying time-averaged fluctuation of centrosomes in embryos expressing SAS-6-GFP injected with buffer (wild type), sodium azide or TEV protease for cleavage of Kin-5[G394tev]-GFP.
Figure 1

**A**

3x TEV cleavage sequence

**B**

Motor

Stalk

**C**

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**D**

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<td>55</td>
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**E**

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<td>α-tub</td>
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Figure 2

A

-TEV

enGal x TEV

+TEV -TEV

Fluorescence intensity

A-P axis

B

ΔKin-5, Kin-5[G394tev]-GFP

+Tev +Water

00:00

00:34

00:40

01:24

01:20

C

ΔKin-5, Kin-5[G394tev]-GFP

+Tev +Water

00:00

00:34

00:40

01:24

01:20

D

before injection after injection

30 min

E

MW [kD] TEV

- - +

Kin-5

α-tub

N=9

250

130

55

Figure 2

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A-P axis

before injection after injection

30 min

enGal x TEV

+TEV -TEV

Fluorescence intensity

A-P axis

ΔKin-5, Kin-5[G394tev]-GFP

+Tev +Water

00:00

00:34

00:40

01:24

01:20

D

before injection after injection

30 min

E

MW [kD] TEV

- - +

Kin-5

α-tub

N=9

250

130

55

Figure 2

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Figure 3

A

Histone

0 min 2 min 3 min 4 min 10 min

B

TEV+Histone

0 min 3 min 7 min 8 min 10 min

0 min 2 min 3 min 4 min 10 min

TEV+Histone
Figure 4

A

B

C

D

E

Kin5-GFP

SAS6-GFP

Fluctuation [µm /s]²

0.025

0.020

0.015

0.010

0.005

wild type

azide

∆Kin-5,

Kin-5-tev-GFP

+TEV

Kin5 cleavage