Microtubules in the fungal pathogen \textit{Ustilago maydis} are highly dynamic and determine cell polarity

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**SUMMARY**

Many fungal pathogens undergo a yeast-hyphal transition during their pathogenic development that requires rearrangement of the cytoskeleton, followed by directed membrane traffic towards the growth region. The role of microtubules and their dynamic behavior during this process is not well understood. Here we set out to elucidate the organization, cellular role and in vivo dynamics of microtubules in the dimorphic phytopathogen \textit{Ustilago maydis}. Hyphae and unbudded yeast-like cells of \textit{U. maydis} contain bundles of spindle pole body-independent microtubules. At the onset of bud formation two spherical tubulin structures focus microtubules towards the growth region, suggesting that they support polar growth in G2, while spindle pole body-nucleated astral microtubules participate in nuclear migration in M and early G1. Conditional mutants of an essential \(\alpha\)-tubulin gene from \textit{U. maydis}, tub1, confirmed a role for interphase microtubules in determination of cell polarity and growth. Observation of GFP-Tub1 fusion protein revealed that spindle pole body-independent and astral microtubules are dynamic, with elongation and shrinkage rates comparable to those found in vertebrate systems. In addition, very fast depolymerization was measured within microtubule bundles. Unexpectedly, interphase microtubules underwent bending and rapid translocations within the cell, suggesting that unknown motor activities participate in microtubule organization in \textit{U. maydis}.

Movies available on-line:

http://www.biologists.com/JCS/movies/jcs1792.html

Key words: Pathogenic fungus, Polar growth, Green fluorescent protein, Microtubule, Motility

**INTRODUCTION**

Many eukaryotic cells are able to follow external stimuli by rearrangement of their cytoskeleton and subsequent membrane traffic towards polarized cellular domains. Filamentous actin and microtubules (MTs) in combination with associated proteins, like molecular motors, support this process by directed delivery of growth supplies during polar growth of some plant cells (Bibikova et al., 1999), vertebrate cells (Nabi, 1999) and fungi (Heath, 1995; Mata and Nurse, 1998). In contrast to actin fibres, MTs support long distance transport processes and thereby determine directionality of cell extension (Nabi, 1999). MTs consist of globular tubulin subunits that slowly polymerize and rapidly depolymerize at their plus-ends (Desai and Mitchison, 1997). This dynamic behavior, in combination with translocation of complete MTs (see Tanaka and Kirschner, 1991; Vorobjev et al., 1997), enables vertebrate cells to react to external stimuli and allows adaptation to specialized functions (reviewed by Joshi, 1998).

In fungi, dynamics of green fluorescent protein (GFP)-tubulin-containing MTs were described for \textit{Saccharomyces cerevisiae} (Carminati and Stearns, 1997; Shaw et al., 1997; Maddox et al., 1999; Maddox et al., 2000; Adames and Cooper, 2000) and \textit{Schizosaccharomyces pombe} (Ding et al., 1998; Mallavarapu et al., 1999; Yamamoto et al., 1999; Drummond and Cross, 2000), and these studies revealed an involvement of MTs in nuclear migration and mitosis.

Many fungi are able to switch between yeast-like and hyphal growth, thereby adapting to special requirements during their life cycle (Carlile, 1995). In particular, yeast-hyphal dimorphism is a common feature of many human and plant pathogens including \textit{Candida albicans, Histoplasma capsulatum} and \textit{Ustilago maydis} (reviewed in Gow, 1995). Dimorphic transitions require the perception of external stimuli that induce the reorganization of the cytoskeleton followed by polar delivery of vesicles containing biosynthetic enzymes, membrane components and wall ingredients towards the expanding growth region. Therefore, it is thought that cytoskeleton-based membrane transport plays a key role in pathogenic development of these fungi (Gow, 1995).

The dimorphic fungus \textit{Ustilago maydis} is the causative agent of corn smut disease and causes considerable losses of grain yield (Agrios, 1999). Outside the plant the fungus exists as haploid yeast-like cells that grow by budding at the poles of the elongated cell (Jacobs et al., 1994). On the plant epidermis two cells of opposite mating types can fuse and give rise to a filamentous dikaryotic hypha, which advances over the plant surface and penetrates the host tissue. Inside the plant the fungus proliferates and tumor formation is induced, followed by karyogamy and sporogenesis (reviewed in Banuett, 1995).
Under laboratory conditions *U. maydis* can be easily cultured and is accessible to both genetic and molecular methods. Therefore, this fungal pathogen is well suited for the investigation of the molecular basis of pathogenicity and dimorphism (reviewed in Banuett, 1995). However, an understanding of the cellular organization of this pathogen and the role of the cytoskeleton in growth, dimorphism and pathogenicity is only at its beginning. These initial studies have revealed that microtubule-dependent organelle transport plays a key role in polar growth (Wedlich-Söldner et al., 2000) and hyphal elongation (Lehmler et al., 1997).

In this study we have investigated the organization and dynamic behaviour of MTs in *U. maydis*. Immunofluorescence studies and mutant analysis indicate a role of MTs in polar budding, morphogenesis and nuclear migration. In vivo analysis of GFP-Tub1 fusion protein demonstrate that MTs are dynamic during interphase and mitosis. Moreover, similar to vertebrate cells, MTs show bending and motility within the cell.

**MATERIALS AND METHODS**

**Strains and growth conditions**

*U. maydis* strains FB1 (a1b1) and FB2 (a2b2) have been described previously (Banuett and Herskowitz, 1989). Strain SG200 (a1 mfa2 b2W2 bE) is a derivative of SG100 (Bölker et al., 1995a) that grows filamentous on charcoal-containing solid medium and was used to investigate the hyphal stages of *U. maydis*. FB2rGFPTub1 and SG200rGFPTub1 were generated by integration of plasmid prGFPTub1 into the succinate-dehydrogenase locus of strain FB2 and SG200, respectively. FB1oteGFPTub1 and FB2oteGFPTub1 contained plasmid potefGFPTub1 integrated into the succinate-dehydrogenase locus of wild-type strain FB1. In strain FB2rTub1 the *tub1* gene is under the control of the *otef*-promoter (Bottin et al., 1996) and integrated by homologous recombination into the *tub1* locus of FB2. All transformation was done as described (Schulz et al., 1990). If not otherwise mentioned strains were grown at 28°C in 2.5% potato dextrose (PD) or complete medium (CM; Holliday, 1974) supplemented with 1% glucose (CM-G) or 1% arabinose (CM-A), respectively. Solid media contained 2% (w/v) bacto-agar. To generate FB2rTub1 mutants, cells were grown on PD or CM-G and incubated in a shaker at 28°C, 200 r.p.m.

**Identification of tub1**

The *tub1* gene was isolated via PCR using degenerate primers and genomic DNA of *U. maydis*. Primers were directed against conserved regions that are specific for α-tubulins. PCR was done using 30 cycles of 95°C, 1 minute; 52°C, 1 minute; 72°C, 2 minutes and primers TubA2 (AACACATGGTAAARGYAYCC) in combination with the reverse primers TubA3 (TIRAACCIGTNGRCACARTC) or TubA4 (ACRTACCTGIAAGCAANGDCG). DNA fragments of 156 bp and 342 bp were sequenced and used to identify a full-length clone in a cosmid library (Bölker et al., 1995b). All subsequent cloning was done using *Escherichia coli* K12 strain DH5α (Bethesda Research Laboratories) following standard protocols (Sambrook et al., 1989).

**Plasmid constructions**

**pTub1E**

A 3.5 kb EcoRI fragment, which contained the full-length *tub1* gene, was cut out of a cosmid and cloned into pUC18 to give pTub1E. **pGFPTub1**

The N-terminal eGFP-Tub1 fusion under the control of the *crg*-promoter was generated in three steps. The open reading frame of *tub1* was amplified from pTub1E generating an *Ndel* site at the start codon and an *EcoRI* site 385 bp downstream of the 3′ end of *tub1* (Pl-polymerase, primers: TubA15 (CGGAATTCCACAGATGGCGAATC-GAT), TubA16 (GGCATATGCAGTGAAGTTCCTTCC)). This fragment was cloned into pCR®2.1-TOPO (Invitrogen) and sequenced, eGFP (Clontech) was excised from p630 (A. Hartmann, unpublished) as a *NcoI*-Ndel fragment. The *crg*-promoter was obtained as a *KpnI*-NcoI fragment of 3.6 kb from p194 (A. Brachmann and R. Kahmann, unpublished). In a four-fragment ligation these DNA fragments were cloned into the *KpnI*-EcoRI digested backbone from p123 (C. Aichinger, unpublished) that contained the carboxin resistance cassette. After digestion with *SspI* the construct was integrated by homologous recombination into the succinate-dehydrogenase locus.

**potefGFPTub1**

To put the eGFP-*tub1* fusion construct under the control of the constitutive *otef*-promoter (Spellig et al., 1996) an *SbfI*-NcoI fragment from p123 containing the *otef*-promoter was used to replace the *crg*-promoter in prGFPTub1 (ligated to an *NcoI*-EcoRI fragment and an *EcoRI*-SbfI fragment from prGFPTub1).

**pcrgTub1**

This plasmid was used for the integration of the *crg*-promoter in front of the *tub1* gene to create conditional mutants of *tub1*. The construction was done in two steps. The 5′ flanking region, containing 556 bp from the *tub1*-promoter up to the ATG, was amplified by PCR from pTub1E, thereby generating two restriction sites (*EcoRI*-KpnI). This fragment, together with a 3.6 kb *crg*-promoter fragment (*KpnI*-Ndel) from prGFPTub2 (G. Steenberg and M. Brill, unpublished) and a 1506 bp 3′ flank (*Ndel*-NcoI) containing the 5′ region of *tub1* from prGFPTub1, was ligated into pSL1.180 digested with *MunI* and *NcoI*. In a second step a *BamHI*-PmlI fragment from pSLHyg+ (A. Brachmann, unpublished) containing a hygromycin resistance cassette was inserted in front of the *crg*-promoter into *BamHI*-Smal.

**Western analysis**

For preparation of protein extracts logarithmic cultures were harvested by centrifugation at 3000 r.p.m. Cell pellets were washed in cold AP100 (100 mM Pipes, 2 mM MgCl₂, 1 mM EGTA, 1 mM EDTA, protease inhibitor; Boehringer, Ingelheim) and disrupted in a French Press at 1.100 psi. Protein concentration was measured according to Bradford, 1976, and equal amounts of protein were loaded and separated by SDS-PAGE. Subsequent western blots were done as described (Steinberg and Schliwa, 1995).

**Light microscopy and image processing**

Cells from logarithmically grown cultures were harvested and immediately observed using a Zeiss Axioskop microscope. For motion analysis 60-frame sequences of single cells were taken at 500 millisecond intervals using a cooled CCD camera (Hamamatsu, C4742-95). Epifluorescence was observed using standard FITC, DAPI and Rhodamine filter sets. For colocalization studies eGFP fluorescence was detected by a specific filter set (BP 470/20, FT 493, BP 505-530, Zeiss). Image processing and measurements were done with Image-Pro Plus (Media Cybernetics) and Photoshop (Adobe). Statistical analyses using unpaired *t*-tests or Mann-Whitney test that do not assume a Gaussian distribution were performed using PRISM (GraphPad).

**Quantitative analysis of MT dynamics**

For quantification of the parameters of MT dynamics, sequences of 60 images at intervals of 500 milliseconds were taken. From these sequences growth and shrinkage rates, catastrophe and rescue values
Microtubules in *U. maydis*

and motility velocities were determined using Image-Pro Plus (Media Cybernetics). Sliding and MT translocation were typically a rapid and discontinuous motility event. Therefore, measurements of shrinking and growth of MTs in bundles excluded these motions and only continuous length changes spanning more than 3 frames (1 second) were taken into account. In budded cells long MTs run through the cell (a). At the onset of budding two ‘paired tubulin structures’ (PTS, arrow in b) appear at the growth region, and are in contact with most MTs. During early bud growth the PTS remain close to the bud neck (arrow in c) and, based on the staining intensity, appear to nucleate bundles of MTs into the growing bud, whereas presumably single MTs extend into the mother cell (c). Prior to mitosis the PTS are positioned one behind the other (arrow in d) and originate arrays of long MTs into mother and daughter cell. At the onset of mitosis cytoplasmic MTs disappear and short spindles are formed (e). During mitosis two asters of microtubules emanate from the nuclei (arrowheads in f) and nuclei migrate to the center of mother and daughter cell, respectively. Finally, cytoplasmic MTs reappear (g), and the cell undergoes cytokinesis. Bars, 3 µm. (C) MTs in dikaryotic hyphae. MTs were decorated with anti-α-tubulin antibodies. Hyphae contain 2-5 long bundles of MTs (a), which often end in the middle of the tip cell (b). Most MTs have no contact with the hyphal tip (arrow in c) or with the septum (arrow in d). Bars, 3 µm (a); 2 µm (b-d).

**Staining procedures**

For indirect immunofluorescence of MTs, formaldehyde (EM-grade, Polyscience) was added to growing cultures to a final volume of 4% and cells were fixed for 30 minutes, washed with phosphate-buffered saline (PBS, pH 7.2) and applied to coverslips precoated with poly-L-lysine. This was followed by washes with PBS and 30 minutes of treatment with 3 mg/ml Novozyme (NovoNordisk). Subsequently cells were washed and incubated in 1% Triton X-100 for 1 minute, followed by additional washes and incubation in blocking reagent (2% milk powder, 2% BSA in PBS, pH 7.2) for 10 minutes. Antibodies against α-tubulin (N356, Amersham) and Cy3- and Cy2-conjugated secondary antibodies (Dianova/Jackson Laboratories), were diluted in 0.2% milk, 0.2% BSA, 0.01% azide in PBS, pH 7.2 and applied for 60 minutes. For colocalization studies of GFP-MTs with tubulin antibodies, cells were fixed with 1-2% formaldehyde for 30 minutes and prepared for immunofluorescence as described above, except that 0.2% Triton X-100 was applied for 15 seconds.

Cytological studies of living cells were performed on logarithmically growing liquid cultures. For microscopic observation 5 µl cells were mixed with 5 µl 1% low melting agarose that was prewarmed at 36°C. Nuclear staining was done using cells fixed with 1% formaldehyde for 15 minutes followed by incubation with 1 µg/ml DAPI (Sigma) in PBS for 15 minutes at 65°C and subsequent PBS washes.

Fig. 1. Nuclear movement and microtubule organization in *U. maydis*. (A) Overlay of epifluorescence of nuclear DAPI-staining and light microscopy of haploid *U. maydis* cells. During most stages of budding the nucleus remains positioned in the middle of the mother cell (a-c). Prior to mitosis, the nucleus migrates into the bud (d) and mitosis occurs (e). This is followed by the separation of the condensed daughter nuclei (f) and subsequent positioning in mother and daughter cell, respectively (g). Bars, 3 µm. (B) Antibody staining of MTs at stages of the cell cycle that correspond to Fig. 1A. In un Budded cells long MTs run through the cell (a). At the onset of budding two ‘paired tubulin structures’ (PTS, arrow in b) appear at the growth region, and are in contact with most MTs. During early bud growth the PTS remain close to the bud neck (arrow in c) and, based on the staining intensity, appear to nucleate bundles of MTs into the growing bud, whereas presumably single MTs extend into the mother cell (c). Prior to mitosis the PTS are positioned one behind the other (arrow in d) and originate arrays of long MTs into mother and daughter cell. At the onset of mitosis cytoplasmic MTs disappear and short spindles are formed (e). During mitosis two asters of microtubules emanate from the nuclei (arrowheads in f) and nuclei migrate to the center of mother and daughter cell, respectively. Finally, cytoplasmic MTs reappear (g), and the cell undergoes cytokinesis. Bars, 3 µm. (C) MTs in dikaryotic hyphae. MTs were decorated with anti-α-tubulin antibodies. Hyphae contain 2-5 long bundles of MTs (a), which often end in the middle of the tip cell (b). Most MTs have no contact with the hyphal tip (arrow in c) or with the septum (arrow in d). Bars, 3 µm (a); 2 µm (b-d).
Inhibitor studies
All inhibitors were purchased from Sigma, and stocks of 1 mM benomyl, nocodazole, thiabendazole, taxol and cytochalasin D in DMSO were prepared and stored at −20°C. For inhibitor studies these drugs were added to growing cultures to 10 μM final concentration and cells were incubated for 30 minutes at room temperature under gentle shaking; effects on MTs were observed at the microscope. In control experiments, corresponding amounts of DMSO were added to the cultures. For halo assays 10 ml CM-A plates were overlaid with 5 ml 0.8% agarose, precooled to 50°C and 100 μl cell suspension added. Filter papers saturated with inhibitors were put on top of the plates. After 15 hours at 28°C cells had formed a lawn within the agarose, but due to the toxic effect of the inhibitors a clear halo around the filters was observed. The diameters of the halos were analyzed statistically.

RESULTS
The microtubule cytoskeleton undergoes polarization during bud growth
We analyzed MT organization during the cell cycle of haploid U. maydis cells by using a monoclonal antibody against α-tubulin. In growing liquid cultures most cells were unbudded and contained a single nucleus that was positioned in the cell center (Fig. 1Aa). At this stage antibodies detected 2-7 long tubulin structures running through the length of the cell (mean=4.1±1.6, n=20 cells; Fig. 1Ba), which were most likely bundles of MTs, as confirmed by in vivo observations of GFP-tubulin fusion protein (see below). At the onset of polar budding (Fig. 1Ab, arrow) a pair of two spherical tubulin structures appeared at the growth region, which focussed the MTs towards the bud (Fig. 1Bb, arrow). These paired tubulin structures (PTS) appeared to result from a gradual concentration of tubulin during early budding (Fig. 2Aa-e). Each spherical unit of these structures had a defined diameter (0.48±0.04 μm, n=10) and distance from each other (0.61±0.09 μm, n=10), and were found in 40-70% of all growing cells. While the bud elongated (Fig. 1Ac) the PTS remained in the proximal part of the daughter cell close to the neck (arrow in Fig. 1Bc). At this stage they radiated long MTs into the mother cell and 2-3 brightly stained bundles of MTs into the medium sized bud (Figs 1Bc, 2Af), whereas no MTs were nucleated by the nuclear spindle pole body (SPB). Prior to mitosis the nucleus migrated into the bud (Fig. 1Ad), while the PTS were still located in the neck region, radiating two arrays of MTs into both mother and daughter cell, respectively (Figs 1Bd, 2Ah). Interestingly, during these stages the PTS varied in shape, ranging from almost spherical (Fig. 1Bd) to more elongated (Fig. 2Ag,Bc-f), and often consisted of globular domains from which MTs were extruded out into the mother and daughter cell (Fig. 2Ba-f). At the onset of mitosis the DNA condensed within the bud (Fig. 1Ae). Cytoplasmic MTs and the tubulin structures disappeared and short spindles of approximately 1 μm were formed (Fig. 1Be), which elongated up to 4 μm before two condensed nuclei could be detected using DAPI staining. Only occasionally, short astral MTs were visible (Fig. 1Be).
preserved during this stage (not shown). In contrast, asters of MTs were observed in late mitosis that appeared to emanate from the SPBs (Fig. 1Bf, arrowheads), while nuclear DNA was still condensed (Fig. 1Af). Finally, nuclei were positioned in the center of both cells (Fig. 1Ag) and SPB-independent MT bundles reappeared in mother and daughter cell, respectively (Fig. 1Bg).

Dikaryotic hyphae contained 2-5 bundles of MTs (Fig. 1Ca), which often ended in the middle of the hyphal tip cell (Fig. 1Cb). MTs usually had no contact with the apex (arrow in Fig. 1Cc) or with the septum of the growing tip cell (arrow in Fig. 1Cd).

Microtubules determine the polarity of the cell

The concentration of MT ends at one cell pole prior to bud formation suggested a role of the tubulin cytoskeleton in bud initiation and growth. To gain further insight into the cellular role of MTs we isolated an $\alpha$-tubulin gene, tub1, by PCR using primers that were specific for $\alpha$-tubulin (see Materials and Methods). The predicted amino acid sequence of Tub1 spans 448 amino acids and shares highest sequence similarity with $\alpha$-tubulin from Schizophyllum commune (82% identity, not shown). We did not find indications for additional $\alpha$-tubulin genes, either by PCR or low stringency Southern analysis (not shown). Therefore, it appeared that tub1 was the only $\alpha$-tubulin gene in U. maydis and its deletion was expected to be lethal.

Consequently, we generated a conditional mutant strain, FB2rTub1, in which the $\alpha$-promoter (Bottin et al., 1996) was fused to tub1 by homologous recombination (see Materials and Methods). This construct allowed expression of tub1 in complete medium containing 1% arabinose (CM-A) and repression of tub1 in CM supplemented with 1% glucose (CM-
Western blot analysis revealed that shifting FB2rTub1 from CM-A to CM-G resulted in a drastic decrease of Tub1 within the first 4 hours (Fig. 3A), which lead to fragmentation of MTs after 2 hours (Fig. 4Ab) and a drastic decrease in MT number within 4-6 hours (Fig. 4Ac, Ae), while MT organization was normal in CM-A (Fig. 4Aa).

Interestingly, the anti-tubulin antibody detected a punctuate cytoplasmic staining of α-tubulin after 4-6 hours in CM-G (Fig. 4Ac, detail) that disappeared at 10 hours in CM-G (Fig. 4Ad, detail). This coincided with the reappearance of single MTs (Fig. 4Ad,e), whose number remained constant during extended growth in CM-G (not shown). The disappearance of MTs was accompanied by morphological changes of cells. In CM-A cells showed normal morphology (Fig. 4Ba), but became thicker after the shift to CM-G (Fig. 4Bb, 5 hours after shift; Fig. 4Bc, 8 hours after shift) until almost spherical cells were observed (arrow in Fig. 4Bd, 10 hours in CM-G). In addition, after 10 hours in CM-G FB2rTub1 cells showed lateral budding (Fig. 4Be), while this was not observed in the wild-type control. Such a role of MTs in bud formation was also confirmed by benomyl treatment, which disrupts MTs within 30 minutes (not shown). After 3 hours of drug treatment 43% of all cells (n=107) formed two or more buds that appeared mainly at the neck region. In addition, many buds grew at the opposing poles or at the lateral region of the cell (not shown). Such growth defects were not observed in the wild-type control cells treated with 0.5% DMSO, suggesting that, in the absence of MTs, both polar growth and growth site selection were impaired.

Expression of GFP-tub1 has no effect on cell growth and microtubule organization

We fused GFP to the amino terminus of tub1 and integrated this fusion construct into the succinate-dehydrogenase locus (Keon et al., 1991) of wild-type strains FB1, FB2 and the solopathogenic strain SG200, which grows filamentous on charcoal-containing plates and occasionally in liquid culture. Expression of the fusion construct either under the control of the crg-promoter (strains FB2rGFPTub1; SG200rGFPTub1) or the constitutive otef-promoter (strain FB1oteFGFPTub1; for details see Materials and Methods) led to an incorporation of GFP-Tub1 fusion protein into MTs. The ratio of
GFP-Tub1 did not negatively affect the cell, and, therefore, incorporation into the polar tubulin structures, expression of these experiments demonstrate that, although not sensitive between wild-type control strains and nocadazole, thiabendazole or taxol revealed no difference in similar for the number of MTs in FB2 and FB2rGFP-Tub1 compared to wild-type strains (Fig. 3B) and Tub1 was not incorporated into the paired tubulin structures and the two bundles that passed into the medium sized bud (Fig. 5A, compare Fig. 2Af). Moreover, the fusion protein could not complement for the depletion of Tub1 in FB2rTub1 under restrictive conditions, suggesting that the fusion protein is not fully active. Therefore, we thoroughly analyzed the growth and shape of GFP-Tub1 expressing strains at a statistical level. In addition, we compared the organization of MTs in FB1 as seen by antibody staining with the MT organization visualized by GFP-Tub1 fusion protein. Finally, we analyzed whether incorporation of GFP-Tub1 has any influence on MT stability by applying MT inhibitors that are known to modify MT dynamics. These experiments revealed that the expression of GFP-Tub1 affected neither cell shape nor doubling time of FB2rGFPTub1 or SG200rGFPTub1 compared to wild-type strains (P=0.2624 and 0.4023), and similarly for the number of MTs in FB2 and FB2rGFPTub1 (P=0.317). Moreover, in halo assays, treatment with benomyl, nocadazole, thiabendazole or taxol revealed no difference in sensitivity between wild-type control strains and GFP-Tub1 expressing strains to any of these drugs (not shown). In summary, these experiments demonstrate that, although not incorporated into the polar tubulin structures, expression of GFP-Tub1 did not negatively affect the cell, and, therefore, could be used to study in vivo dynamics of MTs.

**Free and astral microtubules show dynamic instability**

In vivo observation of GFP-Tub1 fusion protein in strain FB2rGFPTub1 and FB1otefGFPTub1, respectively, revealed four classes of microtubules: (1) long SPB-independent interphase MTs that were usually part of MT bundles, (2) short SPB-independent interphase MTs of 1-5 µm, which had no contact with other MTs and often showed rapid translocations within the cell, (3) SPB-bound and unbundled MTs that radiated into the cytoplasm during mitosis (astral MTs), and (4) spindle MTs. Interphase cells and hyphae contained 2-5 long filamentous structures that contained GFP-Tub1 (Fig. 5A,B). Shrinkage of these structures resulted either in their disappearance or in less intense staining (end of MTs marked by arrows in Fig. 5A,B), suggesting that these filaments either are single MTs or MT bundles. The rate of elongation of interphase MTs was approximately 10 µm/minute, which was not significantly different from that of astral MTs during mitosis in U. maydis (Tables 1, 2; P=0.6553). In addition, the frequency of switching from growth to shrinkage (catastrophe rate) was comparable for both MT types and similar to other values reported for fungi and vertebrate systems elsewhere (Tables 1, 2). On the other hand, shrinkage rates of long interphase MTs were significantly higher than those of astral MTs (P<0.0001). This was due to rapid shortenings at rates of 60-100 µm/minute (Fig. 6), which accounted for 15-18% of all depolymerization events in interphase and occurred almost exclusively within bundles (100% in FB1otefGFPTub1, n=80 in 45 cells; 91.7% in FB2rGFPTub1, n=72 in 35 cells). Interestingly, the difference did not depend on the cell cycle stage, as shrinkage rates of short unbundled interphase MTs and astral MTs in mitosis did not significantly differ (P=0.1833; Tables 1, 2). This suggests that there is a causal relationship between rapid depolymerization and MT bundling. Interestingly, switching from depolymerization to growth
(rescue) was almost never observed for interphase MTs, resulting in a remarkably low rescue rate (Table 1.).

In hyphae rapid depolymerization in opposite directions occurred at comparable frequency and velocity (Fig. 5B,C). Moreover, MTs grew towards both tip and septum of the hyphal cell. Neither polymerization rates nor rapid shrinking velocities differed with direction (elongation rate not different; $P = 0.1953$; shrinking rate not different; $P = 0.5781$). In addition, growth and shrinkage rates are not significantly different from that of haploid sporidia ($P = 0.1076$ and 0.6998, respectively). Assuming that rapid depolymerization occurred at the ‘plus-end’ of the MTs towards their ‘minus-ends’ these data suggest that MTs have an anti-polar orientation within the MT bundles in hyphae. A similar situation was observed in unbudded U. maydis sporidia. In contrast, more than 90% of rapid shortenings were directed towards the neck region of budding cells (arrow in Fig. 5A), while MTs elongated towards the rear cell pole and into the bud (arrowhead in Fig. 5A). In budded cells MT ends that were close to the neck and presumably attached to the PTS did not show detectable activity. This indicates that during bud growth MT ‘minus ends’ are oriented towards the bud neck and that the PTS might support their formation and stabilization. However, this MT reorganization did not influence MT growth and shrinkage rates, as mean values did not differ between unbudded and budded cells in strains FB1otefGFPTub1 and FB2otefGFPTub1 (growth rate in unbudded cells: 11.14±4.14, $n=39$ MTs; growth rate in budded cells: 10.50±2.87, $n=13$; data sets not different, $P = 0.7998$; shrinkage rate in unbudded cells: 40.08±19.92, $n=171$ MTs; shrinkage rates in budded cells: 40.89±19.84, $n=65$ MTs; data sets not different, $P = 0.6433$). Besides the described changes in length, single MTs and MT bundles frequently underwent bending and thereby often changed their location within the cell. Usually, entire bundles were bent (Fig. 7A,B). Occasionally, the polar focus point of

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**Table 1. Dynamics of non-centrosomal/SPB-independent microtubules**

<table>
<thead>
<tr>
<th></th>
<th>Growth rate (µm/minute)</th>
<th>Shrinkage rate (µm/minute)</th>
<th>Catastrophe rate (transitions/second)</th>
<th>Rescue rate (transitions/second)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Squid axon</td>
<td>1.14±0.42</td>
<td>7.98±1.08</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>A498</td>
<td>6.9±2.5</td>
<td>8.4±5.5</td>
<td>0.031</td>
<td>0.114</td>
</tr>
<tr>
<td>Xenopus neuron</td>
<td>10.6±1.91</td>
<td>9.7±2.28</td>
<td>0.012</td>
<td>n.d.</td>
</tr>
<tr>
<td>Ptk1</td>
<td>11.0±3.7</td>
<td>14.8±7.5</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
<tr>
<td>Schizosaccharomyces pombe</td>
<td>3.00±1.64</td>
<td>4.47±2.93</td>
<td>0.01667</td>
<td>0.031</td>
</tr>
<tr>
<td>Ustilago maydis (long MTs)</td>
<td>10.77±2.41 (14)</td>
<td>37.76±17.7 (72)</td>
<td>0.04027 (10)</td>
<td>0.00736 (42)</td>
</tr>
<tr>
<td>Ustilago maydis (short MTs)</td>
<td>3.95±2.30 (7)</td>
<td>20.12±15.65 (25)</td>
<td>n.d.</td>
<td>n.d.</td>
</tr>
</tbody>
</table>

All growth and shrinkage rates are given as means ± s.d. For definitions of catastrophe and rescue rates, see Materials and Methods.


Long and mainly bundled MTs of FB2otefGFPTub1 cells in interphase; number of MTs in parentheses.

Short (<5 µm) and unbundled MTs of FB2otefGFPTub1 cells in interphase; catastrophe and rescue rates were combined; note that all shrinking-data follow a non-Gaussian distribution; number of MTs in parentheses.
MTs was displaced (arrowhead in Fig. 7A), suggesting that MTs were not only oriented towards the cell pole but were held together by unknown structural components. In addition, presumably single MTs were bent by sliding across other MTs at a velocity of approximately 30 µm/minute (Fig. 7C, Table 3). This sliding often involved a single contact point (arrows in Fig. 7C) and, therefore, might reflect some MT bound motor activity that applied force on the neighboring MT and, thereby, caused bending of the polymer. MTs that underwent bending or sliding were excluded from any measurements of MT elongation or shrinking (see above). Moreover, elongation rates of randomly taken long and unbundled MTs did not differ from those organized in bundles ($P=0.6128$), suggesting that sliding within bundles is not a significant source of error for our polymerization/depolymerization data given in Tables 1 and 2.

In addition to sliding and bending we observed a third type of MT motility. In 10-15% of all haploid cells investigated (80 cells in both FB2rGFP/Tub1 and FB1otetGFP/Tub1) motion of short and unbundled MTs occurred (Figs 7D, 8A). Motile MTs had no contact with other MTs and moved over several µm at rates of approximately 40 µm/minute (Table 3), with maximal velocities up to 100 µm/minute. They showed normal depolymerization rates; however, elongation only rarely occurred at much slower rates (Table 1). MT motion usually followed defined cellular tracks that were occasionally visible as a dark line against the cytoplasmic background of GFP-Tub1 fusion protein (Fig. 7D, arrowheads), suggesting that a filamentous cytoskeletal element might support MT motion. Occasionally, several types of MT motion occurred within a sequence of 30 seconds (Fig. 8A). While one end of a newly formed MT kept in contact with a long MT (Fig. 8A, asterisk) the other end elongated (Fig. 8A, arrowhead). During growth the small MT slid along another MT at approx. 48 µm/minute (Fig. 8A, frames 2-5). It then remained stationary and the free end flipped direction, while the MT still elongated. This was followed by the detachment of the small MT and motion along the cortex at a rate of approx. 77 µm/minute (frames 8-10). We consider this complex moving behaviour to be most likely due to an unknown motor activity that underlies the MT motion in U. maydis. However, to exclude the possibility that MT motion results from rapid polymerization at one end and rapid shrinkage at the other end of a MT, a process called

**Fig. 8.** Motion of free MTs and fluorescence speckle experiments. (A) Formation, sliding and cortical motion of a short free MT. A MT elongates while it remains attached to another MT (attached end marked by an asterisk). The attached end slides along this MT, while the other end (arrowhead) still elongates. As it gains contact with the cortex the short MT changes direction and subsequently detaches and rapidly moves along the cortex. Bar, 1 µm. Time in seconds is given in the upper right corner. (B) Depolymerization in a small budded cell of FB2rGFP/Tub1 after 6 hours under restrictive conditions. Under GFP-Tub1 depletion the fusion protein is unevenly incorporated in the MTs, resulting in dark areas and bright speckles (asterisk). The speckles remain stationary while depolymerization at a rate of 42 µm/minute towards the small bud occurs (bud marked by arrow in overview). Time in seconds is given in the upper right corner. Bars, 5 µm in overview and 2 µm in sequence. (C) Motion of a free MT in FB2rGFP/Tub1 after 6 hours under restrictive conditions. GFP-Tub1 is almost depleted and dark patches are formed along the length of the MT (arrowheads). These patches remain at a constant distance from the end while the MT is moving at a rates of approx. 11 µm/minute, suggesting that the complete MT is translocated. Time in seconds is given in the upper right corner. Bars, 5 µm in overview and 2 µm in sequence.
treadmilling, we performed fluorescence speckle microscopy (Waterman-Storer et al., 1998). For this purpose we made use of very low expression levels of GFP-Tub1 protein in FB2rGFP-Tub1 grown for several hours under restrictive conditions (for details see Materials and Methods). Under these conditions, the cellular GFP-Tub1 level is drastically reduced, resulting in a very weak staining and an uneven incorporation of the remaining GFP-Tub1, leading to unlabelled patches within the MT (Fig. 8B,C). In case of treadmilling, these ‘landmarks’ on the MT should be stationary during length changes while the MT apparently moves. We observed this situation in MTs that rapidly depolymerized towards the neck region of small budded cells (Fig. 8B, bud in overview marked by an arrow). The bright speckle (asterisk) remained stationary while the MT depolymerized at a rate of approx. 42 μm/minute. In contrast, dark patches usually remained at a defined distance to the ends and moved with the motile MTs through the cell (Fig. 8C; asterisk marks the end of the cell), although these short MTs occasionally showed elongation and shrinkage (Table 1). This, again, strongly indicates that MT motion in U. maydis is an active transport process rather than a result of rapid length changes.

**Nuclear migration is supported by microtubules**

The nucleus remained positioned in the mother cell until it migrated into the bud prior to mitosis (Fig. 1A). At the onset of mitosis the SPB started to nucleate dynamic and short astral MTs. Occasionally, these MTs kept in contact with the cell cortex, suggesting that they might serve as an anchor for the spindle. After the mitotic spindle had elongated up to 2-3 μm

![Fig. 9. Microtubules and nuclear migration.](image)

(A) In FB1oGFP-Tub1 a rigid bridge of spindle MTs connects two asters during anaphase B. Astral MTs have contact with the cortex and appear to exert a pulling force on the spindle poles, resulting in lateral motion of the aster (arrows). These forces most likely support spindle elongation during mitosis. Bar, 3 μm. Time in seconds given in the upper right corner. (B) Motion of a MT aster during late stages of mitosis within a mother cell (daughter cell to the right). Long astral MTs radiate into the cell and have contact with the cortex (arrowhead). This interaction appears to support movement of the aster and seems to pull the nucleus towards its final position within the cell center. Note that a flexible bridge between both nuclei still exists (arrow, bud neck to the right). Bar, 3 μm. Time in seconds is given in the upper right corner. (C) Prior to cytokinesis, nuclei are positioned within mother and daughter cell, respectively. Dynamic astral MTs kept contact with the cortex and prominent short range motion of the asters was observed. Bar, 3 μm.

### Table 2. Dynamics of centrosomal/astral microtubules

<table>
<thead>
<tr>
<th>Species</th>
<th>Growth rate (μm/minute)</th>
<th>Shrinkage rate (μm/minute)</th>
<th>Catastrophe rate (transitions/second)</th>
<th>Rescue rate (transitions/second)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A498</td>
<td>6.17±3.5</td>
<td>6.59±5.2</td>
<td>0.024</td>
<td>0.112</td>
</tr>
<tr>
<td>BSC-1</td>
<td>7.6±3.6</td>
<td>15.0±12.5</td>
<td>0.043</td>
<td>0.071</td>
</tr>
<tr>
<td>Ptk1</td>
<td>11.9±6.5</td>
<td>19.8±10.8</td>
<td>0.0606</td>
<td>0.1299</td>
</tr>
<tr>
<td>CHO1</td>
<td>19.7±8.1</td>
<td>32.2±17.7</td>
<td>0.0541</td>
<td>0.1961</td>
</tr>
<tr>
<td>S. cerevisiae</td>
<td>4.6±5.7</td>
<td>4.8±2.9</td>
<td>0.026</td>
<td>0.019</td>
</tr>
<tr>
<td>S. cerevisiae</td>
<td>0.49±0.16</td>
<td>1.35±0.71</td>
<td>0.006</td>
<td>0.002</td>
</tr>
<tr>
<td>Ustilago maydis</td>
<td>10.22±4.51 (19)</td>
<td>24.87±13.94 (79)</td>
<td>0.0324 (22)</td>
<td>0.0303 (23)</td>
</tr>
</tbody>
</table>

All growth and shrinkage rates are means ± s.d.

1Yvon and Wadsworth, 1997; 2Dhamodharan and Wadsworth, 1995; 3Shelden and Wadsworth, 1993; 4Adames and Cooper, 2000, s.d. calculated from given s.e.m. and given sample size; 5Carminati and Stearns, 1997, s.d. calculated from given s.e.m. and given sample size

6Based on observation of FB1oGFP-Tub1 cells in telophase; number of MTs in parentheses.
Fig. 10. MT organization and nuclear migration during the cell cycle of U. maydis. Cell cycle stages of U. maydis are taken from Snetselaar and McCann, 1997. During G1 and second phase, unbudded cells contain bundles of anti-parallel MTs (a). At early G1 the cytoskeleton becomes polarized and PTS appear at the growth region (b). While the bud enlarges these structures appear to nucleate MTs and radiate arrays into the mother cell and the growing bud, whereas the SPBs are inactive (c). Prior to mitosis, the nucleus migrates into the bud (d). After it reached the daughter cell the SPBs begin to nucleate spindle as well as astral MTs (e,f), which interact with the cortex and appear to anchor the nucleus during early mitosis. In anaphase the spindle migrates into the neck, where it rapidly elongates (g). This process appears to be supported by astral MTs that pull on the spindle poles. Prior to cytokinesis nuclei migrate towards the cell centers (h), and are supported by unknown motors that interact with astral MTs. Finally, both nuclei are positioned in the middle of mother and daughter cell, respectively (i), and bundles reappear (k). Note that the graph does not indicate the duration of each cell cycle stage.

Table 3. Motility rates of tubulin structures in U. maydis

<table>
<thead>
<tr>
<th>motility event</th>
<th>Velocity (μm/minute)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Motion of single short MTs*</td>
<td>41.19±24.40 (19)</td>
</tr>
<tr>
<td>Sliding/loopping of MTs*</td>
<td>31.80±11.56 (8)</td>
</tr>
<tr>
<td>Spindle translocation in early anaphase</td>
<td>9.26±5.78 (6)</td>
</tr>
<tr>
<td>Spindle elongation in late anaphase</td>
<td>4.14±3.10 (6)</td>
</tr>
<tr>
<td>Aster motion in telophase</td>
<td>21.56±5.56 (15)</td>
</tr>
</tbody>
</table>

Values are means ± s.d. (n), based on the observation in FB1otetGFP-Tub1 and FB2rGFPTub1 expressing strains; however, GFP-Tub1-containing MTs depolymerized towards the neck region, suggesting that the minus-ends of MTs are concentrated at or near the PTS. Interestingly, early Pts colocalized with minus-end directed cytoplasmic dynein (G. Steinberg, unpublished), supporting the conclusion that Pts mark the minus-ends of MTs. Consequently, the Pts might represent a SBP-independent microtubule-organizing centre (MTOC) that nucleates MTs during those stages of the cell cycle in which spindle poles are

it migrated towards the neck constriction at approx. 9 μm/minute (Table 3). This coincided with the appearance of long astral MTs, which radiated into the daughter and mother cell, respectively (Fig. 9A). The nucleus, stained with DAPI, elongated up to 4.67±0.84 μm (n=20) before the DNA separated (not shown). However, a rigid bridge of MTs, probably the anaphase B spindle, still connected the daughter nuclei (Fig. 9A), while astral MTs appeared to pull on the spindle thereby supporting lateral motion of the asters (arrows in Fig. 9A). At this stage the spindle continued to elongate slowly at a rate of approx. 4 μm/minute (Table 3). With increasing distance between the nuclei the MT bridge between them became thinner and more flexible (arrow in Fig. 9B). During this stage, the MT aster showed prominent to-and-fro motion, which gave way to sudden translocations over several μm at a mean velocity of approx. 21 μm/minute (Fig. 9B; Table 3). This motion appeared to be mediated by astral MTs that extended into the cytoplasm (Fig. 9B, arrowheads). Finally, the nuclei were repositioned in the cell center (Fig. 9C). At this stage, astral MTs still had contact with the cell cortex, resulting in short-range motion, again indicating that forces were exerted on the nuclei via astral MTs.

To confirm a role for MT in nuclear migration and positioning we shifted FB2rTub1 to restrictive conditions and subsequently analyzed the position of nuclei. Under permissive conditions 91% of the large budded FB2rTub1 cells (n=100) contained either condensed mitotic nuclei, which were located in the bud, or post-mitotic nuclei positioned in the mother and daughter cell, respectively, and there was no obvious difference from wild-type strains (not shown). In contrast, after 4 hours in CM-G, more than 65% of the large budded cells (n=102) contained condensed nuclei that were incorrectly positioned within the mother cell, suggesting that in the absence of MTs nuclear migration towards the bud was impaired, although DNA condensation occurred. In agreement, benomyl treatment for 45 minutes occasionally resulted in abnormal positioning of post-mitotic nuclei, and after 3 hours after incubation this treatment drastically inhibited nuclear migration into the bud, with 97% of all large budded cells (n=100) containing condensed nuclei in the mother cell. In summary, these results indicate that nuclear migration in G2 is supported by SBP-independent MTs, while astral MTs mediate nuclear motility during mitosis.

**DISCUSSION**

MT-dependent membrane trafficking is essential for fungal growth (Heath, 1995) and is thought to participate in the morphological transition between yeast and hyphal growth of pathogenic fungi (Akashi et al., 1994; Yokoyama et al., 1990). Here we demonstrate that MTs are focussed towards the growing bud and support polar growth of haploid U. maydis cells. During mitosis and early G1 (Fig. 10), astral MTs that emanated from the SBPs supported nuclear migration and positioning. Interestingly, in vivo observation of GFP-MTs demonstrated that haploid cells as well as hyphae contained bundles of dynamic MTs, which had an anti-polar orientation and showed bending and translocation.

**The paired tubulin structures resemble microtubule organizing centers**

Prior to bud formation, the cytoskeleton of unbudded cells underwent polarization. MTs started to focus towards one cell pole and tubulin accumulated there, finally leading to the appearance of a pair of spherical tubulin structures (PTS), which appeared to organize MTs towards the growth region (Fig. 10). Unfortunately, PTS were not visible in GFP-tub1 expressing strains; however, GFP-Tub1-containing MTs depolymerized towards the neck region, suggesting that the minus-ends of MTs are concentrated at or near the PTS. Interestingly, early Pts colocalized with minus-end directed cytoplasmic dynein (G. Steinberg, unpublished), supporting the conclusion that Pts mark the minus-ends of MTs. Consequently, the Pts might represent a SBP-independent microtubule-organizing centre (MTOC) that nucleates MTs during those stages of the cell cycle in which spindle poles are
not active. In S. pombe, spindle poles are also inactive during interphase (Masuda et al., 1992), and two cytoplasmic MTOCs, so-called post-anaphase arrays (PAA), appear at the end of anaphase, where they nucleate MTs that support nuclear migration (Hagan and Hyams, 1988). Therefore, the PAA and the PTS described here might be functional analogues, although electron microscopy revealed that PAA in S. pombe are ring-like and might release MTs to give rise to MT asters (reviewed in Hagan, 1998). Further ultrastructural studies will be needed to understand the detailed organization of PTS in U. maydis.

Interestingly, growth of yeast-like cells of C. albicans might also be supported by polar MTOCs. Immunostaining of C. albicans indicated that SPBs are inactive in unbudded cells and SPB-independent MTs are present in the unbudded cell (Barton and Gull, 1988). Similar to U. maydis, these MTs are focussed towards the growth region at the onset of bud formation. However, as no polar MTOCs are described for C. albicans it remains to be seen whether polar MT organizers are a common feature of the budding form of dimorphic fungi.

**Microtubules support cell polarity**

The polar organization of MTs and defects in polar budding and growth of tub1-mutant strains argue for an involvement of MTs in vectorial transport towards the growth region, which is in agreement with recent findings that MT-dependent trafficking of endosomes is required for polar growth of U. maydis (Wedlich-Söldner et al., 2000). Such a function of MTs corresponds to recent reports on MT function in polar growth. Disruption of MTs with thiabendazole (Sawin and Nurse, 1998), as well as defects in the MT cytoskeleton in mutant strains, led to abnormal growth and the establishment of lateral growth sites (reviewed in Hagan, 1998; Mata and Nurse, 1998) that was probably due to impaired MT-dependent transport of polarity determining components, like Tea1 (Mata and Nurse, 1997) or F-actin (Sawin and Nurse, 1998), towards the cell poles. In agreement, disruption of MTs led to lateral budding in U. maydis, suggesting that MTs serve similar functions in fission yeast and U. maydis. However, the organization of MTs by polar MTOCs during bud growth of U. maydis is fundamentally different from S. pombe.

Surprisingly, although repression of tub1 in FB1rTub1 led to rapid fragmentation of MTs, the antibody still recognized α-tubulin dots in the cytoplasm. These dots disappeared after 8-10 hours under restrictive conditions, which coincided with the reappearance of some MTs. At present we consider it most likely that repression of tub1 disturbed some equilibrium of tubulin and associated factors, which led to rapid MT disruption when α-tubulin was repressed; later, the system recovered, thereby recruiting the residual Tub1 to newly formed MTs.

**GFP-Tub1 is incorporated into microtubules but not PTS**

GFP-Tub1 fusion protein was incorporated into MTs and did not influence cell shape, growth and MT stability. This is in agreement with previous studies showing that GFP-α tubulin was fully biologically active and its expression had no detectable effect on S. cerevisiae (Carminati and Stearns, 1997) and S. pombe (Ding et al., 1998). However, U. maydis GFP-tubulin fusion could not complement for the deletion of endogenous α-tubulin and, in addition, did not incorporate itself into PTS of U. maydis. We find it likely that steric hindrance account for the latter defect. Therefore, cells that only produce GFP-Tub1 fusion protein might not be able to form these structures. As discussed above, PTS have a key role in organizing the MT cytoskeleton during budding. Therefore, we find it most likely that an inability to form PTS accounts for the restricted activity of GFP-Tub1 fusion protein, while incorporation into MTs has no detectable effect on the cell.

**Microtubule dynamics of U. maydis are similar to vertebrate cells**

Growth and shrinkage rates of MTs are reported for a wide spectrum of cell types. In vertebrate cells growth rates are 8-20 μm/minute, while shrinkage occurs at higher velocity with mean rates of 10-32 μm/minute (see Tables 1 and 2). In contrast, recent studies on S. cerevisiae (Carminati and Stearns, 1997; Adames and Cooper, 2000) and S. pombe (Drummond and Cross, 2000) indicated that fungal MTs might exhibit significantly slower elongation and shortening velocities. In U. maydis interphase and astral MTs showed polymerization at mean rates of approx. 10 μm/minute and rapid depolymerization at mean rates of approx. 38 and 25 μm/minute, respectively. These values are most similar to Ptk1 and CHO1 cells (Sheleden and Wadsworth, 1993; see Tables 1, 2) and clearly demonstrate that neither slow growth nor reduced shrinkage rates are a feature of all fungal MTs. On the other hand, interphase MTs of U. maydis showed unusual rapid depolymerization events with rates up to 100 μm/minute that were restricted to MTs within bundles. Moreover, once depolymerization has taken place, switching to polymerization, a process reflected by the rescue rate, almost never occurred (Table 1). These results indicate that the mechanisms and control of MT depolymerization of bundled interphase MTs are specific for U. maydis and might differ from other fungal and vertebrate cell systems. On the other hand, we cannot completely rule out the possibility that sliding between MTs within a bundle might add to rapid depolymerizations. However, elongation rates of randomly taken bundled and long unbundled MTs did not differ, indicating that sliding within bundles is not significantly falsifying our growth and shrinkage data.

**Interphase cells and hyphae contain bundles of anti-polar microtubules**

Rapid depolymerization and slower polymerization of MTs occurs most often at the plus-end of MTs while the minus-ends are often stabilized by components of the MTOC (see Maddox et al., 2000; Wiese and Zheng, 2000). MT-dependent motor proteins utilize the polarity of MTs with most kinesins moving towards the ‘plus ends’, whereas dynein-mediated transport is ‘minus end’ directed. Therefore, knowledge of MT polarity is important for understanding intracellular transport processes in fungal hyphae (see Morris et al., 1995).

Our in vivo observations indicate that hyphae and unbudded haploid cells contain SPB-independent MT filaments. Rapid shortening within these structures leave less intensively stained MTs behind. Therefore, we consider it most likely that these filaments are MT bundles and it would be interesting to see whether ultrastructural data support this conclusion. Within these bundles rapid depolymerization occurs in both directions, indicating that MTs have an antipolar orientation. This
conclusion is supported by former studies on MTs in hyphae of U. maydis (Steinberg et al., 1998). In this study, anti-tubulin antibodies and MT inhibitors were used to show that recovery of MTs starts at both the apex and the basal region of the hyphal tip cell and it was speculated that SPB-independent hyphal MTs are organized in an antiparallel fashion (Steinberg et al., 1998). Such a mixed polarity of MTs was described for dendrites (Baas et al., 1988) raising the possibility that neurons and fungal hyphae are similar with respect to MT-dependent long-distance transport.

Similar motors might support microtubule motility and mitotic nuclear migration

In U. maydis mitosis occurs within the bud and chromosome segregation coincides with rapid nuclear migration. Our in vivo observations show that the SPB does not nucleate MTs during pre-mitotic nuclear motion, although nuclear migration into the bud is a MT-dependent process. Ultrastructural studies on U. maydis demonstrated that the SPB localizes to the leading edge of the migrating nucleus (O’Donell and McLaughlin, 1984), suggesting that the SPB might slide along stationary MTs, a mechanism also discussed for S. pombe (Hagan, 1998). In contrast, later stages of nuclear migration in U. maydis are supported by astral MTs emanating from the SPB that have contact with the cortex. Forces applied to these MTs appear to pull on the nuclei, thereby supporting anaphase B and nuclear migration during mitosis. Such a mechanism is well established for fungi (see Hagan and Hyams, 1996; Inoue et al., 1998; Adames and Cooper, 2000), and it is most likely that cortical dynein is involved in this process (reviewed in Steinberg, 2000).

Interestingly, MTs showed rapid movement and bending during interphase of haploid U. maydis cells. Such motility was not described for fungal cells; also bending of single GFP-MTs occurs upon contact with the cell poles in S. pombe, a process which most likely results from compression of the elongating MT (Drummond and Cross, 2000). However, bending of interphase MTs is a common feature of animal cells (Cassimeris et al., 1988; Seitz-Tutter et al., 1988; Tanaka and Kirshner, 1991). Remarkably, MT translocation in axons appears to be a MT independent mechanism, suggesting that cortical actin could be involved (Ahmad and Baas, 1995), which appears to anchor cytoplasmic dynein and thereby supports MT transport in neurons (Ahmad et al., 1998). Moreover, cortical MT sliding supports nuclear migration in S. cerevisiae and dynein is assumed to participate in this process (Carminati and Stearns, 1997; Adames and Cooper, 2000). In U. maydis, MTs move along unknown filamentous tracks, suggesting that similar transport mechanisms involving F-actin and dynein might account for this process. This raises the intriguing possibility that dynein participates in motion, sliding and looping of MTs in U. maydis; however, the molecular details of MT motility remain to be elucidated.

What is the cellular function of MT motility in U. maydis cells? It was argued that MT translocation supports axon elongation and neuronal growth (Tanaka and Kirshner, 1991) and helps to organize the MT cytoskeleton in epithelial cells (Mogensen, 1999). In contrast to polar axons, however, translocation of MT in U. maydis was often observed in unbudded cells that are not growing. It is important to note that most MT translocations were found in cells that contained a disrupted MT cytoskeleton. Therefore, short motile MTs might be MT fragments, rather than being a defined class of MTs, a notion that is supported by the rare and unusual slow polymerization of these short MTs (Table 1). Consequently, cortical translocation of short MTs might be without biological relevance, but reflects a cytoplasmic motor activity that participates organization of MT bundles in interphase and rearrangement during polar growth.

Conclusion

In contrast to S. cerevisiae, where MTs almost exclusively function in nuclear migration and chromosome segregation (Madden and Snyder, 1998), MTs play a key role in morphogenesis and cell polarity of U. maydis. Such a role of MTs is reminiscent of S. pombe, and similarities in MT organization with that of C. albicans suggest that our results for U. maydis reflect the significance of MTs for fungal morphogenesis and cellular organization in general. The dynamic parameters provided in this study demonstrate that MT dynamics in fungal cells could be within the range of vertebrate systems. Moreover, the existence of MT bending and cortical MT sliding in U. maydis will provide a new means for investigating the molecular basis and cellular role of these types of intracellular motility.

We are grateful to Drs R. Kahmann and M. Feldbrügge for stimulating discussions and helpful suggestions on the manuscript. We wish to thank M. Artmeier for technical help. This work was supported by the Deutsche Forschungsgemeinschaft (SFB 413). The tub1 nucleotide sequence reported in this paper has been submitted to the GenBank/EMBL Data Bank with the accession number Hx975089759.

REFERENCES


