Molecular mechanism of microtubule nucleation from gamma-tubulin ring complex

Akanksha Thawani¹, Howard A Stone², Joshua W Shaevitz³,⁴, Sabine Petry⁵.*

¹Department of Chemical and Biological Engineering, Princeton University
²Department of Mechanical and Aerospace Engineering, Princeton University
³Lewis-Sigler Institute for Integrative Genomics, Princeton University
⁴Department of Physics, Princeton University
⁵Department of Molecular Biology, Princeton University, United States

* Correspondence to: Sabine Petry (spetry@princeton.edu)
Abstract

Determining how microtubules (MTs) are nucleated is essential for understanding how the cytoskeleton assembles. Yet, half a century after the discovery of MTs and αβ-tubulin subunits and decades after the identification of the γ-tubulin ring complex (γ-TuRC) as the universal MT nucleator, the underlying mechanism largely remains a mystery. Using single molecule studies, we uncover that γ-TuRC nucleates a MT more efficiently than spontaneous assembly. The laterally interacting array of γ-tubulins on γ-TuRC facilitates the lateral association of αβ-tubulins, while longitudinal affinity between γ/αβ-tubulin is surprisingly weak. During nucleation, 3-4 αβ-tubulin dimers bind stochastically to γ-TuRC on average until two of them create a lateral contact and overcome the nucleation barrier. Although γ-TuRC defines the nucleus, XMAP215 significantly increases reaction efficiency by facilitating αβ-tubulin incorporation. In sum, we elucidate how MT initiation occurs from γ-TuRC and determine how it is regulated.
Introduction

Microtubules (MTs) enable cell division, motility, intracellular organization and transport. MTs were found to consist of αβ-tubulin dimers fifty years ago, yet, how MTs are nucleated in the cell to build the cytoskeleton remains poorly understood\(^1\)–\(^3\).

MTs assemble spontaneously from αβ-tubulin subunits \textit{in vitro} via the cooperative assembly of many tubulin dimers and hence this process displays a kinetic barrier\(^4\)–\(^8\). Consequently, spontaneous MT nucleation is rarely observed in cells\(^9\),\(^10\). Instead, the major MT nucleator γ-tubulin is required \textit{in vivo}\(^9\)–\(^11\). γ-tubulin forms a 2.2 megadalton, ring-shaped complex with γ-tubulin complex proteins (GCPs), known as the \(\gamma\)-Tubulin Ring Complex (\(\gamma\)-TuRC)\(^12\)–\(^16\). \(\gamma\)-TuRC has been proposed to template the assembly of αβ-tubulin dimers into a ring, resulting in nucleation of a MT\(^15\)–\(^21\). However, kinetic measurements that provide direct evidence for this hypothesis have been lacking and several important questions about how \(\gamma\)-TuRC nucleates MTs have remained unanswered.

In the absence of purified \(\gamma\)-TuRC and an assay to visualize MT nucleation events from single \(\gamma\)-TuRC molecules in real time, recent studies used alternative MT assembly sources, such as spontaneous MT assembly or stabilized MTs with blunt ends hypothesized to resemble the \(\gamma\)-TuRC interface. Based on these alternatives, competition between growth and catastrophe of the nascent plus-end was proposed to yield the nucleation barrier in the cell\(^22\),\(^23\), but this has not been examined with the nucleator \(\gamma\)-TuRC. Recently, the MT polymerase XMAP215 was identified as an essential MT nucleation factor \textit{in vivo}, which synergistically nucleates MTs with \(\gamma\)-TuRC\(^24\)–\(^26\). Yet, the specific roles of XMAP215 and \(\gamma\)-TuRC in MT nucleation have yet to be discovered.

To explore the mechanism of MT nucleation, we reconstituted and visualized MT nucleation by \(\gamma\)-TuRC live with single molecule resolution. We uncover the molecular composition of the
MT nucleus, and determine the roles XMAP215 and γ-TuRC in MT nucleation.
Results

Reconstituting and visualizing microtubule nucleation from γ-TuRC

To study how γ-TuRC nucleates MTs (Fig. 1A), we purified endogenous γ-TuRC from Xenopus egg extracts and biotinylated the complexes to immobilize them on functionalized glass (Fig. S1A-C). Upon perfusing fluorescent αβ-tubulin, we visualized MT nucleation live with total internal reflection fluorescence microscopy (TIRFM). Strikingly, MT nucleation events occurred specifically from single γ-TuRC molecules (Fig. 1B; Fig. S1D and Movie S1-2). Kymographs revealed that attached γ-TuRC assembled αβ-tubulin into a MT de novo starting from zero length within the diffraction limit of light microscopy (Fig. 1C), ruling out an alternative model where MTs first spontaneously nucleate and then become stabilized via γ-TuRC. By observing the fiduciary marks on the MT lattice (Fig. 1C) and generating polarity-marked MTs from attached γ-TuRC (Fig. S1E), we showed that γ-TuRC caps the MT minus-end, while only the plus-end polymerizes. Altogether, our results show that γ-TuRC directly nucleates MTs.

Defining the microtubule nucleus on γ-TuRC

To determine how γ-TuRC nucleates MTs, we measured the kinetics of MT nucleation for a constant density of γ-TuRC and increasing αβ-tubulin concentration (Fig. 1D and Movie S3). Surprisingly, γ-TuRC nucleated MTs starting from 7 µM tubulin (Fig. 1D), which is higher than the minimum tubulin concentration ($C^*$) needed for growth at pre-formed MT plus-ends ($C^* = 1.4$ µM, Fig. 1E). Furthermore, the number of MTs nucleated from γ-TuRC increased non-linearly with tubulin concentration as opposed to the linear increase in MT’s growth-speed with tubulin concentration (Fig. 1E). By measuring the number of MTs nucleated over time with varying αβ-
tubulin concentration (Fig. 1F), we calculated the rate of MT nucleation. The power-law dependence on tubulin concentration (Fig. 1G) yields the number of αβ-tubulin dimers, 3.7 ± 0.5, that initiate MT assembly from γ-TuRC (Fig. 1G). Thus, the cooperative assembly of 3-4 tubulin subunits on γ-TuRC represents the most critical, rate-limiting step in MT nucleation.

Efficiency of γ-TuRC-mediated nucleation

Based on the traditional, fixed, end-point assays for MT nucleation with large error margins, γ-TuRC was believed to be a poor nucleator\textsuperscript{14}. To measure the efficiency of γ-TuRC-mediated MT nucleation, we compared it with spontaneous MT nucleation in our live TIRFM assay (Fig. 1H).

In contrast to γ-TuRC-mediated nucleation, a high concentration of 14 µM tubulin was required for spontaneous assembly of MTs, after which both the plus- and minus-ends polymerize (Fig. 1H, Fig. S1F and Movie S4). The number of MTs assembled as a function of the αβ-tubulin concentration displayed a power-law dependence with the exponent of 8 ± 1 (Fig. 1I), indicating a highly cooperative process that requires 8 αβ-tubulin dimers in a rate-limiting intermediate, in agreement with previous reports (Fig. 1H schematic, refs 4,8). In conclusion, γ-TuRC nucleates MTs significantly more efficiently (Fig. S1G), because its critical nucleus requires less than half the number of αβ-tubulin dimers compared to spontaneous assembly.

Does γ-TuRC nucleate a microtubule via the blunt plus-end model?

It has been widely proposed that the γ-tubulin ring on γ-TuRC resembles the blunt plus-end of a MT formed by a ring of αβ-tubulins\textsuperscript{20,22,27}. To test this proposition, we generated stabilized MT seeds with blunt ends as described recently\textsuperscript{22} and observed MT assembly from αβ-tubulin dimers (Fig. 2A). At a minimum concentration of 2.45 µM, approaching the critical concentration needed
for polymerization, a large proportion of pre-formed MT seeds assembled MTs immediately (Fig. S2A-B, Fig. 2A and Movie S5). The measured reaction kinetics (Fig. 2B) as a function of the αβ-tubulin concentration was used to obtain a power-law of the nucleation rate, 1.2 ± 0.4 (Fig. 2C). This demonstrates that blunt MT seeds assemble tubulin dimers into a lattice in a non-cooperative manner, where a single αβ-tubulin dimer suffices to overcome the rate-limiting step resembling the polymerization of a MT. Thus, the kinetics of γ-TuRC-mediated MT nucleation does not resemble a blunt MT plus-end.

**Molecular insight into microtubule nucleation by γ-TuRC**

We hypothesized that γ-tubulin’s binding properties with αβ-tubulin at the nucleation interface γ-TuRC could provide insight into the mechanism of nucleation. We purified γ-tubulin, which assembles into higher order oligomers in physiological buffer 24 and strikingly, into filaments at high γ-tubulin concentrations (Fig. S2C). Because γ-tubulins have been shown to arrange laterally, as observed previously in its crystallized form28, a plus-ends outward orientation of γ-tubulin molecules could form a nucleation interface.

Surprisingly, the γ-tubulin oligomers efficiently nucleated MTs from αβ-tubulin subunits (Fig. 2D and Movie S6) and even more strikingly, capped MT minus-ends while allowing newly generated MT plus-ends to polymerize (Fig. 2E). This activity is similar to that of γ-TuRC, suggesting that lateral γ-tubulin arrays on the nucleation interface of γ-TuRC are sufficient to nucleate MTs.

Knowing that lateral γ-tubulin arrays in purified γ-tubulin oligomers and within γ-TuRC nucleate MTs, we hypothesized that the longitudinal affinity between γ-tubulin and αβ-tubulin at the interface of γ-TuRC could be critical in regulating its nucleation efficiency. Using biolayer
interferometry, we compared the interaction of αβ-tubulin dimers with themselves versus with γ-tubulin. Specific interactions between probe-bound αβ-tubulin and increasing concentrations of unlabeled αβ-tubulin were measured (Fig. 2F), which must be longitudinal based on the observed protofilaments in the αβ-tubulin sample by EM (Fig. S2D). In contrast, no significant binding between monomeric γ-tubulin and αβ-tubulin was detected (Fig. 2F), suggesting that the heterogenous longitudinal affinity between γ-tubulin and αβ-tubulin on the nucleation interface may be weaker compared to αβ-tubulin with another αβ-tubulin molecule that occurs when the MT lattice polymerizes. In sum, the difference in interaction strength is the basis for the kinetic barrier we observed with γ-TuRC but not with a blunt MT plus-end, which we summarize with an interface interaction model (Fig. 2G).

We next asked how 3-4 tubulin dimers formed the rate-limiting species during γ-TuRC nucleation. In stochastic simulations, the 13 available binding sites on γ-tubulin molecules within γ-TuRC were allowed to be occupied at random with αβ-tubulin subunits. We then assessed how many αβ-tubulin dimers need to assemble on γ-TuRC to obtain two αβ-tubulin molecules on neighboring sites and form a favorable configuration with a lateral contact between the two αβ-tubulins (Fig. 2H). The simulations show that 3.7 ± 1 tubulin dimers assemble on γ-TuRC to form the first lateral contact between two αβ-tubulins (Fig. 2H), in striking agreement with the critical nucleus size we measured. In sum, our data shows that a lateral γ-tubulin array positioned by γ-TuRC promotes MT nucleation. The low γ-tubulin:αβ-tubulin affinity requires binding of 3-4 αβ-tubulin dimers to γ-TuRC to form the first lateral contact between two αβ-tubulin dimers and overcome the kinetic barrier before entering the MT polymerization phase. This nucleation barrier, in turn, provides the ability to further modulate MT nucleation via other factors.
Regulation of γ-TuRC mediated nucleation by microtubule associated proteins

Recent work suggested that MT-associated proteins (MAPs), which stabilize or destabilize MT plus-ends, influence MT nucleation in an analogous fashion.\textsuperscript{7,22,23} We assessed this hypothesis for MT nucleation by γ-TuRC. The protein TPX2 functions as an anti-catastrophe factor \textit{in vitro} \textsuperscript{22,23} and has been suggested to directly stimulate γ-TuRC-mediated nucleation.\textsuperscript{21,29–31} Strikingly, although TPX2 binds along the MT lattice, it does not increase nucleation activity of γ-TuRC (Fig. 3A and Movie S7). Similarly, the catastrophe factor EB1 does not decrease the nucleation activity of γ-TuRC (Fig. S3A and Movie S8). Thus, in agreement with our previous results (Figs. 1 and 2A-B), destabilization of MT plus-ends and a competition between polymerization/depolymerization is not sufficient to explain the properties of MT nucleation from γ-TuRC. Not surprisingly, decreasing the net rate of incorporation of tubulin into a MT using Stathmin, which sequesters tubulin dimers\textsuperscript{32,33}, or MCAK, which removes tubulin dimers from the MT lattice and prevents polymerization\textsuperscript{34,35}, decreased the number of MTs generated from γ-TuRC (Fig. S3B).

How do γ-TuRC and XMAP215 synergistically nucleate microtubules?

At low tubulin concentration of 3.5 µM and 7 µM, where either none or very little MT nucleation occurs from γ-TuRCs alone respectively, the addition of XMAP215 induced many surface-attached γ-TuRCs to nucleate MTs resulting in significant increase in MT nucleation rate (Fig. 3B-C and Movie S9). XMAP215 effectively decreases the minimal tubulin concentration necessary for MT nucleation from γ-TuRC to 1.6 µM, which is very close to that needed for plus-end polymerization. What is the sequence of events that leads to synergistic MT nucleation? By directly visualizing γ-TuRC and XMAP215 molecules during the nucleation reaction, we found
that XMAP215 and γ-TuRC molecules first formed a complex from which a MT was nucleated (Fig. 3D and Movie S11). For 76% of the events (n=56), XMAP215 visibly persisted between 3 to over 300 seconds on γ-TuRC before MT nucleation, and with a 50% probability XMAP215 remained on the minus-end together with γ-TuRC (n=58).

Could XMAP215 accelerate nucleation by altering the critical tubulin nucleus that assembles during γ-TuRC-mediated nucleation? Titrating tubulin at constant γ-TuRC and XMAP215 concentrations (Fig. S4A and Movies S10) yielded a similar power-law dependence between the MT nucleation rate and tubulin concentration (Fig. 3E). The resulting critical nucleus size of 3.2 ± 1.2 is very similar to that for γ-TuRC alone (Fig. 3E). Moreover, the C-terminus of XMAP215 (TOG5 and C-terminal domain), which directly interacts with γ-tubulin but not with αβ-tubulin24, does not enhance MT nucleation from γ-TuRC (Fig. S4B). Altogether, γ-TuRC determines the critical nucleus of αβ-tubulin dimers for MT nucleation (Fig. 2H). XMAP215, which directly binds to γ-tubulin via its C-terminal domain, does not appear to activate γ-TuRC via a conformational change, but likely relies on N-terminal TOG domains to increase αβ-tubulin incorporation by effectively increasing the local αβ-tubulin concentration, and thereby promoting MT nucleation.
Decades after the discovery of αβ-tubulin and MTs and the identification of γ-TuRC as the universal MT nucleator\(^7\)–\(^9\), it has remained poorly understood how MTs are being nucleated\(^7\)–\(^10\). Here, we show that γ-TuRC-mediated MT nucleation is more efficient than spontaneous MT assembly, requiring fewer tubulin dimers to form the rate-limiting reaction intermediate. This explains why MTs do not form spontaneously in the cell and why γ-TuRC is essential, addressing a long debate on γ-TuRC’s MT nucleation activity and requirement\(^36–38\). Spontaneous MT assembly requires higher tubulin concentrations and occurs due to stronger longitudinally-interacting αβ/αβ-tubulin and weaker lateral interactions. In contrast, γ-TuRC-mediated nucleation, driven by the lateral adjacency of the γ-tubulins on the nucleation interface, is sufficient to overcome the intrinsically very weak αβ-tubulin lateral interaction, thereby potentiating MT nucleation. Thus, we propose that, in metazoans analogous to the S. cerevisiae γ-TuSC rings\(^15,16\), GCPs within γ-TuRC restrict the number of laterally-arranged γ-tubulin subunits, and position them in the right geometry to template 13-pf MTs. Finally, our results show that 3–4 αβ-tubulin form the critical nucleus on γ-TuRC, not 1 or 13 which would have been expected from previous mechanistic hypotheses\(^20\). We find that on average 3–4 αβ-tubulin dimers assemble on γ-TuRC to form the first lateral αβ/αβ-tubulin contact and overcome the kinetic barrier that results from low longitudinal affinity between γ-αβ-tubulin on γ-TuRC. However, alternative reaction intermediates during nucleation from γ-TuRC may exist. In the future, it will be important to visualize the nucleation intermediates on γ-TuRC, develop molecular simulations with experimentally derived affinities at various interaction interfaces and evaluate whether additional effects from tubulin straightening play a significant role in MT nucleation in the cell.
The intermediate level of MT nucleation efficiency afforded by γ-TuRC allows other factors to further modulate its efficiency. As such, XMAP215 accelerates MT nucleation from γ-TuRC, while not altering the geometry of the αβ-tubulin nucleus on γ-TuRC or directly activating γ-TuRC. Future studies will be necessary to define the modes by which XMAP215 contributes to γ-TuRC-mediated MT nucleation, such as increasing the probability of the γ/αβ-tubulin interaction or promoting straightening of incoming tubulin dimers. Our findings suggest that influencing γ/αβ-tubulin interaction favorably or unfavorably may underlie a dominant mechanism for regulating nucleation in the cell by other, yet unidentified nucleation factors. Additionally, γ-TuRC’s activity is further regulated via accessory proteins such as CDK5RAP2, and NME72,20,39,40. While the mechanisms of these additional regulation layers are yet to be defined, the insights on MT nucleation by γ-TuRC and XMAP215 provide an essential basis to build upon. Finally, this work opens the door to reconstitute cellular structures in vitro using MT nucleation from γ-TuRC/XMAP215 to further our understanding of how the cytoskeleton is generated to support cell function.
Supplementary Information

Supplementary Information includes four figures and ten videos.

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Author contributions

A.T. designed and performed research, analyzed the data and wrote the manuscript. S.P., J.W.S. and H.A.S. supervised research and wrote the manuscript.

Competing financial interests

The authors declare no competing financial interests.

Abbreviations List

- Microtubule (MT)
- Microtubule associated protein (MAP)
- Gamma-tubulin (γ-tubulin) and Gamma-tubulin ring complex (γ-TuRC)
- Gamma-tubulin complex protein (GCP)
240  Protofilament (pf)

241  Electron microscopy (EM)
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Methods

Purification of recombinant proteins

C-terminal GFP was replaced with mCherry tag in the pET21a vector carrying EB1\textsuperscript{41}. Full-length TPX2 with N-terminal Strep II-6xHis-GFP-TEV site tags was cloned into pST50Tr-STRHISNDHFR (pST50) vector\textsuperscript{42} using Gibson Assembly (New England Biolabs). N-terminal 6xHis-tagged, \textit{Xenopus laevis} Stathmin 1A was a gift from Christiane Wiese (University of Madison). N-terminal tagged 6xHis-TEV MCAK plasmid was a gift from Ryoma Ohi\textsuperscript{43}. Wild-type XMAP215 with C-terminal GFP-7xHis plasmid was a gift from Simone Reber\textsuperscript{44} and was used to clone XMAP215 with C-terminal SNAP-TEV-7xHis-StrepII tags, first into pST50 vector and further into pFastBac1 vector. TOG5-CT truncation of XMAP215 was produced by cloning amino acids 1091-2065 into pST50 vector with C-terminal GFP-7xHis-Strep tags. Human $\gamma$-tubulin TEV-Strep II-6xHis tags was codon-optimized for Sf9 expression, synthesized (Genscript), and further cloned into pFastBac1 vector.

EB1, TPX2, Stathmin and XMAP215 TOG5-CT used in this study were expressed in \textit{E. coli} Rosetta2 cells (EMD Millipore) by inducing with 0.5-1 mM IPTG for 12-18 hours at 16°C or 7 hours at 25°C. Wild-type XMAP215, MCAK and $\gamma$-tubulin were expressed and purified from Sf9 cells using Bac-to-Bac system (Invitrogen). The cells were lysed (EmulsiFlex, Avestin) and \textit{E. coli} lysate was clarified by centrifugation at 13,000 rpm in Fiberlite F21-8 rotor (ThermoFisher) and Sf9 cell lysate at 50,000 rpm in Ti70 rotor (Beckman Coulter) for 30-45 minutes.

EB1 and Stathmin were purified using His-affinity (His-Trap HP, GE Healthcare) by first binding in binding buffer (20mM NaPO\textsubscript{4} pH 8.0, 500mM NaCl, 30mM Imidazole, 2.5mM PMSF, 6mM BME) and eluting with 300mM Imidazole, followed by gel filtration (HiLoad 16/600
Superdex, GE Healthcare) into CSF-XB buffer (100mM KCl, 10mM K-HEPES, 5mM K-EGTA, 1mM MgCl₂, 0.1mM CaCl₂, pH 7.7 with 10% w/v sucrose).

TPX2 was first affinity purified using Ni-NTA beads in binding buffer (50mM Tris-HCl pH 8.0, 750mM NaCl, 15mM Imidazole, 2.5mM PMSF, 6mM BME) and eluted with 200mM Imidazole. All protein was pooled and diluted 4-fold to 200mM final NaCl. Nucleotides were removed with a Heparin column (HiTrap Heparin HP, GE Healthcare) by binding protein in 250mM NaCl and isocratic elution in 750mM NaCl, all solutions prepared in Heparin buffer (50mM Tris-HCl, pH 8.0, 2.5mM PMSF, 6mM BME). Peak fractions were pooled and loaded on to Superdex 200 pg 16/600, and gel filtration was performed in CSF-XB buffer.

MCAK was first affinity purified by binding to His-Trap HP (GE Healthcare) in binding buffer (50mM NaPO₄, 500mM NaCl, 6mM BME, 0.1mM MgATP, 10mM Imidazole, 1mM MgCl₂, 2.5mM PMSF, 6mM BME, pH to 7.5), eluting with 300mM Imidazole, followed by gel-filtration (Superdex 200 10/300 GL, GE Healthcare) in storage buffer (10 mM K-HEPES pH 7.7, 300 mM KCl, 6mM BME, 0.1 mM MgATP, 1mM MgCl₂, 10% w/v sucrose).

XMAP215-GFP was purified using His-affinity (His-Trap, GE Healthcare) by binding in buffer (50mM NaPO₄, 500mM NaCl, 20mM Imidazole, pH 8.0) and eluting in 500mM Imidazole. Peak fractions were pooled and diluted 5-fold with 50mM Na-MES pH 6.6, bound to a cation-exchange column (Mono S 10/100 GL, GE Healthcare) with 50mM MES, 50mM NaCl, pH 6.6 and eluted with a salt-gradient up to 1M NaCl. Peak fractions were pooled and dialyzed into CSF-XB buffer. SNAP-tagged XMAP215 was first affinity purified with StrepTrap HP (GE Healthcare) with binding buffer (50mM NaPO₄, 270mM NaCl, 2mM MgCl₂, 2.5mM PMSF, 6mM BME, pH 7.2), eluted with 2.5mM D-desthiobiotin, and cation-exchanged (Mono S 10/100 GL). Peak fractions were pooled, concentrated and reacted with 2-molar excess SNAP-substrate Alexa-488.
dye (S9129, NEB) overnight at 4°C, followed by purification via gel filtration (Superdex 200 10/300 GL) in CSF-XB buffer. Approximately 70% labeling efficiency of the SNAP-tag was achieved.

γ-tubulin was purified by binding to HisTrap HP (GE Healthcare) in binding buffer (50 mM KPO$_4$ pH 8.0, 500 mM KCl, 1 mM MgCl$_2$, 10% glycerol, 5mM Imidazole, 0.25 µM GTP, 5 mM BME, 2.5mM PMSF), washing first with 50 mM KPO$_4$ pH 8.0, 300 mM KCl, 1 mM MgCl$_2$, 10% glycerol, 25 mM imidazole, 0.25 µM GTP, 5 mM BME), and then with 50 mM K-MES pH 6.6, 500 mM KCl, 5mM MgCl$_2$, 10% glycerol, 25 mM imidazole, 0.25 µM GTP, 5 mM BME) and eluted in 50 mM K-MES pH 6.6, 500 mM KCl, 5mM MgCl$_2$, 10% glycerol, 250 mM imidazole, 0.25 µM GTP, 5 mM BME. Peak fractions were further purified with gel filtration (Superdex 200 10/300 GL) in buffer 50 mM K-MES pH 6.6, 500 mM KCl, 5mM MgCl$_2$, 1 mM K-EGTA, 1 µM GTP, 1 mM DTT.

All proteins were flash-frozen and stored at -80°C, and their concentration was determined by analyzing a Coomassie-stained SDS-PAGE against known concentration of BSA (A7906, Sigma).

**Purification, biotinylated and fluorescent labeling of γ-TuRC**

Endogenous γ-TuRC was purified from *Xenopus* egg extracts and labeled with the following steps at 4°C. 7-8 ml of meiotic extract from *Xenopus* laevis eggs, prepared as described previously$^{45,46}$, was first diluted 5-fold with CSF-XBg buffer (10mM K-HEPES, 100mM KCl, 1mM MgCl$_2$, 5mM K-EGTA, 10% w/v sucrose, 1mM DTT, 1mM GTP, 10 µg/ml LPC protease inhibitors, pH 7.7), centrifuged to remove large aggregates at 3500 rpm (Thermo Sorvall Legend XTR) for 10 minutes, and the supernatant filtered sequentially with 1.2 µm and 0.8 µm Cellulose Acetate filters.
γ-TuRC was precipitated by incubating with 6.5% w/v PEG-8000k (Sigma) for 30 minutes and centrifuged at 17,000 rpm (SS-34 rotor, ThermoScientific) for 20 minutes. γ-TuRC-rich pellet was resuspended in CSF-XB buffer with 0.05% v/v NP-40 using a mortar & pestle homogenizer, PEG was removed via centrifugation at 136,000 xg for 7 minutes in TLA100.3 (Beckman Ultracentrifuge), and supernatant was precleared by incubating with Protein A Sepharose beads (GE LifeSciences #17127901) for 20 minutes. Beads were removed, γ-TuRC was incubated with 4-5 mg of a polyclonal antibody custom-made against C-terminal residues 413-451 of X. laevis γ-tubulin (Genscript) for 2 hours on gentle rotisserie, and further incubated with 1ml washed Protein A Sepharose bead slurry for 2 hours. γ-TuRC-bound beads were washed sequentially with 30 ml of CSF-XBg buffer, 30 ml of CSF-XBg buffer with 250 mM KCl (high salt wash), 10 ml CSF-XBg buffer with 5mM ATP (removes heat-shock proteins), and finally 10 ml CSF-XBg buffer before labeling. For biotinylation of γ-TuRC, beads were incubated with 25 μM NHS-PEG4-biotin (A39259, ThermoFisher) in CSF-XBg buffer for 1 hour at 4°C, and unbound biotin was removed by washing with 30 ml CSF-XBg buffer prior to elution step. For combined fluorescent and biotin labeling of γ-TuRC, the wash step after ATP-wash consisted of 10 ml of labelling buffer (10mM K-HEPES, 100mM KCl, 1mM MgCl₂, 5mM K-EGTA, 10% w/v sucrose, 0.5mM TCEP, 1mM GTP, 10 μg/ml LPC, pH 7.2) and fluorescent labelling was performed by incubating the beads with 1 μM Alexa-568 C₅ Maleimide (A20341, ThermoFisher). Unreacted dye was removed with 10 ml CSF-XBg buffer, beads were incubated with 25 μM NHS-PEG4-biotin (A39259, ThermoFisher) in CSF-XBg buffer for 1 hour at 4°C, and unreacted biotin removed with 30 ml CSF-XBg buffer. Labeled γ-TuRC was eluted by incubating 2-3ml of γ-tubulin peptide (residues 413-451) at 0.4mg/ml in CSF-XBg buffer with beads overnight. After 10-12 hours, γ-TuRC was collected by adding 1-2ml
CSF-XBg buffer to the column, concentrated to 200 µl in 30k NMWL Amicon concentrator (EMD Millipore) and layered onto a continuous 10-50 w/w % sucrose gradient prepared in a 2.2 ml ultracentrifuge tube (11x34 mm, Beckman Coulter) using a two-step program in Gradient Master 108 machine. Sucrose gradient fractionation of γ-TuRC was performed by centrifugation at 200,000xg in TLS55 rotor (Beckman Coulter) for 3 hours. The gradient was fractionated from the top in 11-12 fractions using wide-bore pipette tips and peak 2-3 fractions were identified by immunoblotting against γ-tubulin with GTU-88 antibody (Sigma). γ-TuRC was concentrated to 80 µl in 30k NMWL Amicon concentrator (EMD Millipore) and fresh purification was used immediately for single molecule assays. Cryo-preservation of γ-TuRC molecules resulted in loss of ring assembly and activity.

Assessment of γ-TuRC with protein gel, immunoblot and negative stain electron microscopy

To assess the purity of γ-TuRC, 3-5 µl of purified γ-TuRC was visualized on an SDS-PAGE with SYPRO Ruby stain (ThermoFisher) following the manufacturer’s protocol. Biotinylated subunits of γ-TuRC were assessed by immunoblotting with Streptavidin-conjugated alkaline phosphatase (S921, ThermoFisher). γ-TuRC purification was also assessed by visualizing using electron microscopy. 4 µl of peak sucrose gradient fraction of γ-TuRC was pipetted onto CF400-Cu grids (Electron Microscopy Sciences), incubated at room temperature for 60 seconds and then wicked away. 2% uranyl acetate was applied to the grids for 30 seconds, wicked away, and the grids were air-dried for 10 minutes. The grids were imaged using Phillips CM100 TEM microscope at 64000x magnification.

Preparation of functionalized coverslips
22x22 mm, high precision coverslips (170±5 µm, Carl Zeiss, catalog # 474030-9020-000) were functionalized for single molecule assays based on a recent protocol\textsuperscript{23,47} with specific modifications. Briefly, coverslips were labelled on the surface to be functionalized by scratching “C” on right, bottom corner, placed in Teflon racks, sonicated with 3N NaOH for 30 minutes, rinsed with water and sonicated in piranha solution (2 parts of 30 w/w % hydrogen peroxide and 3 parts sulfuric acid) for 45 minutes. Coverslips were rinsed thrice in water, and all water was removed by spin drying completely in a custom-made spin coater. Pairs of coverslips were made to sandwich 3-glycidoxypropyl trimethoxysilane (440167, Sigma) on the marked sides, placed in glass petri dishes, and covalent reaction was performed in a lab oven at 75°C for 30 minutes. Coverslips were incubated for 15 minutes at room temperature, the sandwiches were separated, incubated in acetone for 15 minutes, then transferred to fresh acetone and quickly dried under nitrogen stream. Coverslip sandwiches were prepared with a small pile of well mixed HO-PEG-NH\textsubscript{2} and 10% biotin-CONH-PEG-NH\textsubscript{2} (Rapp Polymere) in glass petri dishes, warmed to 75°C in the lab oven until PEG melts, air bubbles were pressed out and PEG coupling was performed at 75°C overnight. The following day, individual coverslips were separated from sandwiches, sonicated in MilliQ water for 30 minutes, washed further with water until no foaming is visible, dried with a spin dryer, and stored at 4°C. Functionalized coverslips were used within 1 month of preparation.

Imaging chambers were prepared by first assembling a channel on glass slide with double sided tape strips (Tesa) 5 mm apart, coating the channel with 2mg/ml PLL(20)-g[3.5]-PEG(2) (SuSOS) in dH\textsubscript{2}O, incubating for 20 minutes, rinsing out the unbound PEG molecules with dH\textsubscript{2}O and drying the glass slide under the nitrogen stream. A piece of functionalized coverslip was cut
with the diamond pen and assembled functionalized face down on imaging chamber. The prepared chambers were stored at 4°C and used within a day of assembly.

Microtubule nucleation assay with purified γ-TuRC, microscopy and data analysis

The imaging channel was prepared as follows. First, 5% w/v Pluronic F-127 in dH$_2$O was introduced in the chamber (1 vol = 50 µl) and incubated for 10 minutes at room temperature. The chamber was washed with 2 vols of assay buffer (80mM K-PIPES, 1mM MgCl$_2$, 1mM EGTA, 30mM KCl, 0.075% w/v methylcellulose 4000 cp, 1% w/v D-(-)-glucose, 0.02% w/v Brij-35, 5mM BME, 1mM GTP) with 0.05 mg/ml κ-casien (casein buffer), followed by 1 vol of 0.5 mg/ml NeutrAvidin (A2666, ThermoFisher) in casein buffer, incubated on a cold block for 3 minutes, and washed with 2 vols of BRB80 (80mM K-PIPES, 1mM MgCl$_2$, 1mM EGTA pH 6.8). 5-fold dilution of γ-TuRC in BRB80 was introduced in the flow chamber and incubated for 10 minutes. Unattached γ-TuRC molecules were washed with 1 vol of BRB80.

During the incubations, nucleation mix was prepared containing desired concentration of αβ-tubulin (3.5-21 µM) purified from bovine brain with 5% Cy5-labeled tubulin along with 1mg/ml BSA (A7906, Sigma) in assay buffer, centrifuged for 12 minutes in TLA100 (Beckman Coulter) to remove aggregates, a final 0.68 mg/ml glucose oxidase (SERVA, catalog # SE22778), 0.16 mg/ml catalase (Sigma, catalog # SRE0041) was added, and reaction mixture was introduced into the flow chamber containing γ-TuRC.

Total internal reflection fluorescence (TIRF) microscopy and analysis of microtubule nucleation from γ-TuRC
Nucleation of MTs was visualized with inverted Nikon TiE TIRF microscope using a 100X, 1.49 NA TIRF objective. An objective heater collar was attached (Bioptechs, model 150819-13) and the temperature set-point of 33.5°C was used for experiments. Time-lapse videos were recorded for 10 minutes at 0.5-1 frame per second using Andor iXon DU-897 camera with EM gain of 300 and exposure time of 50-200 ms each frame. Reference time-point zero (0 seconds) refers to when the reaction was incubated at 33.5°C on the microscope, and for most reactions, imaging was started within 30 seconds.

Growth speed of the plus-ends of MTs nucleated by γ-TuRC was measured by generating kymographs in ImageJ. Region of interest (ROI) for individual MTs were selected and resliced to generate length-time plot, a line was fit to the growing MT, the slope of line represents growth speed. The kinetics of MT nucleation from γ-TuRC was measured as follows. A kymograph was generated for every MT nucleated in the field of view. For most nucleation events, the time of nucleation of the MT was obtained from observing the kymograph and manually recording the initiation time point (see Fig. 1C for examples). For MTs where nucleation occurred before the timelapse movie began or where the initiation was not clearly observed in the kymograph, the shortest length of the MT that was clearly visible in the timelapse was measured and measured average growth speed of MTs was used to estimate the time of nucleation. We verified that this procedure accurately estimates the nucleation time for test case MTs where the nucleation event was visible. The measurement of number of MTs ($N(t)$) nucleated versus time was generated from a manual log containing the nucleation time for all MTs observed in the field of view, and a representative set of curves is displayed in Fig. 1F. A straight line was fit to the initial (linear) region of each $N(t)$ versus $t$ curve, rate of nucleation was obtained slope of each linear fit, and its power-law relation with tubulin concentration was obtained and reported (Fig. 1G).
Spontaneous microtubule nucleation and data analysis

Spontaneous MT assembly was visualized similar to γ-TuRC-mediated nucleation with the following changes. The pluronic, casein and NeutrAvidin incubations were performed identical to γ-TuRC nucleation assay but instead of attaching γ-TuRCs, sucrose-based buffer (of the same composition as used for γ-TuRC elution) was diluted 5-fold with BRB80, introduced in the flow chamber and incubated for 10 minutes. Washes were performed with 1 vol of BRB80, nucleation mix was added, and imaging was performed as described above. MTs nucleate spontaneously in solution fall down on the coverslip due to depletion forces during the 10 minutes of visualizing the reaction. The number of MTs nucleated in the field of view were counted manually and plotted in Fig. II.

Preparation and microtubule assembly from blunt microtubule seeds

Blunt MTs were prepared with GMPCPP nucleotide in two polymerization cycles as described recently22. Briefly, a 50 µl reaction mixture was prepared with 20 µM bovine brain tubulin with 5% Alexa-568 labeled tubulin and 5% biotin-labeled tubulin, 1mM GMPCPP (Jena Bioscience) in BRB80 buffer, incubated on ice for 5 minutes, then incubated on 37°C for 30 minutes to polymerize MTs, and MTs were pelleted by centrifugation at 126,000 xg for 8 minutes at 30°C in TLA100 (Beckman Coulter). Supernatant was discarded, MTs were resuspended in 80% original volume of BRB80, incubated on ice for 20 minutes to depolymerize MTs, fresh GMPCPP was added to final 1mM, incubated on ice for 5 minutes, a second cycle of polymerization was performed by incubating the mixture at 37°C for 30 minutes, and MTs were pelleted again by centrifugation. Supernatant was discarded and MTs were resuspended in 200 µl warm BRB80,
flash frozen in liquid nitrogen in 5µl aliquots, stored at -80°C and found to be stable for months. To verify that these MT seeds have blunt ends, frozen aliquots were quickly thawed at 37°C, diluted 20-fold with warm BRB80, and incubated at room temperature for 30 minutes to ensure blunt ends as described previously. MTs were pipetted onto CF400-Cu grids (Electron Microscopy Sciences), incubated at room temperature for 60 seconds and then wicked away. 2% uranyl acetate was applied to the grids for 30 seconds, wicked away, and the grids were air-dried for 10 minutes. The grids were imaged using Phillips CM100 TEM microscope at 130000 x magnification and most MT ends were found to be blunt.

To assay MT assembly from blunt MT seeds, MT assembly experiments similar to γ-TuRC nucleation assays were performed with the following variation. A lower concentration 0.05 mg/ml NeutrAvidin (A2666, ThermoFisher) was attached, and washes were performed with warm BRB80 prior to attaching MTs. One aliquot of MT seeds was thawed quickly, diluted to 100-fold with warm BRB80, incubated in the chamber for 5 minutes, unattached seeds were washed with 1 vol of warm BRB80, and the slide was incubated at room temperature for 30 minutes to ensure blunt MT ends. Wide bore pipette tips were used for handling MT seeds to minimize the shear forces that may result in breakage of MTs. Nucleation mix was prepared as described above and a low αβ-tubulin concentration (1.4-8.7 µM) was used. MT assembly from blunt seeds was observed immediately after incubating the slide on the objective heater. Imaging and analysis were performed as described above for to γ-TuRC nucleation assays. However, the probability curves for MT assembly were obtained (Fig. 2B) by normalizing for the total number of seeds observed in the field of view. Rate of assembly was plotted against [tubulin concentration – C*], where C* represents the critical tubulin concentration below which MT ends do not polymerize obtained directly from experimental measurements (Fig. S2A-B).
Electron microscopy of γ-tubulin filaments in vitro

Purified γ-tubulin was observed to form higher order oligomers previously using analytical gel filtration. γ-tubulin filaments were prepared by diluting pure γ-tubulin to 1-5 µM to the buffer 50mM K-MES pH 6.6, 5mM MgCl₂, 1mM EGTA, 100mM KCl. γ-tubulin mixture were pipetted onto CF400-Cu grids (Electron Microscopy Sciences), incubated at room temperature for 60 seconds and then wicked away. 2% uranyl acetate was applied to the grids for 30 seconds, wicked away, and the grids were air-dried for 10 minutes. The grids were imaged using Phillips CM100 TEM microscope at 130000 x magnification and γ-tubulin filaments were seen to form. At 500 mM KCl, γ-tubulin filaments were not seen.

Nucleation of microtubules from purified γ-tubulin

MT assembly experiments from purified γ-tubulin was performed similar to γ-TuRC nucleation assays described above with following variation. No avidin was attached to the coverslips, and varying concentration of γ-tubulin was prepared by diluting purified γ-tubulin in a high salt buffer (50mM K-MES pH 6.6, 500mM KCl, 5mM MgCl₂, 1mM EGTA), centrifuging to remove aggregates separately for 12 minutes in TLA100 before adding to the nucleation mix containing 15 µM αβ-tubulin (5% Cy5-labeled) with BSA, glucose oxidase and catalase as described above. The reaction mixture was introduced into the flow chamber and imaged via TIRF microscopy. A large number of MTs get nucleated immediately in the presence of 250 nM-1000 nM γ-tubulin.

Measurement of affinity between purified γ-tubulin and αβ-tubulin
Interaction assays between αβ-tubulin and γ-tubulin were performed with biolayer interferometry using Octet RED96e (ForteBio) instrument in an 8-channel plate format. The plate temperature was held at 33°C and the protein samples were shaken at 400 rpm during the experiment. First, Streptavidin or anti-His antibody coated biosensors (ForteBio) were rinsed in interaction buffer (50mM K-MES pH 6.6, 100mM KCl, 5mM MgCl₂, 1mM EGTA, 0.05% Tween20, 1mM GTP). 100 nM biotin-labeled αβ-tubulin, or blank buffer, was bound to Streptavidin sensor, or 200 nM His-tagged γ-tubulin to anti-His sensor until loaded protein results in a wavelength shift (Δλ) of 3 nm. Unbound protein was removed by rinsing the sensor in interaction buffer, and interaction with αβ-tubulin was measured by incubating the sensor containing αβ-tubulin, γ-tubulin or buffer with 0-35 µM unlabeled αβ-tubulin in interaction buffer for 5 minutes. Δλ (nm) was recorded as a measure of the amount of unlabeled αβ-tubulin that binds to the sensor. Longitudinal interaction occurs between αβ-tubulin dimers and the resulting protofilaments were verified by visualizing the αβ-tubulin sample stained with 2% uranyl acetate using electron microscopy as described above (Fig. S2D).

Simulation of site occupation on γ-TuRC by αβ-tubulin dimers

A simulation was performed in MATLAB for occupation of sites on γ-TuRC by αβ-tubulin dimers. A circular grid was simulated with 13 empty positions that were occupied one per unit time stochastically such that a new position was selected by uniform random number generator and filled. If a previously filled position was selected, a different position was selected by the random number generator. The sequence in which the sites were occupied was followed. For each simulation, the total number of sites that were occupied when the first two neighboring sites are
filled was recorded. The simulation was repeated 10,000 times and the probability of occurrence of first neighbor contact versus number of sites occupied is displayed in Fig. 2H.

**Measuring the effect of microtubule associated proteins on γ-TuRC’s activity**

Effect of microtubule associated proteins (MAPs) was measured on γ-TuRC’s nucleation activity. γ-TuRC was attached on the coverslips using the setup described above and a control experiment was performed with identical reaction conditions for each protein tested. Nucleation mix was prepared containing 10.5 µM αβ-tubulin concentration (5% Cy5-labeled tubulin) as specified along with 1mg/ml BSA and oxygen scavengers, and either buffer (control), 10nM GFP-TPX2, 100nM EB1-mCherry, 5 µM Stathmin or 10nM MCAK was added. To test MCAK’s effect, the assay buffer additionally contained 1mM ATP. The reaction mixture containing tubulin and MAP at specified concentration was introduced into the flow chamber containing γ-TuRC, and MT nucleation was visualized by imaging the Cy5-fluorescent channel at 0.5-1 frames per second. For TPX2 and EB1, fluorescence intensity of the protein was simultaneously acquired. The number of MTs nucleated over time was measured as described above and the effect of protein on γ-TuRC’s nucleation activity was assessed by comparing nucleation curves with and without the MAP.

A similar set of experiments were performed to study the effect of XMAP215 on γ-TuRC-mediated nucleation with the single molecule assays with the following differences. 20 nM of XMAP215-GFP was added to nucleation mix prepared with 3.5-7 µM αβ-tubulin concentration (5% Cy5-label) in XMAP assay buffer (80mM K-PIPES, 1mM MgCl2, 1mM EGTA, 30mM KCl, 0.075% w/v methylcellulose 4000 cp, 1% w/v D-(+)-glucose, 0.007% w/v Brij-35, 5mM BME, 1mM GTP). MTs nucleated from attached γ-TuRC with and without XMAP215 were measured to assess the efficiency of nucleation induced by XMAP215 (Fig. 3C). To assess if C-terminal of
XMAP215 increases nucleation efficiency, wild-type XMAP215 was replaced with a C-terminal construct of XMAP215: TOG5-Cterminus-GFP in the described experiment.

To measure the kinetics of cooperative nucleation XMAP215 and γ-TuRC, a constant density of γ-TuRC was attached as described above and nucleation mix nucleation mix was prepared with a range of αβ-tubulin concentration between 1.6-7 μM (5% Cy5-label) with 20 nM of XMAP215-GFP in XMAP assay buffer, introduced into reaction chamber and MT nucleation was imaged immediately by capturing dual color images of XMAP215 and tubulin intensity at 0.5 frames per second.

Triple-color imaging of XMAP215, γ-TuRC and microtubules

For triple-color fluorescence assays (Fig. 3D), Alexa-568 and biotin-conjugated γ-TuRC was first attached to coverslips as described above with the following variation: 0.05 mg/ml of NeutrAvidin was used for attaching γ-TuRC. Nucleation mix was prepared with 7 μM αβ-tubulin (5% Cy5-label), 10 nM Alexa-488 SNAP-tagged XMAP215 with BSA and oxygen scavengers in XMAP assay buffer (80mM K-PIPES, 1mM MgCl₂, 1mM EGTA, 30mM KCl, 0.075% w/v methylcellulose 4000 cp, 1% w/v D-(+)-glucose, 0.007% w/v Brij-35, 5mM BME, 1mM GTP) and introduced into the reaction chamber containing attached γ-TuRC. Three-color imaging per frame was performed with sequential 488, 568 and 647 nm excitation and images were acquired with EMCCD camera at 0.3 frames per second.
Figure 1. Microtubule nucleation by γ-TuRC.

(A) Schematic of γ-TuRC mediated nucleation based on template model. (B) Purified, biotinylated γ-TuRC molecules were attached and time-lapse of MT nucleation is shown. Arrows point to nucleation sites. Representative kymographs of MTs nucleated from γ-TuRC are displayed in (C). The experiment and analyses in (B-G) were repeated at least thrice with independent γ-TuRC preparations. (D) Titrating tubulin concentration with constant the density of γ-TuRC. MT nucleation from γ-TuRC begins at 7 µM tubulin. (E) MT plus-end growth speed increases linearly with tubulin concentration. Linear fit (red line) with shaded 95% confidence intervals is displayed, with critical concentration for polymerization as $C^* = 1.4$ µM. Inset: Number of MTs nucleated by γ-TuRCs within 120 seconds varies non-linearly with tubulin concentration. (F) Number of MTs nucleated ($N(t)$) over time ($t$) is plotted for varying tubulin concentration to obtain rate of nucleation as the slope of the initial part of the curves. (G) Number of tubulin dimers ($n$) in the critical nucleus on γ-TuRC was obtained as $3.7 \pm 0.5$ from the equation $\frac{dN}{dt} \bigg|_{t=0} = kC_{tub}^n$ displayed on a log-log plot. Data from two independent experiments was pooled and reported. (H) Spontaneous MT nucleation (schematized) was measured with increasing tubulin concentration and high concentrations. 14 µM tubulin is required. (I) Number of MTs ($N(t=\tau)$) nucleated spontaneously were plotted against tubulin concentration ($C_{tub}$). Power-law curve was fit as $N(t=\tau) = kC_{tub}^n$ and tubulin cooperativity (exponent) of $n = 8 \pm 1$ was obtained. Experiments were repeated twice independently with many supporting results and all data were pooled. Scale bars, 10 µm. See Figure S1 and Movies S1-S4.
Figure 2. Molecular mechanism for γ-TuRC-mediated microtubule nucleation.

(A) Schematic and a micrograph of pre-formed, blunt MT seeds is shown and MT assembly from them was observed (right) with varying tubulin concentration. (B) Cumulative probability of MT assembly from seeds ($p(t)$) over time ($t$) is plotted and rate of nucleation was obtained as the slope from initial part of the curves. (C) Tubulin dimers ($n$) needed for MT assembly from seeds was from the relation $\frac{dp}{dt}\bigg|_{t=0} = k(C_{tub} - C^+)^n$ displayed on a log-log plot. $n = 1.2 \pm 0.4$ showing non-cooperative assembly of tubulin. (D) MTs nucleate from purified γ-tubulin oligomers efficiently and (E) minus-ends of γ-tubulin-nucleated MTs remain capped while the plus-ends polymerize. (F) Molecular interaction between γ/αβ-tubulin was probed with bio-layer interferometry. Buffer (left), biotin-tagged αβ-tubulin (middle), or His-tagged γ-tubulin (right) were loaded on the probe as bait and untagged αβ-tubulin at 0-35 μM as prey. Wavelength shift, $\Delta \lambda$ (nm) indicated no binding between empty probe and αβ-tubulin or γ/αβ-tubulin, while that between αβ-/αβ-tubulin was observed and confirmed to be longitudinal (protofilament-wise, Fig. S2D). (G) Interface interaction model determines MT nucleation by γ-TuRC where lateral γ/γ-tubulin promote nucleation while low γ/αβ-tubulin affinity tunes nucleation. (H) Simulations were conducted where 13 sites on γ-TuRC were stochastically occupied by αβ-tubulins. For two αβ-tubulin subunits to form lateral bond by occupying neighboring sites, $3.7 \pm 1$ subunits bind on average on γ-TuRC, predicting the size of critical nucleus. Experiments and analyses were repeated at least twice independently with multiple supporting results. Scale bars, 10 μm. See Figure S2 and Movie S5-6.

Figure 3. Regulation of microtubule nucleation by TPX2 and XMAP215.
(A) A constant density of γ-TuRC molecules were attached and 10.5 μM tubulin ± 10nM GFP-TPX2 were added. MTs were counted (right plot) and TPX2 was did not affect γ-TuRC-mediated nucleation. Scale bar, 10 μm. (B) γ-TuRCs were attached and low concentration 3.5-7 μM ± 20nM XMAP215 was added. XMAP215 induces MT nucleation from γ-TuRC efficiently. (C) MT nucleation events were counted and plotted. Scale bar, 10 μm. (D) Sequence of events during cooperative MT nucleation by γ-TuRC and XMAP215 was visualized using labeled γ-TuRC (blue), XMAP215 (red) and tubulin (green). Time-lapse: γ-TuRC and XMAP215 form a complex prior to MT nucleation. XMAP215 variably resides on γ-TuRC for long (>100 seconds, kymograph 1) or short times (~3-10 seconds, kymograph 2) before MT nucleation and remains at the minus-end with 50% probability. Scale bar, 5 μm. (E) Titrating tubulin with constant γ-TuRC and XMAP215 concentration. XMAP215/γ-TuRC nucleate MTs from 1.6 μM tubulin. Number of MTs nucleated \( N(t) \) over time \( t \) is plotted (inset) and rate of nucleation was obtained. Tubulin dimers \( n \) in critical nucleus was obtained as 3.2 ± 1.2 and displayed on a log-log plot. The experiment was performed once for all concentrations denoted and supported by a number of additional experiments. The remaining experiments were repeated more than twice with independent γ-TuRC preparations with additional supporting results. See Figure S3-4 and Movies S7-10.
Supplementary Figure legends

Supplementary Figure 1. Controls for γ-TuRC-mediated and spontaneous microtubule nucleation.

(A-B) Protein gel (left) of purified γ-TuRC was stained with SYPRO Ruby stain and biotinylated sites on γ-TuRC visualized with alkaline phosphatase conjugated to avidin (right). Major, known γ-TuRC components were detected in the purified protein and GCP2/3 are heavily biotinylated during purification. Purified and biotinylated γ-TuRC was stained with uranyl acetate and visualized with transmission electron microscopy. Scale bar, 100nm. The experiments were repeated at least thrice with independent γ-TuRC preparations.

(C) Covalent-reaction of biotin with γ-TuRC does not affect the nucleation activity, as measured by attaching γ-TuRC with anti-Mozart1 antibody and comparing the number of MTs nucleated by untagged and biotinylated γ-TuRC. Scale bar, 10µm.

(D) Control reactions for γ-TuRC-mediated nucleation. MTs were nucleated by attaching purified γ-TuRC (left), adding control buffer (middle) or missing avidin in the reaction sequence (right). Robust MT nucleation only occurs with γ-TuRC attached to coverslips and not in control reactions. Scale bar, 10µm. See Movie S2.

(E) MTs were first nucleated from γ-TuRC with Alexa 568-labeled tubulin (cyan), followed by introduction of Cy5-labeled tubulin (magenta). New tubulin only incorporates on the freely growing, plus-end but not at the nucleated minus-end. Scale bar, 10µm. The experiment was performed more than three times.
(F) Two representative kymographs of spontaneously nucleated MTs are displayed, demonstrating that MTs grow from both the minus-end (dotted line) and the plus-end (solid line). Scale bar, 10 µm. See Movie S4.

(G) MTs nucleation from γ-TuRCs or spontaneously were compared at two tubulin concentrations: 10.5 µM and 14 µM. γ-TuRC nucleates 10-fold higher number of MTs than spontaneous assembly. The experiment was performed twice with many supporting results. See also Figure 1.

Supplementary Figure 2. Microtubule assembly from blunt seeds and filament formation by purified γ-tubulin.

(A) MT assembly (magenta) from MT seeds with blunt ends (cyan) was assayed. Tubulin concentration was titrated, and MT plus-end assembles starting from 2.45 µM tubulin, which is only slightly above the critical concentration of polymerization. Scale bar, 10 µm.

(B) Growth speed of MT plus-ends was measured from kymographs and critical concentration ($C^*$ = 1.4 µM) was determined from the linear fit (red line) with shaded 95% confidence intervals. The experiment and analyses in were repeated twice on independent days along with other supporting data. See also Figure 2 and Movie S5.

(C) γ-tubulin self-assembles into filaments at high concentration and low-salt (100 mM KCl) as imaged with negative-stain electron microscopy, whereas γ-tubulin filaments were not observed at high-salt (500 mM KCl). Scale bar, 100 nm.

(D) Transmission electron microscopy of bio-layer interferometry assay of Fig. 2F show that protofilaments of αβ-tubulin form. The experiment was repeated twice. Scale bar, 100 nm.
Supplementary Figure 3. Effect of microtubule associated proteins on γ-TuRC-mediated nucleation.

(A) γ-TuRC molecules were attached to coverslips and either tubulin alone (pseudo-colored as magenta, left) or tubulin with 100nM EB1-mCherry (pseudo-colored as cyan, right) was added to the reaction. Number of MTs nucleated were measured (right plot) and EB1 was observed to neither increase nor decrease γ-TuRC-mediation nucleation despite functioning as a catastrophe factor in vitro. The experiment was repeated at least twice with independent γ-TuRC preparation. See also Movie S8. Scale bar, 10µm.

(B) γ-TuRC molecules were attached to coverslips and either tubulin alone (left images), tubulin with 10nM MCAK (top right) or tubulin with 5µM Stathmin (bottom right) was added to the reaction. Both MCAK and Stathmin were observed to decrease the number of MTs nucleated because of their role in decreasing the net polymerization of a MT. The experiment was repeated at least twice with independent γ-TuRC preparations. Scale bar, 10µm.

Supplementary Figure 4. Cooperative microtubule nucleation XMAP215 and γ-TuRC.

(A) γ-TuRC molecules were attached and increasing concentration of tubulin was added with 20nM XMAP215. XMAP215 was found to induce MT nucleation from γ-TuRC molecules at even low tubulin concentration of 1.6-3.5 µM where γ-TuRCs alone do not nucleate MTs. See Figure 3E. Scale bar, 10µm.

(B) The role of C-terminal region of XMAP215 was tested in cooperative nucleation with γ-TuRC. MTs nucleated by γ-TuRC alone (left), γ-TuRC with 20nM full-length XMAP215 (middle) or γ-TuRC with 20nM C-terminal domain of XMAP215 were visualized. The C-terminal domains of
XMAP215 do not stimulate MT nucleation from γ-TuRC. The experiment was repeated twice with independent γ-TuRC preparations. Scale bar, 10µm.
Movie Legends

Movie 1. Microtubule nucleation from γ-TuRC complexes

γ-TuRC was attached to functionalized coverslips and MT nucleation was observed upon introducing fluorescent αβ-tubulin (gray). MTs nucleated from individual γ-TuRC molecules from zero length at 15μM αβ-tubulin and the plus-end of nucleated MTs polymerized, but not its minus-end. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 μm.

Movie 2. Microtubule nucleation from γ-TuRC is specific

γ-TuRC was immobilized on coverslips (leftmost panel) and MT nucleation was observed upon introducing fluorescent αβ-tubulin (gray). Control reactions where either no γ-TuRC was added (middle panel) or γ-TuRC was not specifically attached (rightmost panel) did not result in MT nucleation. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 μm.

Movie 3. γ-TuRC molecules nucleated microtubules efficiently

Constant density of γ-TuRC was attached while concentration of fluorescent αβ-tubulin was titrated (3.5-21μM) and MT nucleation was observed. γ-TuRC molecules nucleated MTs starting from 7μM tubulin and MT nucleation increased non-linearly with increasing tubulin concentration. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 μm.
Spontaneous microtubule nucleation occurs at high tubulin concentration

Concentration of fluorescent αβ-tubulin was titrated (7-21µM) and spontaneous MT nucleation was assayed. MTs nucleated spontaneously starting from high concentration of 14µM tubulin and MT nucleation increased non-linearly with tubulin concentration. Both plus- and minus-ends of the assembled MTs polymerize. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 µm.

Movie 5. Microtubule assembly from blunt plus-ends resembles polymerization

MTs with blunt ends (seeds, cyan) were generated and attached to functionalized coverslips. Varying concentration of fluorescent αβ-tubulin was added (1.4-8.7µM, pseudo-colored as magenta) and MT assembly from seeds was assayed. MTs assembled at concentration above 1.4µM tubulin, which is the minimum concentration needed for polymerization of MT plus-ends (C*). MT assembly from seeds increased linearly with the concentration of assembly-competent tubulin (C-C*). Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 µm.

Movie 6. Arrays of purified γ-tubulin nucleate microtubules

Purified γ-tubulin nucleated MTs. Fluorescent αβ-tubulin (10.5µM, colored as gray) was added to purified γ-tubulin at increasing concentration, and MT nucleation was assessed. MTs assembled from 250-1000 nM γ-tubulin, where γ-tubulin alone self-assembled into higher order oligomers and filaments in lateral γ/γ-tubulin arrays. Minus-ends of γ-tubulin-nucleated MTs did not polymerize, while the plus-ends did. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 µm.
Movie 7. TPX2 does not increase γ-TuRC’s microtubule nucleation activity

γ-TuRC was immobilized on coverslips and MT nucleation was observed upon introducing fluorescent αβ-tubulin (10.5µM, pseudo-colored as magenta) without or with 10nM GFP-TPX2 (pseudo-colored as cyan) in the left and right panels respectively. TPX2 bound along the nucleated MTs but did not increase the MT nucleation activity of γ-TuRC molecules. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 µm.

Movie 8. EB1 does not decrease the microtubule nucleation activity of γ-TuRC

γ-TuRC was immobilized on coverslips and MT nucleation was observed upon introducing fluorescent αβ-tubulin (10.5µM, pseudo-colored as magenta) without or with 100nM EB1-mCherry (pseudo-colored as cyan) in the left and right panels respectively. EB1 binds the plus-ends of nucleated MTs but did not decrease the MT nucleation activity of γ-TuRC molecules. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 µm.

Movie 9. XMAP215 increases microtubule nucleation activity of γ-TuRC

γ-TuRC was immobilized on coverslips and MT nucleation was assayed with low concentration of fluorescent αβ-tubulin (3.5µM and 7µM) without (top panels) or with 20nM XMAP215-GFP (bottom panels). XMAP215 decreased the minimal concentration of tubulin necessary to induce MT nucleation from γ-TuRC. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 µm.
Movie 10. Synergistic microtubule nucleation by γ-TuRC and XMAP215

Triple-color fluorescence microscopy was performed to observe the molecular sequence of events during MT nucleation from γ-TuRC and XMAP215. γ-TuRC (blue) and XMAP215 (red) formed a complex before MT nucleation occurred (pseudo-colored as green). For 50% of these events, XMAP215 remains on the nucleated minus-end. Elapsed time is shown in seconds, where time-point zero represents the start of reaction. Scale bar, 10 μm.
Figure 1

A γ-TuRC-mediated microtubule nucleation

B Microtubule nucleation from γ-TuRC in vitro

C Kymographs of γ-TuRC nucleated microtubules

D Titrating tubulin concentration in γ-TuRC-mediated nucleation

E Growth speed of plus-end and number of microtubules

F Kinetics of microtubule nucleation from γ-TuRC

G Cooperativity of tubulin dimers for γ-TuRC-mediated nucleation (n)

H Spontaneous microtubule nucleation

I Cooperativity of tubulin dimers for spontaneous nucleation (n)
**Figure 2**

A Microtubule assembly from stable seeds *in vitro*

Blunt plus-end  ➔  Growing end

B Kinetics of assembly from seeds

![Graph showing the probability of microtubule assembly from seeds over time.](Image)

C Tubulin dimers for microtubule generation from seeds (*n*)

![Graph showing the log(nucleation rate) vs. log(tubulin concentration-C*) relationship.](Image)

D Nucleation of microtubules by purified γ-tubulin

![Images of 10.5μM tubulin, buffer, 125nM γ-tubulin, 500nM γ-tubulin, 1000nM γ-tubulin, 250nM γ-tubulin, 100s, 450s, and Cy5-labeled tubulin.](Images)

E Kymographs of γ-tubulin nucleated microtubules

![Kymographs of γ-tubulin nucleated microtubules.](Images)

F Probing interaction between γ-tubulin and αβ-tubulin

![Images showing the Δλ (nm) over time for different tubulin concentrations.](Images)

G Interface interaction model

![Diagram showing the interface interaction model of γ-tubulin and αβ-tubulin.](Image)

H Simulation of αβ-tubulin binding on γ-TuRC sites

![Diagram showing the simulation of αβ-tubulin binding on γ-TuRC sites.](Image)
A TPX2 does not influence γ-TuRC-mediated nucleation

B XMAP215 decreases the nucleation barrier from γ-TuRCs

C Kinetics of microtubule nucleation from γ-TuRC with XMAP215

D XMAP215 binds γ-TuRC prior to microtubule nucleation

E Cooperativity of tubulin dimers for co-nucleation by γ-TuRC and XMAP215 (n)
Figure S1 γ-TuRC purification and quality control reactions

A Purified, biotinylated γ-TuRC

B Negative stained γ-TuRC under electron microscope

C Biotinylation of does not decrease the nucleation activity of γ-TuRC

D Control reactions for in vitro nucleation by γ-TuRC

E Tubulin swap on γ-TuRC nucleated MTs

F Kymographs of spontaneously nucleated microtubules

G Nucleation kinetics from γ-TuRCs versus spontaneous assembly
Figure S2

A  Titrating tubulin on stable microtubule seeds

B  Growth speed of microtubule plus-ends

C  γ-tubulin filaments form under physiological salt at high concentration *in vitro*

D  αβ-tubulin protofilaments form under biorlayer interferometry
Figure S3

A Effect of EB1 on γ-TuRC-mediated nucleation

10.5 μM Cy5-tubulin

10.5 μM Cy5-tubulin + 100 nM EB1-mCherry

340 s

Tubulin

Number of microtubules nucleated

Time (seconds)

0 100 200

γ-TuRC + EB1

γ-TuRC

B Effect of MCAK and Stathmin on γ-TuRC-mediated nucleation

10.5 μM Cy5-tubulin

10.5 μM Cy5-tubulin + 10 nM MCAK or 5 μM Stathmin

316 s

γ-TuRC

γ-TuRC + 5 μM Stathmin

Number of microtubules nucleated

Time (seconds)

0 100 200

γ-TuRC

γ-TuRC + 10 nM MCAK

γ-TuRC + 5 μM Stathmin
Figure S4

A  Titration of tubulin concentration in γ-TuRC/XMAP215-mediated co-nucleation

Avidin

PEG

γ-TuRC

+ XMAP215 + [tubulin]

Glass

[Images showing micropipette data with varying concentrations of tubulin (1.6μM, 2.4μM, 3.5μM, 5.2μM, 7μM, 8.7μM) for 270 s]

B  C-terminal region of XMAP215 does not stimulate nucleation from γ-TuRC

[Images showing micropipette data with γ-TuRC and XMAP215 C-terminus at 540 s]

γ-TuRC

Avidin

PEG

Glass

10.5μM tubulin + XMAP215

γ-TuRC

+ XMAP215 WT

γ-TuRC

+ XMAP215 C-terminus