Structural Transitions at Microtubule Ends Correlate with Their Dynamic Properties in *Xenopus* Egg Extracts

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**Abstract.** Microtubules are dynamically unstable polymers that interconvert stochastically between growing and shrinking states by the addition and loss of subunits from their ends. However, there is little experimental data on the relationship between microtubule end structure and the regulation of dynamic instability. To investigate this relationship, we have modulated dynamic instability in *Xenopus* egg extracts by adding a catastrophe-promoting factor, Op18/stathmin. Using electron cryomicroscopy, we find that microtubules in cytoplasmic extracts grow by the extension of a two-dimensional sheet of protofilaments, which later closes into a tube. Increasing the catastrophe frequency by the addition of Op18/stathmin decreases both the length and frequency of the occurrence of sheets and increases the number of frayed ends. Interestingly, we also find that more dynamic populations contain more blunt ends, suggesting that these are a metastable intermediate between shrinking and growing microtubules. Our results demonstrate for the first time that microtubule assembly in physiological conditions is a two-dimensional process, and they suggest that the two-dimensional sheets stabilize microtubules against catastrophes. We present a model in which the frequency of catastrophes is directly correlated with the structural state of microtubule ends.

**Key words:** microtubules • dynamic instability • *Xenopus* egg extracts • electron cryomicroscopy • protofilament sheets

**Introduction**

Microtubules are dynamic polymers that switch stochastically and infrequently between growing and shrinking states (Walker et al., 1988). This unusual behavior, called dynamic instability (Mitchison and Kirschner, 1984; Horio and Hotani, 1986), allows rapid spatial changes of the microtubule cytoskeleton during the cell cycle. A particularly striking example of such a rearrangement is the dramatic reorganization of microtubules during the interphase-mitosis transition (Hyman and Karsenti, 1996).

Many studies have been performed with pure tubulin to investigate the basic mechanism underlying dynamic instability. Microtubules elongate by the addition of tubulin dimers, which rapidly hydrolyze one of their two bound GTP molecules (Carlier, 1989). The energy coming from tubulin-GTP hydrolysis is essential to destabilize the microtubule lattice and allow its fast depolymerization (Hyman et al., 1992). For many years, the most popular model proposed that growing microtubules are stabilized by a terminal cap of unhydrolyzed GTP subunits (for review see Erickson and O’Brien, 1992), the loss of which would result in a sudden change between growing and shrinking states (termed a catastrophe). However, no GTP-tubulin has been detected at the present in the lattice of dynamic microtubules, and the GTP cap model remains controversial.

More recently, structural approaches using EM analysis of pure tubulin polymerization have shown that the regulation of both microtubule assembly and dynamics involves changes in their end structure. Two-dimensional sheets of tubulin are observed at the end of growing microtubules, whereas shrinking microtubules display curved protofilaments peeling out from their ends (Erickson, 1974; Kirschner et al., 1974, 1975; Simon and Salmon, 1990; Mandelkow et al., 1991; Chrétien et al., 1995; Tran et
and buffer A with 10 mM imidazole, the bound protein was eluted with the supernatant was incubated with a metal affinity resin (Talon™ IMAC lysis.

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Recombinant Op18/stathmin with a 6-histidine tag was purified from molecules with dynamics, it is important to look at a population of mi-

crotubules undergoing dynamic instability. In a population of microtubules growing in vitro, there are very few catastro-

phases, making it difficult to correlate growing and shrinking microtubules with their end structure (Chrétien et al., 1995). In vivo, microtubules are much more dynamic (Sammak and Borisy, 1988; Belmont et al., 1990; Simon et al., 1992), but to date no studies of microtubule end structure have been performed under physiological conditions.

To investigate the structural basis of dynamic instability under physiological conditions, we analyzed microtubule end structure and dynamics in X. laevis egg extracts. We find that physiological microtubule assembly occurs by the growth of two-dimensional sheets of tubulin, which later close into a tube. To correlate potential changes in end structures with dynamics, we increased the catastrophe frequency by adding the destabilizing factor Op18/stathmin (Belmont and Mitchison, 1996) to extracts. The results show that the increase in the catastrophe frequency induced by Op18/stathmin is accompanied by a decrease in both the length and proportion of the sheets and a concomitant increase in blunt and frayed ends. These results allow us to propose a structural model to explain dynamic instability and its possible relationship with GTP hydro-

Materials and Methods

Purification of Recombinant Op18/Stathmin

R recombinant Op18/stathmin with a 6-histidine tag was purified from E. coli as follows. 5 h after induction by 0.2 mM isopropyl-

β-β-thiogalactopyranoside at 37°C, the cells were pelleted by centrifuga-

tion at 4°C and resuspended in buffer A (20 mM TRIS and 100 mM NaCl, pH 6.8) supplemented with PM SF (1 mM) and protease inhibitor (leupep-

tin, pepstatin, and aprotinin, 100 μg/ml). The cells were lysed using the French Press, the extract was clarified at 17,000 rpm for 30 min at 4°C, and the supernatant was incubated with a metal affinity resin (Talon™ IMAC C R esin; CLONTECH) for 1 h at 4°C. A filter washing the resin with buffer A and buffer A with 10 mM imidazole, the bound protein was eluted with 100 mM imidazole and dialyzed against BRB 80 buffer (80 mM potassium-

Pipes, 1 mM MgCl2, and 1 mM EGTA, pH 6.8). Protein concentration was measured using a Bradford assay with BSA as a reference (calculated mo-

lecular mass of recombinant Op18/stathmin, 18,359 kD).

Preparation of Extracts

CSF X. laevis extracts (arrested in metaphase II of meiosis) were pre-

pared as described before (Murray, 1991) and cycled into interphase by addition of 0.4 mM calcium and 200 μg/ml cycloheximide and incubation for 40 min at room temperature.

Video Microscopy and Data Analysis

Microtubule dynamics were measured in 10,000 g frog egg extracts supple-

mented with human purified centrosomes (2 × 10⁶ centrosomes/ml; Bor-

nens et al., 1987), rhodamine tubulin (Hymans et al., 1990), saturated he-

moglobin, and anti-F-actin solution as described before (Tournebize et al., 1997). Dynamics were recorded using a Zeiss a xioskop, a 100× A pochro-
mate lens (NA 1.4), and a rhodamine filter cube. Images were recorded every 4 s on an 8-bit black and white camera (Sony SSC-M 370CE) with an image processor (model A rgs 10; H amamatsu) and stored on a M acin-

tosh using NIH -Image software. Data analysis was done with a M icrosoft E xcel Macro to determine the growth rate, the shrinkage rate, and the cata-
tastrophe frequency of microtubules (Tournebize et al., 1997). For the ex-

periments with Op18/stathmin, extracts supplemented with 3.6, or 7.5 μM of the recombinant protein were similarly analyzed. Four experiments (0.4 μM Op18/stathmin) and two experiments (3, 6, and 7.5 μM Op18/stath-

min), which were made in different extracts, were averaged. Differences between experiments with and without Op18/stathmin were evaluated us-

ing a t test with a confidence level of 95%. The percentage of time spent in the growing and shrinking states was determined by dividing the time re-

corded in growing (or shrinking) state by the total time recorded.

Specimen Preparation and Electron Cryomicroscopy

Vitreous ice-embedded samples were prepared as reported previously (Dubochet et al., 1985). 15 μl of interphase extract were mixed on ice with 5 μl of purified human centrosomes (2 × 10⁶ centrosomes/ml). 2-μl samples were pipetted onto a holey carbon grid and incubated for 20 s in a humid atmosphere at 25–26°C (Chrétien et al., 1992). The droplet was blotted and plunged quickly into liquid ethane. For the experiments with Op18/stathmin, we followed the same procedure using interphase extracts mixed on ice with centrosomes and different amounts of recombinant Op18/stathmin. The assembly time on the holey carbon grid was the same (20 s) except with 7.5 μM Op18/stathmin, for which the sample was incu-

bated for 1 min (because of the shorter microtubules obtained in this case; see Results). Specimens were stored in liquid nitrogen and observed in a Philips CM 20 operating at 160 kV. Images were recorded under low dose conditions at a magnification of 27,500 and ∼2.5 μm underfocused.

Image Analysis

Micrographs were printed and microtubule end structures were analyzed directly on these prints. Three types of extremities were observed: (1) blunt ends (straight protofilaments or very short protofilament protrusions <30 nm); (2) extensions between 30 nm and 1.5 μm; and (3) spread out ends (separated protofilaments). The percentage of each type of microtu-

bule ends were calculated with respect to the total number of extremities that we could identify (235 with 0.2 μM Op18/stathmin, 84 with 3 μM Op18/

stathmin, 63 with 6 μM Op18/stathmin, and 90 with 7.5 μM Op18/stath-

min). A lot of extremities could not be classified because of the high back-

ground of granules and vesicles, which often hide tips of microtubules. E xtension lengths were measured on the prints. We used the contrast present on microtubule images to localize the beginning of the extension, i.e., the point where the constant contrast and width of the complete cylinder is interrupted because of the apparition of one-sided curved extensions (Chrétien et al., 1995). The contrast of complete microtubules observed by electron cryomicroscopy was also used to determine the protofilament number (Chrétien and Wade, 1991).

Results

Two mechanisms have been proposed for the elongation of microtubules. O ne involves the helical addition of sub-

units to the tip of a growing microtubule (Chen and Hill, 1985; B aley et al., 1990; A mos, 1995), whereas the other proposes the formation of two-dimensional protofilament sheets, which subsequently close into a tube (E rickson, 1974; Kirschner et al., 1975; Simon and Salmon, 1990; Chrétien et al., 1995). To determine the structural nature of microtubule ends elongating under physiological condi-
tions, we examined microtubules nucleated off centrosomes in interphasic Xenopus egg extracts. Extracts supplemented with purified human centrosomes were frozen in liquid ethane after 20 s at room temperature, and the frozen specimens were observed by electron cryomicroscopy. As shown in Fig. 1 a, the microtubules of the centrosomal aster are visible despite the high background from vesicles and granules present in these dense cytoplasm.

Figure 1. (a) Vitreous ice-embedded microtubules observed in interphasic extracts supplemented with isolated centrosomes and incubated for 20 s at 25°C. E, extensions observed at microtubule ends; and C, centriole. (b and c) Vitreous ice-embedded microtubules observed in interphasic extracts in the presence of 7.5 μM recombinant Op18/stathmin. Addition of Op18/stathmin in this condition induces the disappearance of sheets at microtubule ends (black arrows). F, frayed end; and B, blunt end. Bar, 200 nm.
mic extracts. All microtubules observed have 13 protofila-
ments. Some of the microtubules show long extensions at
their plus ends, which are easily recognized because they
are curved and only one edge is visible, unlike the two par-
allel dark edges of the complete tube. These end structures
are similar to those previously observed at the ends of mi-
crotubules assembled off centrosomes using pure tubulin
and represent two-dimensional sheets of tubulin protofila-
ments (Chrétien et al., 1995).

Fig. 2 shows more detailed views of different end struc-
tures observed in extracts: microtubule extremities vary
from blunt ends (Fig. 2 b) to extensions of variable lengths
(Fig. 2, c–e). The average length is ~440 nm with a distri-
bution similar to the one described in vitro (Chrétien et
al., 1995). Microtubules with frayed ends (split protofila-
ments) are also observed in interphase extracts (Fig. 2 a).
Microtubules with such end structures are most likely in a
shrinking state (discussed further below; see Fig. 4). No
microtubule ends were observed with tubulin oligomers,
rings, or long curled protofilaments in extracts. These

Figure 2. Detailed views of microtubule end structures in interphasic extracts. (a) Microtubule end with peeling protofilaments (frayed end). (b) Blunt end. (c–e) Extensions with variable lengths. Bar, 100 nm.
types of structures were mostly reported at the end of microtubules induced to shrink rapidly by a high calcium or magnesium concentrations or by cooling the sample (Kirschner et al., 1974; Simon and Salmon, 1990; Mandelkow et al., 1991; Tran et al., 1997a; Müller-Reichert et al., 1998).

These results establish that under physiological conditions, microtubules grow by elongation of a two-dimensional sheet that closes into a tube as previously proposed on the basis of experiments performed with pure tubulin (Erickson, 1974; Kirschner et al., 1975; Simon and Salmon, 1990; Chrétien et al., 1995).

To analyze the relationship between the structure of a microtubule end and its dynamic nature, we manipulated microtubule dynamics and assessed the consequences on end structure. Xenopus extracts are an excellent system to pursue such an investigation because the growing rate and catastrophe frequency can be independently manipulated by the exogenous addition of regulatory factors. To vary the catastrophe frequency, we added Op18/stathmin, which increases the catastrophe frequency of pure microtubules (Belmont and Mitchison, 1996). The addition of Op18/stathmin reduces the average length of microtubules nucleated off centrosomes in interphase extracts in a dose-dependent manner (Fig. 3 a). Specifically, the addition of 3 μM Op18/stathmin does not change any of the dynamic properties of microtubules (Fig. 3 b, Table I). In contrast,

| Op18/stathmin (μM) | Growth rate (μm/min) | Shrinkage rate (μm/min) | Catastrophe frequency (events/min) |
|--------------------|----------------------|------------------------|-----------------------------------|
| 0                  | 11 ± 4 (95)          | 11.51 ± 4.9 (29)       | 0.34 ± 0.05 (34)                  |
| 3                  | 10.87 ± 3.44 (48)    | 13.3 ± 6.4 (19)        | 0.33 ± 0.07 (20)                  |
| 6                  | 9.57 ± 3 (57)        | 12.94 ± 5.68 (38)      | 0.69 ± 0.09 (48)                  |
| 7.5                | 7.57 ± 2.7 (59)      | 10.0 ± 5.4 (49)        | 1.43 ± 0.17 (66)                  |

Values represent the mean value ± SD of the mean (number of microtubules for the growth and shrinkage rate, number of events for the catastrophe frequency). The rescue frequency is not reported because the number of events recorded was too low in most of the conditions.
The addition of 6 μM Op18/stathmin and 7.5 μM Op18/stathmin increases the catastrophe rate twofold and fourfold, respectively. The shrinkage rate is not significantly different from unperturbed extracts at either 6 or 7.5 μM Op18/stathmin, whereas the growth rate is slightly decreased for the highest concentration used. We analyzed microtubule end structure by electron cryomicroscopy in the presence of 0, 3, 6, and 7.5 μM Op18/stathmin (Figs. 1 b and 4 a). Increasing the Op18/stathmin concentration reduces the proportion of sheets and increases the proportion of blunt and frayed microtubule ends (Fig. 4 a). The proportion of blunt ends increases transiently for intermediate concentrations of Op18/stathmin (3 and 6 μM) and decreases as more microtubules exhibit frayed ends. This result suggests that blunt ends are more unstable compared with the ends with sheets at their extremities. Blunt ends could either grow or transit to shrinkage with a certain probability.
ity, whereas all microtubules with visible sheets at their ends are in a stable growing state. In apparent contradiction with this idea, at 3 μM Op18/stathmin, the proportion of blunt ends is greater than at 0 μM, but there is no significant increase in the catastrophe frequency. One possibility is that the probability of blunt ends to transit to the shrinking state is different at 3 and 6 μM Op18/stathmin despite their similar proportion. This is reflected in the greater number of frayed ends at 6 μM Op18/stathmin. A nother possibility is that we may have classified as blunt ends some very short sheets made of a few protofilaments (as Op18/stathmin decreases the sheet length). In this case, a structural transition would occur between 3 and 6 μM Op18/stathmin, allowing the complete closure of the tube and increasing the probability of a shrinking event.

**Discussion**

Our results show that under physiological conditions, all microtubules with visible sheets at their ends are in the growing state and all microtubules with frayed ends are in the shrinking state. These observations conclusively demonstrate that physiological microtubule assembly occurs by the extension of two-dimensional protofilament sheets at microtubule ends and not by helical polymerization. The structure of depolymerizing ends observed in extracts suggests that under normal conditions of dynamic instability, microtubules shrink by loss of protofilaments that peel away from the tube and dissolve almost immediately into subunits. It seems likely that the rams horns, which are seen after microtubules are forced into shrinking by calcium or magnesium, are caused by stabilization of these structures by the agents used to trigger shrinking (Tran et al., 1997a). Less certain is the dynamic nature of microtubules with blunt ends. A possibility that we favor is that blunt ends represent a structural state that is a metastable intermediate between the growing and shrinking states (Tran et al., 1997b). Thus, microtubules with blunt ends may either grow or transit to shrinking with a certain probability. A similar metastable state has been postulated on the basis of kinetic studies of severed MT ends (Tran et al., 1997b).

One interesting question that arises from this study concerns the mechanism by which Op18/stathmin induces the disappearance of tubulin sheets at the end of growing microtubules and why the elimination of the sheets tends to result in an increase in the catastrophe frequency (as suggested in Fig. 4 a). Op18/stathmin could either prevent the elongation of sheets or increase the rate of tube closure. Each model has different predictions concerning the molecular mechanisms involved. The first model would imply that the addition of tubulin subunits at microtubule ends is slowed down by the presence of Op18/stathmin because of the sequestering of free tubulin dimers (Howell et al., 1999). The second model would predict that Op18/stathmin binds to the protofilaments and favors the curvature leading to tube formation over the outward curvature of the sheet (for review see Chrétien et al., 1999). Because Op18/stathmin does not significantly affect the rate of microtubule elongation under conditions in which catastrophe increases (specifically at 6 μM Op18/stathmin), and because of the evidence suggesting that it can act directly at microtubule ends (Howell et al., 1999; Larsson et al., 1999), we favor the second model. However, the tubulin-sequestering activity of Op18/stathmin could also contribute to the increase in the catastrophe frequency at a higher concentration (7.5 μM) since we observed a slight decrease in the growth rate in this condition.

Why should the elimination of sheets make microtubules more prone to undergo a catastrophe? The tubulin dimer has an intrinsic curvature that causes tubulin protofilaments to have an outward curvature (Howard and Timasheff, 1986; M elki et al., 1989; Mandelkow et al., 1991; Hyman et al., 1995; Tran et al., 1997a; Müller-Richert et al., 1998). When 13 protofilaments associate laterally and close into a tube, their outward curvature puts the microtubule wall under tension. Sheets at the end of a microtubule, therefore, make microtubules stable because they introduce a cap of relaxed interactions between protofilaments at the end of the mechanically strained tube polymer (Chréti et al., 1995, 1999). Loss of the sheet as a consequence of tube closure catching up with sheet extension generates a blunt end that may either transit to shrinking or continue growing. What may influence the probability with which a blunt end transits to a shrinking phase? It seems likely to be associated with GTP-tubulin hydrolysis (Fig. 5). Only a few GTP-tubulin subunits are required to stabilize the ends of microtubules (Drechsel and Kirschner, 1994; Caplow and Shanks, 1996). We speculate that when a blunt end is generated by tube closure catching up with the sheet extension, the chemical composition of tubulin subunits at the blunt end will influence whether the end will transit to shrinking or continue growing. If GTP-tubulin subunits are at the blunt end, then the microtubule is more likely to grow. In contrast, if GDP-tubulin subunits are present, then the microtubule is more likely to transit to the shrinking state. Interestingly, it was suggested that Op18/stathmin triggers catastrophes by stimulating GTP hydrolysis (Larsson et al., 1999; Howell et al., 1999). This could also increase the probability of a blunt end to transit to the shrinking state in our model (Fig. 5). Nevertheless, such a mechanism does not explain the structural changes observed at the end of microtubules in the presence of Op18/stathmin, i.e., the disappearance of two-dimensional tubulin sheets. A definitive resolution of this issue will require the development of techniques capable of directly assessing the chemical nature of tubulin subunits in the end structures observed by cryo-E M.

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