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Simulations of Tubulin Sheet Polymers as Possible Structural Intermediates in Microtubule Assembly

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Abstract

The microtubule assembly process has been extensively studied, but the underlying molecular mechanism remains poorly understood. The structure of an artificially generated sheet polymer that alternates two types of lateral contacts and that directly converts into microtubules, has been proposed to correspond to the intermediate sheet structure observed during microtubule assembly. We have studied the self-assembly process of GMPCPP tubulins into sheet and microtubule structures using thermodynamic analysis and stochastic simulations. With the novel assumptions that tubulins can laterally interact in two different forms, and allosterically affect neighboring lateral interactions, we can explain existing experimental observations. At low temperature, the allosteric effect results in the observed sheet structure with alternating lateral interactions as the thermodynamically most stable form. At normal microtubule assembly temperature, our work indicates that a class of sheet structures resembling those observed at low temperature is transiently trapped as an intermediate during the assembly process. This work may shed light on the tubulin molecular interactions, and the role of sheet formation during microtubule assembly.

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Introduction

Microtubules are one of the three major cytoskeleton components in eukaryotic cells [1,2]. They are hollow cylinders consisting of about 13 parallel protofilaments (PF) formed by the head-to-tail assembly of αβ-tubulin heterodimers. Microtubules play important roles in many eukaryotic cellular processes, including intracellular transport, cell motility, mitosis and meiosis. Microtubule dynamic instability, the phenomenon by which a microtubule switches stochastically between assembly and disassembly phases, is known to be a key property for microtubule function. The regulation of microtubule dynamics has been shown to be both of great biological significance during cell division, and of outstanding pharmaceutical value in tumor therapy. For example, Taxol®, the most widely used anticancer agent, targets tubulin and alters microtubule dynamics resulting in mitotic arrest. Therefore, studying the microtubule assembly/disassembly processes is of great relevance for both biological and pharmaceutical purposes.

To explain the process and mechanism of microtubule assembly, various models have been proposed by both experimentalists and theorists [3,4,5,6,7]. In the most simplistic textbook model, during the microtubule assembly process αβ-tubulin heterodimers add one by one onto the growing end of a microtubule. Most of the existing theoretical work is based on this model [4]. However, a number of experimental observations challenge this view. In 1970s Erickson reported an intermediate sheet structure during microtubule assembly (see also Fig. 1a) [6]. He proposed that tubulins first form a two-dimensional open sheet, which in turn closes into tubes (see Fig. 1a). Several other groups observed that fast growth of existing microtubules occurs via the elongation of a gently curved sheet-like structure at the growing end both in vitro and in vivo [6,7,8]. Using cryo-electron microscopy, Wang and Nogales reconstructed the structure of a curved sheet assembly of GMPCPP-tubulin stabilized by low temperature and high concentration of magnesium [7,8,9]. The use of GMPCPP avoids the complexity due to GTP hydrolysis. This assembly could then directly convert into microtubules by raising the temperature. The authors proposed that it corresponds structurally to the sheet at growing microtubule ends observed by Chrétién and others [7,9]. In this structure tubulin molecules form slightly curved PFs, in the same head-to-tail manner as those in microtubules. However the PFs are paired, with lateral interactions within one pair being indistinguishable from those in microtubules, but with distinct contacts between pairs [5]. Importantly, relative longitudinal displacements between neighboring PFs (“stagger”) are the same as in microtubules, indicating that no longitudinal sliding is needed during the sheet-microtubule transition, in agreement with the direct conversion from one to the other. In the remaining of the paper we call this polymer form “ribbon”, and reserve the term “sheet” for the observed structure.
at the end of a growing microtubule. In this work we suggest that the sheet may contain a class of tubulin structures that include the ribbon, all of which contain alternative lateral bonds different from those observed in microtubules. It is important to mention that in the literature the expression “sheet structure” has been used to refer to a protruding end of an incomplete microtubule [4], with no structural difference in the individual dimers or their interactions with respect to that in the microtubule itself, unlike the two-dimensional sheet of Chrétién and coworkers or the stable ribbon assembly of Wang and Nogales.

Wang and Nogales obtained the sheet structure by stabilizing it at low temperatures. An increase in temperature results in the direct conversion of these structures into microtubules. On decreasing the temperature a GMPCPP microtubule converts into the ribbon structure through peeling (Wang and Nogales, unpublished result; also in [10]). This observation implies that the sheet is thermodynamically more stable than the MT at low temperature, but is less stable at higher temperature (Fig. 1b). The conversion resembles a phase transition, which explains the observed sharp temperature dependence [11]. However, the sheet structure is short-lived in conditions under which MTs are formed, suggesting it as a kinetic intermediate [6,7].

The structural observations of Wang and Nogales raised several questions. How can a ribbon structure with alternating lateral interactions be formed during the assembly of tubulins? What is the relation between the ribbon structure and the sheet structures observed at the growing end of a microtubule at physiological conditions? What is the mechanism of the sheet-to-microtubule transition? If the sheet structure is indeed an intermediate in microtubule assembly in vivo, is there any biological function for it?

Due to the lack of detailed, atomic formation for the sheet, the ribbon, or the microtubule, as well as detailed kinetic studies, in this work we take an inversed problem approach. First we find out a set of minimal requirements for the system properties to reproduce the experimental observations, specifically the structures of Wang and Nogales. Then we assume that similar properties are applicable to the assembly process at physiological conditions as well, examine the consequent dynamics, and make testable predictions.

**Methods**

1. The model

We assume the $\alpha\beta$-tubulin heterodimer to be the microtubule building block, and neglect direct association/disassociation of larger filaments, whose contributions are expected to be very small [2]. This assumption is adopted in most existing models. In this work we focus on the GMPCPP tubulins, therefore will not include GTP hydrolysis in the model. We consider three types of reactions (Fig. 2a &b):

1) A dimer can longitudinally add or dissociate from the ends of a PF (Fig. 2a, process 1). The reaction rates for plus and minus ends are different by a constant ratio $d$ [12,13]. This ensures that the equilibrium constants are the same for the reactions at both ends, as required by thermodynamics. For convenience in this work we call the noncovalent (longitudinal or lateral) interaction between two tubulins a “bond”.

2) A dimer can laterally associate with or dissociate from a PF from either side (Fig. 2a, process 2). The ribbon structure of Wang and Nogales (Fig. 2b) reveals that two neighboring PFs can form...
two types of lateral bonds [6,7,9]. We call one the tube bond as it closely resembles that present in closed, cylindrical microtubules. The other one we called the sheet bond, corresponding to that newly observed by Wang and Nogales between PF pairs.

As suggested by our cryo-EM analysis [9], the main sequence regions involved in lateral interactions between PFs in microtubules are the M-loop (Residues 274–286: PVISAEKAYHEQL in α-tubulin; PLTSRGSOQYRAL in β-tubulin) and the N-loop (Residues 52–61: FFSETGAGKH in β-tubulin; YYNEAAGNKY in β-tubulin) [14,15], whereas the lateral sheet bond interactions between two PFs involve site 1 (Residues 336–342 (H10-S9 loop); KTKRTIQ in α-tubulin; QKNNSSY in β-tubulin) and site 2 (Residues 158–164 (H1-S5 loop); SVDYGGK in α-tubulin; RIEVPDR in β-tubulin) (Fig. S1a). We identified these stretches of residues based on our low-resolution (18 Å) cryo-EM reconstructions, and thus as a coarse approximation to the actual physical interface. Interestingly, the residues involved in the sheet bond are more conserved than those in the tube bond (see Fig. S1b) [16]. It is important to mention that two types of lateral bonds are present in nature in the stable structure of the microtubule doublet, where some PFs need to interact laterally with two neighboring ones simultaneously [17]. The recent doublet structure by Sui and Downing shows a non-MT lateral interaction between PFs B10 and A5 (in their notation) [18]. The doublet and the ribbon structures show that the non-MT interactions in both structures are obtained by rotating one PF relative to another laterally (Fig. S1c). The doublet structure shows even larger rotation angle than the sheet bond, possibly further distorted by other binding proteins in this structure [18]. We also noticed that the various structures obtained by Burton and Himes at slightly basic pHs are easily explained by the existence of alternative types of lateral bonds, but molecular details of their structures are lacking [19]. Physically, the existence of two types of lateral bonds means that the potential of mean force between two neighboring tubulins dimers along the lateral rotational angle assumes a double-well shape. This situation is similar to the lateral interactions along the longitudinal direction, where calculations of electrostatic interactions by Sept et al. show a double-well shaped potential, corresponding to the A- and B-type microtubules [20].

One additional, reasonable assumption is that the formation of the sheet bond is dynamically faster than that of the tube bond. When two protein molecules (or complexes) encounter each other to form a larger complex, it is unlikely that all the mutual interactions between the two surfaces form all at once. Most likely the two protein surfaces form some partial contacts, then gradually adjust to a favorable matching conformation for their mutual interaction, and during the process some residues may need to reorganize slightly. The cryo-EM reconstruction of the low-temperature stabilized ribbons revealed a larger contact surface for the tube bond than for the sheet bond (see Fig. 3). While a larger contact surface may lead to stronger interaction, it may be slower to form. Consequently, a tube bond might be slower to form than a sheet bond does. However, all these discussions are only suggestive, and further experimental studies are needed. As discussed later, a faster sheet bond formation rate is not a necessary assumption in our model, but it increases the percentage of transient ribbon structures, and facilitates formation of the sheet structures.

3) We further propose that the two types of lateral bonds can interconvert (Fig. 2b, process 3). Furthermore, two neighboring lateral bonds can mutually affect each other’s stability and the inter-conversion rates. This assumption is necessary to reproduce the observed low temperature sheet structure. Physically, it is likely that two consecutive lateral bonds affect each other via allosteric changes in the intervening tubulin molecule. Allosteric effects on the tubulin monomers/dimers have already been proposed to play an important role during the microtubule assembly process, although details are unclear [3,9,21]. For simplicity, in our modeling studies we assume the mutual interaction energy between two sheet bonds ΔG_{sheet-sheet} > 0, and other types of interactions ΔG_{tube-sheet} ~ ΔG_{tube-tube} ~ 0, with Sh and Tu referring to the sheet and tube bond, respectively. We will discuss alternative choices later. Below we will show that with these choices one can reproduce the observed low temperature ribbon structure. For a lateral bond conversion reaction, a tubulin dimer needs to rotate about 60 degrees around the longitudinal axis of the neighboring PF [9]. In our simulations of the assembly kinetics and thermodynamic analysis, we do not consider the case in which tubulins within one PF form different types of lateral bonds with their lateral neighbors. Such defects (that tubulins within one PF form different types of lateral bonds with their neighbors) would disrupt the longitudinal and lateral interaction network within the structure, thus be energetically unfavorable, and exist only transiently. This resembles a large class of Ising-type models. For example, protein folding kinetics can often be described by two states without referring to the intermediate transition step. Consequently, our simulation assumes that the tubulin molecules within a PF would rotate collectively and cooperatively. As a consequence, the longer the PF, the harder the rotation is. Also, when a tubulin dimer adds to a PF longitudinally in our kinetic model, it engages in the same lateral bond as the rest of the precedent subunits in the same PF. This approximation greatly simplifies the simulation.

Wang et al. observed the temperature-driven ribbon-microtubule conversion using the GTP analogue GMPcPP [9,11]. Therefore GTP hydrolysis is not a requirement for ribbon/sheet conversion into a microtubule, and thus we did not consider the GTP hydrolysis reaction in this study. We enforce the detailed balance condition by relating the rate constants to the corresponding standard free energy change (ΔG°). For example, the on rate constant k_{(+)} and off rate constant k_{(-)} for a tubulin addition

![Figure 3. Course inspection of the electron density map of the ribbon structure. It reveals a clearly larger buried interface for the tube bond than for the sheet bond.](doi:10.1371/journal.pone.0007291.g003)
reaction, is given by [22]

\[
\frac{k_{(+)}}{k_{(-)}} = \exp\left(-\frac{\Delta G^0}{k_B T}\right),
\]

where \(k_B\) is Boltzmann’s constant, \(T\) is the absolute temperature. Following Erickson and others [2,23,24], we divide the standard free energy \(\Delta G^0\) into two terms, an entropic energy \(\Delta G_{\text{Entropy}}\) accounting for the subunit translational and rotational entropic loss due to bond formation—not the overall entropy contribution, and the remaining free energy change \(\Delta G\). The separation allows proper inclusion of \(\Delta G_{\text{Entropy}}\) while multiple bonds form simultaneously. For instance, the longitudinal binding/dissociation reaction from the plus (upper) end in Fig. 2a gives

\[
\Delta G = \Delta G_{\text{long}}^0 + \Delta G_{\text{Sh}}^0 + \Delta G_{\text{Tu}}^0 - 2\Delta G_{\text{Entropy}},
\]

where \(\Delta G_{\text{long}}^0\) is standard free energy for longitudinal association, \(\Delta G_{\text{Sh}}^0\) the standard free energy change of forming a sheet bond, \(\Delta G_{\text{Tu}}^0\) the standard free energy change of forming a tube bond, and the term \(-2\Delta G_{\text{Entropy}}\) compensates for overcounting of the entropic free energy loss. Detailed description of the rate constant and entropic term calculations can be found in the Supporting Text S1A and B.

2. Simulation details

The assembly process was stochastically simulated with the Gillespie algorithm [25]. At each step, we recorded all the species in the system and their numbers. A reaction was randomly selected from a list of all the possible reactions of all the species in the system. We only simulated the early stage of the microtubule assembly process starting from tubulin dimers. All the simulation parameters were provided in Table 1 and figure captions. There are four energy terms in the model. In our simulations, the binding energy for the longitudinal bond \(\Delta G_{\text{long}}\), and that of the tube lateral bond \(\Delta G_{\text{Tu}}\) were assigned values \(-19 k_B T\) and \(-15.5 k_B T\), respectively, close to what used in the literature after taking into account the entropy term \(\Delta G_{\text{entropy}}\) [2,20,26] (see Supporting Text S1B). Currently there is no direct experimental information to determine the values of the other two terms, the sheet-type lateral bond energy \(\Delta G_{\text{Sh}}\) and the allosteric energy term \(\Delta G_{\text{Allo}}\). In this work we will examine how the assembly dynamics is affected by changing the values of these terms. Future experimental results may suggest possible parameter value ranges by comparing with our simulations. All the results reported here were averaged over 60 independent simulations.

In most calculations we used constant free tubulin dimer concentrations. That is, we started the simulations with tubulin dimers only and kept free tubulin dimer concentration at a fixed value throughout the simulations. Experimentally the total tubulin concentration is fixed. However, here we only examine the very early assembly stage where the percentage of tubulins forming assembly clusters is negligible, so the free tubulin concentration is approximately the same as the total tubulin concentration. Using a constant free tubulin concentration provided us the advantage to increase the simulation efficiency with a limited computational resource. It also allowed us to examine the effect of free tubulin concentrations on the assembly process more easily. Exceptions are Fig. 4f, where the total number of tubulin dimers was kept constant, and the results were averaged over 2000 independent simulations. In this case we kept the system in a small size so we could run simulations for a prolonged time until the system reached equilibrium.

At each sampling step, we took a snapshot of the tubulin assembly clusters. Different clusters have different shape, length and width. To characterize the structural properties of each cluster, we examined the following joint probabilities (or percentages): 1) \(P(\text{Tu-Tu})\)—both of the two neighboring lateral bonds lying between three neighboring PFs being tube type; 2) \(P(\text{Tu-Sh})\)—one tube type, and one sheet type; 3) \(P(\text{Sh-Sh})\)—both being sheet type, with \(P(\text{Tu-Tu}) + P(\text{Tu-Sh}) + P(\text{Sh-Sh}) = 1\). We call the local structure formed by three tubulin dimers in lateral contact as a Tu-Tu, Tu-Sh, or Sh-Sh 3-mer structure. The

### Table 1. Parameters used in the simulation.

| Parameters                                | Values                  | References                     |
|-------------------------------------------|-------------------------|--------------------------------|
| Longitudinal bond strength extracting part of the entropy term \(\Delta G_{\text{long}}\) | \(-19 k_B T\)         | [4,26]                        |
| Sheet bond strength extracting part of the entropy term \(\Delta G_{\text{She}}\)  | Scheme 1: \(-13.5 \sim -17.5 k_B T\), Scheme 2: \(-13 k_B T\) | varying parameter             |
| Tube bond strength \(\Delta G_{\text{Tu}}\)    | Scheme 1: \(-15.5 k_B T\), Scheme 2: \(-16.5 k_B T\) | [4,26]                        |
| Energy barrier \(\Delta G_{\text{ENT}}\)    | \(-9.5 k_B T\)         | estimated                     |
| Entropy loss for two dimer assembly \(\Delta G_{\text{Entropy}}\) \(\sim 2\) | \(5.5 k_B T\)          | [2,4]                         |
| Mutual interaction energy for sheet-sheet bonds \(\Delta G_{\text{She}}\) | Scheme 1: \(0 \sim 6 k_B T\), Scheme 2: \(0 k_B T\) | varying parameter             |
| Mutual interaction energy for tube-tube bonds \(\Delta G_{\text{TuTu}}\) | Scheme 1: \(0 k_B T\), Scheme 2: \(0 \sim 6 k_B T\) | varying parameter             |
| Rate constant for longitudinal assembly at plus end \(k_{\text{long}}\) | \(2 \times 10^6 \mu \text{M}^{-1} \text{s}^{-1}\) | [4,26,47,48,49,50]            |
| Rate constant for longitudinal assembly at minus end \(k_{\text{long}}\) | \(k_{\text{Long}} \times \delta\) | [12,13]                       |
| Assemble ratio between minus and plus ends \(\delta\) | \(1/3\)                | [12,13]                       |
| Rate constant for lateral assembly with tube bond \(k_{\text{Tu}}\) | \(5 \times 10^5 \mu \text{M}^{-1} \text{s}^{-1}\) | [4]                           |
| Rate constant for lateral assembly with sheet bond \(k_{\text{Sh}}\) | \(1 \times 10^5 \mu \text{M}^{-1} \text{s}^{-1}\) | estimated                     |
| Rate constant for conversion between sheet and tube bonds \(k_{\text{ST}}\) | \(5 \times 10^6 \mu \text{M}^{-1} \text{s}^{-1}\) | estimated                     |
| Tubulin concentration \(c\) | \(25 \mu \text{M}\) unless specified otherwise | [9]                           |

*Derived quantities. See Supporting Text S1 B for explanation.
*The entropy term for processes other than \(1\sim 2\) is discussed in Supporting Text S1 B.
The percentage of Tu-Sh structures in the system is calculated as the ratio between the total number of Tu-Sh structures and the total number of 3-dimer structures in all clusters with three or more PFs. A cluster is defined as a ribbon cluster only if $P(\text{Tu-Sh}) = 1$ (Fig. 2c). Therefore a higher value of $P(\text{Tu-Sh})$ means that the cluster is closer to a ribbon structure. A ribbon cluster must have 3 or more PFs by definition. The percentage of ribbon structures in the system is calculated as the ratio between the total number of Tu-Sh structures in the ribbon clusters and the total number of 3-dimer structures in all clusters with three or more PFs. To calculate the population of clusters with certain number ($N$) of PFs, we simply count the total numbers of those $N$-PF clusters at certain steps. The average PF length of an $N$-PF cluster is calculated as the total number of dimers in the cluster divided by $N$.

Currently there is no quantitative experimental data available on the assembly rates at the initial stage we studied here. Therefore all the results are reported with a relative time unit, which can be easily scaled to the experimental rates once available.

**Results**

1. **Effect on tubulin assembly of a difference in binding energy between sheet- and tube- lateral bonds**

Fig. 4 gives the dependence of the assembly process on the value of the $\Delta G_{\text{Sh}} - \Delta G_{\text{Tu}}$ (binding energy difference between the sheet- and tube-type lateral bonds), with fixed values of $\Delta G_{\text{Tu}} = -15.5 k_B T$ and $\Delta G_{\text{Sh}} = 6 k_B T$ for all. The percentage of Sh-Sh structures is negligible for all simulations (data not shown). The percentage of ribbon structures is calculated as the ratio between the total number of Tu-Sh structures and the total number of 3-dimer structures in all clusters with three or more PFs. A cluster is defined as a ribbon cluster only if $P(\text{Tu-Sh}) = 1$ (Fig. 2c). Therefore a higher value of $P(\text{Tu-Sh})$ means that the cluster is closer to a ribbon structure. A ribbon cluster must have 3 or more PFs by definition. The percentage of ribbon structures in the system is calculated as the ratio between the total number of Tu-Sh structures in the ribbon clusters and the total number of 3-dimer structures in all clusters with three or more PFs. To calculate the population of clusters with certain number ($N$) of PFs, we simply count the total numbers of those $N$-PF clusters at certain steps. The average PF length of an $N$-PF cluster is calculated as the total number of dimers in the cluster divided by $N$.
structures and that of Tu-Sh structure decreases on increasing $\Delta G_{\text{SSh}}$ (see Fig. 4a & b). For $\Delta G_{\text{Sh}}=\Delta G_{\text{Tu}}>0$ (the sheet bond stronger than the tube bond, simulating the low-temperature condition) the percentage of ribbon structures stays at a high plateau (top curves in Fig. 4a). For $\Delta G_{\text{Sh}}=\Delta G_{\text{Tu}}>0$ (the tube bond is stronger than the sheet bond, simulating the high-temperature condition) the percentage of ribbon structures starts with a relative high value, then decreases with time. This observation indicates that initially formed sheet bonds transform into tube bonds at a later time. Fig. 4c supports this idea by showing that (for $\Delta G_{\text{Sh}}=\Delta G_{\text{Tu}}=1$ k_BT) the percentage of Tu-Sh structures in 3-PF clusters is higher than that of later formed larger clusters. Fig. 4d gives (also for $\Delta G_{\text{Sh}}=\Delta G_{\text{Tu}}=1$ k_BT) the average PF lengths (as number of dimers) for different cluster sizes. Small clusters with one or two PFs quickly reach steady-state with average longitudinal length of about 4 tubulin dimers. Experimentally, a large amount of small single- and double-PF clusters with length 4–5 tubulin dimers are observed at the initial stage of the assembly process [11]. The longitudinal length of larger clusters increases continuously within the simulation time. From a thermodynamic point of view the explanation for this result is that the lateral bonds within larger clusters stabilize the clusters, but the single and double-PF clusters lack sufficient lateral bonds and cannot grow long [2]. We performed a simulation with the lateral bond addition turned off so only one PF structures can be formed. The observed average single PF structure length quickly reaches a plateau at a slightly larger value (about 10 dimers, data not shown). From a kinetic point of view, the smaller clusters may disappear also by growing in width and thus transforming into larger clusters before growing long. Similarly shown in Fig. 4e, the populations of single- and double-PF clusters reach a plateau, while the numbers of larger clusters increase continuously within the time of simulation.

In Fig. 4b we examined how the percentage of Tu-Sh structures evolves with time. The results show that all the curves reach a plateau. It is unclear whether the system reaches equilibrium or a dynamic steady-state. The latter would mean that newly formed sheet bonds compensate the loss of the Tu-Sh structure population due to Sh→Tu conversion, so the percentage of Tu-Sh structures remains unchanged. If this is the case, the apparent percentage of Sh→Tu conversion should be less than the real value. Therefore, we performed additional simulations with constant total number of tubulin dimers. This time, we used a smaller size system (100 dimers), which allowed us to perform sufficiently long simulations for the system to reach true thermodynamic equilibrium. Fig. 4f shows the evolution of the percentage of Tu-Sh structures with different values of $\Delta G_{\text{Sh}}=\Delta G_{\text{Tu}}$. In the case of $\Delta G_{\text{Sh}}=\Delta G_{\text{Tu}}>0$, thus when the tube bond is thermodynamically more stable, the Tu-Sh structures start at relatively high percentage, then convert after the first few thousand steps. This result is due to the faster formation of sheet bonds versus tube bonds, with the former being transiently trapped as the PFs grow longer. The sheet bonds eventually convert to the thermodynamically more stable tube bonds and the system reaches equilibrium. Compared to Fig. 4b, we did observe larger percentage of Sh→Tu transition in Fig. 4f, indicating that the curve plateaus in Fig. 4b are due to a dynamic steady-state. In the case of $\Delta G_{\text{Sh}}=\Delta G_{\text{Tu}}<0$, where a sheet bond is more stable than a tube bond, in addition to the effect of the positive $\Delta G_{\text{SSh}}$, the Tu-Sh structures are more stable thermodynamically (the top lines of Fig. 4f).

2. Effect on tubulin assembly of mutual allosteric interaction between two adjacent sheet bonds

If formation of a new lateral bond is not affected by the existing PFs ($\Delta G_{\text{SSh}}=0$), one would expect randomly distributed lateral bond types between PFs. The allosteric term $\Delta G_{\text{SSh}}$ is necessary for reproducing the dominating ribbon structures experimentally observed at low temperature ($\Delta G_{\text{S}}=\Delta G_{\text{Tu}}>0$). Fig. 5 shows that, for $\Delta G_{\text{S}}=\Delta G_{\text{Tu}}=-1.5$k_BT, the percentage of ribbon structures and that of T-S structures is sensitive to the value of $\Delta G_{\text{SSh}}$. As $\Delta G_{\text{SSh}}$ increases from 0 to 6 k_BT, the percentage of ribbon structure increases from 20% to around 90% (Fig. 5a). The percentage drops slightly as time evolves. This is because some newly formed small ribbon structures grow to hybrid forms upon adding more PFs. Fig. 5b-d show the 3-mer structure distribution. For $\Delta G_{\text{SSh}}=0$, Fig. 5b shows that the S-S structure is dominating, reflecting the fact that the sheet bond is stronger than the tube bond. While there are still about 20% Tu-Sh structures, the Tu-Tu structures are negligible. On increasing $\Delta G_{\text{SSh}}$ to 2 k_BT (Fig. 5c), the free energy difference between a sheet and a tube bond (−1.5 k_BT) cannot compensate the unfavorable term $\Delta G_{\text{SSh}}$, and more Tu-Sh structures than the Sh-Sh structures are formed. As we further increase $\Delta G_{\text{SSh}}$ to 6 k_BT (Fig. 5d), T-S structures become dominating, while the other two structures are negligible. In the case where $\Delta G_{\text{S}}=\Delta G_{\text{Tu}}>0$, a positive value of $\Delta G_{\text{SSh}}$ maintains its effect on producing higher percentage of newly formed Tu-Sh arrangement, with the ribbon structures dominating the population, but these gradually transform into the more stable microtubule structures (see Fig. S2).

3. The effect of free tubulin concentration on the assembly process

The free tubulin concentration is another factor affecting the assembly kinetics. Fig. 6a and b examine the effect of free tubulin concentration on the assembly process in the case where $\Delta G_{\text{S}}=\Delta G_{\text{Tu}}>0$ (high temperature scenario in which tubulin polymerizes into microtubules). On increasing the free tubulin dimer concentration from 5, to 25, to 125 μM, both the ribbon and T-S structures increase. At higher dimer concentration the population of the ribbon structure forms starts at a high percentage, then drops quickly to the similar level as that at lower dimer concentrations. A possible explanation for this phenomenon is that some of the ribbon structures transform into larger hybrid structures upon PF addition. This is supported by the persistence of the high percentage of Tu-Sh structures at high tubulin concentration (Fig. 6b). The steady-state average length of the single-PF clusters increases as the tubulin concentration goes up (Fig. 6c, curves marked with grey circles), reflecting the fact that increasing the tubulin concentration favors bond formation both thermodynamically and kinetically. The lateral bond formation is apparently favored by high dimer concentrations due to a higher assembly rate, so the multi-PF clusters grow even faster at higher dimer concentration (Fig. 6c, curves marked with open circles). The population of larger clusters (5-PF in the case shown) also increases faster at higher dimer concentrations (Fig. 6d). Overall, our simulations suggest that the sheet intermediates are more likely to be observed at high free tubulin concentrations. This agrees well with the experimental result that larger and more abundant sheet structures are observed during the initial, exponential phase of tubulin of polymerization when free tubulin concentrations are high (>100 μM) [7,27]. Physically, increasing the free dimer concentration increases the cluster growth rates, which effectively allows less time for the internal Sh→Tu transition, and thus increases the percentage of ribbon and Tu-Sh structure, as shown in Fig. 6a & b.

Discussion

Erickson and Pantaloni performed thermodynamic analysis on the initial stages of polymer assembly [24], with the assumption...
\[ \Delta G_{\text{S-S}} - \Delta G_{\text{Tu}} = -1.5 \kappa B T, \quad \Delta G_{\text{S-S}} = 0, 2, 4, 6 \kappa B T \ (b: 0, c: 2, d: 6) \]

**Figure 5. Effect of variable \( \Delta G_{\text{S-S}} \) on the assembly structures for fixed \( \Delta G_{\text{S}} = -17 \kappa B T \) and \( \Delta G_{\text{Tu}} = -15.8 \kappa B T \)**

(a) Percentage of ribbon structures as a function of time (\( \Delta G_{\text{S-S}} = 0, 2, 4 \) as indicated by circled numbers). (b) Trimer-structure distribution v.s. simulation step for \( \Delta G_{\text{S-S}} = 0 \). The three possible trimer structures, T-T (tube-tube), T-S (tube-sheet) and S-S (sheet-sheet), are indicated in the figure. (c) Trimer structure distribution v.s. simulation step with \( \Delta G_{\text{S-S}} = 2 \kappa B T \). (d) Trimer-structure distribution v.s. simulation step with \( \Delta G_{\text{S-S}} = 6 \kappa B T \).  

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that only one type of lateral bond exists. In their model, the sheet is not structurally different from the microtubule structure. In the present study, and while incorporating recent structural information, we are trying to simulate the very early stages of tubulin polymerization at both low and high (physiological) temperature, making a minimal number of assumptions that will reproduce existing experimental observations. The main conclusions from this exercise follow.

**Thermodynamic analysis**

Let’s consider a structure with \( 2n \) PFs of length \( n \) dimers. At low temperatures (less than 15°C), the sheet bond is more stable than the tube bond (\( \Delta G_{\text{S}} - \Delta G_{\text{T}} < 0 \)). Therefore, the thermodynamically most stable structure tends to form as many sheet bonds as possible. However, the term \( \Delta G_{\text{S-S}} \) disfavors a sheet structure with all sheet bonds. One can show that, instead, the most stable structure is the one with alternating lateral bonds, provided \( \Delta G_{\text{S-S}} > |\Delta G_{\text{S}} - \Delta G_{\text{T}}| \). The free energy difference between the structure with neighboring sheet bonds and the one with alternating lateral bonds is \( n(2m-1)(\Delta G_{\text{S-S}} - \Delta G_{\text{T}} + \Delta G_{\text{S}}) \). The difference between a sheet bond-only structure and an unclosed tube bond-only structure is \( n(2m-1)(\Delta G_{\text{S}} - \Delta G_{\text{T}}) \). When \( n \) and/or \( m \) are large, a small difference in the bond energy leads to a large difference in the Boltzmann weight. The structure with alternating lateral bonds is thus the dominating form. Above a certain temperature, the tube bond becomes more stable than the sheet bond (\( \Delta G_{\text{S}} - \Delta G_{\text{T}} > 0 \)), and the microtubule becomes the most stable polymer form. These thermodynamic considerations explain the results in Fig. 4 and Fig. 5. There are several possible origins on the temperature dependence of \( \Delta G_{\text{S}} - \Delta G_{\text{T}} \). We discussed them in Supporting Text S1C.

For the allosteric effect represented by the term \( \Delta G_{\text{S-S}} \), we suggest two possible mechanisms. First lateral interactions have been proposed to straighten a tubulin dimer (this is referred to as the lattice effect) [9,28,29]. Consequently, the lateral interaction surface is in general coupled to straightening, and the allosteric effect proposed here and the lattice effect are closely related and coupled. This effect may exist even if each tubulin monomer is treated as a rigid body. While this is the mechanism we favor, a second alternative scenario is that, as tubulin molecules are flexible, lateral interactions on one side could affect the lateral surface on the other side of the protein.

A sheet structure is a common morphology for biological molecule self-assembly [30,31,32]. Tubulin assembly shares some common features. For example, the ribbon structures are helical, and the tubulins are arranged in a microtubule in a helical manner [9]. Therefore, due to asymmetric off-axis interactions between tubulins these structures are chiral [32]. The general theory discussed by Aggeli et al. may be applied to a more detailed analysis of the tubulin assembly model.

**How is the sheet bond kinetically trapped during the assembly process?**

At physiological temperatures, where \( \Delta G_{\text{S}} - \Delta G_{\text{T}} > 0 \), the microtubule is thermodynamically at the most stable polymer form. However, Fig. 5 shows that a large population of structures
with the sheet bonds can still be observed transiently at the initial assembly stage. The steady state population of ribbons will depend on the actual value of $D_{G_{Sh}} - D_{GTu}$, $\Delta G_{Sh} = 1.5 k_B T$, $c = 5, 25, 125 \mu M$

$$\Delta G_{Sh} - \Delta G_{Tu} = 1.5 k_B T$$

Figure 6. Effects of tubulin dimer concentrations on the assembly process (for $\Delta G_{SS} = 6k_BT$, $\Delta G_{SS} = -14k_BT$, $\Delta G_{Tu} = -15.5k_BT$, i.e., $\Delta G_{SS} - \Delta G_{Tu} = 1.5k_BT$) (dimer concentration $c = 125, 25, 5\mu M$, as labeled in the figure with corresponding circled numbers). (a) Percentage of ribbon structures as a function of time. (b) Probability of finding neighboring T-S structures as a function of time. (c) Average PF lengths of structures with 1 PF (dashed lines with grey circled numbers indicating concentrations) and 5 PFs (solid lines with open circled numbers indicating concentrations) v.s. time. (d) Occurrence of clusters with 5 PFs v.s. time for different tubulin concentration as labeled. doi:10.1371/journal.pone.0007291.g006

Our model also predicts the existence of some hybrid structures between the sheet and the MT forms, where the lateral bond pattern is not so regular (e.g., some of the structures in Fig. 2d and 2e). The cryo-EM images of Chretien et al. revealed a distribution of the sheet bending angles [7], which may correspond to different sheet structures with different ratios of sheet versus tube bonds. It is tempting to speculate that at the tip of the growing structure Sh-Tu alternating bonds predominate (see Fig. 4), but as the structure gets closer to the growing microtubule, more and more sheet bonds have converted to tube bonds, until eventually all lateral contacts are tube contacts (an alternative explanation is that at any given point along the length of the sheet, all lateral bonds are the same, but that they change in synchrony along the length, asymptotically reaching that of the tube bond when the structure finally closes into a tube). The process of conversion was not covered in the present
study, where we focused on the very early stage of the assembly process. In this case the formed structures all have small sizes and therefore the conversion process itself is very fast. Instead, it is the initiation of the conversion that is rate-limiting. To mathematically model the conversion process and the sheet curvatures explicitly at the growing tip of a preformed microtubule, one needs to include more details of the mechano-chemical properties of the system. This is an ongoing effort in our labs.

In our model we choose \( \Delta G_{\text{SS}} \sim \Delta G_{\text{TT}} \sim 0 \), and \( \Delta G_{\text{SSSh}} > 0 \). These are roughly based on steric constraints imposed by the competing strains of two distinct curvatures—the longitudinal curvature along the length of a protofilament, and the curvature of the lateral interactions that give rise to a close structure for the microtubule. Our model also assumes that the value of \( \Delta G_{\text{SS}} - \Delta G_{\text{TT}} \) vary with temperature (Fig. S3a). It is important to point out that this scheme (Scheme 1) is not the only one that can reproduce the observed low and high temperature structures (ribbons and microtubules, respectively, at steady state). For example, an alternative scheme (Scheme 2) could be that \( \Delta G_{\text{SS}} - \Delta G_{\text{TT}} > 0 \) (so the tube bond is always stronger than the sheet bond), \( \Delta G_{\text{SSTT}} \sim \Delta G_{\text{SSSS}} \sim 0 \) (which are unnecessary but for simplicity), but \( \Delta G_{\text{TtTT}} > 0 \), which decreases with temperature (Fig. S3b). Also see Supporting Text S1C, which provides some theoretical analysis with a reaction path Hamiltonian [33] on a possible origin for a hypothetical temperature dependence of \( \Delta G_{\text{TtTT}} \). Our stochastic simulations confirm that this scheme can reproduce the low temperature ribbon structures and the high temperature transient sheet structures (see Fig. S4 and Supporting Text S1C for details). Compared to Scheme 1, which is the focus of this work, and where the Sh-Sh structure is negligible (with \( \Delta G_{\text{SSSS}} > 0 \)), Scheme 2 suggests that a larger percentage of Sh-Sh structures should be observable if one chooses \( \Delta G_{\text{SSSS}} \sim 0 \). A specific way to distinguish the two schemes would be to examine the population difference of 2-PF clusters with sheet bond and tube bond. Fig. S5 shows that, in Scheme 1, the sheet-type 2-PF clusters are dominant at low temperature and the tube-type 2-PF clusters become more at high temperature; in Scheme 2, the tube-type 2-PF clusters are always dominant at both high and low temperature. Experimentally determining the 2-PF cluster structures at both low and high temperatures would allow us to estimate the value of \( \Delta G_{\text{SSSS}} \). This will also help on evaluating the two schemes discussed here and the proposal by Chrétien as well. However, no matter which scheme is correct, our main conclusion remains: the existence of the sheet tubulin structures is due to thermodynamics at low temperatures, but kinetics at higher (physiological) temperatures.

Fygenson et al. carried out variability-based alignment of \( \alpha \) - and \( \beta \)- tubulin sequences [16]. More conserved residues usually have functional importance. In Fig. S1 we reproduced their result, and indicated the above-mentioned residues involved in lateral interactions. It is clear that those residues (especially several charged ones) involved in the sheet bond formation are generally more conserved than those for the tube bond. In addition, there are a smaller number of residues involved in the interface of the former, which can be visualized in a simple fashion by examination of the ribbon electron density map showing a smaller contact surface for the sheet bond than for the tube bond (see Fig. 3). These observations may explain why the sheet bond would be faster to form than the tube bond. The former involves less residues but strong electrostatic interactions, which can guide the approaching tubulins to interact. On the other hand, to form a tube bond more residues need to align (and reorganize) properly with each other, which may result in a high barrier for the reaction. Is it possible that tubulin evolved a sheet→tube, two-step processes to increase the tubulin lateral assembly rate: a free tubulin dimer would first be captured by the fast-forming sheet bond, and this would serve as a primer to guide the complex to form the more stable but slower-to-form tube bond. In a direct tube-bond formation mechanism, the interaction between the loosely formed contact pairs may be too weak to hold the newly added tubulin dimer for sufficiently long time before necessary conformational reorganization takes place to form the stable tube bond, which would result in very low lateral association rate.
How biologically relevant is the proposed sheet bond?

The ribbon structure obtained by Wang and Nogales shows two types of lateral bonds. In our model, we assume that the same types of lateral bonds exist during the assembly process of both GMPCPP and GTP tubulins at physiological conditions. One may argue that the observed ribbon structure is not physiological, as it is obtained at low temperature and high magnesium ion concentrations. High magnesium ions are typically used for the stabilization of all forms of tubulin assembly, and are hypothesized to shield the acidic C-terminal tails of tubulin (E-hooks), perhaps in a manner similar to that proposed for classical MAPs. These MAPs are highly basic, poorly structured, and generally have also a stabilizing effect on different tubulin assembly forms (e.g. they stabilized both microtubules, and tubulin rings). Cold temperature, on the other hand, is known to have a destabilizing effect on microtubules, and tubulin rings). GMPCPP and GTP tubulins at physiological conditions. One may types of lateral bonds exist during the assembly process of both microtubules–to microtubules in vitro renders the polymers cold-stable, just like the anticancer drug taxol does (K. Patel, R. Heald, and E. Nogales, unpublished results)). The formation of the ribbon structure, in the presence of GMPCPP, at low temperatures, was therefore a surprise. A working hypothesis to explain the assembly of the ribbons, in conditions where GTP tubulin would not be able to assemble into microtubules, is that temperature slows down tubulin interactions, with less of an effect on the rate of hydrolysis once a tubulin-tubulin contact has formed. Thus, under low temperature conditions little assembly occurs, and when it does hydrolysis quickly follows, before tubulin has a chance to make a microtubule closure and store the energy as lattice strain. When the hydrolysis step is eliminated, the slow polymerization of GTP tubulin (GMPCPP) can continue without the conformational change, and the destabilization effect that hydrolysis brings on tubulin. Under this simple assumption, we propose that the ribbon assembly conditions shed information on the process of microtubule assembly taking place before microtubule closure. This idea is supported by the structure of the ribbon itself, which shows alternating lateral contacts between protofilaments, that otherwise preserve the precise stagger between protofilaments seen in the microtubule. This suggests that the ribbons would be able to convert directly into microtubules, as it was experimentally confirmed [9]. Concerning the rotation of the lateral sheet bond, it is important to mention that this type of arrangement, or at least one involving alternative lateral contacts without longitudinal displacements between protofilaments, could have been deduced directly from the extended sheets observed by Chretien and colleagues growing at the end of microtubules, unless extreme deformability is otherwise hypothesize for the tubulin subunit, which is beyond reason.

An alternative model for the experimentally observed sheet structures at the end of growing microtubules is that they involved tubulin interactions are not different from those observed in a MT. A sheet structure is simply an incomplete protruding MT structure. However, the stochastic modeling results of VanBuren show that with this model it is very unlikely to form long incomplete structures at a MT growing end. The structures are energetically unfavorable, and are precursors for disassembly rather than assembly [9]. They didn’t examine dependence of the sheet length on the tubulin concentrations. One would expect weak or inverse dependence, since low tubulin concentrations would favor disassembly. This is contrary to the observation that the sheet structures under observed under growth conditions, and become longer (up to several hundred nanometers) upon increasing tubulin concentrations [7].

In conclusion, although there is yet no direct evidence of the presence of the sheet-type lateral bond described here under physiological conditions (the transient character preventing structural characterization, but see discussions on the doublet below), there is very compelling evidence that alternative lateral interactions do exist in a transient intermediate, the sheets at the end of fast growing microtubules. All our analyses indicate that the ribbon structure is the best candidate in existence to describe such intermediates. A somehow similar, and stable structure has been observed in the doublet form, which demonstrates that the alternative lateral bonds do exist in vivo. As discussed below, the unusual high conservation of the residues proposed to participate the sheet bond formation strongly suggest the functional importance of these residues. We put forward the proposal that existence of (at least) two types of lateral bond naturally explains the sheet and microtubule forms observed in vitro, and the interconversion between them.

The situation in vivo is more complex, where various microtubule-associated-proteins (MAPs) may modify the microtubule assembly/disassembly process. While more in vivo studies are necessary to address the functional relevance of the sheet structure observed in vitro, it will also be very informative to study the microtubule assembly process in the presence of purified microtubule-binding proteins. It is important to notice that all structural studies of microtubules with binding partners have been carried out by adding the partners to preassembled (usually taxol-stabilized) microtubules. The effect on the assembly process of +TIPS, for example, should come a lot closer to reproduce what goes on inside the cells, than the analyses carried out to date with purified tubulin alone.

We also suggest that the existence of alternative lateral bond types may have functional importance. Nogales and Wang proposed that the ribbon structure (and the sheet structure in general) could provide a novel surface for microtubule-binding proteins that could recognize surfaces unique to the sheet bond to track microtubule growing ends [5]. It has also been proposed that the sheet structure could constitute a structural cap at the end of growing microtubules [7] of essential importance in dynamic instability (notice that both functions would most likely be linked). Additionally, if the MT lateral bond is indeed stronger than the sheet lateral bonds, free energy would be stored in the lateral bonds of the sheet structure, released upon closure, which could result in mechanical force generation. We provided a more detailed discussion of this idea in the Supporting Text S1.

The nature of lateral interactions also affects the microtubule mechanical properties. Even if only one type of tubulin lateral interactions exists under normal conditions, microtubules in a cell are constantly under mechanical stress due to protein motors and other microtubule associated proteins [34]. There is a certain probability that some of the lateral bonds within a microtubule may convert to another type of interactions under extreme conditions (e.g. buckling under large mechanical force), as implied by recent atomic force microscope studies [35,36]. The new type of lateral bond provides a way of releasing local mechanical stress without breaking the MT. We expect that the mechanical property of a MT with and without this new type of lateral interaction would be dramatically different, and can be tested experimentally. It remains to be examined if these conditions are biologically relevant.

What could be the function of the sheet intermediate?

In addition to the artificially generated ribbon structure of Wang and Nogales, cryo-EM studies have more directly shown the presence of sheet intermediates during microtubule growth, both
in vitro [6,7] and in vivo [8]. Theoretically, the sheet structures and the conversion into microtubules could play several important functional roles. Nogales and Wang proposed that the ribbon structure (and the sheet structure in general) could provide a novel surface for microtubule-binding proteins that could recognize surfaces unique to the sheet bond to track microtubule growing ends [5]. It has also been proposed that the sheet structure could constitute a structural cap at the end of growing microtubules [7] of essential importance in dynamic instability (notice that both functions would most likely be linked).

The sheet bond involves fewer residues but strong electrostatic interactions, which can guide the approaching tubulins to interact. On the other hand, to form a tube bond more residues need to align (and reorganize) properly with each other, which may result in a high barrier for the reaction. Is it possible that tubulin evolved a sheet–tube, two-step processes to increase the tubulin lateral assembly rate: a free tubulin dimer would first be captured by the fast-forming sheet bond, and this would give the formed cluster longer time to adjust to the more stable but slower-to-form tube bond. In a direct tube-bond formation mechanism, the interaction between the loosely formed contact pairs may be too weak to hold the newly added tubulin dimer for sufficiently long time before necessary conformational reorganization takes place to form the stable tube bond, which would result in very low lateral association rate.

We would like to propose here that there could be also a mechanical function for a preformed sheet that eventually closes into microtubule structure. Terrell Hill first proposed that assembly and disassembly of cytoskeletal filaments could generate mechanical force [37]. Subsequent theoretical studies and experimental measurements confirmed this idea [38,39,40,41]. Oster and coworkers proposed a ratchet mechanism and its variations to explain how elongating polymers like microtubules can generate force and push an object forward (see Fig. 8a) [42,43]. Thermal motions of the object and the polymer can produce space between them sufficiently large for a building unit (a tubulin dimer in this case) to add to the polymer’s end. Addition of the new unit prevents the object from moving back. Therefore, the random thermal motion of the object is ratcheted into directional motion at the expense of free energy released from unit addition. Most published work uses the ratchet model to explain force measurements during microtubule assembly [39,44]. With the sheet intermediate, the ratchet effect can generate force at the growing tip or at the zipping front, depending on the location of the load. Interestingly, it could also provide another active force generating mechanism in addition to the passive ratchet model. If the MT lateral bond is indeed stronger than the sheet lateral bonds, free energy would be stored in the lateral bonds of the sheet structure. Transformation to the MT structure is a cooperative process. When many lateral bonds transform together, they would release free energy much larger than that stored in a single lateral bond, thus enable them to push against larger loads. (Fig. 8b) In this way the energy accumulation step (tubulin bond formation) and the work-performing step (tube closure) are temporally and spatially separated. A similar mechanism of performing mechanical work using prestored energy has been proposed for the extension of the Limulus polyphemus sperm actin bundle [45]. Which mechanism dominates would depend on where the contact point between the MT and the load is and on the free energy difference between two types of lateral bonds.

**Discussion**

In this study, using the single assumption that there are nearest-neighbor interactions between two consecutive PFs, together with existing structural information, we were able to generate a simple model to explain a large number of observations concerning the mechanism of microtubule assembly. We suggest that the sheet structure observed during microtubule growth may be a kinetically trapped intermediate, and that it is related to the ribbon structure stabilized at low temperature. Our model predicts that the sheet structures are more likely to be observed at high free tubulin concentrations. Structural studies of 2-PF clusters during the assembly process could provide information to discriminate among several possible mechanistic schemes.

Our current analysis has focused only on the initial stage of in vitro microtubule assembly. A future study should provide a more detailed description of the assembly process, especially the interface between the sheet bonds and the tube bonds along the longitudinal direction within the growing end of a microtubule. Our current treatment that all the lateral bonds within a pair of PFs are identical is clearly only an approximation. In this work we focused on the assembly dynamics of GMPcPP tubulins. We didn’t include GTP hydrolysis dynamics and the resulting tubulin dimer conformational changes. We assume that the structural information extracted from the GMPcPP sheet structure can be extrapolated to the normal assembly process. While supported by several other independent experimental evidences, this assumption requires further scrutiny. Especially we propose that at physiological conditions tubulins can form alternative lateral bond type other than that observed in microtubules, as evidenced in the doublet structure. If being confirmed, it would greatly modify our understanding on the mechanical properties of microtubules, and possible mechanisms of interactions between microtubules and microtubule association proteins (MAP) [2,34,46].

Our current model is essentially a two-dimensional model. The current simple model already provides many new insights on the very initial stage of the assembly process with only small cluster structures formed. Both the sheet and the MT forms are actually three-dimensional manifolds. More structural details are needed to fully account for the helical shape of the sheet and the microtubule structure. In the future a three-dimensional mechano-chemistry model parallel to what have been developed for the direct dimer-addition model would be needed [3,4].

**Supporting Information**

**Supporting Text S1** A. Rate constants in the model. B. Calculation of the entropic contribution. C. Physical origins of the temperature dependence of the free energy terms.

Found at: doi:10.1371/journal.pone.0007291.s001 (0.32 MB DOC)

**Figure S1** Structural basis for the two types of lateral bonds. (a) Structure of the αβ-tubulin dimer with residues involved in lateral interactions indicated. Blue: residues engaged in lateral tube bonds.
(274–286, 52–61). Red: residues engaged in lateral sheet bond (336–342, 158–164) these residues have been identified by docking the high-resolution tubulin structure into the 18 A reconstruction of the ribbon [6], and therefore are correct within the constraints of the limited resolution. Pink and yellow: possible surface residues (100–108, 209–225, 300–311) along the tube–sheet conversion pathway. (b) Variability-based sequence alignment of α and β tubulin performed by Fygenson et al. [7]. The blue and red boxes indicate the residues involved in the tube and sheet bond formation given in (a), respectively. The figure is adapted from Fig. 2 of Fygenson et al. [7] with permission. (c) Comparison of the non-MT lateral interactions observed in the microtubule doublet of axonemes (top) [8] (PDF file provided by Sui and Downing) and the ribbon structures (bottom) [6].

**Figure S2** Effect of variable ΔG_SSSh on the assembled structures with ΔG_S = –14.5 kBT and ΔG_T = –15.5 kBT (ΔG_SSSh = 1 kBT) as indicated. The figure shows the percentage of ribbon structures as a function of the time for ΔG_SSSh = 0, 1, 2 and 3 kBT, as indicated. Found at: doi:10.1371/journal.pone.0007291.s002 (1.54 MB TIF)

**Figure S3** Schematic Illustration of the physical origins of the temperature dependence of the free energy terms. (a) ΔG_S and ΔG_T have different temperature dependence and their difference changes sign over T. (b) The dependence of ΔG_GT on the conformational coordinate describing the necessary collective conformational change upon forming two neighboring lateral tube bonds varies with temperature.

**Figure S4** Effects of variable ΔG_GT on the assembly structures using the Scheme 2 described in Fig. S3b. (0, 2, 4, and 6 kBT, as indicated by corresponding circled numbers). Different ΔG_GT correspond to different temperatures as showed in Fig. S3b and supporting text C. ΔG_S = –13 kBT and ΔG_T = –16.5 kBT were used for all simulations. Other parameters are the same as in the Scheme 1 described in detail in the main text. The final results are averaged over 60 independent simulations. (a) Percentage of ribbon structure v.s. simulation step. (b) Percentage of T-S structure. (c) Average PF length for clusters of different size (1 to 6 PFs as indicated by circled numbers), with ΔG_GT = 2 kBT. (d) Cluster population for clusters of different size (1 to 6 PFs as indicated by circled numbers), with ΔG_GT = 2 kBT.

**Figure S5** Population ratio of tube-cluster versus sheet-cluster for 2-PF structures as a function of time. Solid and dashed lines with triangles correspond, respectively, to Scheme 1 (ΔG_SSSh > 0, ΔG_GT = 0), ΔG_S = 1.5 kBT, ΔG_SSSh = 6 kBT and to Scheme 2 (ΔG_GT > 0, ΔG_SSSh = 0). ΔG_GT = 3.5 kBT, ΔG_GT = 2 kBT), both at high temperature . The lines without triangles are for Scheme 1 (solid line, ΔG_SSSh = ΔG_GT = 1.5 kBT, ΔG_S = 6 kBT) and Scheme 2 (dashed line, ΔG_SSSh = ΔG_GT = 3.5 kBT, ΔG_GT = 6 kBT) at low temperature.

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**Author Contributions**

Conceived and designed the experiments: JX. Performed the experiments: ZW HWW EN JX. Analyzed the data: ZW HWW EN JX. Contributed reagents/materials/analysis tools: HWW WM ZO EN. Wrote the paper: ZW HWW EN JX.

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