**CM1-driven assembly and activation of Yeast γ-Tubulin Small Complex underlies microtubule nucleation**

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**ABSTRACT**

Microtubule (MT) nucleation is regulated by the γ-tubulin ring complex (γTuRC), conserved from yeast to humans. In Saccharomyces cerevisiae, γTuRC is composed of seven identical γ-tubulin small complex (γTuSC) sub-assemblies which associate helically to template microtubule growth. γTuRC assembly provides a key point of regulation for the MT cytoskeleton. Here we combine cross-linking mass spectrometry (XL-MS), X-ray crystallography and cryo-EM structures of monomeric and dimeric γTuSC and open and closed helical γTuRC assemblies in complex with Spc110p to elucidate the mechanisms of γTuRC assembly. γTuRC assembly is substantially aided by the evolutionarily conserved CM1 motif in Spc110p spanning a pair of adjacent γTuSCs. By providing the highest resolution and most complete views of any γTuSC assembly, our structures allow...
phosphorylation sites to be mapped, suggesting their role in regulating spindle pole body attachment and ring assembly. We further identify a structurally analogous CM1 binding site in the human γTuRC structure at the interface between GCP2 and GCP6, which allows for the interpretation of significant structural changes arising from CM1 helix binding to metazoan γTuRC.

**INTRODUCTION**

The microtubule (MT) cytoskeleton plays an essential role in the spatio-temporal control of eukaryotic cellular organization, cytoplasmic transport and chromosome segregation during mitosis (Desai and Mitchison, 1997). The organization and function of the cytoskeletal network is tightly controlled by regulating the rate and location of nucleation, as well as MT polymerization kinetics and stability (Akhmanova and Steinmetz, 2015; Howard and Hyman, 2009; Teixidó-Travesa et al., 2012).

In most cells, MT nucleation occurs primarily at microtubule organizing centers such as centrosomes or spindle pole bodies and is dependent on the universally conserved γ-tubulin ring complex (γTuRC) (Luders and Stearns, 2007). In budding yeast, only homologues of GCP2 and GCP3 (Spc97p and Spc98p) and γ-tubulin (Tub4p) are present (Vinh et al., 2002). Spc110p, a distant pericentrin homologue, recruits this complex to the nuclear face of the spindle pole body (SPB), while Spc72p recruits it to the cytoplasmic face (Knop and Schiebel, 1997, 1998; Nguyen et al., 1998).

In metazoans and plants, the γTuRC is recruited to MT nucleation sites as a large, pre-formed ring-shaped 2.2 MDa complex (Teixidó-Travesa et al., 2012). The metazoan γTuRC is composed of 14 γ-tubulins, and a smaller number of the γ-tubulin binding proteins, GCP2-6, as well as other accessory proteins. GCP2 and GCP3 along with 2 copies of γ-tubulin form a stable 300 KDa subcomplex (γTuSC). Recent cryo-EM structures have revealed that five copies of the GCP2/3 γTuSC are integrated into the metazoan γTuRC along with a GCP4/5 and a GCP4/6 pseudo-γTuSC as well as the other accessory proteins (Liu et al., 2020; Murphy et al., 2001; Oegema et al., 1999; Wieczorek et al., 2020). While sharing only ~15% homology and varying in size from 70 kDa to 210 kDa, GCP2-6 share a conserved core of 2 grip domains (Guillet et al., 2011). Structural and biochemical studies
have shown that the N-terminal domain drives lateral association between GCPs, while the C-terminal domain binds to γ-tubulin (Choy et al., 2009; Farache et al., 2016; Greenberg et al., 2016; Guillet et al., 2011; Kollman et al., 2015).

Previous moderate-resolution cryo-EM structural studies (8 Å) had shown that γTuSCs complexed with the N-terminal domain of Spc110p self-assemble into filaments (hereafter γTuRC) having 6.5 γTuSCs/turn, thereby presenting 13 γ-tubulins to template 13-protofilament MTs (Kollman et al., 2010, 2015). Although close to MT symmetry, the γ-tubulins within each γTuSC were too far apart to correctly match the MT lattice, adopting an open conformation. The relevant in vivo conformation was determined by cryo-tomography and sub-volume averaging, clearly showing a MT-matching geometry at the yeast SPB, suggesting that γTuSC closure might be an important regulatory step (Kollman et al., 2015). To validate this hypothesis, γ-tubulin was engineered with disulfides to stabilize a closed MT-like conformation (γTuRCSS), resulting in significantly enhanced MT nucleation (Kollman et al., 2015). This also had the benefit of improving the cryo-EM map (6.5 Å) such that an initial pseudoatomic model (Greenberg et al., 2016; Kollman et al., 2015) could be built based on the crystal structure of human γ-tubulin (Aldaz et al., 2005; Rice et al., 2008) and the distant and much smaller (75 kDa vs 97 or 98 kDa) human GCP4 (Guillet et al., 2011; Kollman et al., 2015). These structures suggest a hierarchical model of γTuSC activation, with γTuSC assembling at the SPB into an open slightly active conformation prior to closure.

Biochemical studies on the role of Spc110p in γTuRC assembly revealed that higher-order oligomerization of Spc110p and its binding to γTuSCs was required to overcome the intrinsically weak lateral association of γTuSCs at physiologically relevant γTuSC concentrations (Kollman et al., 2010, 2015; Lyon et al., 2016). Deletion studies identified that independent of oligomerization, removal of Spc110p residues 1-111 (Spc110p1-111) was lethal in vivo but only slightly compromised γTuRC assembly in vitro, perhaps suggesting an essential regulatory function. By contrast, deletion of the subsequent centrosomin motif 1 (CM1, Spc110p117-146) additionally abolished γTuRC assembly in vitro. Supporting the need for precise regulation of γTuRC assembly and function, all the components of the γTuSC, as well as Spc110p and Spc72p are phosphorylated at multiple
sites in a cell-cycle dependent manner (Keck et al., 2011). Many of these phosphorylation events, particularly in γ-tubulin or Spc110p, have been implicated in cellular viability, spindle morphology, or shown to effect γTuRC assembly (Fong et al., 2018; Huisman et al., 2007; Keck et al., 2011; Lin et al., 2011, 2014; Lyon et al., 2016; Peng et al., 2015; Vogel et al., 2001). Together these data suggest a hierarchical model of γTuSC activation; with γTuSC assembling at the SPB into an open γTuRC that would be further activated by closure. However, owing to the lack of structural data, the molecular mechanisms by which Spc110p facilitates γTuRC assembly and activation have remained unclear.

Here we use chemical crosslinking coupled with mass spectrometry (XL-MS) and X-ray crystallography to identify and determine the structure of the N-terminal coiled-coil of Spc110p-bound in γTuSC filaments previously observed in cryo-EM reconstructions. The combined data show that a unique pose of the coiled-coil satisfies most of the XL-MS restraints. Furthermore, integrative modeling indicates that only residues N-terminal to the coiled-coil from a single protomer were required to satisfy the majority of the crosslink restraints, suggesting an asymmetric mode of Spc110p binding to γTuRC.

We present cryo-EM structures of monomeric and dimeric γTuSCs at near-atomic resolution, as well as higher-resolution cryo-EM structures obtained from γTuRC filaments in the open and closed conformations. These have allowed de-novo model building of unknown regions and reinterpretation of significant portions of γTuSC structure. Our atomic models of γTuSCs in different assembly and conformational states provide insights into the assembly and activation mechanisms of γTuRC required for microtubule nucleation and reveal how N-terminal regions of Spc110p, notably CM1, facilitate γTuRC assembly. Many of the annotated phosphorylation sites had fallen in regions of γTuSC not represented in the previous homology model, thus the new structure provides a powerful atomic framework for understanding the importance and mechanism of regulatory modifications.

RESULTS

Defining Spc110p:γTuSC interactions by XL-MS
Our previous 6.9 Å cryo-EM reconstruction derived from helical filaments of Spc110p bound to an engineered closed conformation of γTuRCSS revealed an ~40 residue long segment of coiled-coil density contacting the N-terminal region of Spc97p. The limited resolution prevented rigorous assignment of this density to any particular portion of Spc110p. Given its importance for γTuRC assembly, the coiled coil seemed likely to correspond to Spc110pCM1 or alternatively to the 45-residue segment (Spc110p164-208) predicted with high probability to be a coiled-coil (the N-terminal coiled-coil, or Spc110pNCC; see Fig. 1A). Beyond this ambiguity, previous maps also lacked any density for the non-coiled-coil regions of Spc110p1-220 known to be biochemically important and absolutely required for viability (Lyon et al., 2016).

To define the important interaction interfaces between Spc110p and γTuSC, we utilized XL-MS with the same Spc1101-220 construct used for the cryo-EM as well as a longer Spc1101-401 construct (Fig. S1). We observed a significant number of EDC and DSS crosslinks between the N-terminal portions of Spc97p and Spc110pNCC (Fig. S1). Thus, this region and not CM1 was responsible for the coiled-coil-γTuSC interaction apparent in the cryo-EM map. As shown below, CM1 binds at an inter-TuSC cleft that spans adjacent TuSCs.

**The Spc110p NCC164-208 binds to γTuSC at the N-terminal regions of Spc97p**

Due to the limited resolution of the previous cryo-EM reconstruction, the derived atomic model of the coiled-coil contained only the peptide backbone (Greenberg et al., 2016; Kollman et al., 2015). Motivated by the crosslinks observed between the Spc110pNCC and γTuSC, we sought a higher-resolution structure of the NCC region via X-ray crystallography. Previous work indicated that Spc1101-220 is only weakly dimeric (Lyon et al., 2016). Using the proven strategy of fusing weakly interacting coiled-coils with stabilizing domains (Andreas et al., 2017; Frye et al., 2010; Klenchin et al., 2011), we found that an N-terminal fusion of Spc110p164-207 with Xrcc4 produced high yields of soluble protein. The Xrcc4-Spc110p164-207 construct crystallized in a variety of conditions, diffracted to 2.1 Å, and enabled phases to be obtained by molecular replacement using Xrcc4 as a search model. As expected, the electron density map was consistent with a coiled-coil, with interpretable density for Spc110p164-203 residues (Fig. 1B). When docked into the 6.9 Å cryo-EM map, the X-ray model occupied most of the alpha-helical cryo-EM
density. Importantly in the docked conformation the majority of the unique DSS (4/5) and EDC(7/9) crosslinks (Figs. 1C, S1E) were satisfied.

To better understand where the non-coil regions of Spc110p might interact, we used integrative modeling. (Figs 1D, S2, S3, Supplementary Computational Methods) (Alber et al., 2007; Greenberg et al., 2016; Kollman et al., 2015; Rout and Sali, 2019; Russel et al., 2012).

We first considered a single γTuSC bound to an Spc110p1-220 dimer (Fig. S2A). Using a combination of the previous cryo-EM-based γTuSC pseudoatomic model (Greenberg et al., 2016; Kollman et al., 2015), the X-ray structure of Spc110p164-207, and representing the rest of γTuSC and Spc110p1-220 by flexible strings of beads representing the amino-acid chain, approximately three thousand good-scoring models were obtained satisfying the crosslinks and stereochemistry (excluded volume and sequence connectivity). These models were clustered based on structural similarity (Fig. S4) and ~98% of the models were well represented by a single cluster. This cluster satisfied (see Supplementary Methods) 90.9% and 92.8% of the γTuSC-Spc110p1-220 GCN4 EDC and DSS crosslinks, respectively. While the modeling results described below were based on the γTuSC-Spc110p1-220 GCN4 crosslinks, similar results were obtained in all cases using the γTuSC-Spc110p1-401 GST crosslinks.

Consistent with visual inspection of the crosslinks, the localization probability density map from the most occupied cluster (Fig. S2A) indicated that Spc110p1-163 extended from the Spc110pNCC along the Spc97p-Spc98p interface towards γ-tubulin and the C-termini of Spc97p/98p. The precision of the model was insufficient to distinguish separate paths for the non-coiled coil regions of each protomer within the Spc110p dimer. Consequently, we also considered a model containing Spc110p1-163 from a single protomer which almost equally satisfied the crosslink restraints and indicated a similar path (Fig. S2B). As the localization probability map suggested that the two Spc110p protomers might follow different paths, with one path extending towards the adjacent γTuSC, we also modeled two adjacent γTuSCs, each bound to an Spc110p1-220 dimer (Fig. 1D). By considering adjacent γTuSCs, the predicted path spans from the N-terminus of Spc97p of one γTuSC before proceeding towards the Spc98p from the adjacent γTuSC and binding in the space between the two γTuSCs. There is also a component that extends towards the
Spc97p C-terminus and γ-tubulin (Fig. 1D). Together these results suggest a complex path interacting with multiple γTuSCs taken by at least one of the two Spc110p N-termini.

**High resolution filament structures reveal previously uninterpretable regions of γTuSC.**

The observed binding site between Spc110pNCC and γTuSC explains how Spc110p oligomerized at spindle poles can stimulate γTuRC assembly by increasing the local γTuSC concentration. However, this fails to explain the critical biochemical and in vivo functional importance of residues N-terminal to the Spc110pNCC, such as the Spc110pCM1 region for γTuRC assembly and microtubule nucleation (Lyon et al., 2016). While the crosslinking and integrative modeling data suggested a physical basis for these observed functional roles, the actual path and interactions taken by Spc110p1-163 were unknown. Realizing that this would require much higher resolution of Spc110p-γTuSC interactions, we focused on reevaluating both the “open” (γTuRCWT) and disulfide trapped “closed” (γTuRCSS) filaments containing Spc110p (Kollman et al., 2010, 2015) with modern cryo-EM methods. As with previous studies, a combination of local helical and conformational inhomogeneities led to significantly worse resolution in the Spc97p/Spc98p C-terminus/γ-tubulin region compared to the N-terminal and middle domains of Spc97p/Spc98p, particularly for the γTuRCWT filaments. To improve the resolution, we performed symmetry expansion followed by focused classification of segments containing 3 adjacent γTuSCs. The resulting reconstructions were at a resolution of 3.6 Å and 3.0 Å for the γTuRCWT and γTuRCSS filaments, respectively (Figs. 2AB,S5,S6, Table S1). The significantly increased resolution (Figs. S5,S6) allowed us to greatly improve upon previously published models of Spc97p, Spc98p, and γ-tubulin. Overall, we were able to build 712 a.a of Spc97p (Fig. S7A) (87%) 674 a.a of Spc98p (Fig. S7B) (80%), 453 a.a. of γ-tubulin (96%), and 95 a.a of Spc110p1-220 (43%).

Previous high-resolution crystal structures of γ-tubulin have shown that it adopts a bent-like state when not in complex with GCPs, independent of its nucleotide state (Aldaz et al., 2005; Rice et al., 2008). This raised the possibility that γ-tubulin might change
conformation upon assembly into γTuSC. While the changes are small, in our structures, γ-tubulin adopts a conformation distinct from the previously observed bent human γ-tubulin or the yeast tubulin straight conformations (Fig. S8A). In the assembled state, the C-terminal portion of the γ-tubulinH6-H7 loop that most defines the interface with the incoming α-tubulin adopts a conformation similar to a straight yeast β-tubulin, likely potentiating MT formation (Fig. S8A).

In looking for a potential cause for the altered γ-tubulinH6-H7 loop conformation, there was one notable difference in the Spc/γ-tubulin interface. The γ-tubulinT7 loop in assembled γ-tubulin moves such that it now more closely resembles the β-tubulinT7 loop of an assembled β-tubulin (Fig. S8B). The γ-tubulinT7 loop is pinned between a loop (Spc98pH15-16/Spc97pH16-17) located at the N-terminus of a small domain in Spc97 and Spc98p and the adjacent C-terminal helical bundles (Spc98pH22-23/Spc97pH26-27). These results suggest that although subtle, assembly of yeast γ-tubulin into a γTuSC may help promote a more MT-like conformation, facilitating nucleation.

Previous structures of yeast GCPs and their assemblies suggested that the interface between the GCPs was largely formed from the two N-terminal helical bundles. Our high-resolution structures allow us to resolve large divergent N-terminal insertions and extensions present in both Spc97p and Spc98p, but absent in the shorter GCP4 “core” structure, which contribute to the intra- and inter-TuSC interfaces.

The GCP intra-γTuSC interface extends the entire length of the two N-terminal helical bundles of Spc97p and Spc98p, and also features significant contacts by the newly resolved N-terminal extensions (Fig. 2C). Of the residues newly modeled, Spc97p1-54,81-89 and Spc98p163-179 contribute an additional ~3600 Å² of buried surface area to the N-terminal interface. In addition, a previously unmodeled 33-residue insertion in the middle of Spc98p (Spc98p672-704), between helices Spc98pH23 and Spc98pH24, folds into a pair of strands, contributing an additional ~1900 Å² of surface area. In the closed state, there is a small contact patch between the N-terminal region of Spc98pH27 and Spc98pH19. Thus, while the much shorter GCP4 structure, that formed the basis of previous modeling efforts suggested well-conserved N-terminal interactions, it is clear that a very large part of intra-γTuSC stabilization (~5400 out of ~8000 Å², total interface) arises from insertions in
Spc97p and Spc98p, suggestive of very tight binding. This suggests γTuRC assembly, particularly of smaller GCPs, is stabilized using non-TuSC components in metazoans. (Liu et al., 2020; Wieczorek et al., 2020)

In contrast, the inter-γTuSC interface is much more limited in scope (total surface area ~2900Å²) and is mainly composed of two smaller, largely hydrophilic contact patches located at the three N-terminal helical bundles, in addition to a small contact between Spc97pK790 and Spc98pY510 (Fig. 2D). The set of contacts at the N-terminal helical bundle forms a small hydrophobic core, whereas the other γTuSC interface involves very few hydrophobic residues. The limited inter-γTuSC interface explains why γTuSCs fail to assemble under physiological concentrations (Kd ~2 µM), and thus must rely on a combination of CM1 interactions (see below) and avidity effects provided by Spc110p oligomerization (Lyon et al., 2016).

**Spc110p CM1 facilitates γTuRC assembly by binding at the inter-γTuSC interface**

As before (Fig. 1B), we observed coiled-coil density for the Spc110pNCC in our higher resolution maps. Given the observed pitch of the coiled-coil in the crystal structure, as well as density for larger side chains, we were able to assign the register of the NCC (Fig. S9). To assess the path of Spc110p, we generated a difference map between our experimental density maps and an atomic model for γTuRC which did not include Spc110p atoms. This difference map should contain density for Spc110p, and any regions not included in the atomic model. Indeed, the difference map revealed clear density extending from the NCC to a helical density that spans the inter-γTuSC interface and beyond (Fig. 3A). Based on the side-chain features, we were able to unambiguously assign CM1117-141 to the helical inter-γTuSC density (Fig. 3B). In addition, we were able to model additional preceding and following residues (Spc110p112-150), with the N-terminal 8 amino acids making extensive contacts with Spc98p strand Spc98pS3 and the Spc98pS3-S4 loop (Figs. 3C, S10A). While the density connecting the Spc110pCM1 helix with Spc110pNCC was at lower resolution, we were able to build the connecting loop. Interestingly, a pair of helix-dipole/hydrogen bond interactions augment binding of the CM1 helix with Spc98p, with Spc98pD542 hydrogen bonding with the N-terminus of the Spc110pCM1 helix, and Spc110pK120 hydrogen bonding.
with the C-terminus of helix Spc98p ${}^{169}$ (Fig. S10A). On Spc97p, the C-terminus of the Spc110p $^{CM1}$ helix interacts with helices Spc97p $^{123}$ and Spc97p $^{128}$ and the loop C-terminal to Spc97p $^{121}$, as well as the insertion between Spc97p $^{17}$ and Spc97p $^{9}$ at the N-terminus of Spc97p $^{88}$ (Figs. 3C, S10B). While we were unable to trace residues Spc110p 1-111 in our structure, numerous crosslinks map to the region between Spc97p and Spc98p and γ-tubulin at the intra-γTuSC interface (Fig. S2C). These residues may therefore be involved in contacts facilitating activation and closure.

Together, our data reveal that one protomer of Spc110p $^{112-207}$ within each Spc110p $^{1-220}$ dimer adopts a complex path across two γTuSCs, while the Spc110p $^{112-165}$ region of the second protomer is unresolved. For the first time, this defines the molecular role of the conserved CM1 motif. Beginning with Spc110p $^{NC}$ (Spc110p $^{164-208}$) bound to the N-terminus of Spc97p near the intra-γTuSC interface, it next interacts with Spc98p and then weaves a path along the surface of Spc97p. From there, the CM1 helix binds across the inter-γTuSC interface to Spc98p on the adjacent γTuSC. After that, it continues along the surface of Spc98, then turns towards the Spc97p C-terminus ending near γ-tubulin. (Fig. 3AC, S2D). Integrating these data generates a hypothetical continuous path across two γTuSC subunits (Fig S2D). This is in quite good agreement with modeling predictions.

To assess generality of the observed CM1 binding mode, we mapped conservation of the CM1 motif and its binding sites on Spc97p and Spc98p (Figs. S11AB). Of note, the more C-terminal portion of CM1 that binds to Spc97p is better conserved than the N-terminal portion that binds to Spc98p (Fig. S11C). In keeping with this, the CM1 binding site on Spc97p is also highly conserved (Fig. S11B). However, despite the limited conservation of the N-terminal portion of CM1, its binding site on Spc98p is well conserved in Spc98p/GCP3 homologues throughout eukaryotes (Fig. S11B), attesting to its importance. Close inspection of the structure provides a molecular explanation: many of the interactions in this region are via CM1 backbone contacts and are thus less dependent on the precise CM1 sequence.

**Conformational changes of Spc97p and Spc98p during assembly**
To better resolve fundamental questions about the molecular basis for γTuRC assembly and activation, we determined the cryo-EM structure of unassembled γTuSC, without Spc110p, from images of frozen-hydrated single particles. At the concentration of ~1 µM used in data collection, micrographs and 2D classes show a mixture of γ-TuSC monomers and dimers, with a small number of larger oligomers (Fig. S12). We were able to obtain a structure of the γTuSC monomer at ~3.7Å, and of a γ-TuSC dimer at ~4.5Å resolution (Figs. S13, S14, Table S1). The γTuSC dimer is formed from two γ-TuSCs in lateral contact using the same interface as observed in the γTuRC:Spc110p filament structures, but as expected lacks density for both the Spc110pNCC and the Spc110pCM1 helix.

In order to assess the changes that occur during assembly of monomeric γTuSCs into the γTuRC and the subsequent closure, we aligned the N-terminal two helical bundles of Spc97p and Spc98p (residues Spc97p52-276 and Spc98p170-342). This alignment allows for a concise description of the joint conformational changes in both proteins that occur as γTuSCs assemble into γTuRCs and the subsequent closure required for microtubule nucleation (Fig. S15, Movies S1,2,3).

During the transition from monomer to assembled open state (as seen in γTuRCWT), the γ-tubulins move in the same overall direction, approximately orthogonal to the plane of the Spc97p/Spc98p contact interface (Fig. S15A). The center of mass of the γ-tubulins shifts ~13.9 Å and ~15.6 Å when bound to Spc97p and Spc98p, respectively as a result of twisting the helical bundles in Spc97p and Spc98p. All of the conserved contacts in Spc97p and Spc98p observed in assembled γTuSC filaments occur in the N-terminal three helical bundles. Notably, much of the bottom three helical bundles show only minor changes when assembling to the open state. The dominant changes occur on loop Spc98pH10-S1 in the middle contact, which moves ~4.1 Å, and at the N-terminus of helix Spc98pH11, involved in the top contact, which moves ~6.6 Å. The large conformational change in Spc98p required to create these contacts appears to play a significant role in driving the conformational changes during assembly.

*The transition from the open γTuRCWT to the closed γTuRCCSS*
During the transition from the assembled open $\gamma$TuRC\textsubscript{WT} to the engineered $\gamma$TuRC\textsubscript{SS} closed conformation (Figs. 4, S15B), the $\gamma$-tubulins on Spc97p and Spc98p slide past each other in roughly opposite directions relative to the Spc97p/Spc98p interface, undergoing translations of $\sim$6.9 Å and $\sim$7.6 Å respectively (Fig. S15B). In addition, the Spc98p bound $\gamma$-tubulin undergoes a twisting motion of $\sim$5-6 degrees. During these conformational changes, the inter-$\gamma$TuSC contacts make only minor alterations, most of which are limited to the N-terminal three helical bundles of Spc97p and Spc98p. These undergo complex tilting and twisting motions. Overall, these conformational changes alter the pitch and twist of the $\gamma$TuRC assemblies from $\sim$140Å/turn and 54.5° in the open state to $\sim$132Å/turn and 55.1° in the closed state (Fig. 4AB,DE).

Excising a full turn in our $\gamma$TuRC filament structures containing seven $\gamma$TuSC subunits provides a good model for an isolated $\gamma$TuRC as it might bind at the SPB. This reveals that within each $\gamma$TuRC there are only six complete CM1 binding sites, the last one being interrupted at the end of the ring. This in turn suggests that only six Spc110p molecules need to be bound to a $\gamma$TuRC in vivo to stabilize the full ring. This helps explain the apparent symmetry mismatch between the underlying hexameric organization of Spc42p within the SPB and the heptameric $\gamma$TuRC. The geometry is such that the Spc110p\textsuperscript{NCC} binding site most proximal to the SPB would be empty (Fig. 4C).

Finally, by local 3D classification we observed that a closed state is populated in our $\gamma$TuRC\textsubscript{WT} data (Fig. S5, Table S1). While not identical to the disulfide cross-linked closed state, the differences are minimal indicating that the conformational changes observed at highest resolution in $\gamma$TuRC\textsubscript{SS} are representative of those occurring in the closed WT $\gamma$TuRCs (Fig. S15C). The fact that the WT closed state can occur spontaneously and is sampled in our open population suggests that, in the presence of Spc110p, the energy differences between the open and closed states are not large.

**Mapping phosphorylation sites on the $\gamma$TuRC suggests regulatory roles**

The $\gamma$TuSC is heavily phosphorylated in a cell-cycle dependent manner, and perturbing phosphorylation has been shown to affect spindle morphology (Fong et al., 2018; Keck et al., 2011; Lin et al., 2011; Peng et al., 2015; Vogel et al., 2001). To better
understand the potential role of phosphorylation in γTuRC assembly, regulation, and function, we mapped a recently determined set of phosphorylation sites, including a re-analysis of previously determined data and newly acquired data from SPBs (Fong et al., 2018), onto a dimer of our γTuRCSS structure (Fig. 5A). Here we focus on the serine/threonine sites, given the minimal tyrosine kinase activity in yeast. Surprisingly, phosphorylation at the majority of the mapped sites would seem to destabilize the assembled γTuRC and thus may help keep unassembled or partially assembled components inactive. Phosphorylation at two sites would likely stabilize assembly, indicating the complex modulatory role played by phosphorylation.

Many of these phosphorylation sites map to potentially important interfaces: the Spc110p/Spc97p interface (at the Spc110pNCC and at the Spc110pNCC-CM1 loop), the inter-γTuSC interface, the γ-tubulin/α-tubulin interface, as well as a cluster of sites at the Spc97/98p: γ-tubulin interface. These are also a large number of unmapped phosphorylation sites, the majority of which are located on low-resolution or unresolved regions in the N-termini of Spc98p and Spc110p.

Strikingly, a cluster of phosphorylation sites, with many exhibiting in vivo phenotypes, maps near the Spc110p:γTuRCSS interface in the Spc110pNCC region and near the loop connecting Spc110pNCC and Spc110pCM1. Of particular note are a set of sites on Spc110p (Spc110pT182, Spc110pT188) and the adjacent interface on Spc97p (Spc97pS84, Spc97pT88). Together these would add numerous negative charges in a portion of the Spc97p/Spc110p interface that is already highly negatively charged, especially the Spc110pNCC. Phosphorylation at two of these sites (Spc110pT182 and Spc97pS84) would likely negatively impact Spc110p binding, whereas Spc97pT88 is adjacent a positively charged patch; phosphorylation at this site would likely promote Spc110p binding.

Three sites on Spc97p (Spc97pS130, Spc97pS208, Spc97pS209) and two sites on Spc110p (Spc110pS153, Spc110pS156) map onto or near the loop connecting Spc110pCM1 with the Spc110pNCC and its interface with Spc97p. Mutation of Spc97pS130 exhibited a temperature-sensitive phenotype, and the Spc97pS208A/S209A, Spc97pS208D/S209D double mutants were lethal, consistent with phosphorylation of this region potentially having a regulatory role (Fong et al., 2018; Lin et al., 2011). While the loop has a lower resolution...
than other portions of the map, the backbone approximately tracks with a long negatively-charged patch along Spc97p and Spc98p (Fig. S5C). Furthermore, the Spc110p\textsuperscript{150-161} loop has two negative charges and one positive charge. Although phosphorylation at Spc110p\textsuperscript{S153} and Spc110p\textsuperscript{S156} was not consistently observed (Fong et al., 2018), phosphorylation at these sites, as well as on the opposite Spc97p interface would likely destabilize Spc110p binding.

One site on Spc97p maps near the inter-γTuSC dimer interface. The interface overall is rearranged only by a few Ångstroms during activation, so any effect of phosphorylation would presumably only impact assembly. Spc97p\textsuperscript{S797} mutations produce a mild phenotype (Fong et al., 2018), and it is unresolved in all of our structures, but it is likely on a flexible loop near a positive patch in Spc110p and Spc97p and may thus favor assembly.

Finally, γ-tubulin\textsuperscript{S71} localizes near the γ-tubulin:α-tubulin interface, likely decreasing binding affinity, and perhaps even interfering with GTP binding. γ-tubulin\textsuperscript{S71} and γ-tubulin\textsuperscript{S74} mutants (A or D) both exhibit phenotypes, likely reflecting the importance of proper hydrogen bonding near the γ-tubulin GTP binding site (Fig. 5D).

Comparison of yeast γTuRC with metazoan TuRC structures

Recent efforts by several labs have been successful in providing the first models for the more complex metazoan γTuRCs (Liu et al., 2020; Wieczorek et al., 2020). These new structures provide much needed clarity on the stoichiometry of the five different GCPs (GCP2-6) and how they are organized within the γTuRC ring. They also reveal unexpected structural roles for numerous accessory components, with several components remaining unidentified due to limited resolution. Of interest to us was the role CM1-containing accessory proteins might have in metazoan γTuRC assembly and conformation.

Further support for a conserved role for CM1 is apparent in the recently published structure of the human γTuRC purified by affinity with γ-TuNA, an N-terminal truncation of CDK5RAP2, which includes its CM1 motif. Although not interpreted by the authors, we note their density maps clearly show that a helix binds in an identical position to our assigned yeast CM1 helix at the interface between GCP2 and GCP6 (Wieczorek et al., 2020)(Fig. S16A). Notably, a separate structural study of Xenopus γTuRC, where the γTuRC was
purified by affinity against γ-tubulin, showed no density at the same interface (Liu et al., 2020). Furthermore, when the human map is filtered to low resolution, density similar to that observed in our yeast γTuRCss difference map continues from the N-terminus of the CM1 helix along the surface of GCP6 towards GCP4 (Figs. S16B,C). Taken together, these results stress the broad conservation and importance of CM1 binding. These compositional differences led us to wonder whether CM1 binding might also drive conformational rearrangements in the metazoan γTuRCs, analogous to the changes we observed during yeast γTuRC assembly. Perhaps surprisingly, both metazoan γTuRC structures show a very poor match to MT symmetry, and would require substantial γ-tubulin motions to match the microtubule (Fig. 6). The γ-tubulins in the metazoan γTuRC structures are displaced up to ~46 Å from their ideal MT-like positions, as opposed to the 9 Å observed in our closed yeast γTuRCss structure, suggesting that the metazoan γTuRCs may be even more strongly dependent upon additional factors or PTMs to achieve an active conformation than the yeast γTuRCs.

While the human and Xenopus GCPs overlay very well at γ-tubulin positions 1-10, the terminal four positions show a different twist and pitch (Fig. S17A). We suggest here that these differences arise from CM1 binding at GCP2:GCP6 interface in the human γTuRC (Fig. S17A). The relative position of the γ-tubulins bound to GCP2 and GCP6 changes upon CM1 binding to much more closely match what we observe. That is, during the “transition” from a CM1-absent γTuRC (Xenopus) to a CM1-present γTuRC (human γTuRC), the GCP6-bound γ-tubulin moves by ~10 Å to better match the position observed in our closed TuRCss structure (Figs. S17B,C). From this we speculate that binding of CM1-containing accessory proteins at other sites within the γTuRC would further optimize their conformation and MT nucleating ability.

In contrast to the yeast structures, having CM1 bound in the human γTuRC seems to correlate with breaking the GCP2/6 N-terminal interface (Figs. S17D,E). This is a consequence of a lateral translation of ~10 Å of the N-terminal helical bundles of GCP2 away from GCP6 in the human structure. Interestingly, this separated interface appears to be stabilized by an unknown protein or domain which forms contacts with the external face of the γTuRC. The dissociation of this N-terminal interface may be due to weaker GCP2/6
interactions, enhancing the role that additional factors that bind at CM1 or the inter-GCP interface could play in regulating MT nucleation.

Discussion

Using a combination of single-particle and filament cryo-EM data, we have determined structures for monomeric and dimeric γTuSCs, along with assembled open and closed state γTuRCs at near-atomic and atomic resolutions. Our structures complement existing structural and biochemical data with high-resolution snapshots of the yeast γTuSC and γTuRC. Together with previous work, these provide a framework for understanding the molecular basis for MT nucleation and regulatory processes likely necessary to ensure that microtubules are only nucleated at the SPB. We provide the first molecular understanding for the critical role of the conserved Spc110p CM1 region in γTuRC assembly.

The structures suggest that nucleation is positively controlled in at least three ways: i) assembly of γ-TuSCs into an open ring mediated by Spc110p oligomers and Spc110p CM1, ii) closure of each γTuSC from an open state to a closed state to fully align the γ-tubulins to the MT lattice, and iii) PTMs that can affect Spc110p binding directly impact γTuRC assembly or affect γ-tubulin’s affinity for αβ-tubulin.

Although minor, we also observe conformational changes in γ-tubulin upon assembly into γTuSCs that mimic aspects of the bent-to-straight transition in αβ-tubulin and would thus be expected to enhance MT nucleation. Unresolved, is to what extent these differences arise from differences in the protein sequence from yeast to metazoans or represent an assembly-driven enhancement in γ-tubulin conformation. There is at least some role for sequence as we know that there is a strong species barrier such that yeast γTuRC is hundreds-fold more potent at stimulating yeast tubulin polymerization than mammalian tubulin (Kollman et al., 2015).

As initially observed in negative stain EM (Choy et al., 2009), our new cryo-EM structures of monomeric and dimeric γTuSCs show that Spc97p and Spc98p intrinsically adopt an open conformation at the intra-γTuSC interface such that the attached γ-tubulins fail to make microtubule-like lateral contacts. Our structures of open and closed assembled
γTuRCs show that Spc97p and Spc98p undergo large conformational changes during assembly into rings, bringing them much closer to MT geometry. Only smaller conformational changes occur as they transition from the open to the closed state during activation. The observation that a population of γTuRCs in the WT filaments adopts a locally closed conformation, indicates a small energetic barrier to closing. In the absence of other factors microtubule nucleation and lateral association may occur as the open and closed states are spontaneously sampled. However, even if a single γTuSC readily samples the closed state, the likelihood of an entire ring doing so could be quite low. Thus, the addition/removal of PTMs or the binding of other factors could allosterically drive a more ideal template state. Indeed, we know that the yeast CK1δ kinase, Hrr25, is needed for proper spindle formation in vivo and that it binds to γTuRCs and stimulates MT assembly in vitro (Peng et al., 2015), indicating that it is one such activator. γTuRC closure may also be stabilized by the process of microtubule assembly.

Our structures also resolve a long-standing mystery: how the six-fold symmetric Spc42p layer at the SPB could facilitate the formation of a γTuRC containing specifically seven γTuSCs. This is resolved by recognizing that Spc110pCM1 within each dimer extends from one γTuSC to another and cannot bridge across the large gap between the last and first γTuSCs in the ring. Thus, we suggest that six Spc110p dimers are symmetrically bound to the Spc42p lattice at the SPB. These would thus present the six CM1 motifs required to bind at the six complete CM1 binding sites formed within γTuRC heptamer (Fig. 4C). Given the observed pattern of connectivity where Spc110p CM1 extends across the interface in the same direction as the helical rise (Fig. 3A), this would leave the terminal NCC site nearest to the SPB unoccupied.

Our high-resolution structures are further poised to help inform on the mechanism of activation of the stable metazoan γTuRC complexes. Both of the published γTuRC structures would require large conformational changes in pitch and rise to efficiently nucleate microtubules (Fig. 6). Further, the inter-γTuSC interface is also less ideal in the metazoan structures compared to yeast, requiring a 7Å translational shift in the GCP3 C-terminus to match ours. In the human γTuRC we observe a CM1-like helix at the GCP2:GCP6 interface, and the distance between the GCPs at this interface closely matches that
observed in our γTuRC structures indicating a conserved and more optimal spacing upon CM1 binding (Figs. S16, S17B). This suggests the mode of interaction of the Spc110pCM1 helix with GCPs is broadly conserved. The fact that both structures have low resolution density extending past the CM1 N-terminus towards the adjacent GCP suggests there may also be a conserved functional role for the residues N-terminal to CM1 (Fig. S16BC).

Simple addition of a CDK5RAP2 homologue during purification did not yield observable CM1 density in the Xenopus γTuRC complexes (Liu et al., 2020) suggesting that other factors could play a regulatory role in this process. Combined with our additional observation of a change in the local twist and pitch of GCP: γ-tubulin conformation near the GCP2:GCP6, the data suggests that binding of a CM1 helix at the five GCP2:GCP3 inter-γTuSC interfaces could cooperatively rearrange the γTuRC to much better match the microtubule pitch and spacing, leading to activation of microtubule nucleation. The fact that these sites were empty when pulled down from cells suggests a relatively low affinity for the relevant factors or a low abundance of soluble protein. This further suggests a functional role for increasing the local concentration of CM1-containing proteins through either an ordered oligomerization processes, as with Spc110p, or through a more colligative phase-condensate mechanism. For example, if the additional CM1s were to come from pericentrin, which is tightly bound within the PCM, the affect would be to couple γTuRC activation to PCM localization, similar to Spc110p confining yeast γTuRC function to the SPB. Despite these major advances, significant gaps remain in our understanding of the how binding of regulatory proteins and PTMs act to modulate activation of yeast and metazoan γ-TuRCs.

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**Author Contributions**

AB purified proteins complexes and optimized sample preparation for cryo-EM, performed EM imaging experiments, cryo-EM image analysis and built atomic models. ASL and AM created expression constructs, purified proteins, and performed biochemical analyses. AZ performed protein crosslinking and mass spectrometry. SV performed integrative modeling. AB, ASL, SV, and AZ wrote the paper with input from EM, TND, DAA, and AS. TND, EM, MM, AS, and DAA supervised research.

**Conflicts of Interest**
The authors declare that they have no conflicts of interest.

Materials and Methods

γTuSC Purification

γTuSC was prepared essentially as described (Vinh et al, 2002, Lyon et al, 2015).

Cross-linking and mass spectrometry (XL-MS)

XL-MS was carried out as described by (Zelter et al, 2015). All γTuSC-Spc110 reactions were in 40 mM HEPES pH 7.0, 150 mM KCl and contained a final concentration 0.4 µM γTuSC and 0.8 µM Spc110. DSS reactions were carried out at room temperature (RT) for 3 min using 0.44 mM DSS prior to quenching with 100 mM ammonium bicarbonate. EDC reactions were carried out at RT for 30 min using 5.4 mM EDC plus 2.7 mM Sulfo-NHS prior to quenching with 100 mM ammonium bicarbonate plus 20 mM 2-mercaptoethanol. After quenching, reactions were reduced for 30 min at 37°C with 10 mM dithiothreitol (DTT) and alkylated for 30 min at RT with 15 mM iodoacacetamide. Trypsin digestion was performed at 37°C for 4 or 6 hours with shaking at a substrate to enzyme ratio of 17:1 or 30:1 for EDC and DSS reactions, respectively, prior to acidification with 5 M HCl. Digested samples were stored at -80°C until analysis. Mass spectrometry and data analysis were performed as described (Zelter et al., 2015). In brief 0.25 µg of sample was loaded onto a fused-silica capillary tip column (75-µm i.d.) packed with 30 cm of Reprosil-Pur C18-AQ (3-µm bead diameter, Dr. Maisch) and eluted at 0.25 µL/min using an acetonitrile gradient. Mass spectrometry was performed on a QExactive HF (Thermo Fisher Scientific) in data dependent mode and spectra converted to mzML using msconvert from ProteoWizard (Chambers et al, 2012).

Proteins present in the sample were identified using Comet (Eng et al, 2013). Cross-linked peptides were identified within those proteins using Kojak versions 1.4.1 or 1.4.3 (Hoopmann et al., 2015) available at http://www.kojak-ms.org. Percolator version 2.08 (Käll et al, 2007) was used to assign a statistically meaningful q value to each peptide spectrum match (PSM) through analysis of the target and decoy PSM distributions. Target databases consisted of all proteins identified in the sample analyzed. Decoy databases consisted of the corresponding set of reversed protein sequences. Data were filtered to show hits to the target proteins that had a Percolator assigned peptide level q value ≤ 0.01 and a minimum of 2 PSMs. The complete list of all PSMs and their Percolator assigned q values are available on the ProXL web application (Riffle et al, 2016) at https://proxl.yeastrc.org/proxl/p/cm1-tusc along with the raw MS spectra and search parameters used.

Xrc4-Spc110 Purification and X-ray Crystallography

DNA encoding residues 2-132 of H. sapiens Xrc4 (UniProt ID Q13426) fused in frame with residues 164-207 of Spc110 was synthesized by GeneArt (ThermoFisher Scientific) and cloned into pET28a expression vector with N-terminal 6His tag, 3C protease cleavage site, and six-residue linker with sequence GSGGSG. Xrc4-Spc110 was expressed in E. coli BL21-CodonPlus-RIL (Agilent). Cells were harvested by centrifugation then resuspended in lysis buffer (50 mM potassium phosphate pH 8, 300 mM NaCl, 5 mM EDTA, 1 mM DTT,
0.3% Tween-20, 1x Complete protease inhibitor, EDTA-free (Roche). Cells were lysed by Emulsiflex C3 (Avestin). Lysate was cleared by ultracentrifugation at 40,000 x g for 30 min in a Type 45 Ti rotor (Beckman-Coulter). Xrcc4-Spc110<sup>164-207</sup> was then purified by NiNTA affinity chromatography followed by addition of 3C protease overnight at 4°C to cleave the 6His tag. Xrcc4-Spc110<sup>164-207</sup> was further purified by size exclusion chromatography (Superdex 75; GE Healthcare Life Sciences), anion exchange chromatography (MonoQ; GE Healthcare Life Sciences), with a final size exclusion polishing and buffer exchange step (Superdex 75). Crystals of Xrcc4-Spc110<sup>164-207</sup> were obtained with by hanging drop vapor diffusion with 8 mg/mL protein and a well solution containing 13% PEG3350 and 0.2 M magnesium formate. Crystals were cryo-protected by rapid transfer to well solution with 30% PEG3350. Diffraction data was collected under cryogenic conditions at Advanced Light Source beamline 8.3.1. Diffraction data was processed with XDS (Kabsch, 2010) and indexed in space group P1. Phases were obtained by molecular replacement using Phaser within the Phenix package (Adams, et al., 2010; McCoy, et al., 2007). The search model was the PDB ID 1FU1 residues 1-150, with the coiled-coil residues 133-150 mutated to alanine. The S-(dimethylarsenic)cysteine at position 130 in 1FU1 was modified to cysteine. The majority of the structure was built with phenix.autobuild (Terwilliger, et al., 2008) with the remainder built manually in Coot (Emsley, Lohkamp, Scott, & Cowtan, 2010) and refined with phenix.refine (Afonine, et al., 2012). The final structure contains Spc110 residues 164-203, along with the Xrcc4 fusion domain.

**Filament Purification**

Filaments were prepared essentially as described with slight modifications. The buffer used during purification was modified to contain 40 mM HEPES pH 7.5, 100 mM KCl, 1 mM EGTA, 2 mM MgCl<sub>2</sub>, 10% glycerol, and 1 mM DTT. Samples were concentrated and buffer exchanged to obtain a final glycerol concentration of 2.5% glycerol. Oxidation of γTuSC<sup>88</sup> filaments was performed overnight at 4°C by dialysis into 1 mM oxidized glutathione, removing DTT.

**Grid Preparation - γ TuSC**

Prior to grid preparation γ–TuSC aliquots were centrifuged in a benchtop centrifuge (Eppendorf5415D) at 16’000 g for 15 minutes and transferred to a new tube. The sample concentration was assessed on a nanodrop, and diluted to a final concentration of ~1 μM (O.D. at 280 nm wavelength of 0.28-0.35) such that the final buffer conditions were 40 mM HEPES pH 7.5, 2 mM MgCl<sub>2</sub>, 1 mM EGTA, 1 mM GDP, 100 mM KCl and 2.5% v/v glycerol. Data used for initial model generation and refinement had final buffer conditions of 40 mM HEPES pH 7.5, 1 mM MgCl<sub>2</sub>, 1 mM EGTA, and 100 mM KCl. C-flat 1.2-1.3 4C grids were used for sample freezing and glow discharged for ~30s at 20 mA immediately prior to plunge-freezing. Grids were frozen on a Vitrobot Mark II or Mark IV, with the humidity set to 100%, and using Whatman 1 55 mm filter papers.

**Grid Preparation - γTuSC Filaments**
Quantifoil 1.2-1.3 400-mesh grids were used for sample freezing and glow discharged for
~30 s at -20 mA immediately prior to plunge-freezing. Grids were frozen on a Vitrobot Mark IV, with the humidity set to 100%, and using Whatman 1 55 mm filter papers.
The final conditions used for γTuSC (WT) filament freezing was 40 mM HEPES pH 7.5, 2 mM MgCl₂, 1 mM EGTA, 2 mM GDP, 100 mM KCl, 1 mM DTT and 2.5% v/v glycerol.
The final conditions used for γTuSCSS filament freezing was 40 mM HEPES pH 7.5, 2 mM MgCl₂, 1 mM EGTA, 0.5 mM GTP, 100 mM KCl, 1 mM oxidized glutathione and 2.5% v/v glycerol.

Electron Microscopy – γTuSC Single-particle Data

Micrographs used in γTuSC initial model generation were collected using an FEI Tecnai F20 operated at 200 kV at a nominal magnification of 29’000X (40’322X at the detector). The data was collected with a 20 μm C2 aperture, and a 100 μm objective aperture with a target underfocus of ~1-2.5 μm. UCSF Image4 (Li et al., 2015) was used to operate the microscope. Dose-fractionated micrographs were collected on a Gatan K2 Summit camera in super-resolution mode at a dose rate of ~8.5-9.5 electrons per physical pixel per second for 12 seconds, with the dose fractionated into 40 frames.

Micrographs included in the final model were collected using an FEI Tecnai Polara operated at 300 kV at a nominal magnification of 31’000X (39’891X at the detector). Data was collected with a 30 μm C2 aperture and a 100 μm objective aperture inserted with a target underfocus of ~1-3 μm. Leginon (Suloway et al., 2005) or SerialEM (Mastronarde, 2005) were used to operate the microscope. Dose-fractionated micrographs were collected on a Gatan K2 Summit camera in super-resolution mode at a dose rate of ~6 electrons per physical pixel per second for 20 seconds, with the dose fractionated into 100 frames.

Electron Microscopy – γTuSCWT Filament Data

Data was collected in two sessions on a Titan Krios operated at 300 kV at a nominal magnification of 22’500X (47’214X at the detector). The data was collected with a 70 μm C2 aperture, and a 100 μm objective aperture with a target underfocus of ~0.9-2.0 μm. Dose-fractionated micrographs were collected on a Gatan K2 Summit camera in super-resolution mode at a dose rate of 6 electrons per physical pixel per second for 15 seconds, with the dose fractionated into 75 frames.

Electron Microscopy – γTuSCSS Filament Data

Data was collected in two sessions on a Titan Krios operated at 300 kV at a nominal magnification of 22’500X (47’214X at the detector). The data was collected with a 70 μm C2 aperture, and a 100 μm objective aperture with a target underfocus of ~0.6-2.0 microns.
Dose-fractionated micrographs were collected on a Gatan K2 Summit camera in super-resolution mode at a dose rate of 6.7 electrons per physical pixel per second for 12 seconds, with the dose fractionated into 120 frames.

Image Processing – γTuSC Single Particle Initial model generation
Dose-fractionated image stacks were corrected for drift and beam-induced motion as well as binned 2-fold from the super-resolution images using MotionCor (Li et al., 2013). CTF estimation was performed using CTFFIND4 (Rohou and Grigorieff, 2015). Particle coordinates were semi-automatically picked from filtered and binned images using the e2boxer “swarm” tool (Tang et al., 2007). Particles were extracted using Relion (Scheres, 2012) with a box size of 384 physical pixels resampled to 96 pixels for initial processing. A dataset of ~50,000 particles from 217 micrographs was used to generate 300 2D classes using Relion 1.3. 23 classes were selected and used in the generation of a γ-TuSC monomer initial model using the e2initialmodel.py function in EMAN2. This model was then used as a reference in Relion 1.3 for 3D classification into 4 classes of a 115,701 particle dataset from 507 micrographs with a 384 pixel box. Particles from the best γ-TuSC monomer class were then used for further processing and classification into 4 classes in FREALIGN (Grigorieff, 2016). The best class, with a resolution of ~9 Å was then used as a 3D reference for processing of the Polara data.

**Image processing – γTuSC Single Particle Polara Data**

Images were drift-corrected and dose-weighted using MotionCor2 (Zheng et al., 2017). Initial processing to generate monomer and dimer reconstructions was performed with CTFFIND4 (Rohou and Grigorieff, 2015), relion (Scheres, 2012), and FREALIGN (Grigorieff, 2016). Processing leading to the final reconstructions was performed in cisTEM (Grant et al., 2018). Particles were automatically picked from 7381 images in cisTEM, yielding 3,210,917 initial particle coordinates. 2D classification was performed to eliminate junk and ice particles, with 1,187,292 particles being included in initial 3D classification. During 3D classification, Particles were extracted from unbinned super-resolution micrographs with a box size of 376.02 Å (600 pixels).

Classification and alignment were performed using the cisTEM “Manual Refine” tool, as delineated in Figure S13.

**Image processing – γTuSCWT Filaments**

Images were drift-corrected, dose-weighted and binned two-fold using MotionCor2 (Zheng et al., 2017). Filaments were manually picked using e2helixboxer (Tang et al., 2007) from 2204 micrographs. Filaments were extracted in Relion2 (Kimanius et al., 2016) and boxed approximately every 3 asymmetric units, using a rise of 21 Angstroms with a box size of 635.4 Angstroms (600 pixels on the micrographs, rescaled to 448 pixels), yielding 28,753 boxed filament images. 2D classification was performed to eliminate junk particles and filament ends, with 28,648 filament images remaining after culling. These images were initially aligned in Relion2 (Kimanius et al., 2016), while allowing for the refinement of helical parameters. Particle alignments were exported into FREALIGN (Grigorieff, 2016) for additional helical refinement. FREALIGN alignments were used for helical symmetry expansion as implemented in Relion2. Symmetry expanded alignment parameters were then imported into cisTEM (Grant et al., 2018) for local alignment and classification. A user-generated mask was supplied for these refinements, with the final mask containing approximately 3 γTuSC subunits with a total molecular weight of approximately 900 kDa.
Prior to classification, the defocus was refined in cisTEM. Focused classification was performed in cisTEM, as delineated in Fig. S5.

**Image processing – γTuSCSS Filaments**

Images were drift-corrected, dose-weighted and binned two-fold using MotionCor2 (Zheng et al., 2017). Filaments were manually picked using e2helixboxer from 3024 micrographs. Filaments were extracted in Relion2 (Kimanius et al., 2016) and boxed approximately every 3 asymmetric units, using a rise of 21 Å with a box size of 635.4 Angstroms (600 pixels), yielding 175,500 boxed filament images. 2D classification was performed to eliminate junk particles and filament ends, with 152,798 filament images remaining after culling. These images were initially aligned in Relion2, while allowing for the refinement of helical parameters. Particle alignments were exported into FREALIGN (Grigorieff, 2016) for additional helical refinement. FREALIGN alignments were used for helical symmetry expansion as implemented in Relion2. Symmetry expanded alignment parameters were then imported into cisTEM (Grant et al., 2018) for local alignment and classification. A user-generated mask was supplied for these refinements, with the final mask containing approximately 3 γTuSC subunits with a total molecular weight of approximately 900 kDa. Prior to classification, the defocus was refined in cisTEM. Focused classification was performed in cisTEM, as delineated in Fig. S6.

**Difference Map Generation**

The γTuSCSS reconstruction was resampled to 400 pixels using resample.exe included in the cisTEM package. A molecular map of a trimer of γTuSCSS that did not include Spc110p was generated in chimera using the molmap command with a resolution of 3.3 Å. A difference map was generated using the diffmap.exe software obtained from the Grigorieff lab website (https://grigoriefflab.umassmed.edu/diffmap). The difference map was sharpened with a bfactor of -40 Å² and filtered to 5.5 Å with a 5-pixel fall-off using the bfactor software obtained from the Grigorieff lab website (https://grigoriefflab.umassmed.edu/bfactor).

**Initial Atomic Model Generation – Monomers**

To generate an initial atomic model, the crystal structure of human GCP4 and a previously generated pseudo-atomic model were used as templates. Prior to fitting, the GCP4 structure was threaded with the Spc97p and Spc98p sequence, and the human γ-tubulin was threaded with the Tub4p sequence. These initial models were fitted into preliminary structures into segmented density using Rosetta’s relax function. Missing residues were built using RosettaCM density-guided model building (DiMaio et al., 2015), with the human GCP4, γ-tubulin threaded models and the pseudo-atomic model being sampled separately. Well scoring structures were then compared to the density, assessing the quality of the fit to determine the register. In cases where the register was poorly fit and the correct register was clear, the register was manually adjusted to fit map details. Certain regions were built using the RosettaES algorithm (Frenz et al., 2017). This procedure was iterated, with occasional manual modification of the structure in Coot (Emsley et al., 2010).
As a final step, final half-maps were used in the refinement, with the best preliminary models were relaxed and refined through iterative backbone rebuilding (Wang et al., 2016) into one half-map reconstruction, and iteratively refined using Rosetta. This model was used as a starting point for atomic model building into the higher resolution γTuSCSS filament structure.

**Atomic Model Generation – γTuSCSS**

The initial model from monomer fitting was relaxed into the γTuSCSS structure using Rosetta’s relax function, and refined using iterative backbone rebuilding as previously described. Poorly fitting and missing regions were either built in Coot or using the RosettaES algorithm. Residues Spc110p112-150 were manually built in Coot. Finally, the models were iteratively refined using a procedure that involved using Rosetta to relax the models into one half-map and iterative backbone rebuilding, with the best models as assessed using the FSC to the second half-map being combined using the phenix combine_models function, followed by Phenix (Adams et al., 2010) real-space refinement (Afonine et al., 2018) and manual modification. This model was used as the basis for the single-particle monomer model, and the γTuSCWT models. Models were further iteratively refined using Rosetta, Coot, and Phenix. Finally, the Spc110p164-208 crystal structure was relaxed into γTuSCSS density, with the residues 151-164 built manually in Coot. Spc110p was iteratively relaxed into density using Rosetta to relax the models into one half-map and iterative backbone rebuilding, with the best models being visually inspected and manually modified in Coot. A final round of manual refinement of Spc110p was performed in ISOLDE (Croll, 2018) using a density-modified map generated in Phenix (Terwilliger et al., 2020).

**Atomic Model Generation – γTuSCWT open state**

The initial model from γTuSCSS fitting was relaxed into the γTuSCWT structure using Rosetta’s relax function, and refined using iterative backbone rebuilding, with the best models as assessed using the FSC to the second half-map being combined using the phenix combine_models function. Models were further iteratively refined using Rosetta, Coot, and Phenix.

**Atomic Model Generation – γTuSCWT closed state**

The initial model from γTuSCSS fitting was relaxed into the closed γTuSCWT structure using Rosetta’s relax function, and refined using iterative backbone rebuilding, with the best models as assessed using the FSC to the second half-map being combined using the phenix combine_models function, followed by Phenix real-space refinement and manual modification. Models were further iteratively refined using Rosetta, Coot, and Phenix.

**Model Generation – γTuSC monomer**

The initial model from γTuSCSS fitting was relaxed into the γTuSC monomer structure using Rosetta’s relax function, and refined using iterative backbone rebuilding, with the best models as assessed using the FSC to the second half-map being combined using the phenix
combine_models function, followed by Phenix real-space refinement and manual modification. Models were further iteratively refined using Rosetta, Coot, and Phenix. The final round of Phenix real-space refinement was performed against the full map.

Model Generation – γTuSC dimer

The dimer model was generated by using Rosetta’s relax function to fit two γTuSCSS models generated as above into dimer density. The nucleotide was subsequently modified to GDP and poorly fitting regions were deleted. This model was used solely for the segmentation shown in Fig. S14.

Surface Area calculations

Surface area calculations were performed using NACCESS (Hubbard and Thornton, 1993).

2D Classification – Figure S1

Monomer and dimer stacks (384 pixel stacks used in final reconstruction generation) were separately classified using cisTEM (Grant et al., 2018). Classes showing high resolution features were extracted for figure generation using IMOD (Kremer et al., 1996).

Wiring diagrams

Wiring diagrams were generated using the PDBSum online portal (Laskowski, 2009).

Sequence alignments for conservation surfaces.

Sequences for Spc97p and Spc98p homologs from Homo sapiens, Mus musculus, Danio rerio, Xenopus laevis, Drosophila melanogaster, Arabidopsis thaliana, Glycine max, Dictyostelium discoideum, and Saccharomyces pombe were aligned to the sequence from Saccharomyces cerevisiae using the MAFFT algorithm (Katoh and Standley, 2013) implemented on the MPI bioinformatics website (Zimmermann et al., 2018). The Spc110p sequence from Saccharomyces cerevisiae was similarly aligned to orthologs from Homo sapiens, Mus musculus, Danio rerio, Xenopus laevis, Drosophila melanogaster, Dictyostelium discoideum, and Saccharomyces pombe. Sequence alignments were imported using the Multalign Viewer in Chimera (Pettersen et al., 2004), which was subsequently used to color the surfaces and ribbons by conservation.

Figure Generation

Structural figures were generated in UCSF Chimera (Pettersen et al., 2004) or ChimeraX (Goddard et al., 2018). FSC plots were generated in Excel from Part_FSC estimates in Frealign. Map-to-model FSCs were generated in Phenix (Adams et al., 2010), with default parameters. Figures panels were compiled into figures in Affinity Designer.

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**Figure 1: The Spc110p\textsuperscript{NCC} binds near the N-terminus of Spc97p**

A. Spc110p N-terminal region secondary structure prediction, showing lack of predicted secondary structure for the first 111 residues. Also shown are Spc110p\textsuperscript{CM1(117-146)} and the Spc110p\textsuperscript{NCC(164-208)} regions.

B. Structure of Xrcc4-Spc110p\textsuperscript{164-207}, where Spc110p\textsuperscript{NCC} residues 164-203 are resolved.

C. Spc110p\textsuperscript{NCC} structure fit into \(\gamma\)TuSC cryo-EM density map (grey surface, EMDB ID 2799) along with \(\gamma\)TuSC pseudo-atomic model (PDB ID 5FLZ) (Kollman, et al., 2015; Greenberg, et al., 2016). The majority of XL-MS distance restraints are satisfied by this model. Satisfied and violated DSS crosslinks are shown in cyan and purple, respectively. Satisfied and violated EDC crosslinks are shown in blue and red, respectively. Crosslinks that are satisfied by either Spc110p monomer are shown twice, one for each monomer.

D. Localization density map for the ensemble of integrative models consisting of two adjacent \(\gamma\)TuSCs, each bound to an Spc110p\textsuperscript{1-220} dimer. The map shows the positions of different parts of the complex in the ensemble of models from the top cluster; maps for all components are contoured at 2.5% of their respective maximum voxel values. The modeling results shown are based on the \(\gamma\)TuSC-Spc110p\textsuperscript{1-220} GCN4 crosslinks; similar results were obtained using \(\gamma\)TuSC-Spc110p\textsuperscript{1-401} GST crosslinks (see Supplementary Methods).
Figure 2: Structure and Assembly interfaces of WT γTuSC and γTuSCSS

A-B Segmented density of (A) open WT γTuSC and (B) closed γTuSCSS. γTuSC subunits are colored as in the figure inset. Density was segmented within 4.5 Å of the atomic model, showing one Spc110p copy. Disconnected density smaller than 5 Å was hidden using the "Hide Dust" command in Chimera. Spc110pNCC is not visible at this threshold due to heterogeneity.

C-D. Representation of the intra- (C) and inter-γTuSC (D) interfaces of Spc97p/98p illustrated on a γTuSCSS dimer. Interface atoms are shown as spheres and colored by their hydrophobicity according to the Kyte-Doolittle scale.
Figure 3: The γ-tubulin GCP binding interfaces

A. Filtered segmented difference map between experimental density and the fitted atomic model without Spc110p overlaid on a γTuSC$_{SS}$ surface lacking Spc110p. The difference map was segmented to show density near a γTuSC$_{SS}$ monomer, and colored to attribute densities to their putative chains.

B. Density for the helical CM1 density of γTuSC$_{SS}$ showing clear side-chain features unambiguously defining the register. Density was zoned near the atoms in Chimera with a radius of 2.6 Å.

C. View of the binding site for CM1 and the strands preceding and following the CM1 helix.
Figure 4: Conformational Changes of γTuSC During Assembly and Activation

Top and side views of open (A,D) and Closed (SS) (B,E) γTuSC assemblies. Panel C shows a bottom view of an assembled γTuSC heptamer. The arrow indicates the seventh Spc110p\textsuperscript{NCC} binding site in a γTuSC heptamer which is likely not to have bound Spc110p, given the 6-fold symmetry observed in Spc42p at the SPB, and the lack of a CM1 binding site at the adjacent inter-γTuSC interface.
Figure 5: Phosphorylation sites visualized on the γTuSC structure

A. γTuSC tetramer, colored as in Figure 1A, with phosphorylation sites from (Fong et al., 2018) marked with red balls (no known phenotype) or purple balls (phenotype previously reported).

B. View of phosphorylation site at the Spc97p Spc110p NCC binding site.

C. View of the path of the Spc110p loop between the Spc110p NCC and Spc110p CM1 domain. This loop shows 2 phosphorylation sites opposite an acidic path.

D. Phosphorylation sites mapped on the γ-tubulin:α-tubulin interface, illustrating the position of the phosphorylation sites in relation to the interface with α-tubulin, Spc98p-bound γ-tubulin is in khaki, while α-tubulin is in light green.
Figure 6: Metazoan γTuRCs require large motions to template microtubules

A. Yeast closed (This work), B. Xenopus (PDB ID 6TF9), C. Human γTuRC (PDB ID 6V6S) structures placed adjacent to a microtubule to illustrate the motions required to properly template microtubules. For each structure, two γ-tubulins (positions 2,3 for Xenopus and Human and positions 13,14 for yeast) were aligned with two β-tubulins docked in microtubule density to approximate binding of γ-tubulins to a microtubule.