Pivot-and-bond model explains microtubule bundle formation

Marcel Prelogović¹, Lora Winters², Ana Milas³, Iva M. Tolić*²,³, Nenad Pavin*¹

¹ Department of Physics, Faculty of Science, University of Zagreb, Bijenička cesta 32, 10000 Zagreb, Croatia

² Max Planck Institute of Molecular Cell Biology and Genetics, Pfotenhauerstr. 108, 01307 Dresden, Germany

³ Division of Molecular Biology, Ruđer Bošković Institute, Bijenička cesta 54, 10000 Zagreb, Croatia

(Received: 18 April 2017)

*These authors contributed equally to this work.

*Corresponding authors: N.P. (npavin@phy.hr) and I.M.T. (tolic@irb.hr).
ABSTRACT

During mitosis, bundles of microtubules form a spindle, which is responsible for proper segregation of the genetic material. A key question is what are the physical principles underlying the formation and stability of microtubule bundles. Here we show, that random angular movement of microtubules around the spindle pole and forces exerted by passive cross-linking proteins are sufficient for the formation of stable microtubule bundles. We test these predictions by experiments in wild-type and ase1Δ fission yeast cells. In conclusion, the angular motion drives the alignment of microtubules, which in turn allows the cross-linking proteins to connect the microtubules into a stable bundle.
During mitosis the cell forms a spindle, a complex self-organized molecular machine composed of bundles of microtubules (MTs), which segregates the chromosomes into two daughter cells [1]. MTs are thin stiff filaments that typically extend in random directions from two spindle poles [2]. MTs that extend from the same pole can form parallel bundles, whereas MTs originating from opposite spindle poles form anti-parallel bundles [3-5]. Stability of MT bundles is ensured by cross-linking proteins, which bind along the MT lattice, connecting neighboring MTs. Cross-linking occurs only if the distance between the MTs is comparable with the size of a cross-linking protein. These proteins can be divided into two classes: (i) proteins that cross-link MTs without directed movement along the MT, such as Ase1/PRC1 (ref. [6]); (ii) motor proteins that walk along the MT either towards the plus end of the MT, such as Cut7/Eg5 (ref. [7,8]), or towards the minus end, such as Ncd (ref. [9,10]).

Spindle self-organization was studied in different biological systems and several theoretical models were proposed. Formation of antiparallel bundles of MTs in somatic cells of higher eukaryotes was investigated by computer simulations, which include MTs that grow in random directions from two spindle poles and motor proteins that link them [11]. Further, several studies have explored the forces generated in the antiparallel overlaps in vitro [12-14] and in Drosophila embryos [15-17]. Spindle formation was studied in Xenopus eggs, using the “slide and cluster” [18,19] and liquid crystal models [20,21]. In budding yeast, it is suggested that MTs growing in arbitrary directions from the opposite spindle poles can change their direction due to minus end directed kinesin-14 motors bound to both MTs and get aligned, forming anti-parallel bundles [22]. During spindle positioning, myosin motors walking along actin cables accelerate pivoting of astral MTs when they search for cortical anchor sites [23]. Studies in fission yeast have shown that passive (thermal) pivoting motion of MTs around the spindle pole body accelerates kinetochore capture [24-26], together with dynamic instability of MTs [27]. MT rotational motion about a pivot at the SPB was also included in the model
for spindle formation [28] and in vitro studies [29]. However, observation of the dynamics of bundle formation in vivo and a corresponding physical model are required to understand the formation and stability of MT bundles.

In this paper, we combine experiments and theory to explore the formation of parallel MT bundles. We introduce the pivot-and-bond model for the formation of parallel MT bundles, which includes random angular motion of MTs around the spindle pole [24], along with the attractive forces exerted by cross-linking proteins. The model predicts faster bundle formation if MTs diffuse faster and the density of cross-linking proteins is higher, which we tested experimentally. We conclude that the angular motion drives the alignment of MTs, which in turn allows the cross-linking proteins to connect the MTs into a stable bundle.

*Experimentally observed bundle formation.* The process of MT bundle formation can be observed experimentally in the fission yeast *Schizosaccharomyces pombe* because of a small number of MTs in the spindle. At the onset of mitosis, two spindle pole bodies nucleate MTs that form the spindle and additional MTs grow from the spindle pole bodies performing angular motion [24]. In our experiments, we observed that MTs growing at an angle with respect to the spindle eventually join the bundle of spindle MTs (Fig. 1a, Supplementary Movie 1). Such events are also accompanied by an increase in the tubulin-GFP signal intensity in the spindle, suggesting an increase in the number of MTs in the spindle and arguing against the scenario in which one of the MTs depolymerized (Fig. 1b). Additionally, we used cells with GFP-labeled Mal3, a protein that binds to the growing end of the MT [30]. Here we observed MT bundling at a finer time resolution (Fig. 1c, Supplementary Movie 2 and the increased Mal3 signal in the spindle after bundling (Fig. 1d). Aside from MTs joining the already formed spindle, we also observed bundling between pairs of MTs which were both freely pivoting (see Supplementary Figure 1a). We did not observe un-bundling events after
the bundles were formed. In all cases, MTs performed angular motion around the spindle pole body, which allowed them to approach each other and form a bundle.

**Theory.** To explore the physical principles underlying the formation and stability of MT bundles, we introduce the pivot-and-bond model. We describe two MTs as thin rigid rods of fixed length with one end freely joint at the spindle pole body, based on experimental observations both in vivo [24,25] and in vitro [31]. The orientation of the first MT at time $t$ is described by a unit vector $\mathbf{r}(t)$ (Fig. 2b). The orientation of the unit vector changes as

$$\frac{d\mathbf{r}}{dt} = \mathbf{\omega} \times \mathbf{r},$$

(1)

ensuring that the magnitude of $\mathbf{r}$ does not change. The vector $\mathbf{\omega}$ denotes angular velocity of the MT. The other MT has a fixed orientation along the $z$-axis in the direction of unit vector $\mathbf{z}$.

In the overdamped limit, the angular friction is balanced by the torque, $\mathbf{T}$, experienced by the MT:

$$\gamma \mathbf{\omega} = \mathbf{T}. \quad (2)$$

Here, $\gamma$ denotes the angular drag coefficient. We calculate the total torque as $\mathbf{T} = \mathbf{\tau} + \sigma (\mathbf{r} \times \mathbf{n}(t))$, where the first and the second term represent the deterministic and the stochastic components, respectively. If the noise is caused by thermal fluctuations, as in fission yeast [24], $\mathbf{n} = (\eta_i), \ i = 1, 2, 3$ is a 3-dimensional Gaussian white noise where $i$-th and $j$-th components for times $t$ and $t'$ obey $\langle \eta_i(t), \eta_j(t') \rangle = \delta(t - t') \delta_{i,j}$, with $\delta(t - t')$ being the Dirac delta function and $\delta_{i,j}$ is the Kronecker delta function. The magnitude of the noise is related to the angular drag coefficient, following the equipartition theorem, as $\sigma = \sqrt{2k_B T \gamma}$, with $k_B T$ being the Boltzmann constant multiplied by the temperature. We introduce the angular diffusion coefficient, $D = k_B T / \gamma$, and the equation (2) now reads
\[ \omega = \frac{D}{k_B T} \tau + \sqrt{2D} \mathbf{f} \times \eta(t). \]  

(3)

In our model, the torque \( \tau \) in equation (3) is the consequence of forces exerted by cross-linking proteins connecting both MTs. If we denote the positions along the MTs as \( r = r \mathbf{f} \) and \( z = z \mathbf{z} \) respectively, the torque contribution from cross-linking proteins is

\[ \mathrm{d}\tau(r, z, t) = r \times f \mathrm{d}N(r, z, t), \]  

(4)

with \( \mathrm{d}N \) being the number of cross-linking proteins connecting the MT elements \( [z, z + \mathrm{d}z] \) and \( [r, r + \mathrm{d}r] \) while \( f \) is the force exerted by a single cross-linking protein. The elastic force is calculated as \( f = -k(y - y_0 \mathbf{y}) \). Here \( k \) is the Hookean spring constant, \( y = r - z \) is the elongation of the protein linking positions \( r \) and \( z \), with magnitude \( y \) and direction \( \mathbf{y} = y/y \), and \( y_0 \) is the relaxation length of the cross-linking protein. We describe the distribution of cross-linking proteins along the MTs by introducing the density, \( \rho \), which obeys \( \mathrm{d}N(r, z, t) = \rho(r, z, t) \mathrm{d}r \mathrm{d}z \). To calculate the total torque we summed up all the attached cross-linking proteins:

\[ \tau = k \int_0^R \mathrm{d}r \int_0^\infty \mathrm{d}z \rho r \times z \left( 1 - \frac{y_0}{|z - r|} \right), \]  

(5)

where we used \( r \times (z - r) = r \times z \) and allowed the fixed MT to span the entire positive \( z \)-axis. When the total number of cross-linking proteins is large we can use the mean field limit and consider them continually distributed along the MT. In this limit, the cross-linking protein density is given by:

\[ \frac{\partial \rho}{\partial t} = -\frac{\partial j_r}{\partial r} - \frac{\partial j_z}{\partial z} + k_a c(r, z) - k_d(r, z) \rho. \]  

(6)

Here, the currents describe the redistribution of cross-linking proteins along the MTs, \( j_{r,z} = v_{r,z} \rho - D_m \partial_{r,z} \rho \), where the two terms correspond to the drift and the diffusion of cross-linking proteins, respectively. In general, equation (6) can be used for motor proteins and
passive cross-linkers. In the latter case, the velocities are calculated from the balance of the elastic force and friction of cross-linking proteins moving along the MT, \( v_{r,z} = f_{r,z}/\gamma_m \), where the components of the elastic force are calculated as \( f_r = \mathbf{f} \cdot \mathbf{r} \) and \( f_z = -\mathbf{f} \cdot \mathbf{z} \). The friction coefficient is calculated using the Einstein relationship, \( \gamma_m = k_B T / D_m \). We assume that the attachment rate \( k_a \) is constant and that the detachment rate depends on the force experienced by the cross-linking proteins \([32]\), \( k_d(r,z) = k_{d0} \exp[f(y(r,z))/f_c] \), with \( f_c \) being the critical force required for rupturing the MT-protein bond. The extensions of cross-linking proteins in the nucleoplasm are in thermodynamic equilibrium, hence they obey the Boltzmann distribution. Thus, the distribution of cross-linking proteins in the nucleoplasm, with respect to their extensions, is given by

\[
c(r,z) = c_0 \sqrt{k/(2\pi k_B T)} \exp[-k(y(r,z) - y_0)^2 / 2k_B T],
\]

where the constant \( c_0 \) is the linear concentration of cross-linkers in the nucleoplasm. Equations (1)-(6) provide a complete description of angular movement for the MT in the presence of cross-linking proteins.

To obtain the time course of the MT orientation, we parameterize the orientation of the MT given by the unit vector by \( \mathbf{f}(\theta, \phi) = (\sin \theta \cos \phi, \sin \theta \sin \phi, \cos \theta) \), where \( \theta \) and \( \phi \) denote the polar and azimuthal angle, respectively. In this parameterization, the equation of motion for the polar angle reads

\[
\partial_t \theta = -D \partial_{\theta} U(\theta) + \sqrt{2D} \eta(t) \tag{7}
\]

The normalized potential describing the interaction between the MTs, \( U(\theta) \), is implicitly defined as \( -\partial_{\theta} U(\theta) = \tau / k_B T + \cot \theta \), where \( \tau \) denotes the magnitude of the torque and \( \cot \theta \) is the spurious drift term \([33]\) (see Supplementary Note for more details and derivation). This equation is sufficient to describe the bundling process, because the angle between the MTs is given only by the polar angle, the dynamics of which is independent of the azimuthal angle.

In the adiabatic approximation, \( \partial \rho / \partial t = 0 \), equation (6) yields a one-dimensional cross-linker
density profile $\rho_r(r)$ (see Supplementary Note for exact expression). Integrating the density profile over the entire length of the MT allows us to calculate the torque exerted by cross-linking proteins, which in turn allows us to calculate the generalized potential. The normalized potential has an analytical expression $U(\theta) = -\theta_{\text{max}} \left( \frac{\theta(\theta-\theta_{\text{min}})}{\theta_{\text{min}}} + \frac{\theta(\theta_{\text{min}}-\theta)}{\theta_{\text{min}}} \right) - \ln(\sin \theta)$. Here, $\theta_{\text{max}} = \frac{k_a c_0 y_0}{k_{do}}$ is the local maximum of the potential, $\theta_{\text{min}} = \frac{y_0}{R}$ is the local minimum and $\Theta(\theta - \theta_{\text{min}})$ is the Heaviside step function. The depth of the potential well in our model, and therefore the strength of the MT-MT interaction, depends on the attachment and detachment rates of cross-linkers, their concentration in the nucleoplasm and the MT length.

By numerically solving equations (7) for the polar angle, for a large initial angle, we found that the MT performs random movement and spans a large space (Fig. 2c). However, the movement can become abruptly constrained in the vicinity of angle zero. These small angles correspond to a bundled state. Our numerical solutions also show that, in rare cases, constrained MT movement in the vicinity of angle zero can suddenly switch back to free random movement (Fig. 2d). The constrained movement near angle zero is a consequence of short range attractive forces exerted by the cross-linkers that accumulate in larger densities when MTs are in close proximity (compare green and magenta lines in Fig. 2e). The density is constant up to $r_0$ because in that region, the cross-linkers can always attach in a relaxed configuration, while for $r > r_0$, the cross-linkers will always be under tension and their density will drop off dramatically as $r$ increases further (see inset in Fig. 2e and Supplementary Note).

After showing numerically that our model reproduces experimentally observed MT bundling processes (Fig. 2f), we proceed with an analytical approach to provide a complete picture of the stability of MT bundles and their formation. We start by examining the MT-MT
interaction potential. The shape of the normalized potential depends on the parameters of the system, as shown in Fig. 3a for different values of MT length, $R$, and nucleoplasm cross-linker concentration, $c_0$. Because MT bundles can be assembled and disassembled, as shown in Fig. 2c and 2d, we treat the MT as a two state system. The position of the maximum of the normalized potential defines two regions, $\Theta_B \equiv [0, \theta_{\text{max}}]$ and $\Theta_U \equiv [\theta_{\text{max}}, \pi]$, which we term as the bundled and unbundled state, respectively.

Next, we explored the stability of the bundles. We define bundles as stable if the probability of finding the MT in the bundled state, $P_B$, is greater than the probability of finding it in the unbundled state, $P_U$,

$$P_B > P_U. \quad (8)$$

The bundling and unbundling probabilities are calculated as $P_{B,U} \propto \int_{\Theta_{B,U}} e^{-U(\theta)} d\theta$ (bundling probability is shown in Fig. 3b). Here we used the stationary probability distribution [33], $e^{-U(\theta)}$, and normalized using $P_B + P_U = 1$. The sharp transition from zero to one around the value $P_B = 1/2$ introduces a natural boundary between stable and unstable bundles. We find that the bundles are unstable if MTs are too short or if there are not enough cross-linking proteins in the nucleoplasm (see Fig. 3c).

Finally, we calculate how the MT bundling time depends on the parameters of the system. In the case of an isotropic distribution of initial MT orientations, we calculate the bundling time as $\langle t_B \rangle = \frac{1}{2} \int_{\theta_{\text{min}}}^{\pi} \langle t \rangle_{\theta, \theta_{\text{min}}} \sin \theta \, d\theta$, where $\langle t \rangle_{\theta, \theta_{\text{min}}}$ is the first passage time from an initial angle $\theta$ to the angle $\theta_{\text{min}}$ (ref. [34]). After solving these integrals numerically, we found that the bundling time normalized by the diffusion coefficient, $\langle t_B \rangle D$, decreases as the cross-linker concentration increases (solid black line in Fig. 3d), but is not significantly affected by the MT length (solid green line in Fig. 3d). Note that the bundling time, $\langle t_B \rangle$, is
inversely proportional to $D$, which decreases with MT length [24], thus we expect that the bundling time increases with MT length. We also calculate the unbundling time as $\langle t_u \rangle = \frac{1}{2} \int_{\theta_{\min}}^{\pi} d\theta \sin \theta \langle t \rangle_{\theta_{\min} \theta}$, where $\langle t \rangle_{\theta_{\min} \theta}$ is the first passage time from $\theta_{\min}$ to $\theta$. Unbundling time becomes longer than bundling time if the condition for bundle stability, $P_B > P_U$, is fulfilled. Once this condition is satisfied, the bundling time greatly increases (dashed line in Fig. 3d).

In order to compare the theory with experimental observations, we measured the bundling time as the total observation time of MTs divided by the number of observed bundling events, $\langle t_B \rangle = t_{\exp}/n$. Along with the wild type cells, we also performed the measurements on the mutant in which the Ase1 cross-linker was knocked out (denoted $ase1\Delta$, ref. [35,36]), in which we also observed MT bundling (see Supplementary Figure 1b and Supplementary Figure 2). We observed that the bundling time increases with MT length (Fig. 3e), and that the bundling time is significantly longer in $ase1\Delta$ cells (compare green and magenta line in Fig 3e). We normalized bundling time by the diffusion constant (see Supplementary Note) and found a weak dependence on MT length, but a significant increase in $ase1\Delta$ cells compared to wild type (inset of Fig 3e). The theory reproduces the weak dependence on MT length and implies that the deletion of ase1 decreases the effective cross-linker concentration roughly five fold.

In conclusion, our work implies that only passive processes, thermally driven motion of the MTs and passive cross-linkers, are sufficient to describe the formation of parallel MT bundles. By introducing the pivot and bond model we gain a deeper understanding of the mesoscopic properties of the bundling process, such as bundle stability and average bundling time, as well as predict their dependence on biological parameters such as MT length and
cross-linker concentration. This approach is complementary to more exhaustive and detailed methods such as large-scale numerical simulations (for example [28,37]).

Along with parallel MT bundles, mitotic spindles also contain bundles of anti-parallel MTs, which are made of MTs extending from the opposite spindle poles. The theory developed here could be generalized to describe MTs extending from two spindle poles by adding additional angular variables and including directional movement of cross-linking proteins. Just like here, such a model will give insight into the minimal requirements for the formation of anti-parallel MT bundles and therefore shed additional light on the physics of spindle formation.


[38] M. T. Valentine, P. M. Fordyce, and S. M. Block, Eg5 steps it up!, Cell Div 1, 31 (2006).


FIGURE LEGENDS

**FIGURE 1**

**a** Tubulin-GFP Sid4(SP)-GFP

**b** Mal3-GFP Sid4(SP)-GFP

Fig. 1. Formation of MT bundles in *S. pombe* cells. (a) Time-lapse images and the corresponding drawings showing the formation of a parallel MT bundle at the lower spindle pole body in an *S. pombe* cell expressing tubulin-GFP and Sid4-GFP. (b), Measurement of the tubulin-GFP signal intensity of MTs before bundling (green curve in the graph, measured along the green line in the inset) and after bundling (magenta curve in the graph, measured along the magenta line in the inset). The measurements were done on the first and the last image in panel a, respectively. (c), Time-lapse images and the corresponding drawings showing a MT joining the bundle of spindle MTs in an *S. pombe* cell expressing Mal3-GFP.
and Sid4-GFP. (d), Measurements of the Mal3-GFP signal intensity of the spindle and MT before bundling (green curve in the graph, measured along the green line in the inset) and after bundling (magenta curve in the graph, measured along the magenta line in the inset). The measurements were done on the first and the last image in panel e, respectively. Scale bars in panels a and b are 1 μm. In the drawings, microtubule orientations from the previous images are marked with white dashed lines.
Fig. 2. Scheme of the model and numerical solutions. (a), Cartoon representation of the bundling process. Left, a MT (green rod) pivots around the spindle pole body (grey ball). Right, when the MTs come into close proximity, cross-linking proteins (grey springs) attach and cause the MTs to from a bundle. (b), Scheme of the model. The orientations of two MTs are represented by the unit vectors $\hat{r}$ and $\hat{z}$. Cross-linking proteins attach to and detach from MTs at rates $k_a$ and $k_d$, respectively. The elongation of the attached cross-linking protein is denoted $y$ and their relaxed length is $y_0$. The angle between the MTs is denoted $\theta$. (c), A sample path for the starting angle $\theta = 0.9 \text{ rad}$, which shows a bundling event (around the 2.5 min mark). (d), A sample path for an unbundling event (around the 2 min mark). (e), Cross-linking protein density profiles along the MT for two points on the path shown in c, denoted by arrowheads in corresponding colors (large image). The point $r_0$, at which cross-linker density is no longer constant, is shown for the magenta line. The inset shows a schematic of the orientations of attached cross-linkers. The cross-linkers attached within the distance $r_0$ from the spindle pole body (boundary shown by dashed line) are always relaxed, while those attached at larger distances from the spindle pole body are elongated. (f), A sample of MT angle time series obtained using light microscopy on cells with the Mal3-GFP label. All calculations are done with $c_0 = 300 \mu m^{-1}$, $R = 1.5 \mu m$, $D = 0.001 \text{ rad}^2 \text{s}^{-1}$ and other parameters shown in Table 1.
Fig. 3. Solutions of the model in the adiabatic approximation. (a), The effective potential as a function of the polar angle, for angles between 0 rad and 0.4 rad. The inset shows the potential for all angles. Dashed square represents the region plotted in the large image. (b), Bundling probability as a function of $c_0$ for three different values of $R$. The colors of the dots correspond to the color code of the parameters used in a. (c), Phase diagram. The gray area represents the region where bundles are stable. The arrowheads represent the values of $R$ used.
in b. (d), Normalized bundling and unbundling times. The black lines represent the normalized bundling (solid line) and unbundling time (dashed line) as a function of $c_0$ for $R = 1.5 \mu m$. The green line represents the normalized bundling time as a function of $R$ for $c_0 = 250 \mu m^{-1}$. (e) Experimentally measured bundling times for wild type (green) and $ase1\Delta$ cells (magenta). Inset shows the comparison between measured normalized bundling time (dots with error bars) and theoretical curves for $c_0 = 500 \mu m^{-1}$ (solid green line) and $c_0 = 100 \mu m^{-1}$ (solid magenta line).
Table 1. Values of the constant parameters used in this paper.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$</td>
<td>$0.1 , pNnm^{-1}$</td>
<td>Value for Eg5 (ref. [14,38])</td>
</tr>
<tr>
<td>$f_c$</td>
<td>$3 , pN$</td>
<td>Value for kinesin-1 (ref. [39])</td>
</tr>
<tr>
<td>$y_0$</td>
<td>$40 , nm$</td>
<td>Value for Prc1 (ref. [40,41])</td>
</tr>
<tr>
<td>$D_m$</td>
<td>$0.05 , \mu m^2s^{-1}$</td>
<td>Value for Ase1 (ref. [13,42])</td>
</tr>
<tr>
<td>$k_{on}$</td>
<td>$0.01 , s^{-1}$</td>
<td>Value for Ase1 (ref. [13])</td>
</tr>
<tr>
<td>$k_{off}$</td>
<td>$0.1 , s^{-1}$</td>
<td>Value for Ase1 (ref. [13])</td>
</tr>
<tr>
<td>$A$</td>
<td>$3 , \mu m$</td>
<td>Fission yeast nuclear diameter [24]</td>
</tr>
</tbody>
</table>

Author contributions and acknowledgements

N.P. and I.T. designed and supervised the project. M.P. and N.P. developed the theory. L.W. carried out the experiments and A.M. analyzed the experimental data. N.P., M.P. and I.T wrote the manuscript with input from L.W. and A.M. The data that support the findings of this study are available from the corresponding authors upon request. Contact N.P. for information on the theory presented and I.M.T with requests related to the experimental measurements.

We thank D. Radić and V. Despoja for comments on the manuscript and M. Glunčić for valuable discussions and advice regarding our theoretical model; F. Elsner from the Electronics Service of MPI-CBG for building the thermoelectric device; J. Millar for yeast strains; Light Microscopy Facility of MPI-CBG for help with microscopy; J. Brugués, T.M. Franzmann, A. Garcia Ulloa and the rest of the Tolić lab for discussion and advice; and I. Šarić for editing the figures. We acknowledge the Unity through Knowledge Fund (UKF), German Research Foundation (DFG), European Research Council (ERC) and the QuantiXLie Center of Excellence for funding.