Cell-cycle dependent phosphorylation of yeast pericentrin regulates γ-TuSC-mediated microtubule nucleation

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Abstract Budding yeast Spc110, a member of γ-tubulin complex receptor family (γ-TuCR), recruits γ-tubulin complexes to microtubule (MT) organizing centers (MTOCs). Biochemical studies suggest that Spc110 facilitates higher-order γ-tubulin complex assembly (Kollman et al., 2010). Nevertheless the molecular basis for this activity and the regulation are unclear. Here we show that Spc110 phosphorylated by Mps1 and Cdk1 activates γ-TuSC oligomerization and MT nucleation in a cell cycle dependent manner. Interaction between the N-terminus of the γ-TuSC subunit Spc98 and Spc110 is important for this activity. Besides the conserved CM1 motif in γ-TuCRs (Sawin et al., 2004), a second motif that we named Spc110/Pcp1 motif (SPM) is also important for MT nucleation. The activating Mps1 and Cdk1 sites lie between SPM and CM1 motifs. Most organisms have both SPM-CM1 (Spc110/Pcp1/PCNT) and CM1-only (Spc72/Mto1/Cnn/CDK5RAP2/myomegalin) types of γ-TuCRs. The two types of γ-TuCRs contain distinct but conserved C-terminal MTOC targeting domains.

Introduction The budding yeast spindle consists of ~40 microtubules (MTs) that extend between the two opposed spindle pole bodies (SPBs). Because of the closed mitosis in yeast, the SPBs remain embedded in the nuclear membrane throughout mitosis. Cell-cycle regulated spindle assembly begins in S-phase with nucleation of MTs onto the surface of the newly assembled SPB. As soon as MTs emanate from both SPBs, they interdigitate (pole to pole MTs) or attach to the kinetochores (pole to kinetochore MTs) at the end of S phase, forming the bipolar spindle (O’Toole et al., 1999).

The γ-tubulin complex is the core player in MT nucleation. In budding yeast Saccharomyces cerevisiae, two molecules of γ-tubulin (Tub4) assemble together with one molecule of Spc97 (ortholog of human GCP2) and Spc98 (ortholog of human GCP3) into a tetrameric γ-TuSC ring (γ-TuSC), which is conserved in all eukaryotes (Geissler et al., 1996; Knop et al., 1997; Guillet et al., 2011). The purified γ-TuSC of budding yeast is able to self-oligomerize into symmetric ring-like structures under low salt buffer conditions. The diameter and the pitch of the γ-TuSC ring resemble that of MT cylinder, suggesting that the γ-TuSC ring functions as a MT nucleation template. However, the in vitro nucleation activity of the γ-TuSC assemblies remained poor, presumably because of the suboptimal spacing of every second Tub4 within the γ-TuSC ring that blocks interaction with tubulin in the MT cylinder (Kollman et al., 2008, 2010; Choy et al., 2009). The concept of γ-TuSC oligomerization is further supported by in vivo measurements of budding yeast γ-tubulin complex components on detached single MT nucleation sites. The γ-tubulin:Spc97:Spc98 ratio was 2.4:1.0:1.3 with a total of ~17 γ-tubulin molecules per nucleation site (Erlemann et al., 2012), suggesting a slight excess of γ-tubulin and Spc98 molecules over Spc97 in the assembled MT nucleation site.
Microtubules are hollow structures made of proteins that have a central role in cell division and a variety of other important processes within cells. For a cell to divide successfully, the chromosomes containing the genetic information of the cell must be duplicated and then separated so that one copy of each chromosome ends up in each daughter cell. To separate the chromosomes, microtubules extend out from two structures called spindle pole bodies, which are found at either end of the cell, and pull one copy of each chromosome to opposite sides of the cell.

Although the individual proteins that make up a microtubule can self-assemble into tubes, this occurs very slowly, so cells employ other molecules to speed up this process. In yeast cells, a protein called gamma-tubulin is recruited to the spindle pole body by the protein Spc110, where it combines with two other proteins to form a complex called the gamma-tubulin small complex. Several of these complexes then join together to form a ring, which probably acts as the platform that microtubules grow from. Recent observations suggested that Spc110 may help to construct this ring, but without revealing how.

Now, Lin et al. reveal that Spc110 can regulate microtubule formation by controlling how several gamma-tubulin small complexes bind together, and have identified the exact section of Spc110 that interacts with the complexes. However, the Spc110 must become active before it can perform this role, and it is only activated during certain stages of cell division, through phosphorylation. The structures in Spc110 that bind to the gamma-tubulin small complex in yeast are also found in gamma-tubulin binding receptor proteins in human cells. The work of Lin et al. demonstrates that proteins that are assumed to have passive roles within cells, such as Spc110, often play more active roles instead.

DOI: 10.7554/eLife.02208.002

In most eukaryotic organisms, such as fission yeast, Drosophila, Xenopus, and human, multiple γ-TuCs are further assembled with additional GCP family members (GCP4-6) into the larger γ-tubulin ring complex (γ-TuRC) (Gunawardane et al., 2000; Zhang et al., 2000; Murphy et al., 2001; Anders et al., 2006). However, these additional GCP proteins are not encoded in the budding yeast genome.

Various proteins are involved in the recruitment of γ-tubulin complexes to microtubule organizing centers (MTOCs) such as centrosomes and SPBs. The small protein Mozart1 is encoded in most eukaryotic genomes except the one of budding yeast (Teixido-Travesa et al., 2012). In Schizosaccharomyces pombe and Arabidopsis thaliana Mozart1 interacts with the GCP3 subunit of γ-tubulin complexes (Janski et al., 2012; Nakamura et al., 2012; Batzenschlager et al., 2013; Dhani et al., 2013; Masuda et al., 2013). In S. pombe Mozart1 appears important for the γ-TuSC recruitment to SPBs (Dhani et al., 2013; Masuda et al., 2013).

Besides Mozart1, a group of conserved proteins called γ-tubulin complex receptors (γ-TuCRs) are required for targeting γ-tubulin complexes to MTOCs. Most of them carry a highly conserved centrosomin motif 1 (CM1) that interacts with GCP subunits of γ-tubulin complexes (Sawin et al., 2004). How Mozart1 and γ-TuCRs cooperate is not understood. However, in budding yeast cells that lack a Mozart1 gene, γ-TuCRs are the sole factors responsible for γ-TuSC recruitment to SPBs. Spc110 is the budding yeast homolog of pericentrin (PCNT) and functions as γ-TuCRs at the nuclear side of the SPB (Knop and Schiebel, 1997; Sundberg and Davis, 1997). The N-terminal Spc110 encompasses the CM1 that interacts with the Spc98 subunit of γ-TuSC (Knop and Schiebel, 1997; Nguyen et al., 1998; Vinh et al., 2002; Sawin et al., 2004; Zhang and Megraw, 2007; Fang et al., 2008; Samejima et al., 2008). In addition, the N-terminal region of Spc110 is phosphorylated in a cell-cycle dependent manner. Phospho-Spc110 appears as cells progress from S phase, continues accumulating during mitosis, and vanishes at the anaphase onset (Friedman et al., 1996; Stirling and Stark, 1996). Spc110 phosphorylation accounts for the impact of Cdk1 and Mps1 kinases on spindle dynamics (Friedman et al., 2001; Huisman et al., 2007; Liang et al., 2013). However, a clear understanding behind this observation is lacking. Interestingly, when γ-TuSC and an N-terminal fragment of Spc110 (amino acids 1–220 of Spc110; Spc1101–220) were co-expressed in insect cells, a filament-like γ-TuSC-Spc1101–220 complex formed. The nucleation capacity of this purified γ-TuSC-Spc1101–220 complex exceeded that of the γ-TuSC alone (Kollman et al., 2010). Thus, Spc1101–220 influences γ-TuSC properties with yet unclear mechanism.
Here we have tested the possibility that phosphorylation of the γ-TuCR Spc110 regulates MT nucleation by inducing γ-TuSC oligomerization. Single particle analysis of γ-TuSC incubated with phosphomimetic Spc110 mutant proteins showed that Mps1 and Cdk1 promoted MT nucleation through Spc110 phosphorylation. Phosphorylated Spc110 and the interaction with the N-terminal domain of Spc98 induce γ-TuSC oligomerization. In addition, bioinformatic analysis revealed a conserved region around T18, that we named Spc110/Pcp1 motif (SPM). SPM and CM1 motifs are both important for γ-TuSC binding and oligomerization. A comparison of γ-TuCRs for the presence of SPM and CM1 identified SPM-CM1 (Spc110, Pcp1, PCNT) and CM1-only types of γ-TuCRs (Spc72, Mto1, Cnn, CDK5RAP2, myomegalin) in most organisms. While the SPM-CM1 type of γ-TuCRs carries the PACT domain and is targeted only to the centrosome or the nuclear side of the SPB, the CM1-only type of γ-TuCRs has either a MASC (Mto1 and Spc72 C-terminus) (Samejima et al., 2010) or a CM2 motif and is recruited to, centrosomes, the cytoplasmic side of the SPB or acentrosomal MTOCs.

Results

Phosphorylation of N-Spc110 at Mps1 and Cdk1 sites is required for efficient interaction with γ-TuSC

To test whether Spc110\(^{1–220}\) phosphorylation promoted γ-TuSC ring formation, we purified GST-Spc110\(^{1–220}\) (named Spc110\(^{1–220}\)) from both E. coli and the baculovirus expression system. Spc110\(^{1–220}\) encompasses Cdk1 and Mps1 phosphorylation sites and the conserved CM1 motif (Figure 1A). Because of the post-translational modification machinery, Spc110\(^{1–220}\) purified from insect cells harboured phosphorylations on S60/T68 and S36/S91 (Figure 1—figure supplement 1A–D), that correspond to established Mps1 and Cdk1 sites, respectively (Figure 1A; Friedman et al., 2001; Huisman et al., 2007). In contrast, Spc110\(^{1–220}\) was not phosphorylated when purified from E. coli.

We incubated purified γ-TuSC with either the phosphorylated Spc110\(^{1–220}\) from insect cells (Spc110\(^{1–220,P}\) or the non-phosphorylated Spc110\(^{1–220}\) from E. coli and then analyzed the protein complexes by gel filtration. Spc110 dimerizes via the coiled-coli region in the centre of the protein (Kilmartin et al., 1993; Muller et al., 2005). To substitute for the lack of this region, we performed the assays with GST-Spc110\(^{1–220}\) that dimerizes via GST–GST interactions. We used TB150 buffer in our assay instead of the BRB80 buffer used by Kollman et al. (2010). BRB80 induces oligomerization of γ-TuSC without the need for addition of Spc110 (Figure 1—figure supplement 2A, B). In contrast, in TB150 buffer the majority of γ-TuSC was monomeric (Figure 1—figure supplement 2C). TB150 therefore allowed us to monitor the impact of Spc110\(^{1–220}\) on γ-TuSC oligomerization by gel filtration. Addition of Spc110\(^{1–220,P}\) shifted γ-TuSC into fractions that eluted earlier than when γ-TuSC was run on the columns on its own (Figure 1B, shift from fraction 10 to fraction 8). The peak at the void-volume (fraction 8) likely represented γ-TuSC oligomers. In contrast, non-phosphorylated Spc110\(^{1–220}\) expressed in E. coli did not change the γ-TuSC elution profile. The γ-TuSC pool eluted as monomeric γ-TuSC (Figure 1B, fractions 10–11). This result suggests that Mps1 and Cdk1 kinases regulate the interaction between Spc110 and γ-TuSC through phosphorylation of Spc110 N-terminal domain.

Phosphorylation of N-Spc110 regulates the γ-TuSC oligomerization promoting activity

To further confirm that phosphorylation of Spc110\(^{1–220}\) alters the interaction with γ-TuSC, phosphomimetic and non-phosphorylatable Spc110\(^{1–220}\) mutant proteins were purified from E. coli (Figure 2—figure supplement 1A). spc110\(^{2A}\) (Cdk1 sites: S36A, S91A) and spc110\(^{2A}\) (Mps1 sites: S60A, T64A, T68A) have been studied in vivo before and shown to behave as non-phosphorylatable mutants (Friedman et al., 2001; Huisman et al., 2007; Liang et al., 2013). In addition, we analyzed the phosphomimetic Spc110\(^{1–220,P}\) (Cdk1 sites: S36D and S91D), Spc110\(^{1–220,3D}\) (Mps1 sites: S60D, T64D, T68D), and Spc110\(^{1–220,5D}\) (Cdk1 and Mps1 sites: S36D, S60D, T64D, T68D, and S91D). Migration of all purified Spc110\(^{1–220}\) mutant proteins was comparable upon gel filtration, indicating that the mutations did not alter the overall structure of the protein (Figure 2—figure supplement 1B, C).

We used gel filtration chromatography to address whether Spc110\(^{1–220}\) mutant proteins induce higher-order γ-TuSC oligomers. The phosphomimetic Spc110\(^{1–220,P}\) (Cdk1 sites), Spc110\(^{1–220,3D}\) (Mps1 sites), and Spc110\(^{1–220,5D}\) (Cdk1 and Mps1 sites) induced an apparent shift of γ-TuSC towards the void-volume compared to the γ-TuSC only run or γ-TuSC plus non-phosphorylated Spc110\(^{1–220,WT}\) from E. coli (Figure 2A–C, Figure 2—figure supplement 2A, B). The behavior of the phosphomimetic Spc110
mutant proteins reflected that of in vivo phosphorylated Spc110–220\textsuperscript{p} purified from insect cells (Figure 1B), indicating that the mutations to acidic residues did mimic phosphorylation. In contrast, the γ-TuSC shift was not seen upon inclusion of non-phosphorylatable Spc110–220\textsuperscript{5A} (Cdk1 and Mps1 sites) with the γ-TuSC (Figure 2—figure supplement 2A, B). Thus, Spc110–220\textsuperscript{5A} behaved as the non-phosphorylated Spc110–220 purified from E. coli, suggesting that the non-phosphorylatable mutations did not impair the protein. This conclusion was further supported by the observation that Spc110–220 or TB150 buffer only on ice. Oligomerization of γ-TuSC-Spc110–220 was tested by gel filtration chromatography using a Superdex 200 10/300 column. Peak fractions of the chromatograms were analysed by SDS-PAGE and silver staining.

DOI: 10.7554/eLife.02208.004

Figure supplement 2. γ-TuSC does not oligomerize in TB150 buffer.
DOI: 10.7554/eLife.02208.005

Because the upper limit of the separation range of the Superose 6 gel filtration column is ~5000 kDa, the γ-TuSC oligomers in the void-volume contain at least 14 copies of γ-TuSC (14-mers). This indicates the presence of ring-like complexes or even higher-order oligomeric structures in the void-volume. To confirm the existence of structurally organized oligomers, fractions corresponding to the void-volume peak (Figure 2—figure supplement 2A, fraction 7) were subjected to EM analysis (Figure 2D,E, Figure 2—figure supplement 4A). Consistent with the low A\textsubscript{280} absorbance, hardly any oligomeric γ-TuSC species were found in the void fraction of γ-TuSC only or γ-TuSC incubated with the non-phosphorylated Spc110–220\textsuperscript{WT} (Figure 2E, Figure 2—figure supplement 4B). In contrast, oligomeric
Figure 2. Mps1 and Cdk1 phosphorylation of N-Spc110 stimulates γ-TuSC oligomerization. (A) Summary of combination of non-phosphorylatable and phospho-mimicking mutations in Spc110 used in this study. The indicated Spc110<sup>1–220</sup> variants were expressed, purified from E. coli, and then tested for γ-TuSC oligomerization. WT: wild-type; 2A/D: S36A/D, S91A/D; 3A/D: S60A/D, T64A/D, T68A/D; 5A/D: S36A/D, S91A/D, S60A/D, T64A/D, T68A/D
(see Figure 2—figure supplement 1A for SDS-PAGE of purified Spc110–220 proteins and Figure 2—figure supplement 1C for gel filtration chromatograms). (B) Spc110–220 phospho-mimicking proteins induced γ-TuSC oligomerization. Spc110–220 proteins were incubated with γ-TuSC in TB150 buffer. The reconstituted complexes were separated according to size by gel filtration using a Superose 6 column. (C) Bar graph of area ratio of void-volume peak to total area of chromatogram of (B). ** marks statistical significance at p<0.01. Error bars represent SEM. N = 3 to 9 for the number of experiments performed. (D) Void-volume peak fractions of (B) were subjected to negative staining and protein complexes were analyzed with electron microscopy. Representative ring-like structures of γ-TuSC-Spc110–220-2D, γ-TuSC-Spc110–220-3D, and γ-TuSC-Spc110–220-5D. See Figure 2—figure supplement 4A for additional EM pictures. Scale bar: 50 nm. (E) Quantification of (D). Shown is the particle number per field. Particles were categorized based on the morphology. N is indicated on the figure for the number of fields analysed. (F) Quantification of void-volume fractions of γ-TuSC chromatograms. γ-TuSC was incubated with Spc110–220-WT, Spc110–220-5D, Spc110–220-T18A, Spc110–220-T18D, and Spc110–220D-T18D as described in (B). Bar graph of area ratio of void-volume peak to total area of chromatogram was calculated as in (C). ** marks statistical significance at p<0.01. Error bars represent SEM. N = 3 to 9 for the number of experiments performed. (G) Quantification of EM. Void-volume peak fractions of (F) were subjected to negative staining and protein complexes were analyzed with electron microscopy. Particles were categorized based on the morphology. Shown is the particle number per field. Note, the Spc110–220 and Spc110–220D graphs are the same as in (E). N is indicated on the figure for the number of fields analysed. (H) Multiple sequence alignment of SPM element of γ-complex receptors from yeast to human. Residues are marked according to the ClustalX colour scheme. The occurrence of each amino acid in each position of CM1 motif is presented with Weblogo 2.0. The following figure supplements are available for figure 2: DOI: 10.7554/eLife.02208.006

The following figure supplements are available for figure 2:

Figure supplement 1. Purification of Spc110–220 variants.
DOI: 10.7554/eLife.02208.007

Figure supplement 2. Phosphomimetic but not SPM defective N-Spc110 proteins induce oligomerization of γ-TuSC.
DOI: 10.7554/eLife.02208.008

Figure supplement 3. Non-phosphorylatable mutations in Mps1 or Cdk1 sites of Spc110 are neutral to γ-TuSC oligomerization induced by phosphomimetic mutations.
DOI: 10.7554/eLife.02208.009

Figure supplement 4. EM single particle analysis of oligomerized γ-TuSC.
DOI: 10.7554/eLife.02208.010

Figure supplement 5. Mutations of T18 abolish the γ-TuSC oligomerization promoting activity by inactivating the SPM motif.
DOI: 10.7554/eLife.02208.011

Figure supplement 6. Multiple sequence alignment of CM1 motif-containing proteins.
DOI: 10.7554/eLife.02208.012

γ-TuSC ring-like assemblies were observed when Spc110–220D, Spc110–220-5D, and Spc110–220-5D (Figure 2D,E, Figure 2—figure supplement 4A,B) had been added to the γ-TuSC preparation. Interestingly, the percentage of γ-TuSC spirals increased steadily with increasing numbers of phosphomimetic mutations from Spc110–220-2D, Spc110–220-3D to Spc110–220-5D (Figure 2E, Figure 2—figure supplement 4B), while the number of rings per field was relatively constant (Figure 2—figure supplement 4C). The outer diameter of the ring-like assemblies was on average around 40 nm (Figure 2—figure supplement 4D). Thus, combined phosphorylation of Spc110 by Cdk1 and Mps1 kinases enhances the oligomerization activity of Spc110 better than phosphorylation by either kinase alone.

The SPM motif is important for Spc110 oligomerization promoting activity

An additional phosphorylation of N-Spc110 at T18 has been identified in phosphoproteomic studies (Albuquerque et al., 2008; Keck et al., 2011; Lin et al., 2011). However, its precise function remained unclear (Liang et al., 2013). To elucidate the role of Spc110T18 phosphorylation, Spc110–220-T18D, Spc110–220-3D-T18D (Cdk1 plus T18), Spc110–220-5D-T18D (Mps1 plus T18), Spc110–220-5D-T18D (T18D in addition to Mps1 and Cdk1 sites), and Spc110–220-5D-T18A proteins were purified (Figure 2—figure supplement 1A) and tested for their γ-TuSC oligomerization promoting activity. Spc110–220 species with T18D or T18A failed to induce γ-TuSC oligomerization even when combined with the activating 5D mutations (Figure 2F, Figure 2—figure supplement 2C). Similar data were obtained with Spc110–220D-T18E and Spc110–220D-T18B (Figure 2—figure supplement 5A). Moreover, spc110T18, spc110T18E, spc110T18A, and spc110T18V showed identical growth defects at 37°C (Figure 2—figure supplement 5B). Since the effects of T18D/E and T18A/V mutations were indistinguishable, we cannot attribute these mutations as phosphomimetic or non-phosphorylatable. However, our results strongly suggest that these mutations
affect the structure of an important yet unappreciated element around T18 that is important for γ-TuSC oligomerization. Consistently, with EM we observed hardly any oligomeric structures when γ-TuSC was incubated with Spc1101–220-SD-T18D (Figure 2G).

With the sequence alignments of Spc110 orthologs from yeast to human, we observed a conserved motif upstream of CM1, which we designated as Spc110/Pcp1 motif (SPM) (Figure 2H). Spc110T18 sits within this motif. To evaluate the relative importance of CM1 and SPM, we constructed the Spc1101–220-CM1-QA mutant protein with mutations (KE to QA) in two highly conserved residues of the CM1 (Figure 2—figure supplement 6) and the Spc1101–220-ΔSPM lacking the SPM motif (deletion of amino acids 1–20). Spc1101–220 proteins were incubated with γ-TuSC in TB150 buffer to examine their oligomerization capacity. Spc1101–220-SD-CM1-QA and Spc1101–220-SD-ΔSPM mutant proteins failed to activate γ-TuSC oligomerization even when the five activating Cdk1 and Mps1 mutations were present (Figure 2I). Thus, Spc1101–220-SD-ΔSPM behaved as Spc1101–220-SD-T18D (compare Figure 2I with Figure 2—figure supplement 2C), further emphasizing that T18D inactivates the SPM. Taken together, SPM and CM1 of Spc110 are both required for γ-TuSC oligomerization.

The SPM and CM1 motifs and the Cdk1 and Mps1 phosphorylations regulate the affinity of Spc110 to γ-TuSC

Phosphorylation of Spc110 may regulate the γ-TuSC oligomerization by changing its binding affinity. Therefore we performed GST-pulldown assay to measure the binding of the Spc1101–220 variants to γ-TuSC. A constant amount of γ-TuSC was incubated with increasing concentrations of Spc1101–220 variants (0–300 nM). Consistent with the γ-TuSC oligomerization assay (Figure 2), Spc1101–220-SD showed stronger (p = 0.0017 for Spc97/Spc98 and p = 0.0006 for Tub4) γ-TuSC binding than Spc1101–220-WT (Figure 3A,B). Interestingly, while Spc1101–220-2D and Spc1101–220-3D induced comparable levels of γ-TuSC oligomerization (Figure 2B), Spc1101–220-2D showed less γ-TuSC binding than Spc1101–220-3D. A likely explanation is the 30-fold reduction in concentration that was used in the GST-pulldown assay compared to γ-TuSC oligomerization assays. These results suggested that phosphorylation of Spc110 on S36, S60, T64, T68, and S91 promotes γ-TuSC binding to Spc110.

We also analyzed binding of SPM (T18D and ΔSPM) and CM1 (CM1-QA) deficient Spc1101–220 mutant proteins to γ-TuSC. Both Spc1101–220-SD-T18D and Spc1101–220-SD-CM1-QA exhibited reduced γ-TuSC binding relative to Spc1101–220-5D (p < 0.01 for Spc97/Spc98 and p < 0.05 for Tub4) (Figure 3C,D). The γ-TuSC binding capacity of Spc1101–220-2D-5D-CM1-QA was further reduced (p < 0.01, Figure 3C,D), suggesting both CM1 and SPM are important for γ-TuSC binding. Interestingly, while Spc1101–220-SD-T18D, Spc1101–220-SD-CM1-QA, and Spc1101–220-ΔSPM all failed to oligomerize γ-TuSC (Figure 2F,I), they showed different levels of γ-TuSC binding capacity (Figure 3D). The decoupling of γ-TuSC binding capacity from γ-TuSC oligomerization suggests that both processes are not inevitably linked. Taken together, we conclude that both CM1 and SPM motifs contribute to γ-TuSC binding and are required for promoting γ-TuSC oligomerization.

SPM and the phosphorylation of N-Spc110 control MT nucleation activity in vitro

In vitro MT nucleation assays were performed to test the effect of Spc110 variants on γ-TuSC-mediated MT nucleation. Since phosphorylation of Spc110 and the presence of a functional SPM regulate γ-TuSC oligomerization into template rings, we expected to see differences in MT nucleation depending on these parameters (Figures 2 and 3). Compared to the buffer control and γ-TuSC alone, a threefold increase (p < 0.05) in MT nucleation activity was observed for Spc1101–220-2D, Spc1101–220-3D, or Spc1101–220-5D upon incubation with γ-TuSC (Figure 4A,B). In contrast, Spc1101–220-WT showed levels of MT nucleation that were comparable to the buffer control and γ-TuSC alone. Consistent with the proposed role of SPM, the MT nucleation level was reduced to buffer control levels when Spc1101–220-SD-T18D was incubated with γ-TuSC (p = 0.99). These results are in accordance with EM particle quantification assays, as Spc1101–220-2D, Spc1101–220-3D, or Spc1101–220-5D induced similar and significantly more γ-TuSC ring assemblies than Spc1101–220-WT and Spc1101–220-SD-T18D (p < 0.05, Figure 2E, Figure 2—figure supplement 4C).

In summary, these data support the model that Cdk1 and Mps1 phosphorylations of Spc110 regulate γ-TuSC-mediated MT nucleation by controlling template assembly.

Phosphorylation of Spc110 is cell cycle dependent

To analyze cell cycle dependent phosphorylation of Spc110, we generated one phospho-specific antibody against the two Cdk1 sites (Spc110S36 and Spc110S60) (Figure 5A) and another against the Mps1

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sites on Spc110 (Spc110^{560-pT64-pT68}) (Figure 5B). Both anti-Spc110^{pS36-pS91} and anti-Spc110^{560-pT64-pT68} antibodies gave specific signals in vitro and in vivo (Figure 5A–C).

To understand by which Cdk1 kinase complex Spc110^{S36} and Spc110^{S91} were phosphorylated, we co-immunoprecipitated Cdk1^{as1} either in complex with Clb5-TAP (S phase cyclin) or Clb2-TAP (mitotic cyclin) from yeast lysates and performed in vitro kinase assays using Spc110^{1–220-WT} that had been purified from E. coli. Spc110^{S36-S91} was phosphorylated equally by Cdk1-Clb2 and Cdk1-Clb5 as shown in Figure 3.
detected by Spc110<sup>pS36-pS91</sup> antibodies (Figure 5A). It is important to note that both Cdk1 kinase complexes had equal kinase activities toward histone H3 (Figure 5A, bottom).

To confirm the cell-cycle dependent phosphorylation of Spc110 in vivo, we immuno-precipitated (IP) Spc110-GFP with GFP-binder from cells arrested either in G1 phase with α-factor, S phase with hydroxyurea (HU) or prometaphase with nocodazole, and then blotted with phospho-specific antibodies. At G1 phase, no signal was observed with any of the two phospho-specific antibodies (Figure 5D). The phosphorylation of Mps1 was detected at S phase and maintained in mitosis. The Spc110<sup>pS36-pS91</sup> signal also appeared at S phase, and continuously increased in mitosis.
T18 within the SPM motif has been found phosphorylated in vivo and matches the minimal Cdk1 consensus sequence S/T-P (Keck et al., 2011). To elucidate the function of T18 phosphorylation, we raised phospho-specific antibodies against pT18. In vitro phosphorylation assay showed that T18 was only phosphorylated by Cdk1-Clb2 but not by Cdk1-Clb5 (Figure 5—figure supplement 1A). Due to

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**Figure 5.** Cell cycle dependent phosphorylation of Spc110. (A and B) Two phospho-specific antibodies were generated from guinea pigs to recognize phosphorylation of Cdk1 sites (pS36-pS91) (A) and Mps1 sites (p60-p64-pT68) (B). In vitro kinase assays were performed in the presence of recombinant Mps1 or Cdk1α and Spc1101–220, either wild type (WT) or non-phosphorylatable variants as indicated. 3A (Cdk1) indicates T18A-S36A-S91A and 3A (Mps1) S60A-T64A-T68A. In vitro phosphorylated Spc1101–220 was subjected to SDS-PAGE and immunoblot with the corresponding phospho-specific antibodies. The specific kinase activity of Cdk1α-Clb2 and Cdk1α-Clb5 was compared using human histone H1 as substrate (A, bottom). The numbers in the histone H1 experiment represent the relative kinase activity. As negative control of kinase activity, Mps1 was inactivated with chemical inhibitor SP600125 (B), while Cdk1α-Clb2 and Cdk1α-Clb5 overexpressed and purified from budding yeast were inactivated with 1NM-PP1 (A). (C) Phosphorylation of Spc110 and Spc110A in vivo. SPC110-GFP wild type cells (WT) and spc110A-GFP cells were arrested in mitosis with nocodazole. Spc110-GFP was enriched using GFP binder conjugated to beads, and the bound proteins were subject to immunoblotting with the indicated antibodies. (D) SPC110-GFP wild type cells were arrested in G1, S phase, and mitosis as indicated. Spc110-GFP was enriched with GFP-binder and analyzed by immunoblotting with the indicated P-specific antibodies. A non-specific band was used as loading control.

DOI: 10.7554/eLife.02208.015

The following figure supplements are available for figure 5:

**Figure supplement 1.** Phosphorylation of T18 most likely affects γ-TuSC oligomerization promoting activity of Spc110 in a negative manner.

DOI: 10.7554/eLife.02208.016
the low sensitivity of anti-Spc110\(^{\gamma}\), we used mass spectrometry for the analysis of T18 phosphorylation in vivo. Total yeast cell lysates were fractionated into supernatant and pellet that contained SPBs. The pellet fraction was solubilized with high salt buffer and then used for Spc110 enrichment. pT18 was predominately detected in the SPB-containing pellet fraction of prometaphase cells but not in the corresponding fraction of G1 or S phase arrested cells (Figure 5—figure supplement 1B,C). These data suggest that T18 of Spc110 becomes phosphorylated at SPBs in M phase but not S phase, which is consistent with published data (Keck et al., 2011).

Finally, we addressed the effects of T18 phosphorylation on γ-TuSC oligomerization. Because the T18A/V/D/E mutations inactivated the SPM function, we turned to an in vitro phosphorylation approach. Spc110\(^{1-220-SD}\) was incubated with Cdk1\(^{wt}\)-Clb2 in the presence or absence of the Cdk1\(^{1-220-SD}\) inhibitor 1NM-PP1. Only in the absence of 1NM-PP1, Spc110\(^{1-220-SD}\) was phosphorylated on T18 as shown by the phospho-specific anti-T18 antibodies. We incubated Spc110\(^{1-220-SD}\) and Spc110\(^{1-220-SD-pT18}\) with γ-TuSC and analyzed γ-TuSC by gel filtration. The activity of Spc110\(^{1-220-SD-pT18}\) to promote γ-TuSC oligomerization was significantly reduced in comparison with Spc110\(^{1-220-SD}\) (p<0.05) (Figure 5—figure supplement 1D,E). This suggests that in vitro phosphorylation of T18 by Cdk1-Clb2 inhibits γ-TuSC oligomerization.

Taken together, our results are in agreement with previous reports about the timing of Spc110 phospho-dependent mobility shift in SDS-PAGE (Friedman et al., 1996; Stirling and Stark, 1996) and suggest that the Cdk1 (S36, S91) and Mps1 sites become phosphorylated in S and in mitosis. In addition, phosphorylation of T18 within SPM occurs in mitosis and may negatively regulate SPM function.

**Cells with spc110 phospho-, SPM-, and CM1-mutant alleles have SPBs with impaired MT nucleation activity**

To reveal the function of phosphorylation of Spc110 in vivo, we replaced SPC110 with phosphomimetic and non-phosphorylatable mutant alleles. There was no significant difference in the expression levels of the Spc110 variants (Figure 6—figure supplement 1A). spc110\(^{2A/2D}\), spc110\(^{3A/3D}\), spc110\(^{A/5D}\), the SPM deficient spc110\(^{T18D}\), spc110\(^{SPM}\), and the CM1 deficient spc110\(^{CM1-QA}\) cells grew at 23°C comparable to wild-type SPC110 (WT) cells. However, spc110\(^{T18D}\) and spc110\(^{SPM}\) cells displayed a conditional lethal growth defect at 37°C and the SPM/CM1 defective spc110\(^{T18D-CM1-QA}\) double mutant was non-viable at 23°C and 37°C (Figure 6A, left panel). Combining T18D with 2D (Cdk1 sites), 3D (Mps1 sites) and 5D mutations slightly reduced growth at 37°C compared to the 2D, 3D, and 5D mutations (Figure 6A). This shows that, consistent with our in vitro γ-TuSC oligomerization results (Figure 2—figure supplement 2C), the SPM mutation T18D is dominant over the activating Cdk1 and Mps1 phosphomimetic mutations.

The defect of some mutant alleles involved in spindle assembly becomes apparent only after SAC function has been impaired (Wang and Burke, 1995; Daum et al., 2000). In the absence of the SAC gene MAD2 (Li and Murray, 1991), slow growth was observed in spc110\(^{2A}\) and spc110\(^{2D}\) cells at 37°C. This growth defect was even more pronounced in spc110\(^{2A}\) mad2\(^{\Delta}\) cells at 37°C (Figure 6A, right panel). All spc110 mutant alleles with SPM or CM1 inactivating mutations showed reduced growth in the Mad2\(^{\Delta}\) background compared to MAD2 SAC proficient cells (Figure 6A, bottom panel). Together, these growth tests suggest that the absence of phospho-regulation of Spc110 or the lack of SPM or CM1 results in defects that are compensated by a SAC induced delay in cell cycle progression.

Based on our model built from in vitro experiments, phosphorylation of N-Spc110 by Cdk1 (S36, S91) and Mps1 in S phase promotes the binding and oligomerization of γ-TuSC to form the template required for MT nucleation. In addition, the SPM and the CM1 motifs of Spc110 are important for γ-TuSC oligomerization and MT nucleation (Figure 2). Thus, mutations in SPC110 that affect any one of these sites/motifs should display a reduction in their SPB-associated tubulin signal. To test this hypothesis, we expressed GFP-TUB1 in cells and measured the intensity of the tubulin signal at the SPB at different cell cycle phases. Cells were synchronized with α-factor in G1 and then released into prewarmed media at 37°C, the restrictive temperature of some of the mutant cells. In SPM defective spc110\(^{T18A}\) and spc110\(^{T18D}\) cells, the reduction of GFP-tubulin signal at SPBs was already apparent in G1 (Figure 6B, Figure 6—figure supplement 1B, p<0.001). In S phase, the stage where newly synthesized SPBs mature and MT nucleation is initiated, all spc110 mutants except for spc110\(^{2D}\) cells exhibited a marked reduction in MT nucleation at SPBs when compared with WT cells (Figure 6B; for spc110\(^{T18A}\), p<0.05; for spc110\(^{T18D}\), spc110\(^{3A}\), p<0.001; Figure 6—figure supplement 1B). The reduction in tubulin signal was especially pronounced in spc110\(^{T18D}\) and spc110\(^{T18A}\) cells, indicating that SPM mutations blocked MT formation in vivo. spc110\(^{SPM}\) cells also showed MT defects at 37°C (Figure 6—figure supplement 1F).
**Figure 6.** Cells with spc110 phospho-, SPM-, or CM1-mutant alleles have defects in spindle formation. (A) Growth of 10-fold serial dilutions of SPC110 shuffle strains with integration vector encoding SPC110 (WT) or SPC110 mutants with and without the SAC gene MAD2. Growth was tested either on synthetic complete (SC) plates containing 5-FOA or SC dropout plates. (B) The indicated SPC110 wild type cells and spc110 mutant cells carrying
The situation in metaphase was similar to that of S phase although the overall relative differences between WT and spc110 mutants were less pronounced than in S phase (Figure 6B, p<0.001). Consistently, the overall spindle MT signal was reduced in spc110<sup>5A</sup>, spc110<sup>T18D</sup>, and spc110<sup>5D-T18D</sup> cells (Figure 6B,C). Taken together, the in vivo analysis of the tubulin signal at SPBs in spc110 cells is consistent with the in vitro data on γ-TuSC oligomerization and MT nucleation (Figures 2 and 4).

Reduced MT signal intensity at SPBs would be indicative of impaired γ-TuSC activity. In the simplest interpretation, γ-TuSC recruitment to SPBs is impaired. We found that the SPB signal from the γ-TuSC marker Spc97-GFP was reduced in spc110<sup>5A</sup>, spc110<sup>T18D</sup>, spc110<sup>5D-T18D</sup>, and spc110<sup>GSPM</sup> cells in G1 and S phase compared to spc110<sup>WT</sup> and spc110<sup>RD</sup> cells (Figure 6-figure supplement 1C,G). However, this reduction was not observed in mitotic cells. Analysis of Spc97-GFP signal intensity at SPBs in spc110 wild type, spc110<sup>T18D</sup>, and spc110<sup>GSPM</sup> cells revealed that the SPM mutation allowed Spc97 SPB binding but with a delay in cell cycle timing (Figure 6—figure supplement 1D,G). After SPB separation in late S phase wild type, spc110<sup>T18D</sup> and spc110<sup>GSPM</sup> cells eventually carried similar amounts of Spc97-GFP at SPBs.

Analysis of the GFP signal at SPBs in spc110-GFP, spc110<sup>5A</sup>-GFP, spc110<sup>T18D</sup>-GFP, and spc110<sup>5D-T18D</sup>-GFP cells did not reveal any differences between cell types (Figure 6—figure supplement 1E). Thus, the reduction of the Spc97-GFP signal in G1 and S phase most likely reflects a reduction of γ-TuSC binding to the spc110 mutant molecules at SPBs. The observation that all spc110 cells had similar γ-TuSC signals at mitotic SPBs while the MT signal was only reduced in spc110<sup>T18D</sup> and spc110<sup>5A</sup> cells (Figure 6—figure supplement 1C vs Figure 6B), suggests that the SPB-associated γ-TuSC is not fully active in MT organization in these cells. The functions of Spc110 probably extend beyond γ-TuSC binding, most likely to include γ-TuSC oligomerization (‘Discussion’). However, compared with spc110<sup>T18D</sup> or spc110 cells, Spc110-GFP at the SPB was reduced in spc110<sup>GSPM</sup> cells (Figure 6—figure supplement 1H), suggesting the SPM deletion affects incorporation of Spc110 into the SPB. This may contribute to the spc110<sup>GSPM</sup> phenotype and may explain why the growth defect of spc110<sup>GSPM</sup> cells was more pronounced than that of spc110<sup>T18D</sup> cells (Figure 6A).

To further examine the MT nucleation defect in spc110 phosho-mutant cells, we performed an in vivo MT nucleation assay. Cells were first arrested in S phase with HU, then MTs were depolymerized with nocodazole, followed by nocodazole wash out to trigger MT nucleation by the SPB bound γ-TuSC (Erlemann et al., 2012). To prevent cell cycle progression into anaphase we retained the cells in the HU block (Figure 6D). The re-growth of MTs was measured as a change in GFP-Tub1 intensity at SPBs. In comparison to WT and spc110<sup>RD</sup> cells, MT nucleation activity was reduced in the spc110<sup>5A</sup>, spc110<sup>T18D</sup>, spc110<sup>T18D</sup>, spc110<sup>5D-T18D</sup> cells (Figure 6E). Again, these data are consistent with the in vitro data (Figures 1–4) and suggest that Spc110’s ability to stimulate MT nucleation is regulated through stimulatory phosphorylation and requires the SPM motif.

**Spc110 interaction and γ-TuSC oligomerization are facilitated by the N-terminal region of Spc98**

The N-terminal region of Spc98/GCP3 orthologs is conserved. Spc98/GCP3 family members contain five predicted helices followed by an unstructured linker region (Figure 7A, Figure 7—figure supplement 1A). The conserved GRIP1 and 2 motifs that are common to all GCP proteins follow this...
Figure 7. The N-terminus of Spc98 mediates binding to N-Spc110. (A) Alignment of the amino acid sequence of GCP3 homologues from yeast to human. Shown are the putative α-helical regions H1–H5. Residues are marked according to the ClustalX colour scheme. (B) Growth test of spc98Δ1–156 cells with and without the SAC gene MAD2 at 23°C and 37°C. “SPC98” indicates the spc98ΔSPC98 cells while YPH499 is the unmodified wild type strain. (C) GFP-Tub1 signal Figure 7. Continued on next page
N-terminal region (Wiese and Zheng, 2006). Analysis of γ-TuSC by electron microscopy has shown that the N-terminus of Spc98 is in close proximity to the N-Spc110 binding site (Kollman et al., 2010). This topological arrangement raises the possibility that N-Spc98 is involved in binding to N-Spc110 and oligomerization of γ-TuSC. To test this idea, we constructed truncations of N-Spc98 with which to assess the function and biochemical behaviour of γ-TuSC<sup>Δ1–156</sup>. All N-terminal truncations of SPC98 supported cell growth at 23°C (Figure 7B, Figure 7—figure supplement 1B). However, Δ1–156, Δ1–177, and ΔLinker deletions of SPC98 showed reduced or no growth in mad2Δ cells at 37°C.

We analyzed the phenotype of spc98ΔΔN mutants at their restrictive temperature, 37°C. Because γ-TuSC<sup>Δ1–177</sup> formed aggregates already in TB150 buffer and was therefore unsuitable for oligomerization assay (Figure 7—figure supplement 2A, B), we focus our analysis on spc98Δ1–156. Like the wild-type γ-TuSC, the purified recombinant γ-TuSC<sup>Δ1–156</sup> was monomeric in TB150 (Figure 7—figure supplement 2C, D). spc98Δ1–156 cells incubated at 37°C had weaker GFP-Tub1 signal at their SPBs than wild type SPC98 cells (WT, Figure 7C). This is consistent with the observed mild MT nucleation defect of spc98Δ1–156 cells at 37°C (Figure 7—figure supplement 1C, D). Moreover, time-lapse analysis of α-factor synchronized SPC98 SP C42-Cherry and spc98Δ1–156 SP C42-mCherry cells with SPC97-GFP or SPC110-GFP revealed a delay in Spc97-GFP recruitment to SPBs at the time of SPB duplication (Figure 7D). Interestingly, slightly more SPC110-GFP was found at SPBs in spc98Δ1–156 cells, which may help to compensate the spc98Δ1–156 phenotype. Thus, the N-terminal region of SPC98 is important for optimal MT organization and timely γ-TuSC recruitment to SPBs.

We next asked whether γ-TuSC<sup>Δ1–156</sup> can be oligomerized by incubation with SPC110<sup>Δ1–220</sup>. Interestingly, SPC110<sup>Δ1–220</sup> failed to shift γ-TuSC<sup>Δ1–156</sup> to the void volume of the Superose 6 column, as was the case for γ-TuSC (Figure 7E). In agreement with the oligomerization experiment, γ-TuSC<sup>Δ1–156</sup> showed reduced binding to SPC110<sup>Δ1–220</sup> in the pull down assay (Figure 7F, G). Thus, the N-terminus of Spc98 is involved in binding to Spc110. In spite of not changing our conclusion, we noticed that Spc98Δ1–156, 6His was detected less strongly than Spc98-6His by the anti-His antibodies, although in Coomassie Blue stained gels both proteins were present in 1:1 ratio (Figure 7E, upper panel and F). Thus, it is likely that Spc98Δ1–156, 6His transferred less efficiently to the blotting membrane than Spc97-6His, leading to an underestimation of the Spc98Δ1–156, 6His signal relative to full-length Spc98-6His.

Finally, we tested γ-TuSC<sup>Δ1–156</sup> oligomerization in the BRB80 buffer that supports oligomerization without the aid of Spc110. γ-TuSC and γ-TuSC<sup>Δ1–156</sup> shifted to the void volume of the Superose 6 column with equal efficiency (Figure 1—figure supplement 2B, Figure 7—figure supplement 2C). Analysis of the void fraction by EM identified oligomeric γ-TuSC<sup>Δ1–156</sup> rings (Figure 7—figure supplement 2E). Therefore, the N-terminus of Spc98 is important for Spc110-mediated oligomerization but not for Spc110-independent γ-TuSC oligomerization.

Bioinformatic analysis of the C-terminal SPB/centrosomal binding domain of SPM-CM1 and CM1-only γ-TuCRs

Our biochemical and cell biology analyses have identified the novel motif SPM and CM1 (Sawin et al., 2004) as important motifs in Spc110 for γ-TuSC binding and oligomerization. Using bioinformatics
approaches, we investigated the presence of SPM and CM1 motifs in γ-TuCRs from various species (Figure 8A,B). A yet unappreciated CM1 motif was identified in another budding yeast γ-TuCR, Spc72 (Knop and Schiebel, 1998; Figure 8A, Figure 8—figure supplement 1C).

Based on the presence of SPM and CM1 motifs, γ-TuCRs can be divided into the SPM-CM1 and the CM1-only types. Spc110 and fission yeast Pcp1 fall into the SPM-CM1 class. Our analysis also predicts SPM and CM1 motifs, albeit with moderate sequence deviation, in mammalian PCNT (Figure 8B). Another group of γ-TuCRs have only a CM1 motif but no SPM. Human CDK5RAP2 and myomegalin, Drosophila Cnn, budding yeast Spc72, and fission yeast Mto1 fall into this class (Figure 8B).

A further level of category resolution was achieved by the classification based on the C-terminal MTOC targeting domains (Figure 8—figure supplement 1A,B). It was shown that yeast Spc110 (Sundberg et al., 1996; Stirling and Stark, 2000), Pcp1 (Fong et al., 2010), Drosophila D-PLP (Cp309) (Kawaguchi and Zheng, 2004), and mammalian PCNT (Gillingham and Munro, 2000; Takahashi et al., 2002) are targeted to nuclear side of SPBs or centrosomal MTOCs via the conserved PACT domain (Figure 9A). In case of SPBs that are embedded in the nuclear envelope and therefore have a cytoplasmic and a nuclear side, SPM-CM1 γ-TuCRs only function in organizing the nuclear MTs that separate the sister chromatids in mitosis (Figure 9B). In contrast, CM1-only γ-TuCRs of fungi, such as Spc72 in budding yeast (Knop and Schiebel, 1998), Mto1 in fission yeast (Samejima et al., 2010), and Aps8 in Aspergillus nidulans (Zekert et al., 2010), have a MASC motif and function at the cytoplasmic side of the SPB or in the cytoplasm. In case of metazoa, CM1-only type of γ-TuCRs such as CDK5RAP2 (Wang et al., 2010) possess a CM2 domain (Figure 9B, Figure 8—figure supplement 1A), targeting γ-TuCRs to both centrosomal and acentrosomal MTOCs.

In summary, γ-TuCR family proteins can be classified regarding the γ-TuSC interaction motifs and the MTOC targeting domain into three groups.

**Discussion**

**Spc110 is a γ-TuSC-mediated nucleation regulator**

In this study we show that the γ-TuCR Spc110 actively participates in MT formation through the SPM and CM1 elements and Cdk1 and Mps1 phosphorylation sites. Binding of γ-TuSC to Spc110 is a prerequisite for γ-TuSC oligomerization into the ring-like template that initiates MT assembly. Phosphorylation of Spc110 by Cdk1 and Mps1 regulates γ-TuSC binding and thereby MT nucleation. Thus, Spc110 is a MT nucleation regulator. We thereby provide a molecular understanding of the previously observed spindle length phenotypes (the two Cdk1-Clb5 sites) and genetic interaction with SPC97 (three Mps1 sites combined with S36) of SPC110 phosphorylation site mutants (Friedman et al., 2001; Huisman et al., 2007). Our data also explain why an N-terminal fragment of Spc110 influences oligomerization of γ-TuSC when co-expressed in insect cells (Kollman et al., 2010).

SPBs duplicate in G1/S phase of the cell cycle in a conservative manner (Byers and Goetsch, 1975; Adams and Kilmartin, 2000; Pereira et al., 2001; Figure 10A). In late G1, a SPB precursor, named the duplication plaque, assembles on the cytoplasmic side of the nuclear envelope at a specialized substructure of the mother SPB, named bridge (Byers and Goetsch, 1975; Jaspersen and Winey, 2004). In G1/S the duplication plaque then becomes inserted into the nuclear envelope. This allows binding of the Spc110-calmodulin-Spc29 complex from the nuclear side to the central Spc29- Spc42 layer of the embedded duplication plaque (Bullitt et al., 1997; Elliott et al., 1999). In this cell cycle phase Mps1 and Cdk1-Clb5 phosphorylate the N-terminus of Spc110 at five sites to promote γ-TuSC binding to Spc110 (Figure 10B). Consistently, Cdk1-Clb5 associates with SPBs throughout this time window (Huisman et al., 2007). These phosphorylations increase γ-TuSC affinity for Spc110-5P and induce γ-TuSC oligomerization into ring-like complexes that promote MT nucleation (Figure 10B). Phospho-regulation of Spc110 is not absolutely essential for viability as indicated by the growth of spc110Δα cells. However, spc110Δα cells have growth defects in the absence of the SAC gene MAD2 and fail to organize the full set of MTs in S phase and mitosis (Figure 6). We therefore suggest that Mps1 and Cdk1-Clb5 coordinate the timing of MT nucleation through Spc110 phosphorylation with SPB duplication. A SAC induced cell cycle delay can temper any defects in this phospho-regulation as indicated by the genetic interaction between spc110Δα and spc110Δα with mad2Δ (Figure 6A).
Figure 8. Two types of γ-TuCRs defined by the N-terminal γ-TuSC binding motifs: SPM-CM1 and CM1-only. (A) Graphical representations of the patterns of CM1 motif within the multiple sequence alignment of γ-TuCR protein sequences. The CM1 motif sequence logos were shown for γ-TuCR protein sequences retrieved from Pfam database (Microtub_assoc, Pfam id: PF07989), Spc110s and Spc72s from subphylum Saccharomycotina. The Spc72 sequences used to generate the short CM1 logo are shown in Figure 8—figure supplement 1C. The sequence logos were generated with Weblogo Figure 8. Continued on next page.
Spc110\textsuperscript{T18} is an additional Cdk1 in vivo site (Albuquerque \textit{et al.}, 2008; Keck \textit{et al.}, 2011; Lin \textit{et al.}, 2011; \textit{Figure 5—figure supplement 1}). Analysis of T18A, T18V, T18E, and T18D mutations resulted in similar in vitro and in vivo phenotypes. We have to assume that all these mutations inactivate the important SPM motif in which T18 resides. The fact that T18 of Spc110 is phosphorylated only by Cdk1-Clb2 and not by Cdk1-Clb5 and the in vivo phosphorylation of SPB-associated Spc110\textsuperscript{T18} in mitosis but not in S phase (Keck \textit{et al.}, 2011; \textit{Figure 5—figure supplement 1A-C}) indicates that this Cdk1 site is different compared to the two S phase Cdk1 sites S36 and S91 in Spc110. At least in vitro, Cdk1-Clb2 phosphorylation of Spc110\textsuperscript{T18} inactivates in a dominant manner the γ-TuSC oligomerization activity (\textit{Figure 5—figure supplement 1D,E}). Thus, it is well possible that Spc110\textsuperscript{T18} phosphorylation by Cdk1-Clb2 limits the MT nucleation activity of the SPB associated γ-TuSC. Most likely only the surplus of γ-TuSC that is not engaged in MT organization is phosphorylated and affected by Spc110\textsuperscript{T18}. This model fits with the observation that the yeast SPB of haploid cells organizes a constant number of 20 nuclear MTs during mitosis (Winey \textit{et al.}, 1995).

Co-overexpression of SPC97, SPC98, and TUB4 does not induce MT nucleation despite of γ-TuSC assembly (Pereira \textit{et al.}, 1998). Our and other studies have shown that the missing factor promoting γ-TuSC ring assembly is Spc110 (Vinh \textit{et al.}, 2002; Kollman \textit{et al.}, 2010). In vivo data indicate that the binding of γ-TuSC to Spc110, although a prerequisite, is insufficient for MT formation. For example, although the γ-TuSC component Spc97 bound to mitotic SPBs of spc110\textsuperscript{Δ} spc110\textsuperscript{Δ} cells with similar efficiency to wild type cells, the γ-TuSC in the two mutants was insufficient in full MT organization (\textit{Figure 6B, Figure 6—figure supplement 1C}). Consistent with this model, the addition of Spc110\textsuperscript{Δ} spc110\textsuperscript{Δ} to recombinant γ-TuSC induced the formation of γ-TuSC dimers at ~600 kDa (\textit{Figure 2—figure supplement 2A}; note that the protein concentration in this experiment was two-times higher than in \textit{Figure 2—figure supplement 2B}) without the formation of γ-TuSC oligomers. We therefore suggest that γ-TuSC binding to Spc110 and TuSC oligomerization are mechanistically distinct steps (\textit{Figure 10C}).

\textbf{The N-terminus of Spc98 mediates interaction with N-Spc110 and oligomerization}

The N-terminal region of Spc98 exhibits homology to other GCP3 family members including human GCP3 (hGCP3, \textit{Figure 7A, Figure 7—figure supplement 1A}). All homologues contain five predicted helical regions that are followed by an unstructured linker region before the two GRIP domains that are common to all GCP proteins (Guillet \textit{et al.}, 2011). Analysis of the structure of yeast γ-TuSC by electron microscopy localised N-Spc98 at of the base of the Y shaped structure, away from the C-Spc98 that interacts with γ-tubulin. N-Spc98 is close to the N-terminus of Spc97 and N-Spc110 (Choy \textit{et al.}, 2009; Kollman \textit{et al.}, 2010). This is consistent with yeast two-hybrid studies that identified N-Spc98 as an interactor for Spc110 (Knop and Schiebel, 1997; Nguyen \textit{et al.}, 1998). We have analyzed the importance of N-Spc98 by deleting different portions of this domain including the helices, the linker region, and both the helices and the linker. Surprisingly, the N-Spc98 fragment was not essential for the viability at 23°C (\textit{Figure 7B, Figure 7—figure supplement 1B}). The important function of this region, however, was revealed at elevated temperatures and when SAC function was impaired (\textit{Figure 7B}).

Biochemical analysis of γ-TuSC\textsuperscript{Δ} showed that its binding to Spc110\textsuperscript{Δ} and oligomerization capability by Spc110\textsuperscript{Δ} were reduced (\textit{Figure 7E–G}), although in M phase it bound with
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the same affinity for SPBs as WT γ-TuSC (Figure 7D). We suggest that the special SPB arrangement of Spc110 as hexameric units (Muller et al., 2005) compensates in part for the reduced in vitro binding of γ-TuSC SPC98Δ1–156 to Spc1101–220. This compensation may arise from cooperative interactions of γ-TuSC.
Figure 10. Role of Spc110 phosphorylation during SPB duplication. (A) MT nucleation by the SPB during the cell cycle. See 'Discussion' for details. (B) Cell cycle dependent, stimulatory phosphorylations of Spc110 by Mps1 and Cdk1-Clb5 (early S phase to early M). (C) Model for the interaction of γ-TuSC with Spc110. The Spc110 dimer interacts via the SPM and CM1 motifs with the N-terminus of Spc98 (GCP3) and possibly also with other regions of γ-TuSC. DOI: 10.7554/eLife.02208.025
with N-Spc110. Whatever the mechanism, regions of γ-tuSC in addition to the N-terminal 177 amino acids of Spc98 (Figure 7—figure supplement 1B) have to interact with Spc110. Either Spc110 also interacts with the Spc98/GCP3 core or it binds to the N-terminus of Spc97/GCP2.

γ-tuSC°Spc98A–154° was still able to oligomerize under special buffer conditions (Figure 7—figure supplement 2), which is in agreement with the observation that γ-tuSC°Spc98A–156° was functional in vivo and promoted MT nucleation, although defects were seen at elevated temperature and in mad2Δ cells. Thus, the regions that are essential for γ-tuSC oligomerization await identification.

γ-tubulin complex receptors fall into three groups: the SPM-CM1-PACT, CM1-MASC, and CM1-CM2 types

γ-tuCRs from yeast to Drosophila to human carry a conserved CM1 motif. This CM1 motif was first identified in fission yeast Mto1 (Sawin et al., 2004). Subsequently the function of CM1 in γ-tuSC binding was confirmed for Mto1, centrosomin (Drosophila), and CDK5RAP2 (human) (Zhang and Megraw, 2007; Fong et al., 2008; Samejima et al., 2008; Choi et al., 2010). The γ-tuCRs Pcp1 (fission yeast) and Spc110 (budding yeast) also contain putative CM1 motifs (Dictenberg et al., 1998; Takahashi et al., 2002; Zimmerman et al., 2004; Fong et al., 2010; Figure 8B).

In addition, we identified a yet unappreciated CM1 motif in the yeast γ-tuCR Spc72 (Knop and Schiebel, 1998; Figure 8A). Although the putative CM1 motif in Spc72 appears degenerated compared with the full CM1 of other γ-tuCRs, the conserved residues shared by Spc72-CM1 and full CM1 suggest that Spc72 contains a functional CM1. Consistently, Spc72 carries a C-terminal MASC, as is the case for other fungal CM1-only γ-tuCRs (Samejima et al., 2010). Moreover, the predicted CM1 resides in the N-terminal region of Spc72 that is essential for γ-tuSC binding (Knop and Schiebel, 1998; Usui et al., 2003).

Human PCNT falls in terms of structure and function into the γ-tuCR class (Doxsey et al., 1994; Dictenberg et al., 1998; Gillingham and Munro, 2000). Analysis of the PCNT sequence for CM1 in combination with the newly identified SPM identified a potential CM1 in the N-terminus of PCNT (Figure 8B). This is consistent with the γ-tubulin binding ability of human PCNT, which lies within the region that contains the SPM-CM1 (Takahashi et al., 2002). In addition, super-resolution analysis of human centrosomes has mapped the localization of γ-tubulin towards the N-terminus of PCNT (Lawo et al., 2012; Mennella et al., 2012; Sonnen et al., 2012). Experimental evidences are awaited to validate the CM1 motif in Spc72 and PCNT.

Our study has identified SPM as a second conserved motif amongst a subgroup of γ-tubulin complex receptors (Figure 2H), namely the yeast Spc110 homologues, fission yeast Pcp1, and human PCNT (Figure 10; Doxsey et al., 1994; Dictenberg et al., 1998; Flory et al., 2002). The distance between SPM and CM1 is around 90–100 amino acids in the Spc110 and Pcp1 type of receptors. In the mammalian PCNT subfamily it is however only around 70 amino acids. Whether these differences are important for regulation or are indicative of co-adaptation with alterations in the γ-tuSC binding sites remains to be seen.

The SPM is important for Spc110 binding to γ-tuSC and induced oligomerization of γ-tuSC (Figures 3 and 4). Based on these results we propose that SPM co-operates with CM1 to form a bipartite binding element for the γ-tuSC (Figure 10C). Interestingly, however, other γ-tubulin complex receptors such as Spc72, fission yeast Mto1, Drosophila Cnn, and human CDK5RAP2 carry only the CM1 element but no SPM motif (Figure 8). Strikingly, SPM-CM1 type of γ-tuCRs carry a C-terminal PACT domain that targets these proteins to MTOCs (Figure 8B). The CM1-only type of γ-tuCRs either contain a MASC (fungi) or a CM2 C-terminal MTOC targeting sequence (metazoa) (Figure 9B). We also noticed that in some metazoan and the fungi phylum Basidiomycota the PACT domain-containing γ-tuCRs (such as AKAP9 in human and D-PLP in Drosophila) have N-terminal regions distinctive from SPM and CM1 motifs, probably reflecting different binding partners in γ-tuRC. Altogether, we can classify γ-tuCRs in at least three classes: CM1-MASC, CM1-CM2, and SPM-CM1-PACT.

In budding and fission yeast, the CM1-MASC types of γ-tuCRs (Spc72 and Mto1) have specific functions in cytoplasmic MT organization, while Spc110 and Pcp1 (SPM-CM1-PACT) organize only nuclear MTs. This functional division is accompanied by the physical separation of the receptors by the nuclear envelope, which remains intact during the closed mitosis of both organisms. Human CDK5RAP2, myomegalin (CM1-CM2 types), and pericentrin (SPM-CM1-PACT type) are associated with the centrosome (Doxsey et al., 1994; Fong et al., 2008). However, CDK5RAP2 and myomegalin also have
functions in acentrosomal MT nucleation for example MT nucleation by the Golgi (Choi et al., 2010; Roubin et al., 2013). Thus, it may be the acentrosomal/cytoplasmic function that explains the lack of SPM element in Mto1/CDK5RAP2/myomegalin.

Budding yeast organizes MTs only with γ-TuSC and γ-TuCRs. Other organisms have additional building blocks that contribute to MT assembly, such as Mozart1 and GCP4-6 (Dhani et al., 2013; Masuda et al., 2013). However, GCP4 to 6 are not essential for MT nucleation in fission yeast and Drosophila, while GCP2/Alp4, GCP3/Alp6, and Mozart1 are essential for viability (Anders et al., 2006; Verollet et al., 2006). Puzzlingly, fission yeast Mozart1 is essential for γ-TuSC recruitment to SPBs (Masuda et al., 2013). This raises the question how budding yeast compensates for Mozart1’s function. GCP3 and γ-TuCRs of the SPM-CM1 type are conserved between organisms with or without Mozart1. It is likely that small adaptive changes in GCP3 or Spc110 have evolved to compensate for the lack of Mozart1 in budding yeast.

What could be the function of γ-TuCR in organisms that are able to assemble the more stable γ-TuRC? γ-TuCRs may recruit and activate the already assembled γ-TuRC to MTOCs via GCP interactions. In addition, γ-TuCRs could induce γ-TuSC assembly into rings that then either function without the help of additional GCPs or subsequently become stabilized by GCP4-6. In any case, considering the conservation between the SPM and CM1 binding elements of γ-TuRCs and the N-terminal region of Spc98/GCP3, we suggest that the basic principals of MT nucleation are conserved from yeast to human.

Materials and methods
Plasmid and strain constructions
A detailed list of DNA constructs and yeast strains is described in Supplementary file 1 and Supplementary file 2. SPC110 alleles were subcloned into the integration vector pRS304, and SPC98 alleles were subcloned into pRS305K (Sikorski and Hieter, 1989; Christianson et al., 1992; Taxis and Knop, 2006). Point mutations in genes were introduced by PCR-directed mutagenesis and confirmed by DNA sequencing. GST-Spc1101–220 was cloned into pGEX-5X-1 vector (GE Healthcare, UK) and His-tagged Mps1 was cloned into pET28b for expression in BL21 CodonPlus E. coli and into pFastBac1 vector for overexpression in baculovirus-insect cell system. All yeast strains are derivatives of S288c. Gene deletion and epitope tagging of genes at their endogenous loci performed using standard techniques (Knop et al., 1999; Janke et al., 2004). The red fluorescent mCherry was used to mark SPBs through a fusion with SPC42 (Donaldson and Kilmartin, 1996). For SPC97-yeGFP and SPC110-yeGFP strains, the endogenous SPC97 and the integrated SPC110 alleles were tagged with green fluorescent yeGFP. GFP-TUB1 strains were constructed using an integration plasmid (Straight et al., 1997). To remove URA3-based plasmids, transformants were tested for growth on 5-fluoroorotic acid (5-FOA) plates that select against URA3-based plasmids.

Antibodies
Affinity-purified anti-Tub4 (1:1000) and anti-Spc110 (1:1000) antibodies were described previously (Geissler et al., 1996; Spang et al., 1996a, 1996b). Anti-His-tag (1:1000) and anti-GST (1:1000) antibodies were used to detect Spc97 and Spc98 of recombinant γ-TuSC. Secondary antibodies used in semi-quantitative blotting were IRDye800- or Alexa680-conjugated anti-goat, anti-rabbit, anti-guinea pig, and anti-mouse IgGs (1:50,000; Rockland Immunochemicals Inc., Gilbertsville, PA). Phospho-specific antibodies were raised in guinea pigs and purified with immuno-affinity chromatography (1:200; Peptide Specialty Laboratories GmbH, Germany).

Protein purification
Subunits of γ-TuSC and γ-TuSC4NH-Spc98 were expressed with the baculovirus-insect cell expression system. Purification was performed as described (Gombos et al., 2013). Spc1101–220 variants with an amino-terminal GST tag were purified with Glutathione Sepharose 4B (GE Healthcare, UK) as described (Vinh et al., 2002). Proteins were eluted with 5 mM glutathione, concentrated, and run over a Superose 6 column equilibrated in HB100 to remove glutathione. Recombinant Mps1 protein was purified in lysis buffer (50 mM NaH2PO4 pH 7.5, 300 mM NaCl, 10% glycerol, 1 mM DTT, 1 mM PMSF, 1X Complete EDTA-free protease inhibitor cocktail (Roche, Canada)) with column packed with Ni-NTA Sepharose (GE Healthcare, UK). After wash with lysis buffer, protein was eluted with lysis buffer containing...
200 mM imidazole. Small aliquots for kinase assay were snap frozen with liquid nitrogen and stored at −80°C. Cdk1<sup>as1</sup>-Clb2 and Cdk1<sup>as1</sup>-Clb5 complexes were purified from yeast strains kindly given by David Morgan (see Supplementary file 2 for detail) (Loog and Morgan, 2005). CLB2-TAP and CLB5-TAP were encoded on 2-micron plasmid and driven by Gal1 promoter. After overexpression with 2% galactose, cells were harvested and lysed for TAP-tag purification as previously described (Ubersax et al., 2003).

**γ-TuSC oligomerization assay**

Purified γ-TuSC and γ-TuSC<sup>Spc98AN</sup> were mixed with or without Spc110<sup>1–220</sup> in 1:4 molar ratio (2.3:9.3) in TB150 (50 mM Tris–HCl, pH7.0 with 150 mM KCl) or BRB80 (80 mM PIPES pH6.9, 1 mM EGTA, 1 mM MgCl<sub>2</sub>) buffer as indicated in figure legends. After incubating for 1 hr at 4°C, the protein mixture was applied to a gel filtration column (Superdex 200 10/300 or Superose 6 10/300 [GE Healthcare, UK]). Elution profiles were recorded as absorbance at 280 nm. For each fraction 2% of sample volume was analyzed by SDS-PAGE. Proteins were detected by silver staining or immunoblotting.

**In vitro binding assay**

For the measurement of the binding of γ-TuSC to Spc110<sup>1–220</sup>, recombinant GST (300 nM) or GST-Spc110<sup>1–220</sup> proteins (from 0 to 300 nM or fixed 300 nM) were incubated with γ-TuSC (150 nM) in TB150 buffer in a total reaction volume of 30 μl on a rocking platform for 0.5 hr at 4°C. Glutathione-Sepharose 4B bead slurry (20 μl; GE Healthcare, UK) was then added to each reaction, followed by rocking for an additional 1 hr at 4°C. Beads were washed five times with TB150 buffer, 0.1% NP40 followed by heating in 20 μl SDS sample buffer. Input and bound proteins were analysed by immunoblotting. The signal intensities of protein bands on immunoblots were quantified with ImageJ (NIH). Signal intensities were corrected against the membrane background.

**In vitro MT nucleation assay**

Microtubule nucleation assay was performed as described (Gombos et al., 2013) with some modifications. 3 μM γ-TuSC was pre-incubated with 15 μM GST-Spc110<sup>1–220</sup> in HB100 buffer plus 12.5% glycerol for 30 min on ice. An equal volume of 20 μM bovine brain tubulin containing 4% Alexa568-labelled tubulin in BRB80 buffer with 25% glycerol was added and samples were further incubated for 30 min on ice before being transferred to 37°C for 15 min for MT polymerization. Samples were fixed with 1% formaldehyde and diluted with 1 ml cold BRB80 buffer. 50 μl aliquot of reaction mixture was sedimented onto poly-lysine-coated coverslips with a HB-4 rotor at 25,600 g for 1 hr. Samples were post-fixed with pre-cooled methanol, mounted on slides with CitiFluor mounting media, and imaged with a fluorescence microscope as described. Microtubules were counted in 20 random fields.

**In vitro kinase assay**

For the in vitro phosphorylation of Spc110<sup>1–220</sup> variants, 0.5 μg Spc110<sup>1–220</sup> substrate was incubated in kinase buffer (50 mM Tris–HCl pH 7.5, 10 mM MgCl<sub>2</sub>, 2 mM EGTA) with purified Mps1 kinase or Cdk1<sup>as1</sup> kinase co-immunoprecipitated with Clb2-TAP or Clb5-TAP. For autoradiography, 5 μCi [γ-<sup>32</sup>P]-ATP was added to the reaction mixture and the kinase reaction was performed for 30°C. The reaction was stopped by the addition of SDS sample buffer and analysed by SDS–PAGE, Simply Safe Blue staining (Invitrogen, Carlsbad, CA), and autoradiography. For phospho-specific antibody detection, 0.5 mM unlabeled ATP was used in the kinase reaction. For the γ-TuSC oligomerization assay by gel filtration, the phosphorylation reaction was stopped by addition of 15 μM 1NM-PP1 to inhibit Cdk1<sup>as1</sup> or 10 μM SP600125 (Sigma-Aldrich, St Louis, MO) to inhibit Mps1 kinase.

**Electron microscopic analysis of γ-TuSC oligomers**

To visualize γ-TuSC monomers and oligomers, proteins were stained with uranyl acetate and analyzed by electron microscopy. Briefly, we evaporated 300 or 400 mesh copper/palladium grids with a carbon layer on the copper side. To enhance the hydrophilic affinity of the carbon layer, grids were glow discharged for 30–45 s directly before the preparation started. The protein solution was incubated for 30 s on the carbon mesh grids at room temperature. The grids were washed and incubated in 2% uranyl acetate for 4 min and blotted on Whatman 50 paper. The images were taken in low dose modus with an under-focus between 0.8 and 1.5 μm. Particles were viewed with a CM120 electron microscope (Philips Electronics NV, Eindhoven, Netherlands), which was operated...
at 120 kV. Images were captured with a CCD camera (Keen View, Soft Imaging systems, Germany) and viewed with Digital Micrograph Software.

**Phosphopeptide enrichment and mass spectrometry**

Phosphopeptides were identified according to Villen et al. (Dephoure et al., 2008). In brief, SDS-PAGE separated Spc110 was cut out and subjected to trypsin digestion. After digest, phosphopeptides were enriched by IMAC, starting with 25 µl of PHOS-Select beads (Sigma, Germany). Enriched phosphopeptides were eluted and desalted by C18 columns (Stage tips). Peptides were analyzed by LC-MS/MS (Orbitrap Elite, Thermo Fisher Scientific, Waltham, MA) and data were processed using Thermo proteome discoverer software (1.4). Phosphorylation site localization was performed on the Mascot results using PhosphoRS.

**Growth assay**

Yeast cells in the early log phase were adjusted to an OD$_{600}$ of 1 with PBS. 10-fold serial dilutions of cells were spotted onto the indicated plates and incubated as indicated in the figure legends.

**Quantification of the Spc97-GFP, Spc110-GFP, and GFP-Tub1 signal at SPBs**

Asynchronous cells were grown in filter-sterilized YPD with additional 0.1 mg/l adenine (YPAD) to an OD$_{600}$ of 0.3 at 23°C for 3 hr and then shifted to 37°C for 1 hr. Cells were directly sampled for image acquisition. For the image acquisition, z-stack images with 21 0.3 µm steps (2 × 2 binning) were acquired at 37°C with a DeltaVision RT system (Applied Precision, UK) equipped with FITC (fluorescein isothiocyanate), TRITC (tetramethyl rhodamine isothiocyanate), and Cy5 filters (Chroma Technology, Bellows Falls, VT), a 100x NA 1.4 plan Apo oil immersion objective (IX70; Olympus, Japan), and a CCD camera (CoolSNAP HQ; Roper Scientific, Tucson, AZ). Images were processed and analyzed in ImageJ (NIH).

The quantification of the mean background intensity and mean fluorescence intensity of Spc97-GFP and Spc110-GFP signals at SPBs was performed on planes having SPBs in focus. Spc97-GFP or Spc110-GFP intensity within the 3 × 3 pixel-area covering a single SPB or two unsplit SPBs was measured. For GFP-Tub1 at SPBs, GFP-intensity within a 3 × 3 pixel-area surrounding the SPB was measured. Unsplit SPBs were measured together, whereas split SPBs were measured separately. The standard error (SEM) for each data set (n = 50) and level of significance were determined by one-way ANOVA with Turkey’s multiple comparisons’ test.

**Time-lapsed live cell imaging**

Cells were synchronized with α-factor for 1.5 cell cycles and then immobilized onto glass-bottom dishes. Dishes were prepared by incubating them with 100 µl concanavalin A solution (6% concanavalin A, 100 mM Tris-Cl, pH 7.0, and 100 mM MnCl$_2$) for 5 min and subsequently washed with 300 µl of distilled water. Yeast cells were allowed to attach to the dishes for 5–15 min at 30°C. The α-factor blocked cells were then released by two washes with 2 ml of prewarmed SC medium. Image acquisition was started 10–15 min after release from α-factor. Conditions for imaging were as follows: 15 stacks in the FITC channel, 0.1-s exposure, 0.3-µm stack distance, one reference image in bright field channel with a 0.05-s exposure, and 61 frames in total every 3 min. Images were quantified by measuring the integrated density of the sum of projected videos for the region of interest (ROI) around the SPBs and a background region selected from the periphery of the analyzed regions. The mean intensity of the background was subtracted from the ROI. To correct for acquisition, bleaching signal intensities were divided by a bleaching factor. The bleaching factor was determined from the mean of three very short videos that had been generated with the same image acquisition conditions. The data points of bleaching factor were fitted with non-linear one-phase decay equation.

**Nocodazole washout assay (MT regrow assay)**

Cells (5 x 10$^6$ cells/ml) were pre-grown at 23°C in filter-sterilized YPAD. Cells were arrested in G1 by treatment with 10 µg/ml α-factor (Sigma-Aldrich, Germany) for 3 hr at 23°C until >95% of cells showed a mating projection. G1 arrested cells were released into media with hydroxyurea to arrest cells in S-phase for 2 hr. The culture was then shifted to 37°C and nocodazole was added. After 1 hr, nocodazole was removed by exchanging media with YPAD containing hydroxyurea. After removal of nocodazole, cells were sampled at 0, 30, and 60 min and fixed with 4% paraformaldehyde. Image acquisition and GFP-Tub1 quantification were performed as described in previous section.
Trichloroacetic acid (TCA) extraction of yeast cells
To measure the expression level of Spc110 variants and GFP-Tub1 in vivo, whole cell lysates were prepared for SDS-PAGE and immunoblotting (Knop et al., 1999; Janke et al., 2004). 2–3 OD_{600} of late-log phase liquid culture were resuspended in 0.2 M NaOH and incubated on ice for 10 min. 150 μl 55% (wt/vol) TCA was added and the solutions were mixed and incubated for 10 min on ice. After centrifugation the supernatant was removed. The protein pellet was resuspended in high urea (HU) buffer (8 M urea, 5% SDS, 200 mM Tris–HCl pH 6.8, 0.1 mM EDTA, 100 mM DTT, bromophenol blue) and heated at 65°C for 10 min. One-fifth of total sample amount was loaded for SDS-PAGE and western blotting.

Pull-down of recombinant Spc110 protein
For in vivo phospho-Spc110 detection, strains with integrated SPC110-yeGFP or SPC110-TAP alleles were lysed in binding buffer (25 mM HEPES, pH 7.4, 150 mM NaCl, 1 mM EDTA, 1 mM DTT, 50 mM NaF, 80 mM β-glycerophosphate, 0.2 mM Na_{2}VO_{4}, and protease inhibitors) lysed with acid-washed glass beads (Sigma-Aldrich, Germany) in a FastPrep FP120 Cell Disrupter (Thermo Scientific, Germany). Cell lysates were incubated with 0.1% Triton X-100 for 15 min and then clarified by centrifugation at 10,000×g for 10 min. To further solubilize the pellet with insoluble SPBs, binding buffer containing 0.5 M NaCl and 1% Triton X-100 was added to resuspend the pellet and incubated for 40 min. GFP-binder protein (Rothbauer et al., 2008) conjugated Sepharose 4B slurry or IgG-conjugated Dynabeads was added into supernatants and rocked at 4°C for 2 hr, followed by five times of washing with binding buffer. Pull-downed Spc110-GFP or Spc110-TAP was eluted by heating with SDS sample buffer and analyzed by SDS-PAGE and immunoblotting or mass spectrometry.

Bioinformatic analysis
Protein sequences of Spc110 and Spc98 and their homologues in selected organisms were aligned with MAFFT algorithm built in Jalview software (Waterhouse et al., 2009; Katoh and Standley, 2013; Figures 2H and 7A, Figure 2—figure supplement 6, Figure 8—figure supplement 1). The threshold of conservation was set as 20% and highlighted. The relative frequency of each amino acid in each position of SPM and CM1 motifs was visualized using the WebLogo 2.0 (Crooks et al., 2004; Figure 2H, Figure 8, Figure 9, Figure 2—figure supplement 6, Figure 8—figure supplement 1). To demonstrate the occurrence of SPM and CM1 motifs in selected members of γ-TuSC receptor family, 47 protein sequences covering the putative CM1 motifs were subjected to MEME motif scanning analysis (Bailey et al., 2009; Figure 8B).

Jpred3 and Disopred predicted the secondary structure of the N-terminal domain of Spc98 and human GCP3 (Ward et al., 2004; Cole et al., 2008). Domain positions and protein interacting regions of Spc110 and Spc98 (Figure 1A, Figure 7—figure supplement 1A) were illustrated according to γ-TuSC binding studies of Spc72- and Spc110-truncated forms and to the yeast-two-hybrid studies (Sundberg and Davis, 1997; Knop and Schiebel, 1998; Usui et al., 2003).

For the identification of full CM1 motif on Spc110s and degenerative CM1 on Spc72, CM1 motif containing γ-TuCR protein sequences were retrieved from Pfam database (Microtubul_assoc, Pfam id: PF07989) and multiple-aligned with Spc110s and Spc72s from species of subphylum Saccharomycotina by MAFFT multiple sequence alignment algorithm. To define the CM2 motif pattern, CM2 sequences from human CDK5RAP2 and Drosophila Cnn were used to retrieve protein sequences containing putative CM2 with HMMER server (Finn et al., 2011). The retrieved protein sequences were multiple-aligned with CDK5RAP2 and Cnn by MAFFT algorithm and the multiple-aligned CM2 sequences were subjected to Weblogo 3.0 server to visualize the pattern.

Statistical analysis
Statistical analysis of fluorescent intensities, immunoblotting intensities, and in vitro microtubule numbers was performed with GraphPad Prism 6.1. One-way ANOVA with Turkey’s multiple comparisons’ test was used to compare samples and to obtain adjusted p values. The number of repeated experiments and sample size are indicated in figure legends. No statistical method was used to predetermine sample size. The experiments were not randomized. The investigators were not blinded to allocation during the experiments and outcome assessment. In general the data showed normal distribution.

Acknowledgements
This work was supported by a grant of the Deutsche Forschungsgemeinschaft (Schi295-3-2). Dr David Morgen is acknowledged for yeast strains. We thank U Jäkle and S Heinze for excellent technical
support. TL is a member of the international graduate school HBIGS and was supported by a fellowship of the Graduiertenkolleg Regulation of Cell Division.

Additional information

Funding

| Funder                                | Grant reference number | Author                                      |
|---------------------------------------|------------------------|---------------------------------------------|
| Deutsche Forschungsgemeinschaft       | Schi295-3-2            | Tien-chen Lin, Annett Neuner, Yvonne T Schlosser, Elmar Schiebel |
| Deutsche Forschungsgemeinschaft (DFG) | SFB1036                | Annette ND Scharf, Lisa Weber               |

The funders had no role in study design, data collection and interpretation, or the decision to submit the work for publication.

Author contributions

TL, Conception and design, Acquisition of data, Analysis and interpretation of data, Drafting or revising the article; AN, Single particle analysis with electronic microscopy, Acquisition of data, Analysis and interpretation of data; YTS, Characterization of phenotypes of spc98 mutant cells, Acquisition of data, Analysis and interpretation of data; ANDS, LW, MS analysis of phosphorylated peptides, Acquisition of data, Analysis and interpretation of data; ES, Conception and design, Drafting or revising the article

Additional files

Supplementary files

- Supplementary file 1. Plasmids used in this study.
  DOI: 10.7554/eLife.02208.026
- Supplementary file 2. Strains used in this study.
  DOI: 10.7554/eLife.02208.027

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