

Design and Characterization of Modular Scaffolds for Tubulin Assembly^{*[5]}

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Background: Stathmin-like domain (SLD) proteins from vertebrates bind two tubulin molecules.

Results: SLDs that bind tubulin with a programmed stoichiometry are characterized.

Conclusion: Rules are established to design (tubulin)_x-SLD complexes, starting from a 1:1 stoichiometry.

Significance: This work provides new insights into stathmin family member function. The SLDs produced will be useful tools to study interactions of microtubule regulators with tubulin.

In cells, microtubule dynamics is regulated by stabilizing and destabilizing factors. Whereas proteins in both categories have been identified, their mechanism of action is rarely understood at the molecular level. This is due in part to the difficulties faced in structural approaches to obtain atomic models when tubulin is involved. Here, we design and characterize new stathmin-like domain (SLD) proteins that sequester tubulins in numbers different from two, the number of tubulins bound by stathmin or by the SLD of RB3, two stathmin family members that have been extensively studied. We established rules for the design of tight tubulin-SLD assemblies and applied them to complexes containing one to four tubulin heterodimers. Biochemical and structural experiments showed that the engineered SLDs behaved as expected. The new SLDs will be tools for structural studies of microtubule regulation. The larger complexes will be useful for cryo-electron microscopy, whereas crystallography or nuclear magnetic resonance will benefit from the 1:1 tubulin-SLD assembly. Finally, our results provide new insight into SLD function, suggesting that a major effect of these phosphorylatable proteins is the programmed release of sequestered tubulin for microtubule assembly at the specific cellular locations of members of the stathmin family.

Microtubules are dynamic protein assemblies essential for cell morphogenesis, membrane trafficking, and cell division of eukaryotic cells. *In vivo*, typically 13 straight, parallel, protofilaments interact laterally to form a microtubule. Each protofilament is a

longitudinal head-to-tail assembly of $\alpha\beta$ tubulin heterodimers (hereafter referred to as tubulins). *In vitro* experiments with purified tubulin have demonstrated that microtubules switch stochastically between prolonged periods of assembly and disassembly, a phenomenon called dynamic instability (1). *In vivo*, microtubule dynamics is regulated by different classes of proteins. These include polymerases, depolymerases, microtubule stabilizing, and tubulin sequestering proteins (2–5).

The complex events underlying the regulation of microtubule assembly are difficult to observe structurally. This has been achieved on a few occasions, when sufficiently homogeneous samples were obtained. Two strategies have been used. The first exploited the properties of tubulin sequestering proteins to produce homogeneous complexes whose crystal structure has been determined (6, 7). Alternatively, tubulin assemblies have been analyzed using cutting edge transmission electron microscopy (TEM).³ These studies focused on microtubules (8), microtubules decorated with globular proteins (9–11), and a few non-microtubular tubulin species (*e.g.* Ref. 12). But, in most cases, due to the heterogeneity of the assemblies present in solutions of tubulin and of its complexes, obtaining crystals that diffract to atomic resolution remains challenging. Moreover, because of the limitations of the lifetime of the sample in the electron beam (13) and because extensive averaging of images of identical species is not possible, the study of such heterogeneous assemblies by cryo-TEM is also restricted to low resolutions that hardly go beyond the dimensions of globular domains. The availability of new stable and well defined tubulin complexes, including single sequestered heterodimers, would offer new options for crystallization or allow TEM images to be collected that could then be averaged. This would therefore greatly facilitate the study of tubulin assembly regulation structurally and also biochemically.

Stathmin and stathmin-like domains (SLDs) prevent the formation of microtubules (5, 14). The SLDs from vertebrates have

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³ The abbreviations used are: TEM, transmission electron microscopy; DARPin, designed ankyrin repeat protein; SEC-MALLS, size exclusion chromatography coupled to multi-angle laser light scattering; SLD, stathmin-like domain.

been best studied; they bind two tubulins arranged longitudinally, head-to-tail, in protofilament-like complexes (see Fig. 1A) (15, 16). These complexes are homogeneous and stable, but their size (~200 kDa) is at the lower end of the range of species that may be studied at high resolution by cryo-TEM (17). SLDs from *Drosophila* can bind up to four tubulins, in a dynamic association (18). No SLD has been identified that sequesters efficiently a single tubulin, although several attempts at designing such proteins have been made (19, 20). Because vertebrate SLDs allow the binding of other regulatory proteins to their complexes with tubulin (21), they appear to be a useful starting point for the development of stable, well defined, assemblies of tubulin that could be used to study the regulation of microtubule assembly, both biochemically and structurally, including by electron microscopy. But to do so, stable complexes comprising three or four heterodimers should be engineered to be of a size large enough for this methodology to be conveniently applied. The smaller version of these complexes, comprising one tubulin, would extend the range of tubulin complexes that may be crystallized for higher resolution studies beyond T₂R, the ternary complex of two tubulin heterodimers with the SLD of the RB3 protein (RB3_{SLD}). Such platforms will provide stable entities to which regulatory proteins may bind. They may also be used to study the interaction with tubulin of small molecule compounds (6).

Here, we describe the design of SLD-based proteins that make stable complexes with tubulin. The structural characterization of a complex comprising four tubulin heterodimers demonstrates that it consists of longitudinally assembled molecules that have the same overall structure as tubulin in T₂R, strongly suggesting that this applies to all SLD-mediated tubulin assemblies. Moreover, the high resolution structure of a single sequestered tubulin shows that its interactions with the SLD are identical to those in T₂R, thus validating the rules we established for the design of SLDs binding a predefined number of heterodimers assembled longitudinally.

EXPERIMENTAL PROCEDURES

Gene Synthesis, Cloning, Protein Overexpression, and Purification—R4 and R4a genes were purchased from Genscript (Piscataway, NJ). R3 was synthesized according to the method of Stemmer *et al.* (22). R1 was obtained from a plasmid coding for an RB3_{SLD} variant by a modified overlap extension PCR method (23). Its sequence is displayed in Fig. 1. All these constructs have been cloned between the NcoI and XhoI sites in a pET28 plasmid carrying a kanamycin resistance gene and a promoter inducible by isopropyl β-D-1-thiogalactopyranoside. Proteins were overexpressed in *Escherichia coli* BL21 DE3 Star, in LB medium supplemented with kanamycin, using 0.5 mM isopropyl β-D-1-thiogalactopyranoside to induce an expression period of 3 h at 37 °C. Purification was as described (6) except that a first step of nucleic acid precipitation by spermine (24) was added and that the heating step was omitted for R4 and R4a. The concentration of purified SLD was determined by measuring the absorbance at 280 nm, taking advantage of the presence of tryptophan residues in these constructs, as opposed to wild type RB3_{SLD}, whose absorbance at 280 nm is very weak. A

mass spectrometry analysis of R4 showed it has the expected molecular mass, taking into account the removal of the N-terminal methionine and a subsequent Nα-acetylation as is the case for RB3_{SLD} (25). Tubulin was purified from sheep brain by two cycles of assembly-disassembly in a high molarity Pipes buffer (26). Before use, an additional cycle of assembly-disassembly was performed to remove inactive protein. The designed ankyrin repeat protein (DARPin) used in this study and named D2 was selected in the same screen as the D1 DARPin and was produced and purified similarly (27).

Size Exclusion Chromatography (SEC) and Size Exclusion Chromatography Coupled to Multiangle Laser Light Scattering (SEC-MALLS)—For SEC experiments, samples of tubulin alone or mixed with RB3_{SLD}, R1, R3, or R4 were analyzed on a Superdex S200 column (GE Healthcare) equilibrated with 15 mM Pipes-K, pH 6.8, 0.4 mM MgCl₂, and 0.2 mM EGTA (low salt buffer) or with the same buffer containing in addition 60 mM KCl (higher salt buffer). For SEC-MALLS analyses, SEC was carried out on a Prominence HPLC system (Shimadzu) using a KW804 column (Shodex) run in the higher salt buffer. 30-μl samples at 20 or 40 μM tubulin concentrations and containing variable amounts of SLDs were run at a 0.5 ml/min flow rate. Detection was performed using a three-detector static light-scattering apparatus (MiniDAWN TREOS, Wyatt Technology, equipped with a quasi-elastic light-scattering module) and a refractometer (Optilab rEX, Wyatt Technology). Molecular weight calculations were performed with the ASTRA V software (Wyatt Technology) using a dn/dc value of 0.183 ml/g.

Electron Microscopy (EM)—A 5-μl (0.15 mg/ml) tubulin-R4 sample containing Protein A conjugated with 5-nm colloidal gold particles (Cell Microscopy Center, University Medical Center, Utrecht, The Netherlands) as fiducial markers was applied to a glow-discharged carbon-coated Cu 200-mesh grid, allowed to adsorb for 30 s, washed twice with water, and negatively stained for 20 s with 0.5% (w/v) uranyl formate. Specimens were examined in a Fei Tecnai 20 TEM operated at 200 kV. Double tilt tomograms were recorded with a magnification at the detector plane of ×69,000 with a Gatan CCD camera through Inspect3D with the specimen supported by a Fishione double tilt holder. The sample was tilted from -62 to +62° in 0.5° steps. Tomographic reconstructions were made with IMOD (28). The Slicer tool of IMOD was used to combine multiple slices in the beam direction.

Kinetic Analysis of the Tubulin-R1 Interaction—An R71C mutant of R1 was obtained by standard molecular biology techniques and produced and purified as the wild type protein. It was reduced with dithiothreitol and then reacted with acrylodan. Excess acrylodan and protein aggregates were removed by gel filtration on a Superdex S200 column. The resulting protein, named R1*, was used for fluorescence studies (λ_{ex}, 290 nm; λ_{em}, 510 nm). The dissociation constant (K_d) was determined using a fixed concentration of R1* titrated against an increasing amount of tubulin at 20 °C in a Cary Eclipse spectrofluorimeter (Varian). The buffer used consisted of 25 mM Pipes-K, pH 6.8, 0.2 mM EGTA, 0.5 mM MgCl₂, and 10 μM GDP. The data were fitted to a 1:1 binding isotherm with Equation 1,

$$\Delta\text{Fluo} = \text{Fluo}_{\max}$$

$$\times \frac{[\text{R1}^*] + [\text{T}] + K_d - \sqrt{([\text{R1}^*] + [\text{T}] + K_d)^2 - 4 \cdot [\text{R1}^*] \cdot [\text{T}]}}{2 \cdot [\text{R1}^*]} \quad (\text{Eq. 1})$$

where ΔFluo is the variation of the fluorescence signal, Fluo_{\max} is the fluorescence at saturating concentration of tubulin, $[\text{T}]$ and $[\text{R1}^*]$ are the concentrations of tubulin and labeled R1, respectively, and K_d is the dissociation constant.

The dissociation rate constant k_{off} was determined using a Hi-Tech KinetAsyst stopped-flow system (TgK Scientific) at 20 °C in the same buffer. A 30 nM R1* and 50 nM tubulin solution was mixed with either 2.1 or 4.2 μM R1. R1* suffered from some photobleaching in these conditions (data not shown); this was taken into account by fitting the fluorescence decrease with the following mono-exponential decay function,

$$\text{Fluo} = \text{Fluo}_{\min} + \Delta\text{Fluo} \cdot e^{-k_{\text{obs}} \cdot t} + b \cdot t \quad (\text{Eq. 2})$$

where Fluo is the fluorescence signal, Fluo_{\min} is fluorescence at infinite time, ΔFluo is the amplitude of the fluorescence variation, k_{obs} corresponds to k_{off} at saturating concentration of R1, and b is the photobleaching term.

The association rate constant k_{on} was also determined using the same stopped-flow apparatus. We added at least a 6.7-fold excess of tubulin to a fixed concentration of R1* (30 nM), which made the binding reaction pseudo-first-order. The k_{obs} value was fitted with the following exponential equation,

$$\text{Fluo} = \text{Fluo}_{\min} + \Delta\text{Fluo} \cdot (1 - e^{-k_{\text{obs}} \cdot t}) \quad (\text{Eq. 3})$$

where Fluo_{\min} is the fluorescence at time 0. k_{on} was extracted from the plot of apparent rate constant k_{obs} as a function of the tubulin concentration.

Crystallization and Structure Determination—Crystallizations were performed at 293 K by vapor diffusion with the hanging drop method, using 1 ml of well solution and drops formed by 0.8 μl of proteins and 0.8 μl of the crystallization buffer. The tubulin-R4 complex, at a 30 mg/ml concentration in 15 mM Pipes-K, pH 6.8, 0.1 mM GDP, 0.3 mM MgCl_2 , and 0.2 mM EGTA, was crystallized at pH 6.8 using a mix of polyethylene glycol 20,000 (PEG 20,000) and NaCl as precipitants. Most crystals diffracted poorly and anisotropically, leaving ambiguities in the space group and cell parameters. Only one crystal diffracted to almost 4 Å resolution; diffraction data were measured at 100 K on ID29 (European Synchrotron Radiation Facility, Grenoble, France). This crystal was obtained with a crystallization buffer consisting of 1.7 M NaCl and 7% (w/v) PEG 20,000. Before being flash-cooled in liquid nitrogen, it was transferred to the same solution buffered with 15 mM Pipes-K, pH 6.8, and supplemented with 0.1 mM GDP, 1 mM MgCl_2 , 0.5 mM EGTA, and 20% (v/v) glycerol. The data were processed in the C2 space group with XDS (29) (Table 1). The structure was determined by molecular replacement using Amore (30), with $\alpha\beta$ tubulin as a search model (Protein Data Bank code 3RYC (7)). The correlation coefficient between calculated and observed data increased with the number of tubulin heterodimers found (41.9, 49.5, 53.9, and 58.5%). Rigid body refinement in which α and β sub-

units were refined separately resulted in a correlation coefficient of 61%. The R4 starting model was generated from RB3_{SLD} and T₂R. Owing to the moderate resolution of the data, only a few refinement steps with Buster (31) were performed.

In the case of the tubulin-R1 complex (TR1), the C-terminal tail of tubulin was cleaved by subtilisin (7, 32) before tubulin-R1 (TR1) complex formation. TR1 was further complexed with the D2 DARPin. The ternary TR1-D2 complex at 15 to 20 mg/ml, in the same buffer as the one of tubulin-R4 was crystallized in a buffer consisting of 15% (v/v) PEG 550 monomethyl ether and 0.1 M Mes-K, pH 6.5. For data collection, the PEG concentration was raised to 30% before the crystals were flash-cooled in liquid nitrogen. A 2.65 Å data set was collected at 100 K at the Proxima1 beam line (Soleil Synchrotron, Saint Aubin, France) and processed with XDS (29). The intensities were scaled and merged using Scala (33). The structure was solved by molecular replacement with Amore (30) using the structure of a tubulin-D2 complex⁴ as a search model. The structure was refined using the Buster program (31) with iterative model building in Coot (34). Data processing and refinement statistics are summarized in Table 1. Accessible surface areas were calculated with the Areaimol program (33).

RESULTS

Five Modules in Its Sequence Account for the RB3_{SLD} Interaction with the Two Tubulins To which It Binds—To design modular SLD-based constructs that bind any given number of tubulin heterodimers, we used an RB3_{SLD} sequence as a starting point because this protein makes the most stable complex with tubulin among SLDs of other stathmin family members (14). For increased stability, in particular at higher ionic strength, RB3_{SLD} was mutated at four positions compared with the protein used in our previous structural analyses (see supplemental data). The resulting mutant is termed RB3_Q. The boundaries in the RB3_Q sequence of the regions that interact with each tubulin heterodimer were established based on an internal repeat conserved in all SLD sequences (18, 35). Inspection of T₂R structures has indeed shown that the two repeats correspond to two tubulin heterodimer binding regions (16, 36). In RB3_{SLD}, the limits of these repeats, which are displayed in boldface type in Fig. 1B, are Glu⁴⁸–Val⁸² and Glu⁹⁹–Val¹³³ (16) (numbering is in reference to stathmin (14)). The residue immediately downstream the second repeat is an arginine (Arg¹³⁴), which is close to tubulin acidic residues (36); we therefore included it in the second RB3_Q tubulin-binding region. For consistency, the corresponding residue downstream the first repeat (Ile⁸³) was included in the first region (Fig. 1B). To gain more flexibility in the design, we divided each of these two regions into two tubulin subunit binding modules, using a high resolution (2.1 Å) T₂R structure (7). This defines four modules (depicted in different colors in Fig. 1). The N-terminal module (module 1 or M1) comprises residues 4 to 61; it interacts with α 1, the “first” α subunit in T₂R. The following three modules, M2 to M4 (residues 62–83, 84–112, and 113–134, respectively), interact with β 1, α 2, and β 2. When completing this attribution, we noticed

⁴ B. Gigant, L. Pecqueur, B. Dreier, A. Plückthun, and M. Knossow, manuscript in preparation.

TABLE 1
Data collection and refinement statistics

	T ₄ R4	TR1-D2
Data collection^a		
Space group	C2	C2
Cell dimensions		
<i>a</i> , <i>b</i> , <i>c</i> (Å)	639.7, 66.1, 128.1	95.1, 75.6, 155.7
α , β , γ	90.0°, 92.0°, 90.0°	90.0°, 96.3°, 90.0°
Resolution (Å)	50.0–4.17 (4.28–4.17)	48.5–2.64 (2.78–2.64)
<i>R</i> _{sym}	0.112 (0.876)	0.089 (0.627)
<i>I</i> / σ <i>I</i>	8.2 (1.6)	10.5 (2.1)
Completeness (%)	98.6 (90.8)	98.7 (97.5)
Multiplicity	3.7 (3.4)	3.7 (3.7)
Refinement		
Resolution (Å)	4.17	2.64
No. reflections	40,332	31,979
<i>R</i> _{work} / <i>R</i> _{free}	0.241/0.261	0.177/0.201
No. atoms		
Protein	28,901	8446
Ligands/ions	244	180
Waters	0	102
<i>B</i> -factors		
Protein	183	60.2
Ligands/ions	141	65.0
Waters		49.2
Coordinate error (Å) ^b	1.417	0.317
r.m.s.d. ^c from ideal values		
Bond lengths (Å)	0.010	0.010
Bond angles	1.20°	1.20°

^a In both cases, data were collected on a single crystal. There is one complex per asymmetric unit. Values in parentheses are for highest resolution shell.

^b Values shown were estimated from Luzzati plots.

^c r.m.s.d. indicates root mean square deviation.

that the region binding to β 2 (113–141, *i.e.* 29 residues) is significantly longer than the one we had attributed based on the internal repeats and therefore defined a fifth module, M5 (residues 135–141), which caps the β 2 binding region. M5 contributes significantly to T₂R stability as the removal of the last seven residues of RB3_{SLD} (residues 139–145) is detrimental to the stability of the complex, whereas a stop codon at position 142 is not (data not shown). We have used these five modules to design SLDs for binding tubulin with a programmed stoichiometry.

RB3_{SLD}-based Constructs Lead to Programmed Binding of Three and Four Tubulins—To produce a tubulin-SLD 4:1 complex, we designed a construct (R4) consisting of modules M1 to M4 (interacting with two heterodimers in T₂R), followed by M3 (interacting with an α subunit) and ending with the M2–M5 sequence (interacting with a β subunit and a heterodimer) (Fig. 1C). R4 was produced, purified, and characterized both biochemically and structurally. We used SEC-MALLS to determine the mass of the complexes R4 forms with tubulin. In SEC-MALLS conditions, independent of the tubulin-R4 ratio in the sample, the major peak of the complex does not shift, indicating a constant tubulin-R4 stoichiometry (Fig. 2A). The mass determined for the species in this peak (410 kDa, Table 2) is in reasonable agreement with that of a 4:1 tubulin-R4 complex (T₄R4; theoretical mass, 428 kDa). The mass of the controls (tubulin and T₂R) is also fully consistent with the expected values. A second peak is observed (at about 8.6 ml), whose magnitude increases somewhat along with the concentration of R4 but remains small. Because of the small size of this peak and because it is not well separated from the T₄R4 peak, the determination of the mass of the corresponding species is not accurate. Nevertheless, it fits best with a complex containing two tubulins.

Tubulin was also analyzed by classical gel filtration in the presence of varying amounts of R4 (Fig. 2B). Consistent with the SEC-MALLS observations, a species of Stokes radius larger than that of T₂R is formed, most probably indicative of the complex that contains four tubulins. As long as there are more than four tubulin heterodimers per R4 in the sample, the largest Stokes radius species elutes as a well defined and sharp peak. Increasing the concentration of R4 beyond this ratio has two consequences. First, the free tubulin peak disappears. Second, the shape of the fast migrating peak may change, depending on the buffer used for the analysis. In the higher salt buffer used for SEC-MALLS (data not shown) and in a similar one (supplemental Fig. S1), the results are identical to those obtained in SEC-MALLS. By contrast, in a low salt buffer, this peak broadens and its maximum shifts slightly toward lower Stokes radii (Fig. 2B).

We also visualized directly, by electron microscopy, the tubulin complexes formed in presence of R4. Micrographs confirm the presence of complexes with four tubulin molecules, whereas a smaller amount of oligomers with three tubulins can also be seen (Fig. 2C), together with a few presumably uncomplexed tubulin molecules (*e.g.* see bottom left of the right panel of Fig. 2C). Free heterodimers have been commonly observed in micrographs of tubulin-SLD mixtures (*e.g.* see Ref. 18); they might originate from dissociation of the complex during EM sample preparation. To summarize, complexes of 2:1 and 3:1 stoichiometries form between tubulin and R4. These are detected by SEC-MALLS (T₂R4, Fig. 2A) or, in the case of T₃R4, seen by EM (Fig. 2C) and suggested by the SEC experiments (Fig. 2B), implying that R4 molecules compete with each other for tubulin binding. But our results show that T₄R4 is the main complex formed by tubulin and R4. The T₄R4 crystal structure further establishes the organization of tubulin in this complex.

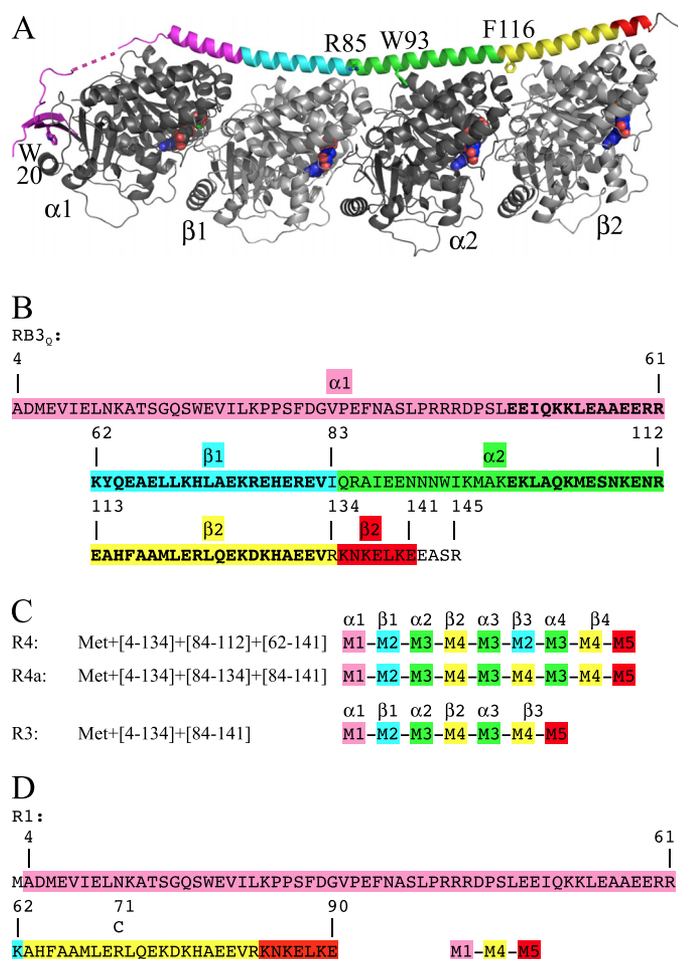


FIGURE 1. The design of RB3_{SLD}-based constructs for binding tubulin with a predefined stoichiometry. *A*, the T₂R structure in which RB3_{SLD} is colored according to the modules used in the design of the new SLDs. The RB3_{SLD} according to which the modules are defined (RB3_Q) contains four mutations that are modeled in the structure, F20W, K85R, F93W, and L116F (numbering is in reference to stathmin), plus the additional C14A mutation. The linker between the N-terminal β hairpin and the C-terminal helix, starting at residue Leu⁴⁷, contains the least ordered region of RB3_{SLD}; part of it is shown as a dashed line. Figs. 1A, 3, and 5 were generated using PyMOL (44). *B*, the RB3_Q sequence colored according to the same modules as in *A*. The residues of the two stretches of the internal repeat (Glu⁴⁸-Val⁸² and Glu⁹⁹-Val¹³³) are highlighted in boldface type. *C*, the design in terms of the modules defined in panel *B* of SLDs engineered to bind three (R3) or four tubulin heterodimers (R4 and R4a). In *A*-*C*, the tubulin subunits interacting with the modules we defined are indicated. *D*, sequence of R1 designed to bind one tubulin molecule. The position of the residue mutated to cysteine (R71C) and used to label R1 for affinity measurements is indicated.

Tubulin in complex with R4 was crystallized, and the structure was determined by molecular replacement. There is one complex consisting of four tubulin heterodimers in the asymmetric unit. The elongated shape of the complex (Figs. 2C and 3A) is also reflected by its large hydrodynamic radius as determined by quasi-elastic light scattering in the SEC-MALLS experiment (Table 2). The R4 main chain was traced in the electron density maps, but consistent with the moderate resolution of the diffraction data set (4.2 Å, see Table 1), the signal for the side chains remained weak (Fig. 3A). As expected from T₂R (16), the tubulin molecules are arranged head to tail as a curved protofilament-like assembly. Also similar to T₂R (36), ~11° rotations superimpose consecutive subunits, *i.e.* those within a tubulin molecule as well as the adjacent ones of con-

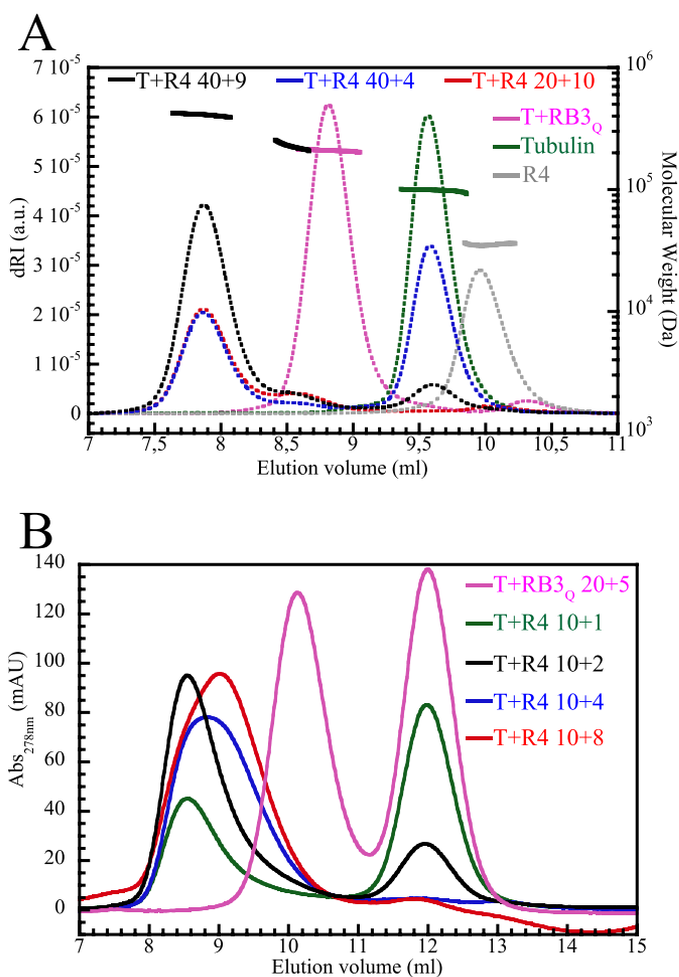


FIGURE 2. R4 forms mainly a T₄R4 complex with tubulin. *A*, SEC-MALLS analysis. The differential refractive index (*dRI*, arbitrary unit (*a.u.*)) on the left axis, dotted lines) and molecular mass (displayed as solid lines for the regions of interest, *i.e.* for the chromatographic peaks, with the scale on the right axis) are plotted as a function of the column elution volume. The samples analyzed were as follows: tubulin (40 μ M, green), R4 (60 μ M, gray), T₂R (40 μ M tubulin and 30 μ M RB3_Q, magenta), and tubulin-R4 (20:10 μ M, red; 40:4 μ M, blue; 40:9 μ M, black). The molecular masses of tubulin-R4 complexes are only displayed in the case of the 40:9 μ M sample. *B*, gel filtration profiles obtained with a low salt buffer. Samples (100 μ l) containing 10 μ M tubulin and increasing concentrations of R4 (1 μ M, green curve; 2 μ M, black; 4 μ M, blue; and 8 μ M, red) were injected on the column. As a control, a sample containing 20 μ M tubulin and 5 μ M RB3_Q was also analyzed (magenta). mAU, milliabsorbance units. *C*, electron micrographs of negatively stained tubulin-R4 complexes. Species comprising four tubulin heterodimers (left) predominate, whereas complexes with three tubulins are also identified (right). Their dimensions (~55 Å \times 355 Å and 55 Å \times 265 Å, respectively) are consistent with those of a smaller SLD complex comprising two tubulins (15). Scale bar, 100 Å.

TABLE 2
 Mass determination of tubulin-SLD complexes by SEC-MALLS

Mass (kDa)	T:R4					
	40:4 μM	40:9 μM	20:10 μM	T:RB3 _Q	Tubulin	R4
Theoretical ^a		428 (T ₄ R4), 228 (T ₂ R4)	409 ^b , 237 ^c	217 (T ₂ R)	100	28.4
Determined by SEC-MALLS	411 ^b , 222 ^c , 99 ^d	410 ^b , 228 ^c , 101 ^d		209	99	35
Rh (nm)	7.7 ^b	7.9 ^b	7.7 ^b	5.5	3.8	ND ^e

^a Assuming 100 kDa for tubulin.

^b Main peak of the complex.

^c Small peak of the complex.

^d Uncomplexed tubulin.

^e ND, not determined; Rh, hydrodynamic radius.

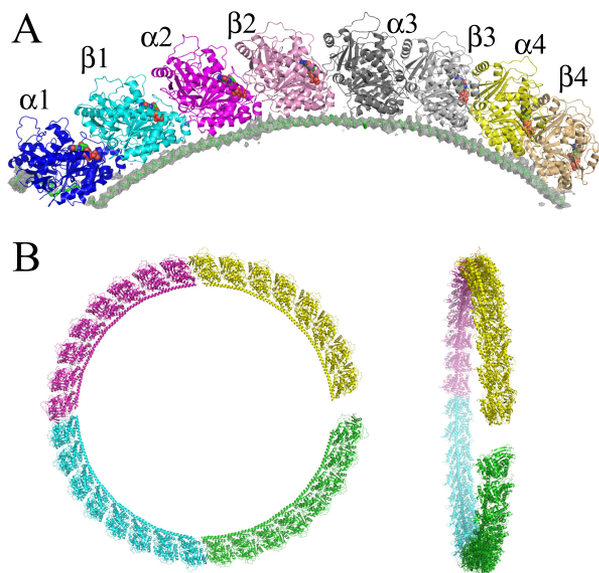


FIGURE 3. The T₄R4 structure. *A*, overview of the complex in which each tubulin is colored differently. The α (β) subunits are in brighter (lighter) colors. The 4.2 Å resolution $2F_{\text{obs}} - F_{\text{calc}}$ electron density map of the R4 molecule, contoured at the 1 σ level, is displayed. *B*, the relative orientations of the tubulin subunits in T₄R4 are close to those in a ring. The model resulting from the repetition of T₄R4 was obtained by superimposing the β 1 moiety of the ($m+1$)th complex onto the β 3 moiety of the m th complex and by keeping in the final model the 1st, 3rd, 5th, and 7th complexes. Each T₄R4 is colored differently. The resulting flat helix is viewed along its axis (*left*) and nearly perpendicularly to it (*right*).

secutive heterodimers. The curvature of the complex is also pictured by a T₄R4 helical superassembly (Fig. 3*B*). A comparison of the two T₂R structures we have determined previously revealed that the intertubulin interface is variable (7) and gives rise to T₂R helical superassemblies with pitches of opposite signs, depending on the crystal form considered. Superassembly of T₄R4 in the crystal form described here yields a helix with a pitch that is close to zero, *i.e.* almost a ring (Fig. 3*B*), suggesting that a wide range of pitches within the limits initially found with T₂R may indeed be observed.

In addition to R4, we designed another construct, termed R4a, which starts with modules M1 to M4, continues with twice a repeat of M3+M4, and ends with M5 (Fig. 1*C*). R4a differs from R4 in the stretch interacting with the β subunit of the third tubulin molecule in the T₄R4 complex. R4a was produced and purified but because the initial characterization of its interaction with tubulin revealed no obvious difference compared with that of R4, it was not characterized any further. Using the same rationale, we designed and produced the R3 construct (Fig. 1*C*),

which also interacts with tubulin, likely giving rise to a T₃R3 assembly (supplemental Fig. S1). The stoichiometries of the new complexes of SLDs with tubulin are as predefined. Unfortunately, the resolution of the structure we determined is not sufficient to demonstrate that in our designs, or at least in the complex crystallized, SLD residues interact with tubulin as expected, *i.e.* as in T₂R. The design, based on the same principles, of an SLD (named R1, Fig. 1*D*) that binds one tubulin and the inspection of the structure of the resulting complex (TR1), has allowed us to validate the rationale of the design of the three modules that constitute this SLD.

R1 Construct Makes a Tubulin-R1 1:1 High Affinity Complex—The R1 N-terminal part consists of an M1 module, the N-terminal end of RB3_Q that interacts with a tubulin α subunit. Its C-terminal part comprises M4 and the capping motif M5, both regions constituting together the C-terminal region of RB3_Q that interacts with a β subunit. In addition, we replaced the first residue of M4 by a lysine, to compensate for the negative charge of the residue that is four residue upstream (Glu⁵⁸), *i.e.* on the same face of the SLD helix. A similar distribution of opposing charges is observed at the M1-M2 and M3-M4 boundaries of RB3_{SLD} (Fig. 1).

The interaction of R1 with tubulin was initially detected in a gel filtration assay. We observed a small shift upon addition of R1 to tubulin (slightly larger than 0.2 ml, see supplemental Fig. S1). Interestingly, a similar shift had been obtained upon addition to tubulin of a protein that makes a 1:1 complex with it and is slightly larger than R1 (113 *versus* 87 residues) (37). The tubulin-R1 interaction was quantified by fluorescence experiments using R1*, an acrylodan-labeled R1. The stepwise addition of tubulin to R1* leads to an increase of the acrylodan fluorescence signal (Fig. 4*A*). Fitting the data with a quadratic equation (Equation 1, see “Experimental Procedures”) yields the value of the equilibrium dissociation constant ($K_d = 1$ nM). This estimate reflects the high affinity of the TR1 complex, but it is probably not accurate, as the R1* concentration used in these experiments was \sim 10-fold higher than the K_d . To confirm the tight interaction and characterize it further, we also determined the dissociation and association rate constants. A chase of R1* from tubulin by unlabeled R1 (Fig. 4*B*) yielded the dissociation rate constant $k_{\text{off}} = 0.016$ s⁻¹. The association rate constant was derived from association kinetics in pseudo-first-order conditions, yielding a k_{on} of 8×10^6 M⁻¹ s⁻¹ (Fig. 4*C*). The K_d deduced ($k_{\text{off}}/k_{\text{on}} = 2$ nM) is consistent with the value determined by titrating tubulin with R1. To identify the interactions responsible for this tight complex, we determined its structure.

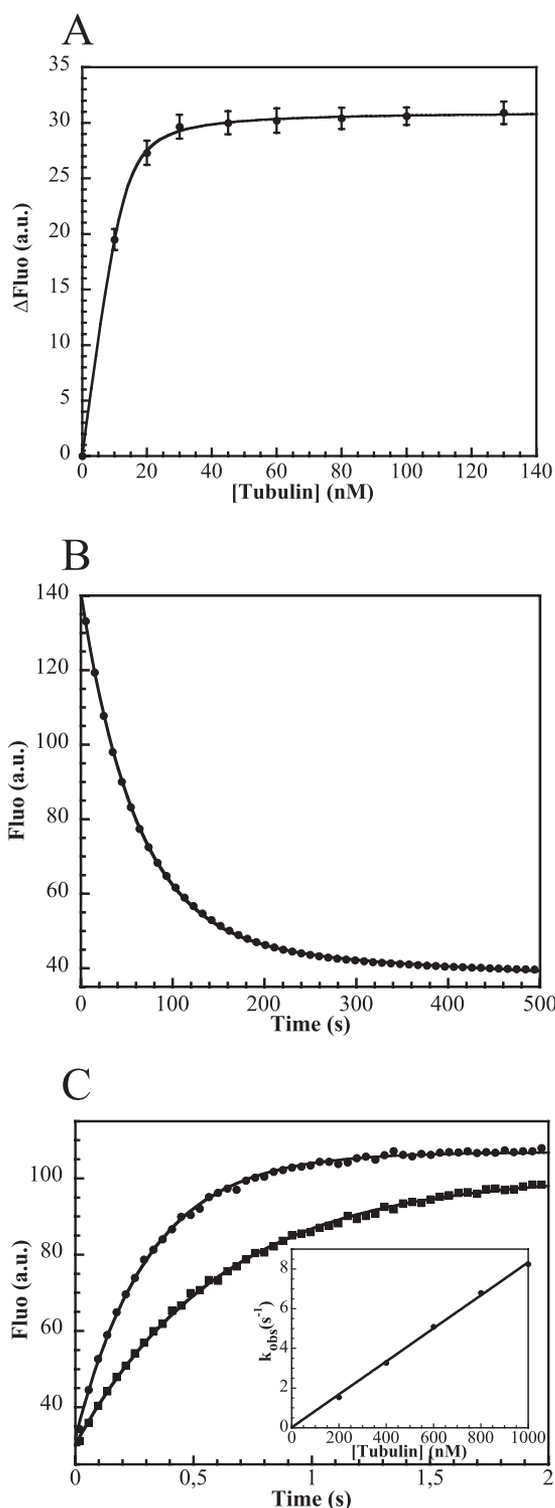


FIGURE 4. The tubulin-R1 interaction monitored by fluorescence spectroscopy. *A*, fluorescence variation of 13 nM R1* upon addition of tubulin. The curve is the fit of the experimental points with Equation 1, from which the K_d (1 nM) is extracted. *Error bars* correspond to the S.D. of the variation of fluorescence signal upon tubulin addition. *B*, dissociation of R1* from tubulin. The fluorescent TR1* complex was formed by mixing 30 nM R1* with 50 nM tubulin. The fluorescence decrease following addition of 4.2 μ M R1 to this sample was monitored in a stopped-flow apparatus. The curve is the fit of the experimental points (5% of which are shown) with Equation 2. The same rate constant was obtained with two R1 concentrations (see "Experimental Procedures") and is interpreted to be the dissociation rate constant of the complex ($k_{\text{off}} = 0.016 \pm 0.003 \text{ s}^{-1}$). *C*, determination of the association rate constant. Tubulin, at concentrations ranging from 200 nM to 1 μ M, was added to a fixed

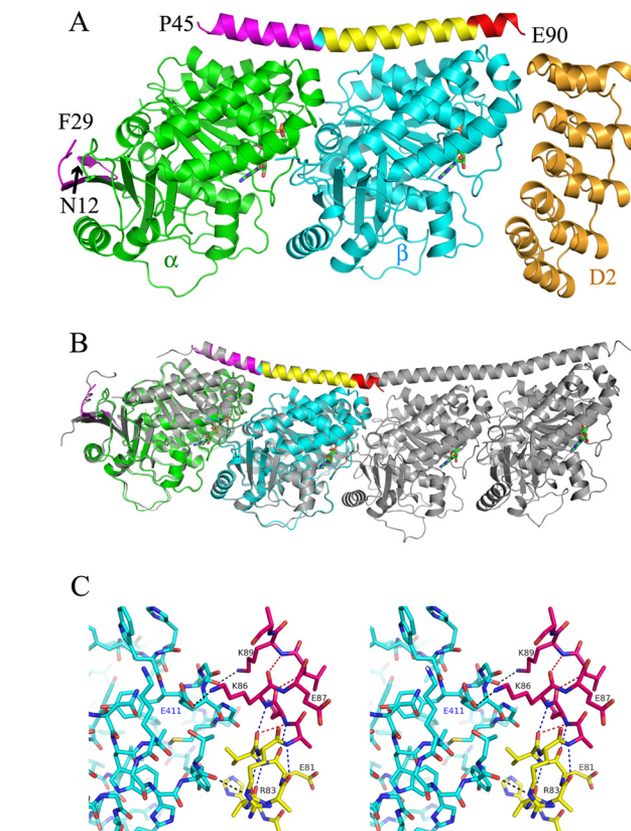


FIGURE 5. The TR1 structure. *A*, overview of the TR1-D2 structure. The tubulin α and β subunits are in green and cyan, respectively. R1 is colored by modules as in Fig. 1. D2, the tubulin-binding DARPIn with which crystals were obtained, is in orange. The nucleotides (GTP on α , GDP on β) are in green. The disordered eight N-terminal residues of R1 as well as the disordered linker between its N-terminal β hairpin and the C-terminal α helix (residues 30 to 44) are not displayed. *B*, comparison of TR1 with T₂R. TR1 is colored as in *A* and superimposed on T₂R (gray). The root mean square deviation after superposition (45) of C α s of α , β , and R1 from TR1 and of α 1, β 1, and RB3_{SLD} from T₂R is 0.691 Å (872 atoms compared). *C*, stereo view of the interaction of the C-terminal end of R1 with tubulin in TR1. R1 and the tubulin β subunit are colored as in *A*. The hydrogen bonds between R1 and tubulin, including the Lys⁸⁶-Glu⁴¹¹ salt bridge, are displayed as black dotted lines. The main chain hydrogen bonds in the R1 helix are displayed as red (3_{10} -helix) or blue (α -helix) dotted lines. R1 residues Glu⁸⁰, Lys⁸⁴, and Glu⁹⁰, whose side chains are not defined, have been modeled as alanines. See supplemental Fig. S2 for the same view with R1 in its electron density map.

Structural Basis for Tubulin-R1 High Affinity—Crystals of TR1 were obtained as a ternary complex with the anti-tubulin DARPIn D2 used as a crystallization chaperone. Data to 2.65 Å were collected from one crystal (Table 1), and the structure was solved by molecular replacement. D2 targets the β tubulin longitudinal interface. Hence, its binding site is distinct from that of R1, and the two proteins do not seem to interfere for tubulin binding (Fig. 5*A*). In the R1 structure, the C-terminal helix is ordered up to the end of the molecule, most of the side chains being defined in the electron density maps. Two R1 stretches could not be traced: the N-terminal half of the β hairpin first strand and the linker between this N-terminal β hairpin and the C-terminal helix. This loop is also mostly disordered in all the

concentration of R1* (30 nM). Fluorescence variations upon addition of 200 nM (square symbols) and 400 nM (dots) tubulin are shown (5% of the experimental points are displayed). The data were fitted according to Equation 3. The variation of k_{obs} as a function of tubulin concentration is linear. k_{on} is the slope ($8 \times 10^6 \text{ M}^{-1} \text{ s}^{-1}$) of that curve (inset). *a.u.*, arbitrary units.

other tubulin-SLD structures determined so far (7, 36), probably reflecting its mobility in the complex. Nonetheless, the accessible surface area buried by the ordered part of R1 is large, giving a rationale for the high affinity of the TR1 complex (38). About 2250 Å² are buried on the R1 side and a similar surface (2130 Å²) on tubulin. Two-thirds of the tubulin buried surface are on the α subunit, and one-third is on β ; this difference reflects the contribution of the R1 N-terminal β hairpin that caps the α longitudinal interface. Tubulin in the TR1 complex is curved and retains the conformation it has in T₂R (and in T₄R4, see above) (Fig. 5B). Moreover, the footprint of R1 on tubulin is very similar to that of RB3_{SLD} on the two distal tubulin subunits in T₂R. Most important, the two R1 moieties make with tubulin in TR1 the exact same interactions as those established by the corresponding modules of RB3_{SLD} in T₂R. This fully validates the design principles used to define M1, M4, and M5 and strongly suggests that M2 and M3, which were similarly defined, also interact with tubulin as planned.

R1 makes a stronger complex with tubulin than previous constructs of similar length (*e.g.* see Refs. 19 and 20). These consisted in a continuous N-terminal region of an SLD, *i.e.* they were based on constructs including the M1 and M2 modules. The tighter interaction established by R1 may be attributed to two main reasons. First, RB3_{SLD} makes fewer contacts with β 1 in T₂R than R1 with β tubulin in TR1 (or than RB3_{SLD} with the β 2 subunit in T₂R). In particular, two salt bridges are mediated by the M4–M5 stretch (between Asp⁷⁶ of R1 and Lys β ¹⁵⁶ of tubulin and, as shown in Fig. 5C, between Lys⁸⁶ and Glu β ⁴¹¹) whereas only one (mediated by Arg⁷⁶ of RB3_{SLD} and Glu β ¹⁵⁹) is contributed by the M2 module (tubulin residue numbering is as in Ref. 12). Secondly, the R1 C-terminal end is derived from an RB3_{SLD} ending at position 141. Glu¹⁴¹ is the residue at the boundary between the helix and the downstream non-helical region (7). In TR1, the R1 helix is not capped in the usual sense (39), but it is stabilized at its C-terminal end by interactions with tubulin of the side chains of two residues: Lys⁸⁶, as mentioned above, and Lys⁸⁹ (respectively equivalent to Lys¹³⁷ and Lys¹⁴⁰ in RB3_{SLD}). In addition, 3₁₀ helix main chain hydrogen bonds are made in the last turn, by the carbonyls of residues 85 and 86 with the amide NH of residues 88 and 89, respectively (Fig. 5C). Constructs with an untimely helix termination probably lacked these interactions and have resulted in proteins that make less stable complexes with tubulin.

DISCUSSION

We have presented the design and construction of artificial SLD proteins that efficiently bind tubulin in a range of pre-defined stoichiometries. This may serve at least two purposes. The first purpose is to provide alternative options for tubulin crystallization. The second purpose is the design of homogeneous and stable tubulin assemblies larger than T₂R for cryo-electron microscopy. Both objectives have been reached. First, new crystals of a single sequestered tubulin have been characterized. Second, as a proof of principle, we designed SLDs that form a T₄R4 complex. Both EM (Fig. 2C) and SEC-MALLS analyses (Fig. 2A) indicate that the designed stoichiometry has been obtained. The stability of the T₄R4 complex, higher than those obtained with SLDs from invertebrates (18), together

with its crystal structure showing the longitudinal arrangements of the tubulins in a curved protofilament-like assembly, make it a well characterized entity for EM studies. In addition to defining the limits in RB3_{SLD} sequence of tubulin subunit binding modules, our study identifies one important feature of strong RB3_{SLD}-based tubulin binders: the presence at their C-terminal end of a capping module, termed here as M5 (Fig. 1). M5 ending at residue 141 confers strong binding to the SLDs we designed, whereas an SLD ending at residue 138 is a weaker binder. Residues up to Lys⁸⁹ in R1 (Lys¹⁴⁰ in RB3_{SLD}) make important contributions to the stability of the SLD helix both through side chain interactions with tubulin and main chain intramolecular hydrogen bonds (see above). Therefore, Glu¹⁴¹ is probably close to the C-terminal end of the shortest M5 module for strong tubulin binding.

In addition to the new SLD constructs described above, our work provides new information on the tubulin-SLD interaction. Most studies of this interaction have been centered on ternary complexes of an SLD with two tubulin heterodimers. It is not straightforward to distinguish biochemically the binding of the two tubulins in these complexes. This has led to discrepancies of the estimates of their affinities and of their binding cooperativity (reviewed in Ref. 40). R1, as it associates with tubulin in a 1:1 complex, represents a simplified starting point for such studies. Both the kinetics and the strength of the association are of interest. We have found that the tubulin-R1 association is fast, its rate constant being close to that estimated from Brownian dynamics simulations (41). By comparison, the k_{on} in T₂R is \sim 400 times slower (42). The suggestion has been made that the slow association kinetic in T₂R may be related to structural rearrangements within the complex (42). As the association rate constants of SLDs other than RB3_{SLD} are significantly faster (14), the slow association rate of T₂R may be due to adjustments of the RB3_{SLD} structure rather than to the tubulin-tubulin association. The question then arises of whether the RB3_{SLD} region where these adjustments take place may be narrowed down. Because the association of R1 with tubulin is fast, it is tempting to suggest that they take place in the RB3_{SLD} modules that are not included in R1, *i.e.* in the modules that interact with the β 1 and α 2 tubulin subunits.

We have also found that the tubulin:R1 affinity is large (K_d in the low nanomolar range). Because no natural tubulin-SLD 1:1 complex has been identified so far, it might have been thought that SLD-based sequesterers are inherently unable to bind one tubulin molecule efficiently. R1 proves that it is not the case as it binds one heterodimer tightly. Therefore, the failure to identify such a natural assembly is not due to an impossibility to produce it using SLD-based sequences. There must be another explanation. In the balance between mechanisms in which SLDs favor either microtubule assembly or disassembly depending on their phosphorylation state, our results give weight to the hypothesis that a main function of SLDs is to store tubulin at specific localizations in the cell (43) to deliver it for microtubule polymerization when required. In this scheme, tubulin becomes available for assembly upon SLD phosphorylation. Obviously, an SLD that releases more than one tubulin heterodimer at a time fulfills this function more efficiently than R1, at least from the point of view of the quantity of kinases

required to release a number of tubulin molecules. The question then arises of whether higher stoichiometry complexes would be even more efficient. It seems that in such complexes the tubulin binding cooperativity is less pronounced. We observed this with R4 which, in certain conditions, gives rise to unsaturated assemblies (T₂R4 and T₃R4) (Fig. 2) and this was also observed with *Drosophila* SLDs (18). SLDs binding two heterodimers may therefore be an optimal choice for efficient sequestration and phosphorylation-inducible tubulin release.

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SUPPLEMENTARY MATERIAL

Supplementary results

RB3q, an improved RB3_{SLD}.

In the course of the structural study of tubulin-SLD assemblies, we screened a set of 25 RB3_{SLD} mutations in order to identify those that strengthen the interaction with tubulin. To assess the effect of these mutations, we used a test based on vinblastine-mediated oligomerization. Vinblastine induces longitudinal tubulin-tubulin associations resulting in curved assemblies (1,2), a process impeded when tubulin is associated with RB3_{SLD}. But in higher ionic strength conditions, T₂R becomes less stable. Hence tubulin is less protected from vinblastine-induced assembly. We identified four mutations (F20W, K85R, F93W, L116F, stathmin numbering (3)) that counteract to some extent the detrimental effect of salt on T₂R stability. Three of them introduce bulkier and aromatic side chains prone to interact with hydrophobic patches of tubulin. In particular, Trp20 targets the same α tubulin pocket as vinblastine does (2). The fourth mutation (K85R) was aimed at reinforcing the RB3_{SLD} interaction with nearby tubulin acidic residues (Asp β 414 and Glu β 417, sequence numbering as in Ref. 4). We have produced an RB3_{SLD} variant named RB3_Q that combines these four mutations plus the C14A substitution and checked that the stabilizing effects are additive, with a predominant contribution of Trp20. We have used RB3_Q as the RB3_{SLD} template for the constructs described in this study.

Supplemental Figure Legends

Figure S1. SEC analysis of tubulin:SLD complexes. Samples (100 μ l) were injected on a Superdex S200 column (GE Healthcare) equilibrated with 80 mM Pipes-K pH 6.8, 2 mM MgCl₂ and 0.5 mM EGTA. They consisted of 10 μ M tubulin (green curve) and either 10 μ M R1 (blue), 10 μ M RB3_Q (magenta), 5 μ M R3 (red), or 3.3 μ M R4 (black).

Figure S2. Stereo close-up of the tubulin-R1 interface as in Fig. 5C, with R1 in its 2F_{obs}-F_{calc} electron density map contoured at the 1 σ level.

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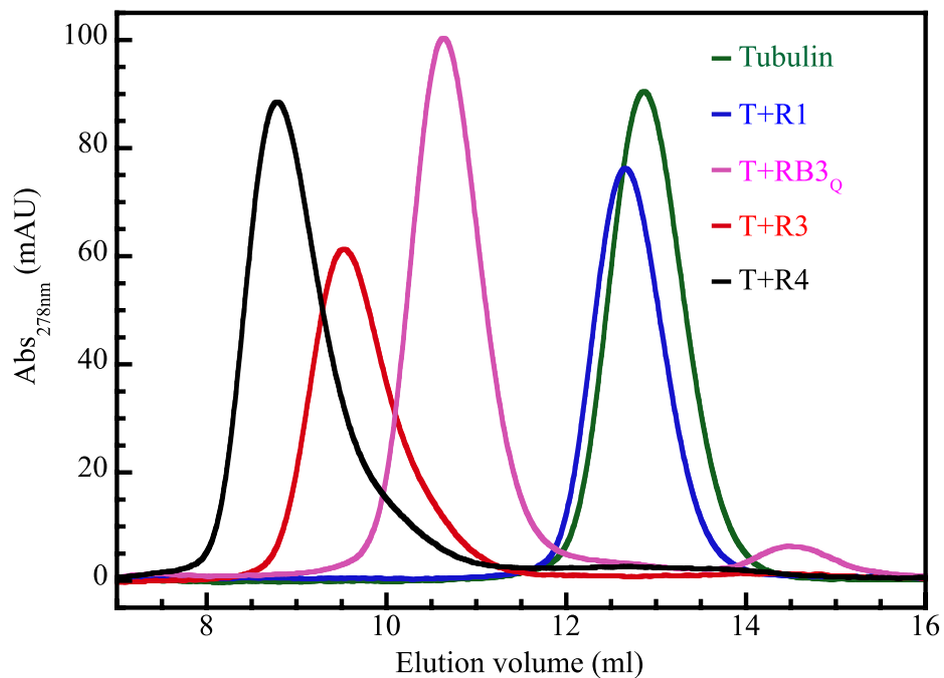


Figure S1

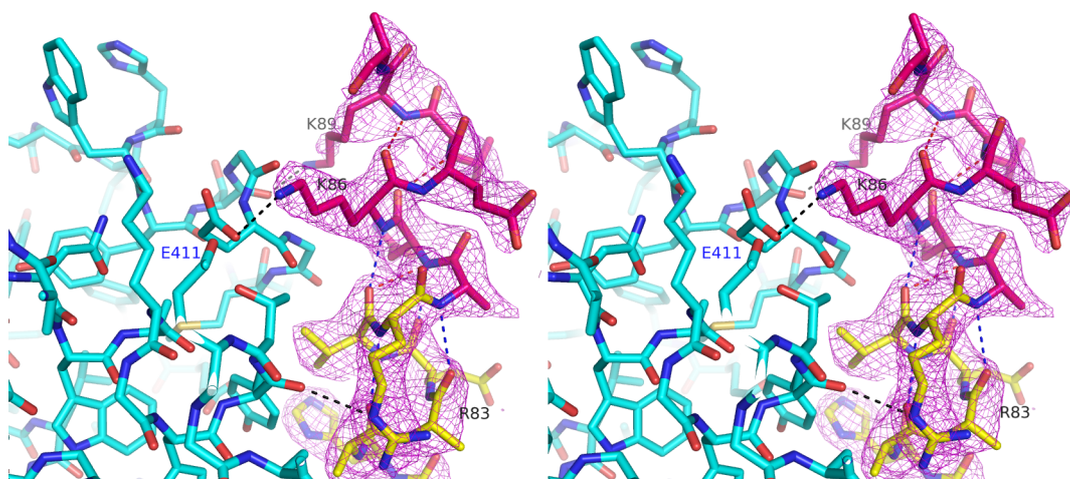


Figure S2