Abstract

αβ-tubulin dimers need to convert between a ‘bent’ conformation observed for free dimers in solution and a ‘straight’ conformation required for incorporation into the microtubule lattice. Here, we investigate the free energy landscape of αβ-tubulin using molecular dynamics simulations, emphasizing implications for models of assembly, and modulation of the conformational landscape by colchicine, a tubulin-binding drug that inhibits microtubule polymerization. Specifically, we performed molecular dynamics, potential-of-mean force simulations to obtain the free energy profile for unpolymerized GDP-bound tubulin as a function of the ~12° intradimer rotation differentiating the straight and bent conformers. Our results predict that the unassembled GDP-tubulin heterodimer exists in a continuum of conformations ranging between straight and bent, but, in agreement with existing structural data, suggests that an intermediate bent state has a lower free energy (by ~1 kcal/mol) and thus dominates in solution. In agreement with predictions of the lattice model of microtubule assembly, lateral binding of two αβ-tubulins strongly shifts the conformational equilibrium towards the straight state, which is then ~1 kcal/mol lower in free energy than the bent state. Finally, calculations of colchicine binding to a single αβ-tubulin dimer strongly shifts the equilibrium toward the bent states, and disfavors the straight state to the extent that it is no longer thermodynamically populated.

Introduction

Microtubules (MTs) are dynamic cytoskeletal polymers formed by the polymerization of αβ-tubulin, a globular heterodimer comprised of two structurally related 55 kDa α- and β-subunits. Microtubules play a vital role in intracellular trafficking and cell division; these functions are influenced by the complex dynamics of the MT plus end, which undergoes stochastic periods of assembly and disassembly. The role of αβ-tubulin conformational changes in the processes of assembly and disassembly has been the subject of great interest and some controversy [1]. Here we investigate the free energy landscape of αβ-tubulin using molecular dynamics simulations, emphasizing implications for models of assembly, and modulation of the conformational landscape by colchicine, a tubulin-binding drug that inhibits microtubule polymerization.

Tubulin has been shown to exist in two extreme conformations: a “straight” conformation observed in antiparallel zinc-induced tubulin sheets [2–6], which is compatible with incorporation into the MT lattice, and a “bent” conformation observed in the structure (T2R complex) of tubulin bound with colchicine, a MT-stabilizing drug, in a complex with the stathmin-like domain of RB3 (SLD-RB3) [7]. (The resolution of this T2R-colchicine structure was recently enhanced to 2.73 Å [8].) X-ray crystallographic studies have also shown similarly bent tubulin structures in a complex with SLD-RB3 [9], bound to other MT-stabilizing drugs (vinblastine [8] and podophyllotoxin), as well as MT-stabilizing drugs (epothilone A and zampanolide) in a complex with tubulin tyrosine ligase (TTL) [10]. Distinguishing the straight and bent tubulin conformations are conformational rearrangements of the intermediate domains and an intra-dimer rotation: rotations of ~8° and ~11°, are required to superimpose the intermediate domains of the α- and β-subunits, respectively. In addition a ~12° intradimer rotation is required to superimpose both subunits of the straight and bent tubulin structures. Finally, the “straight” and bent tubulins are also distinguished by local rearrangements in the M and H1-S2 loop on both subunits, the β-subunit T7 loop and H8 helix, and the α-subunit T5 and H6-H7 loops [4,7,11].

Multiple rotational and translational motions in the α- and β-subunits differentiate the “straight” and “bent” conformations. We describe the ~12° intradimer conformational change of the
Author Summary

Microtubules are composed of αβ-tubulins that play an instrumental role in regulating intracellular trafficking and formation of the mitotic spindle during mitosis and cell division. Structural studies have shown that tubulin exists in a "straight" conformation compatible with that in the microtubule lattice and a "bent" conformation thought to represent the unassembled state. There is current debate as to whether the straight-to-bent conformational change in tubulin is the cause or consequence of tubulin's assembly into the microtubule lattice. Here, we use free-energy molecular dynamics simulations to qualitatively understand the conformational landscape of tubulin in the unassembled state and upon lateral binding. We predict that soluble tubulin exists primarily in a bent conformation; our simulation results show that tubulin primarily adopts an intermediate bent conformation in agreement with structural data. We also show that lateral binding of two tubulins shifts the equilibrium in favor of the "straight" state, supporting the hypothesis that the straight-to-bent conformational change is the consequence of tubulin's incorporation into the microtubule lattice via lateral interactions. We also show that colchicine binding shifts the population of tubulin in favor of a bent state, further implicating our work in drug discovery.

These previously reported molecular dynamics simulations also examined differences between GTP- and GDP-bound tubulin, an issue that we do not consider here.

In this study we utilize free-energy calculations to characterize the conformational landscape of a tubulin heterodimer, interpolating between the "straight" structure from Zn2+-induced protofilaments and "bent" tubulin from the T2R-colchicine complex. In contrast to unrestrained molecular dynamics simulations, we explore tubulin conformations only along a coordinate connecting these two states, which we quantify using an intradimer rotation angle (Figure 1). However, by performing umbrella sampling and analysis using the weighted histogram analysis method (WHAM), we can estimate the free energy associated with deforming tubulin along this coordinate, providing quantitative predictions concerning the relative free energies of the bent and straight states. Specifically, this "potential of mean force" (PMF) predicts that tubulin can exist in a continuum of conformations ranging between straight and bent, but in agreement(27,19),(987,982)
We consider below the relationship of these structures to the conformational coordinate calculated in this work. We selected 1SA0 to define the bent endpoint because (1) it has the same sequence (bovine) as the selected straight tubulin structure, (2) it has a conformation that is somewhat more ‘bent’ than the other structures, and (3) it includes colchicine, making it particularly appropriate for defining changes in the free energy profile caused by colchicine binding. These endpoint structures were processed as follows.

**Unpolymerized tubulin.** For the straight tubulin endpoint structure, taxol and the Zn$^{2+}$ ion were removed from 1JFF while preserving the Mg$^{2+}$ ion-coordinated GTP molecule on the N-site of the $\alpha$-subunit and the GDP molecule on the E-site of the $\beta$-subunit. Missing residues of the $\alpha$-H1-S2 loop (residues \(\alpha:35–60\)) were incorporated using those in Nogales et al.’s earlier Zn$^{2+}$-induced protofilament model (PDB entry 1TUB [6]). For bent tubulin, colchicine was removed from the $\alpha,\beta$-heterodimer in 1SA0 while preserving the Mg$^{2+}$ ion-coordinated GTP molecule on the $\alpha$-subunit N-site and GDP molecule on the $\beta$-subunit E-site. The PRIME 3.1 module [25] from the Schrödinger 2012 suite of programs was used to predict loop conformations for the remaining structurally unresolved residues in the $\alpha$-subunit H1-S2 loop (residues $\alpha:35–60$) and the $\beta$-subunit M loop (\(\beta:271–283\)).

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**Figure 1.** Conformational rearrangements of $\alpha$- and $\beta$-subunits of the straight (PDB entry 1JFF [4], resolution 3.5 Å) and bent (PDB entry 1SA0 [7], resolution 3.58 Å) tubulin heterodimers. The N-terminal nucleotide binding (residues 1–205) and C-terminal (residues 382–437) domains are rendered as grey ribbons. Helices and $\beta$-strands comprising the intermediate domain (residues 206–381) are colored blue (straight) and red (bent). We quantify curvature of the straight and bent heterodimers based on a $\sim 12^\circ$ intradimer rotation between the $\alpha$- and $\beta$-H7 helices, after superimposing the $\alpha$-H7 helices. Figure was generated using PyMOL [49].

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**Figure 2.** Crystal structure of the colchicine-bound soluble tubulin from the bovine tubulin:RB3-stathmin-like domain (SLD) (T2R) colchicine complex. a) The T2R-colchicine complex is a head-to-tail longitudinal assembly of two $\alpha\beta$-tubulin heterodimers. b) $90^\circ$ rotated view of the colchicine binding site. Figure was generated using PyMOL.

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[8,9,24]. We consider below the relationship of these structures to the conformational coordinate calculated in this work. We selected 1SA0 to define the bent endpoint because (1) it has the same sequence (bovine) as the selected straight tubulin structure, (2) it has a conformation that is somewhat more ‘bent’ than the other structures, and (3) it includes colchicine, making it particularly appropriate for defining changes in the free energy profile caused by colchicine binding. These endpoint structures were processed as follows.

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These two modified endpoint structures were then energy-minimized and equilibrated using the GROMACS 4.5.5 package [26]. The GROMOS96 43a1 force field [27] was used in conjunction with the particle mesh Ewald sum method [28] to treat long-range electrostatic interactions. A time step of 2 fs was used for all simulations. Parameters for GTP and GDP were generated from the GlycoBioChem PRODRG2 server [29]. Each endpoint structure was solvated in explicit SPC water molecules [30] in the center of a periodic cubic box (dimensions of 125 Å×125 Å×125 Å) and neutralized with NaCl. The GROMACS 4.5.5 g_mindist module was used to ensure that there was no overlap between the system with its periodic image during equilibration. Each system was energy-minimized using steepest descent while applying Ferguson’s flexible SPC water model [30] constraints on the solvent. Periodic boundary conditions were applied for each system throughout all minimization and equilibration steps. Equilibration was performed in two stages: first, the system was heated to 300 K, through use of velocity rescaling, in the isothermal-isotropic (NVT) ensemble while applying position restraints on the entire system. Each system had attained 300 K by 100 ps, at which point the structure was extracted and further equilibrated in the isothermal-isobaric (NPT) ensemble ([300 K, 1.013 bar] for up to 500 ps using the Parrinello-Rahman barostat [31] implemented in GROMACS 4.5.5.

Laterally-bound tubulin pairs. For the straight pair endpoint, two straight tubulins laterally paired in the α- and β-subunit manner were taken from Wells et al.’s cryo-EM-derived “N” model of the MT lattice [23], of which the atomic coordinates were obtained through rigid fitting to Downing et al.’s 8 Å density map of a 13-protifilament MT [32]. To ensure that the interdimer contacts represent those stabilizing α- and β-lateral contacts, Wells et al. performed a multi-stage flexible fitting and equilibration procedure on the “N” tubulin pair totaling 85 ns. For the bent pair endpoint, we preserved these interdimer lateral contacts when globally aligning the unpolymerized bent tubulins to the positions of the laterally-bound straight tubulins. Both laterally-bound tubulin structures were solvated with explicit SPC water molecules in the center of a periodic box (dimensions of 300 Å×300 Å×300 Å). Energy minimization and equilibration procedures were performed as for the unpolymerized tubulin described previously; the duration of the NVT and NPT equilibration phases were 1 ns and 3 ns, respectively.

Colchicine-bound tubulin. The α1β1 heterodimer from the T2R-colchicine complex was used as the bent endpoint, preserving the colchicine molecule. Solvation, minimization, and equilibration of this structure were performed in analogous fashion as for the unpolymerized bent tubulin. The GlycoBioChem PRODRG2 server [29] was used to generate parameters for colchicine. To create a (hypothetical) structure of colchicine-bound straight tubulin, we used a stepwise procedure involving successive iterations of global alignment of tubulin and superimposition of colchicine for tubulin structures intermediate between the bent to straight endpoints, as described in greater detail in “Generating the morphing path”.

Selecting the reaction path

We chose the path collective variable, or reaction path, to be the angle required for superimposing the secondary structural elements of the α- and β-subunit intermediate domains (helices H6, H7, H8, H9, H10 and strands S7, S8, S9, and S10) of the straight and bent tubulin molecules. The conformational space R relative to this path can be expressed in terms of two collective variables, sR, which represents the progress of the dynamics along the conformational change coordinate, and z/R, which represents the progress away from the reaction path, as follows [33–36]:

\[
s(R) = \frac{1}{P-1} \sum_{i=1}^{P} (i-1) e^{-\beta |R-R(i)|^2} \tag{1}
\]

\[
z(R) = \frac{1}{\lambda} R \left( \sum_{i=1}^{P} e^{-\beta |R-R(i)|^2} \right)^{-2} \tag{2}
\]

where \(i\) is a discrete index ranging from 1 to \(P\), the total number of structures along the reaction coordinate, \(\{R-R(i)\}\) is the mean squared displacement of the 167 Cα atoms of the α- and β-intermediate domain secondary structural elements, and \(\lambda\) is a prefactor in the exponential term that defines both \(s(R)\) and \(z(R)\). (For the laterally-paired tubulins, \(\{R-R(i)\}\) includes the two sets of 167 Cα atoms in both dimers.) The value of \(\lambda\) is chosen to be proportional to the inverse of the mean square displacement between two successive frames along the reaction coordinate; here we set \(\lambda = 2.3 \left( \frac{1}{P} \sum_{i=1}^{P} |R-R(i)|^2 \right)^{-1} \). It is important to emphasize that umbrella sampling along this defined reaction coordinate only allows us to explore the free energy differences for tubulin conformations linearly interpolated along the defined reaction path; this method does not allow us to explore bent tubulin states whose α- and β-subunit intermediate domain rotation deviates substantially from the defined reaction path.

Generating the morphing path

Unpolymerized tubulin and laterally-paired tubulins. The thermally equilibrated straight and bent tubulin endpoint structures were submitted to the Molmov morphing server [37] to generate thirteen equidistantly-spaced, energy-minimized intermediate apo tubulin structures along a linear path. These thirteen apo structures were then liganded with GTP, GDP, and the Mg\(^{2+}\) ion as follows: 1) each of the two endpoint structures was globally aligned with its nearest apo intermediate; 2) GTP, GDP, and the Mg\(^{2+}\) ion were superimposed onto the nearest intermediate structure, which was then solvated, minimized, and equilibrated using the methods described in the previous section. This cycle was iterated from both endpoints for each successive tubulin intermediate. The final snapshots of the fifteen equilibrated tubulin configurations (straight, fully bent, and thirteen intermediate bent structures) were used as the initial \(P=15\) umbrella nodes along the reaction path for the calculating the potential of mean force for unpolymerized tubulin (Figure S2a in Text S1). This procedure was executed in an analogous fashion to produce the final snapshots for the thirteen equilibrated laterally-bound tubulin pair configurations (straight pair, fully bent pair, eleven intermediate bent pairs) for the initial \(P=13\) umbrella nodes in subsequent potential of mean force calculations (Figure S2c in Text S1).

Colchicine-bound tubulin. The absence of a colchicine-bound straight tubulin structure required us to use the equilibrated straight endpoint structure, along with the equilibrated colchicine-bound bent endpoint structure, to generate the colchicine-bound tubulin intermediates. These two structures were submitted to Molmov [37] to produce seven equidistantly-spaced, energy-minimized intermediate apo tubulin structures. These seven apo structures were complexed with colchicine, GTP, GDP, and the Mg\(^{2+}\) ion as follows: 1) the equilibrated colchicine-bound fully bent tubulin complex was globally aligned with its nearest apo
intermediate; 2) colchicine, GTP, GDP, and Mg$^{2+}$ from the equilibrated colchicine-bound fully bent tubulin were then superimposed onto the nearest apo intermediate; and 3) solvation, minimization, and equilibration were performed on the successive *hobo* tubulin intermediate as described above. This cycle was repeated for each successive intermediate from the bent to straight endpoints until the colchicine-bound straight tubulin had been fully equilibrated. The final snapshots for the nine equilibrated colchicine-bound tubulin configurations (straight, fully bent, and seven partially bent intermediates) were used as the initial $P=9$ umbrella nodes along the reaction coordinate for calculating the potential of mean force for colchicine-bound tubulin (Figure S2b in Text S1).

### Umbrella sampling, potential of mean force simulations

Harmonic biasing potentials are placed at each umbrella position along a collective variable. The harmonic biasing potentials for $s(R)$ and $z(R)$ are expressed as:

$$V_s(R) = \frac{1}{2} \kappa_s |s(R) - s(R_\text{i})|^2$$

$$V_z(R) = \frac{1}{2} \kappa_z |z(R) - z(R_\text{i})|^2$$

where $\kappa_s$ and $\kappa_z$ are the force constants for the harmonic restraints for $s$ and $z$, and $R_\text{i}$ is each umbrella position $i$ within the configurational space $R$.

For the unpolymerized tubulin, we initially defined $P=15$ equidistantly-spaced umbrella windows along the $s(R)$ coordinate. However, this discretization proved later insufficient to characterize the free energy along $s(R)$, so we increased the number of umbrella windows to $P=35$ while restraining each umbrella node by $\kappa_s = 10$ kcal/mol. To sufficiently characterize the free energy landscape for the laterally-bound tubulin pair, we increased the initial $P=13$ equidistantly-spaced umbrella windows to $P=50$ (Figure S3b in Text S1) while applying restraints of $\kappa_s = 10$ kcal/mol. Because the domain rearrangements characterizing the straight-to-bent conformational change are fully captured by the $s(R)$ collective variable, we did not place any harmonic restraints on the $z(R)$ path for the unpolymerized tubulin and laterally-paired tubulin dimers.

We used the initial nine colchicine-bound tubulin morphing intermediates to define $P=9$ umbrella windows along $s(R)$. To achieve adequate sampling among successive umbrella windows along $s(R)$, we optimized $\kappa_s$ to 10 kcal/mol for a total number of $P=31$ nodes. To prevent colchicine unbinding, especially from the conformations with lower intradimer curvature ($0^\circ \leq \theta \leq 6^\circ$), we restrained the dynamics along the $z(R)$ coordinate to $\kappa_z = 10$ kcal/mol.

All simulations for the unpolymerized tubulin, laterally-paired tubulins, and colchicine-bound tubulin were performed in the canonical (NVT) ensemble at 1.013 bar and 300 K for 4 ns using the PLUMED 1.3 [38]-implemented version of GROMACS 4.5.5. The last 3 ns of the production runs were used for calculating the free energy profiles.

### Generating free energy profiles using WHAM

The weighted histogram analysis (WHAM) [39,40] approach was used to merge the data from the molecular dynamics simulations and unbiased the umbrella histograms in $s(R)$. For the unpolymerized tubulin and laterally-paired tubulins, we used the *g_chem* module [41] of GROMACS 4.5.5 to calculate the free energy profiles along the $s(R)$ path, based on Eqn. 1, using a resolution of 1000 bins and tolerance of 1e-6. We then estimated the statistical uncertainty $\sigma_{PMF}$ in the $s(R)$ path accordingly [42]:

$$\sigma_{PMF}(\xi) = \left[ N_b^2 - 1 \right]^{-1} \sum_{k=1}^{N_b} \left( W_{h,b}(\xi) - \langle W_{h,b}(\xi) \rangle \right)^2$$

where $\xi$ is the reaction coordinate, $W_{h,b}(\xi)$ is the average of the $N_b$ bootstrapped free energy profiles, and $\langle W_{h,b}(\xi) \rangle$ is the average of the $N_b$ bootstrapped free energy profiles. For each umbrella position $\xi_0$, we generated a new bootstrapped trajectory $\xi_{b,i}(\theta)$ yielding a new histogram $h_{b,i}(\xi)$. WHAM was then executed on this new set of histograms for $N_b$ iterations to compute a bootstrapped free energy profile $W_{b,i}(\xi)$. (The *g_chem* module generates and aligns 1000 bootstrapped free energy profiles $W_{b,i}(\xi)$ at the initial position, $\xi_0 = 0$, in $s(R)$ so that the uncertainty at $\xi_0 = 0$ zero.) The uncertainty values along the $s(R)$ coordinate are reported for the unpolymerized tubulin and laterally-bound tubulin pair in Figures 3b and 4b, respectively.

For colchicine-bound tubulin, we calculated the two-dimensional probability distribution and free-energy surface using the *chem-2d* program (Figures S3a-b in Text S1). Results show that sampling in the $z(R)$ space remained quite close to the $s(R)$ path, even at $s(R)$ positions corresponding to lower tubulin curvature, confirming that the harmonic restraining potential $V_{s(R)}$ was sufficient to limit considerable deviations from the $s(R)$ path. Using GROMACS *g_chem*, we then projected this 2D free energy profile onto $s(R)$ at $z(R) = 0$, using 1000 bins and a tolerance of 0.001, and computed the statistical uncertainty $\sigma_{PMF}$ in the $s(R)$ path using the same bootstrapping methods described above.

### Structural analysis of tubulin conformations

#### Analyzing intradimer curvature of tubulin heterodimers.

All calculations of the intradimer rotation angles of each bent tubulin heterodimer with respect to the straight were performed using UCSF Chimera version 1.7 [43]. The Kabsch and Sander algorithm [44] implemented in UCSF Chimera was invoked to assign secondary structures for PDB structures that lacked secondary structure assignments. Each bent heterodimer was structurally superimposed onto the straight taxol-bound, zinc-induced protofilament tubulin by the $\alpha$-subunit H7 helices. The structural alignment was performed in 3 different ways, using either all atoms, only backbone atoms, or only Ca backbone atoms of the $\alpha$-H7 helix (see Supplemental Table S3 in Text S1 for exact residues). The intradimer rotation was then defined by fitting a plane to the $\alpha$- and $\beta$-subunit H7 helices using least-squares fitting. The intersection angle of the two vector planes represents the intradimer rotation of each bent heterodimer.

To compute buried surface area, the crystal structures were first pre-processed so that each heterodimer is only complexed with GDP and GTP-Mg$^{2+}$; any other protein chains (i.e. RB3-SLD, TTL, D1), solvent molecules, and counter ions other than Mg$^{2+}$ were also removed. The solvent-accessible surface area of each tubulin heterodimer ($\text{SASA}_{\text{ab}}$) and the solvent-accessible surface area of each $\alpha$- and $\beta$-subunit, $\text{SASA}_{\text{a}}$ and $\text{SASA}_{\text{b}}$, excluding the other subunit, were calculated in PyMOL using a solvent probe radius of 1.2 Å and a solvent density of 4. The buried surface area of each tubulin heterodimer was determined accordingly: $\text{BSA} = (\text{SASA}_{\text{a}} + \text{SASA}_{\text{b}}) - \text{SASA}_{\text{ab}}$.

#### Calculating volume of colchicine binding pocket.

For each of the 31 umbrella positions along the conformational change coordinate, five colchicine-bound tubulin structures corresponding to the peak of each histogram were extracted from the last 3 ns of...
Figure 3. The (a) probability distribution, (b) free energy profile, and (c) buried surface area of unpolymerized tubulin as a function of intradimer curvature. Intradimer rotation of the endpoint structures, the straight taxol-liganded, zinc-stabilized protofilament tubulin and bent \( \alpha_1\beta_1 \) heterodimer from the T2R-colchicine complex, are denoted at \( \theta = 0^\circ \) (blue dash) and \( \theta = 12^\circ \) (red dash), respectively. Intradimer rotation of selected intermediately bent tubulins are denoted as dotted lines: \( \theta = 5^\circ \) (GMPCPP-tubulin in double-tube layers), \( \theta = 6^\circ \) (\( \alpha_1\beta_1 \) heterodimer of the T2R-TTL-ADP, T2R-TTL-zampanolide, and T2R-TTL complexes), \( \theta = 6.9^\circ \) (both \( \alpha_1\beta_1 \) and \( \alpha_2\beta_2 \) heterodimers of the GTP-tubulin-D1 DARPin structure), and \( \theta = 10^\circ \) (\( \alpha_1\beta_1 \) heterodimer of T2R-TTL-zampanolide and T2R-TTL epothilone A complexes). Buried surface area values of tubulin heterodimer structures with resolutions up to 3 Å are shown in panel (c); the structures from PMF in (b) (open circles) and crystal structures of the straight (solid triangle) and bent (solid circles) heterodimers are denoted for each tubulin complex.

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the restrained molecular dynamics trajectories. The SiteMap module [45] in the Schrödinger 2012 suite of programs was used to calculate the volume of the colchicine binding pocket, using the OPLS 2005 force field at a 0.7-Å grid resolution and setting a 6 Å buffer region surrounding colchicine. The SiteScore for each colchicine-tubulin complex was confirmed to be greater than 1. For each of the 31 umbrella positions, the arithmetic mean and standard deviation of the colchicine volume \( V_{\text{colc}} \) and binding pocket volume \( V_{\text{pocket}} \) of the five representative structures were calculated.

We also analyzed the hydrogen bonding, hydrophobic interactions, and steric clashes of the colchicine-tubulin complexes at all 31 umbrella positions. For each of the five representative structures from each umbrella position, we used the Maestro command-line “Display Hydrophobic Interactions” script (to identify the atoms of colchicine and tubulin involved in hydrogen bonding and/or hydrophobic interactions. Steric clashes are defined as two atoms separated by a distance that is less than 75% of the sum of their van der Waals radii.

Results/Discussion

Unpolymerized tubulin primarily adopts an intermediately bent conformation

Free energy profile of unpolymerized tubulin. Figure 3 shows the probability distribution, free energy profile, and buried surface area computed from the potential-of-mean force calculations for unpolymerized tubulin (i.e., a single \( \alpha\beta\)-heterodimer). We had hypothesized that the PMF would show two prominent basins, reflecting the straight and bent conformations of tubulin. This hypothesis was only partly confirmed. The PMF does show a free energy basin corresponding to the straight conformation, as defined by 1JFF. This basin is separated by a relatively small free energy barrier at \( \sim 3^\circ \) intradimer rotation from a broad basin that includes, at one end, the bent conformation corresponding to the colchicine-bound structure (1SA0). Integrating the populations, the straight basin \( (\theta < 3.5^\circ) \) accounts for approximately 20% and the various bent states approximately 80%. Although the 1SA0 structure has an intra-dimer rotation that corresponds rather precisely with a local minimum in the PMF, the global minimum in the PMF has an intradimer rotation of \( \sim 6^\circ \). However, we note that the free energy difference between this ‘intermediately bent’ global minimum and the ‘fully bent’ minimum at \( \sim 12^\circ \) is small, \(<1\text{ kcal/mol}\), and only slightly larger than the estimates of the statistical uncertainties (the true errors, which would include limitations of the force field, are of course larger). Thus, we cannot confidently conclude, on the basis of these calculations alone, that the intermediately bent state is lower in free energy than the fully bent state.

However, the broad and relatively featureless basin representing the bent states is in itself a striking prediction, suggesting that tubulin can exist in a nearly continuous range of different bent intradimer rotation angles. Soluble GDP-tubulin was previously thought to exist in only two states, the “straight” and “bent” structures represented by PDB ids 1JFF and 1SA0, respectively. Our calculated PMF results are striking because our PMF suggests a broad range of intermediately bent soluble GDP-tubulins whose curvature correspond to specific energy wells in the PMF; this relatively flat landscape is unexpected within the framework of this two-state model. The number of crystal structures of tubulin has been increasing rapidly in the past few years, and many of the newer structures in fact have intradimer rotation angles, when measured in the same manner as for the PMF, that correspond to our ‘intermediately bent’ states (see Table 1). For example, the cryo-EM structures of tubulin observed in double-layer tubes by Nogales et al. [46], several structures of tubulin in complex with tyrosine tubulin ligase, and a GTP-tubulin-D1 DARPin complex [9] all have intradimer rotation angles in the range of 5–7°, similar to the intermediately bent global free energy minimum observed computationally. The combination of the potential of mean force and the comprehensive analysis of crystal structures suggests that tubulin can adopt a wide range of different ‘bent’ states, and we hypothesize, but cannot currently prove, that tubulin in solution is likely to exist in an ensemble of different states, dominated by a variety of bent states. This hypothesis is also supported by other molecular dynamics studies by Voth and co-workers [12–14], who observed tubulin interconverting between different intradimer rotation angles on the timescale of tens of nanoseconds.

As aforementioned, our metric for calculating tubulin’s curvature focuses on the orientations of the \( \alpha \)- and \( \beta \)-H7 helices. Voth et al. accounted for tubulin’s curvature by the movement of the \( \alpha \)- and \( \beta \)-subunits with respect to one another. To reconcile these two different calculation methods, we utilized Voth’s method to measure the intrinsic bending angle of 15 equidistantly-spaced structures along the reaction coordinate for the free energy landscape of unpolymerized tubulin in Figure 3b. The results, shown in Figure S5 in Text S1, indicate that the two metrics, while not identical, correlate strongly with a slope close to 1.

Buried surface area of unpolymerized tubulin. It is noteworthy that our method of describing conformational change of tubulin is limited to the rotation of the heterodimer. Voth et al. also calculated a twist angle to account for the twisting motions of the \( \alpha \)- and \( \beta \)-subunits with respect to one another, our reaction coordinate does not account for this rotational motion. We recognize limitations in this geometric descriptor, and have also computed the buried surface area (BSA) of various intermediately
### Table 1. Intradimer curvature of “straight” and “bent” tubulin heterodimers.

| Structure                        | Resolution (Å) | PDB ID | Source Organism | Nucleotide State | Heterodimer | Intradimer Rotation (°) | α-subunit Rotation (°) | β-subunit Rotation (°) | Space Group | Ref. |
|----------------------------------|----------------|--------|-----------------|------------------|-------------|------------------------|------------------------|------------------------|-------------|------|
| **“STRAIGHT”**                   |                |        |                 |                  |             |                        |                        |                        |             |      |
| Epothilone A-tubulin*            | 2.89           | 1TVK   | B. taurus       | GDP              | αβ          | 1.9±0.1                | 2.8±1.7                | 1.5±0.6                | P21         | [5]  |
| Docetaxel-tubulin*               | 3.70           | 1TUB   | S. scrofa       | GDP              | αβ          | 4.8±3.3                | 2.3±0.4                | 1.9±0.3                | P21         | [6]  |
| **“BENT”**                       |                |        |                 |                  |             |                        |                        |                        |             |      |
| T2R-TTL-zampalonide              | 1.80           | 444T   | B. taurus       | GDP              | αβ          | 7.2±2.1                | 8.7±1.1                | 13.3±0.9               | P212121     | [10] |
| T2R-TTL-ADP                      | 2.00           | 447J   | B. taurus       | GDP              | αβ          | 9.7±1.5                | 8.7±0.8                | 14.6±0.9               | P212121     | [50] |
| T2R-TTL                          | 2.20           | 455S   | B. taurus       | GDP              | αβ          | 6.1±0.4                | 4.3±1.4                | 12.2±1.0               | P212121     | [10] |
| T2R                              | 2.20           | 3RYC   | O. aries        | GDP              | αβ          | 6.0±0.4                | 9.8±3.5                | 10.4±2.1               | P212121     | [24] |
| GTP-tubulin-D1 DARPin            | 2.22           | 4DRX   | O. aries        | GTP              | αβ          | 6.8±0.3                | 11.3±0.4               | 11.9±1.3               | P21         | [9]  |
| T2R-TTL-epothilone A             | 2.30           | 450    | B. taurus       | GDP              | αβ          | 10.0±2.4               | 8.5±1.0                | 13.8±0.9               | P212121     | [10] |
| T2R                              | 2.40           | 3RYI   | O. aries        | GDP              | αβ          | 6.1±0.4                | 6.0±1.1                | 12.2±0.7               | P212121     | [24] |
| T2R                              | 2.52           | 3RYF   | O. aries        | GTP              | αβ          | 6.3±0.3                | 2.9±0.3                | 11.0±0.5               | P212121     | [24] |
| T2R-colchicine-ustiloxin         | 2.73           | 3UTS   | O. aries        | GDP              | αβ          | 7.7±0.6                | 9.8±0.4                | 10.3±0.3               | P212121     | [24] |
| T2R                              | 2.80           | 3RH    | O. aries        | GMPCPP           | αβ          | 7.7±0.1                | 2.5±0.4                | 11.1±1.0               | P212121     | [24] |
| T2R-vinblastine                  | 3.47           | 4EB6   | O. aries        | GDP              | αβ          | 6.3±0.6                | 3.0±0.2                | 11.1±0.5               | P212121     | [24] |
| T2R-colchicine                   | 3.58           | 1SAO   | B. taurus       | GDP              | αβ          | 12.0±0.1               | 9.2±0.7                | 11.7±1.6               | P65         | [7]  |
| T2R                              | 3.65           | 3KHB   | O. aries        | GDP              | αβ          | 7.7±1.8                | 7.6±1.9                | 13.0±1.8               | P65         | [20] |
| T2R-podophyllotoxin              | 4.20           | 1SA1   | B. taurus       | GDP              | αβ          | 12.1±0.6               | 9.1±0.8                | 12.9±1.3               | P65         | [7]  |

For the RB3-SLD-bound structures, the orientations of the longitudinally-paired αβ and γβ heterodimers with respect to the MT plus (+) and minus (−) ends are shown in Figure S4 of Text S1 along with the relative orientations of the D1-(γβ)1 and D1-(γβ)2 dimers in the GTP-tubulin-D1 DARPin structures. Crystal structures are listed in order of decreasing resolution. Residues comprising the α- and β-H7 helices used for angle calculations are listed in Table S3 of Text S1. Each angle was measured using the backbone N-Cα-C-O, Cα, and all atoms, with the standard deviation shown accordingly.

*Control calculations of the “straight” tubulin were performed on the epothilone A- and docetaxel-bound, Zn2+-stabilized protofilament tubulin with respect to the taxol-bound, Zn2+-stabilized protofilament tubulin (α-H7: 224-242, β-H7: 224-243).

**Outlying values of intradimer tubulin curvature Tα-vinblastine structure are likely due to unraveling of the H7 helix.

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bent tubulin structures, as an alternative way of quantifying
differences between “straight” and “bent” structures (see
Figure 3c). These BSA values in Figure 3c track very closely to
the conventionally defined intradimer rotation angles of our
predicted intermediately bent tubulin structures. Specifically, the
interface between the alpha and beta subunits becomes much
more extensive as tubulin adopts the straight conformation
consistent with incorporation into the microtubule. This suggests
a balance of forces between the energy required to straighten the
bent dimer and the interfacial packing energy gained in the
process. Along the bending coordinate used in our potential of
mean force calculations, the buried surface area varies nearly
linearly with respect to the geometrical parameter we have used to
quantify the intradimer rotation, as might be expected if our
reaction coordinate followed a minimum energy path along the
landscape. Most of the tubulin structures with resolutions better
than 3.0 Å lie close to this line, with the exception of a few
structures in highly bent states. It should be emphasized that we
have only characterized the free energy landscape along a single
coordinate, and other relevant conformational states may exist “off
pathway”. Nonetheless, the fact that the crystal structures that we
characterized as intermediately bent based on the geometric
intradimer angle also have intermediate values of intradimer
buried surface area supports the contention that tubulin can adopt
an intermediately bent state, and a two-state bent/straight model
is thus in certain respects overly simplistic. It is noteworthy that the
various X-ray crystallographic structures represented in Figure 3c
differ in their space group and resolution.

The α- and β-subunits of our bent endotype structure have
intradimer rotations of ~8° and ~11°, respectively. To
to examine whether the rotation of the individual the α- and β-
subunits is related to the rotation of the entire heterodimer, we
calculated the rotation of each α- and β-subunit of the X-ray
crystallographic structures of tubulin shown in Table 1, as well as
for each tubulin intermediate along the reaction coordinate in
Figure 3. (These calculations were performed in accordance with
methods described in Knosov et al. [7].) As shown in Table S1 of
Text S1, the intradimer rotations of the α- and β-subunits of the
interpolated tubulin structures increase monotonically but
nonlinearly with increasing heterodimer curvature. In the X-ray
crystallographic structures of bent tubulins, the β-subunit typically
rotates between ~11–14°, whereas the α-subunit intradimer rotation
varies more widely between ~4–12°. Thus, while our
PMFs do not independently explore the intradimer and the
intradimer rotations, the structural information suggests that the
intradimer rotations in the α- and β-subunits may be partially
independent degrees of freedom.

Having characterized the surprisingly ‘flat’ free energy profile
for tubulin along a bending coordinate, we next examined how
this potential of mean force is modulated by (1) lateral association
of two tubulin heterodimers (with implications for the lattice model
of microtubule assembly), and (2) binding a small molecule drug,
colchicine (with implications for microtubule-binding therapeu-
tics).

**Lateral association of tubulin heterodimers favors the
straight state**

Figure 4 shows the free energy profile and probability
distribution for the laterally-paired tubulins. The region of the
conformational change coordinate ranging from θ = 10° to θ = 12°
was not successfully sampled because the lateral interdimer
contacts were not fully preserved during the NPT equilibration of
the lateral dimers corresponding to this range of curvature,
despite significant effort. These results suggest that unpolymerized
tubulin with >10° intradimer rotation is physically incapable of
forming both α-α and β-β lateral interactions, unless it transitions
into a lower curvature state.

The 1SA0 α,β heterodimer from the T2R-colchicine:tubulin
complex exhibits a ~3° twist angle, which is somewhat large
compared to the degree of twisting exhibited by the newer X-ray
crystallographic structures of tubulins, whose twist angles range
from ~5–7° (see Tables S2a–b in Text S1). We calculated the twist
angles of both heterodimers of thirteen laterally-paired tubulin
structures and found that, in general, they vary from 2–5°, which
are intermediate between the twist angles of the newer crystal
structures of bent tubulin, and the twist angle in the straight (taxol-
bound) structure. Therefore, because our structures of laterally-
paired tubulins along the conformational change coordinate do
not exhibit the extreme twisting motion prevalent in the 1SA0
α,β heterodimer, the lateral contacts in our interpolated
intermediate structures of laterally-paired tubulins should be
stable.

As aforementioned, the role of nucleotide state in affecting
tubulin conformation has been intensely debated in the context of
the lattice and allosteric models of microtubule assembly. The
lattice model postulates that soluble tubulin undergoes conforma-
tional changes only upon incorporation into the microtubule
lattice and independently of nucleotide state, whereas the allosteric
model posits that GTP binding pre-structures soluble tubulin into
the “straight” conformation compatible with that in the MT
lattice. Our PMF of the laterally-paired tubulins suggests that
lateral binding of tubulin heterodimers substantially modifies the
free energy profile, consistent with the lattice model. Consistent
with our expectations, based on the lattice model, lateral binding
strongly shifts the population toward the straight conformation
(~75%), which again is separated from the intermediately bent
conformation by a small free energy barrier (~2 kcal/mol), in this
case at ~4°. The global free energy minimum along the PMF has
an intradimer rotation angle (0.5°) very close to that observed in
structures of zinc-induced protofilament sheets (1JFF). Although
we find it geometrically infeasible for both tubulins to be in a fully
bent state, both can exist in an intermediately bent state when
laterally associated, and this state represents ~25% of the
population. (The two tubulins are constrained to maintain the
same intradimer angle across the PMF, i.e., we have not
investigated laterally associated straight-bent pairs, although
generically we expect these to be unfavorable.) Overall,
comparison of the ‘free’ and laterally-paired tubulin PMFs suggests
that lateral association shifts the bending equilibrium by
~1.5 kcal/mol toward the straight conformation, providing direct
(computational) support for the lattice model of microtubule
assembly.

Specifically, at the earliest stages of tubulin assembly, the lattice
model predicts that lateral contacts will be disadvantaged because
the binding interface is disrupted in the bent conformation. These
results suggest that even forming a single lateral pair is sufficient
to largely shift the equilibrium toward the straight conformation,
which then makes subsequent lateral additions more thermody-
namically favorable.

**Implications of conformational changes in tubulin on MT
assembly.** To begin to explore the functional consequences of an
assembly-dependent conformational change in tubulin that
underlies the lattice model, Rice et al. had used a simple two-step
nucleation-elongation model to approximate the kinetics of the
early stages of MT assembly. In keeping with experimental data,
kinetic parameters were set to capture the low affinity of subunit
addition during nucleation and a high affinity of subunit addition
during elongation: the monomer concentration was set to 10 μM,
the bimolecular on-rate constant to $k_0 = 1 \times 10^6 \text{M}^{-1}\text{s}^{-1}$. Reverse rate constants for nucleation ($k_{\text{nuc}}^d$) and elongation ($k_{\text{Elon}}^d$) such were set such that their respective dissociation constants would be $K_d^{\text{nuc}} = 1 \text{mM}$ and $K_d^{\text{Elon}} = 1 \text{mM}$, respectively, and that the essence of the nucleation-elongation would be captured. The kinetic equations characterizing MT assembly of a nucleus size of $N$ species (in which each species is represented by a heterodimer) have been described. Using these kinetic parameters and a nucleus size of $N$ heterodimers, Rice et al. generated 500-second polymerization profiles for a range of straightening penalties from 0 to 1.6 $k_B T$ ($\sim 1\text{kcal/mol}$). Using our data that the ensemble of bent states are $\sim 1\text{kcal/mol}$ more energetically favorable than the straight conformer, Rice et al.’s model suggests that a $\sim 1\text{kcal/mol}$ straightening penalty increases the time required to assemble into a 4-species nucleus by $\sim 10$-fold. That a modest straightening penalty of 1 kcal/mol is sufficient to decrease the rate of MT assembly highlights the sensitivity of MT assembly kinetics to the initial stages of nucleation and elongation.

**Colchicine binding strongly shifts the population of soluble tubulin toward bent structures.**

**Free energy profile of colchicine-bound tubulin.** Figure 5 shows the calculated free energy profile and probability distribution of the colchicine-bound tubulin complex as a function of the intradimer rotation. The estimated statistical error in the relative free energy profile ranges between 0.3–0.4 kcal/mol, confirming adequate umbrella sampling and overlap throughout all umbrella windows along the reaction coordinate, as shown in Figure S2b in Text S1. The restraints used in generating the PMF prevented colchicine from dissociating from tubulin. As expected, colchicine binding strongly disfavors the straight conformation, and the free energy decreases in a nearly linear manner, with no significant free energy barriers, as the intradimer rotation angle increases, up to the global free energy minimum observed at $\sim 9^\circ$. These results are consistent with previous findings that colchicine binds only to the bent conformer, thus inhibiting tubulin assembly into the microtubule lattice [7,11,20,21]. The intradimer rotation angles observed in crystal structures of colchicine-bound tubulin range from $\sim 8^\circ$–$12^\circ$ (Table I), in good agreement with the PMF. It should be noted that the T2R-colchicine-ustiloxin structure is higher in resolution (2.7 Å) than the T2R-colchicine structure (3.6 Å), which we used to define the most highly bent end-point structure for the PMF. The two structures also have different space groups.

Figure 6 depicts changes in the colchicine-binding pocket as a function of the tubulin intradimer rotation angle. One reason for colchicine strongly disfavoring the straight conformation is simply that the binding pocket becomes much smaller ($235 \text{Å}^3 \pm 29 \text{Å}^3$) than for fully bent tubulin ($937 \text{Å}^3 \pm 31 \text{Å}^3$ at $\theta = 12^\circ$). The pocket cannot collapse entirely in these simulations because colchicine is restrained to remain in the binding pocket. The colchicine-bound tubulin structures with $\sim 8^\circ$ intradimer curvature also appear to be stabilized by hydrogen bonds with the backbone amide of $\alpha$-Val 181 and side chain amine of $\beta$-Lys 350 (Figure S6 in Text S1), in agreement with previous research that Val 181 is essential for colchicine’s biological activity [47].

The results presented here, predicting a continuum of intermediately bent tubulin states ranging from “straight” to “bent”, has potential implications for microtubule-directed drug discovery. In principle, it should be possible to identify compounds that can bind selectively to various “intermediately bent” states, as opposed to the strongly bent state stabilized by colchicine, or the straight state presumably stabilized by taxol. More broadly, our results suggest the potential utility of molecular dynamics methods (including more advanced methods like Markov State Models, which are not used here) to understanding how drugs or drug candidates modulate the free energy landscape of the target proteins. We expect this type of approach to be particularly relevant to understanding and ultimately designing allosteric modulators, and more broadly to how the chemical structure and binding mode of a drug dictates the ways in which it modulates the energy landscape.”

**Conclusions.** Structural and functional studies have previously demonstrated the existence of the straight-bent equilibrium for tubulin, and its relevance to microtubule assembly. Our new contribution here is to compute the free energy along a conformational coordinate connecting two limiting straight and bent structures. We had hypothesized that we would observe two free energy basins, corresponding to the straight and bent states, with a free energy barrier separating these, and that the bent state would correspond to the lowest (most favorable) free energy, consistent with the lattice model of microtubule assembly. The computed “potential of mean force” along the chosen bending coordinate largely supports this hypothesis, but also suggests that the bent state is best understood as an ensemble of states with a significant range of intradimer rotation angles (4–15°). The global free energy minimum in the PMF lies at 6° intradimer rotation, whereas the colchicine-bound structure that we (and others) have used to define the “fully bent” state lies at 12°, and is slightly higher.
in free energy. Our colchicine simulations also support the observation that colchicine biases towards conformations more bent than the $6^\circ$ rotation angle calculated for free tubulin. Recent structures of tubulin co-crystallized with various small molecules and macromolecules support the prediction that tubulin can adopt a wide range of bent states.

Overall, the PMF predicts that the ensemble of bent states, taken as a whole, has a free energy that is lower than that of the straight state by about 1 kcal/mol, a modest value that is nonetheless sufficient to have a significant impact on polymerization kinetics, as represented (crudely) by a simple nucleation-elongation model. We emphasize that the precise value should not

**Figure 6. The colchicine binding pocket is sterically constrained in tubulin conformations with lower curvature.** a) Ratios of the volume of colchicine with respect to the volume of the predicted binding pocket for representative colchicine-bound tubulin heterodimers with curvatures of $0^\circ$, $2^\circ$, $6^\circ$, $8^\circ$, $10^\circ$, and $12^\circ$. b) SiteMap predictions of the colchicine binding site for colchicine-bound tubulin structures with intradimer curvature of $0^\circ$, $2^\circ$, $6^\circ$, $8^\circ$, $10^\circ$, and $12^\circ$. For each colchicine-bound tubulin structure, the representative structure with the maximum SiteScore is displayed. Black dots represent the predicted binding pocket of colchicine. Structural elements comprising the colchicine-binding domain are rendered as cartoon. doi:10.1371/journal.pcbi.1003464.g006
be over-interpreted, because the statistical uncertainty estimated for the PMF is \(\sim 0.5 \text{kcal/mol}\); in addition, the underlying molecular mechanics force field model also has limitations that could lead to systematic errors, the magnitude of which are difficult to estimate. An additional limitation is that the PMF only probes one linear coordinate connecting two structures that we chose as the extreme straight and bent states. There could exist other ‘off-pathway’ states, involving conformational changes orthogonal to and independent of the coordinate probed here, which could also have functional relevance. It is also highly likely that tubulin is capable of adopting bent conformations whose curvature is in the direction opposite of that of 1SA0, as explored by Voth and co-workers [13], and other indirectly bent tubulins in the continuum of structures predicted in our PMF. While unrestrained MD simulations do not have this limitation [12,14,18], it is extremely difficult from them to converge, i.e., to adequately sample the full conformational space. In future work, Markov State Models could potentially be used to more exhaustively characterize the free energy landscape, albeit with substantial computational expense [48].

Nonetheless, the qualitative predictions of the PMFs agree well with a variety of experimental observations, including the fact that the intradimer rotation and surface area buried between the \(\alpha\) and \(\beta\) chains in experimental structures correspond well with low free energy portions of the PMF. We also demonstrate that tubulin binding to the drug colchicine, and binding to another tubulin in a lateral interaction, both perturb the PMF in ways consistent with expectations. Colchicine binding strongly biases the equilibrium toward the bent state, to the extent that the straight state is thermodynamically infeasible; we attribute this shift largely to simple steric considerations, where the binding pocket shrinks as tubulin straightens. Lateral tubulin binding shifts the conformational equilibrium in the opposite manner, such that the most extreme bent states become infeasible and the straight conformation corresponds to the global free energy minimum (although indirectly bent states may still be populated). This result is consistent with the lattice model of assembly, and additionally suggests that a single lateral interaction is sufficient to “pay” the free energy cost of straightening tubulin.

Our PMFs of unpolymerized tubulin and laterally-paired tubulins are based on tubulins bound with GDP at the \(\beta\)-subunit E-site. We had chosen to use GDP-bound tubulin because these were the only X-ray crystallographic structures available at the commencement of this study, and it would have been risky to artificially build in GTP. We have not considered in this work the role of nucleotide state at this site, a factor that has been considered in some prior MD simulations. Although SANS experiments reveal that GTP- and GDP-bound tubulin adopt very similar, bent states in solution and that nucleotide choice does not influence colchicine binding, we cannot rule out the possibility that nucleotide state could modulate the PMF in some way, e.g., GTP shifting the equilibrium of the M-loop in a way that might additional favor lateral interactions. In principle, free energy simulations (PMFs or alchemical changes) could be used to address this issue in future work.

Supporting Information

Text S1 Supporting Information. This section documents the supplementary figures and tables referenced throughout the text. (DOCX)

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Author Contributions

Conceived and designed the experiments: LXP MTH DAA MPJ. Performed the experiments: LXP MTH. Analyzed the data: LXP MTH MB. Contributed reagents/materials/analysis tools: MTH. Wrote the paper: LXP MTH MB DAA MPJ.

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