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Running title: Binding of katanin to microtubules

Arabidopsis katanin binds microtubules using a multimeric microtubule-binding domain

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## **Abstract**

Katanin is a heterodimeric protein that mediates ATP-dependent destabilization of microtubules in animal cells. In plants, the catalytic subunit of *Arabidopsis thaliana* katanin (AtKSS, *Arabidopsis thaliana* Katanin Small Subunit) has been identified and its microtubule-severing activity has been demonstrated in vitro. In vivo, plant katanin plays a role in the organization of cortical microtubules, but the way it achieves this function is unknown. To go further in our understanding of the mechanisms by which katanin severs microtubules, we analysed the functional domains of *Arabidopsis* katanin. We characterized the microtubule-binding domain of katanin both in vitro and in vivo. It corresponds to a poorly conserved sequence between plant and animal katanins that is located in the N-terminus of the protein. This domain interacts with cortical microtubules in vivo and has a low affinity for microtubules in vitro. We also observed that katanin microtubule-binding domain oligomerizes into trimers. These results show that, besides being involved in the interaction of katanin with microtubules, the microtubule-binding domain may also participate in the oligomerization of katanin. At the structural level, we observed that AtKSS forms ring-shaped oligomers.

Keywords: katanin; microtubule-binding domain; oligomerization; severing of microtubules

Abbreviations: AtKSS, *Arabidopsis thaliana* katanin small subunit; GST, Glutathione S-Transferase

## 1. Introduction

Plant microtubules are located in the cell cortex (cortical microtubules during interphase, preprophase band at the entry of mitosis) and in the cytoplasm (nucleus-associated microtubules, spindle and phragmoplast microtubules in mitosis), where they are arranged in highly dynamic and ordered arrays. Upon internal and external stimuli, these arrays reorganize, while acquiring specific functions including the equal segregation of chromosomes, the formation of the cell plate at the end of mitosis, the transport of cargo and the positioning of organelles [1, 2]. The architectural changes of these microtubule arrays are dramatic manifestations of the flexibility and dynamic turnover of the microtubules in plant cells. The precise mechanisms underlying the remodeling of microtubule arrays remain difficult to elucidate, mainly because of the complexity of the plant microtubular cytoskeleton and because of the absence in plant cells of a discrete microtubule-organizing center. When compared with animal microtubules, plant microtubules possess very peculiar properties *in vivo* [3-6], including the fact that cortical microtubules exhibit polymerization-biased dynamic instability at one end and slow depolymerization at the other end [5].

*In vivo*, the control of microtubule assembly and disassembly requires a multitude of adaptors and regulators. These proteins associate with the polymer lattice, with the microtubule ends or with the tubulin dimer. They regulate microtubule stabilization or destabilization in a way to generate the different microtubule arrays observed *in vivo* [7, 8]. Until recently the identification of plant microtubule regulators has been rather elusive because of their low abundance and because most of them show low sequence homology with animal counterparts. Several plant microtubule-stabilizing effectors, also called Microtubule Associated Proteins, have now been identified and their functional characterization is under way [9, 10]. So far, AtKSS (*Arabidopsis thaliana* Katanin Small Subunit) is the only plant microtubule-destabilizing protein that has been functionally characterized.

Katanin is a heterodimer composed of a catalytic subunit of 60 kDa (p60) that hydrolyses ATP to sever microtubules, and a regulatory subunit of 80 kDa (p80) involved in subcellular targeting of katanin [11, 12]. p60 oligomerizes into a ring-shaped hexamer in an ATP-dependent process which is coupled to the disassembly of microtubules [13]. In animal cells, katanin is located at the centrosomes and at spindle poles where it controls the length of the mitotic spindle [14, 15]. In plants, the

Arabidopsis homologue of katanin catalytic subunit, named AtKSS, was identified on the basis of its sequence similarity with human (*homo sapiens*) and sea urchin katanins [16]. In vitro, AtKSS has a microtubule-stimulated ATPase activity and severs microtubules in an ATP-dependent [17]. In vivo, properties and functions of plant katanin are not well understood. Over the last few years, five AtKSS mutants have been described. *botero1* (*bot1*) is a null mutant [18], while the four other mutants, *fragile fiber* (*fra2*; [19, 20]), *ectopic root hair 3* (*erh3*, [21]), *LUC-super-expressor 1* (*lue1*, [22]), *dwarf and gladius leaf1* (*dgl1*, [23]) are mutated in the C-terminal ATPase domain of AtKSS. All these mutants show a common dwarf phenotype associated with an alteration of cellular anisotropic growth. But a closer look to the plants revealed differences between the mutants. For example, both *fra2* and *lue1* show altered composition and organization of cellulose microfibrils, as well as two-branched trichomes, whereas none of these two defects was observed in *bot1*. Similarly, some but not all cells of *bot1*, *dgl1*, *fra2* and *lue1* have disorganized cortical microtubules. Because of the complexity and the diversity of the phenotypes of AtKSS mutants, it has been difficult to get a clear view of the functions of katanin in vivo. By analyzing the induced expression of AtKSS in Arabidopsis, we recently showed that katanin favors bundling of microtubules in the cell cortex [24]. Obviously, AtKSS is required for a proper organization of cortical microtubules in expanding plant cells, but whether the disorganization of cortical microtubule arrays is a direct or indirect consequence of AtKSS's defect is debated.

To further understand the biological roles of AtKSS, we studied the properties of full-length and truncated AtKSS both in vitro and in vivo. We report here that AtKSS interacts with microtubules through a poorly conserved basic domain, both in vitro and in vivo. We observed that AtKSS organized into ring-shape structures, and that the microtubule-binding domain of AtKSS formed trimers. These results suggest that the microtubule-binding domain of AtKSS participate in the oligomerization of plant katanin.

## 2. Results

### 2.1. Identification of the AtKSS microtubule-binding domain

We previously showed that AtKSS binds microtubules in vitro [17]. In a first step to understand the molecular mechanisms by which the plant katanin interacts with microtubules, we analysed the functions of the domains of AtKSS. On the basis of the amino acid sequence alignment of AtKSS (accession number **AF048706**) with other known p60s (*Caenorhabditis elegans*, accession number **S47861**; *Chlamydomonas reinhardtii*, accession number **AAF12877**; *Drosophila melanogaster*, accession number **AAF34687**; *homo sapiens*, accession number **AAC25114**; sea urchin, accession number **AAC15706**; *Xenopus laevis*, accession number **AAD53310**), we distinguished three domains (*figure 1A*). The first domain AtKSS1 corresponds to the 34 first amino acids. It composes a short domain that is highly conserved with vertebrate's p60s (AtKSS1 has 85% sequence similarity with human and *Xenopus* sequences). The domain AtKSS2 (amino acids 35 to 210) delimits an intermediate domain with no significant sequence similarity with known p60s. AtKSS3 (amino acids 211 to 523) delimits a large C-terminus ATPase domain, which is well conserved among all p60s (80% sequence similarity with *Caenorhabditis*, *Drosophila*, human, and *Xenopus* sequences). The proteins corresponding to these domains were produced in fusion with Glutathione S-Transferase GST (GST-AtKSS1, GST-AtKSS2, GST-AtKSS3), as well as AtKSS deleted of one domain (GST-AtKSS12, GST-AtKSS13, GST-AtKSS23; *figure 1B* and *1C*). It has to be noted that whatever expression and purification conditions we used, samples of GST-AtKSS23 contained a C-terminus truncated GST-AtKSS23 (*figure 1C*, lane 6, lower band). As these two proteins can form hetero-oligomers (see results on katanin oligomerization), we could not separate them in native conditions. When purified in denaturing conditions and further refolded, the full-length GST-AtKSS23 loses some of its properties. For these reasons, GST-AtKSS23 we used in this study corresponds to a mixture of the two proteins.

In order to identify the microtubule-binding domain of AtKSS, GST-AtKSS constructs were added to taxol-stabilized microtubules. After centrifugation, analysis of the supernatants and pellets showed that both GST-AtKSS12 and GST-AtKSS2 co-sedimented with microtubules, whereas neither the short N-terminus domain

(GST-AtKSS1) nor the ATPase domain (GST-AtKSS3, GST-AtKSS13) interacted with microtubules (*figure 2*). These results indicate that amino acids 35 to 210 of AtKSS are involved in the interaction of AtKSS with microtubules *in vitro*.

### *2.2. Affinity of the microtubule-binding domain of AtKSS for microtubules in vitro and in vitro*

We performed co-sedimentation assays with microtubules to evaluate the dissociation equilibrium constant of the GST-AtKSS2 and GST-AtKSS12 (*figure 3*). We calculated an apparent  $K_d$  of 5.10  $\mu\text{M}$  for GST-AtKSS2 and 6.87  $\mu\text{M}$  for GST-AtKSS12. The apparent  $K_d$  of GST-AtKSS12 and unfused AtKSS12 had similar values (6.87  $\mu\text{M}$  for GST-AtKSS12 versus 18.03  $\mu\text{M}$  for AtKSS12), showing that in our experimental conditions GST did not significantly interfere with the binding of the proteins to microtubules. These results show that AtKSS binds microtubules with a low affinity, and that the N-terminus part of AtKSS (domain 1) is not involved in the binding of the protein to microtubules *in vitro*.

We wondered whether the microtubule-binding domain of AtKSS interacts with microtubules *in vivo*, where microtubules are covered with microtubule-associated proteins. We transiently expressed AtKSS12 or AtKSS2 fused to DsRed in onion epidermal cells. Sixteen hours after transformation, the AtKSS12-DsRed (*figure 4A*) and AtKSS2-DsRed were detected along filaments. We did not detect any difference between the localization of AtKSS12 and AtKSS2 *in vivo*. Transformed cells were treated with 20  $\mu\text{M}$  oryzalin, a potent plant microtubule destabilizer. This concentration of oryzalin is sufficient to depolymerize all microtubules detectable at the light microscope level *in vivo* [25]. After 20 min of incubation, almost all the stained filaments had disappeared (*figure 4B-D*). These results show that the domain of AtKSS delimited by the amino acids 35 to 210 binds microtubules in living cells.

### *2.3. Oligomerization of AtKSS*

In order to determine whether AtKSS is able to form oligomers, we examined the structure of AtKSS molecules at the electron microscopy level. AtKSS appeared as irregular rings of 25-30 nm diameter composed of globular subunits (*figures 5A-C*). Rings of AtKSS often contained a crack radiating outward. These rings are very similar to hexamers of animal p60 [12].

To identify domains of AtKSS that may be involved in the oligomerization, we studied the oligomeric state of the different AtKSS constructs. First, the molecular mass of the different katanin constructs was estimated by gel filtration. The N-terminus part of katanin (GST-AtKSS1), the ATPase domain of katanin (GST-AtKSS3), and the chimeric protein composed of these two domains (GST-AtKSS13) were eluted from the column as dimers (*table 1*). Because GST exists as a dimer in its native state [26], and because in our experimental conditions GST was eluted as a dimer (*table 1*), we concluded that dimers of GST-AtKSS1, GST-AtKSS3 and GST-AtKSS13 resulted from the dimerization of GST. In contrast the microtubule-binding domain of katanin, either alone (GST-AtKSS2) or combined to the N-terminus domain (GST-AtKSS12) was eluted as trimers. We also performed cross-linking experiments with the different GST constructs using the zero-length cross-linker EDC together with NHS. These experiments could not be done with GST-AtKSS2, as the protein is not stable at room temperature. Incubation of GST-AtKSS12 with the cross-linker yielded to a band of an apparent molecular mass of 160 kDa (*figure 5D*). This result suggests that AtKSS12 forms trimers. None of the chimeric proteins lacking the microtubule-binding domain (GST alone, GST-AtKSS1, GST-AtKSS3 (data not shown) or GST-AtKSS13 (*figure 5E*)) formed oligomers. Trimers of AtKSS12 without GST were also observed after cross-linking (*figure 5F*). All together these data suggest that AtKSS oligomerizes into a ring-shaped oligomers, and that the microtubule-binding domain of AtKSS is involved in the oligomerization process.

#### *2.4. ATPase and microtubule-severing activity of katanin.*

We studied the ATPase activity and severing activity of the different domains of katanin. From all the chimeric AtKSS we studied, only those containing domain 3 (GST-AtKSS13, GST-AtKSS23 and GST-AtKSS3) hydrolyzed ATP (*figure 6A*). GST-AtKSS2 and GST-AtKSS12 had no detectable ATPase activity (data not shown). GST-AtKSS23 had the highest ATPase activity, and its activity was stimulated by microtubules, as observed for baculovirus-expressed full-length AtKSS [17]. Hydrolysis of ATP by GST-AtKSS13 and GST-AtKSS3 was not stimulated by microtubules. These data are in agreement with our results showing that the ATPase domain of katanin is not able to interact with microtubules (*figure 2*).



We also analysed the microtubule-severing activities of different katanin constructs using turbidity experiments and video-microscopy assays. We observed that the chimeric katanin containing the microtubule-binding domain and the ATPase domain (GST-AtKSS23) destabilized microtubules in vitro (*figure 6B and 6E-F*). In contrast, neither GST-AtKSS1, nor the microtubule binding domain alone (GST-AtKSS2) or fused to the domain 1 (GST-AtKSS12), nor the ATPase domain alone (GST-AtKSS3) severed microtubules in vitro (data not shown). Our data did not allow us to quantitatively compare the severing activities of the GST-AtKSS23 and the full-length katanin and thus to determine if the presence of the short NH<sub>2</sub>-terminal domain (AtKSS1) modulates katanin's activities in vitro. Our results show that the microtubule-binding domain and the ATPase domain are sufficient for katanin to sever microtubules in vitro.

### 3. Discussion

Up to now, data obtained from the analysis of four AtKSS mutants [18-20, 22, 23] and from the induced expression of AtKSS in arabidopsis plants [24] clearly indicate that AtKSS is required for a proper organization of cortical microtubules in expanding plant cells. But how katanin achieves this function is still unknown. In this study, we examined some of the molecular properties of katanin, including its interaction with microtubules and its ability to form oligomers.

Our data show that amino acids 35 to 210 of AtKSS are sufficient for AtKSS to bind microtubules both *in vitro* and *in vivo*. Noticeably plant and animal catalytic subunits of katanin bind microtubules via the same domain (domain 2, see *figure 1A*) and with a comparable  $K_d$  (this study, [13, 27]). This may be due to the fact that although the amino-acid sequences of plant and animal microtubule-binding domains of katanin are poorly conserved, they are enriched in basic amino acids that can interact with the acidic tubulin surface.

We found that besides mediating the interaction of katanin with microtubules, the microtubule-binding domain of AtKSS allows intermolecular interactions between katanin subunits. Indeed, the microtubule-binding domain of AtKSS oligomerizes into trimers and AtKSS forms rings similar in size and in shape to sea urchin katanin [12]. These results suggest that the microtubule-binding domain is involved in the oligomerisation of katanin. It has been shown that the microtubule-severing activity of animal katanin is correlated with its ability to oligomerize into a ring-shaped hexamer [13]. Assuming that the active AtKSS is also a hexamer, we propose that the oligomerization of the catalytic subunit of katanins takes place in at least two steps: the formation of a trimer and then the addition of two trimers to form a hexamer. More detailed structural data are now required to understand how katanin subunits interact to form a hexamer, and to evaluate the role of the catalytic state of the ATPase domain in this process.

Our data did not allow us to specify a function to the N-terminus of AtKSS (AtKSS1). This domain does not bind microtubules by itself, neither does it interfere with the interaction of the microtubule-binding domain with microtubules as  $K_d$  values of GST-AtKSS2 and GST-AtKSS12 were similar. It is probably not essential for the oligomerization of AtKSS as GST-AtKSS23 fragments microtubules. It is possible that this domain, although not necessary for severing activity of AtKSS, favors one or

several steps of the microtubule-binding/oligomerization/ATP hydrolysis cycle of AtKSS, and that the procedures we used were not sensitive enough to detect such a fine regulation. The N-terminus part of AtKSS may also be involved in the recruitment of AtKSS-regulatory proteins, as it has been shown for animal katanin [27]. AtKSS-interacting proteins have recently been identified by two-hybrid screen [22]. Characterization of the ability of these proteins to regulate katanin *in vitro* and *in vivo* will be of great help in the understanding of the roles of microtubule severing in higher plants.

Non-ectopic fragmentation of microtubules in plant cells implies that the activity of katanin is regulated *in vivo*. One way to control katanin's activity is to modulate its binding to microtubules. Several lines of evidence suggest that katanin transiently interacts with microtubules. First, anti-katanin antibodies did not detect AtKSS along microtubules [16]. Second, in our hands, co-localization of katanin with microtubules *in vivo* was only observed in transiently transgenic cells in which AtKSS was overexpressed after biolistic transformation (our unpublished data). Neither in transgenic *Arabidopsis* cells expressing AtKSS in an inducible manner [24], nor in tobacco cells expressing AtKSS after agro-infiltration (our unpublished results), could we detect an interaction between katanin and microtubules. Indeed, there is only one study reporting the co-localization of AtKSS with microtubules *in vivo* [22]. In this work, stably overexpressed AtKSS-GFP-GUS decorates microtubules in *Arabidopsis* plants. However the functionality of the tripartite chimeric protein used in the study has been questioned by the authors. Third, the  $K_d$  value we calculated for the microtubule-binding domain of AtKSS is at least ten times higher than the  $K_d$  we estimated for plant MAP65-1b [28]. All these observations are consistent with a transient interaction of plant katanin with microtubules. The absence of co-localization of a microtubule-interacting protein with microtubules *in vivo* is not unique to katanin. Recent studies showed that spastin, an AAA katanin-related protein that destabilizes microtubules does not co-localize with microtubules in animal cells, whereas mutated spastin with altered ATPase activity does [29, 30]. These results show that ATP hydrolysis is involved in the interaction of spastin with microtubules. Whether a comparable mechanism also affects the interaction of plant katanin with microtubules remains to be shown.

The present study provides new insights into the molecular properties of plant katanin. However many questions remain to be answered before we can fully

understand the molecular mechanisms underlying microtubule severing by katanin in plants. Does katanin preferentially associate with a subset of microtubules in vivo? If so, how is this association regulated? Does it depend on the dynamic properties of microtubules, and/or on the presence of microtubule-associated proteins such as MAP65 proteins? What is the role of the homologue of the katanin-accessory protein p80? The ongoing characterization of plant microtubule-interacting proteins should soon provide insights into these challenging issues.

## 4. Methods

### 4.1. Chemicals and antibodies

Unless specified all chemicals were purchased from Sigma (St Louis, MO, USA). GST antibodies were purchased from Amersham Pharmacia Biotech (Little Chalfont, England). Anti-AtKSS antibodies raised against the N-terminus peptide of the protein were previously described [17].

### 4.2. AtKSS cloning and expression

Three domains of AtKSS were defined: the 34 first amino-acids (domain 1), amino acids 35 to 210 (domain 2) and the ATPase domain (aa 211 to 523; domain 3). These domains are further referred to as AtKSS1, AtKSS2 and AtKSS3 respectively. AtKSS1, AtKSS2, AtKSS3, as well as combinations of the domains, AtKSS12 (aa 1 to 210), and AtKSS23 (aa 211 to 523), were cloned into the pGEX-4T-1 plasmid (Amersham Pharmacia Biotech) in frame with the Glutathione S-Transferase GST (Amersham Pharmacia Biotech) using the following primers: for AtKSS1, primers 5HAtKSS60 5' CGCGGATCCGTGGGAAGTAGTAAT 3' and R34 5' CCCGAATTCTTAGAAGAAGATGACAGATGTG 3'; for AtKSS2, primers F34 5' CGCGGATCCTTCGATGGCGCCATTGCTCAG 3' and R210 5' CCCGAATTCTTATGACTTCCCATCTTCAGC 3'; for AtKSS3, primers F210 5' CGCGGATCCAAAAGGGGATTGTATGAGGGAC 3' and 3HAtKSS 5' CCGGAATTCCGTTAAGCAGATCCAAACTC 3'; for AtKSS12, primers 5HAtKSS60 and R210; for AtKSS23, primers F34 and 3HAtKSS. The chimeric AtKSS truncated of the domain 2, AtKSS13 (aa 1 to 34 and 211 to 523) was obtained by PCR using the QuickChange<sup>TM</sup> site-directed mutagenesis kit (Stratagene, CA, USA), the full length AtKSS in pGEX-4T-1 as substrate and the following primers: F-N-AAA 5' CCTATGACACATCTGTCATCTTCTTCAAAGGGGATTGTATGAGGGACC 3' and R-N-AAA 5' GGTCCCTCATAACAATCCCCTTTTGAAGAAGATGACAGATGTGTCATAGG 3'. All sequences were verified by sequencing. All GST-AtKSS proteins were expressed overnight at 15°C in Rosetta (DE3)pLysS bacteria strain (Novagen-Merck

Biosciences, Darmstadt, Germany) except GST-AtKSS23 that was expressed in Rosetta-gami (DE3)pLysS bacteria (Novagen). Proteins were purified on a glutathione column (Amersham Pharmacia Biotech) according to the manufacturer procedures. Purified proteins were frozen in liquid nitrogen in the presence of 10% (v/v) glycerol and stored at  $-80^{\circ}\text{C}$ .

#### 4.3. Expression of AtKSS *in vivo*

For transient expression in onion epidermal cells, full-length AtKSS and truncated AtKSS were cloned into the pOL-LT plasmid [31], under the control of a double strong 35S promoter and upstream the DsRed2 fluorophore (Clontech, BD Biosciences, NJ, USA). Cells were transformed by bombardment with DNA-coated gold beads as described [32]. Cells were observed 12 to 25 hours after transformation using a Leica TCS-SP2 confocal microscope (Leica Microsystems, Wetzlar, Germany). DsRed2 fluorophore was visualized under the following illumination conditions: 543 nm excitation, 580 to 620 nm emission.

#### 4.4. Microtubule cosedimentation assays

Tubulin was purified from beef brain according to [33]. Taxol-stabilized microtubules were prepared using a procedure adapted from [34] according to the recommendations described at the web site of Tim Mitchison's lab (<http://mitchison.med.harvard.edu/protocols/poly.html>). For quantification, 10  $\mu\text{l}$  of taxol-stabilized microtubules were depolymerized for 10 min at  $4^{\circ}\text{C}$  in 90  $\mu\text{l}$  of cold depolymerizing buffer (50 mM KCl, 5 mM  $\text{CaCl}_2$ , 1 mM DTT in cold assembly buffer). The concentration of tubulin dimers was calculated using the absorbance of the sample at 280 nm and an extinction coefficient of  $115,000 \text{ M}^{-1} \text{ cm}^{-1}$ . Considering the quantification procedure, the amount of taxol-stabilized microtubules used in the assays is indicated as concentrations of tubulin dimers (*figures 2 and 3*). To limit adsorption of AtKSS on plastic surfaces, all eppendorf tubes and tips were siliconized [35]. To identify the domains of AtKSS involved in microtubule binding, taxol-stabilized microtubules (0 to 3  $\mu\text{M}$ ) were incubated for 20 min at  $25^{\circ}\text{C}$  with 0.4  $\mu\text{M}$  of the different GST-tagged truncated AtKSS in Binding Buffer (20 mM HEPES pH 7.2, 2 mM  $\text{MgCl}_2$ , 0.1 mM EGTA, 20  $\mu\text{M}$  taxol, 0.5 mg/ml casein). Reactions (50  $\mu\text{l}$ ) were

spun through a glycerol cushion (30 % (v/v) glycerol in binding buffer) for 5 min at 100.000 *g* as described (McNally *et al.*, 2000). Supernatant and the resuspended microtubule pellet were analysed by western blot using an anti-GST antibody and the ECL kit (Amersham Pharmacia Biotech). For  $K_d$  determination, 1  $\mu$ M of taxol-stabilized microtubules and various concentrations of AtKSS constructs (from 0 to 5  $\mu$ M) were used in the assays. The amount of AtKSS constructs free and bound to microtubules were determined by scanning Coomassie blue stained gels and comparing the intensities with standards that were loaded on the same gel using NIH-image software (<http://rsb.info.nih.gov/nih-image/>). The amount of AtKSS bound to microtubules is described by

$$[KMT] = \frac{(K_d + [MT_{tot}][K_{tot}]) - \sqrt{(K_d + [MT_{tot}][K_{tot}])^2 - 4[MT_{tot}][K_{tot}]}}{2}$$

where [KMT] is the concentration of AtKSS bound to microtubules, [MT<sub>tot</sub>] and [K<sub>tot</sub>] are the total concentrations of tubulin dimers and AtKSS respectively. For  $K_d$  calculation, data were analysed using Kaleidagraph (Synergy Software, PA, USA).

#### 4.5. Electron microscopy

AtKSS was purified from insect cells as previously described [17]. Specimens at a concentration of 0,1 mg/ml were imaged by negative stain electron microscopy using a JEOL 1200 EXII microscope operating at 100 kV and a nominal magnification of 40,000x. The stain used was sodium silico-tungstate.

#### 4.6. Gel filtration chromatography

Gel filtration of katanin constructs was performed on a fast protein liquid chromatography system with a Superdex<sup>TM</sup> 200 HR 10/30 column (Amersham Pharmacia Biotech). Column equilibration and chromatography were performed in column buffer (20 mM Tris pH 8, 0.1 M KCl, 2 mM MgCl<sub>2</sub>, 1 mM DTT). The column was calibrated with the following molecular mass standards: ribonuclease A (13.7 kDa), ovalbumin (43 kDa), albumine (67 kDa), aldolase (158 kDa), catalase (232 kDa), ferritin (440 kDa), Dextran Blue 2000 (Amersham Pharmacia Biotech).

#### 4.7. Chemical cross-linking

The zero-length chemical cross-linker 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide hydrochloride (EDC), along with N-hydroxyuccinimide (NHS) were obtained from Pierce (Rockford, IL, USA). Stock solutions of 100 mM EDC and NHS were made fresh in dry DMSO immediately before use. Purified AtKSS proteins (20  $\mu$ g) were incubated with 1 mM EDC and 1 mM NHS at room temperature. After 10 min, 1 mM of EDC and NHS were again added to the samples. After 20 min the reaction was stopped by the addition of 9 mM glycine. The cross-linked AtKSS constructs were observed by Coomassie-blue stained gels and western-blot.



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## TABLE LEGEND

Table I. Number of oligomers of the different AtKSS constructs.

## FIGURE LEGENDS

Fig. 1. AtKSS constructs used in this study. (A) Alignment of amino acid sequences of most known katanin p60 including human (Hs), *Xenopus laevis* (Xl), *Strongylocentrotus purpuratus* (Sp), *Drosophila melanogaster* (Dm), *Arabidopsis thaliana* AtKSS (At), *Chlamydomonas reinhardtii* (Cr) and *Caenorhabditis elegans* (Ce). Identical and similar amino acids are highlighted in black and gray respectively. Dashes represent gaps. (B) Diagram of the different AtKSS constructs corresponding to combination of domain 1 (aa 1 to 34, black box), domain 2 (aa 35 to 210, white box) and domain 3 (aa 211 to 523, gray box). C, Coomassie blue-stained SDS-PAGE gel of the purified AtKSS constructs. Lane 1, GST. Lane 2, GST-AtKSS1. Lane 3, GST-AtKSS2. Lane 4, GST-AtKSS12. Lane 5, GST-AtKSS3. Lane 6, GST-AtKSS23. Lane 7, GST-AtKSS13. Masses of molecular markers are indicated in kDa.

Fig. 2. Binding of AtKSS constructs to microtubules *in vitro*. 0.4  $\mu$ M of GST-AtKSS constructs were incubated with various amount of microtubules. Samples were centrifuged to pellet microtubules and the amount of GST-AtKSS proteins bound to microtubules was evaluated by densitometry. A, Western-blot of supernatant (S) and pellet (P) of GST-AtKSS proteins probed with GST antibodies. Concentrations of microtubules used in each assay (indicated as the concentration of tubulin dimers) are indicated. B, Results are expressed as percentage of total GST-AtKSS constructs that pelleted with microtubules versus tubulin dimers concentration.

Fig. 3. Affinity of the microtubule-binding domain of AtKSS for microtubules. The amount of AtKSS12 (open square, dotted line), GST-AtKSS12 (filled square, solid line) and GST-AtKSS2 (filled circle, solid line), are plotted as a function of the total amount of AtKSS used for co-sedimentation assays with microtubules. The fitting curve was defined using the equation given in Methods, and was used to determine the  $K_d$  of AtKSS12 (18.03  $\mu$ M), GST-AtKSS12 (6.87  $\mu$ M) and GST-AtKSS2 (5.10  $\mu$ M) for microtubules.

Fig. 4. Localization of AtKSS12 *in vivo*. (A) Onion epidermal cell transiently expressing AtKSS12-DsRed and observed 16 hours after bombardment. A dense and disorganized network of filaments is detected in the cortex of the cell. (B to D) Time-lapse observation of an onion cell expressing AtKSS-DsRed and treated with 20  $\mu$ M oryzalin, a microtubule-depolymerizing drug, for 0 min (B), 10 min (C) and 20 min (D). After 20 min of treatment, no microtubules were observed.

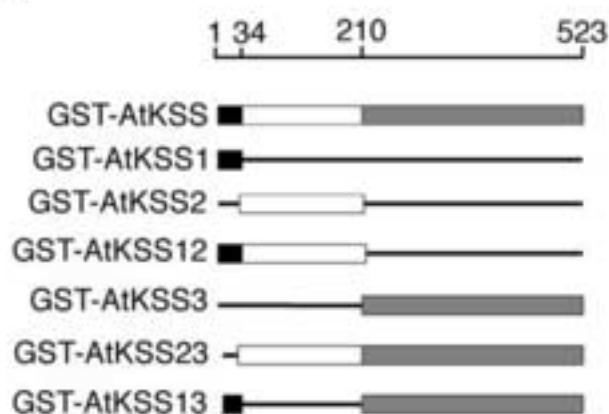
Fig. 5. Structure and oligomerization of AtKSS. (A to C) Electron micrographs of AtKSS purified from baculovirus-infected insect cells. Rings are 25-30 nm diameter. (D to F) Cross-linking of GST-AtKSS12, GST-AtKSS13 and AtKSS12. GST-AtKSS12 (D), GST-AtKSS13 (E) and AtKSS12 (F) were incubated in the absence (lanes -) or in the presence (lanes +) of cross-linker molecules. Samples were analysed by coomassie blue-stained SDS-PAGE (gel) and by western-blot an anti-GST antibody (GST-AtKSS12 and GST-AtKSS13) or anti-AtKSS antibodies (AtKSS12). Masses of molecular markers are indicated in kDa. Bar= 20  $\mu$ m

Fig. 6. ATPase and microtubule-severing activity of GST-AtKSS23. (A) ATPase activity of 0.23  $\mu$ M GST-AtKSS13 (squares) and 0.23  $\mu$ M GST-AtKSS23 (circles) was followed at 340 nm in the absence (open symbols) and in the presence (black symbols) of 0.25  $\mu$ M tubulin dimers. The decrease of the absorbance accompanying ATPase activity shows that GST-AtKSS23 has a basal ATPase activity (open circles) that is two-fold stimulated by microtubules (black circles). GST-AtKSS23 has a very low ATPase activity (open squares) that is not stimulated by microtubules (black squares). (B) Severing of microtubules by GST-AtKSS23 followed by spectrometry at 350 nm. The decrease of the absorbance of taxol-stabilized microtubules incubated GST-AtKSS23 (continuous line) or with baculovirus-expressed full-length *Arabidopsis* katanin (broken line) shows that these two proteins severed microtubules with a similar activity. Microtubules incubated with the buffer alone did not depolymerize (dotted line). (C-F) Severing of fluorescent taxol-stabilized microtubules by GST-AtKSS23 observed by microscopy. Fluorescein-labelled taxol-stabilized microtubules were incubated either with the reaction buffer (C-D) or with GST-AtKSS23 (E-F). After 15 min in the presence of GST-AtKSS23, most microtubules were severed (F). Bar= 10  $\mu$ m

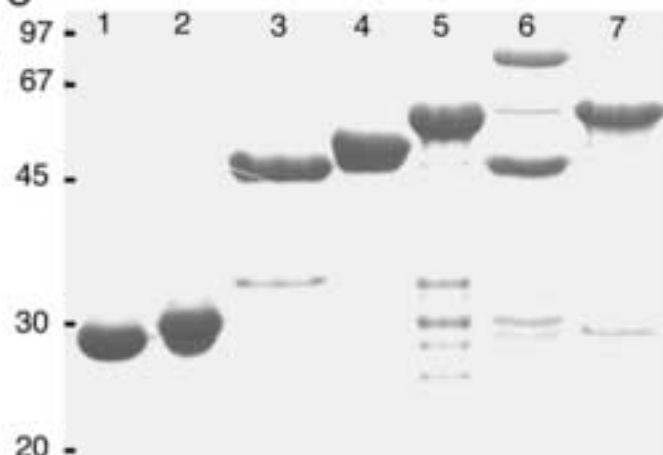
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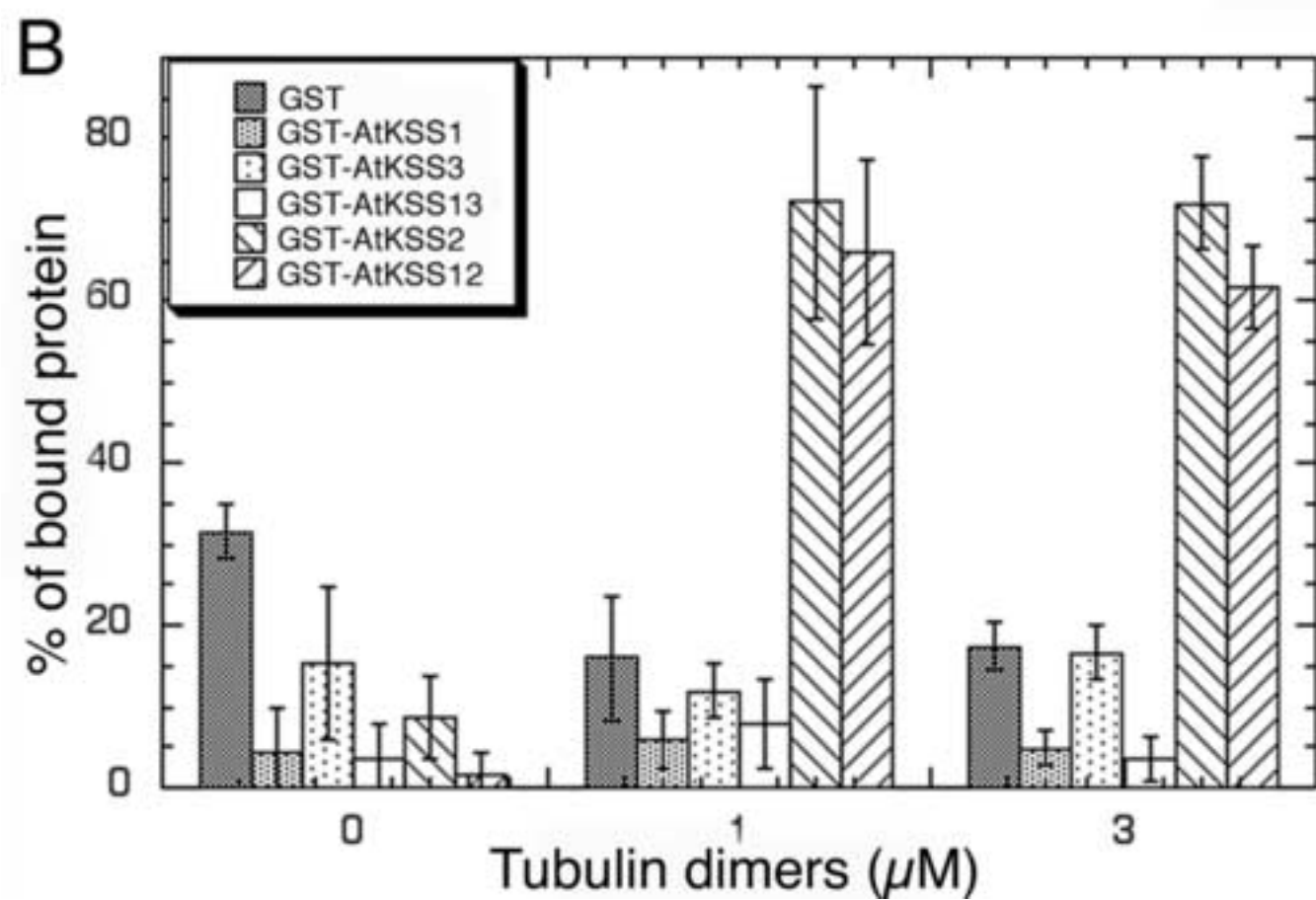
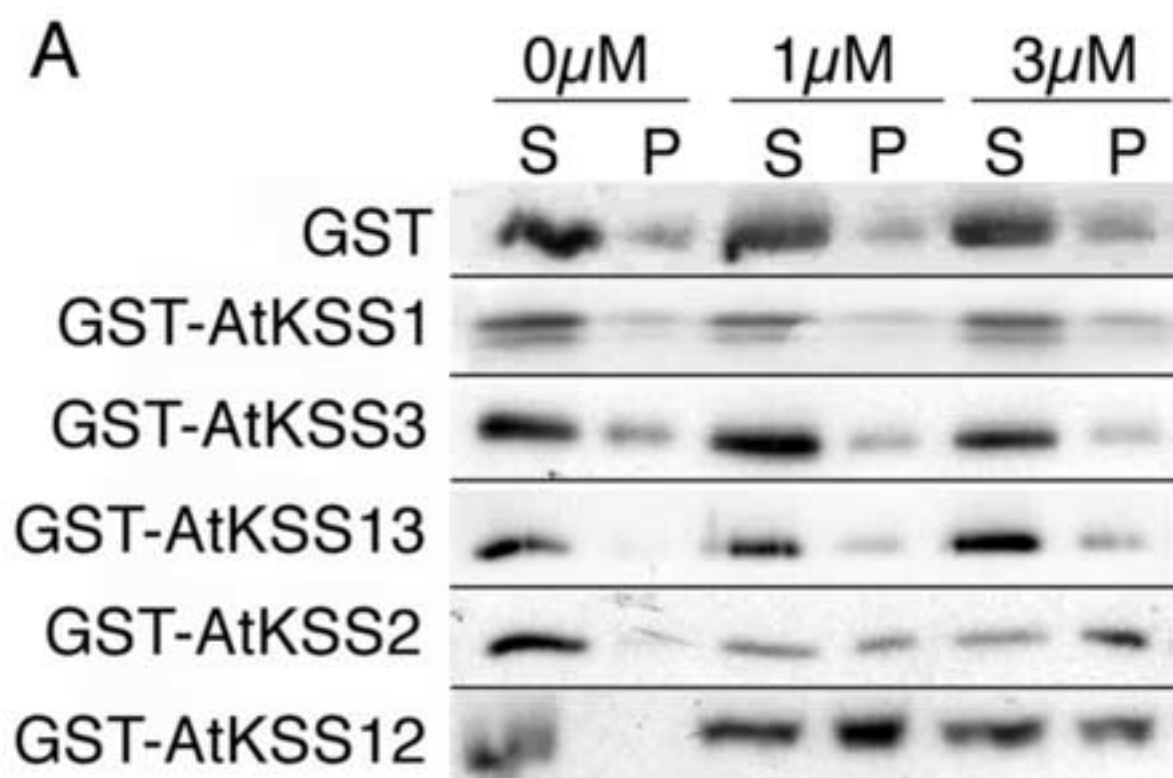
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Xl	.....KSELMLISENVKLAAREYALLDNYDSAMVYVYQGVLDCHNREYEVKDTYLOOKHOOWQOEINVAKHVVDINHTLESF	76
Sp	.....KSVDEICENTENGRYALLDNYEYSLVYVYQGVLDQCELETSVREHPQKKNHOTIRQESSQRYEHVKNITKTLNGF	76
Dm	HSITLLRGGKQKRTTTCVGMVMAKTTFFEEICENAKLARDHALTNYDASCIYYEGLQGLLARQLKATADPLRKGKSHKINOOSSCHAKIKAIQHTLODI	100
At	.....NVGSSNSIAGLQDHLKLAAYALSSYDYSVIFPQGAIAQCNKINLNTDQPLARTKNHNVKKAIMEETEVVWQDAERRAF	81
Cr	.....HGGADVQASTIAVTTGRCCMSDYASACTYENAVELITVYVREIGDPAHQASALLKEVODERSSYWQDAEARGH	79
Ce	.....MNGDVQSVYRQYLIRACVARTHSDAPRNEAGDLLHQEHTDVKSKISASRDEHDARNTFLRAEAMKLVQCNVEDDGL	82
Hs	ELDSFP.....LKAAQH.....DLFASGGVYSHVVF.....VERPSE.....GPRKRQSSQYSDPKS.....HGNRPSTT	133
Xl	ELDSFP.....VKTTQH.....EPPSSDGGVYSLVVF.....VERPSE.....GPRKRQSVQCNDNKS.....HNNRFSAA	133
Sp	SEPAAP.....EPAPNHRAAFPSSHQAAKPAAAEAKDPDVFPPPT.....VDRPSEPYQ...RAARKDPPRSEPSK.....PANKAPGN	153
Dm	SLDLQSTKFAKLRRLRQLSEESTTSKDPKDPDPDINTFPFKDFDVFYGGPKPEPTTQAVGRBAANNRRTTTPATQNSRPSSTIQS.....TARNGPAS	195
At	KEAPFRR.....AASPPINTEKSSVFPQPLDEYFISGGGGMDDFVAKPFR...DVTSEPARAG...QTGTRKSPQOGAWARG.....PTTITGPA	164
Cr	RYDVAR.....SHSYPMNDPRGSAAYDDGYQPIVCHNSGQEDFVYKPRTRGGRSGAPAGAPFRARRSQDNDKLPWANNNHGGGGPPAGNAA	177
Ce	HEAKRQS.....GSPFPADDFYKSSP.....PLPSSSKFGATKKG.....VGAAGPRP	129
Hs	VR.VIRSAQNVNRDRGKA...VRCHEKKEQNKGREEKNKSPAA.....VTEPETNKFFST.GYKDLVEALERDIISQNF.NVHMD	210
Xl	AGPHLP.SARNAMVVMKFP...VVRREKED...ALIKNKSSAD.....VSETEVKRFDGS.GYKDLVEALERDIISQNF.NIHMD	207
Sp	DGGCPCPDRRGOARSGGGRGGGARGSDKDKNRGGKSDKDKKAPS.....GEGDEKKFQPA.GYKDLVENLERDIVQRM.NVHMA	235
Dm	TNSNSSTATAPSGGARTWGRAGGRKLTSTNTNEARDOSTRAGNNGGAAGDGENGDPQAAQEERKQFMNHIAELVDIILERDILQDF.KVHMS	294
At	SGGGGGATSKSTAGARSS...TAGKGAASKSNAERHNGDAE.....DGSKRGLYGP...DEDLAAMLERDVLQSTF.CVRHDD	240
Cr	GPSCHAGAPRRPTGNAPAGNAAAGSYDRPWRQNHQCGQGAAPGGKPGDKGS.....AGEKAKKQYSGP...DQELAAMLERDVLQSTF.SVHMD	266
Ce	REISTSTSHSTNPADVKP.....ANPTQGLP.....QMSAGDSBAS.AYDAYVQAVRGTNATHTNTHTSHLQ	194
Hs	IADLEENKLLBEAVVLPNHWPEYFCGRSPKGVLMVGPVOTGKTLIAGANASDSEKTFPFRVSSSLASFTGQSEIVVRLPSEANFYAPSIIFIDEI	310
Xl	IADLEENKLLBEAVVLPNHWPEYFCGRSPKGVLMVGPVOTGKTLIAGANASDSEKTFPFRVSSSLASFTGQSEIVVRLPSEANFYAPSIIFIDEI	307
Sp	IAGTEENKLLBEAVVLPNHWPEYFCGRSPKGVLMVGPVOTGKTLIAGANASDSEKTFPFRVSSSLASFTGQSEIVVRLPSEANFYAPSIIFIDEI	335
Dm	IADLHDNRLBEAVVLPNHWPEYFCGRSPKGVLMVGPVOTGKTLIAGANASDSEKTFPFRVSSSLASFTGQSEIVVRLPSEANFYAPSIIFIDEI	394
At	VAGLEENKLLBEAVVLPNHWPEYFCGRSPKGVLMVGPVOTGKTLIAGANASDSEKTFPFRVSSSLASFTGQSEIVVRLPSEANFYAPSIIFIDEI	340
Cr	IAGLEENKLLBEAVVLPNHWPEYFCGRSPKGVLMVGPVOTGKTLIAGANASDSEKTFPFRVSSSLASFTGQSEIVVRLPSEANFYAPSIIFIDEI	366
Ce	IIGHNVDVQLBEAVVLPNHWPEYFCGRSPKGVLMVGPVOTGKTLIAGANASDSEKTFPFRVSSSLASFTGQSEIVVRLPSEANFYAPSIIFIDEI	294
Hs	DSICRRCSSSEHNSRRVNSLQVQDQVGGTSENDDPSI.....VVVLAATNPWQIDEALRRRDEERIVIPLPSAKDRRELERISEHELELAD	402
Xl	DSICRRCSSSEHNSRRVNSLQVQDQVGGTSENDDPSI.....VVVLAATNPWQIDEALRRRDEERIVIPLPSAKDRRELERISEHELELAD	399
Sp	DSICRRCSSSEHNSRRVNSLQVQDQVGGTSENDDPSI.....VVVLAATNPWQIDEALRRRDEERIVIPLPSAKDRRELERISEHELELAD	427
Dm	DSICRRCSSSEHNSRRVNSLQVQDQVGGTSENDDPSI.....VVVLAATNPWQIDEALRRRDEERIVIPLPSAKDRRELERISEHELELAD	482
At	DSICRRCSSSEHNSRRVNSLQVQDQVGGTSENDDPSI.....VVVLAATNPWQIDEALRRRDEERIVIPLPSAKDRRELERISEHELELAD	433
Cr	DSICRRCSSSEHNSRRVNSLQVQDQVGGTSENDDPSI.....VVVLAATNPWQIDEALRRRDEERIVIPLPSAKDRRELERISEHELELAD	466
Ce	DSICRRCSSSEHNSRRVNSLQVQDQVGGTSENDDPSI.....VVVLAATNPWQIDEALRRRDEERIVIPLPSAKDRRELERISEHELELAD	382
Hs	DVDEASTAENHNSGSDITVYCPASLMAHNRRIEGLTFEEIN.NLSKEENHM...PTMHDPEMAKKNVSKVSAADIERTEKMEFFPCFC	491
Xl	DVNEICIAENHNSGSDITVYCPASLMAHNRRIEGLTFEEIN.NLSKDDHM...PTMHDPEMAKKNVSKVSAADIERTEKMEFFPCFC	486
Sp	DIDEKSAKKNVSKVSAADIERTEKMEFFPCFC...ESTPAQFLLAGQVSKVGGKEDLVYTMANKKPGV	516
Dm	SVDLTYANELKQSGSDITVYCPASLMAHNRRIEGLTFEEIN.OLATHEVDL...PVHNDPEHNSRCKVSRADLDKTEKMEFFPCFC	571
At	DVNEEDVARTDQSGSDITVYCPASLMAHNRRIEGLTFEEIN.NNSKDDSHD...PVAMCDEHNSRCKVSRADLDKTEKMEFFPCFC	523
Cr	DVNEEDVARTDQSGSDITVYCPASLMAHNRRIEGLTFEEIN.AEAREAGTQGERIISQDFREHNSRCKVSRADLDKTEKMEFFPCFC	558
Ce	KEHYDDIARTDQSGSDITVYCPASLMAHNRRIEGLTFEEIN.TAARESLKAEVRRIDPEHNSRCKVSRADLDKTEKMEFFPCFC	475

### B

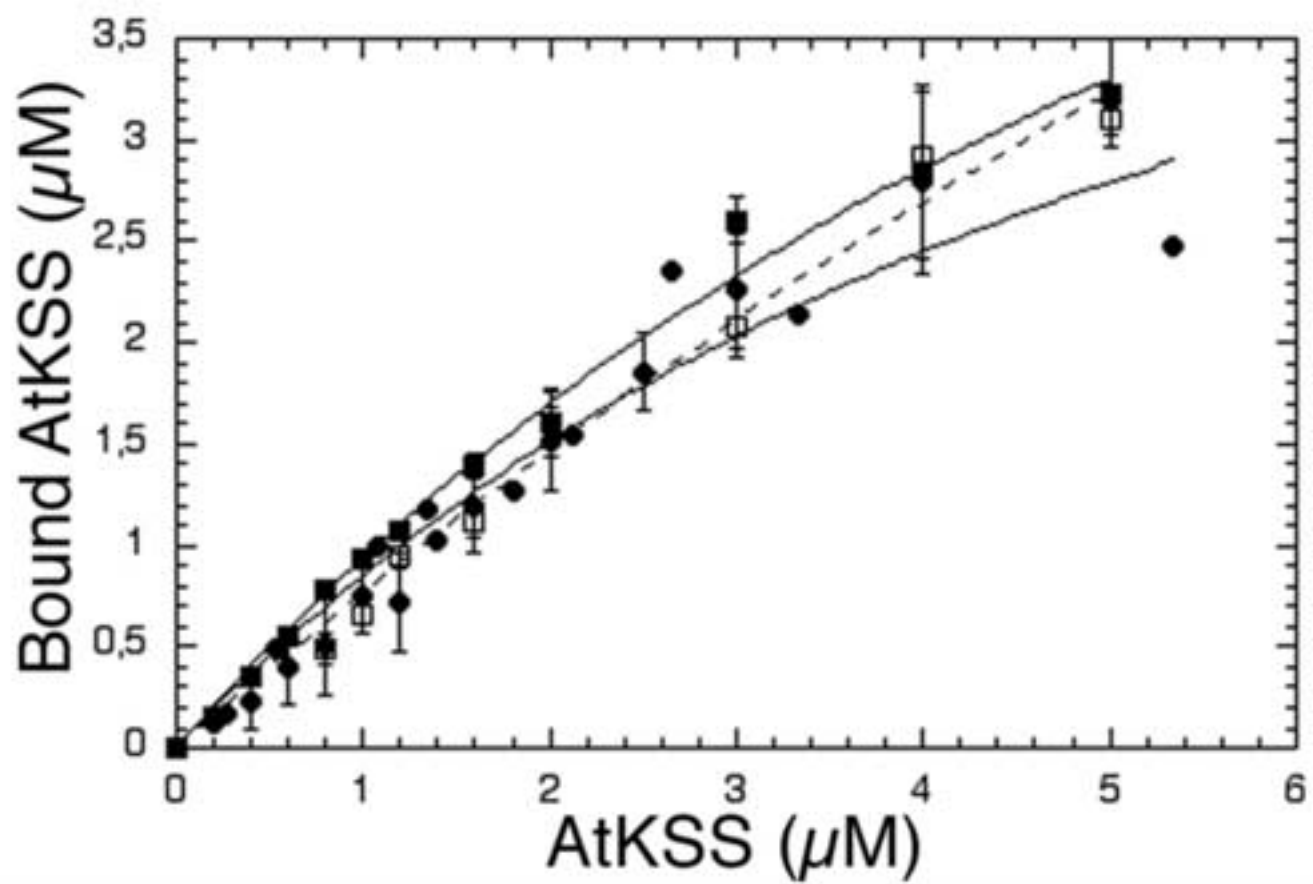


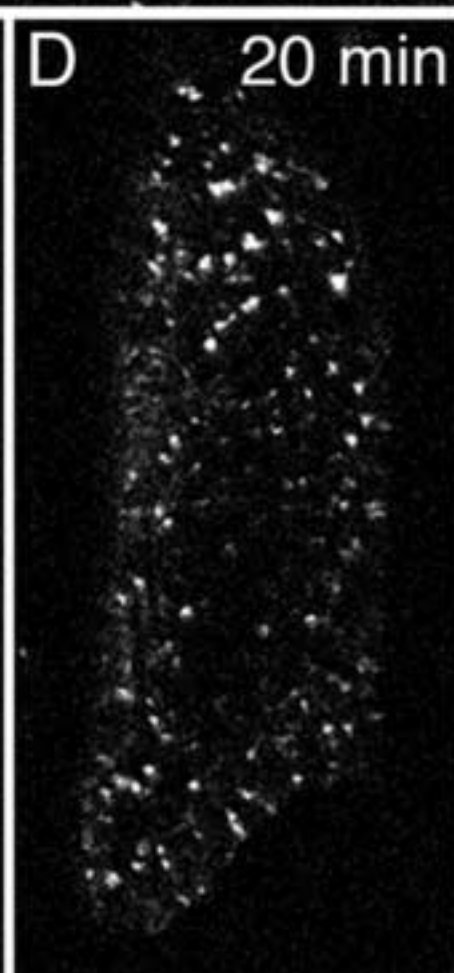
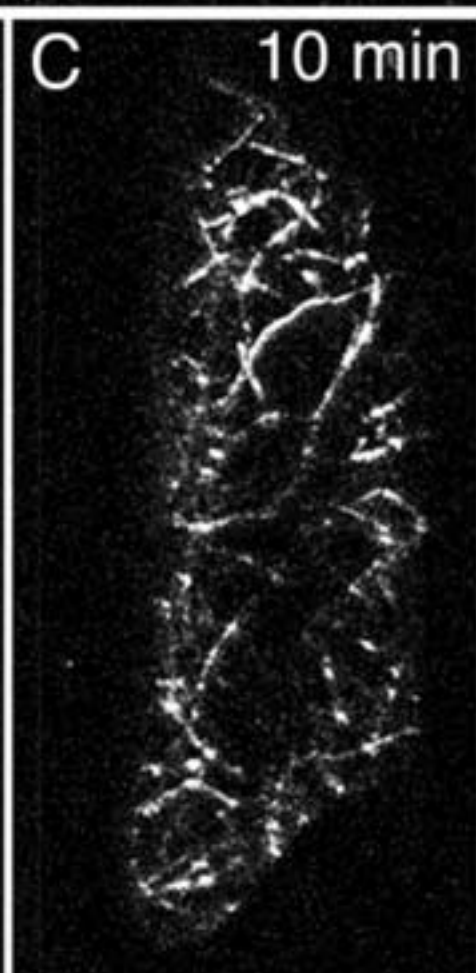
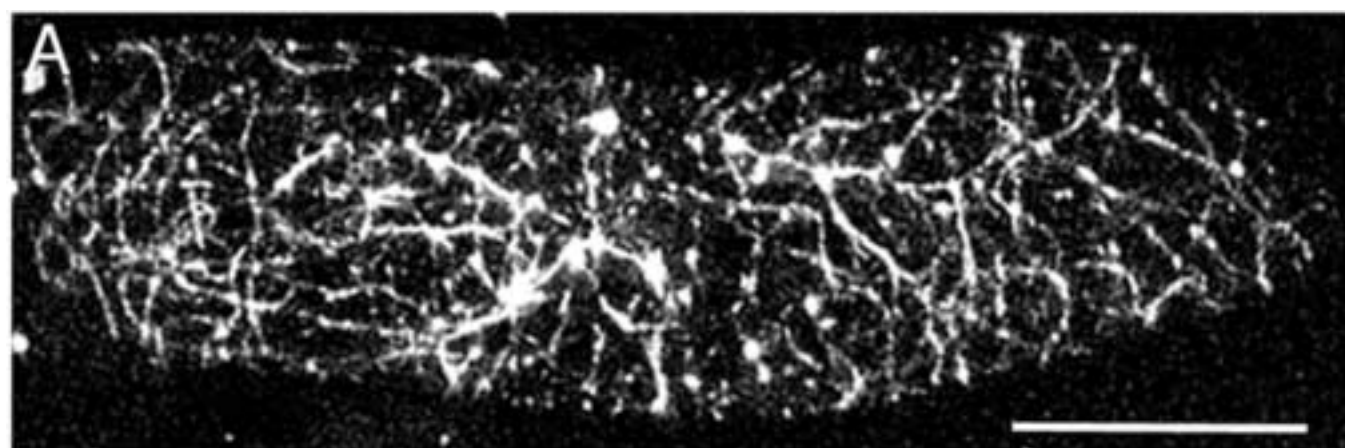
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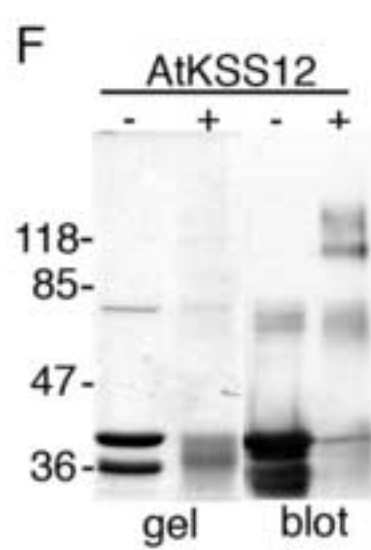
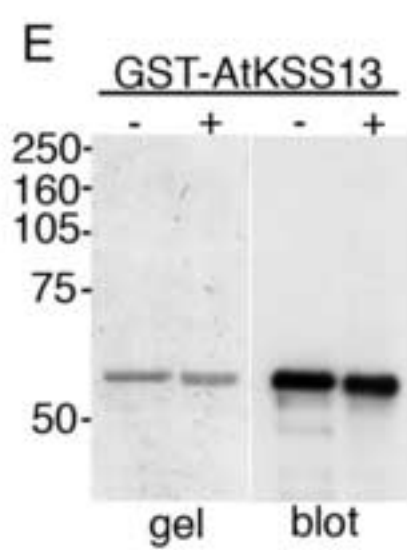
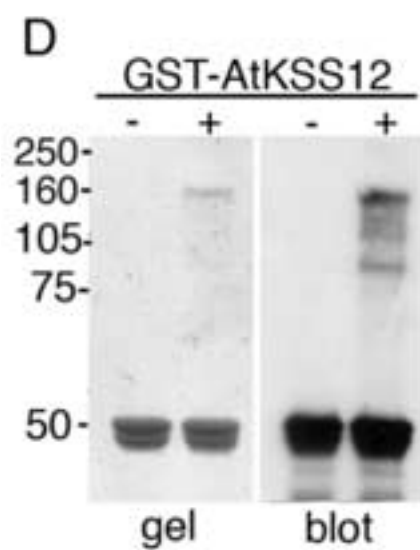
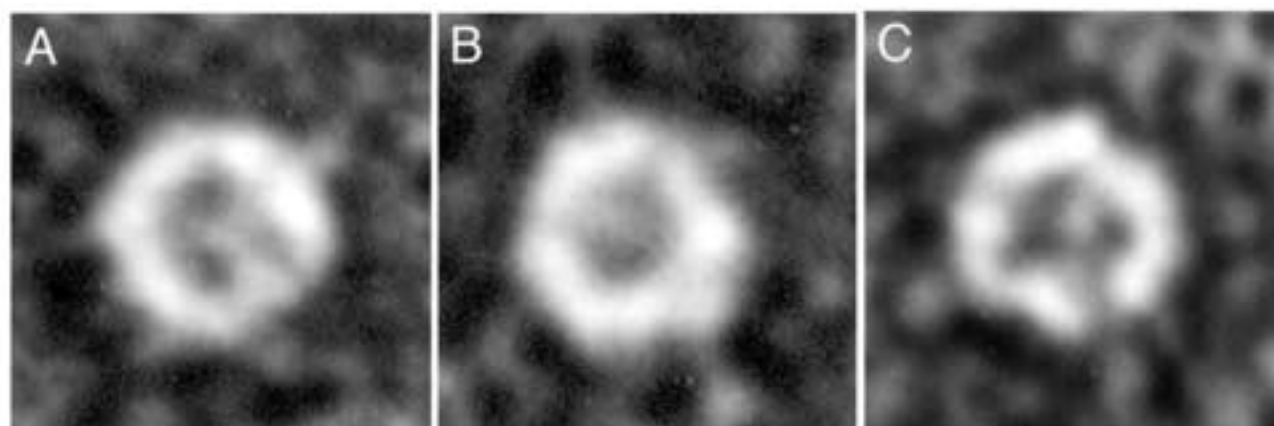


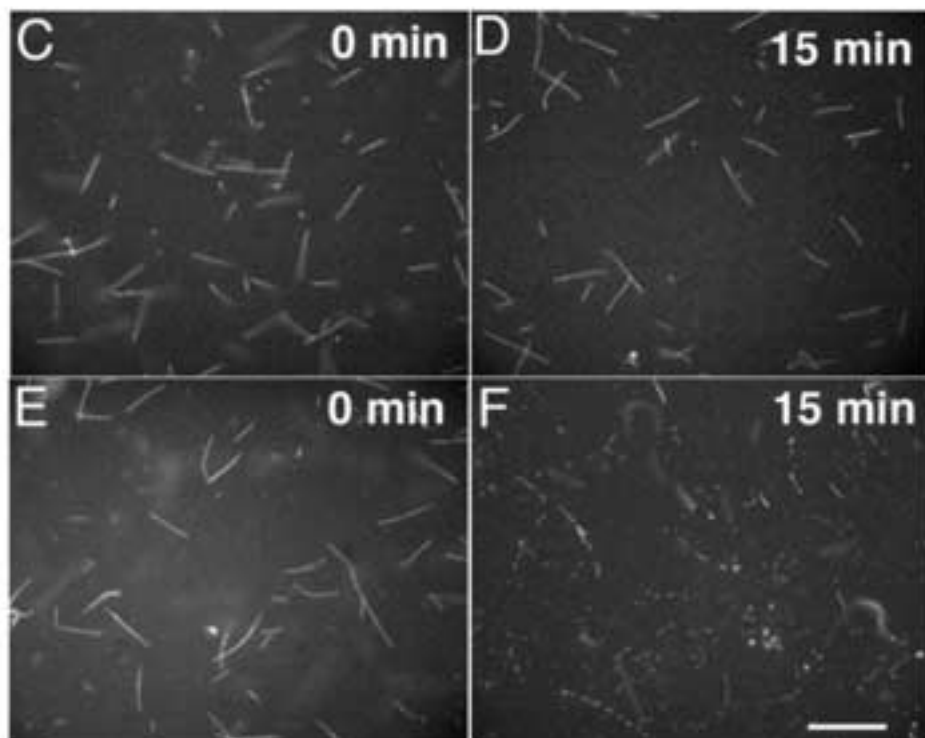
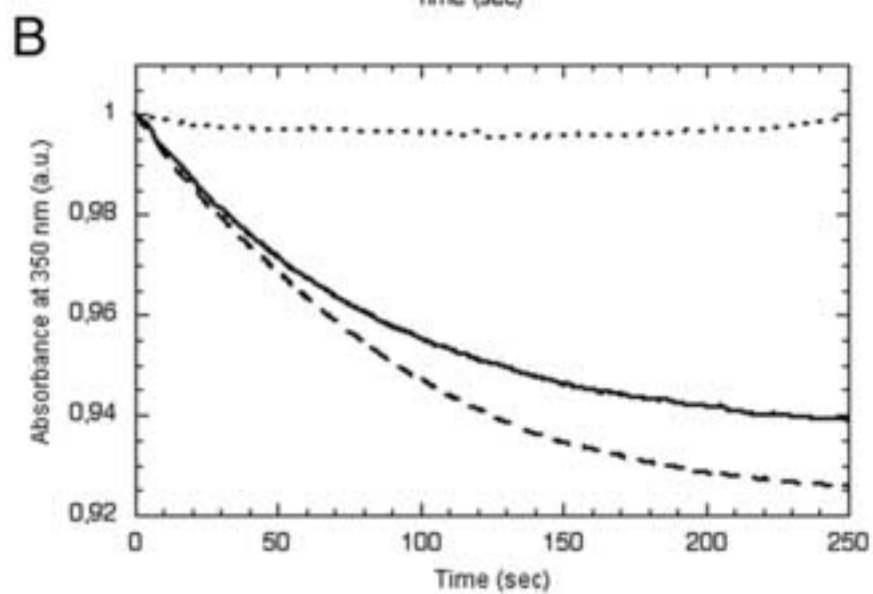
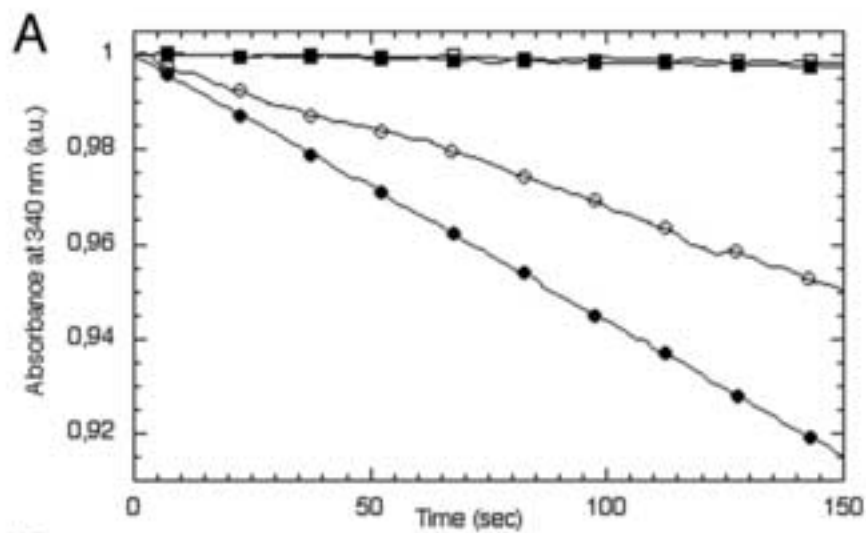












	Predicted molecular mass <sup>a</sup> (Da)	Estimated molecular mass <sup>b</sup> (Da)	number of oligomers
GST	27.783	49.477	1.78
GST-AtKSS1	31.485	57.464	1.83
GST-AtKSS3	62.810	102.058	1.63
GST-AtKSS13	66.512	147.711	2.22
GST-AtKSS2	46.273	126.233	2.73
GST-AtKSS12	49.975	162.465	3.25

<sup>a</sup> Molecular masses predicted from amino acid sequence.

<sup>b</sup> Values calculated from the gel filtration data.