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Polymorphic Dynamics of Microtubules

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Starting from the hypothesis that the tubulin dimer is a conformationally bistable molecule - fluctuating between a curved and a straight configuration at room temperature - we develop a model for polymorphic dynamics of the microtubule lattice. We show that tubulin bistability consistently explains unusual dynamic fluctuations, the apparent length-stiffness relation of grafted microtubules and the curved-helical appearance of microtubules in general. Analyzing experimental data we conclude that taxol stabilized microtubules exist in highly cooperative yet strongly fluctuating helical states. When clamped by the end the microtubule undergoes an unusual zero energy motion - in its effect reminiscent of a limited rotational hinge.

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Microtubules are the stiffest cytoskeletal component and play versatile and indispensable roles in living cells. They act as cellular bones, transport roads [1] and cytoplasmic stirring rods [2]. Microtubules consist of elementary building blocks - the tubulin dimers - that polymerize head to tail into linear protofilaments (PFs). PFs themselves associate side by side to form the hollow tube structure known as the microtubule (MT). Despite a long history of their biophysical study a deeper understanding of MT's elastic and dynamic properties remains elusive to this date. Besides the unusual polymerization related non-equilibrium features like "treadmilling" and the dynamic instability there are a number of other experimental mysteries - in thermal equilibrium - that presently defy coherent explanations, most notably:

(i) The presence of high "intrinsic curvature" [3]-[6] of unclear origin, also identified as a long wave-length helicity [3].
(ii) In various active bending [7] or thermal fluctuation experiments [4][5][8] MTs display length dependent, even non-monotonic apparent stiffness [5].
(iii) They exhibit unusually slow thermal dynamics in comparison with standard semiflexible filaments [5][6].

The most bizarre and controversial feature (ii) has been the subject of much debate and some theoretical explanation attempts based on low shear stiffness modulus have been put forward [9]. However a careful reanalysis of clamped MT experiments, Figs 2, 3 reveals two features not captured by these initial models: the lateral end-fluctuations scale as \( \sim L^2 \) while the relaxation times scale as \( \sim L^3 \). This exotic behavior naively suggests the presence of a limited angular hinge at the MT clamping point. On the other hand artifacts that could trivially lead to a "hinged behavior" (like loose MT attachment and punctual MT damage) were specifically excluded in experiments [4][5]. We will outline here a model based on internal MT dynamics explaining phenomena (i)-(iii). It leads us to the origin of MT helicity (i) implying (ii)-(iii) as most natural corollaries [10]. The two central assumptions of our model are as follows:

(I) The tubulin dimer is a conformationally multistable entity and fluctuates between at least 2 states on experimental time scales.
(II) There is a nearest-neighbor cooperative interaction of tubulin states along the PF axis. We are lead to assumptions I-II from several independent directions: First, the experimentally observed MT helicity [3] implies that there is a symmetry breaking mechanism of individual PF’s conformational properties. In analogy to the classic case of bacterial flagellum the existence of helices in azimuthaly symmetric bundles also necessitates a cooperative longitudinal interaction along protofilaments [11][12]. Second, investigations of single protofilament conformations by Elie-Caille et al [13] reveal that a single taxol- PF can coexist in at least 2 states with comparable free energy: a straight state \( \kappa_{PF} \approx 0 \) and a weakly curved
state with intrinsic curvature $\kappa_{PF} \approx 1/250nm$. These authors also point out the apparent cooperative nature of straight to curved transition within single PFs. Third, when mechanically buckled by AFM tips tubulin dimers occasionally switch back to the initial straight conformation [14]. Fourth, tubulin multistability was inferred from the formation of stable circular MT arcs in kinesin driven gliding assays by Amos & Amos [15]. Unfortunately their clear, seminal observations were subsequently forgotten for decades leading to much of the confusion about MTs we are witnessing today.

**Polymeric MT Model.** Starting from assumptions I – II we model the tubulin dimer state by a two variable state $\sigma_n(s) = 0.1$ (the tubulin dimer in the "straight"/"curved" state, cf. Fig. 1a) at each lattice site with circumferential PF index $n = 1,...,N$ ($N = 11 - 16$ number of PFs) at longitudinal arc length centerline position $s$. The total elastic + conformational energy can be written as $E_{MT} = \int_0^L (e_{cl} + e_{trans} + e_{inter}) ds$ with

$$e_{cl} = \frac{Y}{2} \int \int (\varepsilon - \varepsilon_{pol})^2 r dr ds$$

$$e_{trans} = -\frac{\Delta G}{b} \sum_{n=1}^N \sigma_n(s),$$

$$e_{inter} = -\frac{J}{b} \sum_{n=1}^N (2\sigma_n(s) - 1)(2\sigma_n(s + b) - 1)$$

where the integration in $e_{cl}$ goes over the annular MT cross-section with $r_1 \approx 7.5nm$, $r_2 \approx 11.5nm$ the inner and outer MT radii, with $\Delta G > 0$ the energy difference between the 0 and 1 state and $b \approx 8nm$ the monomer length, $J$ the “Ising” cooperative coupling term along the PF contour and with the polymorphism induced pre-strain $\varepsilon_{pol} \propto \varepsilon_{PF} \sigma_n(s)$ [16] where $\varepsilon_{PF}$ is the strain generated in the curved state. The latter can be estimated from the switched PF curvature $\kappa_{PF} \approx (250nm)^{-1}$ [13] to be $\varepsilon_{PF} \approx \varepsilon_{pol} \kappa_{PF}/2 \approx 10^{-2}$. For an isotropic Euler-Kirchhoff beam, the actual material deformations are related to the centerline curvature via $\varepsilon = -\kappa \vec{r} \cdot \vec{r}$ with $\vec{r}$ the radial vector in the cross-section.

Upon inspection it becomes clear that the phase behavior (straight or curved state stability) is contained in the interplay of the first two terms $e_{cl}$ and $e_{trans}$ while the thermal dynamics is governed by the 3rd $e_{inter}$ which rules over defect behavior (cf. Fig. 1d). To understand the basic behavior we first consider a short MT section along which the PFs are in a uniform state $\sigma_n(s) = \sigma_n(s + b)$ ($e_{inter} = \text{const.}$ can be dropped). Furthermore we resort to the single block ansatz, i.e. at each cross-section there is only one continuous block of switched PFs of length $p$. This ansatz was successfully used by Calladine in modelling bacterial flagellin polymorphic states [11]. In this approximation the energy density becomes

$$e = \frac{B}{2} \left( (\kappa - \kappa_{pol}(p))^2 + \kappa_1^2 \left( \frac{\pi}{N} \phi - \sin^2 \left( \frac{\pi}{N} \phi \right) \right) \right)$$

with the bending modulus $B = \frac{Y r}{12}$ and the polymeric curvature $\kappa_{pol}(p) = \kappa_1 \sin \left( \frac{\pi}{N} p \right)$ with $\kappa_1 = \frac{\kappa_{PF}(r_2 - r_1)^2}{\pi (r_1^2 + r_2^2)}$. The MT phase behavior depends on the polymeric-elastic competition parameter $\gamma = \frac{\kappa_{PF}}{\kappa_1} - 2N \frac{\sigma_1}{Y}$. Physically, $\gamma$ measures the ratio between polymeric energy of tubulin switching and the elastic cost of the transition. For $\gamma < -1$ all the PFs are in the (highly prestrained) state $\sigma = 1$ while for $\gamma > 1$ all them are in the state $\sigma = 0$ - both corresponding to a straight MT. For $-1 < \gamma < 1$ we have coexistence of 2 (locally) stable states: straight ($p = 0$ or $p = N$) and curved state with $p > 0$. For $-\frac{\pi}{2} < \gamma < \frac{\pi}{2}$ the curved state is the absolute energy minimum and the straight state is only metastable. Therefore in this regime, the ground state of a microtubule bearing natural lattice twist will be helical (cf. Fig. 1b). Assuming a stable helical state as observed in [3] we have $p/N \in [1/4,1/2]$ giving us an estimate for the radius of curvature $\kappa_{pol} \approx 9 - 14nm$. This compares favorably with an estimate of observed helices $\kappa^{-1} \approx 11nm$ from [3]. The helical stability and the magnitude of the protofilament curvature $\kappa_{PF} \approx 1/250nm$ [13] with a typical protein Young modulus $Y \approx 1 - 10GPa$, allows us also a simple estimate of the transition energy per monomer $\Delta G \approx +1.1 + +11kT$. In general, the energy in Eqs.1-3 gives rise to a complex behavior and we focus on basic phenomena. It turns out that a most remarkable deviation from standard wormlike chain behavior comes from the change of polymorphic phase that we consider in the following.

**Polymeric Phase Dynamics.** To better understand the central phenomenon, we define at each MT cross-section the complex polymeric order parameter $P(s) = \sum_{n=1}^N e^{i2\pi n/N} \sigma_n(s) = |P(s)| e^{i\theta(s)}$ where $|P(s)|$ denotes the "polymeric modulus" and $\theta$ the "polymeric phase" (cf. Fig. 1b). The polymeric state can then be described by the local (complex) centerline curvature $\kappa_{pol}(p) = \kappa_0 e^{i\theta_0} P(s)$ with $\kappa_0 = \kappa_1 \sin \pi/N$ and $\theta_0$ the natural lattice twist that varies with PF number [17]. This gives rise to a helical MT shape described by the curvature $|\kappa_{pol}| = \kappa_{pol}$ and torsion $\tau \approx \phi' + \theta_0$. For large acting forces both the polymeric phase $\phi$ and amplitude $|P|$ will vary along the contour, however for small (thermal) perturbations the phase fluctuations will be dominant [18]. Based on this and on the observation of stable helical states [3] we will now assume $|P| = \text{const.}$ and write the total energy of the MT whose centerline deflection is described by a complex angle $\theta(s) = \theta_x(s) + i\theta_y(s)$ (deflection angles in x/y direction) as follows:

$$E_{tot} = E_{cl} (\theta, \phi) + E_{pol} (\phi)$$

The first energy term is the "wormlike-chain" bending contribution $E_{cl} (\theta, \phi) = \frac{B}{2} \int (\theta' - \kappa_{pol})^2 ds$. The second term is the polymeric phase energy $E_{pol} (\phi) = \int \frac{Y}{2} (\phi'^2 + \kappa_1^2 \sin^2 \theta) ds$.
where the polymorphic fluctuations induce a strong position / distance dependence - a behavior that could be interpreted as "length dependent persistence length". Indeed for $l_φ \gg s > q_0^{-1}$ the persistence length displays a non-monotonic oscillatory behavior around a nearly linearly growing average value $I_p^*(s) \approx \frac{2}{3} \frac{q_{0}^{2}}{\kappa_{0}^{3}} s + \frac{4}{3} \frac{q_{0}^{2}}{\kappa_{0}^{3}} \sin(q_{0}s)$. This oscillation is related to the helical ground state while the linear growth $l_p^*(s) \sim A^2s$ is associated to the conical rotation of the clamped chain (wobbling mode), cf. Fig. 1c, with an angle $α = 2κ_{0}q_{0}^{-1}$. For $s > l_0$ the saturation regime with a renormalized $I_p^*(\infty) = 1/\left(l_p^{*0} + l_B^0\right)$ with $l_p^{*0} = 2l_0q_0^2\kappa_{0}^{-2}$ is reached. The theory can now be compared with the experimental data [4][5] (cf. Fig. 2) that reveal several interesting characteristics in agreement with predictions. In particular the mean linear growth of $l_p^*(L)$ (single parameter fit $l_p^* \sim L^\delta$ with $δ = 1.05$) and the non-monotonic $I_p^*(L)$ dependence [5] are well captured by the theory. The linearly growing experimental spread of $l_p^*$ with $L$ is likely linked to the spread of $q_{0}$ in the MT lattice populations [17]. The large length plateau $s > l_0$ is not reached even for longest MTs ($\sim 50μm$) in agreement with coherent helices [3]. Our best comparison between theory and experiments (cf. Fig. 2) gives $l_B = 25mm$ corresponding to $Y \approx 9GPa$ (proteins with $Y$ up to $19GPa$ exist [22]) and a helix wave length $λ \approx 7.5μm$. This is close to the expected $6μm$ corresponding to the twist [17] of the predominant 14 PF MTs fraction in the in-vitro MTs preparation of [4][5]. It turns out that $l_B$ is larger than in previous studies $l_B \sim 1-6mm$ where however polymorphic fluctuations were neglected. The absence of the plateau also allows a lower estimate of the coherence length $l_0 > 55μm$ and the coupling constant $J > 4k_BT$.

**Polymorphic Phase Dynamics.** To describe the MT fluctuation dynamics we consider the total dissipa-
tion functional \( P_{diss} = P_{ext} + P_{int} \) which is composed of an internal dissipation contribution \( P_{int} = \frac{1}{2} \xi_\perp \int \dot{\phi}^2 \, ds \) and an external hydrodynamic dissipation \( P_{ext} = \frac{1}{2} \xi_\perp \int |\dot{\rho}|^2 \, ds \) with \( \xi_\perp = 4\pi\eta/(2L/r - 1/2) \) the lateral friction constant, \( \eta \) the solvent viscosity, \( r \) and \( L \) the MT radius and length. The time evolution equation of the phase variable \( \phi(s,t) \) and elastic displacement \( \rho_L(s,t) \) is given by the coupled Langevin equations \( \frac{d\phi}{d\xi} = -\frac{\partial \delta P_{diss}}{\partial \phi} + \Gamma_\phi \) and \( \frac{d\rho_L}{d\xi} = -\frac{\partial \delta P_{diss}}{\partial \rho_L} + \Gamma_\rho \) with \( \Gamma_{\phi/\rho} \) the thermal noise term. In general this dynamics is highly non-linear however in the experimentally relevant regime where the behavior is dominated by the wobbling mode the equations simplify greatly and we end up with a simple diffusive behavior of the wobbling mode \( \frac{d\phi}{d\xi} = \frac{1}{\xi_\perp} L^{-1} \int_0^L \Gamma_L(s,t) \, ds \) with a friction constant given by \( \xi_\perp = \xi_\parallel + \xi_\perp \) where \( \xi_\perp = 2\xi_\perp \kappa_0^2 q_0^{-4} \left( (1 + \cos L q_0) - 4 \sin L q_0 + q_0^2 L^3/3 \right) \). For comparison with the experiment we compute the time correlation of the \( y \) deflection. A short calculation gives \( \langle y(t), y(t) \rangle \propto e^{-t/\tau_y(L)} \) with the relaxation time \( \tau_y(L) \approx \xi_\parallel /k_{BT} \). For small lengths, \( \tau_y(L) \approx L \xi_\parallel /k_{BT} \) is dominated by internal dissipation while for large lengths \( \tau_y(L) \approx \sqrt{\kappa_0/q_0} L^3 \) a careful analysis of the experimental data [5] reveals in fact the latter scaling. An independent single exponent fit gives \( \tau \propto L^\alpha \) with \( \alpha = 2.9 \). Using the value \( \kappa_0/q_0 \approx 4.8 \times 10^{-3} \) from Fig. 2 and \( \xi_\parallel \approx 2\eta \) with \( \eta = 10^{-3} \mu_0 s \) [23] we find the theoretical value \( \tau_{th}/L^3 = 7.9 \times 10^{14} s/m^3 \) that can be compared with the fit of experimental data (Fig. 3) \( \tau_{fit}/L^3 = 6.25 \times 10^{14} s/m^3 \). The excellent agreement of both the exponent and the prefactor leads us again to the strong conclusion that in these experiments the clamped MT is an almost rigid helical polymorphic rotor whose behavior is dominated by the zero energy ("wobbling") mode and hydrodynamic dissipation. For very short MTs the linearly scaling internal dissipation dominates and we could measure \( \xi_\parallel \) from the limit value of \( \tau_y/L \), for \( L \to 0 \). For the available data \( L > 2 \mu m \) [5] this plateau-regime is not yet fully developed and we can only provide an upper estimate from the data \( \xi_\parallel \approx 4 \times 10^{-17} N/s \).

**Conclusion.** The MT fluctuations are well described - both dynamically and statically - by the bistable tubulin model and the reason for appearance of MT helices becomes obvious. The otherwise mysterious lateral fluctuations reflected in \( \rho_p(L) \sim L^3 \) and \( \tau_p(L) \sim L^3 \) scaling are mere consequences of the "wobbling motion" of a polymorphic cooperatively switching helical lattice. We speculate that the implied conformational multistability of tubulin and the allosteric interaction are not just nature’s way to modulate the elastic properties of its most important cytoskeletal mechano-element. It could also be a missing piece in the puzzle of dynamic instability. Another intriguing possibility of using this switch for long range conformational signalling in vivo, could hardly have been overlooked by evolution. I.M.K thanks Francesco Pampaloni for stimulating discussions.

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[19] This mode has a N fold symmetry however for large number of PFs \( N = 11 - 15 \) it can be approximated as continuous.
[20] Another more common definition, from angular correlation \( \langle \cos(\theta(s) - \theta(s')) \rangle \) exhibiting a similarly rich behavior as \( I_p^p \) (yet a distinct functional form) will be discussed elsewhere.
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