Control of microtubule dynamics by the antagonistic activities of XMAP215 and **XKCM1** in Xenopus egg extracts

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Microtubules are dynamic polymers that move stochastically between periods of growth and shrinkage, a property known as dynamic instability. Here, to investigate the mechanisms regulating microtubule dynamics in Xenopus egg extracts, we have cloned the complementary DNA encoding the microtubule-associated protein XMAP215 and investigated the function of the XMAP215 protein. Immunodepletion of XMAP215 indicated that it is a major microtubule-stabilizing factor in Xenopus egg extracts. During interphase, XMAP215 stabilizes microtubules primarily by opposing the activity of the destabilizing factor XKCM1, a member of the kinesin superfamily. These results indicate that microtubule dynamics in Xenopus egg extracts are regulated by a balance between a stabilizing factor, XMAP215, and a destabilizing factor, XKCM1.

he pattern and dynamics of microtubule networks are altered during cell division and differentiation. Despite the importance of these changes, little is known about how they are controlled. Microtubules exist in two states, either growing or shrinking, and they transit stochastically between these two states, a behaviour known as dynamic instability¹. The transition from growth to shrinkage is called a catastrophe, and the transition from shrinkage to growth, a rescue. Thus, microtubule behaviour can be

described by four parameters, namely the growth rate, the shrinkage rate, the catastrophe frequency, and the rescue frequency^{2,3}. In vitro, in pure tubulin solutions, two of these parameters (growth rate and catastrophe rate) are tightly linked and strictly determined by the free tubulin concentration². In contrast, the different parameters of microtubule dynamics are regulated independently in vivo, indicating that the combinatorial action of several regulatory factors may determine the values of each of these parameters and the

a								b	- ch-TOG
MGDDSEWMKL	PIDQKCEHKV	WKARLNGYEE	AVKLFQKIVD	EKSPEWSKYL	GLIKRFVTES	NAVAQLKGLE	70		CIFICO
AALVYVENAH	VAGKTTGEVV	NGVVNKVFNQ	PKARAKELGA	DICLMYVEIE	KAEVVQEELL	KGLDNKNPKI	140		KIAA0097
VVACVETVRK	ALSEFGSKIM	TLKPIIKVLP	KLFESREKAI	RDEAKLLAVE	IYRWIRDALR	PPLQNINPVQ	210		XMAP215
LKELEEEWVK	LPQSAPKQTR	FLRSQQDLKA	KFEQQQAAGD	DGGDDGEEEI	VPQVDAYELL	EAVEILSKLP	280		- XIVIAF 2 15
KDFYDKIEAK	KWQERKEALE	AVEALVKNPK	IEAGDFADLV	KALKTVVGKD	TNVMLVALAA	KCIAGLAAGL	350		msps
RKKFGS YAGH	IVPTILEKFK	EKKPQVVQAL	QEAIDAVFLT	TTLQNISEDV	LAVMDNKNPA	IKQQTSLFLA	420	\neg	A. thaliana (#4263790)
RSFRHCTPST	LPKSL <u>LKPFC</u>	VALLKQINDS	APEVRDAAFE	ALGTAQKVVG	EKAVNPFLAE	VDKLKLDRIK	490		Stu2
ECADKAELAN	GKKGGAAAGE	KKETKA PAAA	PGKPVPNQGA	AAEKDAGKAA	AAPKKAPAAK	PGGPVKKAKA	560	$\langle \langle \langle \rangle \rangle$	
PASSGATAKG	KKAVENKEII	EQELSPEACE	ERAAAVLPAS	CMQQLDSSNW	KERLASMEEF	QKTVESMERN	630		dis1
DIPCQALVKM	LAKKPGFKET	NFQVMQMKLH	IVALIAQKGN	FSKTSACAVL	DGLVDKVGDV	KCGGNAKEAL	700		S. pombe (#4164426)
SGIAEACTLP	WTAEQVVSLA	FAQKNPKNQS	ETLNWLSNAI	KEFGFTGINV	KAFISNVKTA	LAATNPAIRT	770		<pre></pre>
SAITLLGVMY	LYMGAPLRMF	FEEEKPALLS	QIDAEFEKMK	GQTPPVSIRG	SKHGSGRDEG	EEGEEQDEDA	840		ZYG-9
PADVTDLLPR	TDISDKISSD	LVSKIEDKNW	KIRKEGLDEV	TAIINEAKFI	QPSIGELPSA	LKGRLNDSNK	910		-216-5
ILVQQTLTIL	QQLSTAMGHN	IKQHVKNLGM	PIITVLGDSK	ANVRAAALGT	LKSWVDQTGM	KDWLEGEDLS	980		
EELKKENPFL	RQELLGWLAE	KLPSMRTVPS	DLQLCVPYLY	NCLEDRNGDV	<u>RKKAQEALPI</u>	FMMHIGFEKM	1050		
<u>sk</u> atsklkpa	SKDQVVALLE	KAKASMPAKP	AGPPGKASSK	QPPAVAQASA	SPPPAASSDS	GSSTSDYKPD	1120	C Int Mit	Int Mit
PKKTKPGTQA	SKAKTQSVSS	EGNTSENPSN	TSL <u>TREK</u> ANT	SLSKAKPAKQ	72PGKKAPSK	PNAKDEEDKS	1190	M _r (K)	M _r (K)
GPIYIIVPNG	KEQRVKDEKA	LKVLKWNFTT	PRDEYIEQLK	TOMSPCIARW	LQDELFHADF	QRQIKGLAVM	1260	200	200
TEHLESEKEG	VISCLDLVLK	WFTLRFFDTN	TSVLMKCLEY	LKLLFIMLSQ	EEYHLTEMEG	TSFLPYLMLK	1330	200	200
VGEPKDIVRK	DVRAILTKMC	QVYPASKMFN	FVMEGTKSKN	SKQRAECLEE	LGCLVESYGM	NVCQP <u>TPAK</u> A	1400		
LKEIAIHIGD	RDTTVRNAAL	NTIVTVYNVH	GEQVFKLIGN	LSEKDMSMLE	ERIKRAGKKQ	AAAAPAKQVE	1470		
EKPQRVQSAN	ASILRKAPPE	DMSSKLNQAR	NMCGHTEPSH	SVPREFQLDL	DEIENDNGTV	RCEMPALVQH	1540		116
KLDEIFEPVL	IPEPKIRAVS	PHFDDMHSNT	ASTINFVISQ	VASVDINASI	QALAQIDEVL	RQEDKAEAMS	1610	116	
GHIDQFLIAT	FMQLRLAYNT	HMADERLDKD	DIVRLYSCII	GNMISLFQME	SLAREASTGV	LKDLMHGLIS	1680	96 -	96
LMLDARIEDL	EEGQQVVRSV	NLLVVKVLEK	SDQTNIISAL	LMLLQDSLLA	TASSPNFSEL	VMKCLWRMIR	1750		
LLPEAINNLN	LDRILLDIHN	FMRVLPKEKL	KQHKSEMPMR	TLKTLLHTLC	KLKGPKIMDH	LSMIENKHES	1820		
ELEAHLLRVM	KHSIDRTGSK	GDKETEKGAS	CIEDKVGKAN	VSDFLAEMFK	<u>KIGSKENTKE</u>	GLAELYEYKK	1890		66 -
KYSDADIKPF	LKNSSQFFQS	YVERGLRLIE	MEREGKARIA	PNTGMSTHVT	EMTPLPTVTN	TAAPVSNTNG	1960	66	
EEVGPSVYLE	RLKILRQRCG	LDNAKQDERP	PLTSLLSKSS	APAVVSSTDM	LHSKLSQLRE	SREQFQHVEL	2030	S.C. SALE	
DSNQTYPSTT	TSSSASSTNI	DDLKKRLERI	KSSRK				2065		
								N-terminal	C-terminal

Figure 1 XMAP215 amino-acid sequence and homology to other members of the protein family to which it belongs. a, Amino-acid sequence of XMAP215. Red underlining, predicted sites of phosphorylation by the kinase MARK; green underlining, the bipartite nuclear-localization signal; blue underlining, HEAT repeats; red italics, the insertion found in XMAP215 that is absent in ch-TOG/KIAA0097; bold

underlining, a putative site of phosphorylation by Cdc2. b, Phylogenetic tree of XMAP215 relatives. msps, Mini spindles proteins. c, Specificity of anti-ch-TOG antibodies. Immunoblots done using 80µg interphasic (Int) and mitotic (Mit) Xenopus egg extracts show that affinity-purified antibodies raised against the N and C termini of ch-TOG recognize a single protein of M, 215K.

N-terminal

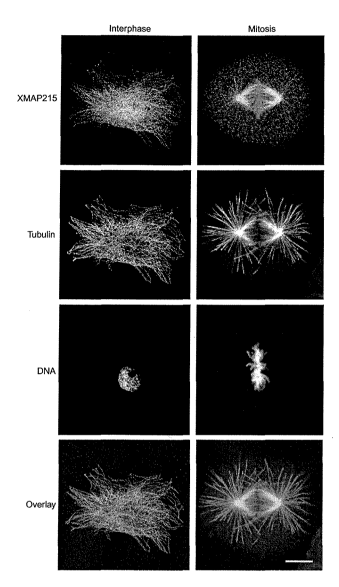


Figure 2 Immunolocalization of XMAP215 protein in XL177 cells. An interphasic cell is shown on the left, and a mitotic cell is shown on the right. In the overlay, microtubules are green, XMAP215 is red, and DNA is blue. Scale bar represents $10 \,\mu$ m.

resulting properties of the microtubule arrays^{2,4}.

Various microtubule-regulating factors have been identified in Xenopus laevis egg extracts. One of these, XMAP215, has the unusual ability to increase both growth and shrinkage rates without affecting the rescue and catastrophe frequencies^{5,6}. In contrast, XMAP230 mostly reduces the catastrophe frequency7 whereas XMAP310 increases the rescue frequency⁸. Two factors that induce catastrophes, albeit by different mechanisms in the presence of pure tubulin, are Stathmin/Op18, a heat-stable protein of relative molecular mass 18,000 (M. 18K), and XKCM1, a member of the kinesin superfamily⁹⁻¹³. Although the identification of these factors represents significant progress, our understanding of the regulation of microtubule dynamics in vivo is still fragmentary. Control of microtubule dynamics is likely to involve a combination of destabilizing and stabilizing factors. The challenge is to identify these important factors and to understand how modulation of one factor in the context of the other factors may be involved in changing the length dis-

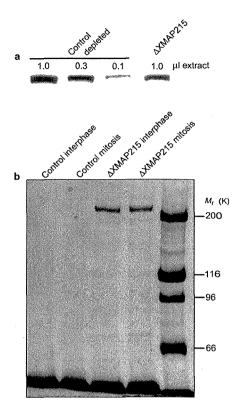


Figure 3 **XMAP215 depletion from Xenopus egg extracts. a**, Immunoblot showing the depletion efficiency. 1 μ l, 0.3 μ l and 0.1 μ l of a control depleted Xenopus egg extract and 1 μ l of a XMAP215-depleted extract were run on a gel and XMAP215 was revealed with an antibody against the ch-TOG C terminus. The amount of XMAP215 remaining in the XMAP215-depleted extract corresponds to 30% of the amount found in the control depleted extract. **b**, Coomassie-blue-stained gel of the XMAP215 immunoprecipitates from interphase and mitotic extracts. Only a band of *M*, 215K, corresponding to XMAP215, was immunoprecipitated in both interphase and mitotic extracts. The bands at the bottom correspond to the lgG heavy chains.

tribution and dynamics of microtubule arrays.

Among the microtubule-associated proteins (MAPs) characterised to date, XMAP215 is of particular interest because, in interphase, it stimulates the growth rate of microtubules at their plus (fast growing) ends¹⁴ without affecting the catastrophe frequency⁵⁶. Furthermore, XMAP215 is a member of a family of proteins that seems to be involved in the control of microtubule organization. This family includes a human homologue, ch-TOG¹⁵, the Mini spindles proteins in *Drosophila melanogaster*¹⁶, ZYG-9 in *Caenorhabditis elegans*¹⁷, Stu2 in *Saccharomyces cerevisiae*¹⁸ and p93^{Dist} in *Schizosaccharomyces pombe*¹⁹⁻²¹. We thus investigated the function of XMAP215 in control of microtubule dynamics in *Xenopus* egg extracts, and found that XMAP215 strongly modulates the catastrophe frequency by opposing the microtubule-destabilizing activity of XKCM1.

Results

Molecular cloning of XMAP215. To investigate the *in vivo* function of XMAP215, we cloned and sequenced the corresponding cDNA, which encodes a protein of 2,065 residues (Fig. 1a). The XMAP215 amino-acid sequence is highly similar to those of the human proteins ch-TOG²² (78.4% identity) and KIAA0097 (ref. 23; 79% identity). Ch-TOG and KIAA0097 are identical with the exception of a 60-amino-acid insertion at position 1,564 of ch-TOG, also present

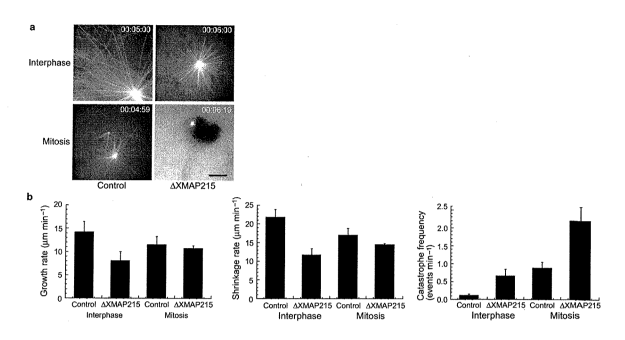


Figure 4 XMAP215 is a major regulator of microtubule dynamics. a, Images of a microtubule aster as observed during recording of movies. Microtubules in interphase and mitotic control depleted extracts have a normal size after 5 min of recording. However, in interphasic XMAP215-depleted extracts, microtubules are shorter than those in control extracts after 6 min of recording. Microtubules are barely visible in mitotic XMAP215-depleted extracts, such that images had to be recorded

for 15 min instead of 5 min. Scale bar represents 10 μm. **b**, Quantification of microtubule dynamics. In comparison with controls, the growth rate and the shrinkage rate decrease after XMAP215 depletion in interphase extracts but not in mitotic extracts. The catastrophe frequency is increased in both interphase and mitotic XMAP215-depleted extracts relative to controls. Data shown are means±s.e.m.

in XMAP215, and two amino-acid changes from alanine to glutamate at positions 1,814 and 1,822 in the KIAA0097 sequence. XMAP215 has another 36-amino-acid insertion at position 1,140 of ch-TOG. XMAP215 is a member of a protein family that includes p93^{Dist} and a putative 809-amino-acid protein (GenBank accession number 4164426) from *S. pombe*, Stu2 from *S. cerevisiae*, ZYG-9 from *C. elegans*, Mini spindles from *D. melanogaster* and a putative 2,021-amino-acid protein from *Arabidopsis thaliana* (GenBank accession number 4263790) (Fig. 1b).

Analysis of the XMAP215 amino-acid sequence showed that it contains two putative sites of phosphorylation by the kinase Cdc2. One is located in the 36-amino-acid insertion (position 1,154) and the other is at position 1,396. XMAP215 also contains two bipartite nuclear-localization signals (positions 544–561 and 556–573) and two HEAT repeats at positions 436–477 and 1,015–1,052 (ref. 24). Interestingly, the amino-acid motif KXGS, which is the target for the kinase MARK²⁵ and is found in the microtubule-binding domains of the MAPs Tau, MAP4 and MAP2B²⁶⁻²⁹, is present once in ch-TOG/KIAA0097 and three times in XMAP215 (Fig. 1a).

XMAP215 localizes to microtubules in interphase and mitosis. To attempt to understand the function of XMAP215 *in vivo*, we raised antibodies against the amino and carboxy termini of XMAP215 and KIAA0097/ch-TOG. In interphase and mitotic *Xenopus* egg extracts, the antibodies raised against both the N and the C terminus of ch-TOG recognized a single band with an M_r value of about 215K (Fig. 1c). Identical results were obtained with the anti-XMAP215 antibodies (data not shown). Thus, antibodies to both ch-TOG and XMAP215 recognize XMAP215 in *Xenopus* egg extracts.

We next determined the localization of XMAP215 in XL177 *Xenopus* tissue culture cells (Fig. 2). Cells were fixed and stained with antibodies against the N terminus of XMAP215. In interphase, XMAP215 co-localized with microtubules. During mitosis, the spindle and the centrosomes were brightly stained whereas astral microtubules were poorly labelled. Similar results were obtained with antibodies to the N terminus of ch-TOG. Thus, XMAP215 has an intracellular distribution that suggests an involvement in the control of microtubule dynamics.

XMAP215 regulates parameters of microtubule dynamics in extracts. To attempt to understand the role of XMAP215 in the control of microtubule dynamics, we used antibodies specific for the C terminus of ch-TOG to deplete *Xenopus* egg extracts. As shown in Fig. 3a, 30% of XMAP215 remained in the extract following depletion. Routinely, between 50% and 70% of XMAP215 was depleted from the extract. In both interphase and mitotic extracts, a single band of $M_r \sim 215$ K was precipitated by the antibody against the ch-TOG C terminus (Fig. 3b). Furthermore, the protein precipitated using antibodies directed against the C terminus of XMAP215 was recognized by anti-chTOG antibodies, showing that both antibodies recognized the same protein (data not shown). These observations indicate that the depletion was specific to XMAP215.

To assay microtubule dynamics, we observed individual microtubules by fluorescence video microscopy of extracts depleted of XMAP215. In both interphasic and mitotic extracts, the depletion of ~60% of XMAP215 was enough to result in much shorter and more dynamic microtubules (Fig. 4a). In interphase, the growth and the shrinkage rates decreased from 14.2 µm min⁻¹ to 8.0 µm min⁻¹ and from 21.7µm min⁻¹ to 11.6µm min⁻¹, respectively (Fig. 4b). This observation correlates well with what is known about the effect of XMAP215 on pure tubulin^{5,6}. To our surprise, we also observed an increase in the interphase catastrophe frequency, from 0.1 events min-1 to 0.7 events min-1 (similar to that observed in mitosis). In mitotic extracts, the effect was even more dramatic, as microtubules were often not observed during the first 15 min of recording in XMAP215-depleted extracts (5 min is normally long enough to record microtubule dynamics). The growth and shrinkage rates in mitotic extracts were not affected by XMAP215 deple-

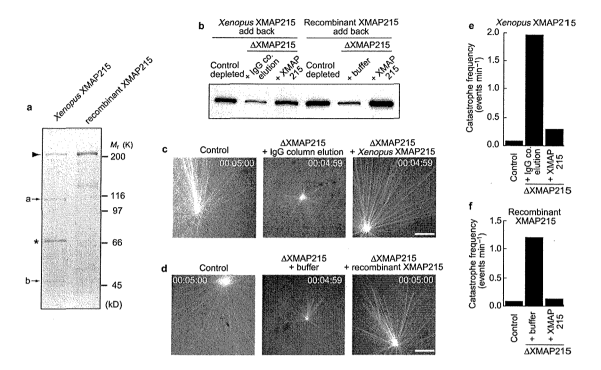


Figure 5 Readdition of purified XMAP215 to depleted extracts. a, Coomassieblue-stained SDS gel showing position of XMAP215. Left, *Xenopus* XMAP215 was purified on an affinity column of antibodies directed towards peptides from the XMAP215 C terminus. The asterisk shows the position of BSA contained in the elution buffer. The associated protein indicated by arrow 'b' is also seen in immuno-affinity purification with control IgG. Variable amounts of another protein, indicated by arrow 'a', were analysed by mass spectrometry and identified as vitellogenin-A2 precursor (Swiss-plot accession number P18709), a common contaminating protein. Right, recombinant XMAP215 was expressed in insect cells and purified on a Mono-S column. **b**, Immunoblot of immunodepleted extracts before and after adding back XMAP215. Interphase extracts were immunodepleted with control IgG (lanes 1, 4) or

tion, being respectively $11.5 \mu m \min^{-1}$ and $16.9 \mu m \min^{-1}$ in control depleted extracts and $10.6 \mu m \min^{-1}$ and $14.4 \mu m \min^{-1}$ in XMAP215-depleted extracts. However, the catastrophe frequency increased from 0.9 events min⁻¹ to 2.1 events min⁻¹ (Fig. 4b).

To confirm that the effects observed were due to XMAP215 removal, we added either immunopurified XMAP215 or XMAP215 produced in baculovirus-infected cells back to the depleted extracts (Fig. 5a). Restoration of the endogenous level of XMAP215 with either protein (Fig. 5b) allowed microtubules to grow long again (Fig. 5c, d), showing that XMAP215 is sufficient to rescue the observed phenotype resulting from the earlier depletion of XMAP215. We next analysed the parameters of microtubule dynamics in these conditions. The catastrophe frequency decreased dramatically to levels similar to that observed in control depleted extracts (Fig. 5e, f), showing that XMAP215 is a major regulator of catastrophes in Xenopus egg extracts. We also analysed the growth and shrinkage rates. Both parameters were increased to some extent. Nevertheless, this change was variable and never reached control levels (data not shown). A possible explanation for this discrepancy is that our purified proteins were not fully active in rescuing the phenotype of the growth and shrinkage rates. However, the main phenotypes characterized, such as the change in microtubule length and in the catastrophe frequency, are fully reverted by the addition of purified XMAP215. Together, these observations indicate that, first, XMAP215 is required for at least half of the growth and the shrinkage rates in interphase but does not contribute to

anti-ch-TOG antibody (lanes 2, 3, 5, 6). An elution fraction from the IgG column (lane 2), *Xenopus* XMAP215 (lane 3), control buffer (lane 5) or recombinant XMAP215 (lane 6) was then added to XMAP215-depleted extracts. **c**, **d**, Images of microtubule asters in control depleted extracts (left), and XMAP215-depleted extracts before (middle) and after (right) adding back immunopurified (**c**) or recombinant (**d**) XMAP215, respectively. Both purified proteins rescued the aster size of XMAP215-depleted extracts. Scale bar represents 10μ m. **e**, **f**, Effects of addition of immunopurified and recombinant XMAP215 on the catastrophe frequency in XMAP215-depleted extracts. Adding back purified proteins caused a 7–10-fold suppression of the catastrophe frequency observed in XMAP215-depleted extracts.

these parameters during mitosis; second, catastrophe factors are also active in interphase; and third, XMAP215 antagonizes the function of catastrophe factors both in interphase and in mitosis. XMAP215 and XKCM1 act antagonistically. We wanted to know which catastrophe factors oppose XMAP215 activity. Depletion of one such factor, Stathmin/Op18, decreases the catastrophe rate about twofold in mitosis³⁰. To assess the degree to which another catastrophe factor, XKCM1, was responsible for the catastrophe rate in mitosis, we added inhibitory anti-XKCM1 antibodies¹⁰ to mitotic extracts and determined the parameters of microtubule dynamics (Fig. 6b). After addition of anti-XKCM1 antibodies to mitotic extracts, the catastrophe frequency declined sevenfold. We conclude that XKCM1 is the major factor regulating the catastrophe rate in mitotic extracts. To determine whether XKCM1 was also responsible for catastrophes in interphase, we studied microtubule dynamics after addition of anti-XKCM1 inhibitory antibodies to interphase and mitotic extracts depleted of XMAP215. This procedure resulted in an increased length of the microtubules in both interphase and mitotic extracts (Fig. 6a). We then quantified the different parameters of microtubule dynamics. In four different depletions of XMAP215 from interphase and mitotic extracts, we observed an increase in the catastrophe frequency, followed by a decrease after inhibition of XKCM1. Values from two similar experiments are shown in Fig. 6b. The catastrophe frequency in an interphase extract increased from 0.1 events per minute to 0.7 events min⁻¹ after depletion of XMAP215, and was reduced to 0.2

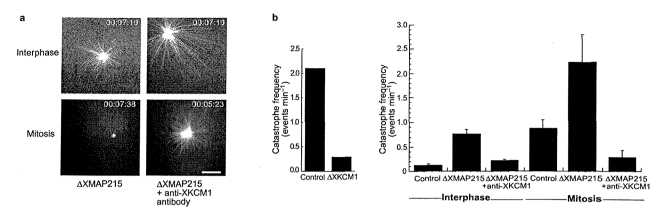


Figure 6 Inhibition of XKCM1 partially rescues the effect of XMAP215 depletion on the catastrophe frequency. a, Images of microtubule asters in XMAP215-depleted extracts before and after inhibition of XKCM1. Microtubules grow longer after XKCM1 inhibition. b, Left, change in catastrophe frequency after inhibition of XKCM1 in mitotic extracts. Right, quantification of microtubule dynamics

in XMAP215-depleted extracts after XKCM1 inhibition. Only a change in the catastrophe frequency was observed; the growth and shrinkage rates were similar to those shown in Fig. 4 and are not shown here. Inhibition of XKCM1 decreases the catastrophe frequency in both interphase and mitotic extracts. Data are shown as means \pm s.e.m.

events min⁻¹ after addition of anti-XKCM1 antibodies. In a mitotic extract depleted of XMAP215, addition of anti-XKCM1 antibodies decreased the catastrophe frequency from 2.2 events min⁻¹ to 0.3 events min⁻¹. We therefore conclude that XMAP215 antagonizes the activity of XKCM1, both in interphase and in mitosis.

XMAP215 is required for spindle assembly. As changes in microtubule dynamics are essential for spindle assembly, and because XMAP215 is a major regulator of microtubule dynamics, we wondered how removal of XMAP215 would affect formation of the mitotic spindle. We added sperm nuclei to mitotic, XMAP215depleted extracts and monitored the formation of bipolar spindles 20 min and 45 min after addition of sperm nuclei. At 20 min, we observed short microtubule asters around chromatin in XMAP215depleted extracts. The length of microtubules depended on the

 20 min
 ΔΧΜΑΡ215

 45 min
 Δ

Figure 7 Spindle formation observed in control and XMAP215-depleted extracts. Microtubule arrays are seen in control (left) and XMAP215-depleted (right) extracts 20 and 45 min after addition of sperm nuclei. Spindles are seen 45 min after sperm addition. Scale bar represents $10 \mu m$.

depletion efficiency. When the depletion was close to 90%, no microtubules were seen growing from the centrosome. In extracts from which 60% of XMAP215 was removed, microtubules were ~10 μ m long (Fig. 7) compared with ~25- μ m-long control microtubules. 45 min after addition of sperm nuclei to control extracts, normal bipolar spindles of width 31.8±4.9 μ m (n=32) were visible. In XMAP215-depleted extracts, short spindle-like structures of width 18.3±3.9 μ m (n=33), containing a few short microtubules and monopolar spindles, were formed (Fig. 7). With higher depletion efficiency, we observed either very short or no microtubules nucleated by the sperm centrosome. We therefore concluded that XMAP215 is required for efficient spindle assembly.

Discussion

XMAP215 is a member of a family of proteins that have been conserved in yeasts, C. elegans, Drosophila, plants and humans. The main regions of homologies representing functional conservation are in their N termini^{16,17}. Mutations in the *C. elegans zyg-9* gene¹⁷ have effects on microtubule organization that are similar to those produced by depletion of XMAP215 from Xenopus egg extracts: microtubules are short and spindle assembly is compromised. We have shown that, in interphase, XMAP215 localizes along the length of microtubules and concentrates at centrosomes. This localization is similar to that of p93^{Dis1}, the S. pombe XMAP215 homologue¹⁹ but different from that of ZYG-9 (ref. 17) and Mini spindle proteins¹⁶, which localize primarily to centrosomes during both interphase and mitosis. It is possible that XMAP215 has more than one function in regulating the microtubule network. It may regulate microtubule stability on the one hand and microtubule nucleation on the other, an idea supported by analysis of the role of XMAP215 in Ran-GTP-dependent microtubule nucleation³

To determine the function of XMAP215 in the context of other regulators of microtubule dynamics, we depleted the protein from *Xenopus* egg extracts. After depletion of XMAP215 from interphase extracts, the average growth rate of microtubules dropped from 14.2 μ m min⁻¹ to 8.0 μ m min⁻¹. This result supports the previous work of Gard and Kirschner¹⁴, who showed that XMAP215 is the major activity stimulating microtubule growth that can be purified from a *Xenopus* egg extract. At the concentration of tubulin present in the extracts (20 μ M), microtubules are expected to grow at a rate of 2.0 μ m min⁻¹ if no other factors are present¹². Thus the residual growth rate of 6.0 μ m min⁻¹ indicates that factors that can stimulate microtubule growth are still present following XMAP215 depletion.

One possibility is that, as we depleted only about 60% of the protein in our experiments, the residual XMAP215 stimulates growth. Alternatively, other factors, such as XMAP230 (refs 7, 32), can stimulate growth in extracts.

Perhaps the most striking effect of XMAP215 depletion from interphase extracts is the dramatic increase in the catastrophe frequency. At first glance, the activity of XMAP215 appears to be different in different circumstances. On addition to purified tubulin, XMAP215 stimulates the growth and shrinkage rates without suppressing catastrophes^{5,6}. In this system, catastrophes are rare and are due to the inherent properties of the tubulin polymer. Clearly, XMAP215 cannot suppress these types of catastrophe. However, Xenopus in egg extracts, XMAP215 both stimulates growth and shrinkage rates and suppresses catastrophes by opposing the activity of XKCM1. It is possible that XKCM1 and XMAP215 compete directly for microtubule ends. Studies of the activity of XKCM1 have shown that it acts on microtubule ends to stimulate microtubule depolymerization¹². Although there is no direct evidence that XMAP215 binds to microtubule ends, early studies of XMAP215 showed that it could stimulate microtubule growth at substoichiometric levels and specifically at plus ends^{5,6,14}. The idea that proteins bind to microtubule ends to modulate their dynamics is attractive and is further supported by work with CLIP-170, which localizes to growing microtubule ends³³⁻³⁵, and by the ability of kinetochores to distinguish ends of microtubules from the lattice³⁶

These results have important consequences for understanding the control of microtubule turnover between interphase and mitosis. Our observations show that XKCM1 is active in mitosis as well as in interphase. However, XKCM1 is prevented from stimulating catastrophes by the action of XMAP215, mainly in interphase. During mitosis, XMAP215 is phosphorylated and less able to promote microtubule growth, although it is still able to bind to microtubules^{5,6,14}. Thus, one likely model that can explain the increase in turnover of microtubules between interphase and mitosis is that XMAP215 promotes microtubule growth in interphase, thus exerting a dominant microtubule-stabilizing activity over the microtubule-destabilizing activity of XKCM1. However, in mitosis, it does not stimulate microtubule growth, allowing catastrophe factors to act more effectively. This hypothesis is supported by the observation that depleting XMAP215 from interphase extracts reduces the growth rate of microtubules to that in mitotic extracts. The finding that immunodepletion of XMAP215 from mitotic extracts leads to an increase in catastrophe frequency in the absence of a change in growth rate is also consistent with this idea. This further indicates that, in mitosis, by binding to microtubules, XMAP215 can still oppose XKCM1 activity to a certain extent. Thus, a precise adjustment of microtubule turnover could be achieved by subtle regulation of the activity of XMAP215.

Although we are beginning to understand how microtubule dynamics are regulated during the cell cycle, we still know little about how this property is modulated to assemble a bipolar spindle. It is thus interesting that depleting XMAP215 from mitotic extracts results in a shorter steady-state length of microtubules. The consequence of this change in microtubule dynamics is that small spindles or no spindles at all form. Similar effects on the relationship between spindle size and microtubule dynamics were seen in previous experiments in which the catastrophe rate was increased by adding Stathmin/Op18 (ref. 37). Thus, the results described here and in ref. 37 indicate that spindle length is determined in large part by the dynamic properties of microtubules, and may be achieved mainly by regulating the frequency of catastrophes, as predicted by a mathematical model that links microtubule dynamics to their average steady-state length³⁸.

Methods Cloning.

Mass spectroscopy results (A. Podtelezhnikov et al., personal communication) indicated that XMAP215 was highly homologous to the human protein ch-TOG¹⁵. We therefore screened a X. laevis oocyte cDNA library (courtesy of J. Shuttleworth) using a clone coding for KIAA0097, obtained from T. Nagase (GenBank accession number D43948). Positive clones contained either a 4.7-kilobase (kb) 3'-terminal part of a gene homologous to *ch*-TOG, or a 2-kb 5'-terminal part that partially overlap ped with the above 4.7-kb clone but lacked the translation initiation codon. The most 5' end of the gene was recovered using 5' RACE (rapid amplification of cloned ends)³⁰ with semi-nested primers (primer 1: 5'-GCCATTGAC-CACTTC; primer 2: 5'-AATTCCTGCCACGTGAGCAT). Southern hybridization with the *ch*-TOG probe highlighted a band of -400 base pairs (bp), which was subcloned and sequenced to reveal high homology to the corresponding 5' terminus of the *ch*-TOG gene. This 381-bp fragment was used to reconstitute the 5' end of the gene on the 2-kb cDNA clone. Next, a single nucleotide replacement was introduced by five cycles of *Pli*-PCR (polymerase chain reaction) into the 28-bp region of the overlap between two cDNAs. This mutation was silent and created a unique *Dra*l restriction site. Finally, to assemble the whole *XMAP215* gene, we carried out a three-way ligation between two cDNA clones (2.1 kb and 4.7kb, cut out using *Ncal+Dral* and *Dral+Awr*II, respectively) and pGEM-T (Promega), linearized with *Ncal+Spel*.

Indirect immunofluorescence microscopy.

XL177 cells were grown on 12-mm coverslips and fixed in 3–4% paraformaldehyde and 0.1% glutaraldehyde plus 0.5%Triton X-100 in BRB80 buffer (80mM PIPES, 1 mM MgCl., 1 mM EGTA, pH 6.8). Free aldehydes were quenched by incubation in 0.1% NABH, in PBS. Cells were blocked in PBS plus 0.2% Tween-20 supplemented with 10% fetal calf serum and then incubated with a 1:200 dilution of affinity-purified rabbit antibody towards the N terminus of XMAP215 and a 1:200 dilution of mouse monoclonal anti-α-tubulin antibodies (Amersham). Anti-mouse IgG coupled to Alexa-488 (Molecular Probes; 1:1,000 dilution) and anti-rabbit IgG coupled to Cy3 (Amersham; 1:500 dilution) were used as secondary antibodies. Stained cells were preincubated in 2µgml⁻¹ Hoechst 3352 to stain the DNA, washed and mounted in the ProLong mounting medium (Molecular Probes). Analysis was done with an LSM 510 confocal microscope (Zeiss). The best localization was obtained using the antibodies towards in interphase mas observed in interphase microtubule asters assembled in *Xenopus* egg extracts (A. P., unpublished observations).

Antibody production.

The first 214 and the last 328 amino acids of KIA0097/ch-TOG were fused to glutathione-S-transferase (GST) in the pGEX-4T vector (Pharmacia) and the recombinant proteins were purified and used to obtain rabbit polyclonal antibodies. Antibodies were affinity-purified according to standard protocols⁴⁰. To remove antibodies towards GST, we passed the affinity-purified antibodies over a GST column. Antibodies to XMAP215 were raised against the first 573 and the last 899 amino acids, fused to a His, tag, and affinity-purified against the same fusion proteins.

Preparation of Xenopus egg extracts and depletion of XMAP215.

Xenopus egg extracts were prepared as described, and spindle assembly and microtubule dynamics were assayed as described^{364,12}. Depletion of XMAP215 and treatment of the immunoprecipitates were done as in ref. 10 using 200µg anti-ch-TOG antibody coupled to 50µl Biorad Affi-Prep beads. Depleted extracts were used to assay spindle assembly or microtubule dynamics. XKCM1 inhibition was achieved using antibodies raised against the N-terminal part of XKCM1 (ref. 10). Antibodies were added to the extract at a final concentration of 170µgml⁻¹ before recording microtubule dynamics. Chromopure normal rabbit IgG (Dianova) was used as a control antibody in amounts equal to that of anti-TOG antibodies. To estimate the depletion efficiency of XMAP215, we loaded 1µl, 0.3µl and 0.1µl of control depleted extracts, and 1µl of XMAP215-depleted extracts, onto SDS gel. The extent of the depletion was deduced from the relative intensity of the XMAP215 band, revealed by immunoblotting using 0.1µgml⁻¹ of antibody to the ch-TOG C terminus.

To confirm the specificity of the ch-TOG antibodies, we performed a depletion using antibodies directed against the C terminus of XMAP215. These antibodies also retrieved a single protein of M, 215K, as observed by Coomassie blue staining (data not shown). Moreover, the anti-ch-TOG antibodies recognized the 215K band immunoprecipitated by the anti-XMAP215 antibody, as determined by immunoblotting (data not shown).

Microtubule dynamics was recorded as described[®] with the following modifications. Recording was normally arbitrarily stopped after 5 min. However, in XMAP215-depleted extracts it was necessary to record for up to 15 min. In one XMAP215-depleted extract in which the depletion efficiency was 90%, we were unable to observe any microtubules even after 30 min of observation. We analysed 128, 106, 91 and 74 microtubules, undergoing 19, 79, 61 and 76 catastrophes, respectively, for the interphase control depleted, interphase XMAP215-depleted, mitotic control depleted and mitotic XMAP215-depleted extracts, respectively.

Purification of XMAP215 and add-back experiments.

Xenopus XMAP215 was purified by immunoaffinity chromatography as described¹⁰. Antibodies towards a peptide from the XMAP215 C terminus were generated against the synthetic peptide (C)DDLKKRLERIKSSRK (synthesized by Genosis, UK), corresponding to the last 15 amino acids of XMAP215. 100µl Affi-Prep beads (Biorad) were coated with 50µg affinity-purified anti-C-terminalpeptide antibody by incubation for 1 h at 4°C. Beads were then washed twice with PBS-T (PBS+0.1% Triton X-100) and then three times with XB-PIs (XB buffer containing 10µgml⁻¹ each of leupeptin, peptida and chymostatin). After removing as much buffer as possible, we added 5 ml interphase egg extract to the beads and rotated them for 1 h at 4°C. The beads were washed three times with XB-PIs and poured into a column. To elute XMAP215, we added an equal volume of XB-PIs containing 400 mM KCI and 1.2 mM XMAP215 C-terminal peptides to the resin. After elution, 0.01% BSA was added as a carrier protein. In controls we used normal rabbit IgG (Dianova), and we treated the controls as above.

To produce recombinant XMAP215, we used the Bac-to-bac baculovirus expression system according to the manufacturer's instructions (Gibco). The full-length *XMAP215* gene was cloned into a pFastBac vector. Hi5 insect cells were then infected to express the recombinant protein. Cells were collected and resuspended in lysis buffer (20mM cation buffer, pH 7.5 (consisting of 6.7 mM each of sodium acetate, MES and HEPES) containing 5% glycerol, 50mM NaCl, 10µgml⁻¹ protease inhibitors (leupeptin, pepstatin and chymostatin), 0.1% Triton X-100, 10mM CaCl, and 10µgml⁻¹ nocodazole). Cells were broken using a Dounce homogenizer with 25 strokes and put on ice for 20min. After adding 1 mM dithiothreitol, we centrifuged cell extracts for 10min at 4°C at 90,000 r.p.m. on a Beckman TLA100.2 rotor. The supernatiant was loaded onto a Mono-S column (Pharmacia) and washed with lysis buffer containing 100 mM NaCl. XMAP215 was eluted with a 100–600 mM NaCl gradient in lysis buffer. XMAP215 eluted at ~400 mM NaCl. The peak fractions were pooled and concentrated 50-fold by Microcon-30 concentrators (Amicon).

In add-back experiments, a 1/10 volume of purified proteins was added to XMAP215-depleted interphase extracts before recording microtubule dynamics. Elution fractions of the IgG immunoaffinity column and lysis buffer were used as controls for add-back of *Xenopus* XMAP215 and recombinant XMAP215, respectively.

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