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Microtubule Regulation in Mitosis: Tubulin Phosphorylation by the Cyclin-dependent Kinase Cdk1

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The activation of the cyclin-dependent kinase Cdk1 at the transition from interphase to mitosis induces important changes in microtubule dynamics. Cdk1 phosphorylates a number of microtubule- or tubulin-binding proteins but, hitherto, tubulin itself has not been detected as a Cdk1 substrate. Here we show that Cdk1 phosphorylates β-tubulin both in vitro and in vivo. Phosphorylation occurs on Ser172 of β-tubulin, a site that is well conserved in evolution. Using a phosphopeptide antibody, we find that a fraction of the cell tubulin is phosphorylated during mitosis, and this tubulin phosphorylation is inhibited by the Cdk1 inhibitor roscovitine. In mitotic cells, phosphorylated tubulin is excluded from microtubules, being present in the soluble tubulin fraction. Consistent with this distribution in cells, the incorporation of Cdk1-phosphorylated tubulin into growing microtubules is impaired in vitro. Additionally, EGFP-β3-tubulinS172D/E mutants that mimic phosphorylated tubulin are unable to incorporate into microtubules when expressed in cells. Modeling shows that the presence of a phosphoserine at position 172 may impair both GTP binding to tubulin and interactions between tubulin dimers. These data indicate that phosphorylation of tubulin by Cdk1 could be involved in the regulation of microtubule dynamics during mitosis.

INTRODUCTION

In eukaryotic cycling cells, interphase microtubules form a dynamic network that is essential for cell polarity and intra-cellular traffic. When cells enter into mitosis, the interphase microtubule network rearranges into a mitotic spindle that is responsible for faithful chromosome segregation between daughter cells. Microtubule dynamics and turnover increase strikingly as cells progress from interphase to mitosis, with a microtubule half-life of 5–10 min in interphase and of 60–90 s in mitosis (Wittmann et al., 2001).

Microtubule dynamics in cells rely in part on the intrinsic properties of the microtubule building block, the α-β-tubulin dimer, and its ability to bind and hydrolyze a GTP nucleotide (Mitchison and Kirschner, 1984). The tubulin dimer is subject to special posttranslational modifications such as glutamylation, tyrosination, and acetylation (MacRae, 1997; Westermann and Weber, 2003). There is little evidence for a role of these modifications in the regulation of microtubule dynamics. Tubulin can also be phosphorylated by several kinases (Westermann and Weber, 2003). However, tubulin phosphorylation has not been connected with the cell-cycle-dependent regulation of microtubule dynamics.

Microtubule dynamics are also regulated by a number of microtubule effectors, including microtubule-associated proteins (MAPs), molecular motors such as kinesins, the Ras-like GTPase Ran-GTP, microtubule plus end-directed proteins and tubulin-binding proteins (Andersen, 1999, 2000; Carazo-Salas et al., 2001; Cassimeris, 2002; Heald and Nogales, 2002; Kinoshita et al., 2002; Galjart and Perez, 2003). These microtubule- or tubulin-associated proteins are themselves under the control of a balance of protein phosphatases and kinases.

The cyclin-dependent kinase Cdk1 (or Cdc2), associated with its cognate partner cyclin B, is a key enzyme for entry in mitosis (Nigg, 2001) and is essential for spindle morphogenesis. Cdk1 up-regulates microtubule dynamics when added in cell-free Xenopus extracts (Verde et al., 1990, 1992). Cdk1 inactivation is necessary for proper anaphase spindle dynamics and for cytokinesis (Wheatley et al., 1997). Large amounts of Cdk1 induce the depolymerization of interphase microtubules when injected into mammalian cells (Lamb et al., 1990) and the destabilization of microtubule arrays when added on lysed mammalian cells (Lieuvin et al., 1994). Among Cdk1 substrates are a number of microtubule effectors (Ubersax et al., 2003). MAP4 has been shown to be phosphorylated by Cdk1 in vivo (Ookata et al., 1997), and other nonneuronal MAPs like E-MAP115 or XMAP215/TOG exhibit consensus sequences for this kinase (Masson and Kreis, 1995; Vasquez et al., 1999; Charrasse et al., 2000). Furthermore, MAP4 and XMAP215/TOG interact with cyclin B and this interaction could target Cdk1 to microtubules (Ookata et al., 1995; Charrasse et al., 2000). Phosphorylation of MAPs either dissociates them from the microtubule lattice.
Mishima et al., 1993, of Op18/stathmin. by Cdk1 at the onset of mitosis (Cassimeris, 2002). Phosphorylation during interphase, is phosphorylated on two serine residues that sequesters tubulin dimers and destabilizes microtubules implicated in different steps of mitosis. Phosphorylation by Mishima et al., France), or on a Leica TCS-SP2 confocal microscope (Wetzlar, Germany).

In the present study, we show that in addition to regulating microtubule effectors, Cdk1 can directly phosphorylate β-tubulin in vitro and in mitotic cells, and that this phosphorylation impairs tubulin incorporation into microtubules. We suggest that the phosphorylation of tubulin by Cdk1 may represent a means by which mitotic cells regulate microtubule dynamics.

MATERIALS AND METHODS

Antibodies

Anti-phospho-peptide P172 polyclonal antibody (Ab) was made by Eurogentech (Seraing, Belgium). Two rabbits were immunized with phosphopeptide Ac-VVpYpSPKSVDTVV-VEC-CONH2 and serum was affinity-purified against phosphopeptide and then depleted against the same unphosphorylated peptide. Anti-α-tubulin AG monoclonal antibody (mAb) and anti-α-tubulin YL1/2 rat mAb were gifts of Laurence Lancache (CEA, Grenoble, France). Anti-β-tubulin TUB2.1 mAb was from Sigma (La Verpillière, France), anti-GFP polyclonal Ab (A11212) from Molecular Probes, and anti-phosphorylated vimentin 4A4 mAb from MBL International (Woburn, MA). Cy3-conjugated anti-rabbit (Jackson ImmunoResearch Laboratories, West Grove, PA) and Alexa 488-conjugated anti-mouse (Molecular Probes) antibodies were used as secondary antibodies for immunofluorescence studies. For Western blots, peroxidase-conjugated goat anti-rabbit IgG and anti-mouse IgG secondary antibodies (Sigma) were used.

Cell Culture and Transfection

HeLa S3 cells were grown at 37°C in suspension cultures in MEM Eagle Joklik’s formula (Cambrey Bioscience, Walkersville, MD), supplemented with 10% horse serum (Invitrogen, Cergy, France) and 1% penicillin/streptomycin (Invitrogen). Adherent HeLa cells were grown in RPMI medium supplemented with 10% fetal calf serum and 1% penicillin/streptomycin on 100-mm plastic dishes for cell extraction preparation or on glass coverslips in 30-mm plastic dishes for transfection and immunofluorescence. Transient transfection was carried out with 1 μg of plasmid DNA using Lipofectamine plus reagent (Invitrogen) according to manufacturer’s instructions. Cells were maintained in the presence of DNA for 30 h. For analysis of GTP constructs by Western blot, transfected cells were scraped in Laemmli buffer. For immunofluorescence, cells were then either lysed in OPT buffer (80 mM Pipes, pH 6.7, 1 mM EGTA, 1 mM MgCl2, 0.5% Triton X-100, 10% glycerol) at 4°C for 5 min before fixation, or fixed directly for 10 min in methanol at −20°C. Cells were then treated for immunofluorescence and visualized either on a Zeiss Axioscop conventional microscope (Le Pecq, France), or on a Leica TCS-SP2 confocal microscope (Wetzlar, Germany).

Native and Recombinant Tubulin Purification

HeLa cell tubulin and bovine brain tubulin were purified as described (Caudron et al., 2000; Fourest-Lieuvin, 2005). Recombinant mouse α2- and β-α tubulins were produced from pET3a plasmids provided by Ronald Melki (LEBS, Gil-sur-Yvette, France). As adapted from Melki et al. (1997), recombinant tubulins were overexpressed in milligram quantities in Escherichia coli BL21(DE3), isolated as inclusion bodies from lysed cells using Bugbuster protein extraction reagent (Novagen, Darmstadt, Germany), dissolved in 20 mM Tris, pH 7.5, 10 mM dithiothreitol, 7.5 M urea, dialyzed against 20 mM Tris, pH 7.5, 0.1 M dithiothreitol, and stored at −80°C until use.

Mutagenesis

Point mutations were performed on mouse β-tubulin-pET3a (same as above) and EGFP-mouse β-tubulin constructs, using the QuickChange kit from Stratagene (La Jolla, CA). The EGFP-mouse β-tubulin construct was a generous gift from Jurgen Wehland and coworkers (Gesellschaft für Biotechnologische Forschung, Braunschweig, Germany).

Phosphorylation Reactions

HeLa cell tubulin (20 pmol), bovine brain tubulin (20 pmol), recombinant α- or β-α tubulin (10 pmol), or purified recombinant protein (25 ng) was phosphorylated in 1 μl of reaction mixture that contained 5 mM Tris, pH 7.5, 1% Microcristalline cellulose, 10 mM MgCl2, 200 U (10 ng) of human Cdk1-cyclin B (New England Biolabs, Beverly, MA) in 20 μl of MEM buffer (100 mM Mes, pH 6.7, 1 mM EGTA, 1 mM MgCl2, 0.1 mM dithiothreitol). Controls were performed with 2 μl of kinase storage buffer instead of kinase. Phosphorylation reactions were stopped with Laemmli buffer supplemented with 10 mM EDTA, and samples were processed for SDS-PAGE and autoradiography or analysis with a phosphorimager (Molecular Imager FX; Bio-Rad, Hercules, CA). Relative amounts of incorporated radioactivity were quantified using Quantity One software (Bio-Rad). For phosphorylation reactions on peptides, peptides mapping the human β1-tubulin sequence, RRNNFTFSVVPPKSVVPPKSVVPPK (10 pmol) and RRNTNSFSVVPPKSVSDTVYEP (pep172-P), were synthesized by Eurogentec. Note that two Arg residues were added at the N-terminus of each peptide to allow their binding onto P81 phosphocellulose (see below). Each peptide (150 μM) was phosphorylated in the same conditions as above. Reactions were stopped and amounts of radioactivity incorporated in peptides were quantified by peptide binding onto P81 phosphocellulose paper as described (Filhol et al., 1991).

Assembly Assays after Tubulin Phosphorylation

To allow proper tubulin assembly, we used a source of γ-ATP without any additives (Perkin-Elmer Cetus; reference BLU/NEG/002A). Bovine brain tubulin (35 μM) was phosphorylated for 20 min at 30°C with 20 μM γ-ATP, 100 μM ATP, 5 mM MgCl2, 10 mM β-glycerophosphate, 0.5 μM microcystine, 200 U (10 μl) Cdk1-cyclin B in 100 μl of PEM buffer (100 mM Pipes, pH 6.7, 1 mM EGTA, 1 mM MgCl2). Controls were performed with 2 μl of kinase storage buffer instead of kinase. Afterward, samples were incubated for 10 min on ice, to depolymerize any tubulin structure that could have assembled during the phosphorylation reaction. GTP (1 mM) was added and each mixture was allowed to polymerize for 30 min at 35°C. For experiments with taxol, taxol was added progressively to a final concentration of 35 μM during the polymerization step. Samples were then layered on PEM-60% glycerol cushions (150 μM) and centrifuged at 100,000 × g for 20 min at 35°C. Supernatants (10 μl), containing nonpolymerized tubulin, were removed from the top of the glycerol cushions and kept for SDS-PAGE. Cushions were discarded, and pellets were washed once with PEM buffer at 35°C before being incubated for 15 min at 4°C in 100 μl of PEM buffer with 10 mM β-glycerophosphate and 0.5 μM microcystine (and supplemented with 5 mM CaCl2 and 50 mM KCl) for experiments with taxol) to depolymerize microtubules. Pellets were then cleared by ultracentrifugation at 200,000 × g for 10 min at 4°C and processed for SDS-PAGE.

Tubulin Extraction from Synchronized Cells

Drugs used were purchased from Sigma, and stock solutions were in dimethyl sulfoxide (DMSO). HeLa S3 or HCT116 cells were synchronized to G1/S phase or to M phase by a 19-h treatment with 0.3 mM nocodazole or 0.3 μM noco decadole, respectively. At this concentration of noco decadole, cells were blocked in mitosis with no net microtubule disassembly (Jordan et al., 1992). Synchronization was controlled by flow cytometry analysis for noco decadole-release experiments, mitotic cells were washed twice with phosphate-buffered saline (PBS) and released into normal medium for 30 or 60 min, until most of them had reached metaphase (Sauer et al., 2005). For experiments with roscovitine and nocodazole, cells were treated for 2 h with 50 μM roscovitine or with DMSO alone and then treated for 3 h with 0.3 μM nocodazole, as adapted from Meijer et al. (1997). To perform tubulin extraction, 106 cells were washed in PBS and lysed with 1 ml of cold OPT buffer supplemented with 10 mM NaF, 10 mM β-glycerophosphate, 0.5 μM microcystine, 1 μM pepstatin, and 400 μM phenylmethylsulfonyl fluoride (PMSF). After a 15-min incubation on ice with shearing through a pipette tip, lysed cells were ultracentrifuged at 200,000 × g for 10 min at 4°C, and supernatant (i.e., cell extract) was stored at −80°C. To perform soluble and insoluble tubulin fractions, the protocol was as above except that cells were extracted with 1 ml of OPT at 35°C for 2 min (soluble fraction) and then washed twice with 40 ml of OPT buffer at 37°C, before being incubated 15 min on ice with 1 ml of cold OPT buffer (insoluble fraction). Eighteen milligrams of cell extraction pellets were incubated for 1 h at 4°C with 300 μl YL1/2-coupled Sepharose 4B previously washed with PEM supplemented with 10 mM NaF and β-glycerophosphate. Bound tubulin was eluted with peptide VGVD-VEGCEEEE(G)EE at 0.1 mg/ml in same buffer, and fractions were analyzed by Western blotting using P172 Ab. For analysis of anti-phosphorylated vimentin 4A4 mAb, 10 treated cells were washed once with PBS, lysed in 100 μl of 1% SDS, and sonicated before SDS-PAGE and Western blotting.

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**Evaluation of the Proportion of Phosphorylated Tubulin in Mitotic Cell Extracts**

Purified bovine brain tubulin was phosphorylated in vitro with Cdk1 in the presence of γ32P-ATP as described in **Phosphorylation Reactions** (see above). The stoichiometry of transferred ATP per mole of β-tubulin was calculated after SDS-PAGE using a phosphorimager and Quantity One software (BioRad). In parallel, aliquots from the same phosphorylation reaction were blotted together with tubulin from mitotic cell extracts. The blot was stained with Ponceau red and then incubated with the P172 Ab. The quantity of cell extract tubulin was evaluated from the Ponceau red staining by comparison with Ponceau red and then incubated with the P172 Ab. The quantity of cell extract tubulin was evaluated from the Ponceau red staining by comparison with the known amounts of bovine brain tubulin, using Quantity One software. Likewise, the P172 signal was quantified. With the in vitro-phosphorylated tubulin, it was possible to link the ratio [P172 staining/amount of tubulin] to a stoichiometry of phosphorylation. Therefore, it was possible to calculate the approximate stoichiometry of phosphorylation in mitotic cells from the ratio [P172 staining/amount of tubulin] for mitotic cell extract tubulin.

**RESULTS**

**In Vitro Phosphorylation of β-Tubulin by Cdk1**

We have previously shown that Cdk1 induces the destabilization of microtubule networks when added on lysed mammalian cells in culture (Lieuvin et al., 1994). Subsequently, we have designed a method for the isolation, in biochemical quantities, of proteins associated with HeLa cell microtubules (Fourrest-Lieuvin, 2005). In a survey of Cdk1 substrates in these cell extracts, we identified a band comigrating with tubulin (unpublished data).

To test directly whether Cdk1 could phosphorylate tubulin, we incubated purified HeLa cell tubulin with or without the human Cdk1-cyclin B1 complex (referred below as Cdk1), in the presence of γ32P-ATP. Results showed a phosphorylation of β-tubulin (Figure 1A). In the same manner, bovine brain β-tubulin was phosphorylated by Cdk1 (see for example, Figure 6A). An additional phosphorylated peptide was observed above β-tubulin (Figure 1A), not comigrating with α-tubulin but apparently corresponding to cyclin B1, which is a known substrate for Cdk1 (Labbé et al., 1989). To test directly whether β-tubulin, and not α-tubulin, was a Cdk1 substrate, we incubated purified recombinant β- and α-tubulins with Cdk1. As a positive control, we included a recombinant form of protein kinase CK2, which is a well-established substrate for Cdk1 in vitro and in vivo (Mulner-Lorillon et al., 1990; Litchfield et al., 1992; Bosc et al., 1995). Both CK2 and β-tubulin were phosphorylated by Cdk1 at similar levels, whereas α-tubulin was not phosphorylated (Figure 1B, right panel). These results indicate that β-tubulin, not α-tubulin, is a bona fide Cdk1 substrate in vitro.

**Determination of the Cdk1 Phosphorylation Site in β-Tubulin Sequence**

Analysis of β-tubulin sequences of diverse origins revealed the presence of two putative sites for phosphorylation by Cdk1. One site, S172PK, was a good candidate for phosphorylation by Cdk1 (Holmes and Solomon, 1996). This site is conserved among β-tubulin genes of different species, from the yeast Saccharomyces cerevisiae to humans (Figure 2A). It is also conserved between human β-tubulin isoforms. The other putative site, T219P, is very short and less conserved (Figure 2A).

To test whether the S172PK site could indeed be phosphorylated by Cdk1, we tested two synthetic peptides containing this site, pep172 and pep172-P. These peptides matched the human β1-tubulin sequence and were indistinguishable, except that a phosphate was incorporated at the Ser 172 residue of pep172-P. When incubated with Cdk1 and γ32P-ATP, pep172 incorporated large amounts of radioactivity, whereas pep172-P did not (Figure 2B). This experiment indicates that the S172PK site, and no other amino-acid in pep172, can be phosphorylated by Cdk1.

To test whether the S172PK site was the preferred phosphorylation site in β-tubulin, we performed point mutations in recombinant β-tubulin. S172A and T219A point mutants and a S172A/T219A double mutant were assayed for phosphorylation by Cdk1 along with WT β-tubulin (Figure 2C). The S172A point mutation strongly inhibited the phosphorylation of β-tubulin by Cdk1 (Figure 2C, lane 6), whereas the T219A point mutation did not (Figure 2C, lane 7). Phosphorylation was also inhibited in the case of the double mutant (Figure 2C, lane 8). These results indicate that the S172PK site
is the main Cdk1 phosphorylation site in recombinant H9252-tubulin.

**Anti-Phospho-Peptide P172 Antibody**

To study H9252-tubulin phosphorylation in cells, an anti-phospho-peptide polyclonal Ab directed against the phosphorylated S172PK site was raised (P172 Ab; see Materials and Methods). We first tested whether the P172 Ab was specific for phosphorylated tubulin, compared with unphosphorylated tubulin, using both recombinant H9252-tubulin and bovine brain tubulin. Western blot analysis showed that P172 Ab reacted with H9252-tubulin phosphorylated by Cdk1 (Figure 3A, lanes 1 and 3), not with unphosphorylated tubulin (Figure 3A, lanes 2 and 4). To test whether the P172 Ab was specific for the phosphorylated Ser172 residue on H9252-tubulin, we tested the antibody on recombinant H9252-tubulin point mutants incubated or not with Cdk1. Results showed that only mutants with phosphorylated Ser172 residue reacted with the antibody (Figure 3B, lanes 5 and 7). We conclude that the P172 Ab reacts specifically with phosphorylated Ser 172, when tubulin is exposed to active Cdk1.

**Phosphorylation on Ser 172 of β-Tubulin in Mitotic Cells**

To examine the β-tubulin phosphorylation status in interphase or mitotic cells, we used cell extracts from aphidicolin- or nocodazole-arrested HeLa S3 cells, respectively. The phoso-specific P172 Ab was blotted on affinity-purified tubulin from these cell extracts. Tubulin from nocodazole-arrested cells was distinctly labeled with P172 Ab, whereas tubulin extracted from cells arrested in G1/S using aphidicolin yielded a barely detectable signal (Figure 4A, bottom panel). The low concentration of nocodazole used in these experiments has been shown to block microtubule dynamics (Jordan et al., 1992). To test whether anomalies in spindle dynamics could interfere with tubulin phosphorylation, nocodazole-arrested cells were released into fresh medium for 30 and 60 min to allow recovery of spindle functionality before extraction (Sauer et al., 2005). The extent of tubulin phosphorylation was comparable in nocodazole-released cell extracts as in nocodazole-arrested cell extracts (Figure 4A). Similar results were obtained for adherent HeLa or HCT116 cells (unpublished data). When cells were pre-treated with roscovitine, an inhibitor of Cdk1 (Meijer et al., 1997), before exposure to nocodazole, the P172 Ab signal was strongly diminished on immunoblots (Figure 4B, middle panel). These results were consistent with β-tubulin being phosphorylated by Cdk1 in mitotic cells, but did not preclude the possibility that β-tubulin was phosphorylated by another kinase at a site cross-reacting with the P172 Ab. To test this possibility, equal amounts of WT and S172A purified recombinant β-tubulins were transferred onto nitrocellulose membranes, which were incubated with mitotic...
cell extracts at 37°C in the presence of ATP and MgCl₂ to activate mitotic kinases. Membranes were then washed and immunoblotted with P172 Ab (Figure 4C). WT/H₉₂₅₂-tubulin reacted strongly with P172 Ab after phosphorylation by mitotic extracts, whereas S₁₇₂₄/H₉₂₅₂-tubulin did not. This showed that P172 Ab was specific of the phosphorylated Ser₁₇₂ in H₉₂₅₂-tubulin, even when tubulin was exposed to a panel of mitotic kinases. Taken together, all these experiments demonstrate that H₉₂₅₂-tubulin is phosphorylated on Ser₁₇₂ in mitotic cells, presumably by Cdk1 itself.

We then evaluated the proportion of phosphorylated tubulin versus total tubulin in mitotic cell extracts (see Materials and Methods). This evaluation indicated that phosphorylated tubulin represented ~0.2–0.5% of the total tubulin in M-phase extracts.

Localization of Phosphorylated Tubulin in Cells
We used both immunofluorescence microscopy and cell fractionation to localize phosphorylated β-tubulin. In immunofluorescence studies, adherent HeLa cells were either labeled with P172 Ab alone (unpublished data) or sequentially double-labeled with P172 Ab and anti-β-tubulin TUB2.1 mAb (Figure 5, A and B). The P172 Ab stained mitotic cells, from metaphase to telophase, but not interphase cells (Figure 5A). Microtubule structures such as spindles or midbodies in mitotic cells were not labeled, as assessed by confocal microscopy analysis (Figure 5B). The P172 labeling was the same with single stain, in every cell type tested (Rat 2, MDCK, and HCT116 cells), whatever the method of fixation used. The P172 Ab may recognize other phospho-proteins than tubulin in whole mitotic cells, but, conservatively, our results demonstrate an absence of detectable incorporation of phosphorylated tubulin in microtubules in cells.

To substantiate this conclusion, we analyzed the soluble (corresponding to the free tubulin) and the insoluble (corresponding to microtubules) tubulin fractions from nocodazole-arrested HeLa S3 cells, by Western blotting with P172 Ab. Results showed that free tubulin was much more labeled by P172 Ab than polymerized tubulin (Figure 5C). A semiquantification of the P172 signal showed that ~77% of the phospho-tubulin was in the free tubulin pool (Figure 5D). Taken together, these results indicate that the bulk of phosphorylated tubulin remains unpolymerized in mitotic cells.

Assembly Properties of Phosphorylated Tubulin
To test the effect of phosphorylation on tubulin polymerization directly, we performed in vitro experiments. Bovine brain tubulin was phosphorylated or not with Cdk1 in the presence of γ³²P-ATP and then allowed to polymerize (see Materials and Methods). After polymerization, samples were centrifuged to separate microtubule pellets from unpolymerized tubulin. Supernatants and pellets were analyzed.
both with a phosphorimager (PI) and by Western blotting with P172 Ab (Figure 6A). Results showed that the bulk (58%) of total tubulin was in pellets (Figure 6A, lane 4 of Coomassie and Ponceau panels, and Figure 6C, top panel, /H11002 taxol histograms). In contrast, the majority (81%) of phospho-tubulin was present in the supernatant and thus was unpolymerized (Figure 6A, lane 3 of PI and P172 panels, and Figure 6C, bottom panel, /H11002 taxol histograms).

The poor incorporation of phosphorylated tubulin in microtubules could reflect either an impairment of tubulin polymerization by phosphorylation or the presence of a large proportion of denatured tubulin among phospho-tubulin molecules. To test these possibilities, taxol was added during the polymerization reaction, to increase total tubulin polymerization. Taxol addition induced an increase in the quantity of phosphorylated tubulin in the microtubule pellet (Figure 6B, P172 panel), which amounted to ~63% of total phospho-tubulin (Figure 6C, bottom panel, + taxol histograms). These experiments indicate that the bulk of phospho-tubulin is not denatured but has an impaired polymerization capacity.

**Behavior of EGFP-β-Tubulin**

To test whether, in cells as in vitro, phosphorylation of tubulin on Ser172 is sufficient to impair tubulin polymerization, we examined the incorporation, in cell microtubules, of various EGFP-tubulin mutants. These mutants mimicked either unphosphorylated tubulin (S172A mutation) or phosphorylated tubulin (S172E or S172D mutations). The WT and mutant constructs were transfected into adherent HeLa cells. Each of the constructs was expressed full-length as assessed in a control experiment (Supplementary Data, Figure S1A). After transfection, cells were either fixed directly (Figure 7A) or lysed before fixation to improve the visualization of microtubules (Figure 7B). Whole cells and lysed cells were either stained only with anti-GFP Ab (unpublished data), or double-stained with anti-GFP Ab and with anti-β-tubulin mAb (Figure 7).

When cells were not lysed before fixation, a GFP staining was observed in the cytoplasm of transfected cells, for every construct tested. However, it was very difficult to distinguish microtubules against the fluorescent background, probably because HeLa cells are not very flat (Figure 7A). When cells were lysed before fixation, we observed a colocalization of GFP staining with the β-tubulin staining in the case of cells transfected with EGFP-β-tubulinWT (Figure 7B, left panels). Concerning EGFP-β-tubulinS172A, a colocalization of the GFP staining with microtubules, although less extensive than in the case of the WT protein, was also evident (Figure 7B, middle panels). In contrast, when cells were transfected with either EGFP-β-tubulinS172E (unpublished data) or EGFP-β-tubulinS172D (Figure 7B, right panels), no GFP staining was observed on microtubules after cell
Phosphorylation of β-Tubulin by Cdk1

In this study, we find that Cdk1 can phosphorylate β-tubulin. We provide evidence that tubulin phosphorylation by Cdk1 occurs in mitotic mammalian cells and impairs tubulin incorporation into microtubules both in cells and in vitro. It is known that protein phosphatases and kinases affect the overall dynamic properties of cellular microtubules during interphase as well as mitosis (Verde et al., 1990; Lieuvin et al., 1994; Yoshida et al., 2003; Schaart et al., 2004; Trinczek et al., 2004). However, the literature stipulates that kinases and phosphatases act through the phosphorylation/dephosphorylation of effector proteins associated with microtubules or with tubulin. In this context, our findings are new, showing a direct interplay between a kinase and tubulin, not mediated by an associated effector.

We find that phosphorylation by Cdk1 mainly occurs on the Ser172 residue of β-tubulin. In the 3-D structure of β-tubulin, the Ser172 residue is located in the T5 loop, which binds the ribose of GDP or GTP, at the exchangeable nucleotide binding site of tubulin (Hesse et al., 1987; Lowe et al., 2001). As shown in our model (Figure 8B), addition of a phosphate group to Ser172 can interfere with nucleotide binding. Nucleotide binding and exchange are important for tubulin assembly, and interference with nucleotide affinity or binding kinetics could logically be involved in the observed inhibition of tubulin assembly capacity by Ser172 phosphorylation. Additionally, residues in the T5 loop are involved in contacts between dimers along protofilaments (Lowe et al., 2001) and modification of these contacts in phosphorylated tubulin might also perturb tubulin assembly. A comprehensive study of the mechanisms through which phosphorylation affects tubulin assembly will require biochemical amounts of isolated phosphorylated tubulin.

Our finding that phosphorylation inhibits tubulin polymerization is consistent with previous observations of microtubule disassembly in cells injected with large amounts of Cdk1 or in lysed cells incubated with Cdk1 (Lamb et al., 1990; Lieuvin et al., 1994). However, we find that <1% of total tubulin is phosphorylated in mitotic cells, and this raises questions concerning the mechanisms through which such a low stoichiometry of tubulin phosphorylation could influence microtubule dynamics.

One possibility is that phosphorylated tubulin interacts with microtubule plus ends in cells. In a mitotic cell, considering that polymerized tubulin represents ~50% of total tubulin, and based on an average microtubule length of 5 μm and on a tubulin dimer size of 8 nm (Caudron et al., 2002), the ratio of the number of microtubule plus ends to the number of microtubule dimers is ~1/16,000. For a stoichiometry of phosphorylated tubulin/total tubulin of 0.2% (20 phosphorylated dimers vs. 10,000 dimers), there is a 30-fold excess of phosphorylated tubulin dimers compared with microtubule plus ends. Additionally, Cdk1 may be specifically targeted to microtubule plus ends. For instance, in the yeast S. cerevisiae, the Cdk1–mitotic cyclin Cln4 complex has been found to be localized to the plus ends of astral microtubules by the protein Kar9, which is related to the mammalian tumor suppressor adenomatous polyposis coli (APC; Maekawa et al., 2003; Maekawa and Schiebel, 2004). In Δclb4 yeast cells, astral microtubule dynamics are altered, showing a moderate increase of growth and shrinkage rates (Maekawa and Schiebel, 2004). Maybe Cdk1 localization mechanisms exist in higher eukaryotic cells. Indeed, in mammalian cells, a subpopulation of Cdk1 is associated with spindles (Bailly et al., 1989; Ribasovol et al., 1989; Rattner et al., 1990; Andreassen and Margolis, 1994). The Cdc14 localization...
Phosphatase also associates with spindles and is essential for microtubule bundling and stabilization during anaphase and exit from mitosis (Cho et al., 2005; Higuchi and Uhlmann, 2005). Therefore, it would be of great interest to examine whether phosphorylated tubulin is a substrate for Cdc14. Phosphorylated tubulin at microtubule ends could conceivably affect microtubule dynamics. In previous studies, Verde et al. (1990) reported that Cdk1 had no effect on pure tubulin polymerization on isolated centrosomes. A requirement for APC or for other factors capable of targeting Cdk1 at microtubule plus ends may account for this negative result.

Because the S172PK site and the T5 loop are both very well conserved in evolution and are present in the yeast S. cerevisiae (Figure 2A), yeast genetics should provide powerful tools to investigate the effects of tubulin Ser172 phosphorylation on the cell cycle and on microtubule dynamics.

Figure 7. Transfected EGFP-β-tubulin<sub>S172D</sub> remains soluble, whereas EGFP-β-tubulin<sub>WT</sub> and EGFP-β-tubulin<sub>S172A</sub> colocalize with microtubules. Adherent HeLa cells were transfected with EGFP-β-tubulin<sub>WT</sub>, EGFP-β-tubulin<sub>S172A</sub>, or EGFP-β-tubulin<sub>S172D</sub>. Cells were either fixed directly (A) or lysed before fixation (B). Whole cells and lysed cells were double-stained with GFP Ab and α-tubulin AG mAb. In the case of EGFP-β-tubulin<sub>S172D</sub> (B, right panel), the GFP staining had been overexposed to visualize lysed transfected cells. The data presented here are representative of an analysis of 100 transfected cells from three independent experiments. Bars, 20 μm.
Figure 8. Ser172 is located near the nucleotide in β-tubulin, and phosphorylated Ser172 might interfere with GTP/GDP binding and turnover. (A) Localization of the Ser172 residue in the 3-D structure of the tubulin heterodimer (accession number PDB#1JFF, Lowe et al., 2001). Note that Ser172 is numbered Ser174 in PDB#1JFF. The serine residue is in red, the GDP nucleotide in β-tubulin is in green, and the GTP nucleotide in α-tubulin is in yellow. (B) Modeling of a phosphoserine (P-Ser) at the position 172, using the graphics package TURBO-FRODO. Some of the possible interferences between the phosphate group and the ribose of GDP are depicted with blue dotted lines.

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